Abstract: A catheter assembly (10) includes a protective tip (14) defining an interior chamber (20) between its proximal and distal ends. A protective lubricious sleeve (24) is positioned within the interior chamber. A catheter (12) of the assembly is configured to be advanced proximally into and through the interior chamber to position at least a portion of the catheter within the protective sleeve, with the protective sleeve being retained upon the catheter as at least a proximal portion of the catheter exits the proximal end of the protective tip and is advanced through a body lumen. A second sleeve (106) may be associated with the protective tip and configured to extend distally from the protective tip to remain outside of the body lumen during use and receive a more distal portion of the catheter.
CATHETER ASSEMBLIES HAVING A PROTECTIVE LUBRICIOUS SLEEVE

RELATED APPLICATION

This application claims the benefit of and priority of U.S. Provisional Patent Application Serial No. 61/925,292, filed January 9, 2014, the contents of which are incorporated by reference herein.

DESCRIPTION

TECHNICAL FIELD

The present disclosure generally relates to catheters. More particularly, the present disclosure relates to catheters provided with a sleeve for lubrication and protection during insertion into a body lumen.

BACKGROUND

Intermittent catheterization is a good option for many users who suffer from various abnormalities of the urinary system. Such catheters are typically provided as single use, individually packaged items and may include a gel-lubricant or hydrophilic coating that may be hydrated to act as a lubricant for reducing friction during insertion into the urethra.

Regarding gel-coated catheters, a user applies a gel-lubricant, such as a water-based gel-lubricant, to the surface of the catheter, which reduces friction for ease of insertion into the urethra. In some instances, the gel-lubricant is supplied with the packaged catheter, in which case the gel-lubricant may be applied to the catheter surface just before or during the packaging operation or as the catheter is being inserted by the user.

When a hydrophilic material is used as a lubricant, a thin coating of hydrophilic material is adhered to the outer surface of the catheter. When this coating is activated by swelling in contact with a hydrating liquid such as water, it provides a surface having an extremely low coefficient of friction. One form of this product provides a sterile, individually packaged, single-use catheter in a dry state or condition. The user opens the package, pours water into the package, waits 30 seconds, and then removes the catheter from the package, now ready for
insertion. Other embodiments provide the amount of liquid water necessary for immersion of the catheter in a separate compartment of the package. In such embodiments, the user must open the separate compartment of the package to allow the liquid immersion water to enter the catheter-containing chamber for direct contact with the hydrophilic coated surface. The catheter is then removed from the package and inserted into the urethra. In yet another embodiment, the catheter is provided in a package that already contains enough loose liquid water to cause it to be immersed. In such an embodiment, the user simply opens the package and removes the catheter therefrom, and then inserts the catheter into the urethra, without the need to add water.

A disadvantage of the gel-coated and hydrophilic coated catheters described above is that the gel-lubricant may get on the hands of the user during handling or the immersion liquid may spill from the package as the user handles the catheter and tries to remove it for subsequent insertion.

SUMMARY

There are several aspects of the present subject matter which may be embodied separately or together in the devices and systems described and claimed below. These aspects may be employed alone or in combination with other aspects of the subject matter described herein, and the description of these aspects together is not intended to preclude the use of these aspects separately or the claiming of such aspects separately or in different combinations as set forth in the claims appended hereto.

In one aspect, a catheter assembly kit includes a protective tip defining an interior chamber between its proximal and distal ends. A protective lubricious sleeve is positioned within the interior chamber. A catheter of the kit is configured to be advanced proximally into and through the interior chamber to position at least a portion of the catheter within the protective sleeve, with the protective sleeve being retained upon the catheter as at least a proximal portion of the catheter exits the proximal end of the protective tip.

In another aspect, a method is provided for applying a protective lubricious sleeve to a catheter. The method includes providing a protective tip defining an interior chamber between proximal and distal ends of the protective tip, with a
protective lubricious sleeve positioned within the interior chamber. At least a proximal portion of a catheter is advanced proximally into the interior chamber of the protective tip via the distal end of the protective tip. At least the proximal portion of the catheter is positioned within the sleeve as it is proximally advanced through the interior chamber. The proximal portion of the catheter is further proximally advanced out of the interior chamber via the proximal end of the protective tip, with the sleeve being retained on the catheter and a portion of the sleeve exiting the interior chamber with the proximal portion of the catheter.

In yet another aspect, a catheter assembly kit includes a protective tip with proximal and distal ends, with a protective lubricious sleeve secured to the protective tip. A catheter of the kit is configured to be advanced into contact with the protective sleeve for advancement through the protective tip from the distal end of the protective tip toward the proximal end of the protective tip to position at least a portion of the catheter within the protective sleeve, with the protective sleeve being retained upon the catheter as at least a proximal portion of the catheter exits the proximal end of the protective tip.

**BRIEF DESCRIPTION OF THE DRAWINGS**

Fig. 1 is a perspective view of a proximal portion of an embodiment of a catheter assembly according to an aspect of the present disclosure;

Fig. 2 is a side elevational view of the proximal portion of the catheter assembly of Fig. 1;

Fig. 3 is a perspective view of the proximal portion of the catheter assembly of Fig. 1, with a catheter thereof in a proximally advanced position;

Fig. 4 is a side elevational view of the proximal portion of the catheter assembly of Fig. 1, with a catheter thereof in a proximally advanced position;

Fig. 5 is a perspective view of a proximal portion of a protective lubricious sleeve of the catheter assembly of Fig. 1;

Fig. 6 is a perspective view of a proximal portion of another embodiment of a protective lubricious sleeve;

Fig. 7 is a perspective view of a proximal portion of yet another embodiment of a protective lubricious sleeve;

Figs. 8 and 9 are side elevational views of a protective tip and protective sleeve assembly.
lubricious sleeve of a catheter assembly according to an aspect of the present disclosure, with the sleeve in a folded configuration;

Figs. 10 and 11 are perspective views of an alternative embodiment of a catheter assembly according to an aspect of the present disclosure;

Fig. 12 is a detail view of an insert and a distal portion of a protective lubricious sleeve of the catheter assembly of Figs. 10 and 11;

Fig. 13 is a side elevational view of an alternative embodiment of a catheter assembly according to an aspect of the present disclosure;

Fig. 14 is a perspective view of the catheter assembly of Fig. 13;

Figs. 15 and 16 are perspective views of an alternative embodiment of a catheter assembly according to an aspect of the present disclosure;

Figs. 17 and 18 are perspective views of an alternative embodiment of a catheter assembly according to an aspect of the present disclosure;

Fig. 19 is a front elevational view of a package containing a protective tip and protective lubricious sleeve of a catheter assembly kit;

Fig. 20 is a side elevational view of an alternative embodiment of a catheter assembly according to an aspect of the present disclosure;

Fig. 21 is a perspective view of the catheter assembly of Fig. 20, with a catheter thereof in an assembled, proximally advanced condition;

Fig. 22 is perspective view of a proximal portion of an alternative embodiment of a catheter and protective lubricious sleeve;

Fig. 23 is a cross-sectional view of the catheter and sleeve of Fig. 22, taken though line 23-23 of Fig. 22;

Fig. 24 is a perspective view of a portion of the catheter of Fig. 22;

Fig. 25 is a perspective view of an alternative embodiment of a catheter assembly according to an aspect of the present disclosure; and

Fig. 26 is a perspective view of an alternative embodiment of a catheter assembly according to an aspect of the present disclosure.

**DETAILED DESCRIPTION OF THE ILLUSTRATED EMBODIMENTS**

The embodiments disclosed herein are for the purpose of providing a description of the present subject matter, and it is understood that the subject matter may be embodied in various other forms and combinations not shown in
detail. Therefore, specific embodiments and features disclosed herein are not to be interpreted as limiting the subject matter as defined in the accompanying claims.

Figs. 1-4 illustrate an embodiment of a catheter assembly 10, such as a urinary catheter assembly. The catheter assembly 10 may be variously configured without departing from the scope of the present disclosure, but in one embodiment, the catheter assembly 10 is provided as a kit, which includes a catheter 12 (such as an uncoated, polymeric urinary catheter) and a separate protective tip 14. The kit may include additional components (e.g., a fluid drainage bag or receptacle) without departing from the scope of the present disclosure.

The protective tip 14 extends between a distal end 16 and a proximal end 18, defining an interior chamber 20 therebetween. The proximal end 18 of the protective tip 14 may optionally be configured for insertion into a body lumen (e.g., a urethral opening) prior to advancement of the catheter 12 into the body lumen. The proximal end 18 of the protective tip 14 may include an aperture or opening 22 that may be moved between a closed configuration (in which there is no object positioned within the opening 22, as in Figs. 1 and 2) and an open configuration (in which the catheter 12 or any other object is partially positioned within or extending through the opening 22, with a portion of the object positioned within the protective tip 14 and another portion positioned outside of the protective tip 14, as in Figs. 3 and 4). In one embodiment, the proximal opening 22 is provided as a slit opening with one or more slits or cuts defining a plurality of deformable petals that may be moved to define the aforementioned open and closed configurations.

In other embodiments, the opening 22 may be differently configured, provided that it is configured to allow passage of the catheter 12 therethrough. The distal end 16 of the protective tip 14 may also include an opening for passage of the catheter 12, with the opening at the distal end 16 being either movable from a closed configuration to an open configuration (similar to the proximal opening 22) or provided in always-open position.

A protective lubricious sleeve 24 is at least partially (but preferably completely) positioned within the interior chamber 20 of the protective tip 14. In use, the catheter 12 is advanced proximally into, through, and out of the interior
chamber 20. As the catheter 12 moves into the interior chamber 20 via the distal opening of the protective tip 14, it enters into the interior of the sleeve 24. Further advancing the catheter 12 proximally through the interior chamber 20 causes the catheter 12 (typically the proximal end 26 of the catheter 12) to engage the sleeve 24 (typically the proximal end 28 of the sleeve 24), with the sleeve 24 being retained on the catheter 12. With the sleeve 24 retained on the catheter 12, further proximal advancement of the catheter 12 with respect to the protective tip 14 causes at least a proximal portion 30 of the sleeve 24 to exit the protective tip 14 with the proximal end 28 of the catheter 12 (Figs. 3 and 4). In a preferred embodiment, the proximal end 18 of the protective tip 14 is positioned within a body lumen (e.g., a urethra) prior to advancement of the catheter 12 and sleeve 24 out of the protective tip 14.

The sleeve 24 provides a barrier which prevents bacteria on the catheter 12 from contacting a body lumen (e.g., a urethra) as the catheter 12 is advanced into and through the body lumen. Additionally, the sleeve 24 provides lubricity to aid catheter insertion. The sleeve 24 may be an inherently lubricious film or may have a lubricious coating applied thereto. For example, the sleeve 24 may be made from a hydrophilic polymer that becomes lubricious when wetted with a wetting agent or fluid. In such an embodiment, a wetting agent or fluid may be placed or located within the interior chamber 20 of the protective tip 14, as will be described in greater detail herein. Similarly, in embodiments in which the sleeve 24 has a lubricious coating, such as a gel-lubricant, the lubricant may be contained within the interior chamber 20 of the protective tip 14, with the coating being applied to the sleeve 24 when the sleeve 24 is placed into the protective tip 14. Alternatively, a sleeve 24 that is pre-coated with a lubricant may be positioned within the interior chamber 20 of the protective tip 14. By providing a sleeve 24 which covers the catheter 12 as it is advanced out of the protective tip 14, the catheter 12 may be uncoated or otherwise omit a lubricious coating and may be directly handled by a user without the risk of bacteria being transferred to the catheter 12 and then from the catheter 12 to the aforementioned body lumen. An uncoated, non-lubricious catheter 12 may be advantageous by being more readily gripped and manipulated than a lubricated catheter, thereby improving and simplifying catheter insertion.
The proximal end 28 of the sleeve 12 may be variously configured, depending on the configuration of the associated catheter 12. In one embodiment, the catheter 12 includes one or more side openings or eyes or drainage portions 32 at a proximal portion 34 of the catheter 12. The eyes 32 allow fluid (e.g., urine) to drain into and through the tubular catheter 12 from the proximal portion 34 of the catheter 12 to a distal portion 36 of the catheter 12, where it may exit the catheter 12 (e.g., via a funnel or opening). In one embodiment, which is shown in Fig. 5, the proximal end 28 of the sleeve 24 is closed to fully encircle and enclose the proximal end 26 and proximal portion 34 of the catheter 12. To allow drainage of fluid from a body location (e.g., the bladder) into and through the catheter 12, the sleeve 24 may include a mesh portion 38 configured to overlay at least a portion of the eyes 32 when the proximal end 26 of the catheter 12 is pressed against and/or adjacent to the proximal end 28 of the sleeve 24.

Fig. 6 shows a variation of the sleeve 24 of Fig. 5. In the embodiment of Fig. 6, the proximal portion 30a of the sleeve 24a includes a perforated portion 40, rather than a mesh portion 38. Other fluid-permeable portions besides a mesh or perforated portion (such as open portions having a size and shape similar to the catheter eyes 32) may also be incorporated into the proximal portion of protective lubricious sleeves according to the present disclosure.

Fig. 7 illustrates another embodiment of a proximal portion 30b of a protective lubricious sleeve 24b. In the embodiment of Fig. 7, the sleeve 24b is configured to be retained upon the catheter 12 at only a portion distal of the eyes 32, such that the eyes 32 remain uncovered by the sleeve 24b. The sleeve 24b may either be configured to be retained upon the catheter 12 upon advancement of the catheter 12 through the protective tip 14 or may be secured (e.g., by a heat seal) to the catheter 12 during manufacturing. In embodiments such as Fig. 7, in which a proximal end 34 of the catheter 12 remains uncovered by the sleeve 24b, it may be advantageous for the proximal end 26 and uncovered portion of the catheter 12 to be lubricated to aid catheter insertion.

It should be understood that the illustrated sleeves are merely exemplary, and other protective lubricious sleeves may be provided without departing from the scope of the present disclosure. For example, in another embodiment, the
proximal portion of a protective sleeve may be similar to the sleeves 24 and 24a of Fig. 5 and 6 to the extent that it covers the entirety of the proximal end of the associated catheter 12, but omits any openings for allowing fluid to pass through the sleeve and enter into the catheter eyes 32. Instead, the sleeve may include a frangible or weakened portion that initially prevents fluid flow through the sleeve, but may be broken or otherwise manipulated to define an at least partially open configuration during use to allow fluid flow into the catheter eyes. In such an embodiment, the catheter and sleeve may be moved through a body lumen to a target location (e.g., moving through a male urethra until the proximal ends of the catheter and sleeve enter into the bladder), at which time the frangible portion may be broken (e.g., by distal relative movement of the sleeve with respect to the catheter) to define an opening or passage or otherwise allow fluid flow through or around the proximal end of the sleeve and into the catheter eyes. For example, the proximal end of the sleeve may have a weakened section, which may be punctured by the proximal end of the catheter upon proximal relative movement of the catheter with respect to the sleeve. In such a configuration, it may be advantageous for the catheter to have a greater length than the sleeve, to allow the catheter to be advanced farther into the body lumen than the sleeve, but with the sleeve having a particular minimum length (e.g., a length approximately equal to the length of the urethra).

In variations of the foregoing design, rather than including a frangible or weakened portion, the distal portion of the sleeve may be otherwise configured to change from a closed configuration to an at least partially open configuration during use to selectively allow fluid flow into the catheter eyes. For example, the proximal end of the sleeve may be defined by an elastic or deformable cuff or endpiece similar to the proximal end 28d of the sleeve 24d of Figs. 15 and 16. By such a configuration, an opening defined in the elastic cuff overlays a closed portion of the catheter during advancement of the sleeve and catheter through a body lumen. When the proximal end of the catheter is in place, the sleeve may be moved distally with respect to the catheter, which causes the elastic cuff to deform and slide distally along the catheter until the opening is positioned distally of the catheter eyes. In this position, with the catheter eyes uncovered by the sleeve, fluid may flow into the catheter eyes and drained out of the body by the catheter.
In other variations, perforated lines, peelable seals, and expandable perforations are among the variety of mechanisms that may be incorporated into a sleeve to allow a portion of its proximal end to fracture or otherwise move from a closed configuration to an at least partially open configuration.

The sleeve 24 preferably has a greater length than the interior chamber 20, to allow the proximal portion 30 of the sleeve 24 to extend outside of the protective tip 14, while a distal portion 42 remains within the interior chamber 20 (as described above). Accordingly, the sleeve may be provided in a folded or bunched configuration. For example, Fig. 1 shows the sleeve 24 in a concertina-style formation within the interior chamber 20 so that the entire sleeve 24 may be positioned within the interior chamber 20. Figs. 8 and 9 illustrate embodiments in which the sleeve 24c is provided in a folded configuration. In the embodiment of Fig. 9, the folded sleeve 24 is positioned within the interior chamber 20a of the protective tip 14a, whereas the protective tip 14a’ of Fig. 8 omits an enclosed interior chamber. Instead, the distal portion 20a’ of the protective tip 14a’ of Fig. 9 is generally tubular, with an open distal end. The associated sleeve 24c may be secured to the open distal end of the protective tip 14a’ or to any other suitable area of the distal portion 20a’ of the protective tip 14a’. The sleeve 24c is inverted in the orientation shown in Fig. 8, with the lubricated surface of the sleeve 24c facing inwardly, such that the sleeve 24c may be handled without contacting the lubricant on the sleeve 24c. In use, the proximal end of the associated catheter is aligned with the proximal end of the sleeve 24c and advanced into and through the protective tip 14a’, entering at the open distal end of the distal portion 20a’. As the sleeve 24c and catheter exit the proximal end of the protective tip 14a’, the lubricated surface of the sleeve 24c (i.e., the surface facing inwardly in the orientation of Fig. 8) will be facing outwardly for advancement into and through a body lumen. Other configurations of the sleeve initially positioned either within the interior chamber of the protective tip or outside of the protective tip may also be practiced without departing from the scope of the present disclosure.

As for the distal end of the sleeve, it may be unsecured to the associated protective tip, but it may be preferred for the sleeve to be secured or connected to the protective tip. In the embodiment of Figs. 1-4, the distal end 44 of the sleeve 24 is secured to the distal end 16 of the protective tip 14. In the embodiment of
Figs. 8 and 9, the distal end 44c is secured to a more proximal portion of the protective tip 14a (Fig. 9). These configurations are merely exemplary and, in other embodiments, a bunched sleeve 24 or a folded sleeve 24c may be secured to the associated protective tip 14, 14a at any suitable location.

Regardless of whether or not the sleeve is secured to the associated protective tip, the distal end of the sleeve may be open or openable, with the distal opening of the protective tip leading into the interior of the sleeve. By such a configuration, a catheter that is proximally advanced into the interior chamber of the protective tip via the distal opening of the protective tip will move into the interior of the sleeve for advancement of the catheter and sleeve into a body lumen, as described above.

Figs. 10-12 show an embodiment of a catheter assembly 10a in which the distal end of the protective tip 14b is defined by a separate grommet or insert 46, with the distal end 44 of the sleeve 24 being secured to the insert 46, such as by heat sealing. Such an embodiment may be preferred to an integrally formed or single-piece protective tip, in terms of ease of manufacturability, because it may be easier to position the sleeve 24 within the interior chamber 20b of the protective tip 14b.

Fig. 12 illustrates the proximal side of the insert 46, with the distal end 44 of the sleeve 24 being secured to the insert 46 around a central opening 48 of the insert 46, such that a catheter 12 being moved proximally into and through the insert opening 48 will pass into the interior of the sleeve 24.

The illustrated insert 46 is generally annular, with a groove or recess 50 along its outer perimeter. In such a configuration, the inner surface of the protective tip 14b may be provided with an inwardly extending, annular projection 52, which allows the insert 46 to be press-fit into place within the protective tip 14b, with the projection 52 being seated within the groove 50 to retain the insert 46. In other embodiments, the insert (if provided) may be differently configured and secured to the protective tip by different means without departing from the scope of the present disclosure. In general, it may be preferred for the insert to include a central opening to allow for passage of a catheter therethrough and into the interior of the sleeve, with an outer perimeter of the insert having a shape that matches the shape of the open distal end of the protective tip. It may be preferred
for the shape of the perimeter of the insert to match the shape of the open distal end of the protective tip in order to provide a sterile seal between the protective tip and the insert. The insert may be fixedly or removably secured to the protective tip by any suitable means, such as a friction fit or an adhesive or the like.

Figs. 13 and 14 illustrate a catheter assembly 10b having the same catheter 12, protective tip 14b, and sleeve 24 as the embodiment of Figs. 10 and 11, but a different insert 46a. In the embodiment of Figs. 13 and 14, the proximal side of the insert 46a includes a generally tubular alignment barrel 54 surrounding the central opening 48. In this embodiment, the distal end 44 of the sleeve 24 may be secured to either the outer surface of the alignment barrel 54 (as illustrated) or to the proximal side of the insert 46a. The alignment barrel 54 helps to guide the proximal end 26 of the catheter 12 into the proximal end 28 and interior of the sleeve 24 as the catheter 12 is advanced into and through the interior chamber 20b of the protective tip 14b.

Figs. 15 and 16 illustrate an embodiment of a catheter assembly 10c having an alternative catheter 12a. The catheter 12a of Figs. 15 and 16 (also shown in Fig. 9) omits side openings or drainage portions, but instead includes an opening or eye or drainage portion 32a at its proximal end 26a. The associated sleeve 24d may also include an opening 56 at its proximal end 28d to become at least generally aligned with the eye 32a of the catheter 12a when the proximal end 26a of the catheter 12a is pressed against the proximal end 28d of the sleeve 24d (Fig. 16). In the illustrated embodiment, the proximal end 28d of the sleeve 24d is defined by an elastic or deformable cuff or endpiece that is separately formed and secured to the body of the sleeve 24d, but it is also within the scope of the present disclosure for the sleeve 24d to be formed as a single-piece device. Preferably, the sleeve opening 56 is larger than the catheter eye 32a so that the sleeve 24d does not obstruct the flow of fluid into the catheter 12a via the eye 32a, but it is also within the scope of the present disclosure for the sleeve opening 56 to be substantially the same size as or smaller than the catheter eye 32a.

Alternatively, rather than a distal opening 56, the distal end 28d of the sleeve 24d may include a mesh portion or a perforated portion to allow fluid flow into the eye 32a at the proximal end 26a of the catheter 12a. Further, other fluid-permeable portions besides a mesh or perforated portion may also be incorporated into the
proximal end 28d of the sleeve 24d.

Figs. 17 and 18 illustrate a catheter assembly 100 having a protective tip 102 of the type shown in Figs. 13-16, with a catheter (not illustrated) and a sleeve 104 of the type shown in Figs. 15 and 16. While the catheter assembly 100 of Figs. 17 and 18 is provided with a catheter and sleeve 104 of the type shown in Figs. 15 and 16 in a preferred embodiment, it is within the scope of the present disclosure for the catheter assembly 100 to have a differently configured catheter and/or sleeve, such as one of the various catheters and sleeves described herein.

The embodiment of Figs. 17 and 18 further includes a distal or auxiliary sleeve 106, which may be provided as a flexible, deformable tubular body formed of a sheet or film material or the like. In contrast to the sleeve 104, the distal sleeve 106 is not configured to be advanced into the urethra, so the same considerations (e.g., lubricity) that affect the design of the sleeve 104 may have lesser importance when designing the distal sleeve 106. Accordingly, the material composition and/or surface treatment of the distal sleeve 106 may be different from composition and/or surface treatment of the sleeve 104. For example, the distal sleeve 106 may be handled by a user to manipulate a distal portion of the catheter during use of the catheter assembly 100, in which case it may be advantageous for the distal sleeve 106 to be non-lubricious. While the distal sleeve 106 may be formed of a different material and/or with a different surface treatment than the sleeve 104, it is also within the scope of the present disclosure for the distal sleeve 106 to be formed of the same material and/or with the same surface treatment as the sleeve 104.

The distal sleeve 106 is secured to the alignment barrel 108 (e.g., secured around the inner surface or perimeter of the alignment barrel 108) and/or to some other portion of the protective tip 102, with at least a portion of the distal sleeve 106 initially positioned within the alignment barrel 108, as in Fig. 17. Preferably, all or at least the majority of the distal sleeve 106 is initially positioned within the alignment barrel 108, but it is also within the scope of the present disclosure for less than half of the distal sleeve 106 to be initially positioned within the alignment barrel 108.

A portion of the distal sleeve 106 is distally advanced out of the alignment barrel 108, as shown in Fig. 18, to surround and receive the distal portion of an
associated catheter, while the proximal portion of the catheter is positioned within the sleeve 104, as shown in Fig. 16. By such a configuration, the entire catheter may be received within the two sleeves 104 and 106, although it is also within the scope of the present disclosure for the proximal end and/or distal end of the catheter to be positioned outside of the sleeves 104 and 106.

The distal end 110 of the distal sleeve 106 may have a fluid drainage funnel 112 or comparable drainage device associated therewith, with the funnel 112 being used to drain fluid from a catheter positioned within the sleeves 104 and 106 into a toilet or other disposal location. If the distal sleeve 106 is provided with a funnel 112, then a simplified catheter omitting a funnel may be provided for use in combination with the protective tip 102. It may be advantageous for the funnel 112 and distal end of the catheter to be provided with mating formations, such that the funnel 112 may be press-fit onto the distal end of the catheter or otherwise secured in place with respect to the distal end of the catheter.

A proximal portion or end 114 of the funnel 112 may be initially fixed with respect to the protective tip 102 (e.g., with part or all of the proximal portion or end 114 received within the alignment barrel 108), as shown in Fig. 17. The funnel 112 may be gripped and pulled distally away from the protective tip 102 to move the distal sleeve 106 from the arrangement of Fig. 17 to the arrangement of Fig. 18. The distal sleeve 106 may be extended away from the protective tip 102 prior to the proximal end of a catheter being proximally advanced into the funnel 112, through the distal sleeve 106, and into the sleeve 104. Alternatively, the proximal end of a catheter may be proximally advanced into the funnel 112, through the distal sleeve 106 (positioned within the alignment barrel 108), and into the sleeve 104 prior to extending the distal sleeve 106 away from the protective tip 102. In either case, with the catheter positioned within the sleeves 104 and 106 (and the distal sleeve 106 in the extended arrangement of Fig. 18), the proximal end of the protective tip 102 may be positioned within the urethra and then the catheter may be advanced proximally out of the proximal end of the protective tip 102 and into the urethra. In another embodiment, the proximal end of the protective tip 102 may be positioned within the urethra prior to advancing the catheter into the funnel 112 and/or prior to moving the distal sleeve 106 into the extended arrangement of Fig. 18. The catheter may pinched or gripped through the distal sleeve 106 by the
user to further advance the catheter through the urethra or, if the funnel 112 is
secured to the distal end of the catheter, the funnel 112 may be gripped and
moved proximally to further advance the catheter through the urethra. After use,
the catheter may be disposed of or cleaned and reused, with the protective tip 102
preferably being discarded.

In one embodiment, the catheter and protective tip are provided as
separate components of a catheter assembly kit, which may be combined to
define a catheter assembly. For example, Fig. 19 shows a protective tip 14
housed within a sealed package 58. The package 58 may include a frangible tear
line 60 to allow a user to open the package 58 and remove the protective tip 14 for
use with a separate catheter. When the protective tip 14 has been removed from
the package 58, the proximal end of a catheter may be proximally advanced into
and through the interior chamber of the protective tip 14, as described in detail
above. After use, the entire catheter assembly may be discarded. Alternatively,
one or more components of the catheter assembly may be reused. For example,
the protective tip may be discarded after use, while the catheter may be reused
(optionally after being washed, rinsed, or re-sterilized) with a new protective tip.
Providing a reusable catheter may promote the use and provision of a catheter
having more advanced features and functionality, as being able to reuse the
catheter may render a more expensive catheter more cost-effective for users with
budgetary restrictions.

In another embodiment, a catheter assembly kit may include a protective
tip with the proximal end of a catheter pre-loaded into the interior chamber of the
protective tip, inside the sleeve. Figs. 20 and 21 illustrate one such embodiment
of a catheter assembly 10d in which a portion of the catheter 12b is pre-loaded
within the protective tip. In the embodiment of Figs. 20 and 21, the catheter 12b
includes separate proximal and distal members 62 and 64. The distal end 66 of
the proximal member 62 and the proximal end 68 of the distal member 64 each
include a fitting or connection point 70, 72 that is configured to mate with the fitting
of the other member of the catheter 12b (Fig. 21). The fittings 70 and 72 may be
variously configured, provided that they form a fluid passage between the
proximal and distal members 62 and 64. Preferably, the fittings 70 and 72 are
configured such that the proximal and distal catheter members 62 and 64 may be
temporarily or removably connected to each other, but it is also within the scope of the present disclosure for the fittings 70 and 72 to fixedly secure the proximal and distal catheter members 62 and 64 together.

Prior to use, at least a proximal end 74 of the proximal member 62 may be pre-loaded within an interior chamber 20 of a protective tip 14 of the catheter assembly 10d, with at least the distal end 66 of the proximal member 62 positioned outside of the protective tip 14 (Fig. 20). The protective tip 14 and proximal catheter member 62 may be provided in a sealed package (similar to the package 58 of Fig. 19) prior to use. When the protective tip 14 and proximal catheter member 62 have been removed from the package (if provided), the distal catheter member 64 may be connected to the proximal member 62 using the fittings 70 and 72. With the catheter 12b fully assembled (as in Fig. 21), the catheter 12b may be proximally advanced to exit the interior chamber 20 of the protective tip 14, with the sleeve 24 retained upon the catheter 12b. After use, the entire catheter assembly 10d may be discarded. Alternatively, one or more components of the catheter assembly 10d may be reused. For example, the protective tip 14 and proximal catheter member 62 may be discarded after use, while the distal catheter member 64 may be reused (optionally after being washed, rinsed, or re-sterilized) with a new protective tip 14 and proximal catheter member 62.

The illustrated configuration of the proximal and distal catheter members 62 and 64 is merely exemplary, and the proximal and distal members 62 and 64 may be differently configured without departing from the scope of the present disclosure. For example, in the illustrated embodiment, the proximal member 62 includes side or lateral eyes 32b, but it is also within the scope of the present disclosure for the proximal member 62 to include a drainage eye at its distal end, as in the embodiment of Figs. 15 and 16. Similarly, the distal member 64 is illustrated with a fluid drainage funnel 76 at its distal end and a gripping aid or handle 78 (which may be separate pieces or part of an integrated unit that is secured to the distal member 64), but either or both of those components could be omitted or additional components may be included (e.g., a fluid drainage bag or receptacle).

In one embodiment, the proximal and distal catheter members 62 and 64
are formed of different materials and/or have different stiffness. For example, it may be advantageous for the distal member 64 to have a greater stiffness or rigidity than the proximal member 62, because the proximal member 62 will often be required to traverse a relatively tortuous body lumen (e.g., a male urethra), whereas the distal member 64 traverses a shorter, less tortuous section of the body lumen, and the enhanced rigidity helps to advance the assembled catheter 12b through the body lumen. However, it is also within the scope of the present disclosure for the proximal and distal catheter members 62 and 64 to be formed of the same material and to have the same stiffness.

Figs. 22-24 illustrate yet another embodiment of a catheter 12c that may be incorporated into catheter assemblies of the present disclosure. In the illustrated embodiment, at least a portion 80 of the catheter 12c is non-tubular, but instead allows for fluid flow along one or more paths or drainage portions 82 defined between an outer surface of the non-tubular portion 80 and an inner surface of the associated protective lubrious sleeve 24e (Fig. 23). The non-tubular portion 80 of the catheter 12c may have any suitable cross-sectional shape, but in the illustrated embodiment has a generally cross- or X-shaped cross-section, which defines four external drainage portions 82. The sleeve 24e associated with the catheter 12c may include at least one fluid-permeable section 84 (e.g., an open section or a mesh or perforated section) to allow fluid flow into the interior of the sleeve 24e. Preferably, as shown in Figs. 22 and 23, the sleeve 24e includes a plurality of fluid-permeable sections 84, with each drainage portion 82 having an associated fluid-permeable section 84 at least partially aligned therewith. A more distal section of the catheter 12c may be generally hollow or tubular, such that the fluid flowing through the external drainage portions 82 eventually enters into the interior of the catheter 12c, or the non-tubular portion 80 of the catheter 12c may extend to the distal end of the catheter 12c, such that fluid never enters into the interior of the catheter 12c.

As described above, the protective lubrious sleeve may be an inherently lubrious thin film or its lubricity may be provided by a coating, such as a gel-, water-, or oil-based layer. Fig. 25 illustrates an example of a protective tip 14c and sleeve 24f of a catheter assembly kit which employs a fluid lubricant for lubricating the sleeve 24f. The protective tip 14c of Fig. 25 is similar to the
embodiment of Figs. 10 and 11, except that a fluid lubricant 86 is placed into the
interior chamber 20c of the protective tip 14c (surrounding the sleeve 24f) prior to
securing the insert 46b to the protective tip 14c. The fluid lubricant 86 maintains
the lubricity of the sleeve 24f for an extended period of time, allowing long-term
storage of the catheter assembly kit prior to use. The lubricant may be a gel-type
lubricant or, when the sleeve is formed of or includes a hydrophilic material, the
lubricant may be a wetting agent or fluid, such as water or saline. The central
opening 48b of the insert 46b may include a fluid-tight seal 88 (which may be
similar to the opening at the proximal end of the protective tip) to prevent leakage
of the fluid lubricant 86 out of the interior chamber 20c. The seal 88 may be
pierceable or otherwise movable from a closed condition to an open condition to
allow a catheter to pass proximally into and through the interior chamber 20c. Any
other protective tip described herein may be provided with a fluid-tight seal, such
as a fluid-tight seal 88 of the type shown in Fig. 25, at its distal end or

In another embodiment, when the protective lubricious sleeve is formed of
or includes a hydrophilic material, the sleeve may be wetted by one or more
hydration sachets 90 positioned within the interior chamber 20d of the protective
tip 14d, as in Fig. 26. In the illustrated embodiment, two substantially identical
hydration sachets 90 are provided, with each inserted into a pocket 92 defined in
or adjacent to the interior chamber 20d of the protective tip 14d prior to securing
the associated insert 46c to the protective tip 14d. The insert 46c may include an
extension 94 associated with each pocket 92, with each extension 94 configured
to press the associated hydration sachet 90 into the pocket 92 and maintain the
hydration sachet 90 therein. Each hydration sachet 90 contains a hydration
substance 96 that acts to hydrate and lubricate the sleeve (not illustrated) within
the interior chamber 20d, thus allowing long-term storage of the catheter
assembly kit prior to use. In one embodiment, the hydration sachets may be fluid-
containing pouches, with the pouches being formed of a water vapor-permeable,
liquid-impermeable material, such as calcium carbonate. The hydration sachets
90 and associated pockets 92 and insert extensions 94 are illustrated as being
elongated, with generally arcuate cross-sectional shapes, but it is within the scope
of the present disclosure for the hydration sachets 90 and associated pockets 92
and insert extensions 94 to be otherwise configured.

Aspects of the present subject matter described above may be beneficial alone or in combination with one or more other aspects. Without limiting the foregoing description, in accordance with one aspect of the subject matter herein, there is provided a catheter assembly kit, which includes a protective tip defining an interior chamber between proximal and distal ends of the protective tip. A protective lubricious sleeve is positioned within the interior chamber. A catheter is configured to be advanced proximally into and through the interior chamber to position at least a portion of the catheter within the protective sleeve, with the protective sleeve being retained upon the catheter as at least a proximal portion of the catheter exits the proximal end of the protective tip.

In accordance with another aspect which may be used or combined with the preceding aspect, the distal end of the protective tip is defined by an insert defining a distal opening configured to allow proximal advancement of the catheter into the interior chamber, with a distal end of the sleeve being secured to the insert and the distal opening in communication with an interior of the sleeve.

In accordance with another aspect which may be used or combined with the preceding aspect, the insert includes a generally tubular alignment barrel surrounding the distal opening and positioned within the interior chamber and within the interior of the sleeve.

In accordance with another aspect which may be used or combined with the preceding aspect, the distal end of the sleeve is secured to an outer surface of the alignment barrel.

In accordance with another aspect which may be used or combined with the any of the preceding two aspects, at least a portion of a distal sleeve is positioned within the alignment barrel. At least a portion of the distal sleeve is configured to be advanced distally out of the alignment barrel to surround a distal portion of the catheter.

In accordance with another aspect which may be used or combined with the preceding aspect, a fluid drainage funnel is secured to the distal sleeve.

In accordance with another aspect which may be used or combined with the any of the preceding aspects, the catheter includes at least one drainage portion defined in a sidewall of the catheter.
In accordance with another aspect which may be used or combined with any of the first six aspects, the catheter includes at least one drainage portion defined in a proximal end of the catheter.

In accordance with another aspect which may be used or combined with any of the first six aspects, at least a portion of the catheter is substantially non-tubular, with a generally cross-shaped cross-section defining at least one drainage portion.

In accordance with another aspect which may be used or combined with any of the preceding three aspects, the sleeve includes a perforated portion configured to overlay at least a portion of the drainage portion as the proximal portion of the catheter exits the proximal end of the protective tip.

In accordance with another aspect which may be used or combined with any of the seventh through ninth aspects, the sleeve includes a mesh portion configured to overlay at least a portion of the drainage portion as the proximal portion of the catheter exits the proximal end of the protective tip.

In accordance with another aspect which may be used or combined with any of the seventh through eighth aspects, the sleeve is configured to be retained upon the catheter at only a portion distal of the drainage portion.

In accordance with another aspect which may be used or combined with the preceding aspect, the sleeve is heat-sealed to the catheter distally of the drainage portion, with the catheter being lubricated proximally of the location at which the sleeve is sealed to the catheter.

In accordance with another aspect which may be used or combined with any of the preceding aspects, the catheter comprises a proximal member and a separate distal member.

In accordance with another aspect which may be used or combined with the preceding aspect, at least a proximal end of the proximal member of the catheter is pre-loaded within the interior chamber and the sleeve, with at least a distal end of the proximal member of the catheter positioned outside of the interior chamber.

In accordance with another aspect which may be used or combined with any of the preceding two aspects, the proximal member of the catheter has a different stiffness than the distal member of the catheter.
In accordance with another aspect which may be used or combined with any of the preceding three aspects, the distal member of the catheter has a greater stiffness than the proximal member of the catheter.

In accordance with another aspect which may be used or combined with any of the preceding aspects, at least one hydration sachet is positioned within the interior chamber.

In accordance with another aspect which may be used or combined with any of the first seventeen aspects, a wetting agent and/or lubricant is positioned within the interior chamber, exteriorly of the sleeve.

In accordance with another aspect which may be used or combined with any of the preceding aspects, the sleeve is provided in a concertina-style formation within the interior chamber.

In accordance with another aspect which may be used or combined with any of the first nineteen aspects, the sleeve is provided in a folded formation within the interior chamber.

In accordance with another aspect, there is provided a method of applying a protective lubricious sleeve to a catheter. The method includes providing a protective tip defining an interior chamber between proximal and distal ends of the protective tip, with a protective lubricious sleeve positioned within the interior chamber. At least a proximal portion of the catheter is proximally advanced into the interior chamber of the protective tip via the distal end of the protective tip to position the proximal portion of the catheter within the sleeve. The proximal portion of the catheter is advanced out of the interior chamber via the proximal end of the protective tip, with the sleeve being retained on the catheter and a portion of the sleeve exiting the interior chamber with the proximal portion of the catheter.

In accordance with another aspect which may be used or combined with the preceding aspect, the protective tip is provided with a distal sleeve secured to it. At least a portion of the distal sleeve is distally advanced to surround a distal portion of the catheter.

In accordance with another aspect, there is provided a catheter assembly kit, which includes a protective tip having proximal and distal ends, with a protective lubricious sleeve secured to the protective tip. A catheter is configured
to be advanced into contact with the protective sleeve for advancement through the protective tip from the distal end of the protective tip toward the proximal end of the protective tip to position at least a portion of the catheter within the protective sleeve, with the protective sleeve being retained upon the catheter as at least a proximal portion of the catheter exits the proximal end of the protective tip.

It will be understood that the embodiments described above are illustrative of some of the applications of the principles of the present subject matter. Numerous modifications may be made by those skilled in the art without departing from the spirit and scope of the claimed subject matter, including those combinations of features that are individually disclosed or claimed herein. For these reasons, the scope hereof is not limited to the above description but is as set forth in the following claims, and it is understood that claims may be directed to the features hereof, including as combinations of features that are individually disclosed or claimed herein.
CLAIMS
1. A catheter assembly kit, comprising:
   a protective tip defining an interior chamber between proximal and distal ends of the protective tip;
   a protective lubricious sleeve positioned within the interior chamber; and
   a catheter configured to be advanced proximally into and through the interior chamber to position at least a portion of the catheter within the protective sleeve, with the protective sleeve being retained upon the catheter as at least a proximal portion of the catheter exits the proximal end of the protective tip.

2. The catheter assembly kit of claim 1, wherein the distal end of the protective tip is defined by an insert defining a distal opening configured to allow proximal advancement of the catheter into the interior chamber, with a distal end of the sleeve secured to the insert and the distal opening in communication with an interior of the sleeve.

3. The catheter assembly kit of claim 2, wherein the insert includes a generally tubular alignment barrel surrounding the distal opening and positioned within the interior chamber and within the interior of the sleeve.

4. The catheter assembly kit of claim 3, wherein the distal end of the sleeve is secured to an outer surface of the alignment barrel.

5. The catheter assembly kit of any of claims 3-4, further comprising a distal sleeve, wherein
   at least a portion of the distal sleeve is positioned within the alignment barrel, and
   at least a portion of the distal sleeve is configured to be advanced distally out of the alignment barrel to surround a distal portion of the catheter.

6. The catheter assembly kit of claim 5, further comprising a fluid drainage funnel secured to the distal sleeve.
7. The catheter assembly kit of any of the preceding claims, wherein the catheter includes at least one drainage portion defined in a sidewall of the catheter.

8. The catheter assembly kit of any of claims 1-6, wherein the catheter includes at least one drainage portion defined in a proximal end of the catheter.

9. The catheter assembly kit of any of claims 1-6, wherein at least a portion of the catheter is substantially non-tubular, with a generally cross-shaped cross-section defining at least one drainage portion.

10. The catheter assembly kit of any of claims 7-9, wherein the sleeve includes a perforated portion configured to overlay at least a portion of said at least one drainage portion as said at least a proximal portion of the catheter exits the proximal end of the protective tip.

11. The catheter assembly kit of any of claims 7-9, wherein the sleeve includes a mesh portion configured to overlay at least a portion of said at least one drainage portion as said at least a proximal portion of the catheter exits the proximal end of the protective tip.

12. The catheter assembly kit of any of claims 7-8, wherein the sleeve is configured to be retained upon the catheter at only a portion distal of said at least one drainage portion.

13. The catheter assembly kit of claim 12, wherein the sleeve is heat-sealed to the catheter distally of said at least one drainage portion, with the catheter being lubricated proximally of the location at which the sleeve is sealed to the catheter.

14. The catheter assembly kit of any of the preceding claims, wherein the catheter comprises a proximal member and a separate distal member.
15. The catheter assembly kit of claim 14, wherein at least a proximal end of the proximal member of the catheter is pre-loaded within the interior chamber and the sleeve, with at least a distal end of the proximal member of the catheter positioned outside of the interior chamber.

16. The catheter assembly kit of any of claims 14-15, wherein the proximal member of the catheter has a different stiffness than the distal member of the catheter.

17. The catheter assembly kit of any of claims 14-16, wherein the distal member of the catheter has a greater stiffness than the proximal member of the catheter.

18. The catheter assembly kit of any of the preceding claims, further comprising at least one hydration sachet positioned within the interior chamber.

19. The catheter assembly kit of any of claims 1-17, further comprising a wetting agent and/or lubricant positioned within the interior chamber, exteriorly of the sleeve.

20. The catheter assembly kit of any of the preceding claims, wherein the sleeve is provided in a concertina-style formation within the interior chamber.

21. The catheter assembly kit of any of claims 1-19, wherein the sleeve is provided in a folded formation within the interior chamber.

22. A method of applying a protective lubricious sleeve to a catheter, comprising:
   - providing a protective tip defining an interior chamber between proximal and distal ends of the protective tip, with a protective lubricious sleeve positioned within the interior chamber;
   - proximally advancing at least a proximal portion of a catheter into the interior chamber of the protective tip via the distal end of the protective tip;
positioning said at least the proximal portion of the catheter within the sleeve as the catheter is proximally advanced through the interior chamber; and advancing said at least the proximal portion of the catheter out of the interior chamber via the proximal end of the protective tip, with the sleeve being retained on the catheter and a portion of the sleeve exiting the interior chamber with said at least the proximal portion of the catheter.

23. The method of claim 22, wherein said providing a protective tip includes providing a protective tip with a distal sleeve secured thereto, and further comprising distally advancing at least a portion of the distal sleeve to surround a distal portion of the catheter.

24. A catheter assembly kit, comprising:
   a protective tip including proximal and distal ends;
   a protective lubricious sleeve secured to the protective tip; and
   a catheter configured to be advanced into contact with the protective sleeve for advancement through the protective tip from the distal end of the protective tip toward the proximal end of the protective tip to position at least a portion of the catheter within the protective sleeve, with the protective sleeve being retained upon the catheter as at least a proximal portion of the catheter exits the proximal end of the protective tip.
**INTERNATIONAL SEARCH REPORT**

**A. CLASSIFICATION OF SUBJECT MATTER**

INV. A61M 25/00, A61M 25/01

ADD.

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

**B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)

A61M

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

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

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**X** See patent family annex.

**X** Further documents are listed in the continuation of Box C.

* Special categories of cited documents:

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

- "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
- "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
- "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
- "A" document member of the same patent family

Date of the actual completion of the international search: 19 March 2015

Date of mailing of the international search report: 12/05/2015

Name and mailing address of the ISA:
European Patent Office, P.B. 5818 Patentlaan 2
NL-2280 HV RIJSWIJK
Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016

Authorized officer: Rodri gues, El odio e

Form PCT/ISA/210 (second sheet) (April 2005)
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INTERNATIONAL SEARCH REPORT

Box No. II  Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. ☑ Claims Nos.: 22, 23
   because they relate to subject matter not required to be searched by this Authority, namely:
   see FURTHER INFORMATION sheet PCT/ISA/210

2. ☐ Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

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

Box No. III  Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

This international searching Authority found multiple inventions in this international application, as follows:

see additional sheet

1. ☐ As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.

2. ☑ As all searchable claims could be searched without effort justifying an additional fees, this Authority did not invite payment of additional fees.

3. ☐ As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.: 

4. ☑ No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-4, 7-17, 20, 21, 24

Remark on Protest

☐ The additional search fees were accompanied by the applicant's protest and, where applicable, the payment of a protest fee.

☐ The additional search fees were accompanied by the applicant's protest but the applicable protest fee was not paid within the time limit specified in the invitation.

☐ No protest accompanied the payment of additional search fees.
Continuation of Box II.1

Claims Nos.: 22, 23

Rule 39.1(iv) PCT - Method for treatment of the human or animal body by surgery. The methods of applying a protective lubricious sleeve claimed in claims 22 and 23 do not explicitly exclude the possibility that such application occurs as the catheter is being introduced into the human or animal body. This possibility appears highly desirable in the light of the description. These methods thus comprise surgical methods.
Further information continued from PCT/ISA/ 210

This international Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1-4, 7-17, 20, 21, 24
   Catheter assembly kit

1.1. Claims: 1-4, 24
   Catheter assembly kit comprising a protective tip

1.2. Claims: 7-9, 14, 15
   Catheter assembly kit comprising a catheter of a particular construction

1.3. Claims: 10-13, 20, 21
   Catheter assembly kit comprising a sleeve of a particular construction

1.4. Claims: 16, 17
   Catheter assembly kit comprising a catheter of a particular stiffness

2. Claims: 5, 6
   Catheter assembly kit comprising a distal sleeve

3. Claims: 18, 19
   Catheter assembly kit comprising wetting elements
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