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(54) Title: ENDOSCOPIC MUCOSAL RESECTION SINGLE STEP HOOD

(57) Abstract: The present disclosure provides systems and methods for cutting mucosal tissue. In particular, the present disclosure provides an endoscopic hood with an integrated wire for single-step tissue cutting. The endoscopic hood is particularly useful for performing repeated cutting, for example EMR resections, without the need for assistance by a nurse or medical technician.
ENDOSCOPIC MUCOSAL RESECTION SINGLE STEP HOOD

Cross-Reference to Related Applications

[0001] This application claims the benefit of U.S. Provisional Application No. 62/164,313, filed May 20, 2015, the disclosure of which is incorporated herein by reference in its entirety.

Technical Field

[0002] The present disclosure relates to the field of endoscopy. Specifically, the present disclosure relates to systems and methods for cutting, for example resecting and/or dissecting, abnormal or diseased mucosal tissue. More specifically, the present disclosure relates to an endoscopic hood designed to create and cut a pseudo-polyp in a single step.

Background

[0003] Organ walls are composed of several layers: the mucosa (surface layer), the submucosa, the muscularis (muscle layer) and the serosa (connective tissue layer). A variety of lesions comprising dead, diseased or abnormal tissue may form on the mucosal walls of the colon, esophagus, stomach and duodenum. For example, gastrointestinal, colonic and esophageal cancers may form within the mucosal layer and manifest as a polyp or tissue mass that extends into the lumen of the respective organ.

[0004] Endoscopic mucosal resection (EMR) is a minimally invasive technique by which cancerous or otherwise abnormal tissues are resected without disrupting the integrity of the organ wall. EMR is generally performed using an endoscope that includes a long narrow tube equipped with a light, video camera and one or more channels to receive a variety of medical instruments. During an EMR procedure, the endoscope is passed down the esophagus or guided through the rectum to the site
of a cancerous or abnormal tissue within the mucosal wall of the target organ. The distal end of the endoscope is equipped with an endoscopic hood that is positioned over the tissue to be resected. Once properly positioned, suction is applied to the endoscope to draw the target tissue into the endoscopic hood, where it is then resected using a variety of techniques known in the art. The excised tissue is then extracted from the endoscope for examination and/or disposal.

[0005] Currently available EMR systems use suction to draw the target tissue into the endoscopic hood and deploy an elastic band around the base of the resulting pseudo-polyp. The suction is then released and the EMR hood and endoscope are pulled proximally to free the pseudo-polyp. A resection snare is passed through the working channel of the endoscope and manipulated to capture and resect the pseudo-polyp from the surrounding tissue. Once freed, the resected tissue is recaptured within the EMR hood using suction, or by introducing a separate grasping element (i.e., basket, forceps etc.) through the endoscope working channel. As multiple elastic bands are provided with such devices, several resections may be performed using the same EMR hood.

[0006] Other EMR systems do not require elastic bands to create the pseudo-polyp. Instead, a snare is deployed around the inside of the EMR hood so that the tissue can be resected immediately following formation of the pseudo-polyp. A major drawback of this system, however, is the minimal recovery afforded by the snare once the resection has been performed. This requires a new snare to be inserted and deployed within the EMR hood for each subsequent tissue resection.

[0007] Although the EMR systems described above use different techniques to form and resect the pseudo-polyp, they are similar in that both require multiple users to simultaneously control the endoscope and resection snare. In a typical EMR
procedure the endoscope is controlled/maneuvered by a physician while the
snaring/resecting steps are performed by a nurse. For example, when the physician
has determined that the target tissue is properly positioned within the EMR hood,
he/she must instruct the nurse to tighten the snare around the pseudo-polyp,
followed by an instruction to apply cauterization energy. Once the target tissue is
fully resected, the physician then instructs the nurse to cease cauterization and
retract the snare. As one might expect, this process limits the physician's tactile
sense for the procedure and requires a significant amount of communication with the
nurse. This disclosure is related to an EMR hood that allows the physician to create
and resect a pseudo-polyp in a single step without the need for additional
assistance.

**SUMMARY**

[0008] Particular aspects of the disclosure are described in the Summary and
Detailed Description, below. Although the disclosure has been described in
connection with specific aspects, it should be understood that the disclosure as
claimed should not be unduly limited to such specific aspects.

[0009] In one aspect, the present disclosure relates to an endoscopic hood
configured to be disposed at a distal end of an endoscope. The endoscopic hood
includes a proximal portion, a distal portion, a lumen extending between the proximal
and distal portions, a wire track disposed about an inner circumference of the distal
portion, and a wire channel extending between the proximal and distal portions. The
wire channel is in communication with the wire track. For example, the wire channel
and the wire track may merge with each other at a gradual angle such that the wire
channel and wire track can receive a wire. The wire channel may extend along an
inner wall of the endoscopic hood. Alternatively, the wire channel may extend along
an outer wall of the endoscopic hood. The wire track may include a planar surface
that is substantially perpendicular to the inner wall of the endoscopic hood. The wire
track may be formed as a groove disposed within the inner wall of the endoscopic
hood. An anchoring element may be positioned adjacent to the site at which the wire
channel merges with the wire track. An end of the wire may be secured (i.e., tied,
anchored, welded etc.) to the anchoring element. The anchoring element may also
include an opening or aperture to slidably receive a portion of the wire. An
elastomeric sleeve may be configured to form an interference fit with the outer
surface of the endoscopic hood. The elastomeric sleeve may include a distal cap
that is coextensive with and substantially parallel to the wire track. The elastomeric
sleeve may enclose the wire channel that extends along the outer wall of the
endoscopic hood. An endoscope may be attached to the proximal portion of the
endoscopic hood. The endoscope may include at least one working channel in
communication with the lumen of the endoscopic hood.

[0010] In another aspect, the present disclosure relates to a tissue cutting
system including an endoscope, an endoscopic hood coupled to the distal end of the
endoscope, an actuation handle coupled to the proximal end of the elongate member
and a wire that forms a loop about the wire track and extends proximally along the
wire channel to the actuation handle. The endoscope includes a proximal end, a
distal end and a lumen extending therebetween. The endoscopic hood includes a
proximal portion, a distal portion and a lumen extending between the proximal and
distal portions. A wire track is disposed about an inner circumference of the distal
portion of the endoscopic hood. A wire channel extends between the proximal and
distal portions of the endoscopic hood, and is in communication with the wire track.
The wire channel and wire track are configured to receive a wire. A protective
elongate member may enclose at least a portion of the wire extends along the length of the endoscope between the endoscopic hood and the actuation handle. The protective elongate member may extend along the exterior surface of the endoscope. Alternatively, the protective elongate member may extend along the lumen of the endoscope. A portion of the wire may be configured to move from an open-loop configuration to a closed-loop configuration. For example, retracting the actuation handle may move a portion of the wire into a closed loop configuration and advancing the actuation handle may move a portion of the wire into an open-loop configuration.

[0011] In yet another aspect, the present disclosure relates to a method of cutting tissue, including inserting, into a body lumen of a patient, a medical device comprising an endoscope, an endoscopic hood coupled to the distal end of the endoscope, an actuation handle coupled to the proximal end of the elongate member and a wire that forms a loop about the wire track and extends proximally along the wire channel to the actuation handle. The endoscope includes a proximal end, a distal end and a lumen extending therebetween. The endoscopic hood includes a proximal portion, a distal portion and a lumen extending between the proximal and distal portions. A wire track is disposed about an inner circumference of the distal portion of the endoscopic hood. A wire channel extends between the proximal and distal portions of the endoscopic hood, and is in communication with the wire track. The wire channel and wire track are configured to receive a wire. The endoscopic hood is positioned over the surface of the target tissue. Once properly positioned, the target tissue is drawn into the lumen of the endoscopic hood. The wire is then moved into a closed-loop configuration around the tissue. Cauterization energy is then applied to the wire to cut the tissue. Once the tissue has been cut, the wire is
returned to an open-loop configuration. Fluid may be injected in or around the target tissue before the target tissue is drawn into the lumen of the endoscopic hood. The endoscopic hood may be repositioned over the surface of a second target tissue after the first target tissue has been cut. These steps may be repeated as necessary to remove multiple target tissues.

**BRIEF DESCRIPTION OF THE DRAWINGS**

[0012] Non-limiting embodiments of the present disclosure are described by way of example with reference to the accompanying figures, which are schematic and not intended to be drawn to scale. In the figures, each identical or nearly identical component illustrated is typically represented by a single numeral. For purposes of clarity, not every component is labeled in every figure, nor is every component of each embodiment of the disclosure shown where illustration is not necessary to allow those of ordinary skill in the art to understand the disclosure. In the figures:

[0013] FIG. 1A is a side perspective view of an endoscopic hood, according to an embodiment of the present disclosure.

[0014] FIG. 1B is a top perspective view of the endoscopic hood of FIG. 1A.

[0015] FIG. 2A is a side view of an outer sleeve configured to slide over the endoscopic hood, according to an embodiment of the present disclosure.

[0016] FIG. 2B is a top perspective view of the outer sleeve of FIG. 2A forming an interference fit with the endoscopic hood, according to an embodiment of the present disclosure.

[0017] FIG. 3 is a side view of the endoscopic hood and outer sleeve of FIG. 2B connected to an actuation handle by a protective elongate member, according to an embodiment of the present disclosure.
FIG. 4A is a side perspective view of the resection wire in an open-loop configuration when the actuation handle of FIG. 3 is in the extended configuration.

FIG. 4B is a side perspective view of the resection wire in a partially closed-loop configuration when the actuation handle of FIG. 3 is in the retracted configuration.

FIG. 4C is a side perspective view of the resection wire in a closed-loop configuration when the actuation handle of FIG. 3 is in the retracted configuration.

FIGS. 5A-5F depict the steps involved in resecting a target mucosal tissue using a single step endoscopic hood, in accordance with an embodiment of the present disclosure.

It is noted that the drawings are intended to depict only typical or exemplary embodiments of the disclosure. It is further noted that the drawings may not be necessarily to scale. Accordingly, the drawings should not be considered as limiting the scope of the disclosure. The disclosure will now be described in greater detail with reference to the accompanying drawings.

DETAILED DESCRIPTION

Before the present disclosure is described in further detail, it is to be understood that the disclosure is not limited to the particular embodiments described, as such may vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodiments only, and is not intended to be limiting beyond the scope of the appended claims. Unless defined otherwise, all technical terms used herein have the same meaning as commonly understood by one or ordinary skill in the art to which the disclosure belongs. Finally, although embodiments of the present disclosure are described with specific reference to
systems and method for cutting mucosal tissue using an endoscope, it should be appreciated that the endoscopic hood of the present disclosure may be applicable to cutting a variety of tissues using a variety of introduction devices, sheaths or systems, such as trocars, catheters, laparoscopes, colonoscopes, ureteroscopes and the like. As used herein, the term "distal" refers to the end farthest away from a medical professional when introducing a device into a patient, while the term "proximal" refers to the end closest to the medical professional when introducing a device into a patient. As used herein, the term "cutting" may include any suitable type of cutting, including resection performed as part of an endoscopic mucosal resection procedure. In some instances, the terms "cutting" and "resection" may be used interchangeably. It should be understood, however, that aspects of the disclosure may also be applicable to other types of cutting, such as dissection as part of an endoscopic submucosal dissection procedure.

[0024] FIG. 1A depicts an endoscopic hood 10 according to one embodiment of the present disclosure. The endoscopic hood 10 is preferably substantially cylindrical and optionally includes a proximal portion 11, a distal portion 12 and a lumen 13 (i.e., working channel) extending therebetween. The lumen 13 is defined by an inner wall 14 with a preferably substantially constant inner diameter. The endoscopic hood further may include an outer wall 15 with a preferably varying (e.g., tapered) outer diameter. The distal portion 12 is configured to permit tissue to be drawn into the lumen 13 of the endoscopic hood 10 using endoscopic suction. A substantially circular resection wire track 18 may be disposed about an inner circumference of the inner wall 14 at the distal portion 12 of the endoscopic hood 10. The resection wire track 18 preferably includes a planar surface that is preferably substantially perpendicular to the inner wall 14 of the endoscopic hood 10. The
resection wire track 18 may be in communication with a resection wire channel 19 that preferably extends along the outer wall 15 of the endoscopic hood 10 between the proximal 11 and distal 12 portions. The resection wire track 18 and resection wire channel 19 are preferably configured to receive at least a portion of a resection wire 40.

[0025] The resection wire 40 is comprised of an electrically conductive material with sufficient flexibility that it slides freely along the resection wire track 18, resection wire channel 19 and protective elongate member 29 (FIG. 3) without kinking, bending or otherwise developing sites of fatigue. In one embodiment, the resection wire 40 includes a diameter of at least 0.25 mm, more preferably at least 0.50 mm and even more preferably at least 1.00 mm to provide sufficient stiffness (i.e., pushability) along its entire length such that advancing one end of the resection wire causes the opposite end of the resection wire to advance without any portion of the resection wire bending or kinking.

[0026] Materials suitable for use as a resection wire include electrically conductive metals or alloys selected, for example, from platinum group metals, particularly platinum, rhodium, palladium, and rhenium, as well as tungsten, gold, silver, tantalum, and alloys of these metals including platinum/tungsten alloys and nickel-titanium alloys (nitinol) among others. The resection wire may be formed from a monofilament material (e.g., monofilament nitinol) or a braided material as are known in the art. In one embodiment, the ability of the resection wire 40 to repeatedly move between an open-loop configuration and closed-loop configuration may be further enhanced by heat treating and/or mechanically shaping a distal portion of the resection wire such that it assumes the shape of the resection wire
track 18 and/or resection wire channel 19 when in the relaxed/unconstrained
configuration.

[0027] In the open-loop configuration, the resection wire 40 is secured to a
distal portion of the endoscopic hood 10, travels around the planar surface of the
resection wire track 18 to form a loop, and continues proximally along the resection
wire channel 19 through a protective elongate member 29 to an actuation handle 30
(see FIG. 3). As best shown in FIG. 1B, a first end 41 of the resection wire 40 is
secured to an anchoring element 20 positioned adjacent to the site at which the
resection wire track 18 and resection wire channel 19 merge. The anchoring element
20 includes an aperture 20a through which the resection wire 40 passes as it
proceeds from the resection wire track 18 to the resection wire channel 19. The
gradual angle at which the resection wire track 18 and resection wire channel 19
merge/intersect allows the resection wire 40 to smoothly transition between an open-
loop configuration and closed-loop configuration as the actuation handle is extended
and retracted, respectively. Referring again to FIG. 1A, the outer wall 15 of the
proximal portion 11 of the endoscopic hood 10 further includes an outwardly
extending tab 16 with an opening 16a therethrough. The tab 16 serves as an
attachment point for a protective elongate member 29 that connects the endoscopic
hood 10 to the actuation handle 30 (as shown in FIG. 3). The resection wire 40
passes through the opening 16a of the outwardly extending tab 16 and extends
through the protective elongate member 29 to the actuation handle 30.

[0028] Still referring to FIG. 1A, the outer wall 15 of endoscopic hood 10
includes a generally decreasing taper from the proximal portion 11 to distal portion 12
to allow the endoscopic hood to navigate body passageways in a minimally invasive
manner. As shown in FIG. 2A, an outer sleeve 22 is configured to slide over the
outer wall 15 of the endoscopic hood 10 to fully enclose the resection wire channel 19. The outer sleeve 22 includes a proximal portion 23, a distal portion 24 and a lumen 25 extending therebetween. The lumen 25 of the outer sleeve 22 is defined by an inner wall with a varying diameter that corresponds to the taper of the outer wall 15 of the endoscopic hood. The inner diameter of the lumen 25 is undersized as compared to the corresponding outer wall of the endoscopic hood. The outer sleeve 22 is comprised of a sufficiently elastomeric material, including, but not limited to, rubber or silicone, such that it expands when positioned over the endoscopic hood, thereby forming an interference fit with the underlying endoscopic hood. The outer sleeve 22 further includes a raised portion 28 that presses against the tab 16 and guides the resection wire into the opening 16a of tab 16.

[0029] Still referring to FIG. 2A, the distal end of the sleeve 22 also includes an inwardly projecting cap 21 that is coextensive with, and substantially parallel to, the resection wire track 18. As best shown in FIG. 2B, when the sleeve 22 is positioned over the endoscopic hood 10, the cap 21 covers, but does not contact, the resection wire track 18 to form a groove within which the resection wire 40 lies. Enclosing the resection wire channel 19 with the outer sleeve 22 prevents the resection wire 40 from jumping out of the resection wire channel 19 as the actuation handle (not shown) is extended or retracted. Similarly, partially enclosing the resection wire track 18 with the cap 21 ensures that repeated movement of the resection wire between the open-loop configuration and closed-loop configuration occurs in a plane perpendicular to the lumen 13 of the endoscopic hood 10 (see FIGS. 4A-B).

[0030] Referring to FIG. 3, the tab 16 serves as an attachment point for a protective elongate member 29 that extends between the endoscopic hood 10 and
the actuation handle 30. A portion of the resection wire 40 passes through the opening 16a of the tab 16 and through the lumen of the protective elongate member 29 to the actuation handle 30. The second end 42 of the resection wire 40 is secured to the actuation handle 30 such that the resection wire 40 moves between an open-loop configuration (FIG. 4A) and a closed-loop configuration (FIGS. 4B-C) as the actuation handle 30 is extended and retracted, respectively. In one embodiment, the protective elongate member 29 may be configured to travel through a working channel of the endoscope (not shown). In another embodiment, the protective elongate member 29 may extend along an exterior surface of the endoscope, thereby providing more room within the endoscope working channel for the introduction of additional medical instruments.

[0031] As will be understood by one of skill in the art, the depth of the cut made by the resecting wire loop is critical. If the cut is too deep the muscularis layer may be injured, possibly leading to a perforation. Conversely, a cut that is too shallow may not remove all of the affected tissue, such that additional procedures are required or, worse, contributing to the development of metastatic cancer. Typically, more than 2.0 mm of target tissue clearance is required to assure complete removal. FIGS. 5A-5F illustrate the use of the endoscopic hood 10 in performing an EMR procedure. As shown, a physician may introduce the endoscopic hood 10 attached to the distal end of an endoscope (not shown) into a patient's body through a natural anatomical opening or an incision. Referring to FIGS. 5A-B, the distal portion 12 of the endoscopic hood 10 is positioned over and in contact with the mucosal layer 34 of the target tissue 31. As depicted in FIG. 5A, the target tissue 31 (shaded area) may include a raised region or polyp that extends outward from the mucosal surface. Polyps, such as pedunculated polyps, may be characterized by a
stalk attached to the mucosal layer. Such polyps are easily drawn into the
endoscopic hood with relatively little suction. Alternatively, as depicted in FIG. 5B,
the target tissue may lie substantially flat, or only slightly raised, along the mucosal
layer 34 of the target tissue. For example, certain other polyps, such as sessile
polyps, may exhibit a broad base that is devoid of any stalk portion such that they lay
substantially flat on the mucosal surface. Such polyps are often difficult to grasp
without applying an amount of suction that draws in part of the underlying muscularis
layer.

[0032] In one embodiment, a fluid such as a gel, saline solution, hypertonic
sugar, indigo carmine, ethylene blue or the like is injected beneath the target
tissue to form a bleb, thereby raising the target tissue. The raised tissue can then be
drawn into the endoscopic hood for resection. In some instances, the target tissue
may be too large to remove in a single step, and must be removed by segmental
resection, in which repeated fluid injections and subsequent tissue removal are
performed along the entire length of the affected area.

[0033] Referring to FIG. 5C, once the endoscope is properly positioned, the
physician applies vacuum suction to draw the mucosal layer 34 of the target tissue
(and a portion of surrounding healthy tissue) into the lumen 13 of the endoscopic
hood 10. In one embodiment, the vacuum suction is provided through the endoscope
(i.e., endoscope suction). In another embodiment, the vacuum source is provided
through the protective elongate member 29. In yet another embodiment, the vacuum
suction is provided by a separate tube running through the working channel of the
endoscope or along an outer surface of the endoscope.

[0034] Referring to FIG. 5D, with the vacuum source maintaining the target
tissue within the lumen 13 of the endoscopic hood 10, the physician retracts the
actuation handle 30 such that the resection wire 40 disposed about the resection wire track 18 in an open-loop configuration moves towards a closed-loop configuration to tighten around the outer surface of the target tissue. Once the physician determines that the resection wire 40 has sufficiently tightened around the target tissue, cauterization energy is applied to the resection wire from an energy source (not shown) to cauterize the tissue. As shown in FIG. 5E, the cauterization energy applied to the tightened loop 33 of the resection wire 40 is sufficient to resect (i.e. cut) the target tissue free from the mucosal layer 34 without cutting either the submucosa 36 or muscularis 38 layers. In the event that the target tissue is not released by the first application of cauterization energy, the physician may further retract the actuation handle 30 such that the resection wire 40 further tightens around the target tissue, followed by a second application of cauterization energy. These steps may be repeated as necessary to cut the target tissue free from the mucosal layer. Alternatively, the physician may apply cauterization energy while the actuation handle 30 is being retracted such that the target tissue is cut as the resection wire 40 moves into an increasingly small closed-loop configuration. It should be appreciated that the cauterization energy applied to the resection wire 40 creates a cauterization zone that extends beyond the diameter of the resection wire. This allows the resection wire 40 to cut entirely through the target tissue without necessarily having to move the resection wire into its smallest possible closed-loop configuration. The ability to simultaneously control the actuation handle 30 and the application of cauterization energy allows the physician to resect tissues with a greater level of precision than can be achieved using conventional EMR systems that require multiple users. Moreover, the greater dexterity afforded by the single step endoscopic hood 10 allows the physician to stop retracting the actuation handle.
30 as soon as the tissue is fully resected. This further enhances the ability to perform multiple/consecutive tissue resections by allowing the resection wire 40 to avoid unnecessarily small closed-loop configurations, thereby minimizing the stress exerted upon the resection wire 40 and the distance required to return to resection wire track 18.

[0035] Referring to FIG. 5F, once the target tissue is released from the mucosal layer, the physician may advance the actuation handle 30 such that the resection wire 40 returns to the open-loop configuration disposed within the resection wire track 18. Once the vacuum suction has been removed, the physician may withdraw the endoscope from the patient. Alternatively, the physician may reposition the distal portion 12 of the endoscopic hood 10 over and in contact with the mucosal surface 34 of another target tissue, which is then resected by repeating the steps outlined above.

[0036] Although the endoscopic hood 10 and outer sleeve 22 described herein are provided as separate pieces that interlock to form an interference fit that encloses the resection wire track 18 and resection wire channel 19, the endoscopic hood 10 may also be formed as a single unitary piece of molded material using techniques known in the art. Alternatively, the endoscopic hood is formed of separate interlocking pieces that are assembled and then irreversibly joined or fused by heating, gluing, soldering welding or the like. Whether formed as separate interlocking components, or as a single unitary piece, a wide range of materials may be used to make the endoscopic hood 10 and/or sleeve 22. Suitable materials may include metals, polymers, metal-polymer composites, and the like. Some examples of suitable metals and metal alloys include stainless steel, such as 304V, 304L, and 316LV stainless steel; mild steel; nickel-titanium alloy such as linear-elastic and/or
super-elastic nitinol, other nickel alloys such as nickel-chromium-molybdenum alloys (e.g., UNS: N06625 such as INCONEL® 625, UNS: N06022 such as HASTELLOY® C-22®, UNS: N10276 such as HASTELLOY® C276®, other HASTELLOY® alloys, and the like), nickel-copper alloys (e.g., UNS: N04400 such as MONEL® 400, NICKELVAC® 400, NICORROS® 400, and the like), nickel-cobalt-chromium-molybdenum alloys (e.g., UNS: R30035 such as MP35-N® and the like), nickel-molybdenum alloys (e.g., UNS: N10665 such as HASTELLOY® ALLOY B2®), other nickel-chromium alloys, other nickel-molybdenum alloys, other nickel-cobalt alloys, other nickel-iron alloys, other nickel-copper alloys, other nickel-tungsten or tungsten alloys, and the like; cobalt-chromium alloys; cobalt-chromium-molybdenum alloys (e.g., UNS: R30003 such as ELGILOY®, PHYNOX®, and the like); platinum enriched stainless steel; titanium; combinations thereof; and the like; or any other suitable material.

[0037] Some examples of suitable polymers may include Poly(methyl methacrylate) (PMMA), polytetrafluoroethylene (PTFE), ethylene tetrafluoroethylene (ETFE), fluorinated ethylene propylene (FEP), polyoxymethylene (POM, for example, DELRIN® available from DuPont), polyether block ester, polyurethane (for example, Polyurethane 85A), polypropylene (PP), polyvinylchloride (PVC), polyether-ester (for example, ARNITEL® available from DSM Engineering Plastics), ether or ester based copolymers (for example, butylene/poly(alkylene ether) phthalate and/or other polyester elastomers such as HYTREL® available from DuPont), polyamide (for example, DURETHAN® available from Bayer or CRISTAM ID® available from Elf Atochem), elastomeric polyamides, block polyamide/ethers, polyether block amide (PEBA, for example available under the trade name PEBAX®), ethylene vinyl acetate copolymers (EVA), silicones, polyethylene (PE), Marlex high-density
polyethylene, Marlex low-density polyethylene, linear low density polyethylene (for example REXELL®), polyester, polybutylene terephthalate (PBT), polyethylene terephthalate (PET), polytrimethylene terephthalate, polyethylene naphthalate (PEN), polyetheretherketone (PEEK), polyimide (PI), polyetherimide (PEI), polyphenylene sulfide (PPS), polyphenylene oxide (PPO), poly
paraphenylene terephthalamide (for example, KEVLAR®), polysulfone, nylon, nylon-12 (such as GRILAMID® available from EMS American Grilon), perfluoro (propyl vinyl ether) (PFA), ethylene vinyl alcohol, polyolefin, polystyrene, epoxy, polyvinylidene chloride (PVdC), poly(styrene-b-isobutylene-b-styrene) (for example, SIBS and/or SIBS 50A), polycarbonates, ionomers, biocompatible polymers, polyisoprene, other suitable materials, or mixtures, combinations, copolymers thereof, polymer/metal composites, and the like. These are just examples and should not be seen as limiting.

[0038] The proximal portion 11 of the endoscopic hood 10 is dimensioned to receive the distal end of an endoscope (not shown). The endoscopic hood 10 may be designed to either permanently or temporarily attach to the distal portion of the endoscope by an attachment mechanism. Permanent attachment mechanisms may include gluing, welding, soldering or the like, while temporary locking mechanisms may include a snap-fit, screw-fit, luer-lock, press-fit using a silicone component or similar device formed into the endoscopic hood 10. In some instances, the endoscopic hood 10 may be integral to the to the endoscope shaft. In some instances, the cross-sectional shape of the proximal portion 11 may be substantially circular, though other shapes may be employed as necessary to receive the distal end of the endoscope (not shown). Inwardly projecting stops 17 are optionally included within the lumen 13 of the endoscopic hood 10 to provide a secure stopping
point against which the distal end of the endoscope may be pressed in an interference fit.

[0039] The dimensions of the endoscopic hood 10 may vary according to a variety of factors, include the desired application and size of the patient. For example, an endoscopic hood designed for rectal insertion may be considerably smaller than an endoscopic hood designed for insertion into the esophagus. The endoscopic hood 10 may be designed for multiple or single uses. As a single-use device, for example, the endoscopic hood 10 may include temporary attachment mechanism and may be stored in hermetically sealed, sterile packaging before use. A multiple-use device, however, may be designed of materials able to withstand high temperature and high pressure sterilization conditions such as those provided by an autoclave.

[0040] The present disclosure is not limited to embodiments in which the resection wire 40 is secured at a first end 41 to an anchoring element 20 located on a distal portion 12 of the endoscopic hood 10. In one embodiment, the first end 41 of the resection wire 40 is not anchored to a distal portion 12 of the endoscopic hood 10, but instead travels proximally along an inner or outer portion of the endoscopic hood and rejoins the resection wire at a more central portion (not shown). In another embodiment, the first end 41 of the resection wire 40 is anchored to a proximal portion 11 of the endoscopic hood (not shown).

[0041] All of the devices and/or methods disclosed and claimed herein can be made and executed without undue experimentation in light of the present disclosure. While the devices and methods of this disclosure have been described in terms of preferred embodiments, it will be apparent to those of skill in the art that variations can be applied to the devices and/or methods and in the steps or in the sequence of
steps of the method described herein without departing from the concept, spirit and scope of the disclosure. All such similar substitutes and modifications apparent to those skilled in the art are deemed to be within the spirit, scope and concept of the disclosure as defined by the appended claims.
We claim:

1. An endoscopic hood configured to be disposed at a distal end of an endoscope, the endoscopic hood comprising:
   - a proximal portion;
   - a distal portion;
   - a lumen extending between the proximal and distal portions;
   - a wire track disposed about an inner circumference of the distal portion; and
   - a wire channel extending between the proximal and distal portions, wherein the wire channel is in communication with the wire track, and wherein the wire channel and wire track are configured to receive a wire.

2. The endoscopic hood of claim 1, wherein the wire channel extends along an inner wall of the endoscopic hood.

3. The endoscopic hood of claim 1, wherein the wire channel extends along an outer wall of the endoscopic hood.

4. The endoscopic hood of any of claims 1 through 3, wherein the wire track includes a planar surface substantially perpendicular to the inner wall of the endoscopic hood.

5. The endoscopic hood of any of claims 1 through 4, wherein the wire track includes a groove disposed within the inner wall of the endoscopic hood.
6. The endoscopic hood of any of claims 1 through 5, wherein the wire channel merges with the wire track at an angle.

7. The endoscopic hood of any of claims 1 through 6, further comprising an anchoring element disposed adjacent to a site at which the wire channel communicates with the wire track.

8. The endoscopic hood of claim 7, wherein the anchoring element defines an aperture.

9. The endoscopic hood of any of claims 7 and 8, wherein the anchoring element is configured to secure an end of the wire.

10. The endoscopic hood of any of claims 8 and 9, wherein the aperture is configured to slidably receive a portion of the wire.

11. The endoscopic hood of any of claims 1 through 10, further comprising an elastomeric sleeve configured to form an interference fit with an outer surface of the endoscopic hood.

12. The endoscopic hood of claim 11, wherein the elastomeric sleeve includes a distal cap that is coextensive with and substantially parallel to the wire track.

13. The endoscopic hood of any of claims 11 and 12, wherein the elastomeric sleeve encloses the wire channel.
14. The endoscopic hood of any of claims 1 through 13, wherein the proximal portion of the endoscopic hood is attached to an endoscope.

15. The endoscopic hood of claim 14, wherein the endoscope includes at least one working channel in communication with the lumen of the endoscopic hood.
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

INV. A61B1/Q0 A61B17/32 A61B17/00

ALLOC.

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)

A61B

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

Electronic database consulted during the international search (name of database and, where practicable, search terms used)

EPO-Internal

C. DOCUMENTS CONSIDERED TO BE RELEVANT

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<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<td>US 2013/158546 AI (TOOMEY CIARAN [IE] ET AL) 20 June 2013 (2013-06-20) paragraphs [0037], [0038], [0046]; figures 5A-C, 6-8</td>
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<td>JP 2010 022697 A (OLYMPUS MEDICAL SYSTEMS CORP) 4 February 2010 (2010-02-04) the whole document</td>
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<td>US 5 897 487 A (OUCHI TERUO [JP]) 27 April 1999 (1999-04-27) figure 3</td>
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Date of the actual completion of the international search: 22 July 2016
Date of mailing of the international search report: 03/08/2016

Name and mailing address of the ISA/European Patent Office, P.B. 5818 Patentlaan 2 NL - 2380 HV Ridderkerk Tel: (31-70) 340-2040, Fax: (31-70) 340-3016

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