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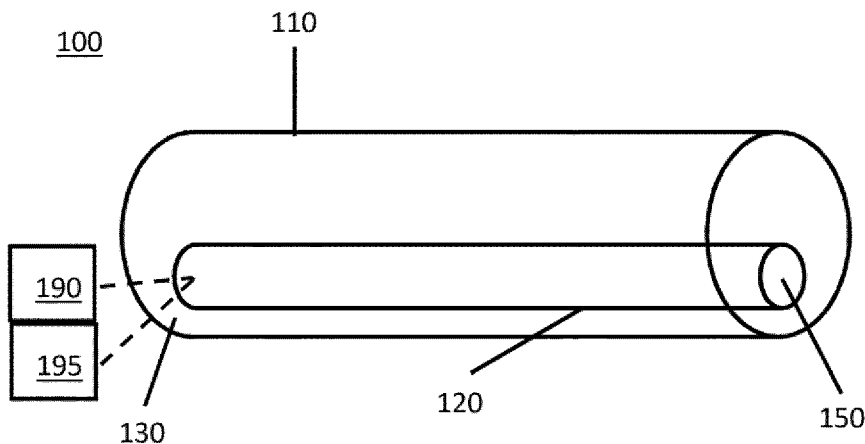


FIG 1A

(57) Abstract: The present invention relates to dispensing instruments and methods for introducing treatment material and fluid-like material with an embryo into a uterus. More specifically, the present invention relates to apparatuses for delivering a fertilized egg or embryo into a maternal uterine endometrium in humans or any other mammalian species and associated computer programs for controlling said apparatuses.



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System for embryo transfer

Field

5 The present invention relates to dispensing instruments and methods for introducing treatment material and fluid-like material with an embryo into a uterus. More specifically, the present invention relates to apparatuses for delivering a fertilized egg into a uterus and associated computer programs for controlling said apparatuses.

10 Background

Human In Vitro Fertilization (IVF) and Embryo Transfer (ET), first successfully performed in 1978, has become a widely practiced procedure to treat infertile couples who have failed with more conventional methods of therapy such as superovulation and
15 intrauterine insemination. The most common indications for IVF and related procedures, such as Gamete In Vitro Fertilization or Gamete Intra-Fallopian Transfer (GIFT) which includes women having blocked or damaged fallopian tubes and includes low sperm and/or egg quality. Related factors include age of the female, and the degree of endometrial receptivity. The procedure may also be used in cases of severe male
20 factor where direct (intracytoplasmic) injection of sperm is an option.

The IVF/ET procedure typically involves the hormonal stimulation of the female to first suppress her ability to ovulate on her own, then stimulate development of follicles in the ovaries with a fertility medication. The mature eggs are removed from the ovary
25 transvaginally using a needle, preferably guided under ultrasound. Following harvesting of the eggs, the eggs are identified and sorted with regard to maturity, and then placed with a sperm sample from the male. Approximately 24 hours after fertilization, the eggs are examined to confirm fertilization, which occurs in approximately 65% to 85% of the eggs harvested.

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After a short development period, the embryos are transferred, along with a volume of fluid, to the uterus using a delivery catheter. The delivery catheter is usually made of a

soft plastic material to avoid damage to the endometrium. There are many potential difficulties in achieving a successful implantation. Because of the soft nature of the standard delivery catheter, in a number of cases, the tip of the catheter may bend back on itself or curve away from the fundus of the uterus. The tip may also accidentally pass
5 between the layer of the endometrium and myometrium. Conversely, a stiffer catheter increases the risk of trauma to the uterus or cervix, with the latter possibly leading to the release of prostaglandins and expulsion of the eggs from the endometrium.

One particular difficulty in achieving successful implantation is the difficulty the
10 surgeon has in visualizing the uterus and the endometrium into which the embryo is implanted.

Summary

15 According to a first aspect of the invention, there is provided an apparatus suitable for delivering a fertilized egg or embryo into a maternal uterine endometrium, comprising a body configured to fit within a lumen of the female reproductive system, the body comprising: a lumen extending from a proximal end of the body to a distal portion of the body and having a distal opening at the distal portion of the body, the lumen
20 configured to slidably receive an inner body having a distal end suitable for penetrating the endometrial epithelium; the apparatus further comprising: a first actuator operable to advance the inner body out from the distal opening of the body and a second actuator operable to expel a fertilized egg from the inner body.

25 In a particular embodiment, the apparatus further comprises a measurement assembly configured to measure whether the apparatus is in a first state indicating that a distance between the distal end of the inner body and the endometrial epithelium is greater than a predetermined distance, or a second state indicating that a distance between the distal end of the inner body and the endometrial epithelium is equal to or less than a
30 predetermined distance; and
an indicating device coupled to the measurement assembly and configured to indicate that the apparatus is in the first state or the second state.

In a particular embodiment, the apparatus further comprises a measurement assembly comprising a measurement portion disposed at the distal portion of the body, the measurement assembly being configured to measure whether the apparatus is in a first state indicating that a distance between the distal portion of the body and the endometrial epithelium is greater than a predetermined distance, or a second state indicating that a distance between the distal portion of the body and the endometrial epithelium is equal to or less than a predetermined distance; and an indicating device coupled to the measurement assembly and configured to indicate that the measurement assembly is in the first state or the second state; wherein the first actuator is operable to advance the inner body out from the distal opening by at least the predetermined distance.

According to a second aspect of the invention, there is provided a computer program product configured to control the apparatus of the first aspect, wherein one or more of the actuators are electronically controlled, the computer program product comprising instructions which, when the program is executed by a computer, cause the computer to carry out the following steps: receive an instruction to actuate one or more electronically controlled actuators; and actuate the one or more electronically controlled actuators.

Brief description of the drawings

To enable better understanding of the present disclosure, and to show how the same may be carried into effect, reference will now be made, by way of example only, to the accompanying schematic drawings, in which:

Fig. 1A shows a perspective view of an apparatus suitable for delivering a fertilized egg or embryo into a maternal uterine endometrium according to one or more embodiments shown and described herein;

Fig. 1B shows a perspective view of another apparatus suitable for delivering a fertilized egg or embryo into a maternal uterine endometrium according to one or more embodiments shown and described herein;

- 5 Fig. 1C shows a perspective view of another apparatus suitable for delivering a fertilized egg or embryo into a maternal uterine endometrium according to one or more embodiments shown and described herein;

10 Fig. 1D shows a perspective view of another apparatus suitable for delivering a fertilized egg or embryo into a maternal uterine endometrium according to one or more embodiments shown and described herein;

15 Fig. 1E shows a perspective view of another apparatus suitable for delivering a fertilized egg or embryo into a maternal uterine endometrium according to one or more embodiments shown and described herein;

20 Fig. 1F shows a perspective view of another apparatus suitable for delivering a fertilized egg or embryo into a maternal uterine endometrium according to one or more embodiments shown and described herein;

Fig. 2A shows a perspective view of measuring portion according to one or more embodiments shown and described herein;

25 Figs. 2B and 2C show views of the imaging device of the measurement assembly of Fig. 2A according to one or more embodiments shown and described herein;

Fig. 3 shows a perspective view of an imaging assembly comprising an imaging device for use as part of a measurement assembly in one or more of the embodiments shown and described herein;

30 Fig. 4 shows a front view of a measuring portion according to one or more embodiments shown and described herein;

Fig. 5A shows a cross-sectional side view of a measuring portion according to one or more embodiments shown and described herein;

- 5 Figs. 5B and 5C show views of the imaging device of the measurement assembly of Fig. 5A according to one or more embodiments shown and described herein;

Fig. 6 shows a front view of a measuring portion according to one or more embodiments shown and described herein;

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Fig. 7A shows a perspective view of a measuring portion according to one or more embodiments shown and described herein;

- Fig. 7B shows a cross-sectional side view of the measuring portion shown in Fig. 7A
15 taken along the line A-A;

Figs. 7C and 7D show views of the imaging device of the measurement assembly of Fig. 7A according to one or more embodiments shown and described herein;

- 20 Fig. 8A shows a perspective view of a measuring portion according to one or more embodiments shown and described herein;

Figs. 8B and 8C show cross-sectional side view of the measuring portion shown in Fig. 8A taken along the line A-A in different configurations;

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Fig. 9A shows a side view of a steering mechanism according to some embodiments shown and described herein;

- Fig. 9B shows a side view of another steering mechanism according to some
30 embodiments shown and described herein;

Fig. 9C shows a top view of the steering mechanisms shown in Figs. 9A or 9B;

Fig. 9D shows a side view of a steering mechanism with a cable in a retracted position according to one or more embodiments shown and described herein;

5 Figs. 9E shows a perspective view of a third actuator according to some embodiments shown and described herein;

Fig. 9F shows a perspective view of the third actuator shown in Fig. 9E with the outer shell depicted as transparent;

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Fig. 9G shows a perspective view of an inner element of the steering mechanism according to some embodiments shown and described herein;

15 Fig. 9H shows a perspective view of a steering mechanism according to some embodiments shown and described herein, with the outer shell illustrated as partially transparent;

Figs. 10A to 10D show perspective views of an apparatus according to one or more embodiments shown and described herein, in various configurations;

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Fig. 11 shows a diagram of an indicating device according to one or more embodiments shown and described herein;

25 Fig. 12 shows a diagram of a computer that may be used to control the apparatus according to one or more embodiments shown and described herein;

Fig. 13A shows a cross-sectional perspective view of a connector according to one or more embodiments shown and described herein;

30 Fig. 13B shows a close-up view of a part of the connector shown in Fig. 13A;

Fig. 13C shows a perspective view of a proximal end of a connector according to one or more embodiments shown and described herein;

5 Fig. 13D shows a perspective view of a distal end of a motion controller according to one or more embodiments shown and described herein;

Fig. 13E shows an exploded view of the connector shown in Fig. 13A;

10 Fig. 13F shows a perspective view of a connection between a body connector and a distal end of a motion controller according to one or more embodiments shown and described herein;

15 Fig. 13G shows a perspective view of a proximal portion of a motion controller according to one or more embodiments shown and described herein;

Fig. 13H shows a perspective view of a motion controller according to one or more embodiments shown and described herein;

20 Fig. 14A shows a cross-sectional side view of a balloon catheter according to one or more embodiments shown and described herein;

Figs. 14B and 14C show close-up cross-sectional side views of the balloon catheter of Fig. 14A;

25 Fig. 14D shows a cross-sectional side view of another balloon catheter according to one or more embodiments shown and described herein;

30 Fig. 14E shows a cross-sectional side view of the balloon catheter of Fig. 14D in a non-everted state;

Fig. 14F shows a cross-sectional side view of the balloon catheter of Fig. 14D in an everted state;

Fig. 14G shows a perspective view of another balloon catheter according to one or more embodiments shown and described herein;

5 Fig. 14H shows a cross-sectional side view of the balloon catheter shown in Fig. 14G;

Fig. 14I shows another cross-sectional side view of the balloon catheter shown in Fig. 14G;

10 Fig. 14J shows a perspective view of another balloon catheter according to one or more embodiments shown and described herein;

Fig. 14K shows a perspective view of the distal end of the balloon catheter shown in Fig. 14G in an everted state;

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Fig. 14L shows a close up cross-sectional side view of the distal tip of the balloon catheter shown in Fig. 14K; and

20 Fig. 15 shows a flow chart of an exemplary method for using a apparatus according to one or more embodiments shown and described herein.

Detailed description

25 Fig. 1A shows a perspective view of an apparatus 100 suitable for delivering a fertilized egg or embryo into a maternal uterine endometrium, wherein the apparatus 100 comprises a body 110 (e.g. a catheter) configured to fit within a lumen of the female reproductive system. The body comprises a lumen 120 extending from a proximal end 130 of the body 110 to a distal portion 140 of the body 110 and having a distal opening 150 at the distal portion 140 of the body. The lumen 120 is configured to slidably
30 receive an inner body 160 having a distal end suitable for penetrating the endometrial epithelium. The apparatus 100 also comprises a first actuator 190 operable to advance

the inner body 160 out from the distal opening 150 of the body 110, and a second actuator 195 operable to expel a fertilized egg or embryo from the inner body.

5 The dimensions of the body 110 are exaggerated in Fig. 1A – 1F for the purposes of clarity, whereas in reality the body 110 is a narrow, elongate body configured to extend into the uterus to the endometrial epithelium.

10 Fig. 1B shows an embodiment of the apparatus 100 of Fig. 1A, wherein the apparatus 100 further includes an inner body 160 having a distal end suitable for penetrating the endometrial epithelium. In this embodiment the inner body 160 is shown located in the lumen 120 of the body 110. A plunger 165 is housed within the inner body 160 and is configured to advance towards the distal end of the inner body 160 when actuated by the second actuator 195 and expel a fertilized egg which is located inside the inner body 160.

15 Fig. 1C shows an embodiment of the apparatus 100 of Fig. 1A, wherein the apparatus further comprises a reference mark 163, present at the body 110, preferably at a distal portion of the body 110. Such reference mark 163 establishes a predetermined position of the inner body 160 within the lumen of the body 110, particularly with respect to the distal end 150 of the body 110.

20 As shown in Fig. 1C, the predetermined position of the distal end of the inner body 160 is located with a predefined distance offset relative to the distal end 150 of the body 110.

25 Fig. 1D shows a perspective view of another apparatus 100 suitable for delivering a fertilized egg or embryo into a maternal uterine endometrium, comprising a body 110 (e.g. a catheter) configured to fit within a lumen of the female reproductive system. The body comprises a lumen 120 extending from a proximal end 130 of the body 110 to a distal portion 140 of the body 110 and having a distal opening 150 at the distal portion 140 of the body. The lumen 120 is configured to slidably receive an inner body 160 having a distal end suitable for penetrating the endometrial epithelium. The apparatus

100 further comprises a measurement assembly comprising a measurement portion 170 disposed at the distal portion 140 of the body 110 (i.e. proximal to the distal opening 150). The measurement assembly 170 is configured to measure whether the apparatus is in a first state indicating the distance between the distal opening 150 of the body 110 and the endometrial epithelium is greater than a predetermined distance (i.e. in the direction that the inner body is advanced from the distal opening 150), or a second state indicating that the distance between the distal opening 150 of the body 110 and the endometrial epithelium is equal to or less than a predetermined distance. The apparatus 100 further comprises an indicating device 180 coupled to the measurement assembly 170 and configured to indicate that the measurement assembly 170 is in the first state or the second state. The measurement assembly may be removable from the body 110. In particular, the measurement portion 170 may be slidably received in a lumen of the body such that it may be extracted from the body 110 through the lumen, meaning the measurement assembly may be reusable. The apparatus 100 also comprises a first actuator 190 operable to advance the inner body 160 out from the distal opening 150 of the body 110 by at least the predetermined distance, and a second actuator 195 operable to expel a fertilized egg from the inner body (for example to advance a plunger 165).

The predetermined distance may be, for example, less than 6mm. The inner body may be slidable to extend from the distal opening by any suitable distance, determined by the predetermined distance to the endometrial epithelium and the desired implantation depth of the embryo. For example, the inner body 160 may be slidable to extend from 0 to 6mm from the distal opening 150 of the body 110.

The body 110 may be an elongate body made of any suitable flexible and biocompatible material, such as PTFE, FEP, PEEK or other flexible lubricious material. Elongate body 110 and lumens may be formed, for example, by extrusion using known methods. The outer diameter of the body 110 may be between 1mm and 1.6mm, more preferably between 1.2mm and 1.4mm, and even more preferably 1.3mm. The total length of the body 110 may be any suitable length for insertion to a position proximal to the endometrial epithelium. For example, the body 110 may have a length of 400mm to 500mm, for example 450mm.

The inner body 160 may be made of any suitable biocompatible material such as PEEK and PI (Polyimide). Such materials enable the walls of the inner body 160 to be made relatively thin whilst maintaining the required rigidity of the inner body 160 for penetrating the endometrial epithelium. The inner body 160 may have any suitable outer diameter which slidably fits within the lumen 120 of the body 110. For example, the lumen 120 may have an inner diameter of greater than 0.45mm and the outer diameter of the inner body may be about 0.45mm.

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10 The plunger 165 may also be made of materials such as PTFE, FEP, PEEK or other flexible lubricious material. The plunger 165 may have a suitable diameter which closely matches the inner diameter of the inner body 160 to effectively aspirate or expel fluid from the inner body 160. For example, the inner diameter of the inner body and the diameter of the plunger 165 may be about 0.28mm.

15 The measurement assembly may be any suitable measurement assembly that enables it to be determined whether the distance between the distal opening 150 of the body 110 and the endometrial epithelium is greater or less than the predetermined distance. It is noted that the measurement assembly may provide a qualitative measurement (for example the measurement assembly may transition between two states) or a quantitative measurement (for example actually measuring a parameter which indicates a value of the distance to the endometrial epithelium and comparing the parameter to a threshold value indicating the predetermined distance). Both possibilities are discussed in the various embodiments disclosed herein. The measuring portion 170 is the part of the measurement assembly which is configured to interact with the endometrial epithelium

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25 in order for the measurement assembly to make the determination and may comprise any suitable element or elements. The elements of the measuring portion 170 may be connected to components at the proximal end of the apparatus 100, such as components of the measurement assembly (not shown) and the indicating device 180 via connections extending through one or more lumens 175 in the body 110. For example,

30 the measuring portion may comprise components optically, acoustically, or electrically connected to proximal components of the measurement assembly such as light or sound emitters or receivers, electrical signal components or electrical power sources.

The indicating device 180 is coupled to the measurement assembly insofar as the difference between the two states of the measurement assembly can be determined (i.e. functionally coupled). The indicating device 180 may be any suitable device which can
5 indicate the difference between the two states. For example, if the measurement assembly provides a qualitative visual difference between the two states, then the indicating device 180 may be a display (e.g. scope display or digital display) configured to display the visual difference to a user of the apparatus, or it may comprise suitable software and/or hardware to receive data (such as image data) from the measurement
10 assembly and perform analysis of the data to determine the state of the measurement assembly (or display the image data). The result of the analysis may be displayed. Alternatively, the measurement assembly may comprise the software/hardware and provide the result of the analysis to the indicating device 180.

15 The indication provided by the indicating device 180 may be provided to a user of the apparatus 100 if the first actuator 190 is to be actuated manually by the user (for example a visual indication on a display or an audio indication from a speaker), or it may be provided as a command signal to an electronic controller (such as the motion controllers disclosed herein) if the first actuator 190 is to be actuated automatically by
20 the electronic controller. The indication provides a means for the operator or controller to verify that the distal opening is close enough to the endometrial epithelium to deliver the fertilized egg through the inner body.

The first and second actuators 190, 195 may be any suitable actuator controllable to
25 advance the inner body 160 and to expel the fertilized egg. For example, the first actuator 190 may be a pusher or other manual tool for advancing the inner body, or it may comprise a motor system configured to advance the inner body. Likewise, the second actuator 195 may be configured to slide a plunger movable in the inner body to pressurize fluid in the inner body distally and expel the egg. The second actuator may
30 comprise a lead screw for advancing the plunger, i.e. the second actuator may be actuated by applying a torque, and the resulting rotary motion converted into linear

motion of the plunger. The rotary actuation of the lead screw allows for a high controllability of the linear motion, and thus the aspirated and dispensed liquid volumes.

5 Figs. 10A to 10D show perspective views of an apparatus wherein the second actuator is a plunger 165. When the indicating device 180 indicates that the distal opening 150 of the body 110 is at or within the predetermined distance to the endometrial epithelium, the inner body 160 may be advanced by the first actuator to penetrate the endometrial epithelium, as shown in Fig. 10B (the endometrial epithelium is not shown). Then, the second actuator 195 may advance the plunger 165 to the distal end of the inner body 10 160, such that the embryo is expelled, as shown in Fig. 10C. Finally, the plunger 165 and inner body 160 may be retracted back into the body 110 as shown in Fig. 10D (or the body 110 may be removed from the uterus without retracting the plunger 165 and/or inner body 160). In some embodiments, the first and second actuators may be configured such that the inner body 160 is retracted as the plunger 165 is advanced. The 15 plunger 165 may be extendable, relative to the inner body 160, to at least the distal end of the inner body 160, which ensures that the embryo is expelled from the inner body 160 (i.e. so that the embryo is prevented from remaining in the inner body 160).

Whilst Fig. 1D shows the measurement portion 170 and distal opening 150 of the body 20 110 at the distal tip of the body 110, in other embodiments the measurement portion 170 and distal opening 150 may be located at a side of the distal portion 140 of body 110, as shown in Fig. 1E. The lumen 120 comprises a gradually curved section proximal to the distal opening 150 of the body 110 in order to guide the inner body 160 laterally.

25 Fig. 1D shows the measurement assembly comprising a measurement portion 170 disposed at the distal portion 140 of the body 110. In other embodiments the measurement portion 170 may be located at a different location (e.g. at a distal portion of the inner body 160) or the measurement assembly may be configured to interact with an element of the apparatus 100 in order to provide a qualitative or a quantitative 30 measurement and may comprise any suitable element or elements, without the need of having a measurement portion within the measurement assembly. This is shown in the embodiment of Fig. 1F, wherein the apparatus 100 comprises an inner body 160 having

a distal portion 161 comprising a distal end suitable for penetrating the endometrial epithelium. The apparatus 100 further comprises a body 110 (e.g. a catheter) configured to fit within a lumen of the female reproductive system. The body comprises a lumen 120 extending from a proximal end 130 of the body 110 to a distal portion 140 of the body 110 and having a distal opening 150 at the distal portion 140 of the body. The lumen 120 is configured to slidably receive the inner body 160. The apparatus 100 of Fig. 1F further comprises a measurement assembly 162. The measurement assembly 162 is configured to measure whether the apparatus 100 is in a first state indicating the distance between the distal end of the inner body 160 and the endometrial epithelium is greater than a predetermined distance (i.e. in the direction that the inner body is advanced from the distal opening 150 of the body 110), or a second state indicating that the distance between the distal end of the inner body 160 and the endometrial epithelium is equal to or less than a predetermined distance. The apparatus 100 of Fig. 1F further comprises an indicating device 180 coupled to the measurement assembly 162 and configured to indicate that the apparatus 100 is in the first state or the second state. The apparatus 100 also comprises a first actuator 190 operable to advance the inner body 160 out from the distal opening 150 of the body 110, and a second actuator 195 operable to expel a fertilized egg from the inner body (for example to advance a plunger 165).

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In an embodiment, the distal portion 161 of the inner body 160 is electrically conductive, and the measurement assembly 162 is configured to make electrical contact with such distal portion 161 of the inner body 160. The measurement assembly thus provides an indication of the capacitance. In an embodiment, the first state is indicated by a capacitance measurement below a threshold value, and the second state is indicated by a capacitance measurement above the threshold value.

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In a particular embodiment, the capacitance varies according to the variation of the distance between the distal portion 161 of the inner body 160 and the endometrial epithelium, i.e. the capacitance value changes depending on the proximity of the distal portion 161 of the inner body 160 with the endometrial epithelium. In an embodiment, the predetermined distance is zero (thus meaning contact between the distal portion 161

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of the inner body 160 and the endometrial epithelium). In an embodiment, the capacitance value is measured by means of the measurement assembly 162, which comprises an electrical system which is electrically connected to the distal portion 161 of the inner body 160 for measuring the capacitance.

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The use of the measurement assembly in combination with the indicating device assists in determining that the distal portion of the apparatus is within a predetermined distance such that the inner body can be advanced into the endometrial epithelium to deliver the fertilized egg. In embodiments wherein the indication provides information regarding the distance between the distal portion of body 110 and the endometrial epithelium, this may assist in avoiding direct contact between the distal portion of body 110 and the endometrial epithelium. Further, prematurely extending the inner body 160 into the uterus whilst the apparatus is still being positioned could cause damage to the inner body 160. Accordingly, the indicating device assists in identifying when it is appropriate to extend the inner body 160 out from the apparatus.

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In embodiments wherein the indication provides information regarding the distance between the distal end of the inner body 160 and the endometrial epithelium, the use of the measurement assembly in combination with the indicating device assists in determining that the distal end of the inner body is within a predetermined distance such that the inner body can be further advanced into the endometrial epithelium to deliver the fertilized egg and/or that the inner body is in a position wherein the fertilized egg can be delivered. Accordingly, the indicating device assists in identifying when it is appropriate to deliver the fertilized egg.

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In an embodiment, the apparatus 100 of figures 1A – 1F further comprises an imaging device, in particular a camera, not shown in the figures. In an embodiment, the camera is attached to the body 110 and moves together, as an integral device, with said body 110. In a particular embodiment, the camera is housed within the body 110.

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Fig. 2A shows a perspective view of measuring portion according to one or more embodiments. The measuring portion comprises one or more light sources 205 proximal

the distal opening 150 of the body 110. The one or more light sources 205 are configured to emit light from the distal portion of the body. The measuring portion further comprises one or more converging lenses 210 configured to focus the one or more light sources at a focal point away from the distal opening 150 of the body 110, and an imaging device 220 having a field of view including the focal point.

The one or more light sources 205 may be any suitable light source. For example, the light sources 205 may comprise one or more optical fibers, each extending through a respective lumen extending from a proximal end of the body (e.g. extending generally parallel to lumen 120) and connected to one or more light emitters at the proximal end of the body.

The one or more converging lenses 210 may be made of any suitable optically clear biocompatible material, such as a biocompatible resin, acrylics or polycarbonates. The lenses 210 may be attached to the body 110 by any biocompatible clear adhesives, such as biocompatible UV clear adhesives and or biocompatible cyanoacrylate adhesive. In the present embodiment, a single lens 210 is used, with a hole formed through the lens 210 such that the imaging device 220 is able to directly view the focal point. However, in other embodiments, the one or more lenses 210 may merely be located away from the imaging device 220. The focal point of the one or more lenses is selected to correspond to the predetermined distance.

The imaging device 220 may be any suitable imaging device. For example, the imaging device may comprise a camera such as a CMOS image sensor. When the imaging device 220 is a camera, the indicating device 180 may be a display configured to display the field of view to a user, or may comprise suitable hardware and/or software configured to receive image data from an imaging device and perform analysis of the image data to determine the state of the measurement assembly. Alternatively, the measurement assembly may comprise the software/hardware and provide the result of the analysis to the indicating device 180. The imaging device may instead comprise a fiberoptic viewable by the user.

Whilst the apparatus 200 comprises three light sources 205a, 205b and 205c, in other embodiments the apparatus may comprise one, two or more than three light sources.

The apparatus 200 may also comprise one or more secondary light sources 230
5 configured to emit unfocussed light onto the endometrial epithelium from the distal portion of the body (i.e. the light does not pass through the lens 210). The colour of the secondary light source may be selected so as to allow the focussed light to be distinguishable.

10 Fig. 2B illustrates a view 240 of the imaging device 220 as the body 110 is advanced towards a surface, wherein a distance of the distal opening 150 of the body 110 is greater than the predetermined distance. The surface is further away than the focal point and so the light sources appear as separated spots on the surface. As the body 110 is advanced, the spots converge, indicated by the arrows. Fig. 2C illustrates the view 240
15 when the body is advanced such that the light converges. If the body is further advanced closer than the focal point, the spots begin to diverge once again. The view of the imaging device 220 therefore serves as the indicating device 180 in indicating to a user that the distal opening 150 of the body 110 is at the predetermined distance. Alternatively, imaging data may be analysed by the indicating device 180 against
20 predetermine criteria to determine if the focal point has been reached. For example, known object tracking algorithms can be implemented to track the location of the three spots of light, to determine when the spots are close enough to be considered as converged.

25 Where two light sources 205, a similar converging behaviour occurs in the view 240 such that the focal point can be observed. The provision of three or more light sources 205 provide easier identification of the focal point in the case that the surface of the endometrial epithelium is uneven, or when the plane containing the three light sources is not parallel to the surface of the endometrial epithelium.

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In an embodiment where only one light source 205 is used, the focal point can still be determined from the view of the imaging device 220. Specifically, the view of the

imaging device 220 can be calibrated such that the xy position of the spot from the light source at the focal point is known. This can be used to provide a comparison point on an eyepiece of a fiberscope or on a digital display, or can be used as criteria in an image analysis algorithm.

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Fig. 3 shows a perspective view of an imaging assembly 300 comprising an imaging device 310 that may be used as part of a measurement assembly in one or more of the embodiments disclosed herein. The imaging device 310 comprises a CMOS chip located inside housing 310a and lens 310b at a distal end of the imaging assembly 300 for receiving focussing light from the endometrial epithelium onto the CMOS chip. The
10 imaging assembly 300 further comprises a plurality of wires or cables 312 for electrically connecting the CMOS chip to a proximal end of the apparatus to provide the necessary power and signal communications (e.g. to a power source and image processor at a proximal end of the apparatus) to operate the imaging assembly 300. For
15 example, the imaging assembly 300 may comprise a plurality (e.g. four) coaxial cables 312, each cable comprising an outer conductor and an inner coaxial conductor inside the outer conductor. The coaxial wiring is preferred to reduce noise in the received signal and therefore to achieve a clearer image. Each cable 312 may individually pass through a separate lumen extending through the body 110 from the distal portion 140 to the
20 proximal end 130. Alternatively, one or more (or all) of the cables 312 may pass through a single insulating shroud 320 that extends through the body 110 from the distal portion 140 to the proximal end 130. The insulating shroud 320 may be any electrically insulating material, for example PVC. The proximal portions of the cables are electrically connected to the various components at the proximal end of the apparatus to
25 provide suitable power and signal connections to the CMOS chip.

Fig. 4 shows a front view of a measuring portion according to one or more embodiments. The measurement assembly of Fig. 4 comprises a distance measuring probe at a proximal end of the apparatus, and configured to emit and receive a signal at
30 the distal portion 140 of the apparatus. The probe could be any distancing probe configured to calculate distance from any suitable signal, for example any suitable optical, acoustic, electrical or other signal. The probe emitter/receiver terminate at

openings 410a and 410b proximal the distal opening 150. In some embodiments, the signal may be emitted and received from a single opening 410 rather than a pair of openings. The probe may be, for example, an optical coherence tomography (OCT) probe including an OCT fiber bundle terminating proximal to the distal opening 150.

5 The OCT scanning probe is configured to calculate the distance of the distal opening to a portion of the endometrial epithelium. For example, the OCT probe may comprise a light signal emitter at the proximal end of the apparatus. The emitter may emit a light signal that is split into a reference path having a known length and a second path that is emitted to (and reflected from) the endometrial epithelium. Light reflected from the
10 reference path and the second path is recombined and detected at the proximal end of the apparatus (i.e. by the indicating device 180). The resulting interference pattern may be processed to determine density changes in the second path which can be interpreted as surfaces. The distance to each of the density changes can therefore be determined, including the distance to the surface of the endometrial epithelium.

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The apparatus 400 may optionally also comprise a fiberscope 420. The indicating device 180 is configured to compare the measured distance to the predetermined distance and to indicate whether the measured distance corresponds to the opening being farther or close than a predetermined distance to the endometrial epithelium.

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In one embodiment, the measuring portion 170 may comprise any suitable acoustic distance-measuring device. The acoustic device may emit acoustic signals to the endometrial epithelium, from the distal portion 140 of the apparatus 100 in response to electrical control signals transmitted through the body 110 from the proximal end of the
25 apparatus. The reflected acoustic signals may be converted into an electrical signal by the acoustic device and transmitted to the indicating device 180 for processing, from which the distance to the endometrial epithelium can be calculated.

In another embodiment, the measurement assembly may comprise a camera proximal
30 the distal opening 150 of the body 110 and configured to view a portion of the endometrial epithelium, wherein the depth of field of the camera has a predetermined value defined by nearest and the farthest objects that are in acceptably sharp focus, such

that when the image is in focus, the distal opening is at the predetermined distance to the endometrial epithelium. The indicating device 180 may comprise a display for displaying an image captured by the camera. Alternatively, the indicating device 180 may comprises suitable software and/or hardware for processing the image data of the camera to determine whether the image is in focus (for example by measuring the sharpness of the image using known image analysis methods). Alternatively, the measurement assembly may comprise the software/hardware and provide the result of the analysis to the indicating device 180.

10 In another embodiment, the measurement assembly may comprise a confocal probe .

In yet another embodiment, the measurement assembly may comprise a laser probe comprising an optical fiber terminating at opening 410 proximal to the distal opening 150 of the body 110. The laser probe is configured to measure the time of flight of laser radiation transmitted to the endometrial epithelium and reflected back to the laser probe through the optical fiber and calculate the distance to the endometrial epithelium from the time of flight value. The indicating device 180 is configured to compare the distance measured by the laser probe (taking into account the distance travelled by the radiation along the optical fiber) to the predetermined distance and to indicate whether the measured distance corresponds to the opening being farther or at/closer than a predetermined distance to the endometrial epithelium.

In another embodiment, the measurement assembly may comprise an ultrasound probe configured to transmit and receive sound waves from one or more openings 410 towards the endometrial epithelium, the measurement assembly configured to calculate the distance of the distal opening to the endometrial epithelium. The indicating device 180 is configured to compare the distance measured by the ultrasound probe (taking into account the distance travelled by the sound waves along the apparatus) to the predetermined distance and to indicate whether the measured distance corresponds to the opening being farther or at/closer than the predetermined distance to the endometrial epithelium.

Fig. 5A shows a cross-sectional side view of a measuring portion according to another embodiment. The measuring portion comprises a light source 510 having a fixed aperture configured to illuminate a shape on the endometrial epithelium (for example a circular region), and an imaging device 530 having a field of view 540 including the illuminated region. As discussed previously, the light source 510 may be any suitable light source and may be connected to a light emitter at a proximal end of the apparatus via an optical fiber extending through a lumen 520 in the body 110. A narrower aperture of the light source may be achieved by providing an opaque cylindrical body 515 which extends distally from the light source 510. The opaque body may be, for example, hypodermic tubing fixed to the light source 510 through which the light may be emitted.

When the shape appearing in the view of the imaging device 530 is bigger than a predetermined size, this indicates the distance between the distal opening and the endometrial epithelium is greater than a predetermined distance. When the viewed shape matches or is smaller than the predetermined size, this indicates the distance between the distal opening and the endometrial epithelium is equal to or less than a predetermined distance. The view 540 of the imaging device may be displayed on a display comprising a comparison overlay 550, as displayed in Figs. 5B and 5C. In Fig. 5B, the distal opening is further from the endometrial epithelium than the predetermined distance. In Fig. 5B, the distal opening is at the predetermined distance to the endometrial epithelium. Alternatively, imaging data may be analysed by the indicating device 180 against predetermined criteria to determine the size of the illuminated shape in the view of the imaging device. For example, known object tracking algorithms can be implemented to track the size of the shape, to determine when the shape is small enough to be considered as corresponding to the predetermined distance being reached.

Fig. 6 shows a front view of a measuring portion according to one or more embodiments. The measuring portion comprises a stereo camera 410a, 410b, and a light source 420. The stereo camera is configured to image the endometrial epithelium. The measurement assembly and/or indicating device may comprise software/hardware configured to perform a stereo algorithm to determine one or more depths of points in the stereo image. The distance to the endometrial epithelium is then calculated from the

one or more depths. In particular, the cameras are positioned at a known angle to each other but pointed at the same target. The software combines and analyses the image data to perform depth measurements of the surface of the endometrial epithelium.

5 Fig. 7A shows a perspective view of a measuring portion according to one or more embodiments. Fig. 7B shows a cross-sectional side view taken along the line A-A of Fig. 7A. The measuring portion comprises a light source 720 configured to illuminate the endometrial epithelium, and a rigid protrusion 730 positioned to cast a shadow onto the endometrial epithelium. The measuring portion further comprises an imaging device
10 710 having a field of view 760 which includes the shadow. The protrusion 730 is positioned relative to the light source 720 and the imaging device 710 such that when an edge of the viewed shadow crosses a predetermined location in the field of view, the measurement assembly transitions from the first state to the second state. The protrusion may be made of any suitable material which remains rigid as the apparatus is advanced
15 through the uterus, for example stainless steel wire or biocompatible plastic such as peek or FEP or PTFE or PI. It may be attached to the distal portion of the body 110 by any suitable means, such as any biocompatible adhesive (for example a UV curable clear or cyanoacrylate adhesive).

20 The light source 720 may be any suitable light source. For example the light source may be optically connected to a light emitter at a proximal end of the apparatus via an optical fiber extending through a lumen 750 of the body 110. Likewise, the imaging device may be any suitable device such as a fiberscope connected to a proximal end via lumen 740 of body 110, or it may comprise a camera electrically connected to a power source
25 and signal receiver at a proximal end of the apparatus through a lumen 740 extending through body 110.

The view may be displayed on a display comprising a comparison overlay 770, as shown in Figs. 7C and 7D, which shows exemplary views of the shadow cast on the
30 endometrial epithelium as the apparatus is advanced. In Fig. 7C, the distal opening is further away from the endometrial epithelium than the predetermined distance. In Fig. 7D, the apparatus has been advanced closer to the endometrial epithelium and the edge

of the shadow crosses the overlay 770 and the measurement assembly is within the predetermined distance. Alternatively, imaging data may be analysed by the indicating device 180 against predetermine criteria to determine the size of the illuminated shape in the view of the imaging device. For example, known object tracking algorithms can be implemented to track the position of the shadow in the field of view (for example using known edge detection algorithms), to determine when the position of the shadow indicates that the predetermined distance has been reached.

Fig. 8A shows a perspective view of a measuring portion according to one or more embodiments. Fig. 8B shows a cross-sectional side view taken along the line A-A of Fig. 8A. The measuring portion comprises a light source 820 configured to illuminate a resiliently deformable protrusion 830 configured to extend towards the endometrial epithelium. The measuring portion further comprises an imaging device 810 having a field of view of the protrusion. The protrusion being undeformed (i.e. in its relaxed state) indicates that the measurement assembly is in the first state and the protrusion being deformed indicates the second state. The indicating device may comprise a display coupled to the imaging device such that the protrusion 830 is viewable by an operator. Alternatively, imaging data may be analysed by the indicating device 180 against predetermine criteria to determine whether the protrusion has deformed. For example, known object tracking algorithms can be implemented to track the shape of the protrusion (for example using known edge detection algorithms), to determine when the deformation of the protrusion indicates that the predetermined distance has been reached.

Fig. 8B shows a configuration wherein the distal opening 150 is farther from the endometrial epithelium than the predetermined distance. Fig. 8C shows a configuration wherein the distal opening 150 is closer to the endometrial epithelium than the predetermined distance, wherein the protrusion is deformed due to abutting the endometrial epithelium.

The light source 820 may be any suitable light source. For example, the light source may be optically connected to a light emitter at a proximal end of the apparatus via an

optical fiber extending through a lumen 850 of the body 110. Likewise, the imaging device may be any suitable device such as a fiberscope connected to a proximal end via lumen 840 of body 110, or it may comprise a camera electrically connected to a power source and signal receiver at a proximal end of the apparatus through a lumen 840
5 extending through body 110.

The protrusion 830 may be made of any suitable material which is deformable as the apparatus is advanced towards the endometrial epithelium as the protrusion 830 abuts the endometrial epithelium, without penetrating the endometrial epithelium. For
10 example, it may be made of a stainless steel wire or biocompatible plastic such as PEEK, FEP, PTFE or PI. It may be attached to the distal portion of the body 110 by any suitable means, such as biocompatible adhesive (e.g. a UV curable clear or cyanoacrylate adhesive). The protrusion 830 may be configured to extend in the direction in which the apparatus is intended to be advance towards endometrial
15 epithelium, and the length of the protrusion 830 may correspond to the predetermined distance.

In yet another embodiment, the measurement portion may comprise a pressure sensor configured to measure the pressure exerted on the distal portion of the body 110 by the
20 endometrial epithelium when the distal portion 140 abuts it. When the pressure signal is above a predetermined threshold, the distal portion is considered to be at the endometrial epithelium. For example, it may be acceptable to exert a low amount of pressure on the endometrial epithelium without causing damage to the endometrial epithelium.. Any suitable pressure sensor may be used, such as an optical pressure
25 sensor configured to measure the optical change in a light signal caused by deformation of a sensor chip at the distal end of the apparatus.

The pressure sensor may have appropriate electrical connections to a proximal end of the apparatus through one or more lumens 175 extending through the body 110. For
30 example, the pressure sensor may be electrically connected to the indicating device 180, and the indicating device may comprise suitable hardware and/or software to analyse whether the electrical signal of the pressure sensor indicates that the endometrial

epithelium has been reached. The indicating device 180 may then indicate to the user or to a controller that the predetermined distance (of zero) has been reached. The indicating device 180 may optionally also be configured to analyse whether the electrical signal of the pressure sensor indicates that the pressure exerted on the endometrial epithelium is higher than a second threshold, indicating that damage may be caused to the endometrial epithelium. The indicating device 180 may then indicate to the user or the controller that the acceptable level of pressure has been exceeded.

In yet another embodiment, the measurement portion may comprise a capacitive sensor configured to measure the capacitance, preferably as the distance between the distal portion of the body and the endometrial epithelium changes. The capacitive sensor is located on the part of the distal portion of the body which is configured to impact the endometrial epithelium first as the body is advanced and the measurement assembly is configured to provide an indication of the capacitance. When the capacitance is above a predetermined threshold, the distal portion of the body is considered to be in contact with the endometrial epithelium. Instead of a capacitance sensor, an electrically conductive element may be used, the electrically conductive element being located at least on the part of the distal portion of the body which is configured to impact the endometrial epithelium first as the body is advanced. The measurement assembly may comprise an electrical system which is electrically connected to the electrically conductive element for measuring the capacitance. The electrically conductive element may be an electrically conductive portion of the body or may be an element attached to the distal portion of the body for capacitance measurement purposes. In an embodiment, the electrically conductive element is an electrically conductive wire that extends along at least a portion of the body, preferably along the whole length of the body.

In yet another embodiment, the measurement assembly may comprise several of the aforementioned embodiments, thus combining different types of measurements based on different measurement techniques.

In yet another embodiment, the measurement assembly comprises a measurement portion disposed at the distal portion of the body, the measurement assembly being

configured to measure whether the apparatus is in a first state indicating that a distance between the distal portion of the body and the endometrial epithelium is greater than a predetermined distance, or a second state indicating that a distance between the distal portion of the body and the endometrial epithelium is equal to or less than a predetermined distance; wherein the measurement assembly is further configured to measure whether the apparatus is in a third state indicating that a distance between the distal end of the inner body and the endometrial epithelium is greater than a predetermined distance, or a fourth state indicating that a distance between the distal end of the inner body and the endometrial epithelium is equal to or less than a predetermined distance; and wherein the indicating device is further configured to indicate that the apparatus is in the third state or the fourth state. In an embodiment, the measurement assembly is configured to make electrical contact with a distal portion of the inner body which is electrically conductive, and to provide an indication of the capacitance, wherein the third state is indicated by a capacitance measurement below a threshold value and the fourth state is indicated by a capacitance measurement above the threshold value; wherein preferably the predetermined distance is zero. This embodiment provides information of both the positioning of the distal portion of the body and the positioning of the distal end of the inner body, thus allowing more precise control of the implantation process.

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In some embodiments, the apparatus 100 may further comprise a steering mechanism configured to steer a distal portion of the body 110 (i.e. a portion of the body which is configured to extend into the uterus), and a third actuator for operating the steering mechanism which may be manually actuated by a user or actuated by one or more motors. Any mechanism suitable for steering the body 110 may be used. The body 110 may be slidable relative to the steering mechanism and third actuator, such that it may be advanced towards the endometrial epithelium without having to advance the steering mechanism.

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Fig. 9A shows a side view of a steering mechanism configured to steer a distal portion of the body 110 according to some embodiments. The steering mechanism comprises a plurality of connected rings 910 forming a tube wall configured to at least partially

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surround a portion of the body 110. Specifically, the rings 910 are sized to receive the body 110 inside the tube wall, and may slidably receive the body 110. The mechanism further comprises at least one cable 920 extending longitudinally along the tube wall and attached to a distal end 930 of the tube wall. For example, the at least one cable

5 may extend longitudinal along the inner or outer side of the tube wall or may extend longitudinally through the tube wall. The mechanism may comprise a plurality of said cables located at different circumferential positions about the tube wall. The connected rings 910 may be connected by the cables 920 alone. The connected rings 910 may further be configured to maintain a predetermined alignment by comprising a set of

10 interlocking features which inhibit the rings 910 from rotating about the longitudinal axis. The steering mechanism shown in Fig. 9A comprises a plurality of connected rings 910 which are pivotable about each other. In other words, the connected rings are undulated such that neighbouring rings abut at a tangential contact point. This enables the rings 910 to roll about each other to steer the body 110 when one of the cables 920

15 are retracted by the third actuator (not shown). The circumferential position of the tangential contact points about which the ring 910 roll may be varied along the length of the steering mechanism. For example, sequential circumferential contact points may be offset by 90 degrees in order to accommodate rolling in two perpendicular directions by a plurality of cables. Preferably, the steering mechanism comprises two pairs of said

20 cables, each pair comprising radially opposing cables, in order to provide steering in a plurality of directions. Such a configuration is shown in Fig. 9C, which is a top view of such a steering mechanism.

The connected rings shown in Fig. 9A may be made of any suitable material such as

25 medical grade injection moldable or 3D printed plastic resin such as ABS, PET, DELRIN or PTFE. The cables 920 may be made of, for example, stainless steel and threaded through holes formed in the walls of the connected rings forming the tube wall. The cables 920 may each extend through a pair of holes in the connected rings and be looped at the distal end 930 as shown in Figs. 9A and 9C. The third actuator may be any

30 suitable actuator for contracting the one or more cables.

Fig. 9B shows a side view of a steering mechanism according to an alternative embodiment. The steering mechanism comprises a plurality of connected rings 915 forming a tube wall configured to at least partially surround a portion of the body 110. Specifically, the rings 915 are sized to receive the body 110 inside the tube wall, and may slidably receive the body 110. The mechanism further comprises at least one cable 920 extending longitudinally through the tube wall and attached to a distal end 930 of the tube wall. The plurality of connected rings 910 are rings of a compressible tube or spring. When one of the cables is retracted by the third actuator (not shown), the tube or spring is compressed on the side of the retracted cable such that the steering mechanism is steered towards the contracted side. As the tube is compressible, the tension force is partly counteracted such that the resulting curve of the mechanism shown in 9B may be less than for the embodiment shown in Fig. 9A for a given tensioning force. Once again, preferably, the steering mechanism comprises two pairs of said cables, each pair comprising radially opposing cables, in order to provide steering in a plurality of directions, as shown in Fig. 9C.

The connected rings shown in Fig. 9B may be a compressible tube or spring made of any suitable material such as FEP, PTFE. The cables may be made of any suitable material such as stainless steel and threaded through holes formed in the walls of the connected rings forming the tube wall. The cables 920 may each extend through a pair of holes in the connected rings and be looped at the distal end 930 as shown in Figs. 9A and 9C. The third actuator may be any suitable actuator for contracting the one or more cables.

Fig. 9D shows a side view of a steering mechanism when one of the cables has been retracted, such that the steering mechanism is steered in the direction illustrated by arrow 925.

Figs. 9E shows a perspective view of the third actuator according to some embodiments. Fig. 9F shows a perspective view of the third actuator shown in Fig. 9E with the outer shell 950 depicted as transparent. The third actuator comprises an inner element 940 and an outer shell 950 housing the inner element 940. Each of the at least one cables 920 are

fixed to the outer shell at a proximal portion 960, for example by welding, adhesive, or about a fixed winch. The inner element 940 is fixedly connected to the tube wall 910 either directly or indirectly via one or more intermediate components. The outer shell 950 is rotatable relative to the inner element 940. Optionally, a handle 970 may be fixed to the inner element 940 through the outer shell 950. Rotating the outer shell 950 about the inner element 940 causes one of the cables 920 to be retracted, which steers the tube wall in a certain direction as discussed above. The body may extend through the handle 970 and the inner element 940 such that it may be advanced through the steering mechanism.

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Fig. 9G shows a perspective view of the inner element according to some embodiments. The inner element 940 may comprise one or more components which are biased to abut the outer shell 950. For example, the inner element 940 may comprise two hemispheres 942a, 942b, which are biased away from each other to abut the inner surface of the outer shell 950. The biasing causes a frictional fit to be obtained between the inner element 940 and outer shell 950, such that they may only slide relative to each other when a sufficient force is applied, meaning that once a desired direction of the steering mechanism is achieved, it may be maintained in position by the third actuator. The inner element may be connected to the handle via a handle connector 946, and may be connected to the tube wall via an intermediate portion 948. The intermediate portion 948 may be incompressible and the cables 920 may be threaded through the intermediate portion and to the tube wall.

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Fig. 9H shows a perspective view of a steering mechanism according to some embodiments, with the outer shell 950 illustrated as partially transparent. The steering mechanism of Fig. 9H also comprises an inner element 940 and an outer shell 950 housing the inner element 940 and rotatable relative to inner element 940, as described above. The steering mechanism may also comprise the handle 970. Further, each of the cables 920 are fixedly connected to the outer shell via a winch 952 (i.e. the cables are wound to the winch). The winch may be wound or unwound using an adjuster 954 (for example a flat head adjuster) which is accessible externally to the outer shell 950, meaning that the default tensions for each cable 920 may be calibrated after assembly of

the steering mechanism. The rings 910 forming the tube wall may be fixedly connected to the inner element 940 via an inflexible tubular intermediate 958. The tubular intermediate 958 may be of any suitable length. For example, in embodiments where the outer catheter is used, the tubular intermediate 958 may be configured to extend through
5 any inflexible parts of the outer catheter.

The cables 920 may extend through channels 956 formed in the inner surface of outer shell 950, and thus avoid contact with the inner element 940, avoiding damage to the cables as the outer shell rotates relative to the inner shell 940. The cables 920 may
10 extend from the channels 956 and through lumens formed in the wall of the tubular intermediate 958, and to the tube wall.

The outer shell 950, inner element 940, handle 970, winches 952 and tubular intermediate 958 may be made of medical grade injection moldable or 3D printed
15 plastic resin such as ABS, PET, DELRIN or PTFE.

The steering mechanism may instead be electronically controlled. For example, the steering mechanism shown in Fig. 9H may comprise an electronic controller fixed to the inner element 940, and comprising motors for actuating movement of the outer shell
20 950 relative to the inner element 940. Alternatively, a steering mechanism may comprise a plurality of connected rings forming a tube wall configured to at least partially surround a portion of the body, and at least two opposing cables extending longitudinally through the tube wall and attached to a distal end of the tube wall, wherein the cables are connected to a motor configured to contract the cables to steer
25 the steering mechanism.

The tube wall of the steering mechanism is sized to slidably receive the body 110. The outer diameter of the tube wall may be about 1.4mm to 1.8mm or about 1.5mm to 1.7mm, for example 1.6mm. In embodiments where the steering mechanism is used, the
30 body 110 may be configured to advance beyond the end of the steering mechanism by about 20mm to 40mm, for example about 30mm.

Fig. 11 shows a diagram of an indicating device 180 which may be used when the output of the measurement assembly is a signal (for example if the measurement assembly comprises a camera or distance measuring probe). The indicating device comprises a device interface 1100 configured to communicate with (i.e. send and receive signals from) the measurement assembly. The indicating device further comprises a memory 1110, processor 1120 and indicator 1130. It will be appreciated than any suitable computing system may be used for the indicating device 180 and that the device shown in Fig. 11 is one of many devices that could be used. For example, a distributed computer system may be used, for example comprising one or more servers or client computing systems, using any known distributed computing technique. In some examples, a general-purpose computer or any other processing system may be used to implement the methods disclosed herein. Furthermore, the steps disclosed below may be implemented in software, hardware, or any combination of these to achieve the same steps.

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It will be appreciated by the skilled person that the indicating device of Fig. 11 may be powered by any suitable powering means. The memory 1110 may comprise one or more volatile or non-volatile memory devices, such as DRAM, SRAM, flash memory, read-only memory, ferroelectric RAM, hard disk drives, floppy disks, magnetic tape, optical discs, or similar. Likewise, the processor 1120 may comprise one or more processing units, such as a microprocessor, GPU, CPU, multi-core processor or similar. The device interface 1100 may comprise wired connections, such as optic, fiber-optic, ethernet or similar, or any suitable wireless communication.

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In the case of the measurement assembly making a quantitative measurement (e.g. comprising a distance sensing probe), the memory 1110 may contain suitable instructions for performing a comparison of the received signal to the predetermined distance, and the processor 1120 may be configured to perform the instructions. For example, the indicating device 180 may receive the raw data signal via device interface 1110, the processor 1120 may convert the raw data into a numerical value for the predetermined distance, and the numerical value may be compared to the predetermined distance. If the numerical value is equal to or lower than the predetermined distance,

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then the processor 1120 may instruct the indicator 1130 to indicate that the predetermined distance has been reached. For example, the indicator 1130 may provide an audio or visual signal to the operator, or may comprise an electronic controller configured to provide a command signal to an electronically controlled motion controller to perform the operation of advancing the needle and expelling the embryo (with the memory 1110 containing suitable instructions for providing control signals to the actuators). The indicating device 180 may also be configured to further indicate to the user when the automatic procedure of advancing the need and expelling the embryo has been completed (for example an audio or visual signal). Alternatively, the indicator 1130 may comprise a display displaying the measured distance alongside the predetermined distance.

In the case of the measurement assembly comprising a camera and the measurement being a qualitative measurement, the indicating device may merely be a display for displaying the real-time image captured by the camera, optionally with a comparison overlay on the display where appropriate. Alternatively, the memory 1110 may store instructions for performing image analysis of the image data to automatically determine if the image indicates that the predetermined distance has been reached. The processor 1120 may be configured to carry out the instructions. The processor 1120 may then instruct the indicator 1130 to indicate that the predetermined distance has been reached. For example, the indicator 1130 may provide an audio or visual signal to the operator, or may provide a command signal to an electronically controlled motion controller to perform the operation of advancing the needle and expelling the embryo.

Whilst Fig. 11 shows the motion controller and the indicating device as separate devices, indicating device and motion controller may be incorporated into a single device. Further, the measurement assembly may instead be connected to the indicating device indirectly via a device interface of the motion controller.

Fig. 12 shows a diagram of a computer 1200 that may be used to control one or more of the actuators disclosed herein. The computer 1200 may comprise one or more of a controller 1210, a processor 1220, a user interface 1230, a display 1240, a network

connection 1245, a device interface 1250, and a memory 1255 storing instructions for programs 1260 and a data repository 1270. It will be appreciated that any suitable computing system may be used for the computer 1200 and that the device shown in Fig. 12 is one of many devices that could be used. For example, a distributed computer system may be used, for example comprising one or more servers or client computing systems, using any known distributed computing technique. In some examples, a general-purpose computer or any other processing system may be used to implement the methods disclosed herein. Furthermore, the steps disclosed below may be implemented in software, hardware, or any combination of these to achieve the same steps.

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It will be appreciated by the skilled person that the indicating device of Fig. 12 may be powered by any suitable powering means. The memory 1255 may comprise one or more volatile or non-volatile memory devices, such as DRAM, SRAM, flash memory, read-only memory, ferroelectric RAM, hard disk drives, floppy disks, magnetic tape, optical discs, or similar. Likewise, the processor 1220 may comprise one or more processing units, such as a microprocessor, GPU, CPU, multi-core processor or similar. The device interface 1250 may comprise wired connections, such as optic, fiber-optic, ethernet or similar, or any suitable wireless communication.

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The device interface 1250 is configured to communicate with (i.e. send and receive control information) one or more electronically controlled actuators. Any of the actuators disclosed herein may be electronically controlled. The device interface 1250 may also be configured to communicate with the indicating device 180. Alternatively, the computer 1200 may comprise all of the features of the indicating device shown in Fig. 11 and may itself be considered as the indicating device 180.

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The computer 1200 provides manual control of the electronically controlled actuators by a user of the apparatus. For example, the memory 1255 may comprise instructions for a program 1260 which when executed by processor 1220 causes a graphical user interface (GUI) to be displayed on display 1240, or remotely via network connection 1245. The GUI may comprise appropriate inputs to enable the user to actuate one or more of the actuators. For example, the user may select a function from the GUI, which

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causes the controller 1210 to send a corresponding control signal to one or more actuators to perform the function.

For example, the user may be able to select a command from the GUI for advancing the
5 body by a certain distance, or at a certain constant speed. The user may be able to select
one or more commands for steering the steering mechanism. The user may be able to
select one or more commands for advancing the inner body and/or plunger. The
commands may correspond to a sequence of commands such that several commands are
10 performed in sequence. For example, the memory 1255 may store instructions for
advancing the body 110 until the predetermined distance is reached (i.e. until the
indicating device indicates that the predetermined distance is reached), and for
subsequently advancing the inner body and plunger to expel the embryo. It may be that
all actuators are actuated via such an automated process, or that only some are actuated
automatically and the rest are actuated manually (i.e. by user control via the GUI).

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The computer 1200 may be configured to receive data from the measurement assembly,
either directly or via indicating device 180, and display the data on the GUI alongside
the command options. For example, the display 1240 may display real-time image data
received from the measurement assembly. The GUI may also display an indication that
20 the predetermined distance has been reached to the user.

In some embodiments, the apparatus may further comprise a motion controller
configured to connect to a proximal portion of the body 100, wherein the controller is
configured to actuate the first actuator, second actuator, and/or a fourth actuator for
25 advancing the body relative to the motion controller. Any suitable motion controller
may be used. For example, the motion controller may comprise one or more linear or
rotary mechanisms for actuating the first actuator, the second actuator and/or the fourth
actuator. In some embodiments, one or more of the first, second and fourth actuator may
comprise a lead screw, each lead screw rotatable to advance the body 110, inner body
30 160 or plunger 165, the motion controller may be configured to actuate the one or more
lead screws. The motion controller may be manually controlled or electronically

controlled. The motion controller may be configured to connect to the body via a connector.

Fig. 13A shows a cross-sectional perspective view of a connector 1300 configured to connect to a motion controller 1350. Fig. 13E shows an exploded view of the connector 1300. The connector 1300 comprises a connector housing 1310 which is insertable into a distal end of the motion controller 1350. The connector 1300 also comprises a body connector 1314 fixedly receiving a proximal portion of the body 110 (e.g. by a frictional fit or by adhesive), an inner body connector 1324 fixedly receiving a proximal portion of the inner body 160 (e.g. by frictional fit or by adhesive) and a plunger connector 1332 fixedly receiving a proximal portion of the plunger 165 (e.g. by frictional fit or by adhesive).

The body connector 1314 is received by the connector housing 1310 and is slidable relative to the connector housing 1310 at a fixed rotational orientation. For example, the connector housing 1310 may comprise a longitudinally extending protrusion or recess and the body connector 1314 may comprise a corresponding recess or protrusion such that the body connector 1314 is only receivable in the connector housing 1310 at a single rotational orientation. The connector 1300 further comprises a body lead screw 1316 having an external screw thread 1320 which corresponds to an internal screw thread 1312 of the connector housing 1310. The body lead screw 1316 abuts the body connector 1314 such that, when turning the body lead screw 1316 to advance the body lead screw distally, the body connector 1314 also advances distally (but in a fixed orientation). In the illustrated embodiment, the body lead screw 1316 is connected to the body connector 1314 by one or more protrusions 1318 which are received by a corresponding recess 1322 in the body connector. The interaction between the protrusion 1318 and the recess 1322 means that the body lead screw 1316 abuts the body connector 1314 when it slides both distally and proximally, meaning the body lead screw 1316 is able to both advance and retract the body connector 1314 and thus the body 110. In some embodiments the protrusion 1318 may instead be located on the body connector 1314 and the recess on the body lead screw 1316.

The inner body connector 1324 is housed by the body connector 1314 and is slidable relative to the connector housing 1310 and the body connector 1314. In some embodiments, the inner body connector 1324 may be slidable relative to the connector housing 1310 and body connector 1314 at a fixed rotational orientation (for example using corresponding protrusions and recessions in the body connector 1314 and the inner body connector 1324 as previously disclosed).

Fig. 13B shows a close up of the inner body connector 1324 and the plunger connector 1332 shown in Fig. 13A. The connector further comprises an inner body lead screw 1328 having an external screw thread 1326 which corresponds to an internal screw thread 1327 of the body connector 1314. The inner body lead screw 1328 abuts the inner body connector 1324 such that, when turning the inner body lead screw 1328 to advance the inner body lead screw 1328 distally, the inner body connector 1324 also advances distally. In the illustrated embodiment, the inner body lead screw 1328 is connected to the inner body connector 1324 by one or more protrusions 1325 which are received by a corresponding recess in the inner body connector 1325. The interaction between the protrusion 1325 and the recess means that the inner body lead screw 1328 abuts the inner body connector 1324 when it slides both distally and proximally, meaning the inner body lead screw 1328 is able to both advance and retract the inner body connector 1324 and thus the inner body 160. In some embodiments the protrusion 1325 may instead be located on the inner body connector 1324 and the recess on the inner body lead screw 1328. In some embodiments, the inner body connector 1324 and inner body lead screw 1328 may be a single unitary body.

The plunger connector 1332 is housed by the inner body connector 1324 and is slidable relative to the connector housing 1310, the body connector 1314 and the inner body connector 1324. In some embodiments, the plunger connector 1332 may be slidable relative to the connector housing 1310, body connector 1314 and inner body connector 1324 at a fixed rotational orientation (for example using corresponding protrusions and recessions in the inner body connector 1324 and the plunger connector 1332 as previously disclosed). The connector further comprises a plunger lead screw 1336 having an external screw thread 1333 which corresponds to an internal screw thread

1334 of the inner body connector 1324. The plunger lead screw 1336 abuts the plunger connector 1332 such that, when turning the plunger lead screw 1336 to advance the plunger lead screw 1336 distally, the plunger connector 1332 also advances distally. In the illustrated embodiment, the plunger lead screw 1336 is connected to the plunger connector 1332 by one or more protrusions 1335 in the plunger connector 1332 which are received by a corresponding recess in the plunger lead screw 1336. The interaction between the protrusion 1335 and the recess means that the plunger lead screw 1336 abuts the plunger connector 1332 when it slides both distally and proximally, meaning the plunger lead screw 1336 is able to both advance and retract the plunger connector 1332 and thus the plunger 165. In some embodiments the protrusion 1335 may instead be located on the plunger lead screw 1336 and the recess on the plunger connector 1332. In some embodiments, the plunger connector 1332 and plunger lead screw 1336 may be a single unitary body.

15 The connector 1300 may comprises an interlocking feature 1340 at a distal end, such as a Luer lock, for securing the distal end of the connector 1300 to the next distal component of the apparatus. For example, the interlocking feature 1340 may be configured to interlock with a corresponding interlocking feature on the balloon catheter disclosed below, and as show in Fig. 14J, or it may be configured to interlocking with a proximal end of the steering mechanism disclosed above, such as handle 970.

Fig. 13C shows a perspective view of a proximal end of the connector 1300. Fig. 13D shows a perspective view of a distal end of the motion controller 1350. The proximal end of the connector 1300 is configured to connect with the distal end of the motion controller 1350. The body lead screw 1316 comprises a toothed proximal end 1360 configured to engage with a body lead screw driver 1352 of the motion controller 1350. The inner body lead screw 1328 comprises a toothed proximal end 1362 which is configured to engage with an inner body lead screw driver 1354 of the motion controller 1350. The plunger lead screw comprises a toothed proximal end (not shown) configured to engage with a plunger lead screw driver 1356 of the motion controller 1350. The proximal end of the connector housing 1310 may further comprise a locking feature such as a protrusion 1364 configured to be received by a recess of the distal end of the

motion controller (not shown) in order to prevent relative rotation of the distal end of the motion controller 1350 and connector housing 1310. The connector 1300 may be connected to the motion controller 1350 by a push-fit, click-fit, frictional-fit, or similar mechanism. The lead screw drivers 1352, 1354 and 1356 may be distally biased by
5 springs 1358 (see Fig. 13A) in order to ensure engagement with the proximal ends of the lead screws during rotation.

Whilst the illustrated embodiment illustrates a connector which actuates the body connector, inner body connector and plunger connector by the lead screw mechanisms,
10 it will be appreciated that in other embodiments the connector only actuates one or more of the body connector, inner body connector and plunger connector by a lead screw mechanism. For example, the connector may comprise only the body connector 1314, actuated by the body lead screw 1316 as described above, and the needle and plunger may be actuated differently (i.e. by a linear motion mechanism). It will also be
15 appreciated that the outermost screw thread of a given embodiment engages the inner screw thread 1312 of the connector housing 1310. For example, in embodiments where the connector 1300 actuates only the inner body 160 and plunger 165 by the screw thread mechanisms disclosed, the outer thread 1326 of the inner body lead screw 1328 engages with the inner thread 1312 of the connector housing 1310, rather than the inner
20 screw thread of the body connector 1314.

The connector housing 1310, connectors 1314, 1324 and 1332 and lead screws 1316, 1328 and 1336 may be made of any suitable material such as plastic and may be molded or 3D printed. Likewise, the lead screw drivers 1352, 1354 and 1356 may be made of
25 any suitable material such as plastic.

Electrical wiring and/or optical fibers extending from the measurement portion 170 through the body 110 from the distal end of the apparatus may terminate at the connector 1300. In particular, a proximal portion of electrical wiring extending from the
30 distal end of the apparatus through the body 110 may be electrically connected to an electrical terminal of the connector. The motion controller may comprise a corresponding electrical terminal biased towards the electrical terminal of the connector

to maintain contact with the electrical terminal of the connector. Similarly, a proximal portion of the optical fibers may be optically connected to an optical terminal of the connector 1300, and the motion controller may comprise an optical terminal configured to form a face seal with the optical terminal of the connector 1300 and biased towards
5 the optical terminal of the connector 1300 to maintain the face seal with the first optical terminal.

Fig. 13F shows a perspective view of the connection between the body connector 1314 and the distal end of the motion controller 1350. The illustrated embodiment shows a
10 proximal portion of electrical cables 312 (see e.g. Fig. 3) which extend from the measurement portion (e.g. camera 310), through the body 110 to the connector 1300. The cables 312 are electrically connected to an electrical terminal (such as a printed circuit board) on the body connector 1314 (e.g. by soldering). The electrical terminal comprises electrical connections which are configured to engage electrical pins 1374 on
15 the motion controller 1350. The electrical pins 1374 are biased distally such that the electrical connection between the pins 1374 and the electrical terminal of the body connector 1314 is maintained as the body connector 1314 is advanced through the connector housing 1300. It will be appreciated that optical connections between the connector 1300 and the motion controller 1350 can be similarly established and
20 maintained.

Fig. 13G shows a perspective view of a proximal portion of the motion controller 1350. The motion controller 1350 comprises a first motor 1381, a second motor 1383 and a third motor 1385. In one embodiment, the first motor 1381 may be configured to drive
25 the body lead screw driver 1352, the second motor 1383 may be configured to drive the inner body lead screw driver 1354 and the third motor may be configured to drive the plunger lead screw driver 1356. The motion controller 1350 may be configured to advance/retract the body 110, inner body 160 and plunger 165 at the same rate by running the first, second and third motors 1381, 1383, 1385 in unison at the rate
30 corresponding to the same advancement speed of the body 110, inner body 160 and plunger 165. The motion controller 1350 may also be configured to advance/retract the inner body 160 and plunger 165 at the same rate relative to the body 110 by running the

second and third motors 1383, 1385 in unison at the rate corresponding to the same advancement speed of the inner body 160 and plunger 165. The motion controller 1350 may also be configured to advance/retract the plunger 165 relative to the inner body 160 by running the third motor 1385 alone.

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Alternatively, the first motor 1381 may be configured to turn each of the lead screw drivers 1352, 1354 and 1356 (e.g. via a first threaded rod) to advance the lead screws 1316, 1328 and 1336 at the same rate (to advance the body 110, the inner body 160 and the plunger 165 at the same rate). The second motor 1383 may be configured to turn the inner body lead screw driver 1354 and the plunger lead screw driver 1356 (e.g. via a second threaded rod) to advance the inner body lead screw 1328 and the plunger lead screw 1336 at the same rate (to advance the inner body 160 and plunger 165 at the same rate). The third motor 1386 may be configured to turn the plunger lead screw driver 1356 (e.g. via a threaded rod) to advance the plunger 165.

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The motion controller 1350 may further comprise an axial motor (not shown) to retract the plunger lead screw driver 1352 before connection or disconnection of the connector 1300, in order to prevent damage to the plunger lead screw driver 1352 during connection or disconnection of the connector 1300.

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The connector 1300 may be assembled in the following steps (the steps need not necessarily be performed in chronologically in the order presented). The proximal portion of the plunger 165 may be fixed to the plunger connector 1332. The plunger connector 1332 may be connected to the plunger lead screw 1336 (for example the connector and lead screw are pushed together until the protrusion falls into the recess to secure the connector and lead screw together). The proximal portion of the inner body 160 may be connected to the inner body connector 1324. The inner body connector 1324 may be connected to the inner body lead screw 1328 (e.g. by pushing the connector and lead screw are together until the protrusion falls into the recess to secure the connector and lead screw together). The proximal portion of the body 110 may be connected to the body connector 1314. The body connector 1314 may be connected to the body lead screw 1316 (e.g. by pushing the connector and lead screw are together

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until the protrusion falls into the recess to secure the connector and lead screw together). The distal end of the plunger 165 may be passed through the inner body connector 1324 from the proximal end and through the inner body 160. The plunger lead screw 1336 may be screwed to the inner body connector 1324. The distal end of the inner body 160
5 may be passed through the body connector 1314 from the proximal end and through the body 110. The inner body lead screw 1328 may be screwed to the body connector 1314. The distal end of the body may be passed through the connector housing 1310 from the proximal end. The body lead screw 1316 may be screwed to the connector housing 1310. The lead screws may then be actuated to moved the body 110, inner body 160 and
10 plunger 165 to the correct relative positions.

When the connector 1300 is assembled, the plunger lead screw 1336 may be actuated to aspirate the fluid containing the embryo. The connector 1300 is then connected to the motion controller 1350.

15 The length from the proximal end of the body connector 1314 to the distal tip of the body 110 may be about 300mm to 400mm, preferably 325mm to 375mm, for example 358mm. The maximum length of the connector 1300 from the proximal end to the distal tip of the body 110 may be bout 400mm to 500mm, preferably 425mm to 475mm, for
20 example 450mm.

Fig. 13H shows a perspective view of a motion controller 1350 according to an embodiment. The motion controller 1350 is configured to connect to a proximal portion of the body 110 and to actuate on said body 110. In the embodiment shown in Fig. 13H,
25 the motion controller 1350 comprises two lead screws 1351, 1353 configured to advance/retract the body 110, and to rotate and advance/retract an outer catheter which houses the body 110 in an embodiment. In the embodiment shown the motion controller 1350 further comprises three actuation mechanisms 1357, 1359, 1361, each actuation mechanism configured to actuate on one of the lead screws 1351, 1353. The actuation
30 of the lead screws 1351, 1353 may be manual or may be motorized. Although in this embodiment two lead screws 1351, 1353 are shown, in other embodiments there may be only one lead screw, depending on the elements intended to be moved by the motion

controller 1350. The motion controller 1350 allows fine and precise control of the movement of the body 110 and of the outer catheter housing the body, as well as fixing the outer catheter in the required position. In an embodiment, the motion controller 1350 may be actuated based on a real time image obtained by an imaging device, such as a camera, present in the apparatus 100.

In some embodiments, the apparatus may further comprise an outer balloon catheter.

Fig. 14A shows a cross-sectional side view of a balloon catheter according to one or more embodiments. The balloon catheter comprises an outer tubular body 1410 which slidably receives an evertable tube 1420. The evertable tube is configured to receive the body 110 or tube wall 910 of the steering mechanism. In the illustrated embodiment, the evertable tube 1420 is attached to the body 110 or tube wall 910 to form a first seal 1430 at one end of the evertable tube 1420. The evertable tube is attached to the inner surface of the outer tubular body to form a second seal 1440 at another end of the evertable tube. At a proximal end of the balloon catheter, the body 110 or tube wall 910 enters the balloon catheter through a slidable seal 1470 formed by, for example, an O-ring, which allows the body 110 or tube wall 910 to advance through the balloon catheter whilst fluidically sealing the proximal end of the balloon catheter from the interior. The balloon catheter is configured to contain a fluid (e.g. water) in the volume formed between the evertable tube 1420 and the outer tubular body 1410 (wherein the volume is fluidically sealed by the seals 1470, 1430 and 1440). The balloon catheter may also comprise a port 1460 for filling the said volume, for example fluidically connected to a reservoir.

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Fig. 14B shows a close-up cross-sectional side view of the balloon catheter of Fig. 14A in a non-everted state. In use, the tip 1450 of the outer tubular body may be placed at the entrance of the cervix of a patient. The body 110 or tube wall 910 may then be advanced through the balloon catheter. The forward pressure of the body 110 or tube wall 910 and the fluid pressure causes the evertable tube 1420 to evert in the distal direction (i.e. through the cervix). The evertable tube 1420 acts as a roller as it everts through the cervix, and the fluid between the evertable tube 1420 and the outer tubular

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body 1410 limits shear forces, thereby facilitating insertion of the body 110 or tube wall 910 into the uterus.

Fig. 14C shows a close-up cross-sectional side view of the balloon catheter of Fig. 14A in a fully everted state. As can be seen, body 110 or tube wall 910 may be advanced through the balloon catheter to a maximal position defined by the most distal position of the seal 1430 (after which the tube 1420 is not able to evert further). The seal 1430 may be positioned on body 110 or the tube wall 910 such that it protrudes from the tube 1420 by a predetermined distance.

Fig. 14D shows a cross-sectional side view of another balloon catheter according to one or more embodiments. Unlike the embodiment shown in Figs. 14A to 14C, the evertable tube 1420 is fully rolled back on itself and is attached to the tip 1450 to form two seals 1440a and 1440b on the outer tubular body. The evertable tube 1420 is configured to receive the tube wall 910 or the body 110, for example by a frictional fit or may be adhesive. The inner volume of the evertable tube may be pre-filled with fluid or the balloon catheter may comprise a port 1460 for filling the evertable tube with fluid. As in the previous embodiment, as the tube wall 910 or the body 110 is advanced, the inner part of the evertable tube 1420 travels forward (distally, see arrow 1445) which creates forces on the leading distal edge of the tube 1420 (indicated by arrow 1446). Accordingly, the tube 1420 everts as the tube wall 910 or body 110 is advanced through the balloon catheter. Fig. 14E shows a cross-sectional side view of the balloon catheter of Fig. 14D in a non-everted state, and Fig. 14F shows a cross-sectional side view of the balloon catheter of Fig. 14D in an everted state.

Fig. 14G shows a perspective view of another balloon catheter according to one or more embodiments. The balloon catheter comprises an outer tubular body formed by proximal section 1410a, distal section 1410b and tip 1450. As in the other embodiments, the balloon catheter further comprises an evertable tube 1420 slidably received in the outer tubular body and configured to receive the tube wall 910 or the body 110. The balloon catheter further comprises a reservoir of fluid 1480, and a port 1460 (e.g a valve) for filling the reservoir 1480 with fluid.

Fig. 14H shows a cross-sectional side view of the balloon catheter shown in Fig. 14G. The reservoir 1480 comprises a piston housing 1484, a piston 1486 and a biasing mechanism (e.g. a spring) configured to bias the piston distally. The biasing mechanism 1482 causes the fluid in the reservoir to be pressurised.

Fig 14I shows another cross-sectional side view of the balloon catheter shown in Fig. 14G. The evertable tube is fully rolled back on itself and is attached to the outer tubular body to form two seals 1440a and 1440b on the outer tubular body. A port 1442 between the seals 1440a and 1440b fluidically connects the interior of the evertable tube 1420 with the fluid in the reservoir 1480 (via a fluid path extending along the interior of the proximal section 1410a and into the reservoir). Accordingly, the pressurised fluid in the reservoir 1480 serves to pressurise the evertable tube 1420 such that it does not deflate.

The balloon catheter may further comprise a releasable locking mechanism for locking the piston in a proximal position. For example, there may be provided a tab 1475 (see Fig. 14G) which is configured to prevent the piston from moving distally when located between an external flange 1488 and a proximal end of the reservoir 1480 (see Fig. 14J). When the tab is removed, the external flange 1488 (and therefore the piston 1486 may move distally to pressurise the fluid. Fig. 14J shows a perspective cross-sectional view of the balloon catheter when the tab 1475 has been removed.

As shown in Fig. 14J, the piston housing 1484 and piston 1486 may be configured to slidably receive the tube wall 910 or body 110. In particular, the piston housing 1484 and piston 1486 may comprise proximal and distal holes for slidably receiving the tube wall 910 or body 110. The tube wall 910 or body 110 may be fluidically separated from the fluid in the reservoir.

As in the previous cases, advancing the tube wall 910 or body 110 creates a forward pressure on the inner surface of the evertable tube 1420 (by friction or by the tube wall 910 or body 110 being attached to the evertable tube 1420). Fig. 14K shows a

perspective view of the distal end of the balloon catheter shown in Fig. 14G in an everted state. Fig. 14L shows a close up cross-sectional side view of the distal tip of the balloon catheter in the everted state.

- 5 The evertable tube may be made of any flexible tubing, for example any suitable thermoplastic elastomer such as polyether block amide. (and the evertable tube may be fixed to the outer tubular body and tube wall 910 or body 110 by any suitable medical adhesive such as any cyanoacrylate adhesive (e.g. a UV curable adhesive).
- 10 The outer diameter of the evertable tube in the everted state may be about 1.9mm to 2.5mm and preferably 2.1mm to 2.3mm, for example 2.2mm.

Fig. 15 shows a flow chart of an exemplary method for using the apparatus 100 according to the present disclosure. In a first step 1510, the distal end of apparatus 100
15 is positioned against the cervix (for example through a speculum).

In a next step 1515, the body 110 (and optionally the steering mechanism) is inserted into the uterus by advancing the apparatus through the cervix. This may be done by the use of any of the balloon catheters as disclosed herein, or the body 110 may be inserted
20 without the use of a balloon catheter.

In a next step 1520, optionally, the body 110 may be steered by a steering mechanism as disclosed herein.

25 Next, in step 1525, the body 110 may be advanced through the uterus. For example, the body 110 may be advanced manually by the operator, or the body 110 may be advanced by an electronic motion controller as part of an automatic process or in response to a command inputted by the user.

30 In step 1530, the distance of the distal opening 150 to the endometrial epithelium may be measured. The measurement may be a qualitative measurement determining whether the distal opening 150 is farther from or at or closer to the predetermined distance (for

example as viewed by the user via a fiberscope or camera display, or as determined from image analysis by the indicating device). The measurement may be a quantitative measurement determining a distance value and comparing the distance value to the predetermined distance.

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If the distance to the endometrial epithelium is still greater than the predetermined distance, then the body 110 is advanced further into the uterus.

10 If the predetermined distance has been reached (as indicated by the indicating portion), then the body 110 is held in place and the inner body 160 is advanced (step 1540). The inner body 160 may be advanced manually by the user via an actuator, or may be advanced using a motion controller, either as part of an automatic process or in response to a command inputted by the user.

15 In step 1545, the embryo is expelled. Again, the embryo may be expelled manually by the user via an actuator, or may be expelled using an electronic motion controller (for example by advancing a plunger), either as part of an automatic process or in response to a command inputted by the user.

20 Optionally, in step 1550, the inner body 160 may be retracted. The inner body 160 may be retracted manually by the user via an actuator, or may be retracted using an electronic motion controller, either as part of an automatic process or in response to a command inputted by the user.

25 In step 1555, the body 110 is removed from the uterus. The body 110 may be removed manually by the user, or may be removed using an electronic motion controller, either as part of an automatic process or in response to a command inputted by the user.

The following also forms part of the present disclosure.

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A steering mechanism configured to steer a distal portion of the body configured to fit within a lumen of the female reproductive system and an actuator for operating the steering mechanism. The body may be part of any apparatus disclosed herein.

- 5 The steering mechanism may comprise a plurality of connected rings forming a tube wall configured to at least partially surround a portion of the body; and at least one cable extending longitudinally along the tube wall and attached to a distal end of the tube wall; wherein the actuator is operable to contract and extend at least one cable to steer the distal portion of the body; preferably wherein the steering mechanism
10 comprises a plurality of said cables located at different circumferential positions about the tube wall.

The actuator may comprise: an inner element; an outer shell housing the inner element, each of the at least one cable being fixedly connected to the outer shell at a proximal
15 portion; the inner element being fixedly connected to the tube wall; the outer shell being rotatable relative to the inner element; and optionally, a handle fixed to the inner element through the outer shell.

The inner element may comprise one or more components biased to abut the outer shell.
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The body may be configured to be slidable relative to the steering mechanism and actuator.

A balloon catheter comprising: an outer tubular body; and an evertable tube slidably
25 received in the outer tubular body and configured to receive a steering mechanism for steering a body configured to fit within a lumen of the female reproductive system or a body configured to fit within a lumen of the female reproductive system; wherein the balloon catheter is configured to contain a fluid between the outer tubular body and the evertable tube; and when the steering mechanism or body is advanced, the evertable
30 tube everts.

The balloon catheter may comprise a fluid port for filling the balloon catheter with fluid.

The fluid port may be fluidically connected to a reservoir of fluid.

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The reservoir may be formed by a piston housing and a piston, and a biasing mechanism configured to bias the piston distally to pressurise the fluid in the reservoir.

10 The balloon catheter may further comprise a releasable locking mechanism for locking the piston in a proximal position.

The piston housing and piston may be configured to slidably receive the steering mechanism or body.

15 The maximum volume of the reservoir may be less than an overfilling volume of the balloon catheter.

The steering mechanism or body may be fixed to the tubular balloon such that the steering mechanism or body is advanceable to a maximum position.

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A connector having a connector housing and configured to connect to a motion controller, the connector housing receiving one or more of: a body connector configured to receive a proximal portion of a body a body configured to fit within a lumen of the female reproductive system, the body connector slidable relative to the connector
25 housing at a fixed orientation, and a lead screw for advancing the body connector; and/or an inner body connector configured to receive a proximal portion of an inner body slidably received in a lumen of a body configured to fit within a lumen of the female reproductive system, the inner body connector slidable relative to the connector housing, optionally at a fixed orientation, and a lead screw for advancing the inner body
30 connector; and/or a plunger connector configured to receive a proximal portion of a plunger slidably received by an inner body slidably received in a lumen of a body configured to fit within a lumen of the female reproductive system, the plunger

connector slidable relative to the connector housing, optionally at a fixed orientation, and a lead screw for advancing the plunger connector.

5 A proximal end of one or more of the lead screws for advancing the body connector, inner body connector or plunger connector may comprise teeth for rotationally locking the one or more lead screws to corresponding lead screw drivers of a motion controller.

The connector may be configured to releasably connect to the motion controller, optionally at a fixed orientation.

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The connector (e.g. body connector) may comprise an electrical terminal configured to connect to an electrical terminal of the motion controller. The connector (e.g. body connector) may comprise an optical terminal configured to form a face seal with an optical terminal of the motion controller.

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A motion controller configured to connect to the connector disclosed above and to drive one or more of the lead screws.

20 The motion controller may comprise one or more lead screw drivers, the one or more lead screw drivers comprises teeth for rotationally locking to the corresponding lead screw. The lead screw drivers may be distally biased towards the connector.

The motion controller may be configured to releasably connect to the connector, optionally at a fixed orientation.

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The motion controller may comprise an electrical terminal configured to connect to an electrical terminal of the connector, wherein the electrical terminal of the motion controller is biased towards the electrical terminal of the connector to maintain electrical contact.

30 The motion controller may comprise an optical terminal configured to form a face seal with an optical terminal of the controller, wherein the optical terminal of the motion

controller is biased towards the optical terminal of the connector to maintain the face seal.

The following clauses also form part of the present disclosure:

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1. An apparatus suitable for delivering a fertilized egg or embryo into a maternal uterine endometrium in humans or any other mammalian species, comprising a body configured to fit within a lumen of the female reproductive system, the body comprising:

10 a lumen extending from a proximal end of the body to a distal portion of the body and having a distal opening at the distal portion of the body, the lumen configured to slidably receive an inner body having a distal end suitable for penetrating the endometrial epithelium;

the apparatus further comprising:

15 a measurement assembly comprising a measurement portion disposed at the distal portion of the body, the measurement assembly configured to measure whether the apparatus is in a first state indicating that a distance between the distal portion and the endometrial epithelium is greater than a predetermined distance, or a second state indicating that a distance between the distal portion and the endometrial epithelium is
20 equal to or less than a predetermined distance;

an indicating device coupled to the measurement assembly and configured to indicate that the measurement assembly is in the first state or the second state; and

a first actuator operable to advance the inner body out from the distal opening by at least the predetermined distance, and a second actuator operable to expel a fertilized egg
25 from the inner body.

2. The apparatus of clause 1, wherein the measurement assembly comprises:

one or more light sources proximal the distal opening, each configured to emit light from the distal portion of the body; and

30 one or more converging lenses configured to focus the one or more light sources at a focal point away from the distal opening; and

an imaging device having a field of view including the focal point;

wherein when the emitted light is incident on the endometrial epithelium at the focal point, the distal opening is at the predetermined distance away from the endometrial epithelium;
optionally wherein the indicating device comprises a display of the imaging device.

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3. The apparatus of clause 2, wherein the imaging device comprises a camera, or a fiberscope.

4. The apparatus of clause 2 or 3, further comprising a secondary light source
10 configured to emit unfocussed light onto the endometrial epithelium from the distal portion of the body.

5. The apparatus of any of clauses 2 to 4, wherein the one or more light sources
15 comprise two or more separated light sources, such that the light sources converge at the focal point, wherein when the incident light on the endometrial epithelium from the light sources converge, the distal opening is within the predetermined proximity; preferably wherein the one or more light sources comprise three or more separated light sources.

20 6. The apparatus of clause 1, wherein the measurement assembly comprises an optical coherence tomography (OCT) scanning probe comprising an OCT fiber bundle terminating proximal to the distal opening, the OCT scanning probe configured to calculate the distance of the distal opening to a portion of the endometrial epithelium.

25 7. The apparatus of clause 1, wherein the measurement assembly comprises a camera proximal the distal opening and configured to view a portion of the endometrial epithelium, wherein the depth of field of the camera has a predetermined value, such that when the image is in focus, the distal opening is at the predetermined distance to the endometrial epithelium; optionally wherein the indicating device comprises a
30 display for displaying an image captured by the camera.

8. The apparatus of clause 1, wherein the measurement assembly comprises a confocal probe.

9. The apparatus of clause 1, wherein the measurement assembly comprises laser probe comprising an optical fiber terminating proximal to the distal opening, the laser probe configured to measure the time of flight of laser radiation transmitted to the endometrial epithelium and reflected back to the laser probe through the optical fiber.

10. The apparatus of clause 1, wherein the measurement portion comprises a light source having a fixed aperture configured to illuminate a shape on the endometrial epithelium, and an imaging device having a field of view including the illuminated region; wherein the viewed shape being bigger than a predetermined size indicates the first state, and the viewed shape matching or being smaller than the predetermined size indicates the second state;

15 optionally wherein the indicating device comprises a display coupled to the imaging device, the display comprising a view of the illuminated endometrial epithelium and a comparison overlay, wherein the comparison overlay is sized and shaped such that the viewed shape being bigger than the comparison overlay indicates the first state, and the viewed shape matching or being smaller than the comparison overlay indicates the second state.

11. The apparatus of clause 1, wherein the measurement portion comprises a light source and a stereo camera configured to image the endometrial epithelium, the measurement assembly configured to calculate a distance of the distal opening to the endometrial epithelium.

12. The apparatus of clause 1, wherein the measurement portion comprises a light source configured to illuminate the endometrial epithelium, a rigid protrusion positioned to cast a shadow onto the endometrial epithelium, and an imaging device having a field of view including the shadow; wherein the protrusion is positioned such that when an edge of the the viewed shadow crosses a predetermined location in the field of view, the measurement assembly transitions from the first state to the second state; optionally

wherein the indicating device is a display coupled to the imaging device and displaying a view of the imaging device, the display further comprising a comparison overlay, wherein the comparison overlay is positioned such that when an edge of the viewed shadow crosses the comparison overlay as the body is advanced, the measurement
5 assembly transitions from the first state to the second state.

13. The apparatus of clause 1, wherein the measurement assembly comprises an ultrasound probe comprising an emitter and receiver proximate the distal opening configured to emit sound waves towards the endometrial epithelium, the measurement
10 assembly configured to calculate the distance of the distal opening to the endometrial epithelium.

14. The apparatus of clause 1, wherein the measurement portion comprises a light source configured to illuminate a resiliently deformable protrusion, the protrusion
15 configured to extend towards the endometrial epithelium, and an imaging device having a field of view of the protrusion; wherein the protrusion being undeformed indicates the first state and the protrusion being deformed indicates the second state; optionally wherein the indicating device comprises a display coupled to the imaging device.

20 15. The apparatus of clause 1, wherein the measurement assembly comprises a pressure sensor located on the part of the distal portion which is configured to impact the endometrial epithelium first as the body is advanced; wherein the predetermined distance is zero and the first state is indicated by a pressure measurement below a threshold value, and the second state is indicated by a pressure measurement above the
25 threshold value.

16. The apparatus of any preceding clause, further comprising a steering mechanism configured to steer a distal portion of the body and a third actuator for operating the steering mechanism.
30

17. The apparatus of clause 16, wherein the steering mechanism comprises:

a plurality of connected rings forming a tube wall configured to at least partially surround a portion of the body; and

at least one cable extending longitudinally along the tube wall and attached to a distal end of the tube wall;

5 wherein the third actuator is operable to contract and extend at least one cable to steer the distal portion of the body;

preferably wherein the steering mechanism comprises a plurality of said cables located at different circumferential positions about the tube wall. .

10 18. The apparatus of clause 17, wherein the third actuator comprises:

an inner element;

an outer shell housing the inner element, each of the at least one cable being fixedly connected to the outer shell at a proximal portion;

the inner element being fixedly connected to the tube wall;

15 the outer shell being rotatable relative to the inner element; and

optionally, a handle fixed to the inner element through the outer shell.

19. The apparatus of clause 18, wherein the inner element comprises one or more components biased to abut the outer shell.

20

20. The apparatus of any of clauses 16 to 19, wherein the body is slidable relative to the steering mechanism and third actuator.

21. The apparatus of any preceding clause, further comprising a balloon catheter, the
25 balloon catheter comprising:

an outer tubular body; and

an evertable tube slidably received in the outer tubular body and configured to receive the steering mechanism or body; wherein

the balloon catheter is configured to contain a fluid between the outer tubular body and
30 the evertable tube; and

when the steering mechanism or body is advanced, the evertable tube everts.

22. The apparatus of clause 21, wherein the balloon catheter comprises a fluid port for filling the balloon catheter with fluid.
- 5 23. The apparatus of clause 22, wherein the fluid port is fluidically connected to a reservoir of fluid.
24. The apparatus of clause 23, wherein the reservoir is formed by a piston housing and a piston, and a biasing mechanism configured to bias the piston distally to
10 pressurise the fluid in the reservoir.
25. The apparatus of clause 24, further comprising a releasable locking mechanism for locking the piston in a proximal position.
- 15 26. The apparatus of clause 23 or 24, wherein the piston housing and piston are configured to slidably receive the steering mechanism or body.
27. The apparatus of any of clauses 23 to 26, wherein the maximum volume of the reservoir is less than an overfilling volume of the balloon catheter.
20
28. The apparatus of any of clauses 21 to 27, wherein the steering mechanism or body is fixed to the tubular balloon such that the steering mechanism or body is advanceable to a maximum position.
- 25 29. The apparatus of any preceding clause, further comprising a plunger slidably received by the inner body, wherein the second actuator is operable to expel a fertilized egg from the inner body by sliding the plunger distally relative to the inner body; optionally wherein the second actuator comprises a lead screw for advancing the plunger.
30
30. The apparatus of clause 29, wherein the first actuator is configured to retract the inner body as the plunger is slid distally relative to the inner body.

31. The apparatus of any preceding clause, further comprising a motion controller configured to connect to a proximal portion of the body, the controller configured to actuate one or more of the first actuator, second actuator, and/or a fourth actuator for
5 advancing the body relative to the motion controller.

32. The apparatus of clause 31, wherein one or more of the first, second and fourth actuators comprises a lead screw, and the motion controller is configured to actuate one or more of the lead screws.
10

33. The apparatus of clause 32, further comprising a connector having a connector housing and configured to connect to the motion controller, the connector housing receiving one or more of:

a body connector fixedly receiving a proximal portion of the
15 body, the body connector slidable relative to the connector housing at a fixed orientation, wherein the fourth actuator comprises a lead screw for advancing the body connector; and/or

an inner body connector fixedly receiving a proximal portion of the inner body, the inner body connector slidable relative to the connector housing,
20 optionally at a fixed orientation, wherein the first actuator comprises a lead screw for advancing the inner body connector; and/or

a plunger connector fixedly receiving a proximal portion of a plunger slidably received by the inner body, the plunger connector slidable relative to the connector housing, optionally at a fixed orientation, wherein the second actuator
25 comprises a lead screw for advancing the plunger connector.

34. The apparatus of clause 33 wherein a proximal end of one or more of the lead screws for advancing the body connector, inner body connector or plunger connector comprises teeth for rotationally locking the one or more lead screws to corresponding
30 lead screw drivers of the motion controller, optionally wherein the lead screw drivers are distally biased towards the connector.

35. The apparatus of clause 33 or 34, wherein the connector is configured to releasably connect to the motion controller, optionally at a fixed orientation.

5 36. The apparatus of any of clauses 32 to 35 wherein the measurement portion is electrically connected to a first electrical terminal of the connector, and the motion controller comprises a corresponding second electrical terminal biased towards the first electrical terminal to maintain contact with the first electrical terminal; and/or wherein the measurement portion is optically connected to a first optical terminal of the connector, and the motion controller comprises a corresponding second optical terminal
10 configured to form a face seal with the first optical terminal and biased towards the first optical terminal to maintain the face seal with the first optical terminal.

37. The apparatus of any preceding clause, wherein the measurement assembly comprises an electronic output indicating the measurement of the measurement portion,
15 and the indicating device is configured to analyse the electronic output to determine whether the measurement assembly is in the first state or the second state.

38. The apparatus of clause 37, wherein the indication is configured to be provided to an electronic controller configured to control one or more actuators.
20

39. The apparatus of any preceding clause, wherein one or more of the actuators are electronically controlled.

40. A computer program product configured to control the apparatus of clause 39,
25 comprising instructions which, when the program is executed by a computer, cause the computer to carry out the following steps:
receive an instruction to actuate one or more electronically controlled actuators; and
actuate the one or more electronically controlled actuators.

30

All of the above are fully within the scope of the present disclosure, and are considered to form the basis for alternative embodiments in which one or more combinations of the

above described features are applied, without limitation to the specific combination disclosed above.

5 In light of this, there will be many alternatives which implement the teaching of the present disclosure. It is expected that one skilled in the art will be able to modify and adapt the above disclosure to suit its own circumstances and requirements within the scope of the present disclosure, while retaining some or all technical effects of the same, either disclosed or derivable from the above, in light of his common general knowledge in this art. All such equivalents, modifications or adaptations fall within the scope of the
10 present disclosure.

CLAIMS

1. An apparatus suitable for delivering a fertilized egg or embryo into a maternal uterine endometrium in humans or any other mammalian species, comprising a body
5 configured to fit within a lumen of the female reproductive system, the body comprising:
a lumen extending from a proximal end of the body to a distal portion of the body and having a distal opening at the distal portion of the body, the lumen configured to slidably receive an inner body having a distal end suitable for penetrating the
10 endometrial epithelium;
the apparatus further comprising:
a first actuator operable to advance the inner body out from the distal opening of the body and a second actuator operable to expel a fertilized egg from the inner body.
- 15 2. The apparatus of claim 1, further comprising:
a measurement assembly configured to measure whether the apparatus is in a first state indicating that a distance between the distal end of the inner body and the endometrial epithelium is greater than a predetermined distance, or a second state indicating that a distance between the distal end of the inner body and the endometrial epithelium is
20 equal to or less than a predetermined distance; and
an indicating device coupled to the measurement assembly and configured to indicate that the apparatus is in the first state or the second state.
3. The apparatus of claim 2, wherein the measurement assembly is configured to
25 make electrical contact with a distal portion of the inner body which is electrically conductive, and to provide an indication of the capacitance, wherein the first state is indicated by a capacitance measurement below a threshold value and the second state is indicated by a capacitance measurement above the threshold value.
- 30 4. The apparatus of claim 3, wherein the predetermined distance is zero.
5. The apparatus of claim 1, further comprising:

a measurement assembly comprising a measurement portion disposed at the distal portion of the body, the measurement assembly being configured to measure whether the apparatus is in a first state indicating that a distance between the distal portion of the body and the endometrial epithelium is greater than a predetermined distance, or a
5 second state indicating that a distance between the distal portion of the body and the endometrial epithelium is equal to or less than a predetermined distance; and
an indicating device coupled to the measurement assembly and configured to indicate that the measurement assembly is in the first state or the second state;
wherein the first actuator is operable to advance the inner body out from the distal
10 opening by at least the predetermined distance.

6. The apparatus of claim 5, wherein the measurement assembly comprises:
one or more light sources proximal the distal opening, each configured to emit light
from the distal portion of the body; and
15 one or more converging lenses configured to focus the one or more light sources at a focal point away from the distal opening; and
an imaging device having a field of view including the focal point;
wherein when the emitted light is incident on the endometrial epithelium at the focal point, the distal opening is at the predetermined distance away from the endometrial
20 epithelium;
optionally wherein the indicating device comprises a display of the imaging device;
optionally wherein the imaging device comprises a camera, or a fiberscope.

7. The apparatus of claim 6, further comprising a secondary light source configured
25 to emit unfocussed light onto the endometrial epithelium from the distal portion of the body.

8. The apparatus of claim 6 or 7, wherein the one or more light sources comprise
two or more separated light sources, such that the light sources converge at the focal
30 point, wherein when the incident light on the endometrial epithelium from the light sources converge, the distal opening is within the predetermined proximity; preferably
wherein the one or more light sources comprise three or more separated light sources.

9. The apparatus of claim 5, wherein the measurement assembly comprises:
- an optical coherence tomography (OCT) scanning probe comprising an OCT fiber bundle terminating proximal to the distal opening, the OCT scanning probe
5 configured to calculate the distance of the distal opening to a portion of the endometrial epithelium, or
 - laser probe comprising an optical fiber terminating proximal to the distal opening, the laser probe configured to measure the time of flight of laser radiation transmitted to the endometrial epithelium and reflected back to the
10 laser probe through the optical fiber, or
 - a pressure sensor located on the part of the distal portion of the body which is configured to impact the endometrial epithelium first as the body is advanced; wherein the predetermined distance is zero and the first state is indicated by a pressure measurement below a threshold value, and the second state is indicated
15 by a pressure measurement above the threshold value, or
 - an ultrasound probe comprising an emitter and receiver proximate the distal opening configured to emit sound waves towards the endometrial epithelium, the measurement assembly configured to calculate the distance of the distal opening to the endometrial epithelium,
20
 - or any combination thereof.

10. The apparatus of claim 5, wherein the measurement assembly comprises a capacitive sensor located on the part of the distal portion of the body which is configured to impact the endometrial epithelium first as the body is advanced; wherein
25 the measurement assembly is configured to provide an indication of the capacitance, preferably as the distance between the distal portion of the body and the endometrial epithelium changes.

11. The apparatus of claim 5, wherein the measurement assembly comprises an
30 electrically conductive element located on the part of the distal portion of the body which is configured to impact the endometrial epithelium first as the body is advanced; wherein the measurement assembly is configured to provide an indication of the

capacitance, preferably as the distance between the distal portion of the body and the endometrial epithelium changes.

12. The apparatus of claim 10 or 11, wherein the predetermined distance is zero and the first state is indicated by a capacitance measurement below a threshold value, and the second state is indicated by a capacitance measurement above the threshold value.

13. The apparatus of any of claims 5 to 12, wherein the measurement assembly is further configured to measure whether the apparatus is in a third state indicating that a distance between the distal end of the inner body and the endometrial epithelium is greater than a predetermined distance, or a fourth state indicating that a distance between the distal end of the inner body and the endometrial epithelium is equal to or less than a predetermined distance; and wherein the indicating device is further configured to indicate that the apparatus is in the third state or the fourth state.

14. The apparatus of claim 13, wherein the measurement assembly is configured to make electrical contact with a distal portion of the inner body which is electrically conductive, and to provide an indication of the capacitance, wherein the third state is indicated by a capacitance measurement below a threshold value and the fourth state is indicated by a capacitance measurement above the threshold value; wherein preferably the predetermined distance is zero.

15. The apparatus of any preceding claim, wherein the body comprises a reference mark which establishes a predetermined position of the inner body within the lumen of the body, wherein in the predetermined position the distal end of the inner body is located with a predefined distance offset relative to the distal end of the body.

16. The apparatus of any preceding claim, further comprising an inner body having a distal end suitable for penetrating the endometrial epithelium.

17. The apparatus of any preceding claim, further comprising an imaging device having a field of view including a focal point away from the distal opening.

18. The apparatus of any preceding claim, further comprising a steering mechanism configured to steer a distal portion of the body and a third actuator for operating the steering mechanism.

5

19. The apparatus of claim 18, wherein the steering mechanism comprises:
a plurality of connected rings forming a tube wall configured to at least partially surround a portion of the body; and
at least one cable extending longitudinally along the tube wall and attached to a distal
10 end of the tube wall;
wherein the third actuator is operable to contract and extend at least one cable to steer the distal portion of the body;
preferably wherein the steering mechanism comprises a plurality of said cables located at different circumferential positions about the tube wall.

15

20. The apparatus of claim 19, wherein the third actuator comprises:
an inner element;
an outer shell housing the inner element, each of the at least one cable being fixedly connected to the outer shell at a proximal portion;
20 the inner element being fixedly connected to the tube wall;
the outer shell being rotatable relative to the inner element; and
optionally, a handle fixed to the inner element through the outer shell.

25

21. The apparatus of claim 20, wherein the inner element comprises one or more components biased to abut the outer shell.

22. The apparatus of any of claims 18 to 21, wherein the body is slidable relative to the steering mechanism and third actuator.

30

23. The apparatus of any preceding claim, further comprising a balloon catheter, the balloon catheter comprising:
an outer tubular body; and

an evertable tube slidably received in the outer tubular body and configured to receive the steering mechanism or body; wherein the balloon catheter is configured to contain a fluid between the outer tubular body and the evertable tube; and

5 when the steering mechanism or body is advanced, the evertable tube everts.

24. The apparatus of claim 23, wherein the balloon catheter comprises a fluid port for filling the balloon catheter with fluid.

10 25. The apparatus of claim 24, wherein the fluid port is fluidically connected to a reservoir of fluid; optionally wherein the maximum volume of the reservoir is less than an overfilling volume of the balloon catheter.

15 26. The apparatus of claim 25, wherein the reservoir is formed by a piston housing and a piston, and a biasing mechanism configured to bias the piston distally to pressurise the fluid in the reservoir.

27. The apparatus of claim 26, further comprising a releasable locking mechanism for locking the piston in a proximal position.

20

28. The apparatus of claim 26 or 27, wherein the piston housing and piston are configured to slidably receive the steering mechanism or body.

25 29. The apparatus of any of claims 23 to 28, wherein the steering mechanism or body is fixed to the tubular balloon such that the steering mechanism or body is advanceable to a maximum position.

30 30. The apparatus of any preceding claim, further comprising a plunger slidably received by the inner body, wherein the second actuator is operable to expel a fertilized egg from the inner body by sliding the plunger in a distal direction relative to the inner body; optionally wherein the second actuator comprises a lead screw for advancing the plunger.

31. The apparatus of claim 30, wherein the first actuator is configured to retract the inner body as the plunger is slid distally relative to the inner body.

5 32. The apparatus of any preceding claim, further comprising a motion controller configured to connect to a proximal portion of the body, the motion controller comprising one or more lead screws configured to advance/retract the body and/or to advance/retract and/or rotate an outer catheter housing the body.

10 33. The apparatus of claim 32, wherein the motion controller comprises one or more actuation mechanisms configured to actuate one or more of the lead screws.

34. The apparatus of any preceding claim, further comprising a motion controller configured to connect to a proximal portion of the body, the controller configured to
15 actuate one or more of the first actuator, second actuator, and/or a fourth actuator for advancing the body relative to the motion controller.

35. The apparatus of claim 34, wherein one or more of the first, second and fourth
20 actuators comprises a lead screw, and the motion controller is configured to actuate one or more of the lead screws.

36. The apparatus of claim 35, further comprising a connector having a connector housing and configured to connect to the motion controller, the connector housing receiving one or more of:

25 a body connector fixedly receiving a proximal portion of the body, the body connector slidable relative to the connector housing at a fixed orientation, wherein the fourth actuator comprises a lead screw for advancing the body connector; and/or

an inner body connector fixedly receiving a proximal portion of the inner body, the inner body connector slidable relative to the connector housing, optionally at a fixed
30 orientation, wherein the first actuator comprises a lead screw for advancing the inner body connector; and/or

a plunger connector fixedly receiving a proximal portion of a plunger slidably received by the inner body, the plunger connector slidable relative to the connector housing, optionally at a fixed orientation, wherein the second actuator comprises a lead screw for advancing the plunger connector.

5

37. The apparatus of claim 36 wherein a proximal end of one or more of the lead screws for advancing the body connector, inner body connector or plunger connector comprises teeth for rotationally locking the one or more lead screws to corresponding lead screw drivers of the motion controller, optionally wherein the lead screw drivers are distally biased towards the connector.

10

38. The apparatus of claim 36 or 37, wherein the connector is configured to releasably connect to the motion controller, optionally at a fixed orientation.

15

39. The apparatus of any of claims 35 to 38 wherein the measurement portion is electrically connected to a first electrical terminal of the connector, and the motion controller comprises a corresponding second electrical terminal biased towards the first electrical terminal to maintain contact with the first electrical terminal; and/or wherein the measurement portion is optically connected to a first optical terminal of the connector, and the motion controller comprises a corresponding second optical terminal configured to form a face seal with the first optical terminal and biased towards the first optical terminal to maintain the face seal with the first optical terminal.

20

40. The apparatus of any preceding claim, wherein the measurement assembly comprises an electronic output indicating the measurement of the measurement portion, and the indicating device is configured to analyse the electronic output to determine whether the measurement assembly is in the first state or the second state.

25

41. The apparatus of claim 40, wherein the indication is configured to be provided to an electronic controller configured to control one or more actuators.

30

42. The apparatus of any preceding claim, wherein one or more of the actuators are electronically controlled.

43. A computer program product configured to control the apparatus of claim 42,
5 comprising instructions which, when the program is executed by a computer, cause the computer to carry out the following steps:
receive an instruction to actuate one or more electronically controlled actuators; and
actuate the one or more electronically controlled actuators.

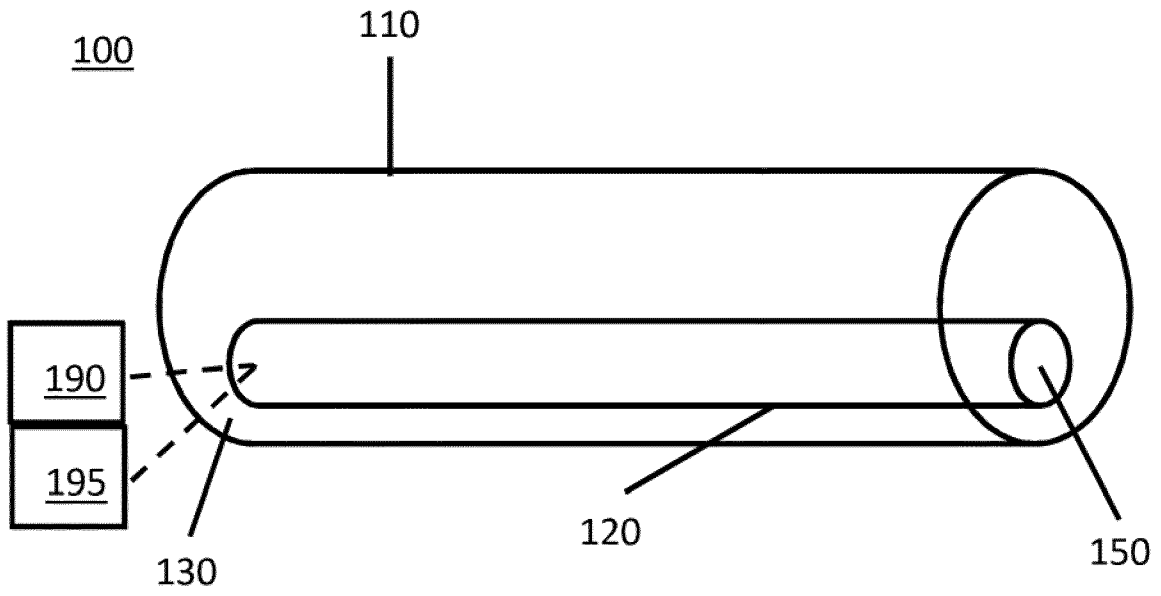


FIG 1A

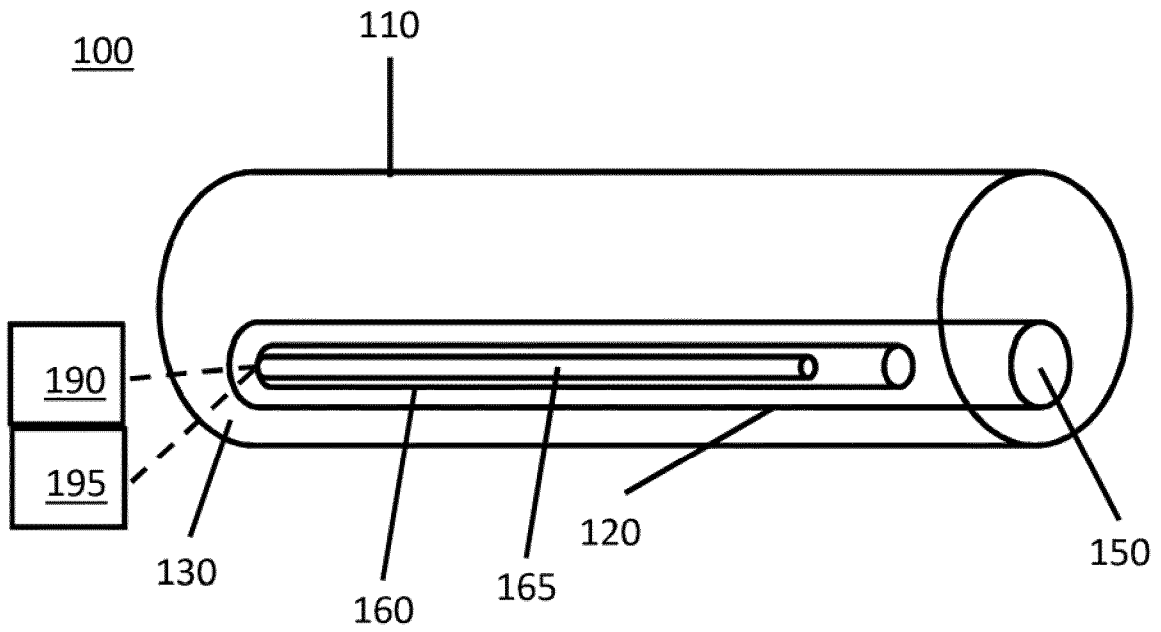


FIG 1B

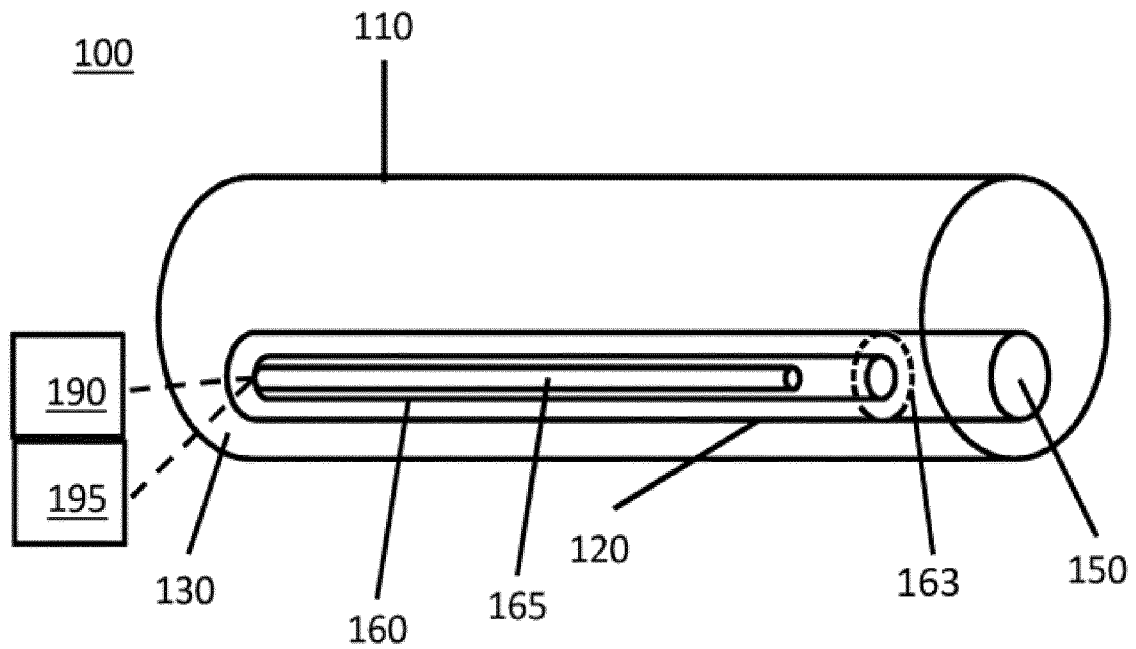


FIG 1C

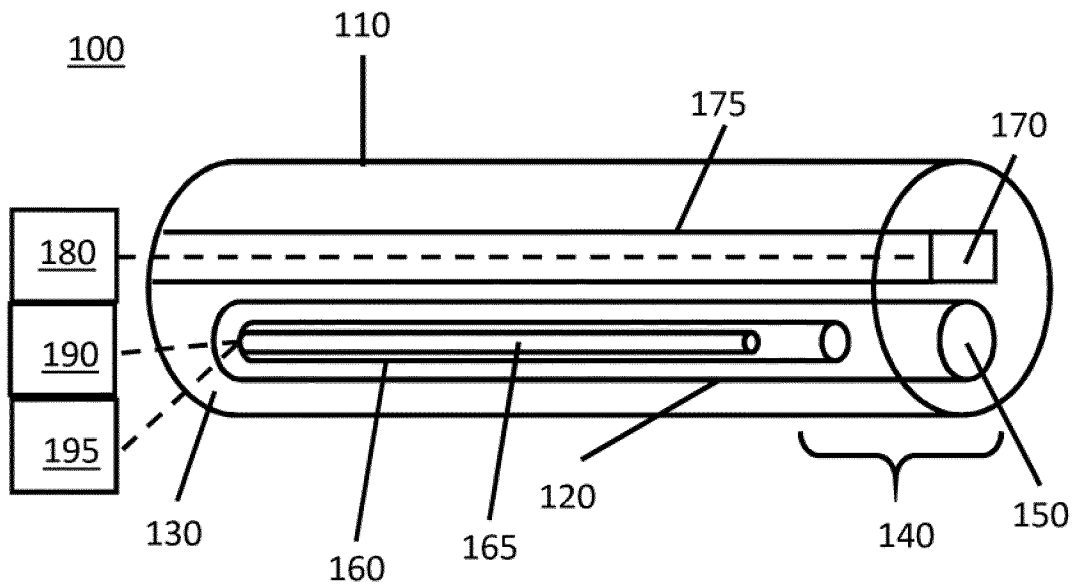


FIG 1D

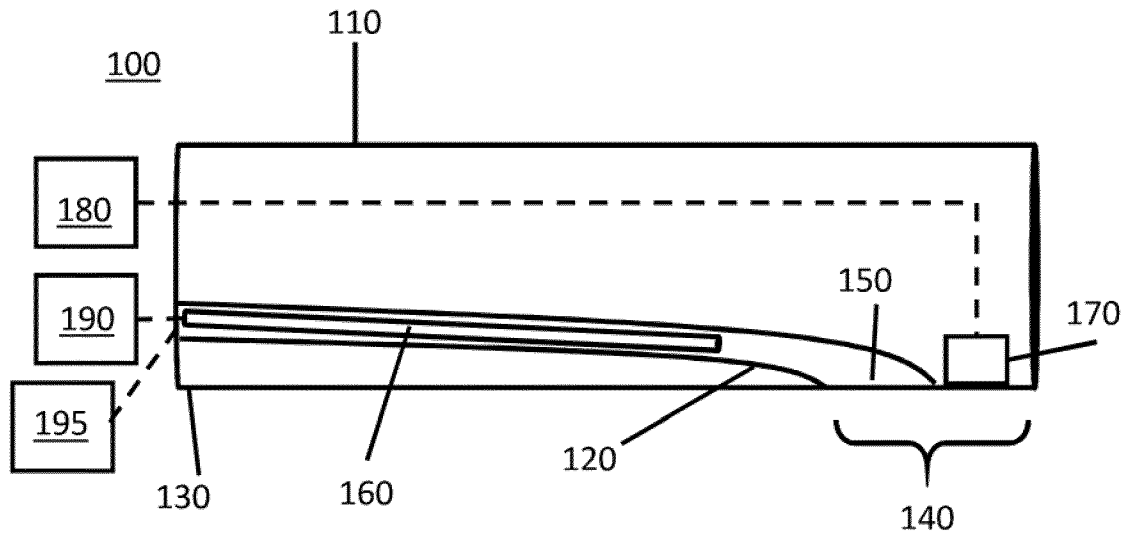


FIG 1E

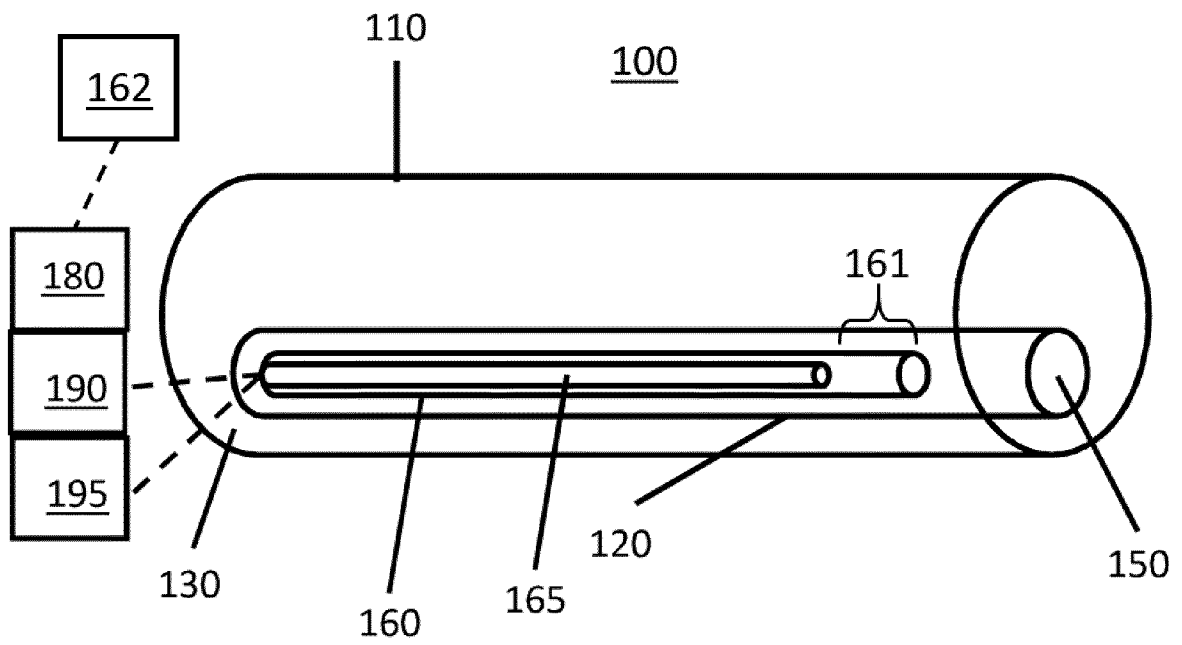


FIG 1F

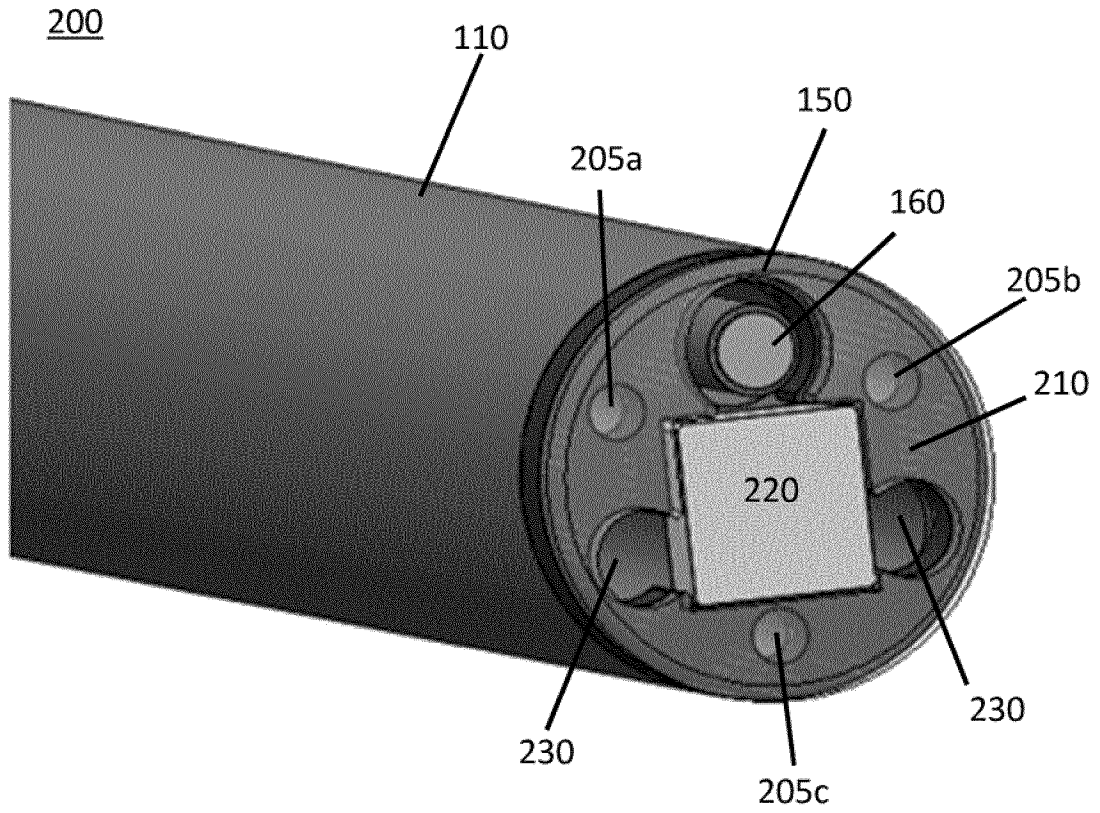


FIG 2A

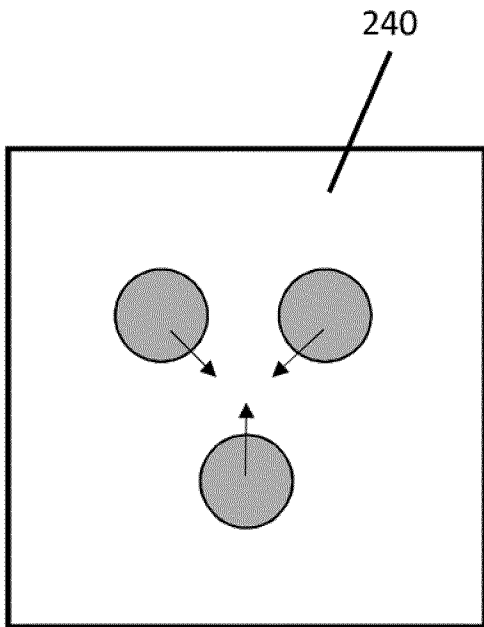


FIG 2B

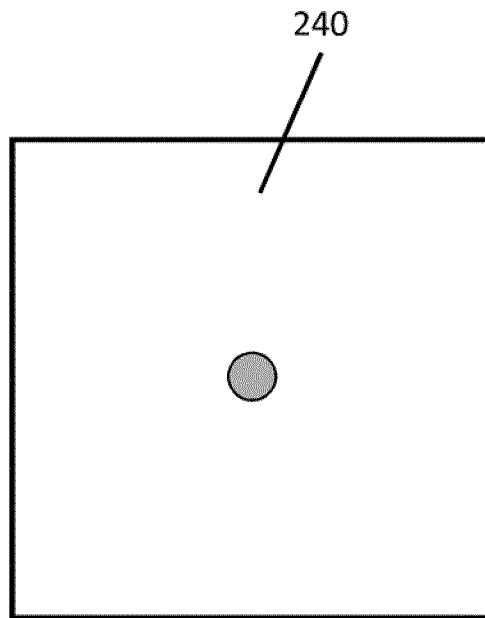


FIG 2C

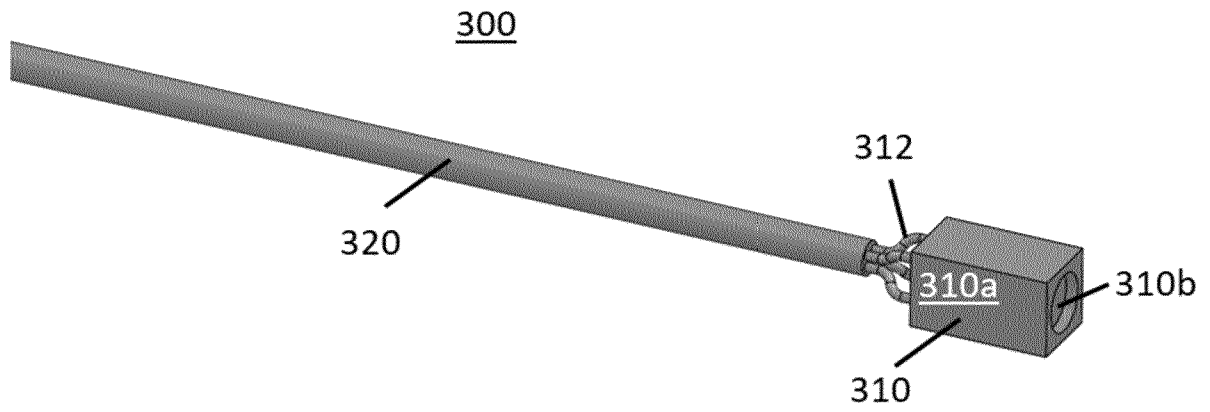


FIG 3

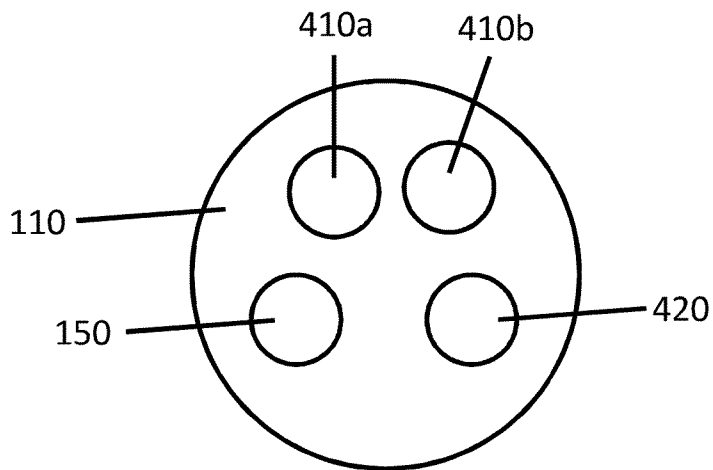


FIG 4

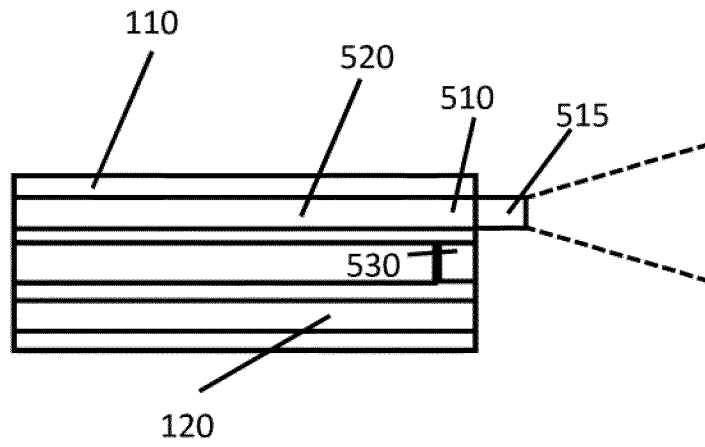


FIG 5A

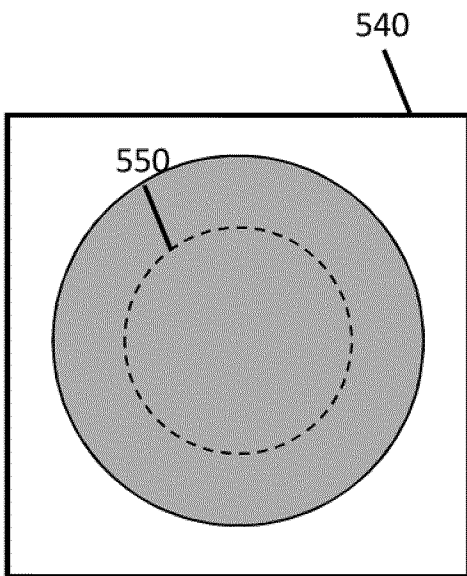


FIG 5B

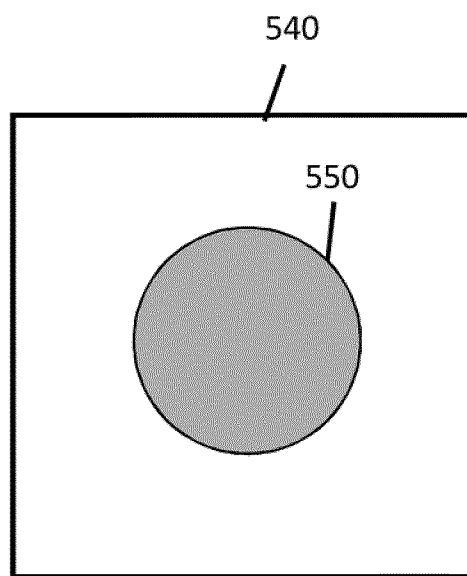


FIG 5C

7/26

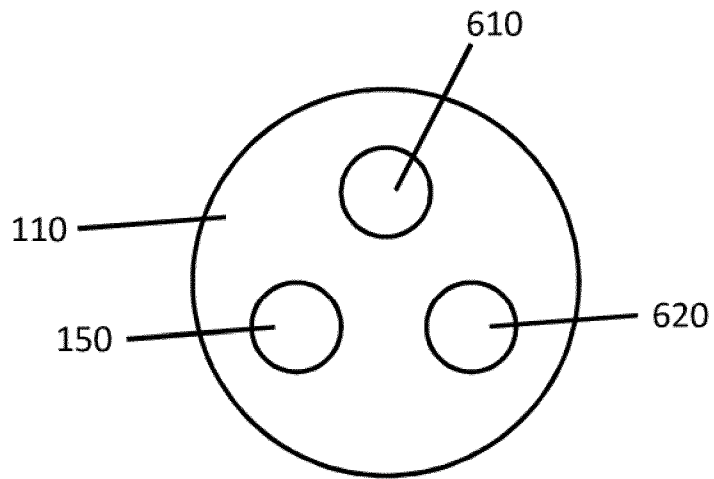


FIG 6

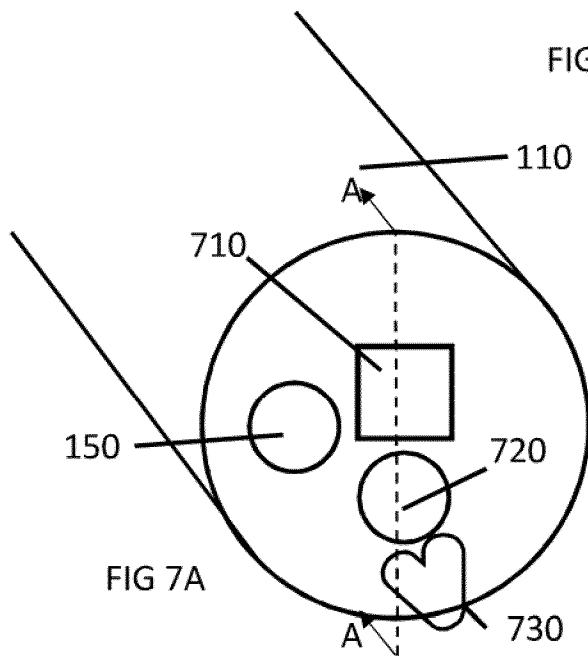


FIG 7A

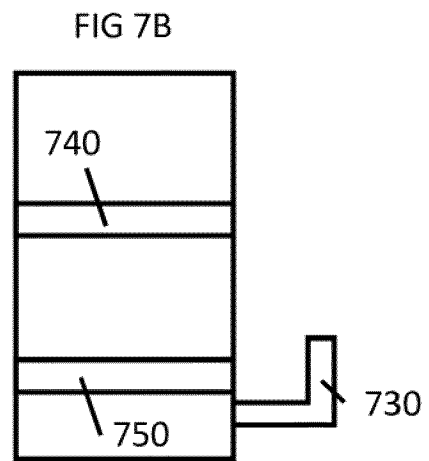


FIG 7B

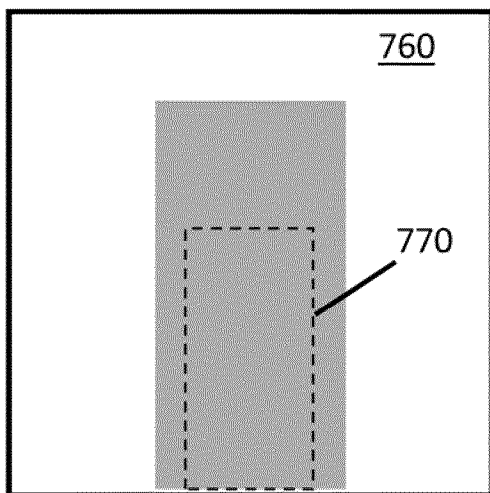


FIG 7C

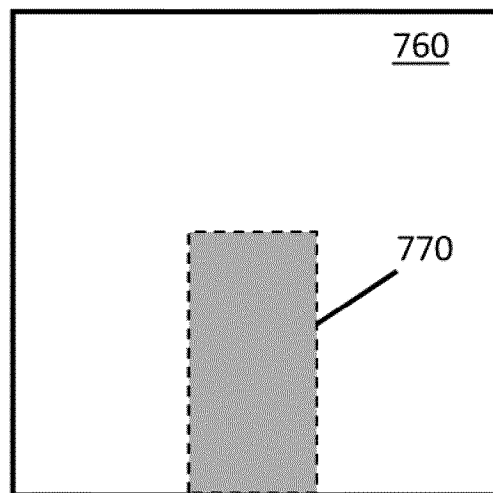


FIG 7D

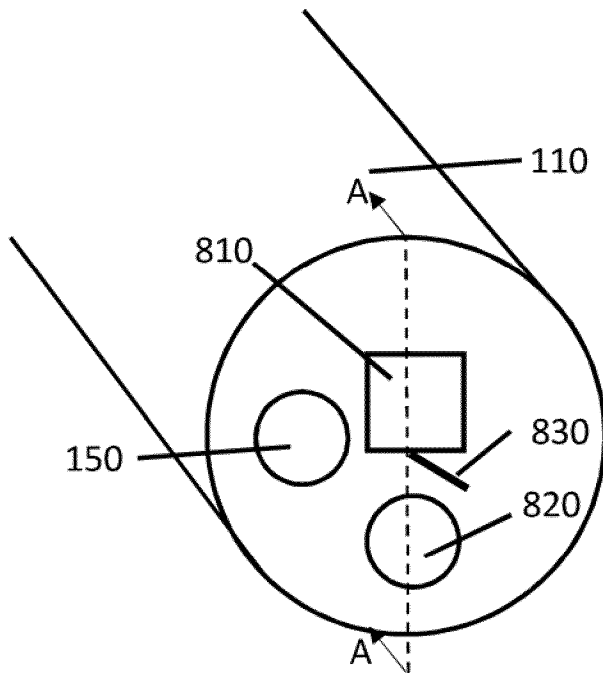


FIG 8A

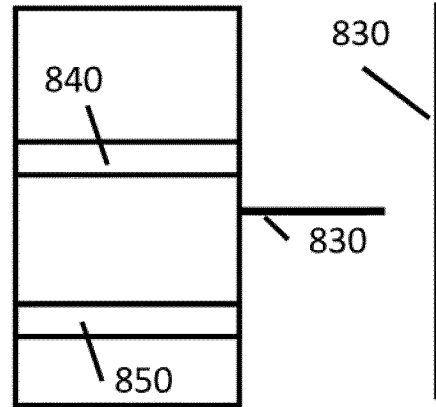


FIG 8B

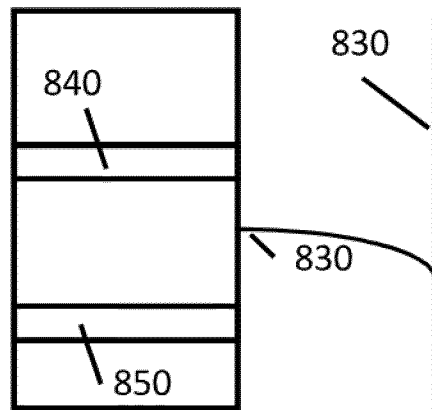


FIG 8C

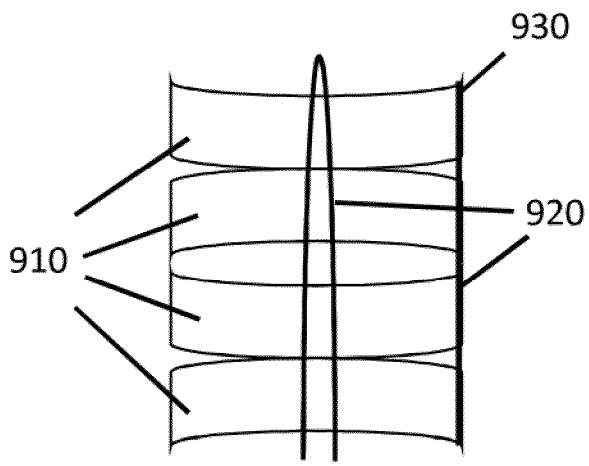


FIG 9A

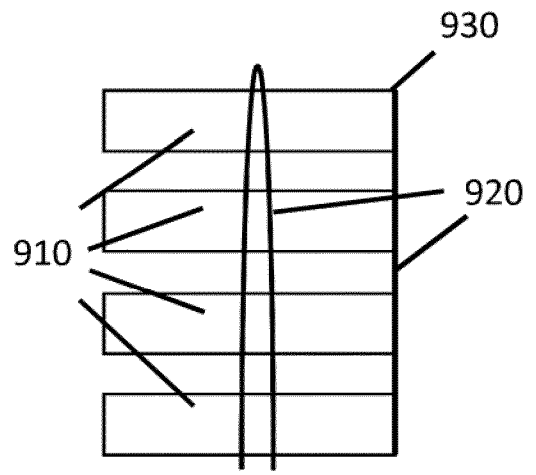


FIG 9B

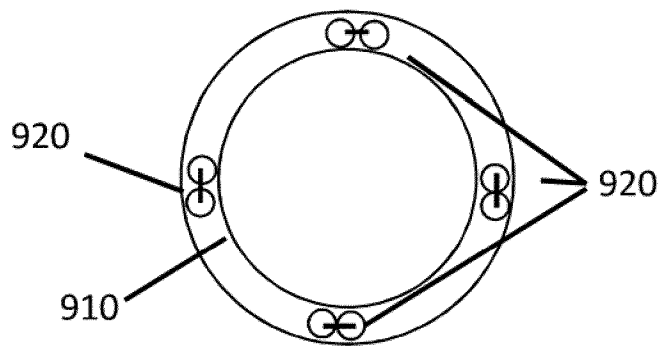


FIG 9C

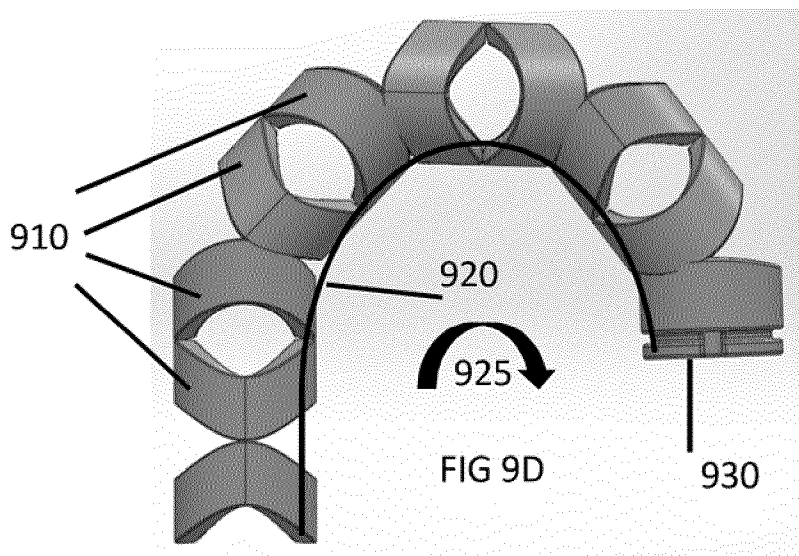
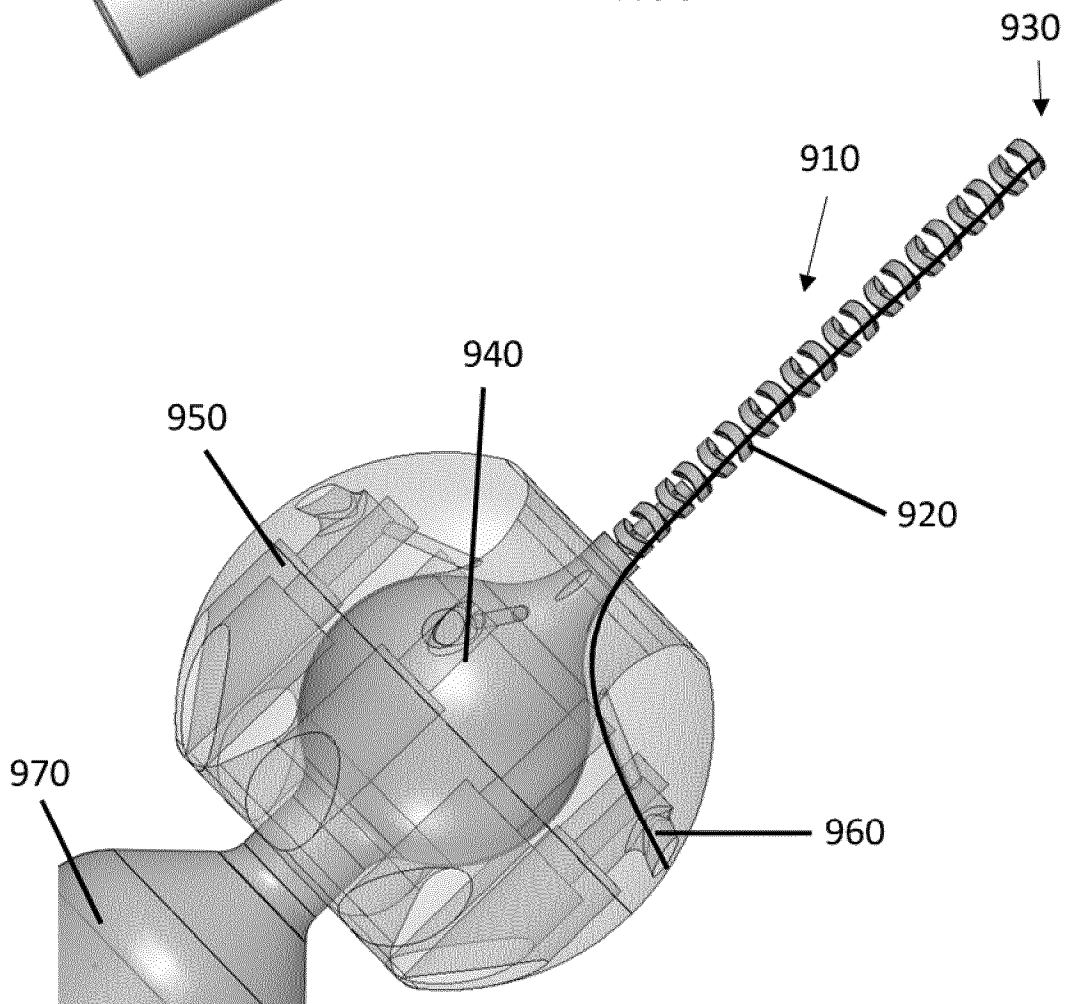
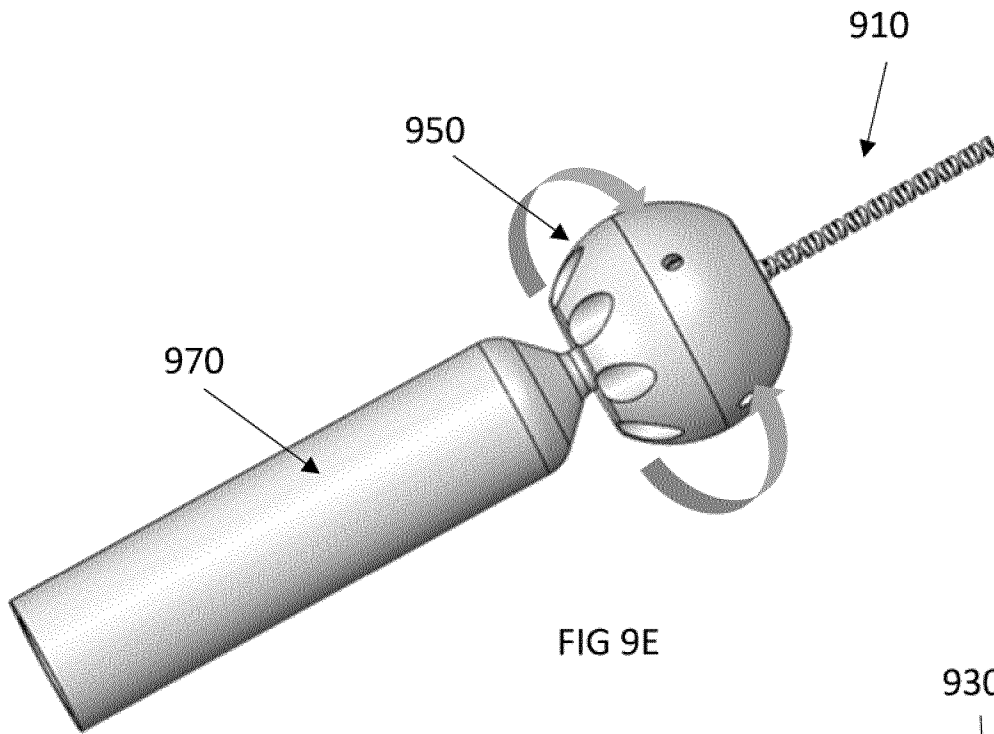


FIG 9D



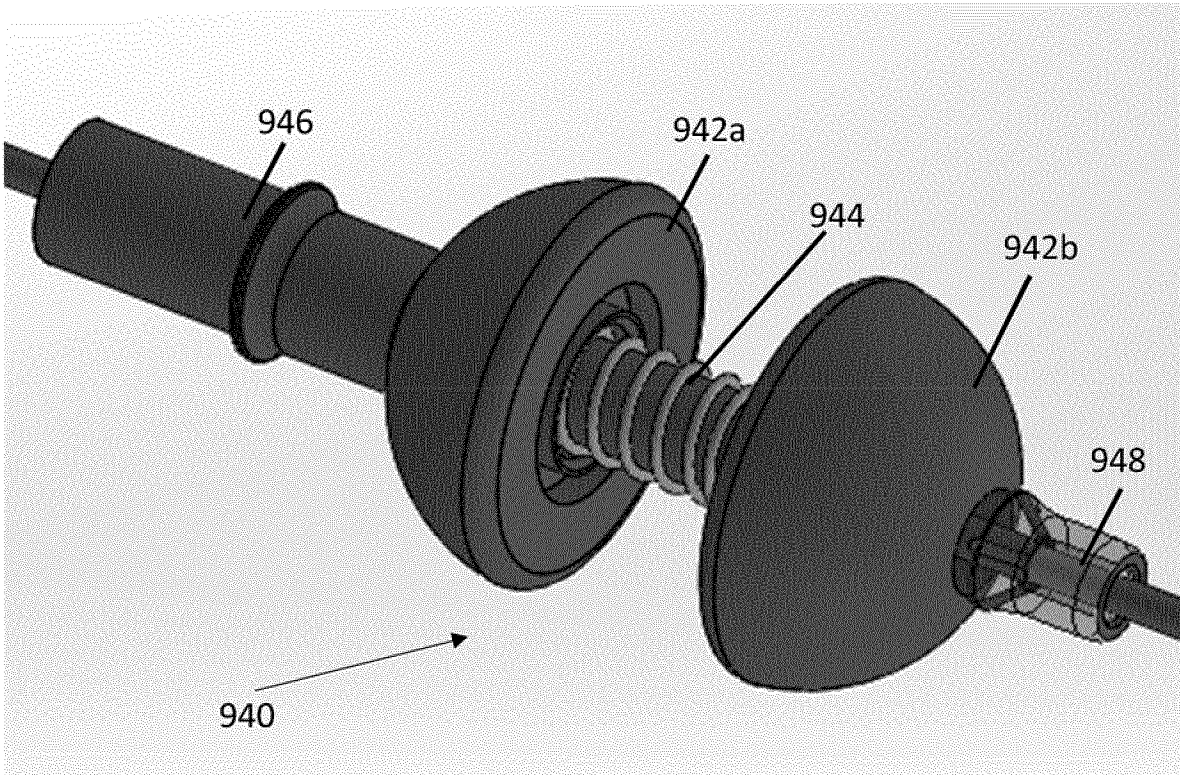


FIG 9G

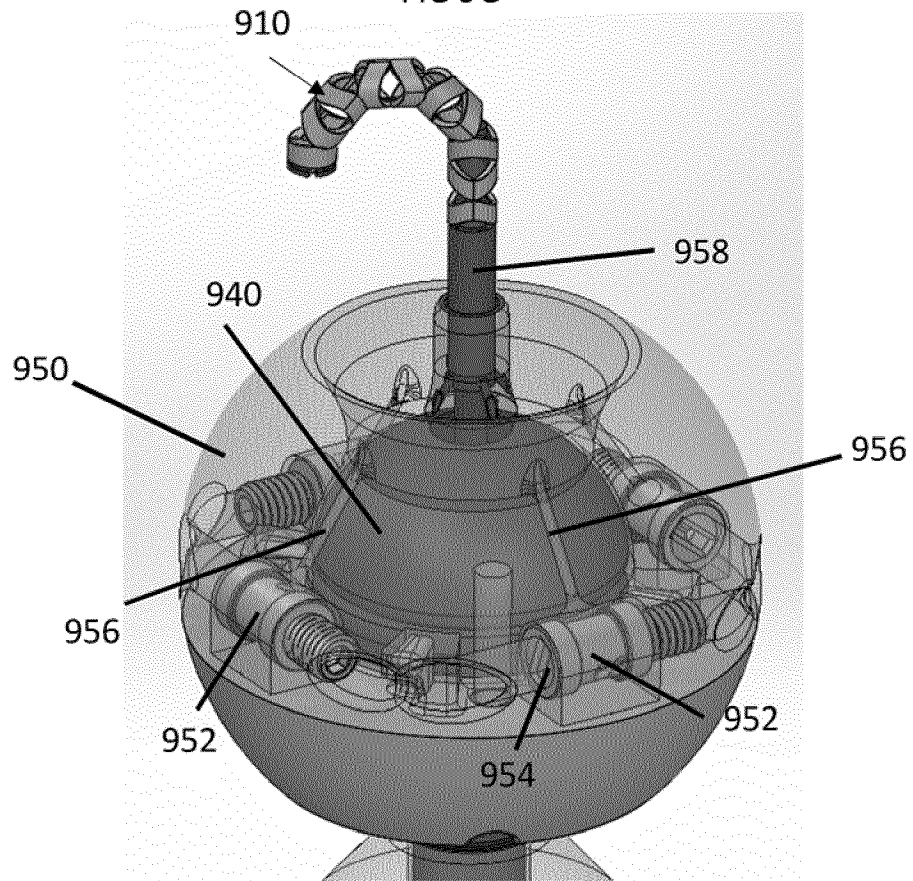


FIG 9H

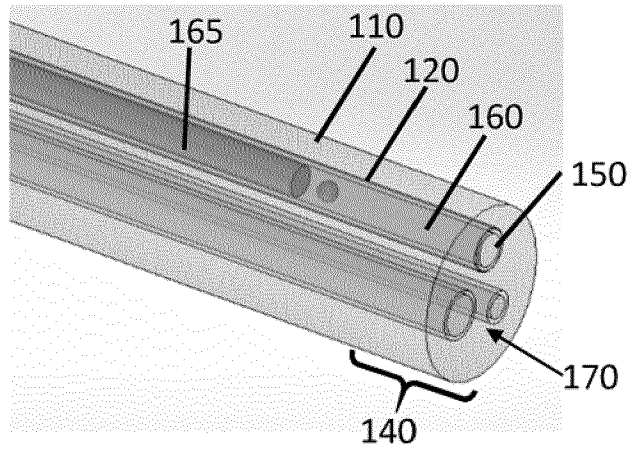


FIG 10A

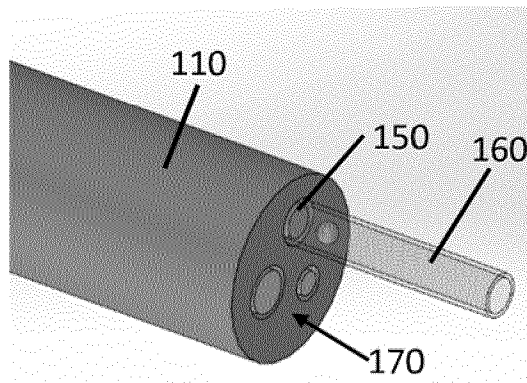


FIG 10B

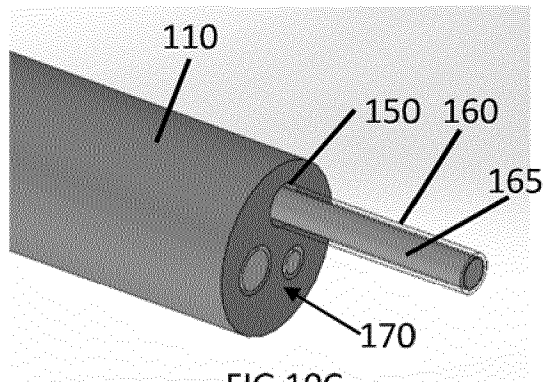


FIG 10C

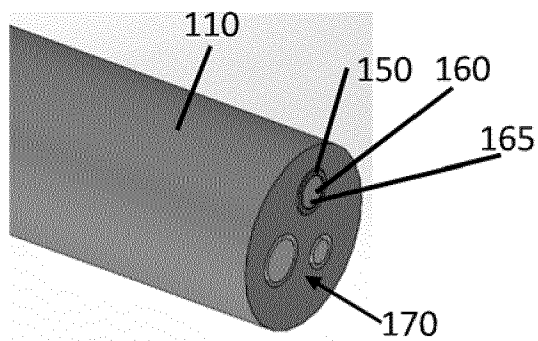


FIG 10D

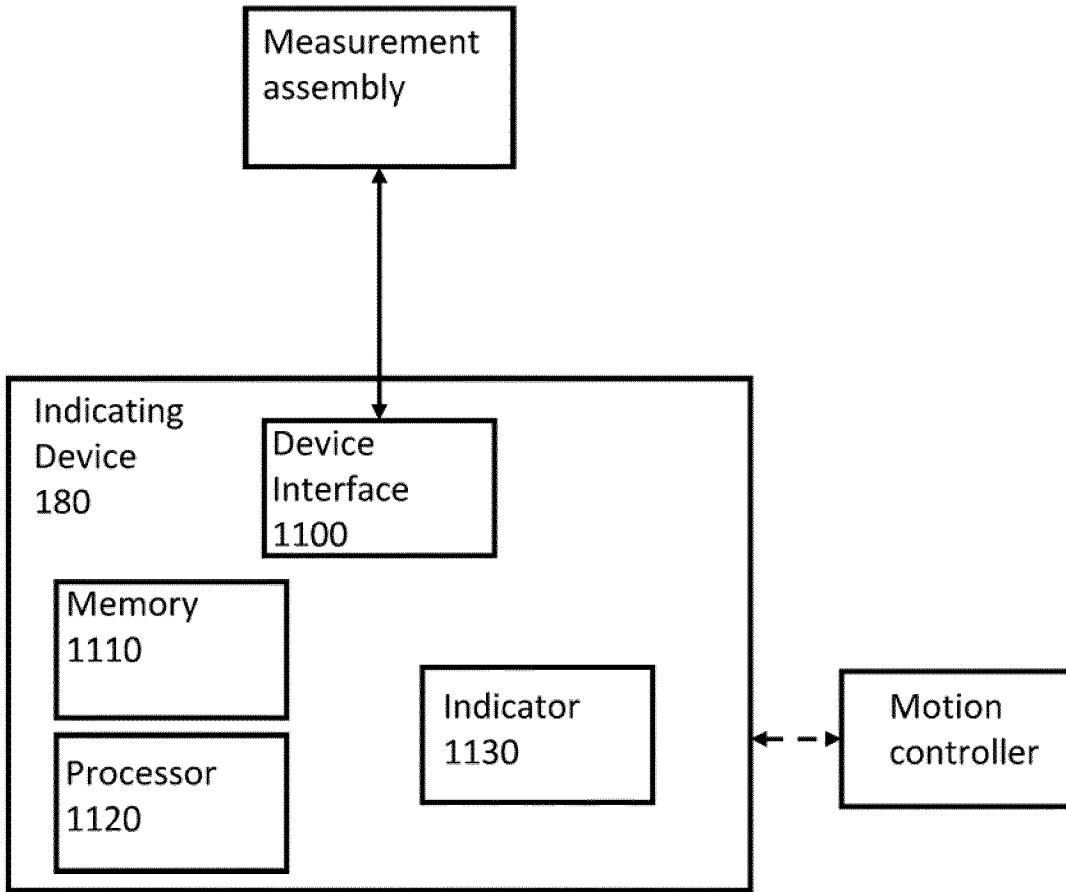


FIG 11

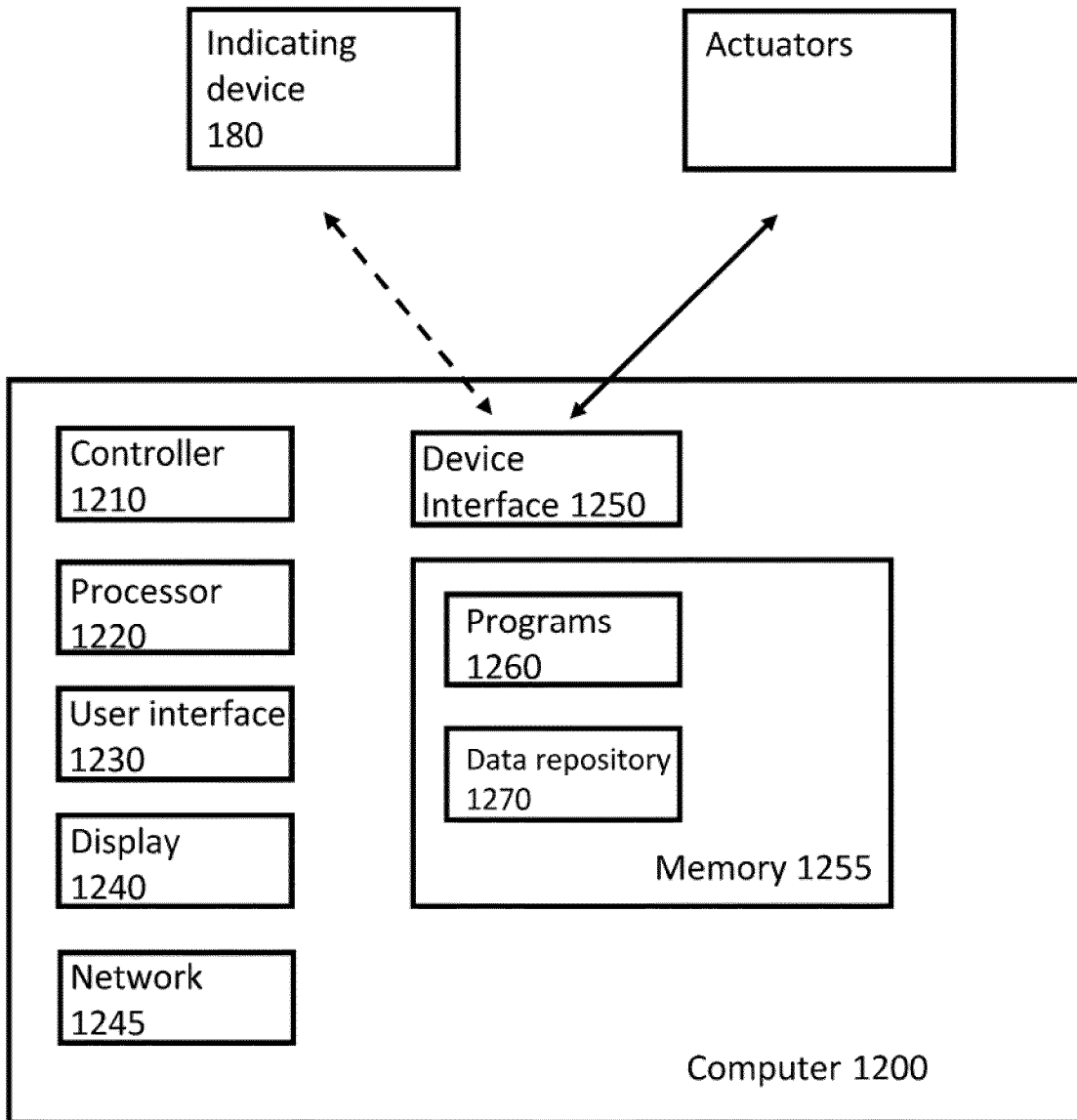


FIG 12

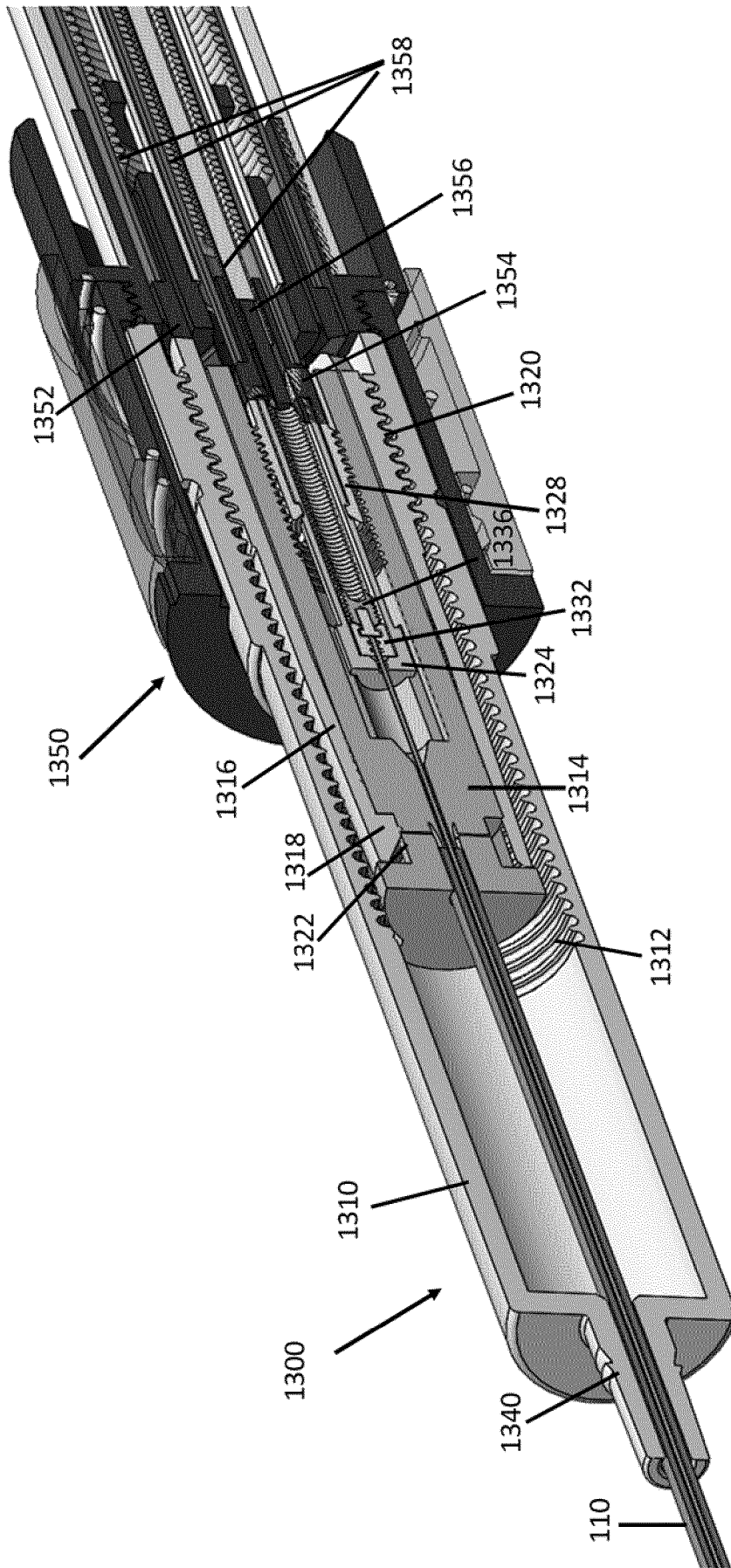
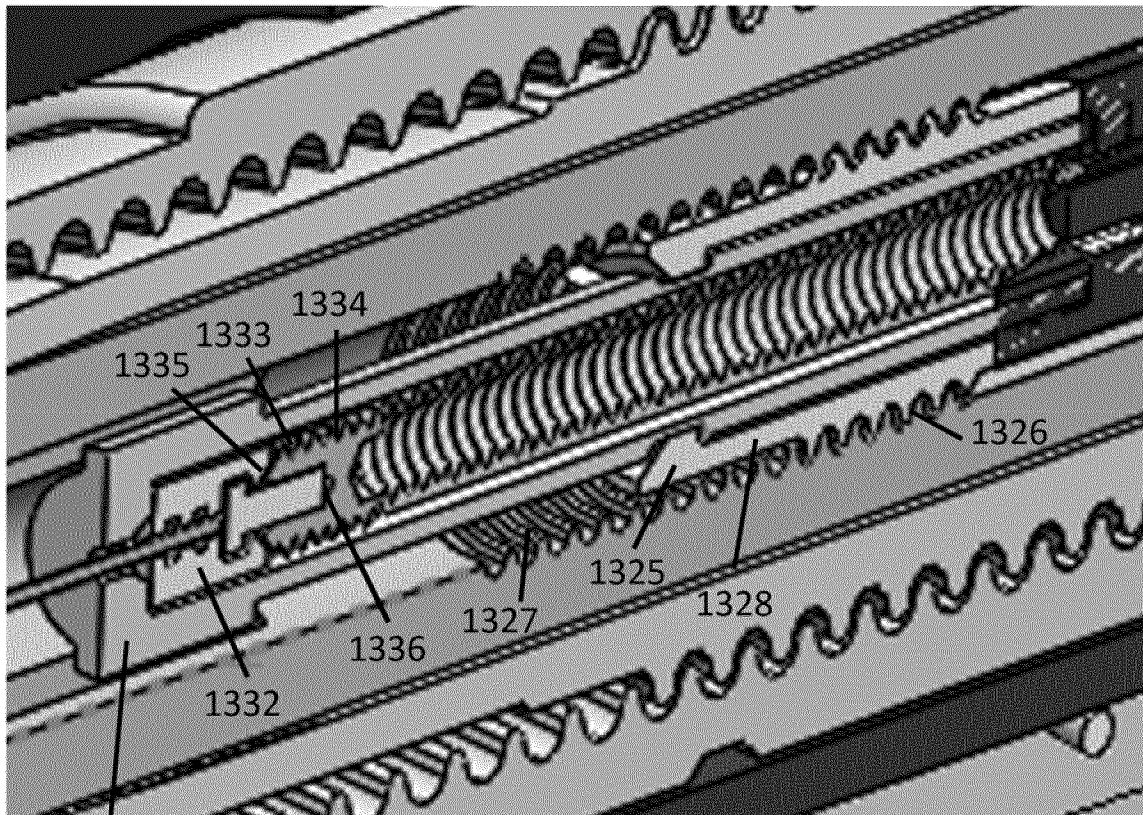


FIG 13A



1324

FIG 13B

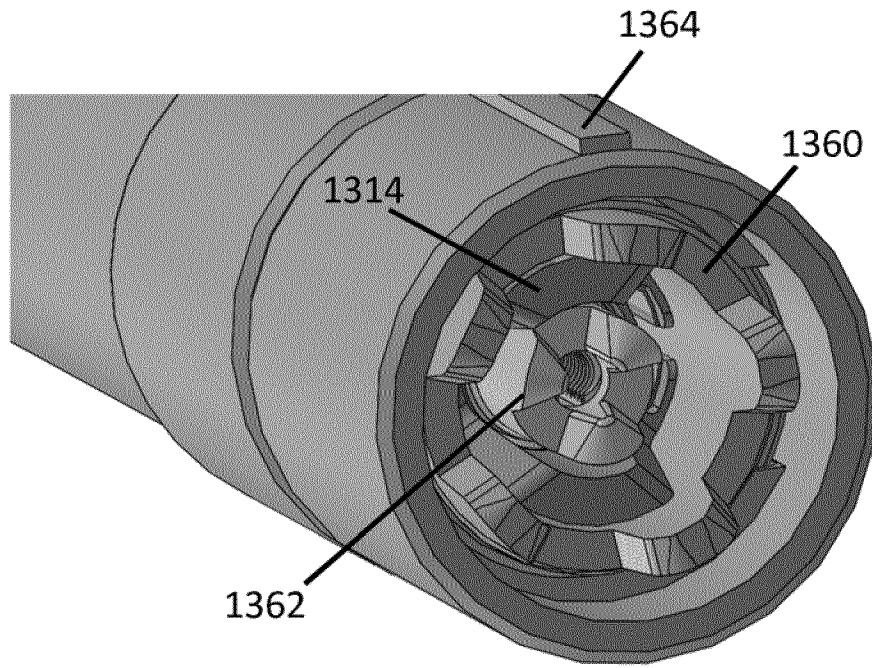


FIG 13C

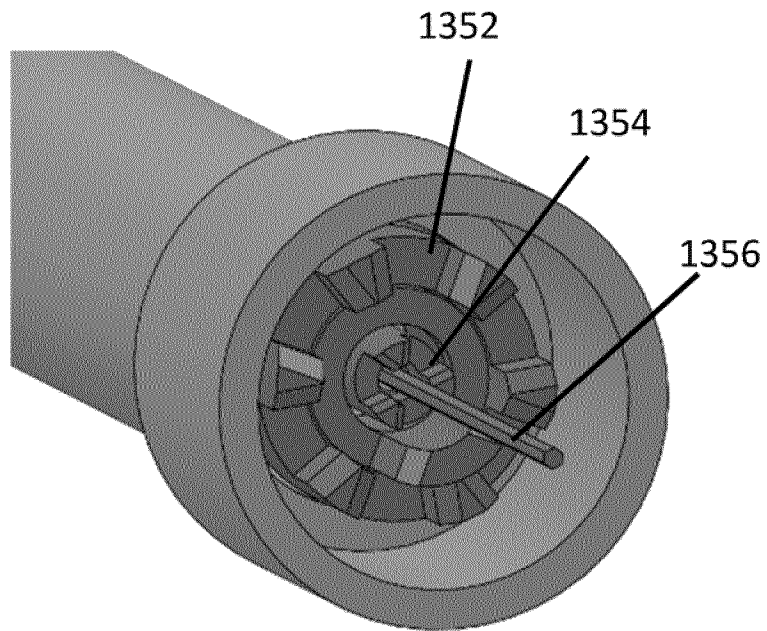


FIG 13D

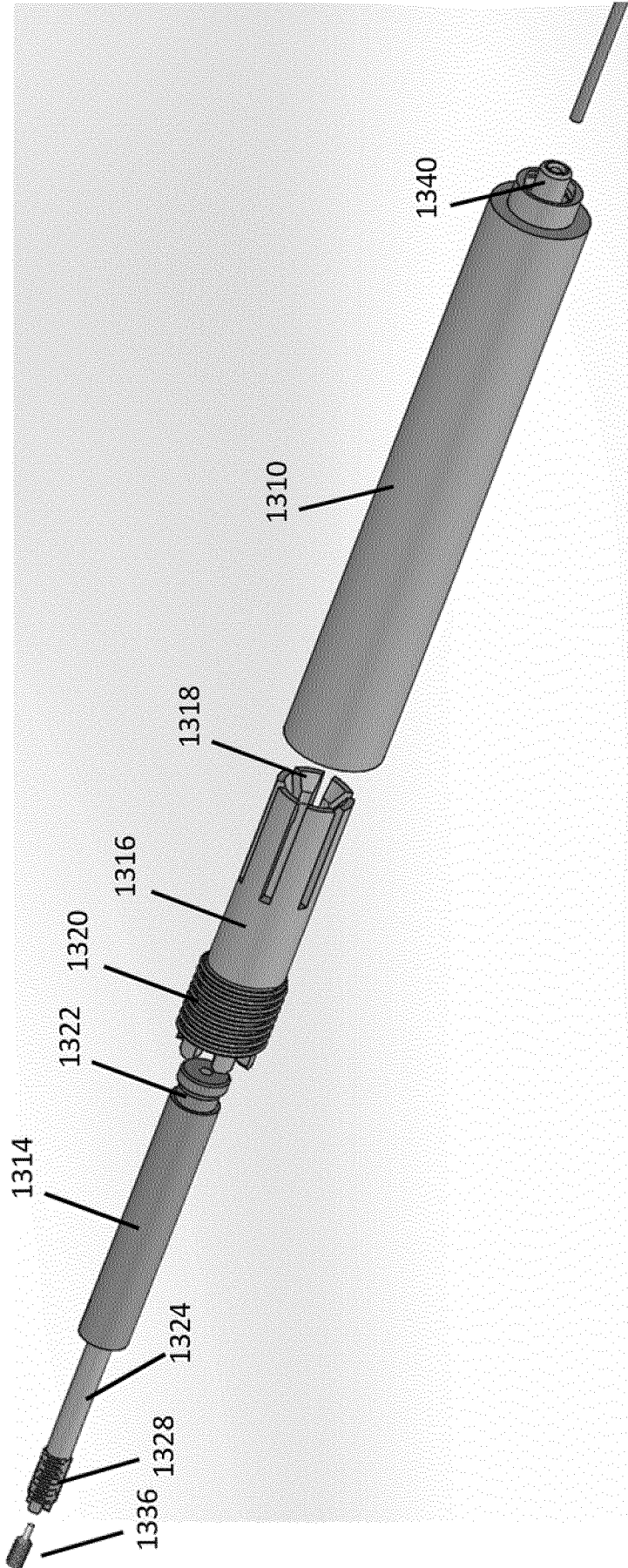


FIG 13E

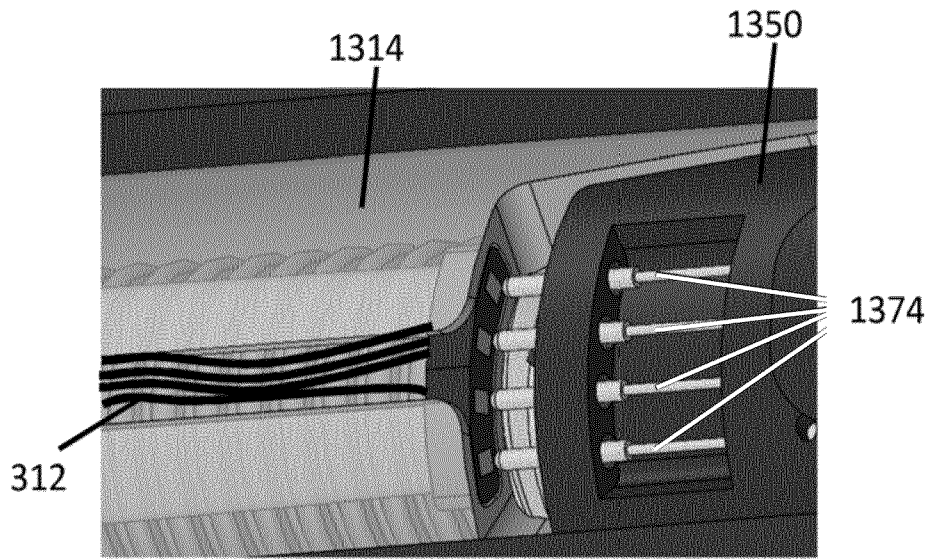


FIG 13F

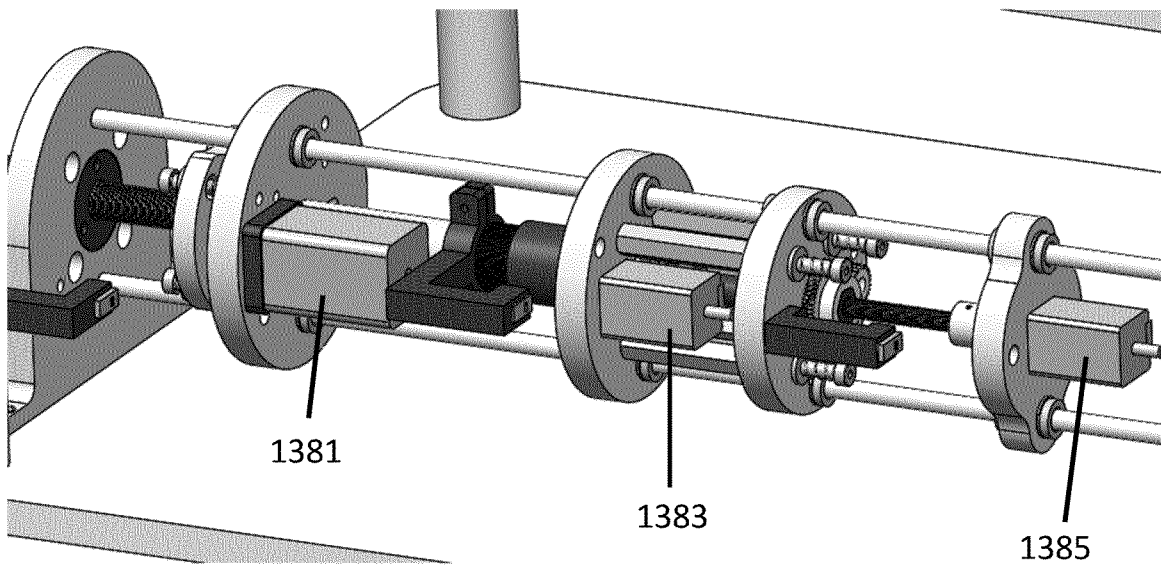


FIG 13G

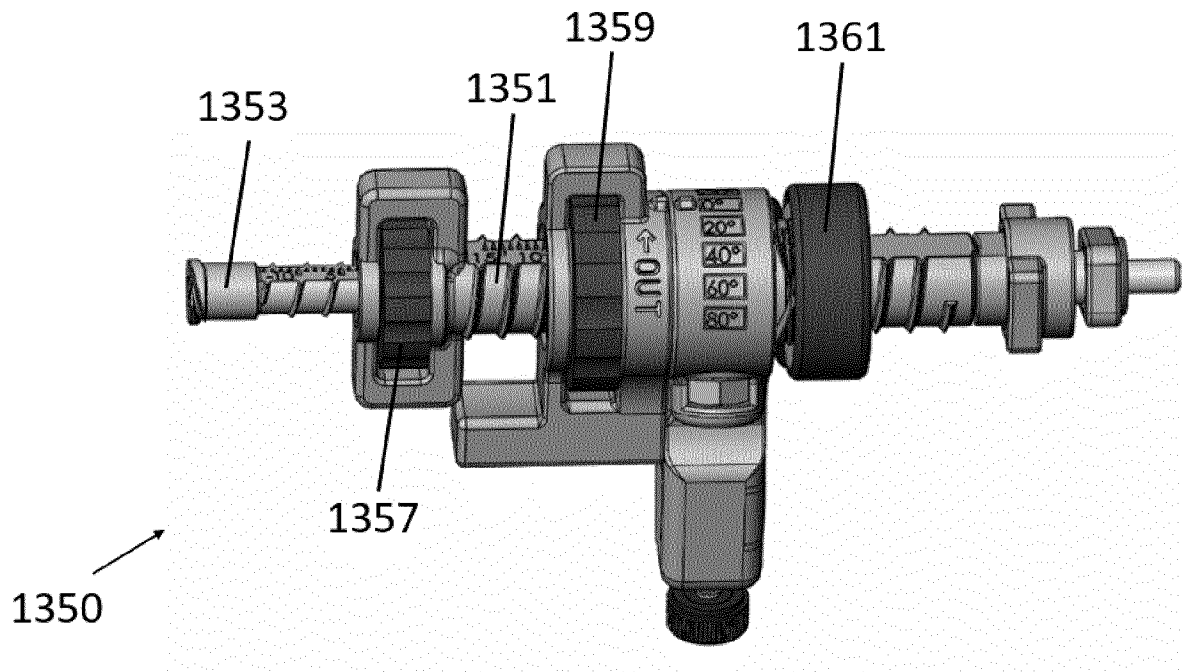


FIG 13H

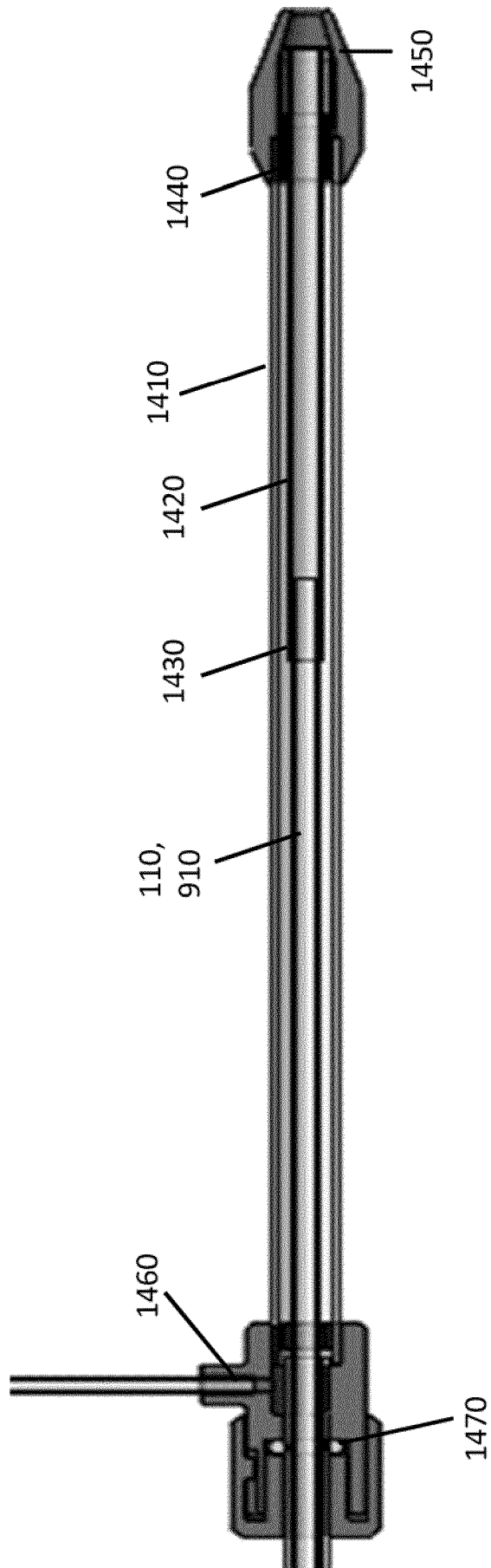


FIG. 14A

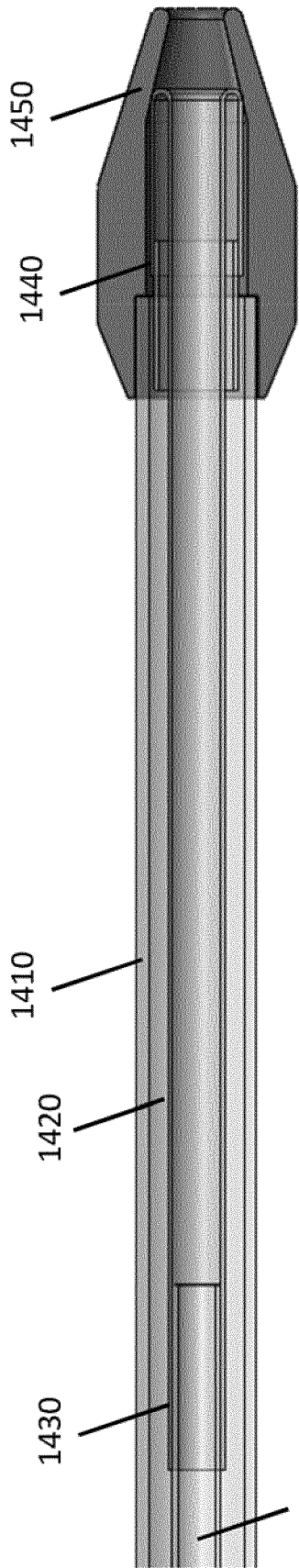


FIG. 14B

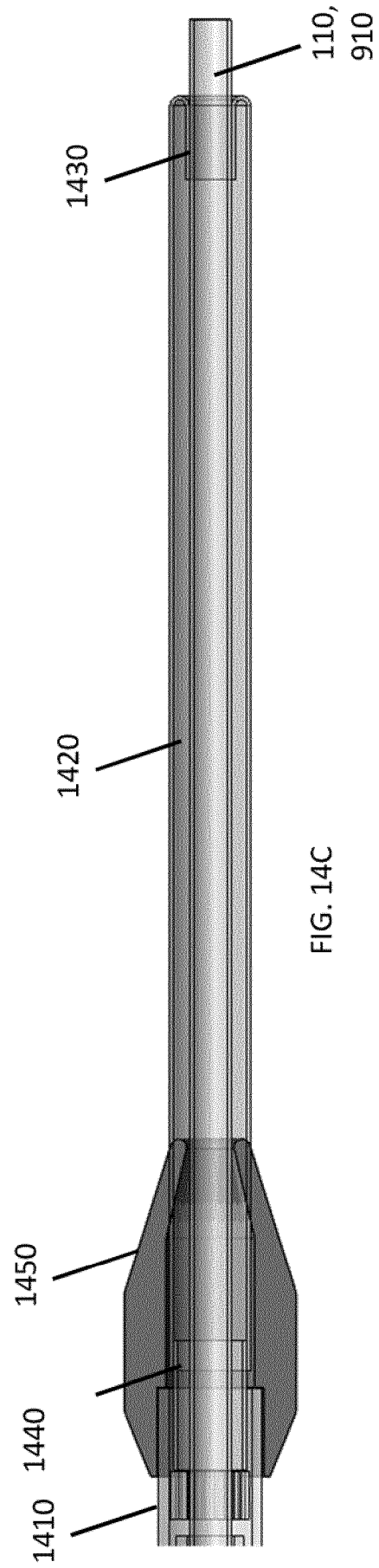


FIG. 14C

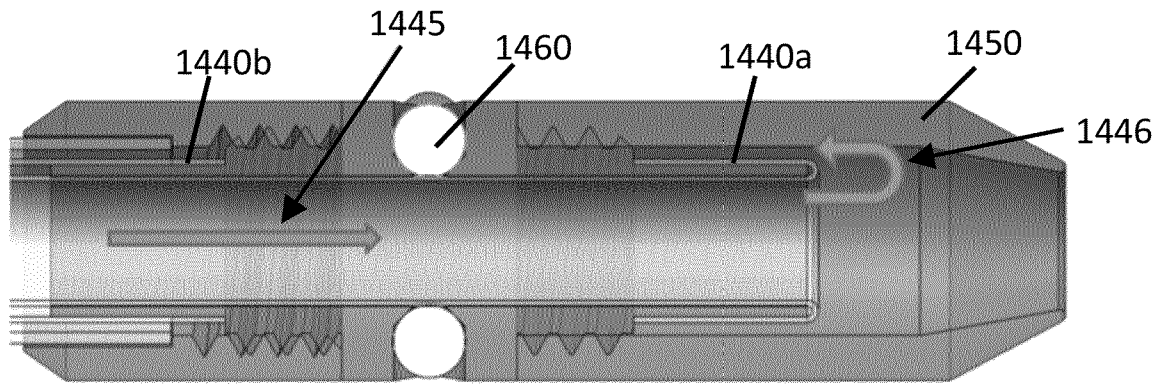


FIG. 14D

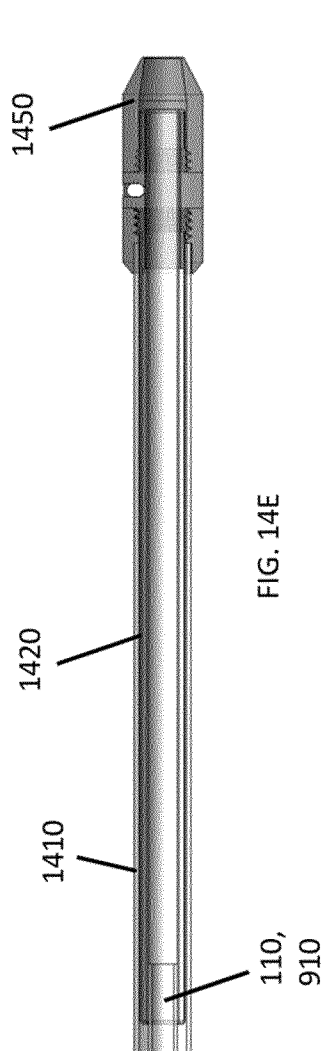


FIG. 14E

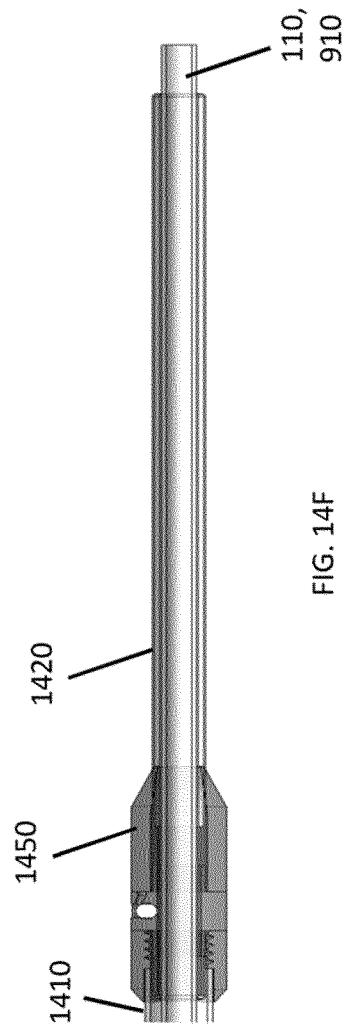


FIG. 14F

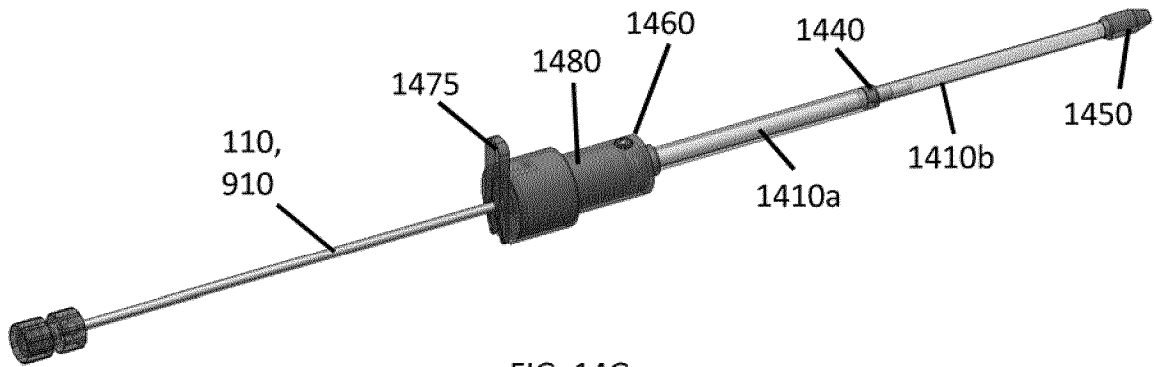


FIG. 14G

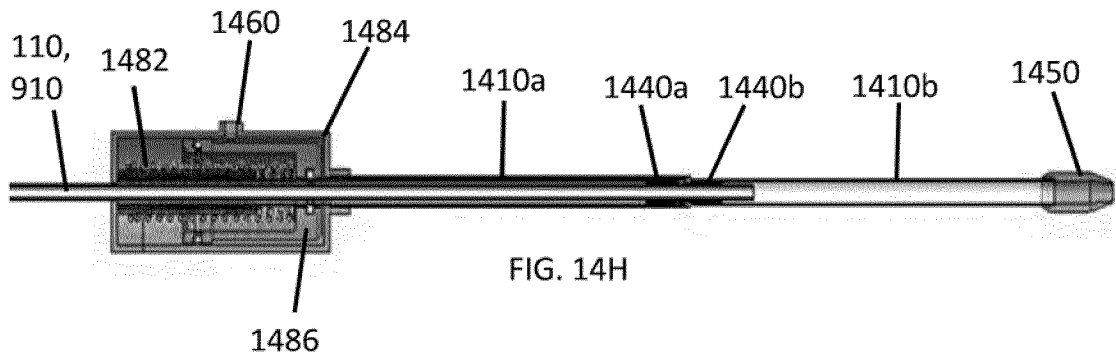


FIG. 14H

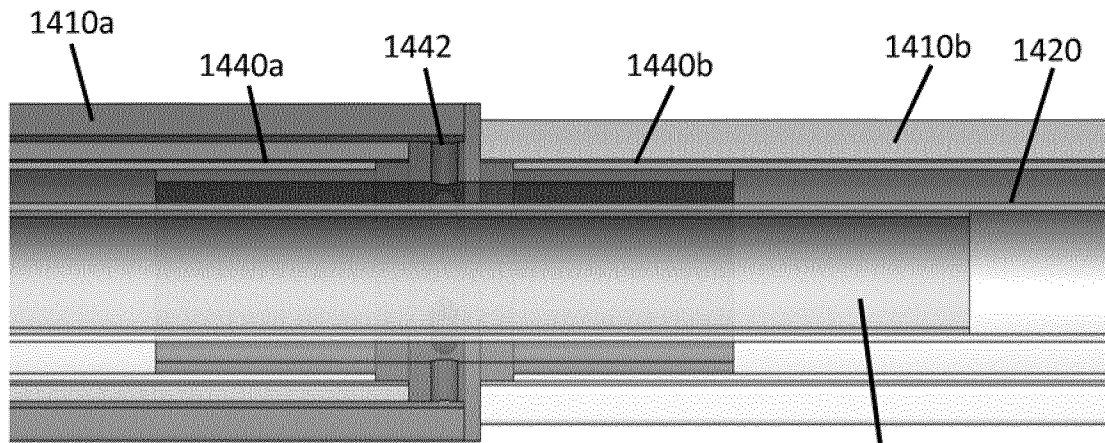
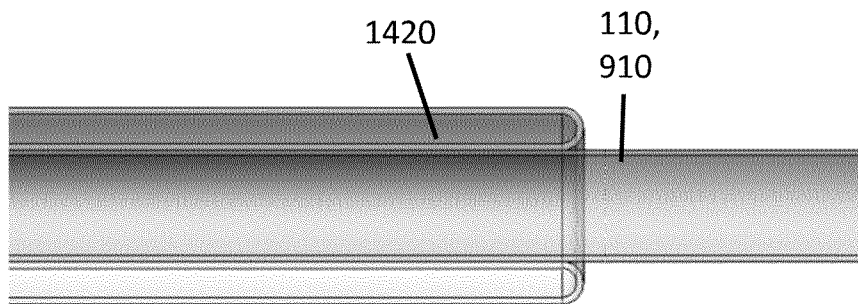
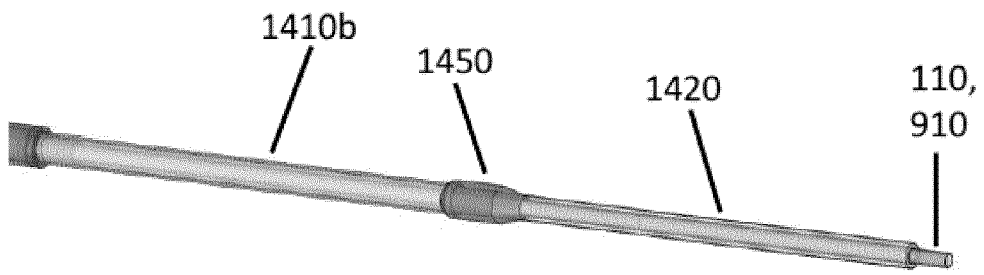
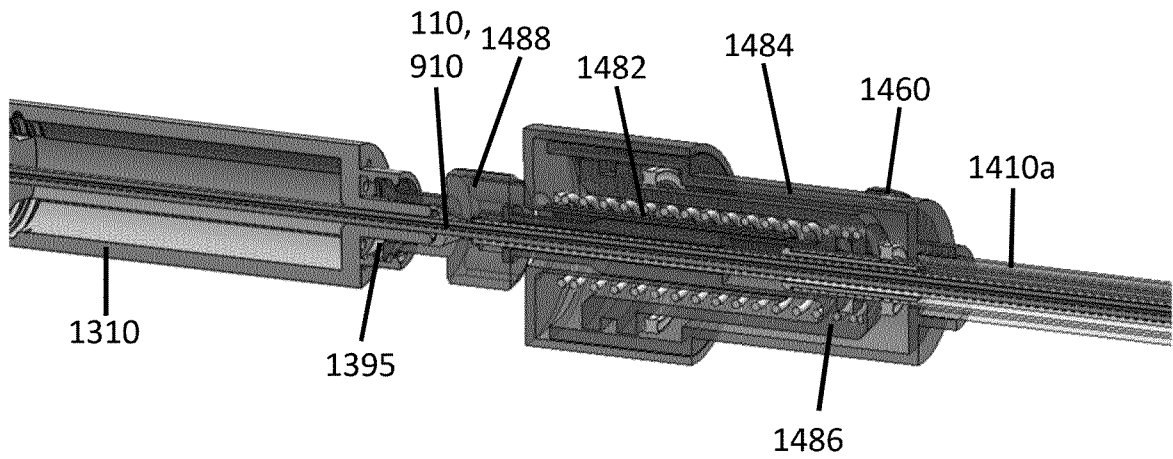


FIG. 14I

110,
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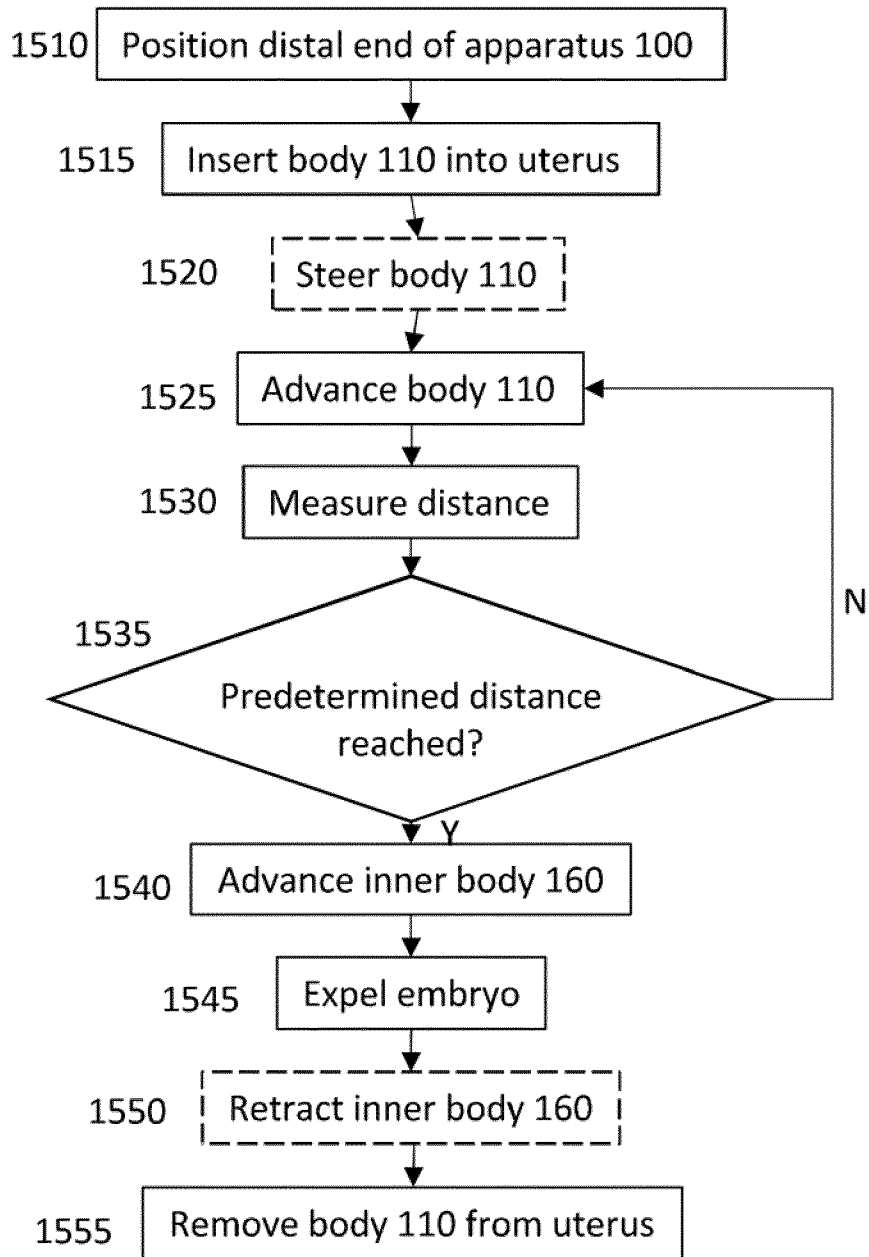


FIG 15

INTERNATIONAL SEARCH REPORT

International application No
PCT/EP2021/078712

A. CLASSIFICATION OF SUBJECT MATTER				
INV. A61B17/435 A61B90/00 A61B90/30 A61B17/00 A61B1/005				
A61B1/06 A61B1/00 A61D19/04 A61B17/34 A61M25/01				
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) A61B A61D A61M				
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched				
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) EPO-Internal, WPI Data				
C. DOCUMENTS CONSIDERED TO BE RELEVANT				
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.		
X	US 2019/159662 A1 (PAPAS RALPH SAID [LB] ET AL) 30 May 2019 (2019-05-30) paragraphs [0036], [0037], [0060] -----	1		
A	US 2019/261846 A1 (OH SEUNG-JUNE [KR] ET AL) 29 August 2019 (2019-08-29) paragraph [0064] -----	1		
<input type="checkbox"/> Further documents are listed in the continuation of Box C. <input checked="" type="checkbox"/> See patent family annex.				
* Special categories of cited documents : <table style="width: 100%; border: none;"> <tr> <td style="width: 50%; border: none;"> "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) or 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 </td> <td style="width: 50%; border: none;"> "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 "&" document member of the same patent family </td> </tr> </table>			"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) or 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 "&" document member of the same patent family
"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) or 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 "&" document member of the same patent family			
Date of the actual completion of the international search	Date of mailing of the international search report			
22 December 2021	03/03/2022			
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 Angeli, Markus			

INTERNATIONAL SEARCH REPORT

International application No.
PCT/EP2021/078712

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.:
because they relate to subject matter not required to be searched by this Authority, namely:

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.:
1-14, 16, 40-43

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.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

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

1. claims: 1-14, 16, 40-43

An apparatus for delivering a fertilized egg directed to a measurement assembly for indicating a distance between the apparatus and an inner member.

2. claim: 15

An apparatus for delivering a fertilized egg directed to a marker for enhancing positioning of an inner body with respect to the apparatus.

3. claim: 17

An apparatus for delivering a fertilized egg directed to an imaging device for enhancing the view to the delivering site.

4. claims: 18-22

An apparatus for delivering a fertilized egg directed to a steering mechanism for enhancing manoeuvring of the apparatus.

5. claims: 23-29

An apparatus for delivering a fertilized egg directed to an evertable tube for facilitating introduction of the apparatus.

6. claims: 30-39

An apparatus for delivering a fertilized egg directed to a lead screw for enhancing actuation of the apparatus.

INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No

PCT/EP2021/078712

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
US 2019159662 A1	30-05-2019	US 2019159662 A1	30-05-2019
		WO 2020055927 A1	19-03-2020

US 2019261846 A1	29-08-2019	US 2019261846 A1	29-08-2019
		US 2020375449 A1	03-12-2020
