The present invention is directed to a syringe assembly (48), particularly useful for use with conventional prefilled pharmaceutical cartridge-needle units (46), including a hollow body (4) housing the cartridge-needle unit or other syringe structure. The substantially hollow body and the syringe structure include mating slits (28, 29, 30) and a ring (26) so that the needle (60) can be exposed for I.M. injection use and then withdrawn into the body for safety. The body is configured so that when the user diametrically squeezes the body adjacent the slit(s), the ring disengages the slit(s) due to the deformation of the body which permits the needle tip (62) to be positioned inside, for safety, or outside, for use, of the body. The internal position is also useful for safely injecting a pharmaceutical into an IV port (88). When the invention is used with a conventional pharmaceutical cartridge-needle unit as the syringe structure, the body can be part of an enclosure unit (2) which acts as the packaging for the pharmaceutical cartridge-needle unit, as the body of the syringe for administration of I.V. and I.M. injections, and as individual integral sharps containment for safe antineedle-stick disposal of the used cartridge-needle unit.
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SAFETY SYRINGE ASSEMBLY WITH RADIAALLY DEFORMABLE BODY

CROSS REFERENCE TO RELATED APPLICATION
This application is related to U.S. Patent Application Serial No. 07/558,878, filed July 27, 1990, the disclosure of which is incorporated by reference.

BACKGROUND OF THE INVENTION
The need for preventing inadvertent needle sticks has been recognized for many years. For example, U.S. Patent No. 2,571,653 to Bastien shows a syringe in which the barrel of the syringe is mounted within a protective sheath. The sheath can be placed at different axial positions relative to the barrel, one exposing the needle for use and one covering the needle for safety. Many other safety syringe assemblies have been developed as well. However, one of the problems with the prior art safety syringe assemblies is that they generally cannot be used with conventional syringes, are awkward to use and make calibrations and volume graduations difficult to see clearly. Also, existing safety syringe assemblies often are not economical to produce; this is especially true when the syringe structures are intended to be disposable.

SUMMARY OF THE INVENTION
The present invention is directed to a syringe assembly, particularly useful for use with conventional prefilled pharmaceutical cartridge-needle units, including a hollow body housing the cartridge-needle unit or other syringe structure. The hollow body and the syringe structure include mating positioning elements to permit the syringe structure to be secured at one or more axial positions within the body. The positioning elements are preferably radially extending recesses and extensions. The hollow body typically has at least two internal recesses sized and positioned to engage a radially extending member carried by the syringe structure so that the
needle can be partially or totally exposed for use and then withdrawn into the body for safety. The body is configured so that when the user squeezes the body at positions diametrically adjacent the recess, the radially extending member becomes disengaged from the recess due to the deformation of the body. This frees the needle permitting it to be repositioned within the body.

The body is constructed so that the body may be deformed to permit the radially extending member to freely move from the recess to permit the syringe structure and body to move axially relative to one another. This may be achieved by constructing the body with an elliptical cross-sectional shape and forming pairs of the recesses through the walls of the body while constructing the radially extending member as an annular, outwardly extending ring carried by the syringe structure. The radially extending member can be disengaged from the recesses by squeezing the body to deform the body from the generally elliptical shape to a generally circular shape. Preferably, three sets (or four sets to permit one molded body to accommodate both short and long needle lengths) of axially spaced-apart recesses are provided. One recess near the distal end of the body for I.M. (IntraMuscular) injections. An intermediate recess (or two to accommodate both long and short needle cannula) such that the tip of the needle is within the interior of the body may be used for safer disposal. Also, this position may be used for I.V. (IntraVenous) port injection. By properly sizing the opening at the distal end of the body, the body may be placed over the injection valve of an I.V. port, permitting the sharpened point of the needle cannula to pierce the membrane port only after this sharpened point has been shielded from human contact. This permits I.V. port injection with a high degree of safety and efficacy. The recess or recesses closest to the plunger end of the body are used to ensure the needle tip is well within the body for safe disposal of the syringe structures.

One of the primary advantages of the invention is that it permits the needle of a syringe structure to be safely housed within the body in an extremely simple and cost
effective manner. In addition, although the invention can be used with conventional syringes, the invention finds particular utility when used with prefilled pharmaceutical cartridge-needle units. In such case, the body can be part of an enclosure unit, the enclosure unit acting as both the protective packaging for the cartridge-needle unit, and also as a part of the operational syringe assembly itself.

Other features and advantages will appear from the following description in which the preferred embodiments have been set forth in detail in conjunction with the accompanying drawings.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a side cross-sectional view of an enclosure unit made according to the invention;

FIG. 2 is an isometric view of a set of the enclosure units of FIG. 1;

FIG. 3 shows the enclosure unit of FIG. 1 after the adapter has been driven into the body section at the needle end of the body section;

FIG. 4 illustrates a syringe assembly made according to the invention by mounting a cartridge-needle unit mounted within the body section of the enclosure unit of FIG. 3 with the cap at the end section used to plug the end section to maintain the cartridge-needle unit within the enclosure unit and help prevent tampering;

FIG. 5 illustrates the syringe assembly of FIG. 4 after the end unit has been separated from the body section and discarded and the stem section has been separated from the adapter and mounted to the piston at the plunger end of the barrel of the cartridge-needle unit;

FIG. 6 illustrates the syringe assembly of FIG. 5 after the sheath has been removed from the needle and the plunger has been depressed, injecting the contents of the barrel through the needle;

FIG. 7A is a cross-sectional view taken along line 7-7 of FIG. 6 showing the elliptical cross-sectional shape of the body section and the engagement of the adapter ring with
the circumferential extending body slots formed in the body section at the needle end of the body section;

FIG. 7B illustrates the syringe assembly of FIG. 7A after the user has deformed the body section by squeezing the body section as indicated to disengage the adapter ring from the body slots to permit the cartridge-needle unit to be moved axially within the body section;

FIG. 8 illustrates the movement of the cartridge-needle unit from the use position of FIG. 6, with the needle exposed, to the safe position of FIG. 8, with the needle fully housed within the body section, and also illustrates how the syringe assembly of FIGS. 6 and 8 could be used to inject a medicine into, or remove a sample from, an injection arm of an IV port while keeping the sharpened tip of the needle in a safe position;

FIG. 9A is an enlarged view taken along line 9A of FIG. 8 illustrating the engagement of a standard adapter ring within a body slot;

FIG. 9B shows an alternative embodiment of the adapter ring of FIG. 9A in which the plunger facing surface of the ring provides a ramp effect to enhance movement of the cartridge-needle unit towards the plunger end of the body section;

FIG. 10 illustrates the syringe assembly of FIG. 6 in which the adapter ring is positioned within the body slots adjacent the plunger end of the body section to accommodate longer needles than used with the embodiment of FIG. 8;

FIGS. 11 and 12a illustrate alternative embodiments of the syringe structure of FIG. 8 adapted for use with different length needles and including different width body slots to accommodate different width adapter rings so the same body section can be used with needles of different lengths and still maintain the proper position of the tip of the needle for IV port use;

FIG. 12b is a partial side view of the syringe structure of FIG. 12a;

FIGS. 13 and 14 illustrate a further alternative embodiment of the invention in which the adapter ring does not
extend completely around the adapter to engage the appropriately positioned body slot for a short needle version of the cartridge needle unit;

FIGS. 15 and 16 show the embodiment of FIGS. 13 and 14 with a long needle version of the cartridge-needle unit and with an adapter ring positioned to the opposite side of the body section as in the embodiment of FIG. 13, so that the adapter ring of the embodiments of FIGS. 13 and 15 engage the appropriately intermediate position body slots located at different axial positions along the body section according to the lengths of the needles;

FIG. 17 illustrates a further embodiment of the invention in which the outer surface of the body section is normally cylindrical with the inner surface having internal flanges which engage a circumferential slot formed in an adapter mounted to the hub of the cartridge-needle unit;

FIG. 18A is a cross-sectional view of the syringe assembly of FIG. 17 taken along line 18-18 illustrating the engagement of the flanges into the annular slot;

FIG. 18B illustrates the syringe assembly of FIG. 18A after the body section has been deformed causing the flanges to disengage from the annular slot to permit the cartridge-needle unit to move axially within the body section;

FIG. 19 illustrates an alternative embodiment of the invention in which the syringe assembly of FIG. 4 has been modified to eliminate the need for a separate adapter by using a cartridge-needle unit in which the hub of the cartridge-needle unit is modified to include its own adapter ring;

FIG. 20 shows a further embodiment of the invention for use with smaller sized cartridge-needle units; and

FIG. 21 is a cross-sectional view taken along line 21-21 of FIG. 20.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

FIG. 1 illustrates a one-piece molded enclosure unit 2, preferably made from clear polypropylene or polyethylene. Enclosure unit 2 includes elongate, substantially hollow body section 4 having a needle end 6, a
plunger end 8 and defining an interior 10 therein. Unit 2 also includes an end section 12 frangibly connected to plunger end 8 at frangible connections 14. End section 12 includes a cap 16 connected to the remainder of end section 12 by an integral hinge 18 to permit the end 20 of end section 12 to be oriented and sealed as suggested in FIG. 4.

Enclosure unit 2 also includes an adapter 22 extending from needle end 6 and attached thereto by frangible connections 24. Adapter 22 has a circumferentially extending adapter ring 26 sized to engage specially positioned pairs of body slots 28, 29, 30 formed in body section 4. The method and purpose for doing so will be discussed below. Finally, enclosure unit 2 also includes a hollow stem section 32 frangibly attached to adapter 22 by frangible connections 34.

Stem section 32 is hollow to accommodate a sheathed (typically sterile) needle (see FIG. 4) and also has a threaded tip 36 to permit stem section 32, once removed from adapter 22, to be secured to a piston to create a plunger as suggested in FIG. 5. Adapter 22 is moved from the position of Fig. 1 to that of Fig. 3 by forcing stem section 32 in the direction of arrow 35, thus severing frangible connections 24, until ring 26 engages slots 28. Body section 4 includes a shoulder 37 at needle end 6. Shoulder 37 helps to keep adapter 22 from inadvertently passing back out of interior 10. Shoulder 37 may be a continuous shoulder, or it may be formed as diametrically opposed segments.

FIG. 2 illustrates a set 38 of enclosure units 2 molded as a one-piece item and secured to one another through frangible connections 40 at needle end 6 of body section 4 and by frangible connections (not shown) extending between the finger grips 42 at plunger ends 8. As discussed in the above-referenced application, set 38 of enclosure units 2 acts as both the protective shipping and storage container for cartridge-needle units 46, shown in Fig. 4, as well as providing other functions as discussed below.

FIG. 4 illustrates enclosure unit 2 with a conventional cartridge-needle unit 46, such as one made by Wyeth-Ayerst Laboratories of Radnor, Pennsylvania, under the
trademark TUBEX, mounted therein to create a syringe assembly 48 in its as-shipped condition. Cartridge-needle unit 46 is of the type including a barrel 50 containing a flowable material, typically a liquid, and having a piston 52, typically initially at the plunger end 54 of barrel 50. A needle assembly 56 is mounted to the needle end 58 of barrel 50. Needle assembly 56 includes a needle 60, having a sharpened tip 62, and a hub 64 used to mount needle 62 barrel 50. An elastomeric sheath 65 is used to cover needle 60 and seal tip 62. Hub 64 is secured within the interior 66 of adapter 22 through the engagement of a lip 68 (see FIG. 3) with the plunger facing edge 70 of hub 64. As suggested by arrow 72, once cartridge-needle unit 46 is mounted within enclosure unit 2, cap 16 is pivoted to close end 20 and is sealed in place, such as with an irreversible mechanical lock which is also tamper-resistant and tamper-evident, an adhesive or with heat or ultrasonic welding techniques. The thickness of cap 16 is sufficient to resist tampering with the contents of barrel 50 while syringe assembly 48 is in its as-shipped condition of FIG. 4. To further inhibit tampering, an internal metal shield could be used.

To use syringe assembly 48, stem section 32 is fractured from adapter 22, typically with a twisting action, and tip 36 is secured to piston 52, typically with a threading action as suggested by arrow 74 in FIG. 5. Next, sheath 65 is removed and the injection is given by driving plunger 76, made up of stem 32, piston 52 and thumb plate 53, as shown by arrow 78 of FIG. 6.

The present invention permits needle 60 to be withdrawn back into body section 4 in a simple, straightforward manner. As shown in FIG. 7A, body section 4 has an elliptical cross-sectional shape with body slots 28 positioned through the walls of body 4. This permits the outer circumference of adapter ring 26 to enter body slots 28 so long as body section 4 is in its normal, typically undeformed condition of FIG. 7A. To disengage adapter ring 26 from body slots 28 thereby permitting its axial movement, the user 79 squeezes body 4 as suggested by arrows 80 in FIG. 7B opposite body slot 28. The appropriate place to squeeze body section 4 is further aided by
the provision of three different sets of tactile bumps 82, 83, 84, shown in FIG. 2, opposite body slots 28, 29 and 30. Once body section 4 has been sufficiently deformed to release adapter ring 26, cartridge-needle unit 46 can be moved axially relative to body section 4 from the I.M. injection position of FIG. 6 to the I.V. injection position of FIG. 8, at which adapter ring 26 engages body slots 29. As can be seen in FIG. 8, body slots 29 are positioned so that tip 62 of needle 60 is within interior 10 of body section 4, but near needle end 6.

Doing so permits syringe assembly 48 to be easily and safely used to inject a liquid into or withdraw a liquid from a membrane valve 86 of a conventional I.V. port 88 as shown in FIG. 8. The opening at needle end 6 is sized to properly guide injector arm 86 while eliminating any substantial possibility of an inadvertent needle stick during such a procedure.

FIG. 9A illustrates the engagement of adapter ring 26 within body slot 29. At FIG. 9B a modified adapter ring 26a is shown with a tapered plunger-facing surface 90. The use of surface 90 keeps adapter ring 26 snug within body slot 29 and also provides a ramp effect to make movement of adapter ring 26a towards plunger end 8 of body section 4 easier.

FIG. 10 illustrates a syringe assembly 48a identical to syringe assembly 48 but with a needle 60a being longer than needle 60. To properly position tip 62 relative to needle end 6, adapter 22 is positioned along body section 4 until adapter ring 26 engages body slots 30 rather than body slots 29.

FIGS. 12a and 11 illustrate further embodiments providing for the proper positioning of tips 62 of needles 60, 60a (the short and long needle versions, respectively). Short needle 60 is used with a conventional width adapter ring 26 as in FIG. 8. However, with syringe assembly 48b, shown in FIG. 11, extra width adapter ring 26b is used. Adapter ring 26b will engage body slots 28a, 29a and 30a but will not engage narrow width body slots 29. However, normal width adapter ring 26 can engage all of the body slots 28a, 29, 29a and 30a. Therefore, adapter ring 26b is repositioned from body slots 28a, past body slots 29 and into engagement with body slots 29a so to properly position tip 62. With syringe assembly 48c, shown
in FIG. 12, adapter ring 26 is initially positioned within body slots 29a (although somewhat loosely). Movement of adapter ring 26 from body slot 28a towards plunger end 8 allows adapter ring 26 to engage body slots 29 and thus properly position tip 62 in this case as well. FIG. 12b shows the use of an I.V. drip visual indicator 91 between body slots 29, 29a to show the user which body slot is to be used for use with an I.V. port 88. Other visual indicators or markings could be used adjacent the body slots used for I.M. injections and safe disposal.

FIGS. 13-16 illustrate another method by which different length needles can be accommodated, and also demonstrate that adapter ring 26 need not be a full circle. Syringe assembly 48d, illustrated in FIG. 13, uses a short length needle 60 while syringe assembly 48e uses long length needle 60a while both use a body section 4b. Body section 4b includes a pair of body slots 28, a single body slot 29b, a single body slot 29c and a pair of body slots 30b formed in a plunger end 8a. As can be seen in FIG. 14, adapter ring 26c is generally C-shaped and is sized to substantially impede or prevent rotation of adapter 22b within interior 10 of body section 4b when engaged within the body slot 29b. The axial and rotary position of body slot 29b is chosen to properly position tip 62 of needle 60. Similarly, with reference to FIGS. 15 and 16, adapter 22c has its adapter ring 26d positioned 180° from adapter ring 26c of FIG. 14 so that adapter ring 26d engages body slot 29c rather than slot 29b so tip 62 of needle 60a is also properly positioned relative to needle end 6 of body section 4b.

In both embodiments of FIGS. 13 and 15, plunger end 8a is thickened or strengthened. This helps to rigidify body section 4b at body slots 30a so that once adapter rings 26c, 26d enter body slots 30a, it will be quite difficult to squeeze plunger end 8a of body section 4b with sufficient force to disengage the adapter ring from the body slot.

FIGS. 17, 18A and 18B illustrates a situation in which the body slots 28, 29 and 30 are replaced by flanges 92, 93, 94 which engage an annular slot 96 formed in adapter 22d. The disengagement of flanges 92 from slot 96 occurs in the same
manner as discussed with reference to FIGS. 7A and 7B. That
is, the user squeezes body section 4c as shown by arrows 80 to
cause the disengagement of flanges 92 from slot 96. This
permits the relative axial movement of adapter 22d within body
section 4c so to reposition needle 60 within the interior 10 of
the body section.

FIG. 19 illustrates a still further alternative
embodiment of the syringe assembly 48 of FIG. 4. Syringe
assembly 48f is similar to syringe assembly 48 except that
adapter 22 is replaced by providing an adapter ring 26e as an
integral part of hub 64a. This eliminates the need for
adapter 22, but likely requires modification of a conventional
cartridge-needle unit.

Turning now to FIGS. 20 and 21, a syringe assembly
48g is shown adapted for use with a smaller sized cartridge-
needle unit 46a. One distinction between syringe assembly 48g
and syringe assembly 48 is the use of an extended length
adapter 22e, a portion 98 of which extends past finger grips 42
to permit user 79 to grasp portion 98 when placing the assembly
into an I.V. injection or a safe disposal condition.

Body section 4c of syringe assembly 48g includes a
pair of inwardly biased fingers 100 having protrusions 102
shaped to conform to the outer surface of portion 98 of adapter
22e. This helps to center adapter 22e, and cartridge-needle
unit 46a therewith, within interior 10 of body section 4c.
Fingers 100 could also be used to center and stabilize
cartridge-needle unit 46 in the other embodiments.

Other modifications and variations can be made to
disclosed embodiments without departing from the subject of the
invention as defined in the following claims. For example, the
invention is particularly adapted for use with a cartridge-
needle unit. However, the invention is also adaptable for use
with syringes and can be used with separately packaged
cartridges and cartridge-needle units as well. While most of
the disclosed embodiments use an elliptically shaped body
section, the body section could, however, be another shape,
such as generally triangular, as well. The deformation of the
body section could also be created by twisting or rotating the
adapter ring (or other radially extending member) to cause the radially extending member to disengage from a body slot. However, doing so would require the appropriate rotary manipulation of the pieces in addition to axial manipulation.

While enclosure unit 2 is, in the preferred embodiment, a one-piece molded structure, it may be desirable to make it from two or more pieces. For example, it may be desirable to make adapter 22 and cap 16 as separate pieces. Also, it may be desired to mount adapter 22 to hub 64 prior to positioning the cartridge-needle unit within the body section.
WHAT IS CLAIMED IS:

1. A syringe assembly comprising:
   a syringe structure;
   the syringe structure including a first positioning element;
   a hollow body housing at least a portion of the syringe structure;
   the body including an inner surface having a second positioning element formed thereat, the first and second positioning elements sized for mating engagement when aligned so to prevent relative axial movement of the body and the syringe structure; and
   the body including means for permitting at least a part of the body to be deformed radially when the radially extending member is to be removed from the recess to allow relative axial movement of the body and the syringe structure.

2. The assembly of claim 1 wherein one of the first and second positioning elements includes a radially extending member and the other of the first and second positioning elements includes a recess sized to accept the radially extending member.

3. The assembly of claim 2 wherein the first positioning element includes said radially extending member.

4. The assembly of claim 2 wherein the second positioning element includes said radially extending member.

5. The assembly of claim 1 wherein the syringe structure includes a cartridge-needle unit having a barrel, a piston mounted within the barrel, and a needle assembly mounted to the barrel.

6. The assembly of claim 5 wherein the syringe structure includes an adapter mounted to the cartridge-needle unit, the adapter including the first positioning member.
7. The assembly of claim 2 wherein the radially extending member includes an annular ring.

8. The assembly of claim 1 wherein the body has a substantially constant wall thickness.

9. The assembly of claim 1 wherein the inner surface is generally elliptical in cross-section.

10. The assembly of claim 9 wherein the outer surface is generally elliptical in cross-section.

11. The assembly of claim 9 wherein the body deformation permitting means includes the body constructed to be resilient and sufficiently deflectable through the application of a squeezing force by a user so to temporarily change the cross-sectional shape of the inner surface at the second positioning element from generally elliptical to generally circular thus releasing the first and second positioning elements.

12. The assembly of claim 3 wherein the recess includes a circumferentially extending slot formed through the body.

13. The assembly of claim 3 wherein the inner surface of the body includes a plurality of axially spaced-apart recesses.

14. A syringe assembly comprising:
   a syringe structure including a needle, the needle having a needle tip, and a radially extending ring;
   a hollow body housing at least a portion of the syringe structure;
   the body including a generally elliptically-shaped inner surface having first and second axially spaced-apart recesses formed thereat, the recesses and ring sized for mating
engagement when aligned so to prevent relative axial movement of the body and the syringe structure;
the first recess positioned so the needle tip is external of the body when the ring is engaged therewith, the second recess positioned so the needle tip is within the body when the ring is engaged therewith; and
the body being constructed to be resilient and sufficiently deflectable through the application of a squeezing force by a user so to temporarily change the cross-sectional shape of the inner surface from generally elliptical to generally circular thus releasing the ring to allow relative axial movement of the body and the syringe structure.

15. A disposable medical needle apparatus, for use with a needle assembly of the type including a radially extending member, comprising: a case having an elongated cavity surrounded by a longitudinally extending wall, the cavity having a first open end at a first longitudinal end of the wall and having a second open end at a second opposite longitudinal end of the wall, the case having a needle-passing opening in the second end, the case having a recess formed within the interior of the wall sized to accept the radially extending member, the case including means for permitting at least a part of the body section to be deformed radially when the radially extending member is to be removed from the recess to allow the needle assembly, after use, to be withdrawn from the needle-passing opening and held within the cavity.

16. The apparatus of claim 15 further comprising a needle assembly, having an elongated tubular needle with a sharp distal end, a barrel and a hub, the hub securing the tubular needle the hub to the barrel, the hub having an outwardly extending flange which acts as the radially extending member.

17. A self-resheathing safety needle comprising: a longitudinally extending case having a first open end and a second open end and having a wall extending between the first
and second ends thereby forming a cavity, a recess extending outwardly from the cavity, a needle assembly having an elongated needle, a barrel and a hub mounting the needle to the barrel, the hub having a radially extending connector for cooperating with the recess to releasably hold the hub in the case, thereby preventing outward egress of the needle assembly from the cavity through the first end of the case, and means for permitting the case to be deformed radially to allow the connector to be withdrawn from the recess to permit the needle assembly to move along within the cavity.

18. The needle of claim 17 including first and second of said recesses, the first recess positioned so that a portion of the needle is external of the cavity when the connector is engaged therewith, the second recess positioned so that the needle is entirely within the cavity when the connector is engaged therewith.

19. A disposable medical needle protection apparatus comprising:

- a tubular outer case having first and second open ends;
- a hub axially movable within the case between first and second positions which are proximate the first and second ends, respectively, the hub being adapted to mount a needle which extends past the first end when the hub is in the first position and which is wholly disposed within the case when the hub is in the second position;
- locking means for securing the hub and therewith the needle to the case when the hub is in the first and second positions and;
- release means operable by applying a force to the case for releasing the locking means to permit axial movement of the hub between the first and second positions;

whereby, following the use of the needle, it can be retracted wholly within the case by activating the release means and moving the hub to the second position to prevent accidental needle stick.
20. Apparatus according to claim 19 wherein the force applied to the release means is applied in a generally radial direction of the case.

21. Apparatus according to claim 20 including means proximate the second end of the case inhibiting the application of the radial force to discourage releasing the locking means after the hub has been retracted to the second position.

22. Apparatus according to claim 21, wherein the inhibiting means comprises a flange affixed to the case adjacent the second case end and projecting generally radially outwardly therefrom.

23. Apparatus according to claim 19, wherein the release means is defined by a tubular wall of the case dimensioned so that a first portion of the wall extends generally radially beyond a periphery of the hub and a second portion of the wall is in contact with the hub to permit relative movements in a radial direction between the hub and the wall when the force is applied to the wall.

24. Apparatus according to claim 23, wherein the wall and the hub define at least one projection and a cooperating slit oriented transverse to an axis of the case, at least one of which is proximate the second end of the case, the projection and the slit being arranged and dimensioned so that they engage each other when in alignment and thereby lock the hub to the case when the hub is in the second position, and so that the locking means is released by applying the force in a generally radially inward direction to the first portion of the wall.

25. A disposable medical needle protection apparatus comprising:

   a tubular outer case having first and second ends;
a hub longitudinally movably disposed within the case and movable between first and second positions proximate the first and second ends of the case, the hub being adapted to firmly hold a needle projecting from the hub in the direction of the first end so that the needle extends past the first end when the hub is in the first position and is wholly disposed within the case when the hub is in the second position; and releasable locking means defined by the case and the hub including first means securing the hub to the case when the hub is in either the first or second position and preventing relative longitudinal movements of the hub in the case, and second means operable by applying a generally radially directed force to an exterior of the casing for disengaging the first means to permit relative longitudinal movements of the hub in the casing between the first and second positions.

26. Apparatus according to claim 25 wherein the first means includes means for securing the hub to the case at a third position intermediate the first and second position, and wherein the second means is operable to disengage the hub from the case when it is in the third position to permit movement of the hub from the third to the second position.

27. A disposable medical needle protection apparatus comprising:

a tubular outer case having first and second ends, a longitudinal axis and an interior space which, in cross-section, has generally transverse major and minor dimensions, the case being constructed of a resiliently deformable material;

a generally cylindrical hub longitudinally movably disposed in the space for movement between first and second positions proximate the first and second ends of the case, the hub being adapted to securely hold a needle projecting from an end thereof so that the needle extends past the first end of the case when the hub is in the first position and is wholly disposed within the space when the hub is in the second position, the hub having a cross-section perpendicular to the
axis of the case which is less than the major dimension of the space and such that a portion of the hub is normally in contact with a portion of the case defining the minor dimension while a remainder of the hub is spaced from another portion of the case defining the major dimension; and

releasable locking means on the hub and the case defined by slits oriented transversely to the longitudinal axis of the case and projections formed to be extendable into the slits, there being at least one each of the slits and the projections and at least two of one of the slits and projections, the at least one of the slits and projections being defined by the hub and the at least two of the slits and projections being defined by the case and positioned proximate the first and second ends thereof so that the hub is in the first or the second position when the corresponding slits and projections are in engagement;

whereby the locking means is released to enable movement of the hub between the first and second positions by manually compressing the case in the direction of the major dimension of the space to thereby reduce the major dimension and correspondingly increase the minor dimension of the space and disengage the projection from the corresponding.

28. Apparatus according to claim 27 wherein there are at least three of either the projection or the slits formed in the case, the third one thereof being located intermediate the first and second positions whereby the hub can be locked to the case in a third, intermediate position.

29. A disposable medical needle protection apparatus comprising:

a generally oval, resiliently deformable outer case having first and second ends, a longitudinal axis and first and second slits formed in a relatively narrower portion of the case, oriented transversely to the axis and located proximate the first and second ends of the case;

a generally cylindrical hub adapted to securely hold a needle projecting from an end thereof, the hub having a
diameter selected so that the hub is in contact with the relatively narrower portion of the case and spaced from a relatively wider portion of the case, the hub further including a radially oriented projection adopted to be received within the slits in the case;

the slits and the projection being arranged so that the needle extends past the first end of the case when the projection engages the slit proximate the first end, and so that the needle is wholly disposed within the case and cannot be contacted from an exterior thereof when the projection engages the slit adjacent the second end of the case;

whereby following use of the needle it can be wholly retracted inside the case by applying external pressure to the relatively wider portion of the case adjacent the first end which correspondingly spreads apart the relatively narrower portion of the case and causes disengagement of the projection from the slit so that, thereafter, the hub can be axially moved towards the second end until the projection engages the slit in adjacent the second end of the case.

30. Apparatus according to claim 29 including means defined by the case proximate the first end preventing axial movement of the hub past the first end.

31. Apparatus according to claim 29 wherein the hub has a generally tubular configuration and includes first and second, axially oriented apertures, a needle holder including the needle and means on the hub engageable with the holder a for substantially immovably securing the holder to the hub.

32. Apparatus according to claim 31 wherein the needle holder includes means for connecting it with an end of a syringe insertable into the case.

33. Apparatus according to claim 32 wherein the connecting means includes means for disengaging the syringe from the needle holder.
34. Apparatus according to claim 32 wherein the case has a length so that an end of the syringe projects past the second end of the case when the projection of the hub engages the slit in the case adjacent the first end thereof, whereby the hub can be moved from adjacent the first end to adjacent the second end by compressing the relatively wider portion of the case and pulling the projecting end of the syringe out of the case.
### INTERNATIONAL SEARCH REPORT

**International Application No.** PCT/US91/06878

**CLASSIFICATION OF SUBJECT MATTER**

According to International Patent Classification (IPC) or to both National Classification and IPC

**IPC(5):** A61M 5/32  
**U.S. CL.:** 644/198

**FIELD SEARCHED**

- **U.S. CL.:** 604/110, 192, 197, 198, 240, 203

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**DOCUMENTS CONSIDERED TO BE RELEVANT**

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**IV. CERTIFICATION**

**Date of the Actual Completion of the International Search:** 07 APRIL 1992  
**Date of Mailing of the International Search Report:** 2 MAY 1992  
**International Searching Authority:** ISA/US  
**Signature of Authorized Officer:** ADÁM J. CERMÁK

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