#### (12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization

International Bureau





(10) International Publication Number WO 2014/182959 A2

(43) International Publication Date 13 November 2014 (13.11.2014)

(51) International Patent Classification:

A61M 25/00 (2006.01) A61B 17/12 (2006.01)

A61M 5/172 (2006.01)

(21) International Application Number:

PCT/US2014/037385

(22) International Filing Date:

8 May 2014 (08.05.2014)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

61/821,058 8 May 2013 (08.05.2013) US 61/915,425 12 December 2013 (12.12.2013) US 61/917,131 17 December 2013 (17.12.2013) US

- (71) Applicant: EMBOLX, INC. [US/US]; 5760 Arboretum Drive, Los Altos, CA 94024 (US).
- (72) Inventors: ALLEN, Michael, P.; 5760 Arboretum Drive, Los Altos, CA 94024 (US). BROMMER, Robert; 5760 Arboretum Drive, Los Altos, CA 94024 (US). HANNON, Trevor; 5760 Arboretum Drive, Los Altos, CA 94024 (US). DE NEVE, Robert; 5760 Arboretum Drive, Los Altos, CA 94024 (US).
- (74) Agents: LIMBACH, Douglas, C. et al.; SHAY GLENN LLP, 2755 Campus Drive, Suite 210, San Mateo, CA 94403 (US).

- (81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AO, AT, AU, AZ, BA, BB, BG, BH, BN, BR, BW, BY, BZ, CA, CH, CL, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IR, IS, JP, KE, KG, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PA, PE, PG, PH, PL, PT, QA, RO, RS, RU, RW, SA, SC, SD, SE, SG, SK, SL, SM, ST, SV, SY, TH, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW.
- (84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LR, LS, MW, MZ, NA, RW, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, RU, TJ, TM), European (AL, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HR, HU, IE, IS, IT, LT, LU, LV, MC, MK, MT, NL, NO, PL, PT, RO, RS, SE, SI, SK, SM, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, KM, ML, MR, NE, SN, TD, TG).

#### Published:

 without international search report and to be republished upon receipt of that report (Rule 48.2(g))



 $\textbf{(54) Title} : \textbf{DEVICE AND METHODS FOR TRANSVASCULAR TUMOR EMBOLIZATION WITH INTEGRATED FLOW REGULATION \\$ 

(57) Abstract: A catheter assembly may be provided with a catheter body and an inflatable balloon. The catheter body has a proximal end, a distal end and a balloon inflation lumen. The inflatable balloon is attachable to the distal end of the catheter body. The balloon has an inner surface that at least partially defines an interior volume. The balloon is configured such that the interior volume can be in fluid communication with the inflation lumen of the catheter body to inflate the balloon. The balloon also has a proximal surface and a distal surface. The balloon is provided with a channel that extends through the balloon. The channel is configured to provide fluid communication between the proximal surface of the balloon and the distal surface of the balloon. Other catheter assemblies and methods of use are also disclosed.

# DEVICE AND METHODS FOR TRANSVASCULAR TUMOR EMBOLIZATION WITH INTEGRATED FLOW REGULATION

#### CROSS REFERENCE TO RELATED APPLICATIONS

5 [0001] This application claims the benefit of U.S. Provisional Applications No. 61/821,058 filed May 8, 2013, No. 61/915,425 filed December 12, 2013, and No. 61/917,131, filed December 17, 2013, each of which is herein incorporated by reference in its entirety.

#### INCORPORATION BY REFERENCE

10 [0002] All publications and patent applications mentioned in this specification are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

#### **FIELD**

[0003] This application relates generally to medical methods and devices. More specifically, the present application discloses various embodiments of occlusion devices adapted to a catheter, and methods for their use in delivering fluids, embolic materials and other therapeutic agents to sites within the body.

20 BACKGROUND

25

30

[0004] There are over one million cases of cancer diagnosed each year in the United States and numerous approaches of therapy including systemic chemotherapy, radiation and surgical resection. Given that systemic chemotherapy and radiation interact with healthy tissue, complications and toxicity often result. Targeted drugs are now being used and produce a lower rate of complications. Ablative approaches, including microwave, radiofrequency and cryogenic therapies have been used; however, these methods are often not selective and tissues and organs surrounding or below the tumor can be affected.

[0005] According to the National Institute of Health, 30,640 people were diagnosed with primary liver cancer (hepatocellular carcinoma, HCC) and 142,820 people were diagnosed with colorectal cancer in the U.S. in 2013. Seventy five percent of these will metastasize to the liver. Liver resection and transplant are the only curative means; however, only small numbers of patients are eligible. Systemic chemotherapy for primary and metastatic tumors in the liver is ineffective, having a response rate of about 20% and a survival benefit of 10.7 months vs. 7.9 months over symptomatic care.

[0006] Catheters are commonly used in medicine for delivery of fluids, therapeutics, and implants, and in sampling tissues and bodily fluids. Catheters can be constructed with balloons or other tools to dilate tissue, block fluid flow or isolate segments of the anatomy, such as in treatment of the cancers described above.

- 5 [0007] Trans-Arterial Embolization therapy is the transvascular injection of drug and/or embolic agents directly into the tumor vasculature using a microcatheter. Embolization therapy causes a shutdown of blood flow and, when drug or radioactivity is present, simultaneous release of high concentrations of drug or radioactivity. The technique is also noted for its very low level of toxicity.
- 10 [0008] In the early 1980's, transarterial chemoembolization (TACE) began to be used as a selective cancer therapy. In this method, chemotherapeutic and embolic agents are injected directly into the vasculature of the tumor, a technique that is most common for the treatment of hepatocellular carcinoma. More recently, transarterial radioembolization (TARE) has been used clinically. In this method, radioactive embolic particles, typically yttrium-90 (y90), are injected rather than chemotherapeutic agents. Although the liver is a common target for TACE and TARE, other organs, including, but not limited to, the pancreas, lung, kidney, prostate, stomach, colon and head and neck have been treated using these methods. Chemoembolization was established as a standard of care for intermediate stage hepatocellular carcinoma in 2006.
  - [0009] Numerous studies have demonstrated transarterial embolization to be effective on a number of primary cancers and to have better performance than chemotherapy for both HCC and metastatic colorectal cancers in the liver; however, studies show inconsistent outcomes with reported tumor responses from 15% to 85%. Although anatomical and individual differences are clearly of significance in between-patient variation, clinical studies, each of which include a range of patients, show very different outcomes, indicating that there currently is little procedural optimization or standardization.

20

25

30

35

[0010] The above procedures are accomplished by inserting a small catheter into the femoral artery at the groin and navigating it into the liver vasculature, typically the hepatic artery, then into the right or left lobe of the liver or more selectively into particular segments of the liver or super-selectively directly into or adjacent to the tumor. Super-selective transarterial delivery of antitumor agents into the tumor vasculature has become state-of-the-art and requires catheters that can reach into small vessels. Presently, standard microcatheters, typically at or about 3 Fr are used to inject antitumor agents into the target vasculature. These standard microcatheters rely on normal blood flow as the means by which the embolic agent moves into the tumor and systolic pressure as the packing force. However, the injection pressure is typically higher than the blood pressure and blood flow can be reversed. When this happens, the cancer agent flows in a

retrograde direction with respect to normal blood flow and away from the tumor, with a concomitant risk of delivery of the anti-cancer therapy to organs that can be damaged by these toxic agents. This situation also results in loss of an unknown amount of drug.

5

10

15

20

25

30

35

[0011] The endpoint of the above procedures is determined by physicians' visual observation and can range between fully embolized to partially embolized with the amount of dose delivered being highly variable. Retrograde and anterograde reflux, distribution, packing, quantity of dose delivered and procedure endpoint are variables that can be highly dependent on the rate and pressure of injection, the selection of the type of endpoint, the patient's systolic pressure and the operator. As such, clinical trials using TACE to treat hepatocellular carcinoma have demonstrated wide variations in tumor response. The most significant problems that occur with the current means of delivery and methods of embolization therapy include inconsistent efficacy and non-target embolization.

[0012] Using standard straight-tip catheters, non-target embolization in the retrograde direction can be caused when the pressure of injection exceeds the systolic blood pressure and the embolic agents flow backwards over the catheter and into the general circulation. Anterograde reflux and non-target embolization occurs when the embolic agents flow into distal vasculature, through arteriovenus shunting and into the venous circulation. This can easily occur because venous blood pressure on average is about 10 to 15 mmHg as compared to arterial diastolic blood pressure of about 80 mmHg.

[0013] When therapeutic agents are delivered into the vasculature of a target structure using the normal anterograde blood flow to carry the therapy to the target, injection rate and pressure of the therapy must be carefully controlled in relation to the flow volume and pressure of blood to avoid retrograde reflux of drug backward over the catheter and into the general circulation. In particular, when injecting embolic agents into the vasculature of a tumor, pressure distal to the catheter tip continues to increase as embolization progresses, causing a resistance that prevents embolic agents from filling the target vasculature and the possibility of retrograde reflux and non-target embolization. It would be desirable to eliminate this retrograde reflux, non-target embolization, and the inconsistent dosages that are delivered to targets with current state of the art procedures. It would be further desirable to eliminate the low levels of particle distribution and density throughout the target vasculature. It would be still further desirable to replace current delivery devices that are not always capable of fully isolating the target vasculature and often do not allow the operator to control pressure, flow rate and other parameters associated with therapeutic delivery.

[0014] The present state-of-the-art embolization therapy for tumors in the liver relies on high volume forward flow from the hepatic artery to deliver embolization agents into the tumor.

However; distal embolization of larger capillaries causes: (1) high intra-tumor vascular pressure, (2) high pressure in arteries feeding the tumor, (3) proximal reflux backwards over the delivery catheter, (4) increased anterograde bypass in distal hepatoenteric arteries and (5) poor filling and distribution of embolic agents in the tumor. This situation results in an uncontrollable number of particles entering the tumor and procedural high variability.

5

10

15

20

25

30

[0015] Problems with the current method of embolization therapy that cause inconsistent outcomes include: variable procedural endpoints, unknown quantity of dose delivered, reflux of embolization agents into the general circulation, anterograde bypass of embolization particles into the general circulation, non-target embolization, rising intra-tumor arterial pressures during the initial stages of embolization and catheter movement during injection. The current delivery catheters are unable to control many of the above mentioned variables, making any standardization of the current procedures difficult or impossible to achieve.

[0016] The following patents and published patent applications provide some examples of the current state of this art. U.S. Pat. No. 5,647,198 describes a catheter with a pair of spaced apart balloons that define an intra-balloon space. A lumen passes through the catheter and exits within the intra-balloon space allowing injection of drugs, emulsions, fluids and fluid/solid mixtures. A perfusion lumen or bypass extends from a location proximal to the proximal balloon and to the distal tip to allow shunting of blood past the inflated balloons. U.S. Pat. No. 5,674,198 describes a two balloon catheter that is designed for treating a solid tumor. The balloons are positioned to isolate the blood flow into the tumor and allow injection of a vaso-occlusive collagen material to block the tumor blood supply. Clifton et al. (1963) Cancer 16:444-452 describes a two balloon catheter for the treatment of lung carcinoma. The four lumen catheter includes a lumen for independent injection in the space between the balloons. Rousselot et al. (1965) JAMA 191:707-710 describes a balloon catheter device for delivering anticancer drugs into the Liver. See also U.S. Pat. No. 6,780,181; U.S. Pat. No. 6,835,189; U.S. Pat. No. 7,144,407; U.S. Pat. No.

U.S. Pat. No. 6,780,181; U.S. Pat. No. 6,835,189; U.S. Pat. No. 7,144,407; U.S. Pat. No. 7,412,285; U.S. Pat. No. 7,481,800; U.S. Pat. No. 7,645,259; U.S. Pat. No. 7,742,811; U.S. App. No. 2001/008451; U.S. App. No. 2001/0041862; U.S. App. No. 2003/008726; U.S. App. No. 2003/0114878; U.S. App. No. 2005/0267407; U.S. App. No. 2007/0137651; U.S. App. No. 2008/0208118; U.S. App. No. 2009/0182227 and U.S. App. No. 2010/0114021.

[0017] What is needed and is not provided by the prior art is a delivery system and method that enable optimization and standardization of treatment delivery, such as by delivering a known quantity of embolic agent to a prescribed target area, and elimination of non-target embolization.

5

10

15

20

25

30

35

SUMMARY OF THE DISCLOSURE

[0018] According to aspects of the present disclosure, devices and methods are provided for partial occlusion with unidirectional bypass flow, designed to be adapted to a catheter for delivery of therapeutic agents to a target site within the body. Such delivery devices may be intended for any medical purpose, but the embodiments described herein are focused on devices intended for performing transarterial delivery of therapeutic agents to a target site within the body. The entry point for the delivery catheter can be any arterial access point, typically the femoral artery located at the groin. The target can be any structure; however, of particular interest are tumors, primary or metastatic, of any organ or tissue that is accessible by a microcatheter through the arterial system. Cancers of particular interest include, but are not limited to, primary and metastatic cancers in the liver, pancreas, colon, rectum, kidney, stomach, lung, bladder, head and neck and uterus. Procedures that can benefit from the access and delivery methods and devices of the present disclosure include, but are not limited to, transarterial chemoembolization using drug eluting beads (DEB TACE), transarterial chemoembolization using Lipiodol (Lipiodol TACE), transarterial radioembolization (TARE) and transarterial embolization (TAE). Other procedures which can benefit from methods and devices of the present disclosure include direct delivery of chemotherapy or targeted drugs to the site of the cancer, the general delivery of drugs, venous or arterial embolization or other substances to specific regions of the body and drainage or aspiration of fluid or tissue. [0019] In some embodiments, microcatheter methods and devices disclosed herein create a partial occlusion (in the hepatic artery) with concomitant reduction of pressure and flow in the vascular compartment distal to the occlusion. This can: (1) eliminate proximal reflux, (2) reverse hepatoenteric arterial flow distal to the occlusion, (3) limit or stop anterograde bypass, (3) reduce flow rate into the tumor, (4) delay the onset of high intra-tumor pressure, (5) increase the extent of embolization and (6) enable a quantitative pressure measurement as an endpoint. [0020] In some embodiments, the device includes an occlusion structure adapted to the distal section of a two lumen catheter, whereby the occlusion structure has one or more channels disposed from its proximal surface to its distal surface allowing flow therethrough. The channels are configured to permit a controllable arterial or venous flow which is less than the un-occluded flow of the artery or vein, and the channel(s) may have one-way valve(s) that allow flow in only one direction. The flow rate can be any amount as compared to the normal un-occluded flow rate from 0% flow up to 100% flow, most typically from 85% flow to 95% flow. The flow channels and/or valves can be configured so as to regulate flow and/or pressure from one side of the occlusion structure to the other. The flow and/or pressure regulation can be static or variable, permitting a single continuous flow rate or multiple flow rates that are pre-calibrated or adjusted

by the operator. Of particular interest in some embodiments is arterial flow regulation to a constant flow rate that is pre-calibrated.

5

10

15

20

25

30

35

[0021] In some embodiments, the flow rate reduction results in a pressure drop in the vascular space distal to the partial occlusion. When this occurs, distal arterial side branches of the occluded artery that are flowing away from the occluded artery, and part of arterial networks, may reverse direction and begin to flow toward the occluded artery. By way of example, embolization of tumors in the right lobe of the liver are accessed by a catheter advanced through the right hepatic artery (RHA) and to the vicinity of the tumor. Typically, the catheter tip does not enter the tumor vasculature and remains proximal to the tumor and within the right hepatic artery or branch thereof. In this example, the artery feeding the tumor is typically a branch of the RHA. However, there other distal hepatoenteric arteries that branch from the RHA and flow away from the RHA and to the liver and gastrointestinal tract. In this instance, when using a standard straight catheter, injection of embolic agents from the distal tip of the catheter results in flow of embolic agents into both the tumor and side branch arteries causing anterograde bypass and non-target embolization of the liver and gastrointestinal tract, a situation that causes toxicity and complications.

[0022] In some embodiments of the present disclosure, the blood pressure drop in the arterial space distal to the partial occlusion device causes the branch hepatoenteric arteries to reverse flow and now flow toward the tumor. This situation prevents anterograde bypass, reduces non-target embolization and focuses substantially all injected drug and/or embolic agents into the tumor.

[0023] In some embodiments of the present disclosure, a partial occlusion device comprises a balloon with flow regulating channels and one-way valves which, when inflated in an artery, permits limited anterograde flow from vascular space proximal to the partial occlusion to the vascular space distal to the partial occlusion and prevents retrograde flow. So constructed, the device will significantly reduce the hepatic arterial flow into the tumor which is the tumor's main source of blood flow. In this instance, injected anti-cancer agents will flow into the tumor at a significantly slower rate and delay the onset of a high pressure in tumor vessels and retrograde blood flow toward the catheter tip. This slower filling of the tumor will improve the amount and distribution of embolic agents in the tumor. The lower pressure distal to the occlusion will cause the reversal of flow in distal hepatoenteric arteries during embolization and limit anterograde bypass, and the one-way valves will eliminate retrograde bypass of embolic agents into proximal branch arteries such as the gastroduodenal artery.

[0024] By isolating the distal arterial space that is adjacent to the tumor from the arterial blood supply, the device of the present disclosure enables pressure measurement to be used to signal a

procedural endpoint at a predetermined pressure or pressures. By way of example, the endpoint of the procedure can occur at a point when systolic pressure (120 mmHg) is first reached or at a point when systolic pressure is stabilized, however any pressure, pressure profile or algorithm can be used to determine an endpoint of the procedure. Such a measurable endpoint can contribute to standardization of the procedure and improved efficacy.

5

10

15

20

25

30

35

[0025] The occlusion structure of the device of the present disclosure may be held within a pocket within the catheter such that the outer diameter of the radially constrained occlusion structure is approximately equal to or less than the outer diameter of the catheter as described in co-pending U.S. provisional patent applications 61/821058, 61/915425 and 61/917,131. The pocket can be a longitudinal space in the catheter and can be formed as a reduction in the catheter diameter of a defined length and a depth equal to or greater than the thickness of the occlusion structure in a radially constrained configuration. Alternately, a pocket can be formed using an extension projecting distally beyond the catheter body, the distal extension having a diameter smaller than the catheter body. In this instance, the distal end of the catheter pocket is defined by the proximal end of a nose-piece. In some embodiments, the nose-piece has a diameter equal to or less than the diameter of the catheter body and is positioned over the distal extension at a defined distance from the distal end of the catheter body.

[0026] The occlusion structure of devices of the present disclosure can be advanced in a radially constrained configuration, to at least the proximity of a target within the body and then placed in its radially expanded configuration. Alternately, the device can be pre-formed in a fully expanded configuration, adapted to the distal end of a catheter and delivered to the target site. The valves are typically configured to allow proximal to distal (anterograde) bypass flow; however the opposite is possible or the valves may be absent allowing two way flow.

[0027] In some embodiments, a catheter assembly may be provided with a catheter body and an inflatable balloon. The catheter body has a proximal end, a distal end and a balloon inflation lumen. The inflatable balloon is attachable to the distal end of the catheter body. The balloon has an inner surface that at least partially defines an interior volume. The balloon is configured such that the interior volume can be in fluid communication with the inflation lumen of the catheter body to inflate the balloon. The balloon also has a proximal surface and a distal surface.

The balloon is provided with a channel that extends through the balloon. The channel is configured to provide fluid communication between the proximal surface of the balloon and the distal surface of the balloon.

[0028] In some embodiments, a device for delivering a therapeutic agent to a target site within a body is provided. The device comprises a catheter body having a proximal end, a distal end, a first axial lumen and a second axial lumen. The first axial lumen extends from the proximal end

of the catheter body to the distal end of the catheter body and provides fluid communication therebetween. The second axial lumen extends from the proximal end of the catheter body to a more distal location on the catheter body. The device further comprises a balloon radially disposed near the distal end of the catheter body. The balloon has a proximal balloon surface, a distal balloon surface, a radially constrained configuration and a radially expanded configuration. The balloon is in fluid communication with the second axial lumen and has at least one channel extending from the proximal balloon surface to the distal balloon surface, thereby providing fluid communication therebetween. The device also comprises a balloon sheath, disposed over an outer surface of the balloon. The sheath has a proximal side and a distal side. The proximal side has an opening that is in alignment with a proximal end of the balloon channel. The distal side of the sheath extends over a distal end of the balloon channel. The distal side of the sheath is capable of deflecting distally when a distally directed fluid pressure is applied through the opening in the proximal side, through the balloon channel and against a proximal surface of the distal side of the sheath, thereby allowing fluid to pass through the balloon channel and into a vascular compartment distal to the balloon and the balloon sheath. The distal side of the sheath is also capable of deflecting proximally onto the distal balloon surface when a pressure is applied to the distal side of the sheath, thereby preventing proximally directed flow in the balloon channel.

[0029] In some embodiments, a method of embolizing a tumor is provided. The method comprises advancing a device including a catheter body and a partial occlusion structure to a target tumor site within the body, and allowing an anterograde blood flow past the partial occlusion structure. The allowed anterograde blood flow is less than a blood flow that would normally be present if the partial occlusion structure were not in place. The method further comprises injecting an embolic substance from the device to allow the anterograde blood flow to carry the embolic substance into a vasculature of the tumor target. The method also comprises monitoring a real time pressure measurement in a vascular space distal to the partial occlusion structure, signaling a procedural endpoint based on the monitoring of the pressure measurement, and withdrawing the device from the body.

[0030] While aspects of the present disclosure will be described with particular reference to delivery of chemotherapeutic agents, radiotherapeutic agents, embolic agents or combinations thereof into the vasculature that supplies blood to a tumor, the same principles apply to the delivery or aspiration of a variety of materials into or from other locations, and through other luminal structures in the body.

5

10

15

20

25

#### BRIEF DESCRIPTION OF THE DRAWINGS

- [0031] The novel features of the disclosure are set forth with particularity in the claims that follow. A better understanding of the features and advantages of the present disclosure will be obtained by reference to the following detailed description that sets forth illustrative embodiments, in which the principles of the disclosure are utilized, and the accompanying drawings of which:
  - [0032] FIG. 1A, 1B and 1C illustrate embodiments of the disclosure herein;
  - [0033] FIG. 2 illustrates a distal cross sectional view of a bidirectional embodiment;
- 10 [0034] FIG. 3 illustrates a unidirectional embodiment;

- [0035] FIG. 4 illustrates a unidirectional embodiment with balloon inflation and guidewire/injection lumens;
- [0036] FIG. 5 illustrates an embodiment of the present disclosure including a distal tip and adapter;
- [0037] FIG. 6 illustrates an embodiment of the present disclosure with a bidirectional channel within the catheter;
  - [0038] FIG. 7 illustrates an embodiment of the present disclosure with a unidirectional channel;
  - [0039] FIG. 8 illustrates a device of the present disclosure with a distal tip;
  - [0040] FIG. 9A shows a linear cross section through the catheter and balloon;
- 20 [0041] FIG. 9B shows a linear cross section through the catheter and balloon and two circular bidirectional channels;
  - [0042] FIG. 9C shows a linear cross section through the catheter and balloon and three circular bidirectional channels;
  - [0043] FIG. 9D shows a linear cross section through the catheter and balloon and two circular unidirectional bypass channels with individual valves on each channel;
  - [0044] FIG. 9E shows a linear cross section through the catheter and balloon and three circular unidirectional bypass channels with individual valves on each channel;
  - [0045] FIG. 9F shows a linear cross section through the catheter and balloon and two circular unidirectional channels with a one-piece valve covering both channels;
- 30 [0046] FIG. 9G shows a linear cross section through the catheter and balloon and two circular unidirectional channels with a one-piece valve covering three channels;
  - [0047] FIG. 9H shows a linear cross section through the catheter and balloon and four bidirectional bypass channels formed from pleats in the balloon;

[0048] FIG. 9I shows a linear cross section through the catheter and balloon and four unidirectional bypass channels formed from pleats in the balloon and a one-piece valve covering all 4 channels;

- [0049] FIG. 9J shows a linear cross section through the catheter and balloon and four
- 5 bidirectional channels which radiate outward from the catheter;
  - [0050] FIG. 9K shows a linear cross section through the catheter and balloon and four unidirectional channels which radiate outward from the catheter and a one-piece valve covering all four valves;
- [0051] Fig. 9L shows a linear cross section through the catheter and balloon and four unidirectional channels which radiate outward from the catheter outward and a single cross shaped valve covering all four channels;
  - [0052] FIG. 10 shows a tumor and associated vasculature;
  - [0053] FIG. 11shows an expanded view of a tumor and its vasculature including blood flow direction;
- [0054] FIG. 12 illustrates a standard micro-catheter over a guidewire inside the tumor vasculature with injection;
  - [0055] FIG. 13 illustrates a catheter of the present disclosure over a guidewire and inside a main artery
  - [0056] FIG. 14 illustrates a catheter of the present disclosure over a guidewire and inside the branch artery;
  - [0057] FIG. 15 illustrates a catheter of the present disclosure with inflated balloon and closed valves;
  - [0058] FIG. 16 illustrates a catheter of the present disclosure with inflated balloon and open valves;
- 25 [0059] FIG. 17 illustrates a catheter of the present disclosure with inflated balloons, open valves and initiation of fluid injection;
  - [0060] FIG. 18 illustrates a catheter of the present disclosure during injection with inflated balloons and valves closed;
- [0061] FIG. 19 illustrates a catheter of the present disclosure at a period of time following the completion of injection with inflated balloon and open valves;
  - [0062] FIG. 20 shows the deflation of the balloon;

- [0063] FIG. 21 shows the withdrawal of the catheter into a main artery;
- [0064] FIG. 22 shows an embodiment of the present disclosure with a channel within the catheter and closed valve;

[0065] FIG. 23 shows an embodiment of the present disclosure with a channel within the catheter and open valve.

- [0066] FIG 24A illustrates a standard microcatheter;
- [0067] 24B, 24C, 24D and 24E illustrate a cross sectional view of a sequential construction of an embodiment of the present disclosure;
  - [0068] FIG. 25 illustrates a cross sectional view of the distal portion of one embodiment of the device according to the present disclosure;
  - [0069] FIG 26 illustrates a view of an embodiment including an occlusion balloon concealed within pocket formed between proximal and distal surfaces;
- 10 **[0070]** FIG. 27A and 27B illustrate a distal catheter with and occlusion balloon unexpanded and expanded;
  - [0071] FIG. 28A, 28B, 28C, and 28D illustrate an embodiment of the present disclosure including a two layer catheter;
  - [0072] FIG. 29 gives examples of balloons used in various embodiments of the present
- 15 disclosure;
  - [0073] FIG. 30 shows a front view of a balloon with one-way bypass channels and valves;
  - [0074] FIG. 31 shows a cross section through the balloon of Fig 30;
  - [0075] FIG. 32 illustrates a cross sectional view of a sequential construction of an embodiment of the present disclosure including a balloon pocket and integral nose cone;
- 20 **[0076]** FIG. 33 illustrates a cross sectional view of a sequential construction of an embodiment of the present disclosure including a balloon pocket and separate nose cone;
  - [0077] FIG. 34 illustrates an embodiment of the present disclosure with two balloons;
  - [0078] FIG. 35 illustrates an embodiment of the present disclosure including a balloon with valves;
- 25 [0079] FIG. 36 illustrates an embodiment of the present disclosure with pressure sensors;
  - [0080] FIG 37A, 37B, 37C and 37D illustrate an embodiment of the present disclosure with a balloon or balloons placed on the circumference of the catheter;
  - [0081] FIG. 38A and 38B illustrate a tumor with vascular anatomy and embolization using a standard straight nose catheter;
- 30 [0082] FIG. 39A and 39B illustrate a tumor with vascular anatomy and embolization using a balloon including channels and valves;
  - [0083] FIG 40A, 40B, 40C and 40D illustrate a tumor with vascular anatomy and embolization using an occlusion balloon;
  - [0084] FIG. 41 illustrates constructions of embodiments of the present disclosure;
- 35 [0085] FIG. 42 illustrates a two layer occlusion structure with unidirectional flow;

- [0086] FIG. 43 illustrates a prototype of the occlusion structure of FIG 42;
- [0087] FIG. 44 illustrates an embodiment of the present disclosure including a catheter with a pocket;
- [0088] FIG. 45 illustrates embodiments of the present disclosure for complete occlusion;
- 5 [0089] FIG. 46 illustrates an embodiment with bidirectional and unidirectional channels;
  - [0090] FIG 47 illustrates valve constructions of an embodiment of a unidirectional occlusion structure of the present disclosure;
  - [0091] FIG. 48 shows a prototype micro-valve;
  - [0092] FIG. 49 shows an alternate embodiment of an occlusion structure;
- 10 [0093] FIG. 50 illustrates a method of operation of one embodiment of a unidirectional occlusion structure of the present disclosure;
  - [0094] FIG. 51 illustrates a method for delivery of embolic particles using a device of the present disclosure;
  - [0095] FIG. 52 illustrates a partial occlusion balloon with channel and valve;
- 15 [0096] FIG. 53 illustrates a side view of constrained occlusion balloon in catheter pocket;
  - [0097] FIG. 54 illustrates a distal end construction including pocket, constrained balloon and expanded partial occlusion balloon with channels and valves in closed position;
  - [0098] FIG. 55 illustrates a serial construction of a full length catheter device;
  - [0099] FIG. 56 illustrates liver vasculature and associated tumor vasculature;
- 20 [0100] FIG. 57 illustrates a tumor embolization method for a standard catheter;
  - [0101] FIG. 58 illustrates a tumor embolization method for the catheter of the present disclosure;
  - [0102] FIGS. 59A, 59B, 59C and 59D illustrate fabrication steps of the distal end of an occlusion catheter according to aspects of the present disclosure;
  - [0103] FIG. 60 illustrates the completed distal end of the occlusion catheter of FIGS. 59A 59D being introduced into small branches of a vascular system;
  - [0104] FIG. 61A shows an entire occlusion catheter constructed according to principles of the present disclosure; and
  - [0105] FIG. 61B shows details of construction features on the distal end of the occlusion catheter of FIG. 61A.

30

35

25

### **DETAILED DESCRIPTION**

[0106] The device of the present disclosure allows improved distribution of anti-cancer agents into target tumor vasculature by reducing arterial flow and pressure during drug and/or embolic agent injection. The present device reduces toxicity and complications by eliminating reflux of embolic materials and/or anti-cancer agents into proximal arterial branches and reduces or

eliminates anterograde bypass of embolic materials and/or anticancer agents into distal arterial branches. Further, the present disclosure enables pressure measurement as a means to a quantitative endpoint of the procedure. Such a device can improve efficacy and reproducibility of the technique and reduce complications.

5

10

15

20

25

30

35

[0107] Referring to FIG. 1A, a longitudinal cross section of device 2 of the present disclosure is shown with catheter body 4, distal tip 6, balloon 8 (unexpanded configuration), balloon inflation tube 10, guidewire and injection tube 12 and fittings 14. Catheter body 4 can have a length of 10 cm to 400 cm, typically 60 cm to 250 cm and a diameter of 0.25 mm to 5 mm, typically 0.5 mm to 1.5 mm. Device 2 may or may not include a distal tip 6, the distal tip having a length of 1 mm to 50 mm, more typically from 5 mm to 30 mm. The balloon inflation tube 10, positioned at the proximal end of catheter body 4, is connected to, and in fluid communication with a balloon inflation lumen that runs longitudinally through the length of catheter body 4 and terminates at, and is in fluid communication with balloon 8. The guidewire and injection tube 12, positioned at the proximal end of catheter body 4, is connected to, and in fluid communication with a lumen that runs longitudinally through the length of catheter body 4 and terminates at the distal end or distal tip of catheter body 4, thereby allowing a guidewire to enter through fitting 14 and exit through the distal end of device 2 through catheter body 4. Fittings 14 are connected to each of balloon inflation tube 10 and guidewire and injection tube 12 and can connect to a syringe, inflation device or any other device or means to inject air, gas, fluid, suspensions, emulsions, contrast, therapeutic agents, embolic agents or any other material capable of being injected through balloon tube 10 or guidewire tube 12 and longitudinal lumens that run to the balloon or distal end of device 2.

[0108] Referring to FIG 1B, a longitudinal cross section of a first embodiment of the present disclosure is shown, with device 2, balloon 8 (in the expanded configuration) having valve 16 in the open position and valve 18, in the closed position. In this embodiment flow channels 20 and 22 are constructed through balloon 8. Valves 16 and 18 allow fluid to flow in only one direction. Balloon 8 has a proximal side 9 and a distal side 11. By way of example, if fluid pressure is higher on the proximal side of balloon 8 and lower on the distal side of balloon 8, both valves 16 & 18 will open in response to the pressure difference and allow fluid to flow distally through the valves. If the pressure is higher on the distal side of balloon 8, valves 16 and 18 will close and prevent fluid from flowing proximally. Alternately, the valves can be position or constructed so that fluid can pass proximally and be prevented from flowing in the distal direction. Valves 16 and 18 are shown as a simple "flap" type valve, however, they can be any type of valve, such as a diaphragm that open and close in response to a pressure differential. Balloon 8 is shown with two channels and two valves; however there can be 1, 2, 3 or more channels and/or valves.

Device 2 of this embodiment may include channels and may or may not include valves. If valves are not included, a bidirectional flow will result.

5

10

15

20

25

30

35

[0109] Referring to FIG. 1C, a longitudinal cross section of another embodiment of the present disclosure is shown with device 24 and channel 26 running through and within catheter body 28.

Balloon 30 has a proximal side 13 and a distal side 15. Channel 26 extends from the proximal side of balloon 30 to the distal side of balloon 30. A valve 32 is illustrated over channel 26 on the distal side of balloon 30, however, if desired the valve can be positioned on the proximal side of channel 26 and balloon 30. The function and operation of valve 32 of this embodiment of the present disclosure is identical to that presented in FIG 1A and FIG 1B. As in this embodiment, if valve 32 is not included, a bidirectional flow will result.

[0110] FIG. 2 illustrates an exemplary embodiment of a longitudinal cross section of the distal end of device 2 with catheter body 4, balloon inflation lumen 34, balloon 8, and channel 38. Balloon inflation lumen 34 extends from the proximal end of catheter body 4 and exits at balloon 8. In this case, a valve in balloon 8 is not included and a bidirectional flow will result. The optimal balloon inflation lumen diameter is 0.1 mm to 0.5 mm; however this lumen can be in the range of 0.25 mm to 1 mm.

[0111] FIG. 3 illustrates an example of a longitudinal cross section of the distal end of device 2 with catheter body 4, balloon inflation lumen 34, balloon 8, channel 38 and valve 16. Valve 16 is shown over the distal opening of channel 38 in the closed position, however if pressure is applied to the proximal valve surface through channel 38, the valve will allow fluid to pass distally. The valve 16 will prevent proximal flow. The valve can be positioned at the proximal or distal opening or anywhere within the cannel. Location and configuration of the valve will determine flow direction.

[0112] Referring to FIG. 4, a longitudinal cross section of the distal end of device 2 is shown with catheter body 4, balloon inflation lumen 34, balloon 8, channel 38 and guidewire/injection lumen 40. Guidewire/injection lumen 40 extends from the proximal end of catheter body 4 and exits at the distal end of catheter body 4 or distal tip 6. The optimal guidewire/injection lumen diameter is 0.1 to 1.0 mm; however, this lumen can be in the range form 0.025 mm to 2 mm.

[0113] Referring to FIG. 5, an example of a longitudinal cross section of the distal end of device 2 is shown with catheter body 4, balloon inflation lumen 34, balloon 8, channel 38, guidewire/injection lumen 40, adapter 42, balloon pocket 44 and distal tip 6. Distal tip 6 has an optimal diameter of 0.3 mm to 1.3 mm; however, distal tip 6 can range from 0.1 mm to 4 mm.

Adapter 42 is adapted to create a smooth transition from the distal tip 6 to the catheter body 4. By way of example, if the distal tip 6 is 1 mm in diameter and the catheter body 4 is 2 mm in diameter, the adapter will taper from a diameter of 1 mm at its distal most point to 2 mm at its

proximal most point to create a smooth transition from the smaller diameter distal tip to the larger diameter catheter body. As shown in FIG. 5, adapter 42 is positioned on the distal tip 6 at a location that is distal to distal end of catheter body 4, such that a balloon pocket 44 is formed between the distal end of catheter body 4 and the proximal end of adapter 42. Balloon pocket 44 holds unexpanded balloon 8 such that its unexpanded profile is minimized. Optimally balloon 8 would conform to an outer diameter that is equal to the diameter of the catheter body 4. By way of example, if the distal tip diameter is 1 mm and the catheter body is 2 mm, a balloon pocket is formed that is 0.5 mm deep. If balloon 8 is no more than 0.5 mm thick when unexpanded, it will lie equal to or below the surface of catheter body 4. This allows facilitation of the movement of the caterer within the artery or vein.

5

10

15

20

25

30

35

[0114] Referring to FIG. 6, a longitudinal cross section of the distal end of device 24 is shown with catheter body 28, balloon inflation lumen 50, guidewire/injection lumen 52, balloon 30, and channel 26 with proximal port 54 and distal port 56. Channel 26 passes through and within catheter body 28 and extends from the proximal side of balloon 30 to the distal side of balloon 30. Port 54, at the proximal end of channel 26 is in fluid communication with the outside of the catheter body that is proximal to balloon 30 and port 56, at the distal end of channel 26, is in fluid communication with the area outside of catheter body 28 that is distal to the balloon 30. In this case, the flow through channel 26 is bidirectional. The optimal channel diameter is 0.1 mm to 1 mm; however this channel can be in the range of 0.05 mm to 2 mm.

[0115] FIG 7 illustrates an exemplary embodiment of a longitudinal cross section of the distal end of device 24 with catheter body 28, balloon inflation lumen 50, guidewire/injection lumen 52, balloon 30, valve 32 and channel 26 with proximal port 54 and distal port 56. Channel 26 passes through and within catheter body 28 and extends from the proximal side of balloon 30 to the distal side of balloon 30. Port 54, at the proximal end of channel 26 is in fluid communication with the outside of the catheter body that is proximal to balloon 30 and port 56, at the distal end of channel 26, is in fluid communication with the area outside of catheter body 28 that is distal to the balloon 30. Valve 32, is shown at the opening of port 56 of channel 26. Valve 32 allows flow in the distal direction and prevents flow in the proximal direction. In this case, the flow through channel 26 is unidirectional. The optimal channel diameter is 0.1 mm to 1 mm; however this channel can be in the range of 0.05 mm to 2 mm. The valve can be positioned at the proximal or distal opening or anywhere within the cannel. Location and configuration of the valve will determine flow direction.

[0116] Referring to FIG 8, a longitudinal cross section of the distal end of device 24 is shown with catheter body 28, balloon inflation lumen 50, guidewire/injection lumen 52, balloon 30, valve 32, channel 26 and distal tip 6.

5

10

15

20

25

30

[0117] Referring to FIG. 9, linear cross sections through catheter body 4 and balloon 8 are shown. Figure 9 illustrates examples of flow channels through a balloon positioned on a catheter, however, various alternatives, modifications, and equivalents may be used. FIG 9A shows catheter 4 and balloon 8 without a channel. FIG 9B shows catheter 4 and balloon 8 with two channels 38 through balloon 8. In this case the flow is bidirectional. FIG 9C shows a catheter and balloon with three bidirectional cannels. FIG 9D shows a catheter 4 with a balloon 8 with two channels, each with a one-way valve. In this case the channel flow in each channel is unidirectional. FIG 9E shows catheter 4 with balloon 8 and three channels 38, each with an independent valve 38. The flow is unidirectional. FIG 9F is a catheter and balloon with two channels and a single circumferential valve 33 that covers both channels 38 and allows flow in only 1 direction. FIG 9G shows a catheter 4 and balloon 8 with three channels and a single circumferential valve 33 that covers all three channels. FIG 9H shows a catheter 5, and four bidirectional channels 39 that are formed by pleating balloon 36 from the outer circumference of the expanded balloon, inward toward the central catheter and securing the inner apex of the balloon to the catheter, thereby forming triangular channels which radiate outward from the central catheter, forming a "V" shape. The lower point of the 'V" is positioned at the catheter surface and the open end of the "V" is positioned at the outer circumference of the circle defined by the largest diameter of the inflated balloon. FIG 9I shows four channels 39, with a single circumferential valve 35 covering all four triangular channels. Although the figure shows four channels, the device of this disclosure can have one, two, three, four or any number of channels. FIG 9J shows a catheter 7 and a balloon 39 with four channels 37 that are formed by pleating the balloon 39 from the inner catheter 7 toward the outer circumference of the expanded balloon 39. FIG 9K shows four channels 37as in FIG 9J with a single circumferential valve 39 that covers all four channels 37. FIG 9L shows four channels 37 as in FIG 9J with a one piece cross-shaped flap valve 41.

[0118] While the above is a complete description of exemplary embodiments of the present disclosure, various alternatives, modifications, and equivalents may be used. Therefore, the above description should not be taken as limiting the scope of the disclosure, which is defined by the appended claims and the claims in any subsequent applications claiming priority hereto.

[0119] FIG. 10 illustrates a tumor and its associated vasculature with tumor 60, main artery 62, side branch artery 64, tumor artery 66, tumor capillary 68 and anterograde arterial flow direction illustrated by arrows 70.

[0120] FIG. 11 is an expanded view of FIG. 10 with tumor 60, main artery 62, side branch artery 64, tumor artery 66 and anterograde arterial flow direction illustrated by arrows 70.

[0121] FIG. 12 illustrates injection of fluid 76 using a standard micro-catheter 72 with tumor 60, main artery 62, side branch artery 64, tumor artery 66, guidewire 74 and retrograde arterial flow direction illustrated by arrows 70. In this instance, injection pressure and flow volume of the fluid 76 that may contain anti-cancer drugs, radioembolic substances, chemoembolic substances, embolic agents or the like, through microcatheter 72 is higher than the tumor vasculature can accept causing a reversal of fluid flow and blood flow in tumor artery 66, side branch artery 64 and main artery 62. This retrograde flow causes the injected fluid 76 to enter the main artery, flowing in both directions and into the general circulation resulting in the injected fluid traveling to non-target tissues and organs. This unintended delivery of fluid 76 to non-target sites is undesirable and must be avoided since it can cause serious complications. The present 10 disclosure solves this problem by preventing reflux and associated non-target delivery of fluid 76.

5

15

20

25

30

[0122] FIG. 13 illustrates catheter 4 of the present disclosure entering the main artery with balloon 8 and distal tip 6. Although the entry point from outside the body is typically through the femoral artery at the groin, any artery or vein from any location on the body can be used for access provided that it creates a pathway to the target vasculature.

[0123] FIG. 14 illustrates catheter 4 with balloon 8, of the present disclosure, following guidewire 74 into side branch artery 64.

[0124] FIG. 15 illustrates catheter 4, inside the branch artery 64 with balloon 8 in the expanded configuration, channels 38 and valves 16. Valves 16 are illustrated in the closed position immediately following the inflation of balloon 8.

[0125] FIG. 16 illustrates catheter 4, inside the branch artery 64 with balloon 8 in the expanded configuration, channels 38 and valves 16. Valves 16 are illustrated in the open position since anterograde blood flow as indicated by arrows 70 and associated blood pressure causes these valves to open and allow the blood to continue to flow in the anterograde direction and into the tumor vasculature.

[0126] FIG. 17 illustrates the initial injection of fluid 76 into side branch artery 64 through catheter 4, channels 38 and open valves 16. When the injection is initiated, the anterograde blood flow carries the injection fluid 76 into the tumor vasculature including tumor arteries 66, and capillaries 68.

[0127] FIG. 18 illustrates injection of fluid 76 at a point when fluid pressure increases within the tumor vasculature and concomitant retrograde arterial blood flow and injection fluid flow in the direction as illustrated by arrows 70. Shown in this figure are catheter 4 of the present disclosure with tumor 60, main artery 62, side branch artery 64, tumor artery 66, and guidewire 74.

Injection fluid 76, may contain anti-cancer drugs, radioembolic substances, chemoembolic 35

substances, embolic agents or the like, which can cause serious complications if delivered to non-target sites. In this case, the retrograde pressure causes valves 16 to close and prevents the reflux of injection fluid into the general circulation, thereby preventing complications associated with delivery of injection fluid to non-target sites.

- [0128] FIG. 19 illustrates a point in time following the completion of fluid injection. At this point, the pressure in the vasculature that is the distal to balloon 8, including side branch 64 and tumor artery 66, is reduced below normal blood pressure due to the gradual uptake of the injected fluid into the tumor vasculature. The blood pressure on the proximal surface of balloon 8 and valves 16 cause them to open allowing anterograde blood flow to be reestablished. When this occurs, the excess fluid 76 distal to balloon 8 and within the side branch artery 64 and tumor vasculature, including tumor artery 66 and tumor capillaries 68, is flushed forward and up into the tumor vasculature, thereby enabling delivery of the entire fluid dose and eliminating fluid reflux and associated complications.
  - [0129] FIG. 20 illustrates the deflation of balloon 8 on catheter 4.
- 15 [0130] FIG. 21 illustrates the withdrawal of catheter 4 into the main artery 62.
  - [0131] FIG. 22 illustrates another embodiment of the present disclosure as described in FIGS. 6, 7 and 8. In this case valve 32 on the distal end of channel 26 of catheter 28 is in the closed position.
  - [0132] FIG 23 illustrates an embodiment of FIG. 22 with valve 32 in the open position.
- [0133] A Method, according to the present disclosure is illustrated by FIGS. 13 through 21; the method applies to both the embodiment illustrated in FIGS. 1B, 2, 3, 4 and 5 and the embodiment illustrated in FIGS. 1C, 6, 7, 8, 22 and 23.
  - [0134] Referring to FIG 24A, a longitudinal cross section of a standard single lumen straight tip catheter 101, having a proximal and distal end, is shown with catheter body 102, and hub 105.
- Hub 105, positioned at the proximal end, further comprises guidewire/injection lumen 110, in fluid communication with a catheter lumen longitudinally oriented and extending from hub 105 and exiting at the distal end of the catheter body 102. The proximal hub connects to a syringe or other means to inject fluids via a luer fitting, thereby allowing injection of a fluid through the longitudinal lumen and exit at the distal end of catheter body 102.
- 30 **[0135]** Referring to FIG. 24B through 24E, longitudinal cross-sections of a sequential assembly of a preferred embodiment of the present disclosure is shown. Referring to FIG 24B, device 103 is shown, having a proximal and a distal end, catheter body 102, catheter extension 104 and hub 106. Hub 106 further comprises handle 109, guidewire/injection luer fitting 110 and balloon fill luer fitting 112. Luer fitting 110 is in fluid communication with a first longitudinal
- 35 guidewire/injection lumen of catheter body 102, extending to the distal end of catheter extension

104, and luer fitting 112, in fluid communication with a second longitudinal balloon fill lumen of catheter body 102, extending to a balloon fill port located near the distal end of catheter body 102. FIG 24C further comprises nose cone 114, positioned on catheter extension 104, forming balloon pocket 116 disposed between the distal end of catheter body 102 and nose cone 114.

5

10

15

25

30

35

- Further, a portion of catheter extension 104 can, if desired, extend distal to nose cone 114, thereby forming distal tip 118. FIG 24C, further illustrates occlusion balloon 120 in a radially compressed configuration and FIG 24E illustrating balloon 120 in a radially expanded configuration. Hubs 106 can be constructed from styrene, polyurethane, polypropylene, lipid resistant polycarbonate, polycarbonate, Pebax (polyether block amide), of any durometer, or any convenient material and can have any configuration, including, but not limited to, a solid structure comprising two lumens or tubular extensions of the lumens of catheter body 102, provided that they are in fluid communication as described above. Catheter body 102 can be formed from any plastic or thermoplastic material including polyurethane, PTFE, polyimide, polypropylene, Pebax or the like, and can comprise a single section or multiple sections of different diameter, durometer, braid or coil reinforcement or any convenient construction with a diameter of between 1Fr and 10 Fr more typically of 2 Fr to 5 Fr. Catheter extension 104 can
- have a diameter of 0.5 Fr to 5 Fr, more typically of 1Fr to 3 Fr and can be absent or can be of any length, typically 2 mm to 30 mm, more typically from 5 mm to 20 mm. If the catheter extension 104 extends beyond nose cone 114, the section distal to the nose cone forms the distal tip 118.

  20 Distal tip 118 is advantageous when injecting deep into the tumor vasculature is desired and will also help tracking of device 103 over a guidewire around sharp corners and through a tortuous vasculature path. Nose cone 114 can be made from any polymer or metal or can be formed from a radiopaque marker band. Balloon pocket 116 can be of any length between 2 mm and 50 mm, more typically between 5 mm and 20 mm. Occlusion balloon 120 has a longitudinal length of 1

mm to 30 mm, more typically of 2 mm to 10 mm and a diameter of 1 mm to 50 mm, typically from 2 mm to 10 mm and can be composed of silicone, polyurethane, polyethylene, PET (polyethylene terephthalate), nylon or the like and can be of any configuration or of any length or shape and can be glued, chemically bonded, heat bonded, RF welded, sonically fused, compressed or crimped under a collar to catheter 102 or catheter extension 104.

[0136] Referring to FIG 25A, a distal section of a device 128 of a preferred embodiment of the present disclosure is shown and includes catheter body 102, catheter extension 104, nose cone 114, balloon pocket 116, balloon fill lumen 124, guidewire/injection lumen 126 and radially compressed balloon 120. Referring to FIG 25B, balloon 120 is shown in its radially expanded configuration. Balloon fill lumen 124 can be of any convenient shape including but not limited to round, semicircular, or crescent or any shape, typically optimized to provide maximum area

and flow rate. Guidewire/injection lumen 126 is typically round, having a diameter of 0.005" to 0.1", more typically from 0.01" to 0.05"; however, it can be of any desirable shape.

[0137] Referring to FIG 26A, the distal end 133 of an embodiment of the present disclosure is shown with catheter body 102, catheter extension 104, nose cone 114, balloon pocket 116, balloon fill lumen 124, guidewire/injection lumen 126 and radially compressed balloon 120. In

5

10

15

20

25

30

35

this instance, the balloon pocket 116 is formed between a proximal collar 130 and a distal collar 114, tapered forward thereby forming a nose cone. The balloon bonding tails 115 can be bonded within the pocket or compressed or bonded under collars 114 and 130. Distal collar 130 can comprise a metal, such as a radiopaque marker band or a plastic such as heat shrink tubing and

can be 1 mm to 20 mm in length, more typically from 2 mm to 10 mm. Balloon fill lumen 124 is shown traveling under balloon pocket 116 and ending at its distal end. Guidewire/injection lumen 126 is shown traveling longitudinally through catheter 102 and catheter extension 104, ending at the distal end of the catheter. Balloon 120 is shown tucked into pocket 116 with outer

diameter substantially no larger than the outer diameter of catheter body 102. Figure 26B shows

the same construction as FIG 26A with balloon 120 in its radially expanded configuration.

**[0138]** Referring to FIG 27A, a view of distal section 133 is shown and further illustrates that the profile of a preferred embodiment of the present disclosure, including the radially compressed balloon 120, the distal collar 114 and the proximal band 130, have an outer diameter equal to or less than that of catheter body 102. FIG 27B shows the same construction as FIG 27A with balloon 120 expanded from pocket 116 and between collars 114 and 130.

[0139] Referring to FIG 28A, a distal section of an alternate embodiment 135 of the present disclosure is shown with outer catheter 119, inner catheter 127, nose cone 125, radially constrained balloon 131 and catheter channels 123. The outer catheter 119 is adapted over inner catheter 121, the catheters configured to provide a radially distributed space between inner and outer catheters extending longitudinally along the length of device 135. Outer and inner catheters can have a length of 10 cm to 250 cm, more typically 50 cm to 150 cm and a diameter of between 0.5 Fr and 10 Fr more typically of 1 Fr to 5 Fr. Inner catheter 127 can have a length less than, equal to, or longer than the outer catheter 119, however in the present figure, inner catheter 127 is shown to be longer than outer catheter 119, its distal end forming the catheter extension 127. Nose cone 125 is disposed along the distal extension of inner catheter 127 at some distance from the distal end of outer catheter 119, the distance being 2 mm to 50 mm, more typically between 5 mm and 20 mm. The balloon pocket is formed between the distal end of catheter 119 and the proximal end of nose cone 125. FIG 28B shows an end view of outer catheter 119, disposed over inner catheter 127, with radially configured channels 123 and stand-

offs 127 disposed between outer catheter 119 and inner catheter 127. Four channels are

illustrated, however device 135 can have 0, 1, 2, 3, 4 or any number of channels and stand-offs, the stand-offs defining the outer edges of the channels 123 and can be formed on either the inner or outer catheter with a height limited only by the diameter of the inner and outer catheters and space there between. Although stand-offs are shown, they are not required, provided that the inner catheter OD is smaller than the outer catheter ID, thereby forming a space between the 5 inner and outer catheters which allow fluid to flow longitudinally along device 135. Device 135 can comprise single layer inner and outer catheters or one or both can have multiple layers. In a preferred embodiment, outer catheter 19 is a three layer construction with an outer Pebax layer, a central polyimide layer including reinforcement such as a coil or braid and an inner Teflon layer. Inner catheter 127 is a single layer of low friction tubing, or tubing of similar construction to that 10 described for the outer catheter 119. FIG 28C shows a longitudinal view of device 135 with outer catheter 119, unexpanded balloon 131, nose cone 125 and a catheter extension of inner catheter 127. Balloon 131 is shown tucked within a pocket formed between the distal end of catheter 119 and the proximal end of nosecone 125. FIG 28D shows a longitudinal cross section of device 135, showing balloon inflation channel 123 disposed between outer catheter 119 and 15 inner catheter 121. In this instance, the distal end of balloon 131 is shown inserted into the proximal end of nose cone 125; however both proximal and distal balloon tails can be bonded directly to inner catheter 127, reflowed into catheter 119 or nose cone 125 or by any means, provided that the balloon tails are positioned approximately below the outer diameter of catheter 119. 20

[0140] Referring to FIG 29A though FIG 29E, examples of balloon configurations that may be used in the device of the present disclosure are shown which can be compliant or noncompliant, dilation or occlusion and can be made from any material including, but not limited to, silicone, polyurethane, polyethylene, PET (polyethylene terephthalate) and nylon.

25 **[0141]** Referring to FIG 30, a surface view of balloon 160 is shown with one-way bypass channels 162 and valves 164, the balloon is described in detail in co-pending patent application number 61/821,058.

30

[0142] Balloon 160 and valves 164 allow flow from the compartment proximal to the proximal surface of balloon 160 to the compartment distal to the distal surface of balloon 160 (anterograde flow) and prevents flow from the compartment distal to the distal surface of balloon 160 to the compartment proximal to the proximal surface of balloon 160 (retrograde flow). Balloon 160 can be disposed on the catheter of the present disclosure and held within a balloon pocket as illustrated in FIGS 24-29 and enable anterograde injection of therapeutic agents from within an artery and into a target while maintaining normal (anterograde) blood flow through channels 162

of balloon 160 and prevent retrograde flow (reflux) of therapeutic agents backward over the catheter, even when pressure distal to balloon 160 is elevated above systolic.

- [0143] Referring to FIG 31, a cross section of balloon 160 is shown with channels 162 and microvalves 164, positioned within channels 162.
- [0144] FIG 32 shows an example of a sequential assembly of an embodiment of the present disclosure. Referring to FIG 32A, catheter 170 of device 168 is shown. FIG 32B illustrates a first step in the construction of device 168 whereby a balloon pocket 172 is formed about a distal section of catheter 170 and a second step, as in FIG 32C, whereby a rounded distal end 174 of catheter 170 is formed and a third step as in FIG 32D whereby a balloon 176, with bonding tails 178 is disposed within pocket 172 of catheter 170 and a fourth optional step whereby catheter 170 or other material is reflowed at position 180 over balloon tails 178. FIG 32F illustrates balloon 176 in its radially expanded configuration with balloon pocket 172 and tails 178 bonded
  - [0145] Referring to FIG 33, three alternate embodiments of the present disclosure are illustrated.
- FIG 10A shows device 173 with catheter 181, balloon pocket 177, radially constrained balloon 178, catheter extension 171 and nose cone 182. FIG 33B shows device 173 with radially expanded balloon 178 and bonding tails 183 bonded within balloon pocket 177. FIG 33C illustrates device 175 with nosecone 182 and proximal bonding tail 191 reflowed into catheter 181 at position 179 and the distal balloon bonding tail 193 reflowed into or under nose cone 182

in pocket 172 without being covered by reflow or other means.

20

- to catheter extension 171 at location 184. FIG 33D illustrates device 179 with catheter 181, balloon 178, nose cone 182 and collar 186. Balloon 178 has a proximal tail 191 positioned under collar 186 and distal balloon tail 193 reflowed or bonded to catheter extension 171 and under nose cone 182 or into nose cone 182. FIG 33E shows device 179 with balloon 178 expanded from within the balloon pocket formed between collar 186 and nose cone 182.
- 25 [0146] FIG 34 shows yet another embodiment of the present disclosure with two balloons 188 and 189, catheter 183, balloon pockets 194 and 190, reflow areas 192 and nose cone 189.

  Although the example of FIG 34 shows both balloon 188 and 189 positioned within pockets 194 and 190, only one balloon need be positioned within a pocket.
  - [0147] FIG 35 shows still another embodiment of the present disclosure with balloon 196 containing channels 195 and 197, valve 198 in the closed orientation, valve 199 in the open orientation, collar 183, nose cone 1102 and reflow area 1100. Although valve 198 is shown closed and valve 199 is shown open, they will typically act in unison and all either be simultaneously open or closed.
- [0148] FIG 36 shows still another embodiment of the present disclosure which includes two pressure sensors, positioned distal and proximal to balloon 1110, although a single pressure

sensor positioned either distal or proximal to the balloon can be used. These pressure sensors can be used to monitor and, in conjunction with a syringe, control injection pressure either manually of by an automated means. Alternately pressure proximal or distal to the occlusion balloon can be measured through the catheter using an external pressure gauge (1113), the distal pressure being measured via the guidewire/injection lumen 1105 or any other catheter lumen or other tube. The pressure gauge can be connected to a pump, via a processor, allowing the pump to achieve a defined pressure or be programed to a specific set of pressures, volumes and/or flow rate as a function of time.

5

10

15

20

25

30

35

[0149] Referring to FIG 37, four embodiments of balloon configurations are shown. FIG 37A, shows device 1115 with balloon 1117 and catheter 1116. Balloon 117 in a radially expanded configuration, occupies only part of the circumference of catheter 1116. FIG 37B illustrates device 1118 with catheter 1119 and balloons 1121 whereby the four balloons 1121, in radially expanded configurations are arranged circumferentially about catheter 1119, each occupying a part of the overall outer circumference of catheter 1119. FIG 37C illustrates device 1123, with catheter 1125 and balloon 1127 in a radially expanded configuration, whereby balloon 1127, in a radially constrained configuration is positioned within a pocket of catheter 1125 and the radially outermost part of balloons 1127 is positioned approximately at or below the outer diameter of catheter 1125. FIG 37D illustrates device 1131 with catheter 1135 and balloons 1137 in a radially expanded configuration, whereby balloons 1137, in a radially constrained configuration are positioned within a pocket of catheter 1135 and the radially outermost part of balloons 1137 are approximately positioned at or below the outer diameter of catheter 1135.

[0150] Referring to FIG 38, an anatomical structure 1120 is shown with main artery 1122, right artery 1124, left artery 1126, right capillaries 1128, left capillary 1129, arterial side branch 1136, vein 1130, arteriovenus shunt 1132, tumor 1133, blood flow directional arrows 1134, standard straight tip catheter 1138, and embolization particles 1125. FIG 38A illustrates the beginning of a transarterial embolization (TAE) procedure wherein the embolization particles 1125 are exiting the distal end of catheter 1138 and are carried by forward (antrograde) blood flow into tumor 1133 in a delivery method that is completely mediated by blood flow and normal blood pressure (flow mediated delivery). Capillary beds 1128 and 1129 of tumor 1133 begin to fill with embolic particles 1125 and arteriovenus shunt 1132 carries particles into vein 1130 causing anterograde reflux and non-target embolization. The flow through the areteriovenous shunt 1132 is rapid since the arterial pressure is significantly higher than venous pressure. Referring to FIG 38B, continued injection of particles 1125 from the distal end of standard straight tip catheter 1138 results in the packing of particles and embolization of the distal ends of capillary beds 1128 and 1129. Distal capillary embolization causes the flow through arteriovenous shunt 1132 to stop

5

10

15

20

25

30

35

and pressure to build in left artery 1126. As embolization progresses, the back pressure in artery 1126 continues to rise until embolic particles reflux in the retrograde direction 1142 causing nontarget embolization of the right artery 1124, arterial side branch 1136 and main artery 1122. This situation can cause non-target embolization, loss of an unknown amount of particles, delivery of an unknown and irreproducible dose and non-optimal distribution of embolic particles in the tumor vasculature. In this instance, both anterograde and retrograded reflux can occur. [0151] Referring to FIG 39, anatomical structure 1120 is shown as in FIG 38. In this instance, a balloon 1141, with channels 1143 and one-way valves (FIGS 30 and 31) is positioned about the distal end of catheter 1139. Balloon 1141, so constructed, will allow only anterograde (normal) flow and prohibit retrograde flow. Referring to FIG 39A, balloon 1141 is shown in its radially expanded configuration and blood is flowing through balloon channels 1143 as indicated by blood flow arrow 1134 and into the vasculature of tumor 1133. Embolic particles 1125 are released from the distal end of catheter 1139 and carried forward by blood flow into capillaries 1128 and 1129. Capillary beds 1128 and 1129 of tumor 1133 begin to fill with embolic particles 1125 and arteriovenus shunt 1132 carries particles into vein 1130 causing anterograde reflux and non-target embolization. The flow through the areteriovenous shunt 1132 is rapid since the arterial pressure is significantly higher than venous pressure. Referring to FIG 39B, continued injection of particles 1125 from the distal end of balloon catheter 1139 results in the packing of particles and embolization of the distal ends of capillary beds 1128 and 1129. Distal capillary embolization causes the flow through arteriovenous shunt 1132 to stop and pressure to build in left artery 1126. As embolization progresses, the pressure in artery 1126 continues to rise, however the valves of balloon 1141 close and prohibit retrograde reflux. In this instance, continued injection will increase the packing pressure of particles 1125 and can increase packing density and increase flow into distal locations in the margins of a tumor or other structure thereby improving particle distribution throughout the target vasculature. As part of the present method, pressure distal to the balloon can be regulated between systolic and any pressure above systolic, provided that it is within a range that is safe for the patient. By way of example, injection pressure can be low at the onset of the embolization procedure and increased at some point thereafter to a pressure greater than systolic. Such a point may, for example, be chosen to coincide with the stoppage of flow through arteriovenous shunt 1132. This method may improve particle distribution and packing. Alternately, the injection pressure through catheter 1139 can be high at the onset, thereby forcing particles rapidly into the distal section of capillaries 1128 and 1129 and hasten embolization of arteriovenous shunt 1132, thereby reducing anterograde reflux. Alternately, according to this method, a low to high pressure gradient or a high to low pressure gradient can be used. The aim for the use of a pressure mediated delivery of particles is

to optimize for a low level of anterograde reflux, substantial elimination of retrograde reflux, high particle distribution and high particle density. A pressure sensor as in FIG 36 can be used on the proximal and/or distal side of balloon 1142 to monitor pressure and enable a selection of a procedural end point based on a definitive pressure reading.

[0152] Referring to FIG 40, anatomical structure 1120 is shown as in FIG 38. In this instance, an occlusion balloon 1142 is positioned about the distal end of catheter 1140. Referring to FIG 40A, balloon 1142 of catheter 1140 is shown in a radially expanded configuration. Since expanded balloon 1142 completely occludes artery 1126, all arteries and capillaries distal to the balloon are isolated from the main artery 1122, right artery 1124 and side branch artery 1136 thereby causing blood pressure distal to the balloon to drop from approximately normal arterial pressure of about 80 mmHg to a pressure in the range of 0-50 mmHg. When this happens, blood flow through the arteriovenous shunt 1132 can reverse as shown by blood flow arrow 1135, or the anterograde flow slowed or stopped. Referring to FIG 40B, initial injection of particles 1125 will be against a pressure, with a minimal anterograde flow or into a flow stasis.

15

20

25

30

35

Retrograde pressure flow against the particle injection can result from the flow of venous blood from yein 1130, through arteriovenus shunt 1135 and into the arterial capillary 1129 or from arteriovenous capillary beds associated with capillaries 1128. As particles 1125 are injected, they fill capillaries 1128 and 1129; however, particles cannot easily flow through arteriovenous shunt 1132 because of the reversal or slowing of flow and pressure. Continued injection can result in embolization of the distal portion of capillary 1129 and blockage of arteriovenous shunt 1132 with concomitant reduction or elimination of anterograde reflux. Increasing injection pressure through catheter 1140 following embolization of arteriovenous shunt 1132, can result in a high levels of particle density and distribution. Alternately, according to this method, a gradient can be used. The profile of the pressure gradient can be any function of time and pressure including, but not limited to, a linear or step function from low to high, high to low, alternating high to low and low to high or any other function and can be administered manually, in a semiautomated manner or using a programmable delivery means. Alternately according to this method, a pressure sensor as in FIG 36 can be used on the proximal and/or distal side of balloon 1142 to monitor pressure and select a procedural end point based on a definitive pressure reading. Referring now to FIG 40D, the injection through catheter 1140 into tumor 1133 can be accomplished using an automated pump/pressure monitor system whereby the pressure distal to occlusion balloon 1142 is measured on gauge 1152, the pressure reading transferred through connection 1154 to pump 1150 which controls the injection of anti-tumor agents from syringe

and/or pressure. The endpoint can be selected at any desirable pressure.

1156. Pump 1150 can be controlled manually or programed to any function of flow rate, time

[0153] The aim of the present method is to eliminate retrograde reflux, reduce or eliminate anterograde reflux, control the particle density and distribution, deliver an optimal dose, enable a defined pressure endpoint, improve efficacy and reduce toxicity.

[0154] Referring to FIG. 41A, a longitudinal cross section of a catheter is shown with proximal and distal ends, catheter body 204, distal tip 203 and proximally disposed hub 206. Catheter body 204 has two lumens that are in fluid communication with hub 206, a first lumen extending from port 208 of hub 6 to the distal tip 203 of catheter 204 whereby fluid can be injected from the proximal hub 206 and exit at the distal tip 203 of catheter 204 and a second lumen extending from port 210 of hub 206 to an intermediate location at some distance from the distal tip 203 of catheter 204, the second lumen adapted to communicate with a balloon for inflation and deflation.

10

15

20

25

30

35

[0155] Referring to FIG 41B, a longitudinal cross section of a first embodiment of the present disclosure is shown with proximal and distal ends, catheter body 204, distal tip 203, and two layered occlusion balloon 214 with channels 205 and valves 207 and proximally located hub 206. Although balloon 214 is shown with two channels, each with a valve, balloon 214 can have 1, 2, 3, 4 or any number of channels and any number of valves or be without valves. In this instance, the valve configuration allows fluid to flow from the proximal side of balloon 214 to the distal side of balloon 214 and to restrict flow from the distal side of balloon 214 to the proximal side of balloon 214; however, the opposite valve orientation and flow direction is also part of the present disclosure. Catheter body 204 can have a diameter of between 1 Fr and 10 Fr, more typically 2 Fr to 5 Fr and a length of 10 cm to 250 cm, more typically 50 cm to 150 cm. Two layered occlusion balloon 214 can be from 1 mm to 30 mm in diameter, more typically 2 mm to 10 mm in diameter, in its radially expanded configuration.

[0156] Referring to FIG 41C, an alternate embodiment of the device of the present disclosure is shown, having catheter body 218, distal tip 209, hub 206 and umbrella shaped occlusion structure 220. When in its radially expanded configuration, the occlusion structure will completely occlude the flow of the vessel. The umbrella shaped occlusion structure 220 is positioned at some distance from the distal end of catheter 216 and forms an umbrella shaped structure disposed circumferentially about catheter 216 with its outer diameter in contact with the vessel. Umbrella shaped occlusion structure 220 can be from 1 mm to 30 mm in diameter more typically 2 mm to 10 mm in diameter when in its radially expanded configuration and a longitudinal thickness of 0.25 mm to 10 mm, more typically 0.5 mm to 2 mm. Umbrella shaped occlusion structure 220 is shown with its closed end attached to the catheter distal to the open end of the V shape; however, it can be positioned in the opposite orientation or it can be positioned at a 90 degree angle with respect to catheter body 18.

[0157] Referring to FIG 41D, device 222 of the present disclosure is shown having catheter 218, distal tip 209, hub 206, and a unidirectional umbrella occlusion structure 224 with channels 230 and valves 228. Occlusion structure 224 will allow proximal to distal flow and prevent distal to proximal flow.

[0158] Referring to FIG 41E, device 232 is shown with catheter body 234, catheter distal extensions 235 and distal tip 211. Catheter extension 235 can have a diameter of 0.5 Fr to 5 Fr, more typically of 1Fr to 3 Fr and can be absent or can be of any length, typically 2 mm to 30 mm, more typically from 5 mm to 20 mm.

10

15

20

25

30

- [0159] Referring to FIG 41F, a preferred embodiment of the present disclosure is shown with catheter body 237, catheter extension 235, distal tip 211, nose-piece 241 and two layered occlusion balloon 243 in its radially expanded configuration. In this instance, two layered occlusion balloon 243 is disposed within a pocket formed on distal catheter extension 235 and between the distal end of catheter body 237 and the proximal end of nose-piece 241. The nose piece can be a tapered nose cone, a distally rounded piece of tubing or catheter, a blunt tube or any structure with a diameter equal to less than the catheter body. When in the radially constrained configuration, the outer diameter of the two layered occlusion balloon 243 has an outside diameter that is about equal to the outer diameter of the catheter body 237.
  - [0160] Referring to FIG 41G, yet another embodiment of device 245 of the present disclosure is shown, having catheter body 247, distal tip 234, nose-piece 241, proximal hub 206 and unidirectional umbrella shaped occlusion structure 224 with channels 230 and valves 228. In this instance, unidirectional umbrella occlusion structure 224 with channels 230 and valves 228 is disposed within a pocket formed on distal catheter extension 234 and between the distal end of catheter body 247 and the proximal end of nose-piece 241. The nose piece 241 can be a tapered nose cone, a radiopaque marker band, a distally rounded piece of tubing or catheter, a blunt tube or any structure of about equal diameter to the catheter body. When in the radially constrained configuration, the unidirectional umbrella shaped occlusion structure 224 has an outside diameter that is about equal to the outer diameter of the catheter body 247.
  - [0161] Referring to FIG 42, four views of a preferred embodiment of the unidirectional occlusion structure of present disclosure is shown. FIG 42A illustrates a two layered unidirectional occlusion structure 236 in its radially expanded configuration (also seen in FIGS 41B and 41F), having a proximal end 238, a distal end 240, balloon 242, balloon sheath 244, channel 246, valve structure 250, outer balloon sheath tail 254, balloon tail 256, flow direction arrow 252 and flow exit 248. When occlusion structure 236 is disposed on a catheter as in FIG 41F, fluid flows in the proximal to distal direction (anterograde) as indicated by arrow 252 through channel 246 and valve 250 and exits out the distal flow exit 248. The anterograde fluid

pressure on the inner surface of balloon sheath 244 at the distal end of channel 246, causes distally directed displacement or deflection of the inner surface of balloon sheath 244 at valve 250, thus allowing fluid to pass through flow exit 248. When flow is reversed, fluid pressure on the outer distal surface of balloon sheath 244 at valve 250 causes the balloon sheath 244 to press against the distal surface of balloon 242, closing valve 250 and preventing retrograde flow. Placing the unidirectional occlusion structure 236 in the opposite direction on the catheter will result in distal to proximal flow and prohibit proximal to distal flow. Although the occlusion structure of FIG 42A is shown with two layers including an inner balloon and an outer sheath, it is understood that the sheath need not be present and a balloon with channels from the proximal surface to the distal surface is considered part of the present disclosure. Balloon 242, including channels 246 can be formed by molding, extruding, vacuum forming or otherwise shaping a material to include the desired number and configuration of channels. Alternately, a standard balloon, including but not limited to, round or oval, can be modified to achieve proximal to distal channels. One method to modify a balloon is by forming longitudinal pleats circumferentially oriented, thereby forming V shaped channels that extend from the proximal surface of the balloon to the distal surface of the balloon. Placing a sheath over such a modified balloon in the same manner as described above would give the same result as the balloon shown in FIGS 42A through 42D.

5

10

15

25

30

35

[0162] Referring to FIG 42B, a side view of the unidirectional occlusion structure 236 of this disclosure is shown with proximal end 238 distal end 240, balloon 242, balloon sheath 244 and flow direction arrow 252.

[0163] Referring to FIG 42C, a proximal view of the unidirectional occlusion structure 236 is shown with proximal end 238, distal end 240 and channels 246.

[0164] Referring to FIG 42D, a distal surface view of the unidirectional occlusion structure 236 is shown with proximal end 238, distal end 240 and flow exit 248. Flow exit 248 is formed as a space between balloon tail 256 and balloon sheath 254. It is also possible to terminate balloon sheath 244 immediately below channels 246 forming a valve 250 that does not include balloon sheath tail 254.

[0165] Referring to FIG 43, an illustration of a prototype of the unidirectional occlusion structure 236 is shown in its radially expanded configuration with proximal end 238, distal end 240, balloon sheath 244, balloon 242 (positioned inside balloon sheath 244), and channel 246. This device was tested and will withstand at least 220 mmHg against its distal surface without retrograde flow.

[0166] Referring to FIG 44, device 60 illustrates the unidirectional occlusion structure 272 in a radially constrained configuration adapted to the distal extension 267 of a catheter 265 with

distal end 262, proximal end 264, proximal collar 266, distal collar 268 (formed into a nose cone) and device pocket 270. Proximal collar 266 and distal collar 268 can comprise a metal, such as a radiopaque marker band, heat shrink tubing or any plastic material such as polyurethane, polyethylene, polystyrene, acetal, PTFE, nylon or the like, and can be 1 mm to 20 mm in length, more typically from 2 mm to 10 mm in length. In this instance, circumferentially oriented occlusion structure 272 is held within pocket 270 of catheter 265, with an outer diameter approximately equal to the outer diameter of catheter 265.

5

10

15

20

25

30

[0167] Referring to FIG 44B, device 260 is shown with unidirectional occlusion structure 272 in its radially expanded configuration with proximal end 264, distal end 262, balloon 242, balloon sheath 244, valve 250, flow exit 248, channel 246, flow arrow 252, proximal collar 266, distal collar 268, catheter 274, balloon fill lumen 270 and guidewire/injection lumen 276. In this instance, there is no distal balloon sheath tail, the balloon sheath terminating on the balloon surface just below channel 246 and above the perimeter of catheter extension 267, thereby positioning the flow exit between the termination of the balloon sheath and the catheter.

[0168] Referring to FIG 45A, device 280 is shown with catheter 282 and an umbrella shaped structure occlusion structure 2284 in its radially expanded configuration, whereby the umbrella shaped occlusion structure 284 is oriented circumferentially about catheter 282 such that its outer circumference is 360 degrees about catheter 282. When device 280 is placed in an artery or vein and umbrella shaped occlusion structure 284 is placed in its radially expanded configuration, the outer perimeter of occlusion device 284 will be at least in contact with the interior of the vessel wall and substantially occlude flow. FIG 45A shows occlusion structure 284 in forward V orientation and FIG 45B shows device 282 with the umbrella shaped occlusion device 288 in a reverse V configuration. The occlusion structure of the present disclosure can also have a 90 degree orientation with respect to the catheter when in its radially expanded configuration.

[0169] Referring to FIG 46A, device 290 is shown with proximal end 292, distal end 294, catheter 296, two-way occlusion structure 299 in its radially expanded configuration, frame 298 and channels 2100, whereby fluid can flow from proximal to distal or distal to proximal through channels 2100. Although two channels are shown, two-way occlusion structure 298 can have 1, 2, 3 or any number of channels.

[0170] Referring to FIG 46B, device 2102 is shown with proximal end 2104, distal end 2106, catheter 2108, and unidirectional umbrella shaped occlusion device 2110 comprising, frame 2111, channels 2112 and radial valve 2114, whereby fluid will flow from proximal to distal (anterograde) only, retrograde flow being prohibited by radial valve 2114. Although device 2102 will allow only anterograde flow, if desired, device 2110 of apparatus 2102 can be

configured to allow only retrograde flow and/or have a forward V configuration as shown, or, if desired, a reverse V configuration or an orientation 90 degrees with respect to catheter 2108.

[0171] The frames 298 and 2111 of occlusion structure 2110 can be made of metal, such as shape memory metals nitinol or elgiloy, or plastic such as polyethylene, polyurethane, polystyrene, PTFE, acetal and nylon or elastic materials such as silicone or fabrics such as cotton

5

10

15

35

and rayon and can include a mesh, a wire frame, a diaphragm and can be pleated or otherwise folded or can be any other convenient structure or material provided that it is of sufficient strength and porosity to occlude elevated vascular pressures and capable of integrating channels and valves. Valve 114 can be made from flexible or rigid plastics including polypropylene and polyurethane, elastomeric materials such as silicone and can have a configuration including a flap, sock, cone, duck bill and diaphragm or the like with a thickness of 1 mil to 50 mil, more typically 2 mil to 10 mil.

[0172] Referring to FIG 47A, a distal surface view of a unidirectional occlusion structure 2120 of the present disclosure is illustrated with the catheter 2122 (extending forward), device frame 2124, radial valve 2125 and channels 2126 disposed under radial valve 2125. As shown, radial valve 2125 extends radially outward from catheter 290 and covers all four valves. Four channels are shown in this example; however, any number of channels can be used. This configuration allows flow from the proximal surface to the distal surface of unidirectional umbrella shaped occlusion structure 2120; however, the reverse flow is also possible.

20 [0173] FIG 47B illustrates another embodiment of the present disclosure comprising unidirectional umbrella shaped occlusion structure 2128 with catheter 2130 (extending forward), device frame 2132, valves 2136 and channels 2134 disposed under valves 2136. In this instance, each channel has a separate valve and although four channels and valves are shown, the device of this disclosure can have any number of channels and valves limited only by the size of the valve and channel and the area of the frames 2124 and 2132. This unidirectional configuration allows flow from the proximal surface to the distal surface of umbrella shaped occlusion structure 2128; however, the opposite flow can be easily achieved by changing the flow direction of the valves or rotating the unidirectional occlusion device 180 degrees on catheters 2122 and 2130.

30 **[0174]** FIG 48 illustrates a prototype micro-valve 2140 configured from 5 mil polyurethane material. This device was tested and will restrain a fluid pressure of at least 220 mmHg applied against its distal surface.

[0175] Referring to FIG 49A, device 2142 is shown with triangular shaped occlusion structure 2145 in its radially expanded configuration and adapted to catheter 2144 whereby the occlusion structure 2145 has frame 2146 oriented circumferentially about catheter 2144 such that its outer

circumference comprises 360 degrees. When device 2142 is placed in an artery or vein, frame 2146 is placed in its radially expanded configuration and the outer perimeter of device frame 2146 will be at least in contact with the interior of the vessel wall and at least substantially occlude flow.

[0176] Referring to FIG 49B, device 2148 is shown with catheter 2150 and a unidirectional 5 triangular shaped occlusion structure 2151 comprising frame 2152, channels 2154 and radial valve 2158. Although reference has been made to a unidirectional occlusion valve with an umbrella shape or a triangular shape, it is understood that any shape including, but not limited to, rectangular, oval, conical, and round can be used. Yet another construction of a unidirectional occlusion structure is a dilation or occlusion balloon or any other medical balloon disposed with 10 channels and valves, the valves extending from a proximal surface to a distal surface. [0177] Referring now to FIG 50A to 50C, a method of deploying an occlusion structure 2191 from a radially constrained configuration to a radially expanded configuration and then returning it to the constrained configuration is shown. FIG 50A shows a longitudinal cross section of device 2180 with a proximal end 2182, a distal end 2184, outer catheter 2186, inner catheter 15 2188, nose cone 2190 and radially constrained unidirectional occlusion structure 2191 with frame 2192, valve 2194, and frame attachment point 2196. Unidirectional occlusion structure 2191 is attached to inner catheter 2188 at attachment point 2196 whereby occlusion device 2191 is preloaded with a force which encourages its distal end to pivot proximally outward at attachment point 2196. In this instance, the outer catheter 2186 constrains occlusion structure 20 2191 against the preloaded force. Device 2180 is first positioned in the vasculature at or in the

[0178] Referring to FIG 50B, outer catheter 2186 is retracted proximally as shown by arrow 2200 while the inner catheter 2188 is held stationary, thereby removing the constraint on occlusion structure 2191, allowing it to pivot outward and in a proximal direction at attachment point 2196 and into its radially expanded configuration. Frame 2192 can be made from a memory metal such as nitinol or elgiloy and pre-formed at attachment point 2196 to the radially expanded configuration thereby pre-loading an outward force on occlusion structure 2191 as it is moved to its radially constrained configuration. If a braided nitinol tube is used, it can be pre-formed into a radially expanded configuration whereby occlusion structure 2191 is oriented circumferentially about catheter 2144 with an outer circumference of 360 degrees. As in this example, the mesh can be coated with polyurethane, PTFE, silicone or the like and channels formed through the mesh and valves placed over the channels.

vicinity of a target structure.

25

101791 Referring to FIG 50C, outer catheter 2186 is retracted distally while holding inner catheter 2188 stationary thereby pivoting frame 2192 distally at attachment point 2198 and placing occlusion structure 2191 in its radially constrained configuration.

[0180] Referring to FIG 51, an anatomical structure 2200 is shown with main artery 2202, right artery 2204, left artery 2206, capillaries 2208, tumor 2209 and blood flow directional arrows 2212. FIG's 51A-51E illustrates a method of the present disclosure wherein a tumor is embolized with drug eluting beads as in Transarterial Chemoembolization (TACE). [0181] In a first step, device 2211, comprising a two lumen catheter 2214 and a radially constrained unidirectional balloon occlusion structure 2216 (also in FIG 42), is advanced over a guidewire 2213 using lumen one (guidewire/injection lumen) of catheter 2214 from an entry point on the surface of the body, usually the femoral artery at the groin, and positioned at, or in the vicinity of, an artery feeding a tumor as in FIG 51A. As indicated by arrows 2212, the blood flows in an anterograde direction over device 2211 and into capillaries 2208 of tumor 2209. [0182] In a second step, the unidirectional balloon occlusion structure 2216 is placed in a radially expanded configuration by inflating the inner balloon of the two layered device of FIG 15 42 using the second lumen of catheter 2214 (balloon inflation lumen) as seen in anatomical FIG 51B. When placed in a radially expanded configuration, normal blood pressure between about 80 mmHg and 130 mmHg urges valves 2217 of occlusion structure 2216 to the open position, thereby allowing anterograde blood flow through channels 2219 and into the capillaries 2208 of

10

20

25

30

tumor 2209.

[0183] Figure 51C illustrates a third step whereby chemoembolization particles 2218 are beginning to be injected into left artery 2206 and capillaries 2208 of tumor 209. At this point, valves 2217 of unidirectional occlusion structure 2216 are in the open position and blood is flowing in the anterograde direction through channels 2219 which continues to carry chemoembolization particles 2218 into the vasculature of tumor 2209.

[0184] Referring to FIG 51D, a fourth step is illustrated whereby chemoembolization particles 2218 begin to embolize the distal ends of capillaries 2208, increasing pressure in the proximal section of capillaries 2208 and left artery 2206. This back pressure causes blood flow and chemoembolization particles 2218 to flow in a retrograde direction; however, the back-pressure in left artery 2206 urges valves 2217 to close, thereby maintaining particles 2218 in the vascular compartment distal to occlusion device 2217. Using currently available straight tip catheters, the chemoembolization procedure would be terminated at this point since particles would reflux backward over the catheter and into the general circulation causing non-target embolization and associated complications.

[0185] FIG 51E illustrates a fifth step, not possible using present catheters, whereby embolization particles continue to be injected, without retrograde reflux, and further fill the vasculature of the tumor with particles 2218. This method can both prevent the complications associated with retrograde reflux and allow more particles to enter the tumor.

- [0186] FIG 51F is a final step in the present method whereby occlusion structure 2216 is placed in a radially constrained configuration and device 2211 is withdrawn from the body over guidewire 2213.
- [0187] Although particular mention has been given to a device that is capable of transitioning from a radially constrained configuration to a radially expanded configuration, such a transition is not required. A unidirectional occlusion structure of the present disclosure can be configured in a permanently expanded configuration. In this instance, the occlusion structure may be a highly flexible material such as a low durometer plastic or rubber or a flexible mesh or any material or construction that provides sufficient strength and flexibility to navigate through vasculature and to a target and provide unidirectional occlusion.
- 15 [0188] Referring to FIG 52, distal section 302 of a device is shown with distal end 303, proximal end 304, catheter body 306, distal tip 308, nose cone 310, partial occlusion balloon 312 in a fully expanded configuration, channel 314 and one-way valve 316. In this embodiment, flow is permitted in the proximal to distal direction through channel 314 and restricted, by one-way valve 316 to flow proximally. Partial occlusion balloon 312 can be any shape and diameters

  20 from 1 mm to 30mm more typically from 2 mm to 10 mm.
  - [0189] Referring to FIG 53, a longitudinal section 320 of a distal section of the device of the present disclosure is shown with catheter body 306 distal tip 308, nose cone 310, radially constrained balloon 322, proximal balloon pocket boundary 324 and distal balloon pocket boundary 326. Radially constrained balloon 322 sits within the pocket defined by the distal end of catheter 306 at boundary 324 and the proximal end of nosecone 310 at boundary 326. The outer diameter of the constrained balloon is approximately equal to the outer diameter of catheter body 306. This allows the balloon to sit within the pocket and maintain the catheter at a desirable minimal diameter.

25

30

35

[0190] Referring to FIG 54A, a distal section 330 of the device of the present disclosure is shown with catheter body 306, catheter extension 307, distal tip 308, nose cone 310, balloon pocket 332, guidewire and injection lumen 334 and balloon inflation lumen 336. The catheter body 306 has a diameter of 0.25 mm to 5 mm, more typically from 0.5 mm to 1.5 mm and a length of 10 cm to 240 cm more typically from 75 cm to 150 cm. The catheter extension 307 has a diameter of 0.25 mm to 3 mm more typically from 0.4 mm to 1 mm and a length of 5 mm to 100 mm more typically from 5 mm to 40 mm. The balloon pocket 332 has a depth equal to the

difference in diameter of the catheter body 306 and the catheter extension 307 and a length of 1 mm to 50 mm more typically from 5 mm to 15 mm. The balloon wall thickness and inner diameter are selected, extruded or molded to fit into balloon pocket 332 with minimal balloon extending above of balloon pocket 332.

5 [0191] Referring to FIG 54B, distal section 330 includes a balloon 338 in a radially constrained configuration held within balloon pocket 332 and having an outer diameter approximately equal to the outer diameter of catheter body 306.

10

- [0192] Referring to FIG 54C, distal section 330 includes balloon 338 in a radially expanded configuration with channels 314 and one-way valves 316 in a closed orientation. Partial occlusion balloon 338 can be any shape and diameters from 1 mm to 30 mm more typically from 2 mm to 10 mm and a length of 1 mm to 50 mm more typically from 5 mm to 15 mm. Channels 314 can be of any shape and configuration and an opening that is calibrated to the desired flow therethrough. In a preferred embodiment, the balloon will have a diameter of 6 mm and a channel diameter of 0.5 mm to 1.5 mm. Balloon 338, including channels 314 can be formed by molding, extruding, vacuum forming or otherwise shaping a material to include the desired number and configuration of channels. Alternately, a standard balloon, including but not limited to, round or oval, can be modified to achieve proximal to distal channels. One method to modify a balloon is by forming longitudinal pleats circumferentially oriented; thereby forming V shaped channels that extend from the proximal end of the balloon to the distal end of the balloon.
- Placing a sheath or film over such a modified balloon results in longitudinal channels and a one way valve as described in co-pending application 61/917,131.
  - [0193] Referring to FIG 55A through 55C, a serial construction of the device of the present disclosure is illustrated.
- [0194] Referring to FIG 55A, a longitudinal view of device construction 350 with catheter body 306, distally located catheter extension 307 and proximally located hub 352 comprising guidewire and injection port 354 and balloon inflation/deflation port 356.
  - [0195] Referring to FIG 55B, a longitudinal view of a device construction 350 is shown with added nose cone 310 and balloon pocket positioned between the distal end of catheter body 306 and the proximal end of nose cone 310.
- 30 [0196] Referring to FIG 55C, a longitudinal view of the device 358 of the present disclosure is shown with catheter body 306, balloon 340 and hub 352. Balloon 340 is shown in a radially expanded configuration with channels 314 and valves 316 in the open position.
  - [0197] Referring to FIG 56, an anatomical structure 360 is shown with tumor 362, main artery 366, distal main artery 367, side branch arteries 370 and 374, tumor capillaries 373, 375, 377 and
- 35 379 and blood flow direction arrows 368, 372, 376 and 378. In the case of a tumor in the right

liver lobe, artery 366 is the right hepatic artery and 367 is the distal right hepatic artery which flow toward the tumor as seen by flow direction arrow 368 and 378. In this instance, artery 370 is the gastroduodenal artery and artery 374 is a hepatoenteric artery such as the superduodenal artery, the normal flow of both is away from the hepatic artery, as shown by flow direction arrows 372 and 376, and into arterial networks which supply both the liver and gastrointestinal tract. Blood from the hepatic artery 366 also flows into the tumor capillaries 373, 375, 377, and 379. Normal blood flow through the right hepatic artery is in the range of 4 ml/sec.

5

10

15

20

25

30

35

[0198] Referring to FIG 57A through 57H, a tumor embolization method according to current medial practice is shown. At least some of the steps shown are used in current catheter based embolization therapy in tumors of the liver

[0199] The first step of the procedure is to advance guidewire 382 from the femoral artery at the groin, through the iliac artery, aorta, celiac artery, hepatic artery and into the right hepatic artery 366 as in anatomical structure 380 of FIG 57A. The diameter of guidewire 382 is typically from about 0.25 mm to 1.25 mm more typically from 0.4 mm to 1 mm.

[0200] In the second step of the procedure, illustrated in FIG 57B, guide-catheter 392 is advanced over guidewire 382 and along the same arterial path as for guidewire 382. Typically, the guide catheter has an outer diameter of about 1.5 mm to 2.5 mm and has a central lumen that can accept a microcatheter with an outer diameter of 0.5 mm to 1.5 mm. The guide catheter is too large to access the vasculature in the vicinity of the tumor and is typically advanced as far along the vascular path toward the tumor as possible. The blood flow follows the same normal pattern as in FIG 56 and flows around the sides of guide catheter 392.

[0201] In the third step shown in FIG 57C, guidewire 382 is optionally removed and replaced with a smaller diameter guidewire 398 that can fit in the central lumen of a microcatheter. Guidewire 398 typically has a diameter of 0.2 mm to 0.75 mm, more typically in the range of 0.25 mm to 0.6 mm.

[0202] The fourth step of the procedure, microcatheter 3104 is advanced over guidewire 398 to a position beyond the distal end of guide catheter 392 and into the vasculature within or in the vicinity of the tumor as shown in anatomical structure 3102 of FIG 57D. Microcatheter 3104 is advanced as close as is practical to the tumor and, if the anatomy allows, into the vasculature of the tumor as in superselective embolization. Microcatheter 3104 typically has a diameter of 0.75 mm to 1.5 mm, more typically at about 1 mm and a total length of 50 cm to 200 cm, more typically from 75 cm to 150 cm. The central lumen microcatheter 3104 is optimized to have an inner diameter as large as possible; however it is usually in the range of about 0.5 mm. [0203] In a fifth step, guidewire 398 is removed from microcatheter 3104 as illustrated in

anatomical structure 3200 of FIG 57E. Removal of guidewire 398 allows the central lumen of

microcatheter 3104 to be used to inject drug and/or embolic materials into the target site within the right hepatic artery and tumor. Blood continues to flow around guide catheter 392 and microcatheter 3104 and into capillaries 373, 375, 377, and 379 of tumor 362, gastroduodenal branch 370 and hepatoenteric branch 374 according to the normal flow pattern shown by arrows 368, 372, 376, and 378.

5

10

15

25

30

35

[0204] In a sixth step illustrated in anatomical structure 3208 of FIG 57F, drug and or embolization agents are injected using a syringe or other means from the proximal end of microcatheter 3104 through guidewire/injection port 354 of hub 52 (FIG 55), longitudinally through guidewire injection lumen 334 (FIG 54), and out the distal end of catheter extension 307. In this instance, embolic particles 3210 are carried by normal blood flow into distal right hepatic artery 3214, tumor capillaries 373, 375, 377 and 379, as illustrated by flow arrow 3212, and into hepatoenteric artery 374 in the direction indicated by flow arrow 376. Drug and/or embolization agents that travel through gastroenteric branch artery 374 or any other arterial branch distal to the distal tip of microcatheter 3104, by normal forward flow, are deposited at non-target sites, including parts of the liver and intestine. This anterograde (to the catheter tip) bypass into distal hepatoenteric arteries can cause serious complications including damage or death to sections of the liver or intestine, gastric ulcers or even the death of the patient. Further, drug and/or embolic agents that travel to non-target sites, fail to enter the tumor; this resulting in a lower than optimal dose to be delivered to the tumor and a lower efficacy than desired.

However, to avoid the aforementioned serious complications, physicians often under-embolize the tumor vasculature.

[0205] Referring to FIG 57G, as forward flow mediated embolization progresses, the distal ends of capillaries 373, 375, 377 and 379 fill with particles 3210 and become embolized. This process dramatically slows the blood flow moving through the tumor and causes a sharp rise in pressure within the tumor vasculature and concomitant retrograde deflection of the high volume blood flow from the hepatic artery. Further, backpressure develops in the distal hepatic artery 3214, resulting in particles to flow in a retrograde direction as in flow direction arrow 3234. This can result in: (1) increased anterograde bypass into hepatoenteric branch 374, reflux over the catheter and into gastrodudenal artery 370 and (3) a signal to the physician that particle injection should stop, even though the tumor is only partially embolized. In this instance, it is possible that larger capillaries become embolized first, due to a larger blood flow while smaller capillaries remain un-embolized. The rapid rise in pressure is in part caused by distal capillary embolization and in part caused by the high volume blood flow from the hepatic artery. Given that embolization is the desired endpoint, it appears that slowing the forward flow of blood from the hepatic artery would allow the tumor to accept the blood and drug and/or embolic agent flow for a longer

period of time and allow more embolization to occur and an improved distribution of particles in the tumor vasculature.

[0206] Referring to FIG 57H, injection of drug and/or embolic agents is complete, microcatheter 3104 and guide catheter 392 are removed and the final embolization distribution in tumor capillaries 373, 375, 377 and 379 is shown where, in this example, lager capillary 379 is embolized to the greatest extent, smaller capillary 377 is embolized to a lesser extent and small capillary 373 remains un-embolized.

5

10

15

20

25

30

35

[0207] Referring to FIG 58, a method of tumor embolization, according to the device of the present disclosure is shown. Steps 1 through step 5, shown in FIG's 57A through 57E, are the same for both a standard catheter as shown in FIG 57 and the device of the present disclosure and are not further illustrated.

[0208] Referring to FIG 58A, the device of the present disclosure is positioned in the distal right hepatic artery 3214, with partial occlusion balloon 3256 in its radially expanded configuration comprising channels 3258 and 3260, and one-way valves 3262 and 3264 that are in the open position. FIG 58A shows two channels, however, one, two, three or any number of channels can be used. With or without valves. The maximum channel size is limited by balloon diameter, but can be as small as is practical. Valves 3262 and 3264 can be flap, duck bill, diaphragm, or any type of valve provided that it permits flow only in one direction. Optional pressure sensors 3266, which provides real time pressure measurement in the vascular space distal to partial occlusion balloon 3256 and pressure sensor 3265 which provides real time pressure monitoring in the vascular space proximal to the partial occlusion balloon 3256 are shown. Pressure sensor 3266 which measures pressure in the distal vascular space can be used to signal a procedural endpoint based on a predetermined or non-predetermined pressure reading. This will, for the first time, allow a quantitative and definitive pressure mediated endpoint rather than the present subjective flow mediated endpoint and will enable the procedure to be reproducible and able to be standardized allowing center to center and physician to physician consistency. This is possible only because the vascular space distal to the partial occlusion balloon 3256 is isolated from the vascular space proximal to partial occlusion balloon 3256, thereby allowing that the arterial pressure in the distal space to be closely related to the intra-tumor arterial blood pressure. Blood flow direction through hepatic artery 368, distal hepatic artery 3214 and proximal artery 370 are normal as seen in flow direction arrows 368, 372 and 378 as is the blood flow in tumor capillaries 373, 375, 377 and 379 and illustrated by flow direction arrows 3212. However, partial occlusion balloon 3256 causes a significant reduction in blood flow in distal right hepatic artery 214 and in tumor capillaries 373, 375, 377 and 379. Blood flow can be regulated by the partial occlusion balloon of the present disclosure such that total flow can range from near 100%

(unconcluded flow) to near 0% as in full occlusion. Of most interest is partial occlusion that results in 1% to 25% flow as compared to the un-occluded artery. Therefore, channels 3258 and 3220 allow only a fraction of normal blood flow to pass distally. Blood pressure distal to partial occlusion balloon 3256 is also dramatically reduced by anywhere from about 5 mmHg reduction to 100 mmHg reduction, depending on the nature of the occlusion. This pressure drop causes branch artery 374 to reverse direction as seen by flow direction arrow 376 and now flow toward the distal main artery 3214 and tumor capillaries 373, 374, 377 and 379. The flow reduction and pressure reduction caused by partial occlusion balloon 256 also reduces the flow and pressure within the tumor capillaries 373, 374, 377 and 379. This is of significance because it allows more drug/embolization agents to enter the tumor before backpressure causes flow stasis and retrograde flow.

5

10

15

20

25

30

35

[0209] Referring now to anatomical structure 3280 of FIG 58B, injection of drug and/or embolic agents is initiated. Blood flow from the proximal main artery 366 is attenuated as it passes through channels 3258 and 3260 and into distal main artery 3214. The anti-cancer agents are carried by the attenuated forward blood flow through distal main artery 3214 and into tumor capillaries 373, 375, 377 and 379. Valves 3262 and 3264 are in the open position as pressure in the distal vascular space is lower than the blood pressure in the vascular space proximal to partial occlusion balloon 3256. Branch artery 374 continues to flow in the reverse direction as indicated by flow direction arrow 376 since blood pressure in the distal right hepatic artery is lower than that of the arterial network connected to the distal end of branch artery 374. In this instance, anterograde drug/embolic agents are prevented from flowing into branch artery 374 and anterograde bypass and non-target delivery does not occur. Optional pressure sensors 3266 and 3265 or pressure measurement through guidewire/injection lumen 334 of catheter 6 (FIG 54A) can be used to monitor real-time pressure.

[0210] Referring to FIG 58C, and looking at anatomical structure 3290, a slower rate of blood flow and lower pressure through distal hepatic artery 3214 allows tumor capillaries 373, 375, 377 and 379 to fill at a slower pace and to a greater distribution than in the current method of full unregulated forward flow. At some point, however, the embolization of tumor capillaries will cause retrograde deflection of blood and anti-tumor agents and a pressure build up in distal hepatic artery 3214 as in FIG 58D. At this point, the increased pressure in distal right hepatic artery 3214 causes valves 3262 and 3264 to close, preventing retrograde flow and non-target embolization through branch artery 370 or any other arteries proximal to partial occlusion balloon 3256. This retrograde deflection and pressure build up will progress at a slower rate as compared to the current standard method of FIG 57. The slower buildup of pressure and retrograde flow allows a larger time window for the physician to terminate the procedure. If

pressure monitoring is done, a defined pressure can be used to terminate the procedure. If the back pressure in distal hepatic artery 3214 exceeds about systolic pressure, branch artery 374 will again follow in the normal flow pattern as illustrated in the flow direction arrow 376 of FIG 56. This situation will allow anterograde drug and/or embolic agents to flow in branch artery 374 and to non-target sites. However, visual observation of contrast movement in branch artery 374 or a defined pressure measurement at or below the flow reversal pressure of branch artery 374 can be used as a procedural endpoint signal.

5

10

15

20

25

30

35

[0211] Referring now to anatomical structure 3300 of FIG 58E, the procedure is complete and catheter 3254, of the present disclosure, and guide catheter 3252 are removed. The distribution and filling of tumor capillaries 373, 375, 377 and 379, using the device of the present disclosure, are improved as compared to the distribution and filling associated with the current microcatheter 3104 of FIG 57H.

[0212] In some embodiments, a pressure feedback loop is implemented to control the rate of fluid introduction (e.g. the rate of embolic bead injection). In such embodiments, a blood pressure sensor may be located at the distal tip of the catheter to monitor the pressure of the vasculature distal to the occlusion or partial occlusion structure. In some embodiments, a predetermined pressure set point may be used. The set point may be an absolute pressure, or a percentage of systolic pressure (such as about 100% of systolic pressure). Such systems can be programmed to maintain the distal vasculature at or below the set point by automatically controlling the infusion rate (e.g. the rate or pressure of an injection pump.) By ensuring that the pressure of the distal vasculature does not exceed systolic pressure, retrograde and/or other undesirable blood flows can be prevented.

[0213] In some embodiments, a rate of pressure change can be used instead of or in addition to a preset pressure in the pressure feedback loop. For example, if the distal vasculature pressure begins to rise faster than a predetermined rate, the rate of embolic substance injection can be slowed, temporarily stopped, or a procedural endpoint may be signaled. The endpoint signal may be an audible, visual, tactile or other signal to persons involved in the procedure, and/or may be a signal that automatically shuts off or changes the state of medical equipment used in the procedure.

[0214] FIGS 59A – 59D show the fabrication steps of the distal end of an occlusion catheter according to aspects of the present disclosure.

[0215] FIG 60 shows the completed distal end of the occlusion catheter of FIGS. 59A - 59D being introduced into small branches of a vascular system. The two enlarged contact points of this design allow the catheter to be navigated into smaller vasculature than can be navigated by conventional catheter tips.

[0216] FIG 61A shows an entire occlusion catheter constructed according to principles of the present disclosure. FIG 61B shows details of construction features on the distal end of the occlusion catheter of FIG 61A.

[0217] It is important to note that the device of the present disclosure regulates flow and pressure in the arterial space distal to the partial occlusion balloon, significantly reduces flow rate and pressure in tumor capillaries and causes flow reversal of distal branch arteries. The aforementioned device and method thereby enable substantial elimination of retrograde and anterograde bypass to non-target sites and a more complete filling of the tumor vasculature with drug and or embolic agents and should improve efficacy and reduce complications over standard devices and methods.

#### **CLAIMS**

What is claimed is:

5

10

15

20

25

30

35

1. A catheter assembly comprising:

a catheter body having a proximal end, a distal end and a balloon inflation lumen; and an inflatable balloon attachable to the distal end of the catheter body, the balloon having an inner surface that at least partially defines an interior volume, the balloon being configured such that the interior volume can be in fluid communication with the inflation lumen of the catheter body to inflate the balloon, the balloon having a proximal surface and a distal surface,

wherein the balloon is provided with a channel that extends through the balloon, the channel configured to provide fluid communication between the proximal surface of the balloon and the distal surface of the balloon.

- 2. The catheter assembly of claim 1, wherein the catheter assembly comprises a distal tip section that extends beyond the distal end of the catheter body.
- 3. The catheter assembly of claim 1 wherein the balloon comprises a valve configured to affect fluid flow through the channel.
- 4. The catheter assembly of claim 3 wherein the valve is positioned at a distal end of the channel.
- 5. The catheter assembly of claim 3 wherein the valve is positioned within the channel.
- 6. The catheter assembly of claim 3 wherein the valve is configured to allow flow in a distal direction and restrict flow in a proximal direction.
- 7. A device for delivering a therapeutic agent to a target site within a body, the device comprising:

a catheter body having a proximal end, a distal end, a first axial lumen and a second axial lumen, the first axial lumen extending from the proximal end of the catheter body to the distal end of the catheter body and providing fluid communication therebetween, the second axial lumen extending from the proximal end of the catheter body to a more distal location on the catheter body;

a balloon radially disposed near the distal end of the catheter body, the balloon having a proximal balloon surface, a distal balloon surface, a radially constrained configuration and a radially expanded configuration, wherein the balloon is in fluid communication with the second axial lumen and has at least one channel extending from the proximal balloon surface to the distal balloon surface, thereby providing fluid communication therebetween; and

a balloon sheath, disposed over an outer surface of the balloon, the sheath having a proximal side and a distal side, the proximal side having an opening that is in alignment with a

proximal end of the balloon channel, the distal side of the sheath extending over a distal end of the balloon channel, wherein the distal side of the sheath is capable of deflecting distally when a distally directed fluid pressure is applied through the opening in the proximal side, through the balloon channel and against a proximal surface of the distal side of the sheath, thereby allowing fluid to pass through the balloon channel and into a vascular compartment distal to the balloon and the balloon sheath, and wherein the distal side of the sheath is also capable of deflecting proximally onto the distal balloon surface when a pressure is applied to the distal side of the sheath, thereby preventing proximally directed flow in the balloon channel.

5

10

15

20

30

35

- 8. The device of claim 7, wherein the interface between the distal side of the sheath and the distal balloon surface forms a unidirectional valve.
  - 9. The device of claim 7, wherein a catheter extension is disposed longitudinally from the distal end of the catheter body.
- 10. The catheter extension as in claim 9, wherein the catheter extension has a diameter less than the catheter body.
- 11. The device of claim 9, wherein the balloon and the balloon sheath are disposed on the catheter extension.
  - 12. The device of claim 9, wherein a nose piece is adapted to the catheter extension at a location distal to the balloon and the balloon sheath.
  - 13. The device of claim 12, wherein the balloon and the balloon sheath are positioned in a pocket between the distal end of the catheter body and a proximal end of the nose piece.
  - 14. The device of claim 9, wherein an outer diameter of the radially constrained balloon and the balloon sheath is less than or equal to an outer diameter of the catheter body.
    - 15. A method of embolizing a tumor, the method comprising:

advancing a device including a catheter body and a partial occlusion structure to a target tumor site within the body;

allowing an anterograde blood flow past the partial occlusion structure, the allowed anterograde blood flow being less than a blood flow that would normally be present if the partial occlusion structure were not in place;

injecting an embolic substance from the device to allow the anterograde blood flow to carry the embolic substance into a vasculature of the tumor target;

monitoring a real time pressure measurement in a vascular space distal to the partial occlusion structure;

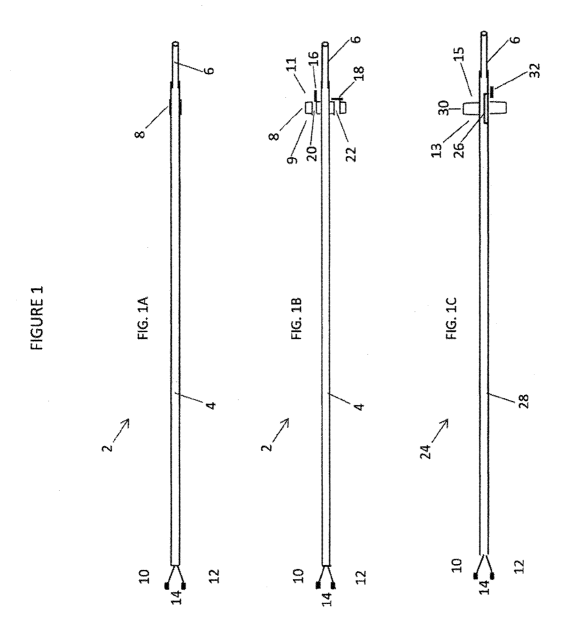
signaling a procedural endpoint based on the monitoring of the pressure measurement; and

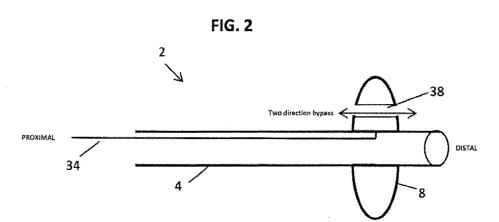
withdrawing the device from the body.

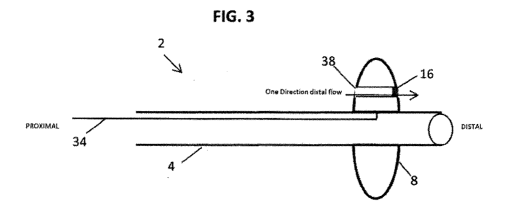
16. The method as in claim 15, wherein the allowed anterograde blood flow is within a range of about 1% to about 25% of a blood flow that would normally be present if the partial occlusion structure were not in place.

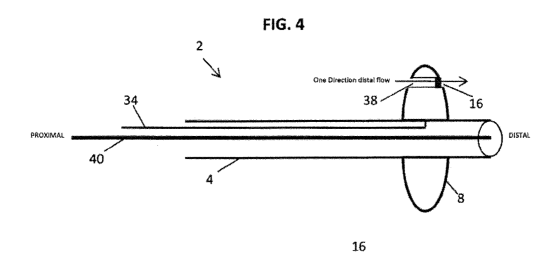
- The method of claim 15, wherein the signaling of the procedural endpoint occurs
   when a predetermined pressure is reached in the vascular space distal to the partial occlusion structure.
  - 18. The method of claim 17, wherein the predetermined pressure is a predetermined percentage of an un-occluded systolic pressure.
    - 19. The method of claim 17, wherein the predetermined percentage is about 100%.

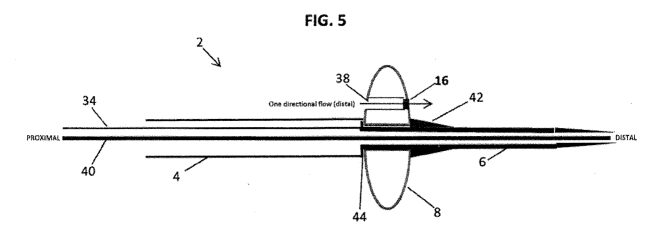
10

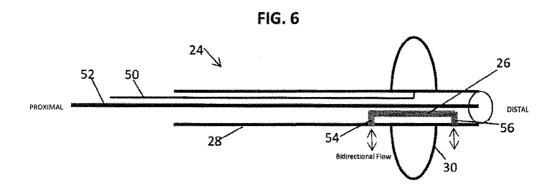


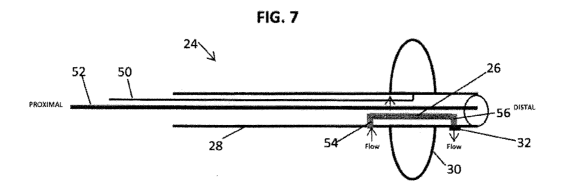


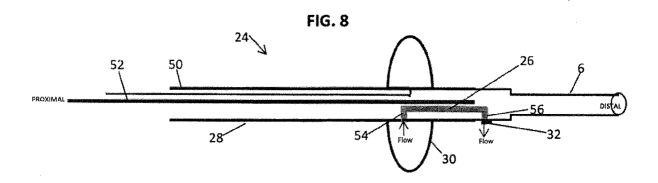






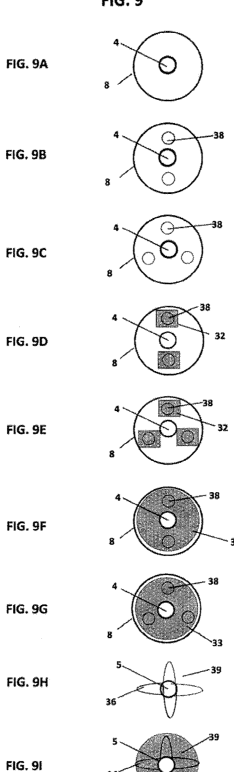






5/47

FIG. 9



6/47



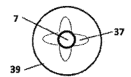


FIG. 9K

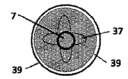
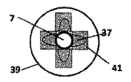
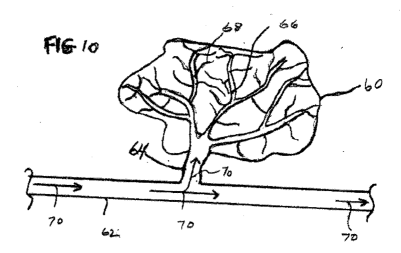
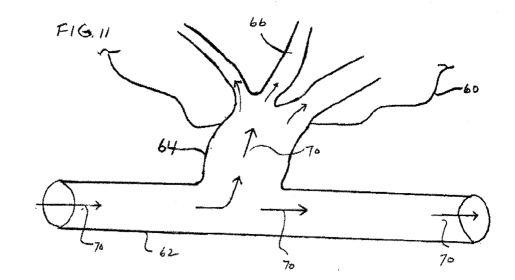
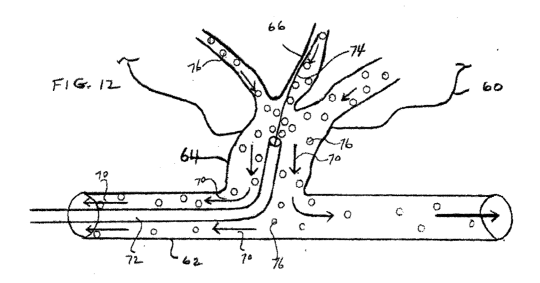


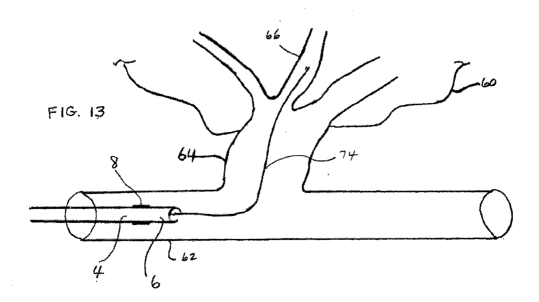
FIG. 9L

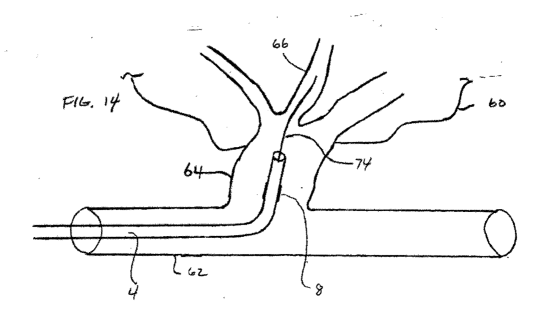


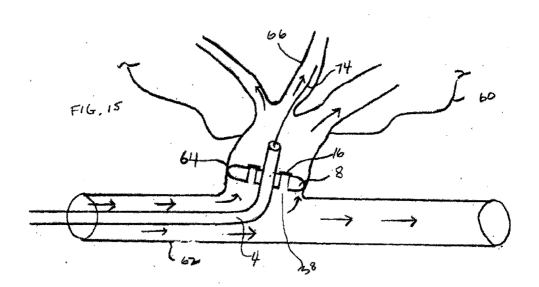


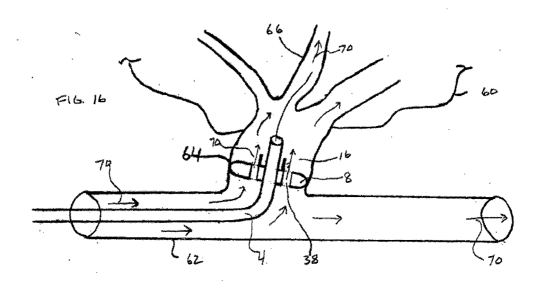


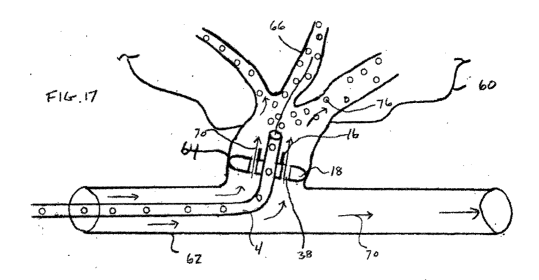


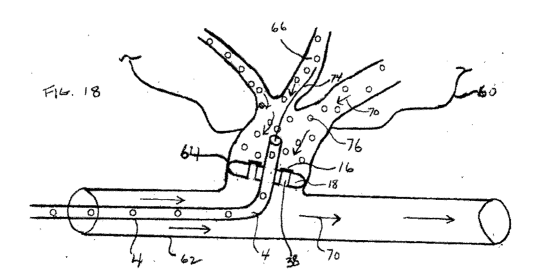


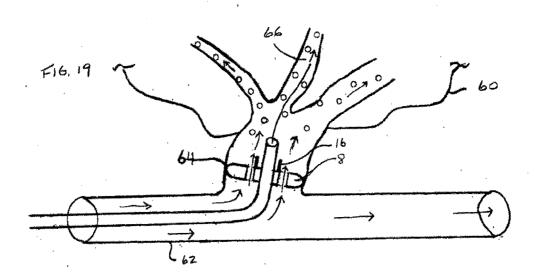


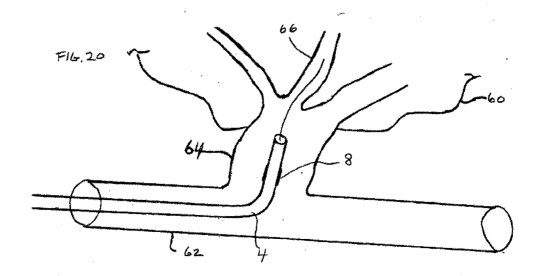


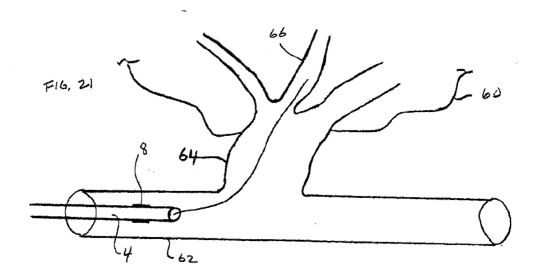


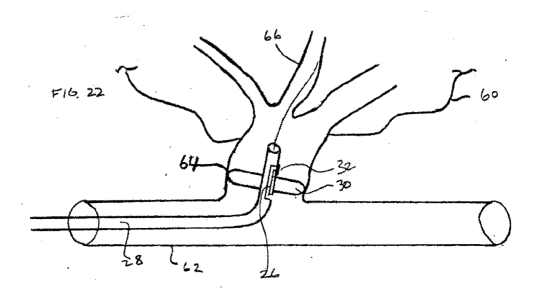


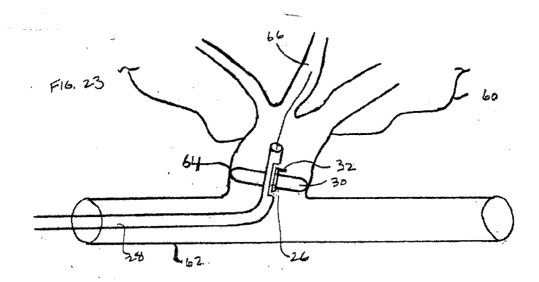


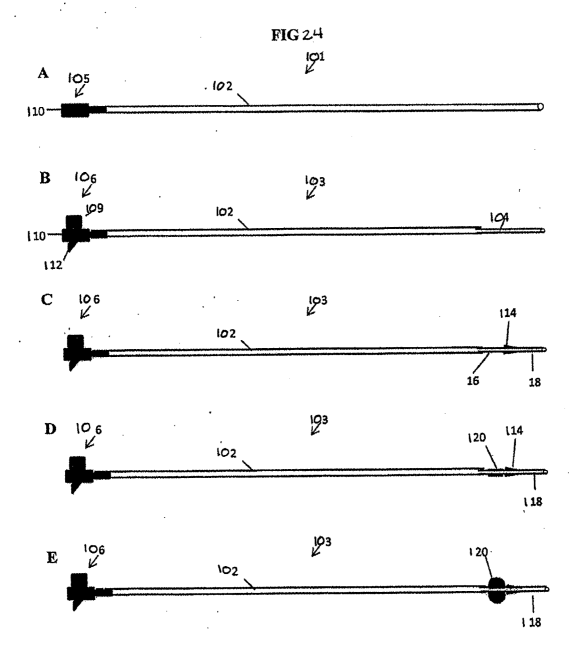












15/47

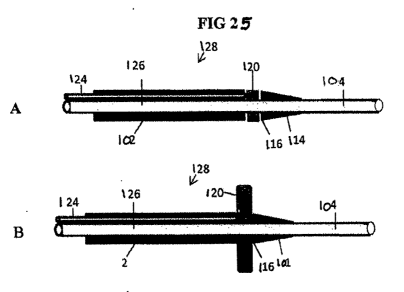
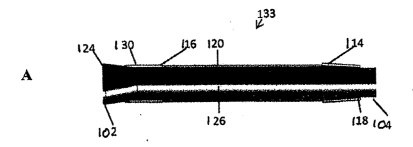
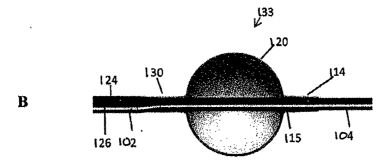
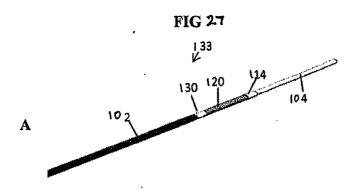
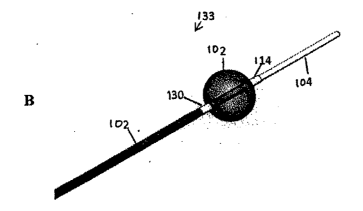


FIG 240

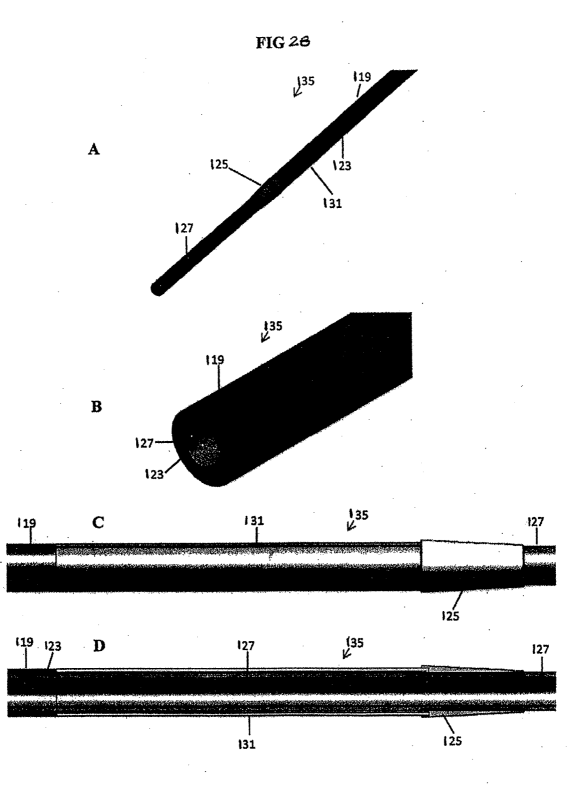








17/47





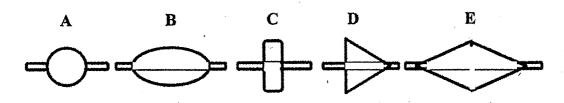
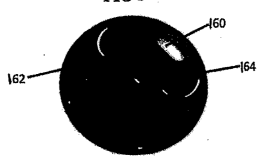


FIG 30



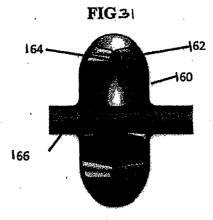


FIG 32

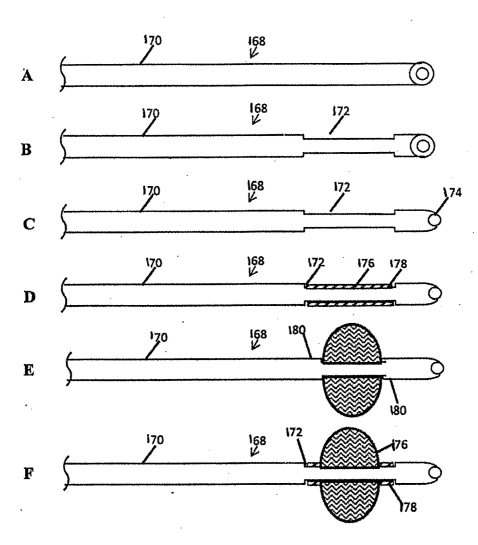


FIG 33

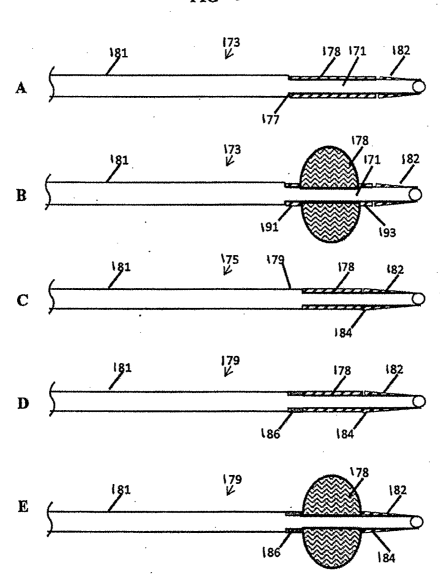
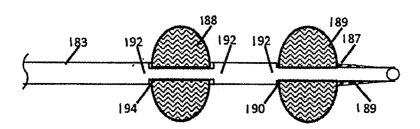
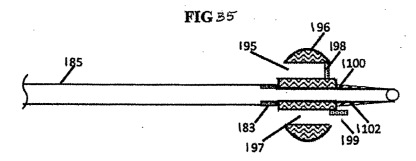
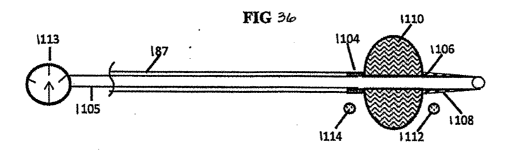


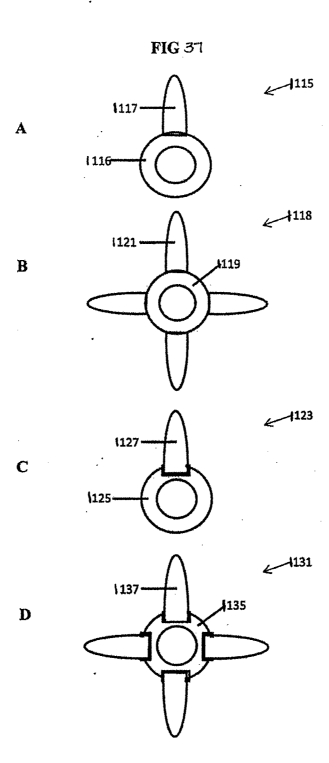
FIG 34







22/47



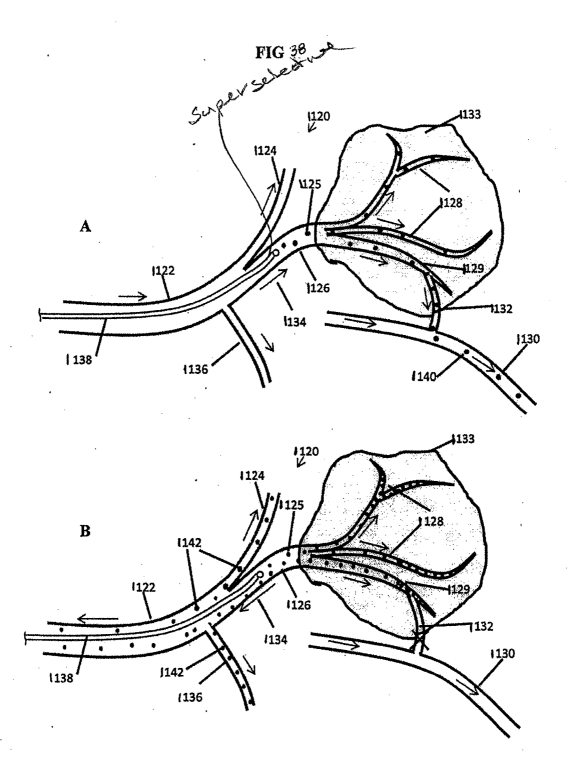
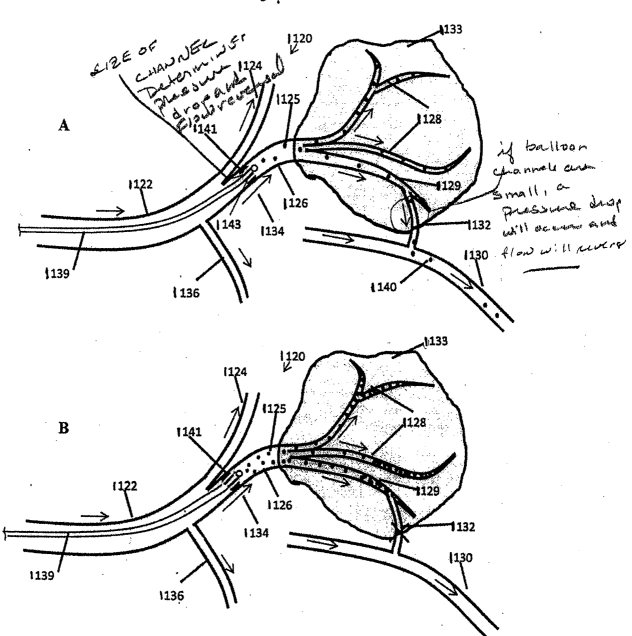
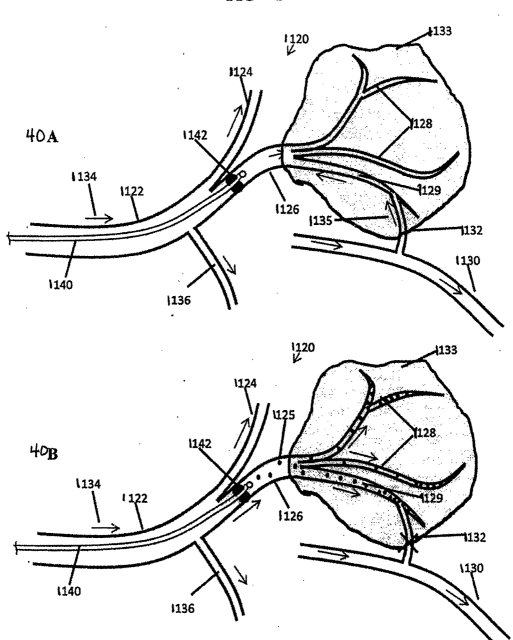
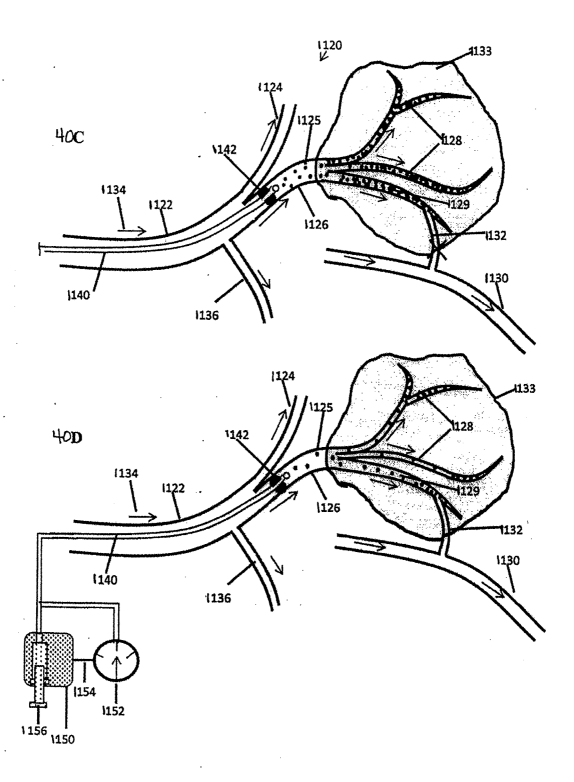


FIG 39









F1G.41

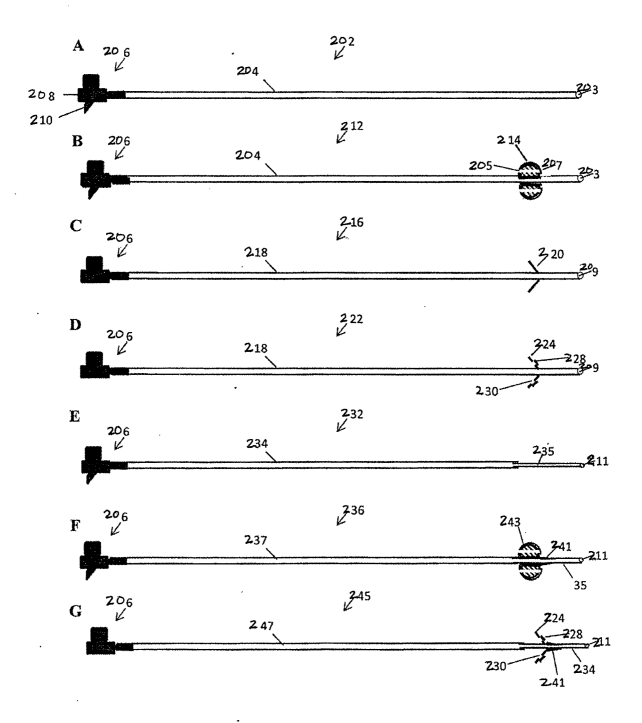
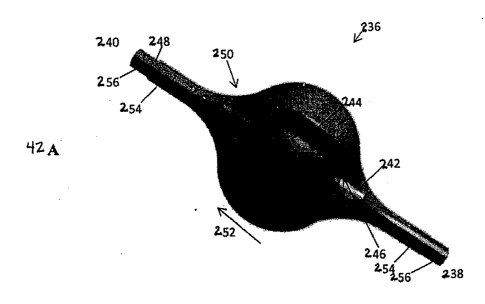
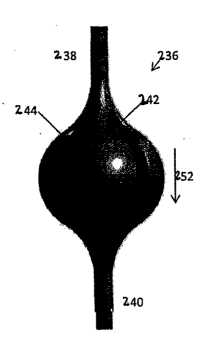


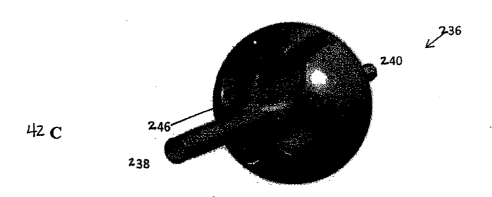
FIG 42



42B



29/47



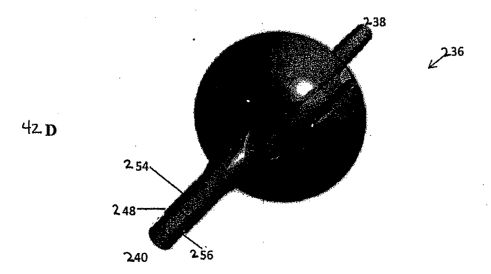


FIG 43

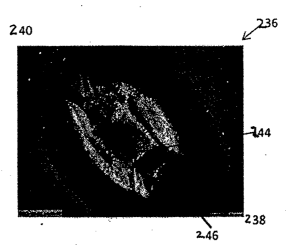
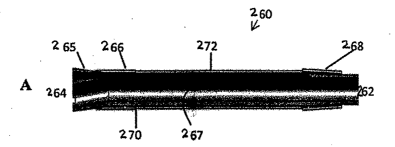


FIG 44



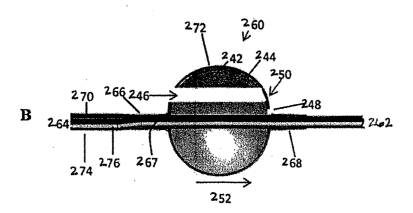


FIG 45

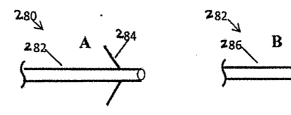


FIG 46

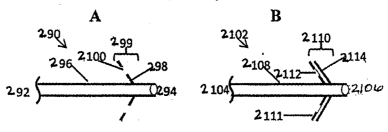


FIG 47

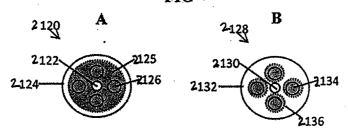


FIG 48

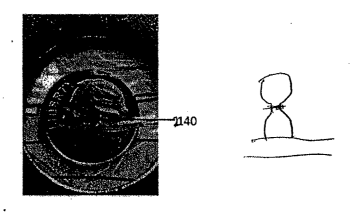


FIG49

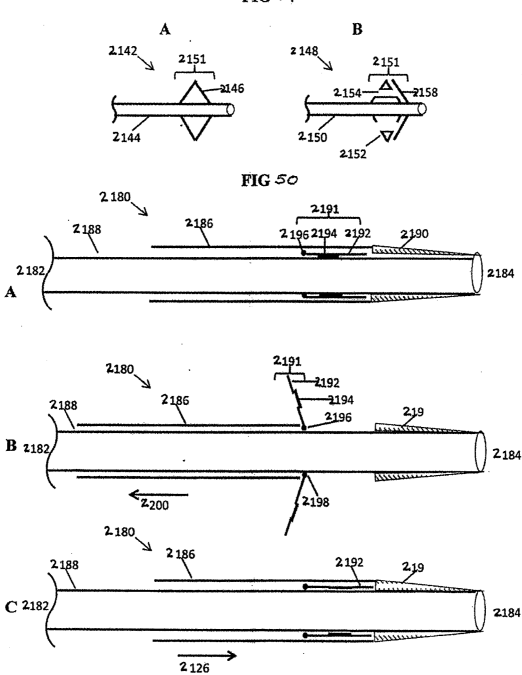
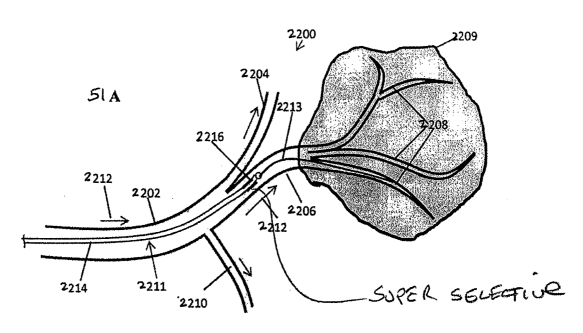
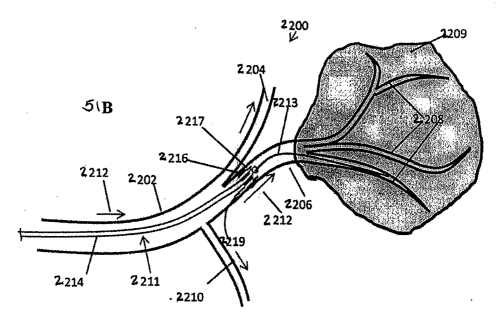
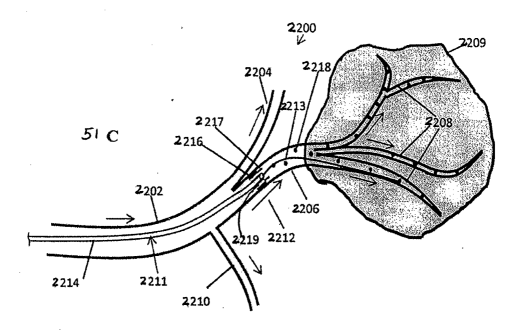
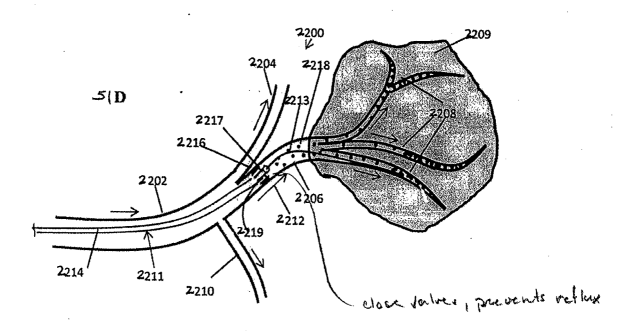


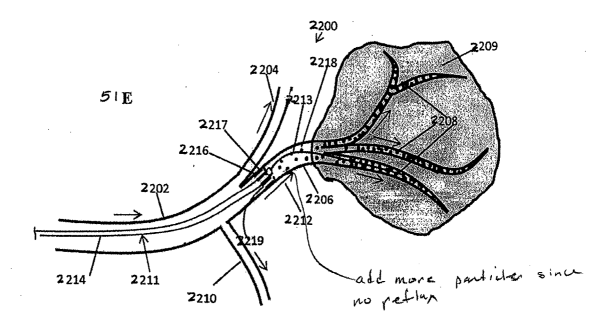
FIG 51

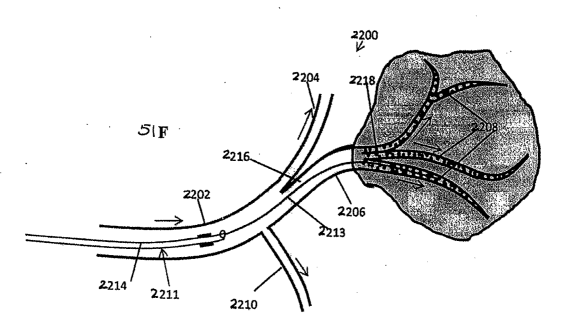


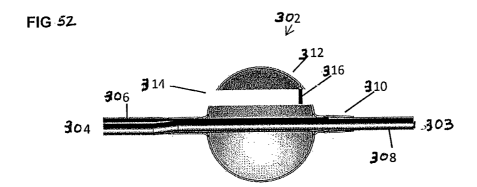


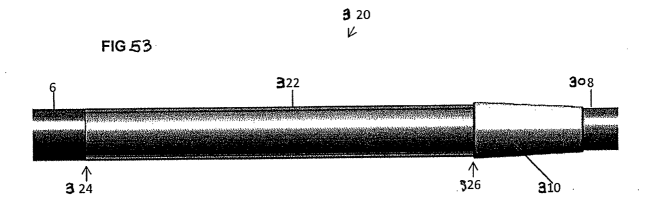












37/47

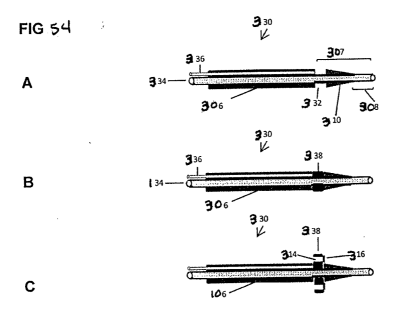
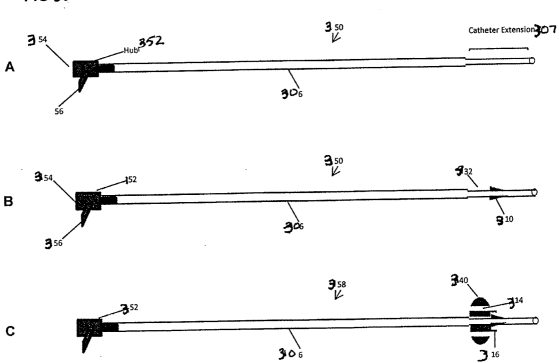
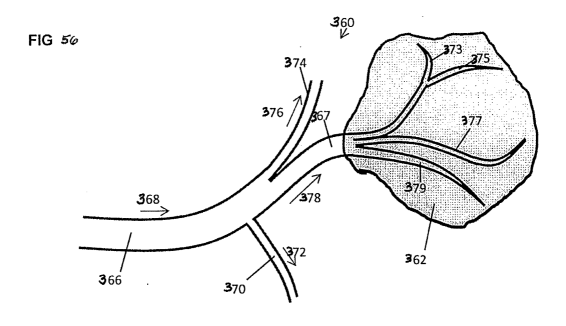


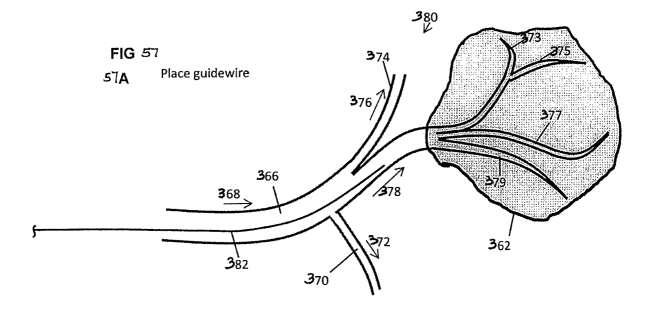
FIG 55

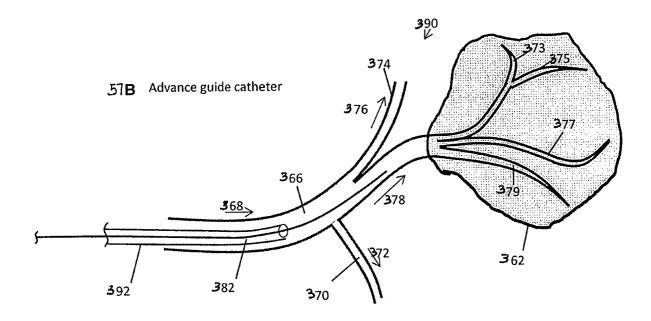


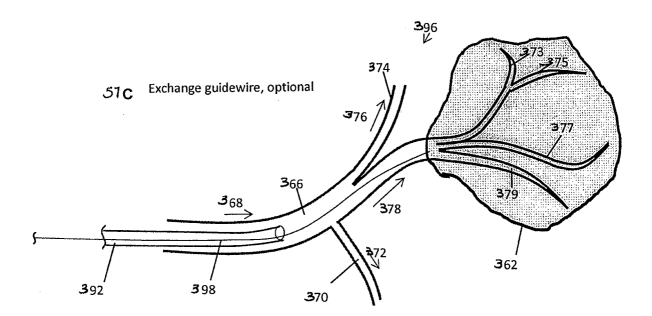
38/47

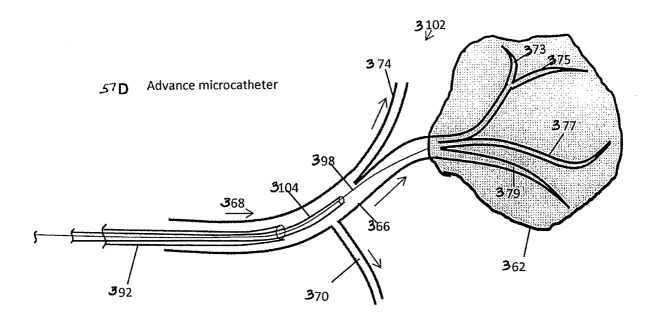
Show normal flow, label arteries, tumor etc.

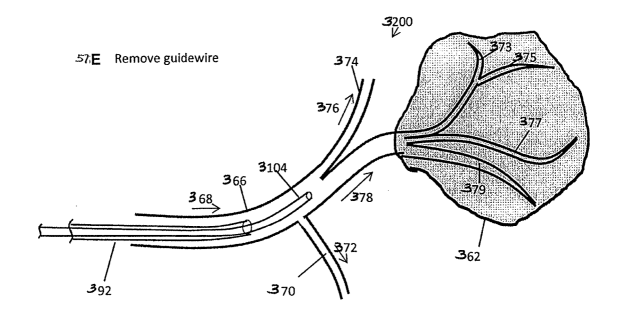




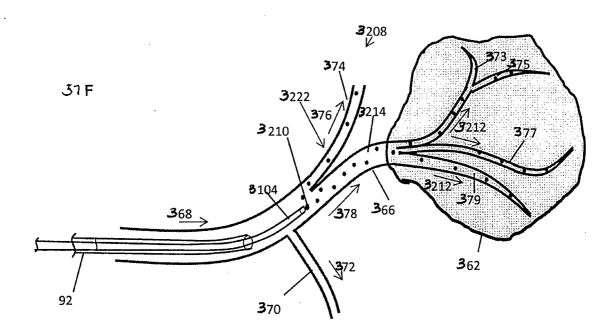


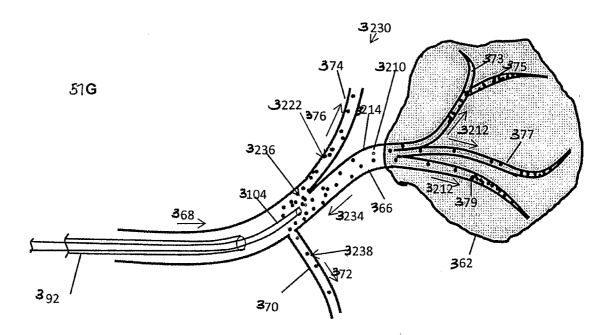






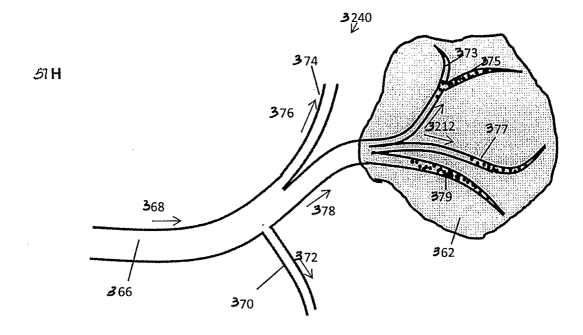
Inject particles

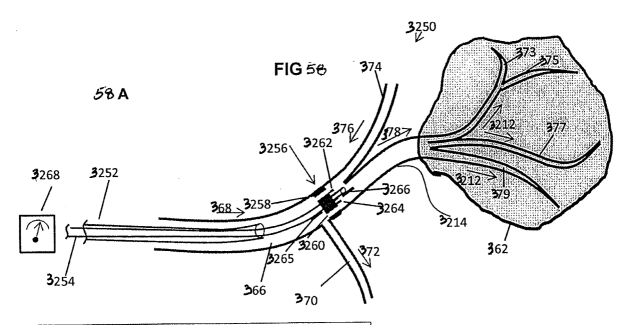




Backpressure signals endpoint, small caps unembolized poor distribution

Stop procedure. Remove guide catheter, microcatheter, poor distribution

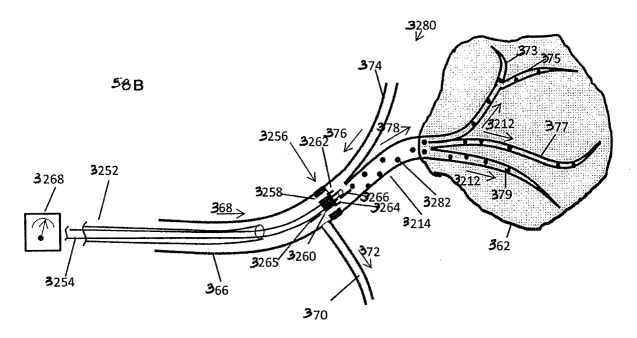


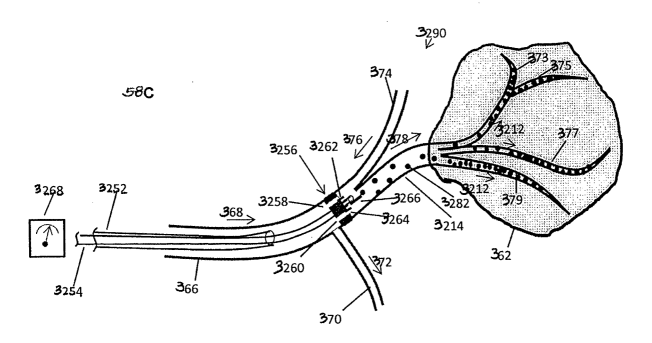


Inflate balloon, partial occlusion, flow regulation, pressure drop, and reversal of flow, check pressure

### 43/47

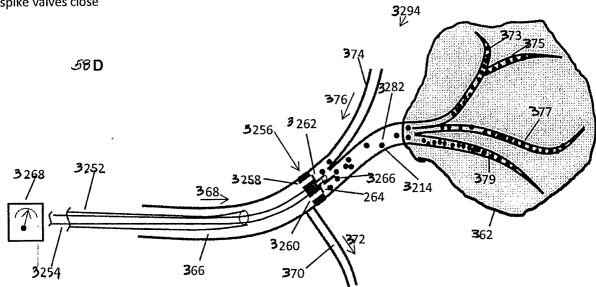
Initiate particle injection, slower particle movement into tumor, monitor pressure

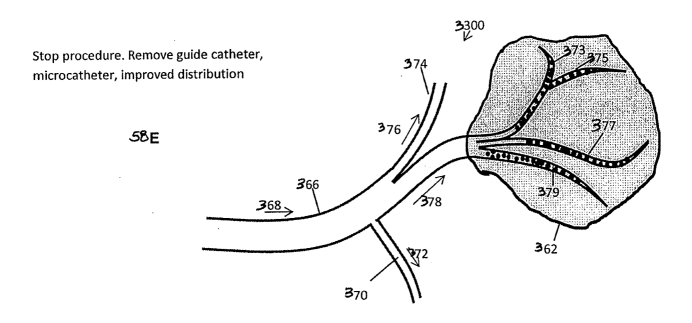




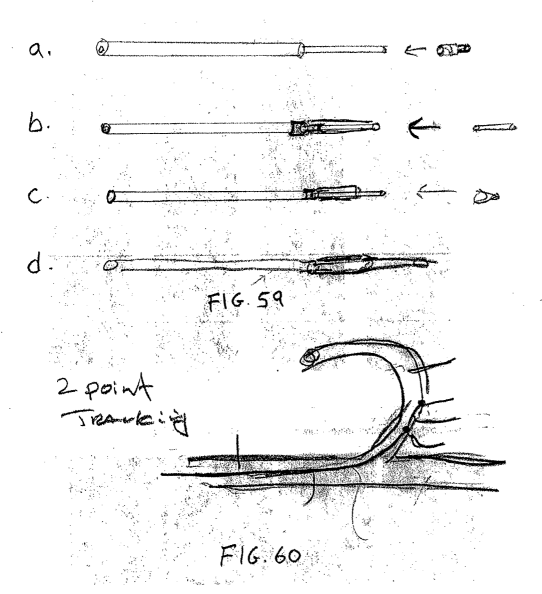
### 44/47

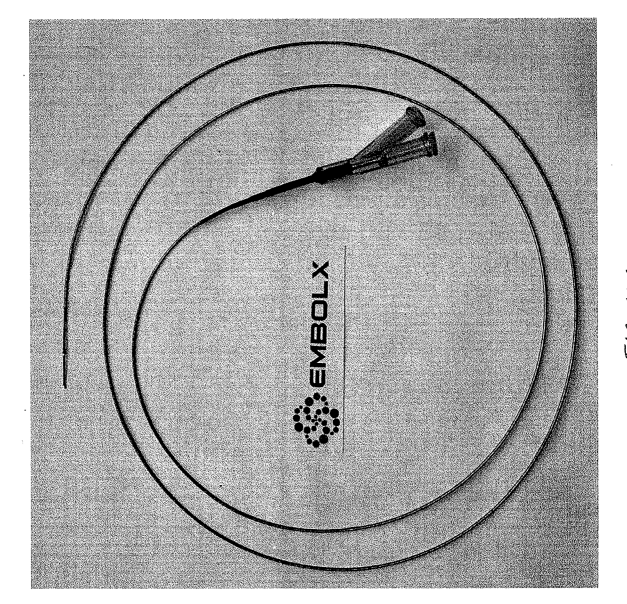
Stable predefined backpressure signals endpoint, large and small caps embolized, improved distribution, monitor pressure during injection keeping below desired level Pressure spike valves close





**END** 





F16.61 A

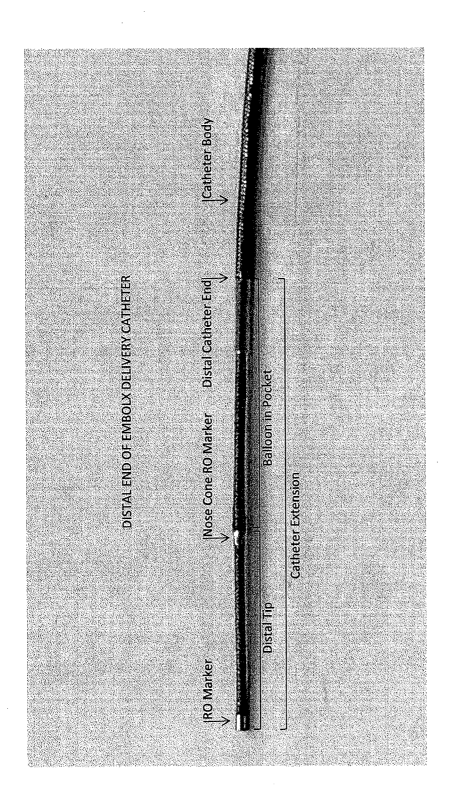


FIG. 618

#### (12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

#### (19) World Intellectual Property Organization

International Bureau

(43) International Publication Date

13 November 2014 (13.11.2014)





(10) International Publication Number WO 2014/182959 A3

- (51) International Patent Classification:
- (21) International Application Number:

PCT/US2014/037385

(22) International Filing Date:

A61M 29/00 (2006.01)

8 May 2014 (08.05.2014)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

61/821,058 8 May 2013 (08.05.2013) US 61/915,425 12 December 2013 (12.12.2013) US 61/917,131 17 December 2013 (17.12.2013) US

- (71) Applicant: EMBOLX, INC. [US/US]; 5760 Arboretum Drive, Los Altos, CA 94024 (US).
- (72) Inventors: ALLEN, Michael, P.; 5760 Arboretum Drive, Los Altos, CA 94024 (US). BROMMER, Robert; 5760 Arboretum Drive, Los Altos, CA 94024 (US). HANNON, Trevor; 5760 Arboretum Drive, Los Altos, CA 94024 (US). DE NEVE, Robert; 5760 Arboretum Drive, Los Altos, CA 94024 (US).
- (74) Agents: LIMBACH, Douglas, C. et al.; SHAY GLENN LLP, 2755 Campus Drive, Suite 210, San Mateo, CA 94403 (US).
- (81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM,

AO, AT, AU, AZ, BA, BB, BG, BH, BN, BR, BW, BY, BZ, CA, CH, CL, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IR, IS, JP, KE, KG, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PA, PE, PG, PH, PL, PT, QA, RO, RS, RU, RW, SA, SC, SD, SE, SG, SK, SL, SM, ST, SV, SY, TH, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW.

(84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LR, LS, MW, MZ, NA, RW, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, RU, TJ, TM), European (AL, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HR, HU, IE, IS, IT, LT, LU, LV, MC, MK, MT, NL, NO, PL, PT, RO, RS, SE, SI, SK, SM, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, KM, ML, MR, NE, SN, TD, TG).

#### Published:

- with international search report (Art. 21(3))
- (88) Date of publication of the international search report:

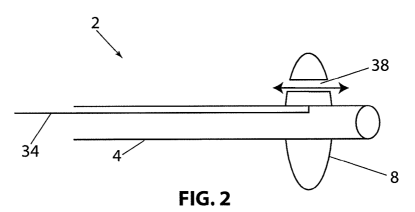
5 March 2015

(15) Information about Correction:

**Previous Correction:** 

see Notice of 15 January 2015

 $\textbf{(54) Title:} \ \ \text{DEVICE AND METHODS FOR TRANSVASCULAR TUMOR EMBOLIZATION WITH INTEGRATED FLOW REGULATION } \\$ 



(57) Abstract: A catheter assembly may be provided with a catheter body and an inflatable balloon. The catheter body has a proximal end, a distal end and a balloon inflation lumen. The inflatable balloon is attachable to the distal end of the catheter body. The balloon has an inner surface that at least partially defines an interior volume. The balloon is configured such that the interior volume can be in fluid communication with the inflation lumen of the catheter body to inflate the balloon. The balloon also has a proximal surface and a distal surface. The balloon is provided with a channel that extends through the balloon. The channel is configured to provide fluid communication between the proximal surface of the balloon and the distal surface of the balloon. Other catheter assemblies and methods of use are also disclosed.



#### INTERNATIONAL SEARCH REPORT

International application No. PCT/US 14/37385

A. CLASSIFICATION OF SUBJECT MATTER IPC(8) - A61M 29/00 (2014.01) CPC - A61M 25/104			
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)			
CPC: A61M 25/104 IPC(8): A61M 29/00 (2014.01)			
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched USPC: 604/102.02, 96.01, 103.01, 103.05, 103.06, 103.07, 103.08; 606/192 (keyword limited; terms below)			
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) PatBase; Google Patents; Google Search Terms Used: catheter, bypass*, passthrough, perfusion, balloon, inflat*, valv*, channel, lumen, conduit, antegrade flow, distal flow			
C. DOCUMENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to claim No.
Х	US 4,909,252 A (GOLDBERGER) 20 March 1990 (20.03.1990) fig 3a, 3b, 3c, col 3, ln 60-62, col 5, ln 10-17, col 6, ln 14-34		1, 3-6
Υ	COT 0, III 10-17, COT 0, III 14-54		2
A			7-14
Y US 5,766,192 A (ZACCA) 16 June 1998 (16.06.1998) f		fig 6	2
Α	US 5,334,154 A (SAMSON et al) 02 August 1994 (02.08.1994) entire document		1-14
	·		
Further documents are listed in the continuation of Box C.			
* Special categories of cited documents:  "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand			
"E" earlier a	to be of particular relevance the principle or theory underlying the invention  "E" earlier application or patent but published on or after the international "X" document of particular relevance; the claimed invention cannot be		
tiling date  "L" document which may throw doubts on priority claim(s) or which is step when the document is taken alone step when the document is taken alone			
special reason (as specified)  "O" document referring to an oral disclosure, use, exhibition or other  "O" document referring to an oral disclosure, use, exhibition or other		step when the document is documents, such combination	
means being obvious to a person skilled in the art  "P" document published prior to the international filing date but later than the priority date claimed document member of the same patent family			
Date of the actual completion of the international search  Date of mailing of the international search report			ch report
18 November 2014 (18.11.2014) 0 1 DEC 2014		:	
	Name and mailing address of the ISA/US  Authorized officer:		
Mail Stop PCT, Attn: ISA/US, Commissioner for Patents P.O. Box 1450, Alexandria, Virginia 22313-1450		Lee W. Young	
Facsimile N	o. <b>571-273-3201</b>	PCT Helpdesk: 571-272-4300 PCT OSP: 571-272-7774	

#### INTERNATIONAL SEARCH REPORT

International application No.

PCT/US 14/37385

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: This application contains the following inventions or groups of inventions which are not so linked as to form a single general inventive concept under PCT Rule 13.1. In order for all inventions to be examined, the appropriate additional examination fees must be paid.			
Group I: Claims 1-14, directed to a catheter assembly			
Group II: Claims 15-19 directed to a method of embolizing a tumor.			
Continued on Supplemental Page			
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 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			
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.			

#### INTERNATIONAL SEARCH REPORT

International application No. PCT/US 14/37385

Continuation of Box III: Observations where unity of invention is lacking

The inventions listed as Groups I-II do not relate to a single general inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons:

Groups I-II are related to an apparatus (Group I) and a method of using the apparatus (Group II), and share the technical features of a catheter body having a proximal end, a distal end and a balloon inflation lumen; and

an inflatable balloon attachable to the distal end of the catheter body, the balloon having an inner surface that at least partially defines an interior volume, the balloon being configured such that the interior volume can be in fluid communication with the inflation lumen of the catheter body to inflate the balloon, the balloon having a proximal surface and a distal surface,

wherein the balloon is provided with a channel that extends through the balloon, the channel configured to provide fluid communication between the proximal surface of the balloon and the distal surface of the balloon.

However, these shared technical features fail to make a contribution over the prior art of US 4,909,252 A (GOLDBERGER), which teaches a catheter assembly comprising:

a catheter body (28) having a proximal end, a distal end and a balloon inflation lumen (15, fig 3b, col 5, ln 10-17)); and an inflatable balloon (36) attachable to the distal end of the catheter body, the balloon having an inner surface that at least partially defines an interior volume, the balloon being configured such that the interior volume can be in fluid communication with the inflation lumen of the catheter body to inflate the balloon, the balloon having a proximal surface and a distal surface (fig 3b, col 5, ln 10-17), wherein the balloon is provided with a channel (32) that extends through the balloon, the channel configured to provide fluid communication between the proximal surface of the balloon and the distal surface of the balloon (fig 3b, col 6, in 14-34).

The special technical feature of Group II is method of embolizing a tumor, the method comprising: advancing a device including a catheter body and a partial occlusion structure to a target tumor site within the body; allowing an anterograde blood flow past the partial occlusion structure, the allowed anterograde blood flow being less than a blood flow that would normally be present if the partial occlusion structure were not in place; injecting an embolic substance from the device to allow the anterograde blood flow to carry the embolic substance into a vasculature of the tumor target; monitoring a real time pressure measurement in a vascular space distal to the partial occlusion structure; signaling a procedural endpoint based on the monitoring of the pressure measurement; and withdrawing the device from the body; not required by the inventions of Group I.

device from the body, not required by the inventions of Group I.

As the common features were known in the art at the time of the invention, they cannot be considered special technical features that would otherwise unify the groups.

Therefore, Groups I-II lack unity under PCT Rule 13 because they do not share a same or corresponding special

#### (12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

#### **CORRECTED VERSION**

(19) World Intellectual Property Organization

International Bureau





(10) International Publication Number WO 2014/182959 A9

(43) International Publication Date 13 November 2014 (13.11.2014)

(51) International Patent Classification: *A61M 25/00* (2006.01) *A61B 17/12* (2006.01) *A61M 5/172* (2006.01)

(21) International Application Number:

PCT/US2014/037385

(22) International Filing Date:

8 May 2014 (08.05.2014)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

61/821,058 8 May 2013 (08.05.2013) US 61/915,425 12 December 2013 (12.12.2013) US 61/917,131 17 December 2013 (17.12.2013) US

- (71) Applicant: EMBOLX, INC. [US/US]; 5760 Arboretum Drive, Los Altos, CA 94024 (US).
- (72) Inventors: ALLEN, Michael, P.; 5760 Arboretum Drive, Los Altos, CA 94024 (US). BROMMER, Robert; 5760 Arboretum Drive, Los Altos, CA 94024 (US). HANNON, Trevor; 5760 Arboretum Drive, Los Altos, CA 94024 (US). DE NEVE, Robert; 5760 Arboretum Drive, Los Altos, CA 94024 (US).
- (74) Agents: LIMBACH, Douglas, C. et al.; SHAY GLENN LLP, 2755 Campus Drive, Suite 210, San Mateo, CA 94403 (US).
- (81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM,

AO, AT, AU, AZ, BA, BB, BG, BH, BN, BR, BW, BY, BZ, CA, CH, CL, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IR, IS, JP, KE, KG, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PA, PE, PG, PH, PL, PT, QA, RO, RS, RU, RW, SA, SC, SD, SE, SG, SK, SL, SM, ST, SV, SY, TH, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW.

(84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LR, LS, MW, MZ, NA, RW, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, RU, TJ, TM), European (AL, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HR, HU, IE, IS, IT, LT, LU, LV, MC, MK, MT, NL, NO, PL, PT, RO, RS, SE, SI, SK, SM, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, KM, ML, MR, NE, SN, TD, TG).

#### Published:

- without international search report and to be republished upon receipt of that report (Rule 48.2(g))
- (48) Date of publication of this corrected version:

15 January 2015

(15) Information about Correction: see Notice of 15 January 2015



(54) Title: DEVICE AND METHODS FOR TRANSVASCULAR TUMOR EMBOLIZATION WITH INTEGRATED FLOW REGULATION

(57) Abstract: A catheter assembly may be provided with a catheter body and an inflatable balloon. The catheter body has a proximal end, a distal end and a balloon inflation lumen. The inflatable balloon is attachable to the distal end of the catheter body. The balloon has an inner surface that at least partially defines an interior volume. The balloon is configured such that the interior volume can be in fluid communication with the inflation lumen of the catheter body to inflate the balloon. The balloon also has a proximal surface and a distal surface. The balloon is provided with a channel that extends through the balloon. The channel is configured to provide fluid communication between the proximal surface of the balloon and the distal surface of the balloon. Other catheter assemblies and methods of use are also disclosed.

# DEVICE AND METHODS FOR TRANSVASCULAR TUMOR EMBOLIZATION WITH INTEGRATED FLOW REGULATION

#### CROSS REFERENCE TO RELATED APPLICATIONS

5 [0001] This application claims the benefit of U.S. Provisional Applications No. 61/821,058 filed May 8, 2013, No. 61/915,425 filed December 12, 2013, and No. 61/917,131, filed December 17, 2013, each of which is herein incorporated by reference in its entirety.

#### INCORPORATION BY REFERENCE

10 [0002] All publications and patent applications mentioned in this specification are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

#### **FIELD**

[0003] This application relates generally to medical methods and devices. More specifically, the present application discloses various embodiments of occlusion devices adapted to a catheter, and methods for their use in delivering fluids, embolic materials and other therapeutic agents to sites within the body.

20 BACKGROUND

25

30

[0004] There are over one million cases of cancer diagnosed each year in the United States and numerous approaches of therapy including systemic chemotherapy, radiation and surgical resection. Given that systemic chemotherapy and radiation interact with healthy tissue, complications and toxicity often result. Targeted drugs are now being used and produce a lower rate of complications. Ablative approaches, including microwave, radiofrequency and cryogenic therapies have been used; however, these methods are often not selective and tissues and organs surrounding or below the tumor can be affected.

[0005] According to the National Institute of Health, 30,640 people were diagnosed with primary liver cancer (hepatocellular carcinoma, HCC) and 142,820 people were diagnosed with colorectal cancer in the U.S. in 2013. Seventy five percent of these will metastasize to the liver. Liver resection and transplant are the only curative means; however, only small numbers of patients are eligible. Systemic chemotherapy for primary and metastatic tumors in the liver is ineffective, having a response rate of about 20% and a survival benefit of 10.7 months vs. 7.9 months over symptomatic care.

[0006] Catheters are commonly used in medicine for delivery of fluids, therapeutics, and implants, and in sampling tissues and bodily fluids. Catheters can be constructed with balloons or other tools to dilate tissue, block fluid flow or isolate segments of the anatomy, such as in treatment of the cancers described above.

- 5 [0007] Trans-Arterial Embolization therapy is the transvascular injection of drug and/or embolic agents directly into the tumor vasculature using a microcatheter. Embolization therapy causes a shutdown of blood flow and, when drug or radioactivity is present, simultaneous release of high concentrations of drug or radioactivity. The technique is also noted for its very low level of toxicity.
- [0008] In the early 1980's, transarterial chemoembolization (TACE) began to be used as a selective cancer therapy. In this method, chemotherapeutic and embolic agents are injected directly into the vasculature of the tumor, a technique that is most common for the treatment of hepatocellular carcinoma. More recently, transarterial radioembolization (TARE) has been used clinically. In this method, radioactive embolic particles, typically yttrium-90 (y90), are injected rather than chemotherapeutic agents. Although the liver is a common target for TACE and TARE, other organs, including, but not limited to, the pancreas, lung, kidney, prostate, stomach, colon and head and neck have been treated using these methods. Chemoembolization was established as a standard of care for intermediate stage hepatocellular carcinoma in 2006.
  [0009] Numerous studies have demonstrated transarterial embolization to be effective on a number of primary cancers and to have better performance than chemotherapy for both HCC and

number of primary cancers and to have better performance than chemotherapy for both HCC and metastatic colorectal cancers in the liver; however, studies show inconsistent outcomes with reported tumor responses from 15% to 85%. Although anatomical and individual differences are clearly of significance in between-patient variation, clinical studies, each of which include a range of patients, show very different outcomes, indicating that there currently is little procedural optimization or standardization.

25

30

35

[0010] The above procedures are accomplished by inserting a small catheter into the femoral artery at the groin and navigating it into the liver vasculature, typically the hepatic artery, then into the right or left lobe of the liver or more selectively into particular segments of the liver or super-selectively directly into or adjacent to the tumor. Super-selective transarterial delivery of antitumor agents into the tumor vasculature has become state-of-the-art and requires catheters that can reach into small vessels. Presently, standard microcatheters, typically at or about 3 Fr are used to inject antitumor agents into the target vasculature. These standard microcatheters rely on normal blood flow as the means by which the embolic agent moves into the tumor and systolic pressure as the packing force. However, the injection pressure is typically higher than the blood pressure and blood flow can be reversed. When this happens, the cancer agent flows in a

retrograde direction with respect to normal blood flow and away from the tumor, with a concomitant risk of delivery of the anti-cancer therapy to organs that can be damaged by these toxic agents. This situation also results in loss of an unknown amount of drug.

5

10

15

20

25

30

35

[0011] The endpoint of the above procedures is determined by physicians' visual observation and can range between fully embolized to partially embolized with the amount of dose delivered being highly variable. Retrograde and anterograde reflux, distribution, packing, quantity of dose delivered and procedure endpoint are variables that can be highly dependent on the rate and pressure of injection, the selection of the type of endpoint, the patient's systolic pressure and the operator. As such, clinical trials using TACE to treat hepatocellular carcinoma have demonstrated wide variations in tumor response. The most significant problems that occur with the current means of delivery and methods of embolization therapy include inconsistent efficacy and non-target embolization.

[0012] Using standard straight-tip catheters, non-target embolization in the retrograde direction can be caused when the pressure of injection exceeds the systolic blood pressure and the embolic agents flow backwards over the catheter and into the general circulation. Anterograde reflux and non-target embolization occurs when the embolic agents flow into distal vasculature, through arteriovenus shunting and into the venous circulation. This can easily occur because venous blood pressure on average is about 10 to 15 mmHg as compared to arterial diastolic blood pressure of about 80 mmHg.

[0013] When therapeutic agents are delivered into the vasculature of a target structure using the normal anterograde blood flow to carry the therapy to the target, injection rate and pressure of the therapy must be carefully controlled in relation to the flow volume and pressure of blood to avoid retrograde reflux of drug backward over the catheter and into the general circulation. In particular, when injecting embolic agents into the vasculature of a tumor, pressure distal to the catheter tip continues to increase as embolization progresses, causing a resistance that prevents embolic agents from filling the target vasculature and the possibility of retrograde reflux and non-target embolization. It would be desirable to eliminate this retrograde reflux, non-target embolization, and the inconsistent dosages that are delivered to targets with current state of the art procedures. It would be further desirable to eliminate the low levels of particle distribution and density throughout the target vasculature. It would be still further desirable to replace current delivery devices that are not always capable of fully isolating the target vasculature and often do not allow the operator to control pressure, flow rate and other parameters associated with therapeutic delivery.

[0014] The present state-of-the-art embolization therapy for tumors in the liver relies on high volume forward flow from the hepatic artery to deliver embolization agents into the tumor.

However; distal embolization of larger capillaries causes: (1) high intra-tumor vascular pressure, (2) high pressure in arteries feeding the tumor, (3) proximal reflux backwards over the delivery catheter, (4) increased anterograde bypass in distal hepatoenteric arteries and (5) poor filling and distribution of embolic agents in the tumor. This situation results in an uncontrollable number of particles entering the tumor and procedural high variability.

5

10

15

20

25

[0015] Problems with the current method of embolization therapy that cause inconsistent outcomes include: variable procedural endpoints, unknown quantity of dose delivered, reflux of embolization agents into the general circulation, anterograde bypass of embolization particles into the general circulation, non-target embolization, rising intra-tumor arterial pressures during the initial stages of embolization and catheter movement during injection. The current delivery catheters are unable to control many of the above mentioned variables, making any standardization of the current procedures difficult or impossible to achieve.

[0016] The following patents and published patent applications provide some examples of the current state of this art. U.S. Pat. No. 5,647,198 describes a catheter with a pair of spaced apart balloons that define an intra-balloon space. A lumen passes through the catheter and exits within the intra-balloon space allowing injection of drugs, emulsions, fluids and fluid/solid mixtures. A perfusion lumen or bypass extends from a location proximal to the proximal balloon and to the distal tip to allow shunting of blood past the inflated balloons. U.S. Pat. No. 5,674,198 describes a two balloon catheter that is designed for treating a solid tumor. The balloons are positioned to isolate the blood flow into the tumor and allow injection of a vaso-occlusive collagen material to block the tumor blood supply. Clifton et al. (1963) Cancer 16:444-452 describes a two balloon catheter for the treatment of lung carcinoma. The four lumen catheter includes a lumen for independent injection in the space between the balloons. Rousselot et al. (1965) JAMA 191:707-710 describes a balloon catheter device for delivering anticancer drugs into the Liver. See also U.S. Pat. No. 6,780,181; U.S. Pat. No. 6,835,189; U.S. Pat. No. 7,144,407; U.S. Pat. No. 7,412,285; U.S. Pat. No. 7,481,800; U.S. Pat. No. 7,645,259; U.S. Pat. No. 7,742,811; U.S. App. No. 2001/008451; U.S. App. No. 2001/0041862; U.S. App. No. 2003/008726; U.S. App. No.

2008/0208118; U.S. App. No. 2009/0182227 and U.S. App. No. 2010/0114021.
 [0017] What is needed and is not provided by the prior art is a delivery system and method that enable optimization and standardization of treatment delivery, such as by delivering a known quantity of embolic agent to a prescribed target area, and elimination of non-target embolization.

2003/0114878; U.S. App. No. 2005/0267407; U.S. App. No. 2007/0137651; U.S. App. No.

### SUMMARY OF THE DISCLOSURE

PCT/US2014/037385

WO 2014/182959

5

10

15

20

25

30

35

[0018] According to aspects of the present disclosure, devices and methods are provided for partial occlusion with unidirectional bypass flow, designed to be adapted to a catheter for delivery of therapeutic agents to a target site within the body. Such delivery devices may be intended for any medical purpose, but the embodiments described herein are focused on devices intended for performing transarterial delivery of therapeutic agents to a target site within the body. The entry point for the delivery catheter can be any arterial access point, typically the femoral artery located at the groin. The target can be any structure; however, of particular interest are tumors, primary or metastatic, of any organ or tissue that is accessible by a microcatheter through the arterial system. Cancers of particular interest include, but are not limited to, primary and metastatic cancers in the liver, pancreas, colon, rectum, kidney, stomach, lung, bladder, head and neck and uterus. Procedures that can benefit from the access and delivery methods and devices of the present disclosure include, but are not limited to, transarterial chemoembolization using drug eluting beads (DEB TACE), transarterial chemoembolization using Lipiodol (Lipiodol TACE), transarterial radioembolization (TARE) and transarterial embolization (TAE). Other procedures which can benefit from methods and devices of the present disclosure include direct delivery of chemotherapy or targeted drugs to the site of the cancer, the general delivery of drugs, venous or arterial embolization or other substances to specific regions of the body and drainage or aspiration of fluid or tissue. [0019] In some embodiments, microcatheter methods and devices disclosed herein create a partial occlusion (in the hepatic artery) with concomitant reduction of pressure and flow in the vascular compartment distal to the occlusion. This can: (1) eliminate proximal reflux, (2) reverse hepatoenteric arterial flow distal to the occlusion, (3) limit or stop anterograde bypass, (3) reduce flow rate into the tumor, (4) delay the onset of high intra-tumor pressure, (5) increase the extent of embolization and (6) enable a quantitative pressure measurement as an endpoint. [0020] In some embodiments, the device includes an occlusion structure adapted to the distal section of a two lumen catheter, whereby the occlusion structure has one or more channels disposed from its proximal surface to its distal surface allowing flow therethrough. The channels are configured to permit a controllable arterial or venous flow which is less than the un-occluded flow of the artery or vein, and the channel(s) may have one-way valve(s) that allow flow in only one direction. The flow rate can be any amount as compared to the normal un-occluded flow rate from 0% flow up to 100% flow, most typically from 85% flow to 95% flow. The flow

channels and/or valves can be configured so as to regulate flow and/or pressure from one side of the occlusion structure to the other. The flow and/or pressure regulation can be static or variable, permitting a single continuous flow rate or multiple flow rates that are pre-calibrated or adjusted

by the operator. Of particular interest in some embodiments is arterial flow regulation to a constant flow rate that is pre-calibrated.

5

10

15

20

25

30

35

[0021] In some embodiments, the flow rate reduction results in a pressure drop in the vascular space distal to the partial occlusion. When this occurs, distal arterial side branches of the occluded artery that are flowing away from the occluded artery, and part of arterial networks, may reverse direction and begin to flow toward the occluded artery. By way of example, embolization of tumors in the right lobe of the liver are accessed by a catheter advanced through the right hepatic artery (RHA) and to the vicinity of the tumor. Typically, the catheter tip does not enter the tumor vasculature and remains proximal to the tumor and within the right hepatic artery or branch thereof. In this example, the artery feeding the tumor is typically a branch of the RHA. However, there other distal hepatoenteric arteries that branch from the RHA and flow away from the RHA and to the liver and gastrointestinal tract. In this instance, when using a standard straight catheter, injection of embolic agents from the distal tip of the catheter results in flow of embolic agents into both the tumor and side branch arteries causing anterograde bypass and non-target embolization of the liver and gastrointestinal tract, a situation that causes toxicity and complications.

[0022] In some embodiments of the present disclosure, the blood pressure drop in the arterial space distal to the partial occlusion device causes the branch hepatoenteric arteries to reverse flow and now flow toward the tumor. This situation prevents anterograde bypass, reduces non-target embolization and focuses substantially all injected drug and/or embolic agents into the tumor.

[0023] In some embodiments of the present disclosure, a partial occlusion device comprises a balloon with flow regulating channels and one-way valves which, when inflated in an artery, permits limited anterograde flow from vascular space proximal to the partial occlusion to the vascular space distal to the partial occlusion and prevents retrograde flow. So constructed, the device will significantly reduce the hepatic arterial flow into the tumor which is the tumor's main source of blood flow. In this instance, injected anti-cancer agents will flow into the tumor at a significantly slower rate and delay the onset of a high pressure in tumor vessels and retrograde blood flow toward the catheter tip. This slower filling of the tumor will improve the amount and distribution of embolic agents in the tumor. The lower pressure distal to the occlusion will cause the reversal of flow in distal hepatoenteric arteries during embolization and limit anterograde bypass, and the one-way valves will eliminate retrograde bypass of embolic agents into proximal branch arteries such as the gastroduodenal artery.

[0024] By isolating the distal arterial space that is adjacent to the tumor from the arterial blood supply, the device of the present disclosure enables pressure measurement to be used to signal a

procedural endpoint at a predetermined pressure or pressures. By way of example, the endpoint of the procedure can occur at a point when systolic pressure (120 mmHg) is first reached or at a point when systolic pressure is stabilized, however any pressure, pressure profile or algorithm can be used to determine an endpoint of the procedure. Such a measurable endpoint can contribute to standardization of the procedure and improved efficacy.

5

10

15

20

25

30

35

[0025] The occlusion structure of the device of the present disclosure may be held within a pocket within the catheter such that the outer diameter of the radially constrained occlusion structure is approximately equal to or less than the outer diameter of the catheter as described in co-pending U.S. provisional patent applications 61/821058, 61/915425 and 61/917,131. The pocket can be a longitudinal space in the catheter and can be formed as a reduction in the catheter diameter of a defined length and a depth equal to or greater than the thickness of the occlusion structure in a radially constrained configuration. Alternately, a pocket can be formed using an extension projecting distally beyond the catheter body, the distal extension having a diameter smaller than the catheter body. In this instance, the distal end of the catheter pocket is defined by the proximal end of a nose-piece. In some embodiments, the nose-piece has a diameter equal to or less than the diameter of the catheter body and is positioned over the distal extension at a defined distance from the distal end of the catheter body.

[0026] The occlusion structure of devices of the present disclosure can be advanced in a radially constrained configuration, to at least the proximity of a target within the body and then placed in its radially expanded configuration. Alternately, the device can be pre-formed in a fully expanded configuration, adapted to the distal end of a catheter and delivered to the target site. The valves are typically configured to allow proximal to distal (anterograde) bypass flow; however the opposite is possible or the valves may be absent allowing two way flow.

[0027] In some embodiments, a catheter assembly may be provided with a catheter body and an inflatable balloon. The catheter body has a proximal end, a distal end and a balloon inflation lumen. The inflatable balloon is attachable to the distal end of the catheter body. The balloon has an inner surface that at least partially defines an interior volume. The balloon is configured such that the interior volume can be in fluid communication with the inflation lumen of the catheter body to inflate the balloon. The balloon also has a proximal surface and a distal surface.

The balloon is provided with a channel that extends through the balloon. The channel is configured to provide fluid communication between the proximal surface of the balloon and the distal surface of the balloon.

[0028] In some embodiments, a device for delivering a therapeutic agent to a target site within a body is provided. The device comprises a catheter body having a proximal end, a distal end, a first axial lumen and a second axial lumen. The first axial lumen extends from the proximal end

of the catheter body to the distal end of the catheter body and provides fluid communication therebetween. The second axial lumen extends from the proximal end of the catheter body to a more distal location on the catheter body. The device further comprises a balloon radially disposed near the distal end of the catheter body. The balloon has a proximal balloon surface, a distal balloon surface, a radially constrained configuration and a radially expanded configuration. The balloon is in fluid communication with the second axial lumen and has at least one channel extending from the proximal balloon surface to the distal balloon surface, thereby providing fluid communication therebetween. The device also comprises a balloon sheath, disposed over an outer surface of the balloon. The sheath has a proximal side and a distal side. The proximal side has an opening that is in alignment with a proximal end of the balloon channel. The distal side of the sheath extends over a distal end of the balloon channel. The distal side of the sheath is capable of deflecting distally when a distally directed fluid pressure is applied through the opening in the proximal side, through the balloon channel and against a proximal surface of the distal side of the sheath, thereby allowing fluid to pass through the balloon channel and into a vascular compartment distal to the balloon and the balloon sheath. The distal side of the sheath is also capable of deflecting proximally onto the distal balloon surface when a pressure is applied to the distal side of the sheath, thereby preventing proximally directed flow in the balloon

[0029] In some embodiments, a method of embolizing a tumor is provided. The method comprises advancing a device including a catheter body and a partial occlusion structure to a target tumor site within the body, and allowing an anterograde blood flow past the partial occlusion structure. The allowed anterograde blood flow is less than a blood flow that would normally be present if the partial occlusion structure were not in place. The method further comprises injecting an embolic substance from the device to allow the anterograde blood flow to carry the embolic substance into a vasculature of the tumor target. The method also comprises monitoring a real time pressure measurement in a vascular space distal to the partial occlusion structure, signaling a procedural endpoint based on the monitoring of the pressure measurement, and withdrawing the device from the body.

[0030] While aspects of the present disclosure will be described with particular reference to delivery of chemotherapeutic agents, radiotherapeutic agents, embolic agents or combinations thereof into the vasculature that supplies blood to a tumor, the same principles apply to the delivery or aspiration of a variety of materials into or from other locations, and through other luminal structures in the body.

5

10

15

20

25

### BRIEF DESCRIPTION OF THE DRAWINGS

- [0031] The novel features of the disclosure are set forth with particularity in the claims that follow. A better understanding of the features and advantages of the present disclosure will be obtained by reference to the following detailed description that sets forth illustrative embodiments, in which the principles of the disclosure are utilized, and the accompanying drawings of which:
  - [0032] FIG. 1A, 1B and 1C illustrate embodiments of the disclosure herein;
  - [0033] FIG. 2 illustrates a distal cross sectional view of a bidirectional embodiment;
- 10 [0034] FIG. 3 illustrates a unidirectional embodiment;

- [0035] FIG. 4 illustrates a unidirectional embodiment with balloon inflation and guidewire/injection lumens;
- [0036] FIG. 5 illustrates an embodiment of the present disclosure including a distal tip and adapter;
- 15 [0037] FIG. 6 illustrates an embodiment of the present disclosure with a bidirectional channel within the catheter;
  - [0038] FIG. 7 illustrates an embodiment of the present disclosure with a unidirectional channel;
  - [0039] FIG. 8 illustrates a device of the present disclosure with a distal tip;
  - [0040] FIG. 9A shows a linear cross section through the catheter and balloon;
- 20 [0041] FIG. 9B shows a linear cross section through the catheter and balloon and two circular bidirectional channels;
  - [0042] FIG. 9C shows a linear cross section through the catheter and balloon and three circular bidirectional channels;
  - [0043] FIG. 9D shows a linear cross section through the catheter and balloon and two circular unidirectional bypass channels with individual valves on each channel;
    - [0044] FIG. 9E shows a linear cross section through the catheter and balloon and three circular unidirectional bypass channels with individual valves on each channel;
    - [0045] FIG. 9F shows a linear cross section through the catheter and balloon and two circular unidirectional channels with a one-piece valve covering both channels;
- 30 [0046] FIG. 9G shows a linear cross section through the catheter and balloon and two circular unidirectional channels with a one-piece valve covering three channels;
  - [0047] FIG. 9H shows a linear cross section through the catheter and balloon and four bidirectional bypass channels formed from pleats in the balloon;

[0048] FIG. 9I shows a linear cross section through the catheter and balloon and four unidirectional bypass channels formed from pleats in the balloon and a one-piece valve covering all 4 channels;

- [0049] FIG. 9J shows a linear cross section through the catheter and balloon and four
- 5 bidirectional channels which radiate outward from the catheter;
  - [0050] FIG. 9K shows a linear cross section through the catheter and balloon and four unidirectional channels which radiate outward from the catheter and a one-piece valve covering all four valves;
- [0051] Fig. 9L shows a linear cross section through the catheter and balloon and four unidirectional channels which radiate outward from the catheter outward and a single cross shaped valve covering all four channels;
  - [0052] FIG. 10 shows a tumor and associated vasculature;
  - [0053] FIG. 11 shows an expanded view of a tumor and its vasculature including blood flow direction;
- [0054] FIG. 12 illustrates a standard micro-catheter over a guidewire inside the tumor vasculature with injection;
  - [0055] FIG. 13 illustrates a catheter of the present disclosure over a guidewire and inside a main artery
  - [0056] FIG. 14 illustrates a catheter of the present disclosure over a guidewire and inside the branch artery;
    - [0057] FIG. 15 illustrates a catheter of the present disclosure with inflated balloon and closed valves;
    - [0058] FIG. 16 illustrates a catheter of the present disclosure with inflated balloon and open valves;
- 25 [0059] FIG. 17 illustrates a catheter of the present disclosure with inflated balloons, open valves and initiation of fluid injection;
  - [0060] FIG. 18 illustrates a catheter of the present disclosure during injection with inflated balloons and valves closed;
- [0061] FIG. 19 illustrates a catheter of the present disclosure at a period of time following the completion of injection with inflated balloon and open valves;
  - [0062] FIG. 20 shows the deflation of the balloon;

- [0063] FIG. 21 shows the withdrawal of the catheter into a main artery;
- [0064] FIG. 22 shows an embodiment of the present disclosure with a channel within the catheter and closed valve;

[0065] FIG. 23 shows an embodiment of the present disclosure with a channel within the catheter and open valve.

- [0066] FIG 24A illustrates a standard microcatheter;
- [0067] 24B, 24C, 24D and 24E illustrate a cross sectional view of a sequential construction of an embodiment of the present disclosure;
  - [0068] FIG. 25 illustrates a cross sectional view of the distal portion of one embodiment of the device according to the present disclosure;
  - [0069] FIG 26 illustrates a view of an embodiment including an occlusion balloon concealed within pocket formed between proximal and distal surfaces;
- 10 **[0070]** FIG. 27A and 27B illustrate a distal catheter with and occlusion balloon unexpanded and expanded;
  - [0071] FIG. 28A, 28B, 28C, and 28D illustrate an embodiment of the present disclosure including a two layer catheter;
  - [0072] FIG. 29 gives examples of balloons used in various embodiments of the present
- 15 disclosure;

- [0073] FIG. 30 shows a front view of a balloon with one-way bypass channels and valves;
- [0074] FIG. 31 shows a cross section through the balloon of Fig 30;
- [0075] FIG. 32 illustrates a cross sectional view of a sequential construction of an embodiment of the present disclosure including a balloon pocket and integral nose cone;
- 20 [0076] FIG. 33 illustrates a cross sectional view of a sequential construction of an embodiment of the present disclosure including a balloon pocket and separate nose cone;
  - [0077] FIG. 34 illustrates an embodiment of the present disclosure with two balloons;
  - [0078] FIG. 35 illustrates an embodiment of the present disclosure including a balloon with valves;
- 25 [0079] FIG. 36 illustrates an embodiment of the present disclosure with pressure sensors;
  - [0080] FIG 37A, 37B, 37C and 37D illustrate an embodiment of the present disclosure with a balloon or balloons placed on the circumference of the catheter;
  - [0081] FIG. 38A and 38B illustrate a tumor with vascular anatomy and embolization using a standard straight nose catheter;
- 30 [0082] FIG. 39A and 39B illustrate a tumor with vascular anatomy and embolization using a balloon including channels and valves;
  - [0083] FIG 40A, 40B, 40C and 40D illustrate a tumor with vascular anatomy and embolization using an occlusion balloon;
  - [0084] FIG. 41 illustrates constructions of embodiments of the present disclosure;
- 35 [0085] FIG. 42 illustrates a two layer occlusion structure with unidirectional flow;

- [0086] FIG. 43 illustrates a prototype of the occlusion structure of FIG 42;
- [0087] FIG. 44 illustrates an embodiment of the present disclosure including a catheter with a pocket;
- [0088] FIG. 45 illustrates embodiments of the present disclosure for complete occlusion;
- 5 [0089] FIG. 46 illustrates an embodiment with bidirectional and unidirectional channels;
  - [0090] FIG 47 illustrates valve constructions of an embodiment of a unidirectional occlusion structure of the present disclosure;
  - [0091] FIG. 48 shows a prototype micro-valve;
  - [0092] FIG. 49 shows an alternate embodiment of an occlusion structure;
- 10 [0093] FIG. 50 illustrates a method of operation of one embodiment of a unidirectional occlusion structure of the present disclosure;
  - [0094] FIG. 51 illustrates a method for delivery of embolic particles using a device of the present disclosure;
  - [0095] FIG. 52 illustrates a partial occlusion balloon with channel and valve;
- 15 [0096] FIG. 53 illustrates a side view of constrained occlusion balloon in catheter pocket;
  - [0097] FIG. 54 illustrates a distal end construction including pocket, constrained balloon and expanded partial occlusion balloon with channels and valves in closed position;
  - [0098] FIG. 55 illustrates a serial construction of a full length catheter device;
  - [0099] FIG. 56 illustrates liver vasculature and associated tumor vasculature;
- 20 [0100] FIG. 57 illustrates a tumor embolization method for a standard catheter;
  - [0101] FIG. 58 illustrates a tumor embolization method for the catheter of the present disclosure;
  - [0102] FIGS. 59A, 59B, 59C and 59D illustrate fabrication steps of the distal end of an occlusion catheter according to aspects of the present disclosure;
  - [0103] FIG. 60 illustrates the completed distal end of the occlusion catheter of FIGS. 59A 59D being introduced into small branches of a vascular system;
  - [0104] FIG. 61A shows an entire occlusion catheter constructed according to principles of the present disclosure; and
  - [0105] FIG. 61B shows details of construction features on the distal end of the occlusion catheter of FIG. 61A.

30

35

25

#### **DETAILED DESCRIPTION**

[0106] The device of the present disclosure allows improved distribution of anti-cancer agents into target tumor vasculature by reducing arterial flow and pressure during drug and/or embolic agent injection. The present device reduces toxicity and complications by eliminating reflux of embolic materials and/or anti-cancer agents into proximal arterial branches and reduces or

eliminates anterograde bypass of embolic materials and/or anticancer agents into distal arterial branches. Further, the present disclosure enables pressure measurement as a means to a quantitative endpoint of the procedure. Such a device can improve efficacy and reproducibility of the technique and reduce complications.

5

10

15

20

25

30

35

[0107] Referring to FIG. 1A, a longitudinal cross section of device 2 of the present disclosure is shown with catheter body 4, distal tip 6, balloon 8 (unexpanded configuration), balloon inflation tube 10, guidewire and injection tube 12 and fittings 14. Catheter body 4 can have a length of 10 cm to 400 cm, typically 60 cm to 250 cm and a diameter of 0.25 mm to 5 mm, typically 0.5 mm to 1.5 mm. Device 2 may or may not include a distal tip 6, the distal tip having a length of 1 mm to 50 mm, more typically from 5 mm to 30 mm. The balloon inflation tube 10, positioned at the proximal end of catheter body 4, is connected to, and in fluid communication with a balloon inflation lumen that runs longitudinally through the length of catheter body 4 and terminates at, and is in fluid communication with balloon 8. The guidewire and injection tube 12, positioned at the proximal end of catheter body 4, is connected to, and in fluid communication with a lumen that runs longitudinally through the length of catheter body 4 and terminates at the distal end or distal tip of catheter body 4, thereby allowing a guidewire to enter through fitting 14 and exit through the distal end of device 2 through catheter body 4. Fittings 14 are connected to each of balloon inflation tube 10 and guidewire and injection tube 12 and can connect to a syringe, inflation device or any other device or means to inject air, gas, fluid, suspensions, emulsions, contrast, therapeutic agents, embolic agents or any other material capable of being injected through balloon tube 10 or guidewire tube 12 and longitudinal lumens that run to the balloon or distal end of device 2.

[0108] Referring to FIG 1B, a longitudinal cross section of a first embodiment of the present disclosure is shown, with device 2, balloon 8 (in the expanded configuration) having valve 16 in the open position and valve 18, in the closed position. In this embodiment flow channels 20 and 22 are constructed through balloon 8. Valves 16 and 18 allow fluid to flow in only one direction. Balloon 8 has a proximal side 9 and a distal side 11. By way of example, if fluid pressure is higher on the proximal side of balloon 8 and lower on the distal side of balloon 8, both valves 16 & 18 will open in response to the pressure difference and allow fluid to flow distally through the valves. If the pressure is higher on the distal side of balloon 8, valves 16 and 18 will close and prevent fluid from flowing proximally. Alternately, the valves can be position or constructed so that fluid can pass proximally and be prevented from flowing in the distal direction. Valves 16 and 18 are shown as a simple "flap" type valve, however, they can be any type of valve, such as a diaphragm that open and close in response to a pressure differential. Balloon 8 is shown with two channels and two valves; however there can be 1, 2, 3 or more channels and/or valves.

Device 2 of this embodiment may include channels and may or may not include valves. If valves are not included, a bidirectional flow will result.

[0109] Referring to FIG. 1C, a longitudinal cross section of another embodiment of the present disclosure is shown with device 24 and channel 26 running through and within catheter body 28.

- 5 Balloon 30 has a proximal side 13 and a distal side 15. Channel 26 extends from the proximal side of balloon 30 to the distal side of balloon 30. A valve 32 is illustrated over channel 26 on the distal side of balloon 30, however, if desired the valve can be positioned on the proximal side of channel 26 and balloon 30. The function and operation of valve 32 of this embodiment of the present disclosure is identical to that presented in FIG 1A and FIG 1B. As in this embodiment, if valve 32 is not included, a bidirectional flow will result.
  - [0110] FIG. 2 illustrates an exemplary embodiment of a longitudinal cross section of the distal end of device 2 with catheter body 4, balloon inflation lumen 34, balloon 8, and channel 38. Balloon inflation lumen 34 extends from the proximal end of catheter body 4 and exits at balloon 8. In this case, a valve in balloon 8 is not included and a bidirectional flow will result. The optimal balloon inflation lumen diameter is 0.1 mm to 0.5 mm; however this lumen can be in the range of 0.25 mm to 1 mm.

15

20

- [0111] FIG. 3 illustrates an example of a longitudinal cross section of the distal end of device 2 with catheter body 4, balloon inflation lumen 34, balloon 8, channel 38 and valve 16. Valve 16 is shown over the distal opening of channel 38 in the closed position, however if pressure is applied to the proximal valve surface through channel 38, the valve will allow fluid to pass distally. The valve 16 will prevent proximal flow. The valve can be positioned at the proximal or distal opening or anywhere within the cannel. Location and configuration of the valve will determine flow direction.
- [0112] Referring to FIG. 4, a longitudinal cross section of the distal end of device 2 is shown with catheter body 4, balloon inflation lumen 34, balloon 8, channel 38 and guidewire/injection lumen 40. Guidewire/injection lumen 40 extends from the proximal end of catheter body 4 and exits at the distal end of catheter body 4 or distal tip 6. The optimal guidewire/injection lumen diameter is 0.1 to 1.0 mm; however, this lumen can be in the range form 0.025 mm to 2 mm.
  [0113] Referring to FIG. 5, an example of a longitudinal cross section of the distal end of device 2 is shown with catheter body 4, balloon inflation lumen 34, balloon 8, channel 38, guidewire/injection lumen 40, adapter 42, balloon pocket 44 and distal tip 6. Distal tip 6 has an optimal diameter of 0.3 mm to 1.3 mm; however, distal tip 6 can range from 0.1 mm to 4 mm. Adapter 42 is adapted to create a smooth transition from the distal tip 6 to the catheter body 4. By way of example, if the distal tip 6 is 1 mm in diameter and the catheter body 4 is 2 mm in diameter, the adapter will taper from a diameter of 1 mm at its distal most point to 2 mm at its

proximal most point to create a smooth transition from the smaller diameter distal tip to the larger diameter catheter body. As shown in FIG. 5, adapter 42 is positioned on the distal tip 6 at a location that is distal to distal end of catheter body 4, such that a balloon pocket 44 is formed between the distal end of catheter body 4 and the proximal end of adapter 42. Balloon pocket 44 holds unexpanded balloon 8 such that its unexpanded profile is minimized. Optimally balloon 8 would conform to an outer diameter that is equal to the diameter of the catheter body 4. By way of example, if the distal tip diameter is 1 mm and the catheter body is 2 mm, a balloon pocket is formed that is 0.5 mm deep. If balloon 8 is no more than 0.5 mm thick when unexpanded, it will lie equal to or below the surface of catheter body 4. This allows facilitation of the movement of the caterer within the artery or vein.

10

15

20

25

30

35

[0114] Referring to FIG. 6, a longitudinal cross section of the distal end of device 24 is shown with catheter body 28, balloon inflation lumen 50, guidewire/injection lumen 52, balloon 30, and channel 26 with proximal port 54 and distal port 56. Channel 26 passes through and within catheter body 28 and extends from the proximal side of balloon 30 to the distal side of balloon 30. Port 54, at the proximal end of channel 26 is in fluid communication with the outside of the catheter body that is proximal to balloon 30 and port 56, at the distal end of channel 26, is in fluid communication with the area outside of catheter body 28 that is distal to the balloon 30. In this case, the flow through channel 26 is bidirectional. The optimal channel diameter is 0.1 mm to 1 mm; however this channel can be in the range of 0.05 mm to 2 mm.

[0115] FIG 7 illustrates an exemplary embodiment of a longitudinal cross section of the distal end of device 24 with catheter body 28, balloon inflation lumen 50, guidewire/injection lumen 52, balloon 30, valve 32 and channel 26 with proximal port 54 and distal port 56. Channel 26 passes through and within catheter body 28 and extends from the proximal side of balloon 30 to the distal side of balloon 30. Port 54, at the proximal end of channel 26 is in fluid communication with the outside of the catheter body that is proximal to balloon 30 and port 56, at the distal end of channel 26, is in fluid communication with the area outside of catheter body 28 that is distal to the balloon 30. Valve 32, is shown at the opening of port 56 of channel 26. Valve 32 allows flow in the distal direction and prevents flow in the proximal direction. In this case, the flow through channel 26 is unidirectional. The optimal channel diameter is 0.1 mm to 1 mm; however this channel can be in the range of 0.05 mm to 2 mm. The valve can be positioned at the proximal or distal opening or anywhere within the cannel. Location and configuration of the valve will determine flow direction.

[0116] Referring to FIG 8, a longitudinal cross section of the distal end of device 24 is shown with catheter body 28, balloon inflation lumen 50, guidewire/injection lumen 52, balloon 30, valve 32, channel 26 and distal tip 6.

5

10

15

20

30

[0117] Referring to FIG. 9, linear cross sections through catheter body 4 and balloon 8 are shown. Figure 9 illustrates examples of flow channels through a balloon positioned on a catheter, however, various alternatives, modifications, and equivalents may be used. FIG 9A shows catheter 4 and balloon 8 without a channel. FIG 9B shows catheter 4 and balloon 8 with two channels 38 through balloon 8. In this case the flow is bidirectional. FIG 9C shows a catheter and balloon with three bidirectional cannels. FIG 9D shows a catheter 4 with a balloon 8 with two channels, each with a one-way valve. In this case the channel flow in each channel is unidirectional. FIG 9E shows catheter 4 with balloon 8 and three channels 38, each with an independent valve 38. The flow is unidirectional. FIG 9F is a catheter and balloon with two channels and a single circumferential valve 33 that covers both channels 38 and allows flow in only 1 direction. FIG 9G shows a catheter 4 and balloon 8 with three channels and a single circumferential valve 33 that covers all three channels. FIG 9H shows a catheter 5, and four bidirectional channels 39 that are formed by pleating balloon 36 from the outer circumference of the expanded balloon, inward toward the central catheter and securing the inner apex of the balloon to the catheter, thereby forming triangular channels which radiate outward from the central catheter, forming a "V" shape. The lower point of the 'V" is positioned at the catheter surface and the open end of the "V" is positioned at the outer circumference of the circle defined by the largest diameter of the inflated balloon. FIG 9I shows four channels 39, with a single circumferential valve 35 covering all four triangular channels. Although the figure shows four channels, the device of this disclosure can have one, two, three, four or any number of channels. FIG 9J shows a catheter 7 and a balloon 39 with four channels 37 that are formed by pleating the balloon 39 from the inner catheter 7 toward the outer circumference of the expanded balloon 39. FIG 9K shows four channels 37as in FIG 9J with a single circumferential valve 39 that covers all four channels 37. FIG 9L shows four channels 37 as in FIG 9J with a one piece cross-shaped flap valve 41. 25

[0118] While the above is a complete description of exemplary embodiments of the present disclosure, various alternatives, modifications, and equivalents may be used. Therefore, the above description should not be taken as limiting the scope of the disclosure, which is defined by the appended claims and the claims in any subsequent applications claiming priority hereto.

10119] FIG. 10 illustrates a tumor and its associated vasculature with tumor 60, main artery 62, side branch artery 64, tumor artery 66, tumor capillary 68 and anterograde arterial flow direction illustrated by arrows 70.

[0120] FIG. 11 is an expanded view of FIG. 10 with tumor 60, main artery 62, side branch artery 64, tumor artery 66 and anterograde arterial flow direction illustrated by arrows 70.

[0121] FIG. 12 illustrates injection of fluid 76 using a standard micro-catheter 72 with tumor 60, main artery 62, side branch artery 64, tumor artery 66, guidewire 74 and retrograde arterial flow direction illustrated by arrows 70. In this instance, injection pressure and flow volume of the fluid 76 that may contain anti-cancer drugs, radioembolic substances, chemoembolic substances, embolic agents or the like, through microcatheter 72 is higher than the tumor vasculature can accept causing a reversal of fluid flow and blood flow in tumor artery 66, side branch artery 64 and main artery 62. This retrograde flow causes the injected fluid 76 to enter the main artery, flowing in both directions and into the general circulation resulting in the injected fluid traveling to non-target tissues and organs. This unintended delivery of fluid 76 to non-target sites is undesirable and must be avoided since it can cause serious complications. The present disclosure solves this problem by preventing reflux and associated non-target delivery of fluid 76.

[0122] FIG. 13 illustrates catheter 4 of the present disclosure entering the main artery with balloon 8 and distal tip 6. Although the entry point from outside the body is typically through the femoral artery at the groin, any artery or vein from any location on the body can be used for access provided that it creates a pathway to the target vasculature.

[0123] FIG. 14 illustrates catheter 4 with balloon 8, of the present disclosure, following guidewire 74 into side branch artery 64.

15

20

25

30

[0124] FIG. 15 illustrates catheter 4, inside the branch artery 64 with balloon 8 in the expanded configuration, channels 38 and valves 16. Valves 16 are illustrated in the closed position immediately following the inflation of balloon 8.

[0125] FIG. 16 illustrates catheter 4, inside the branch artery 64 with balloon 8 in the expanded configuration, channels 38 and valves 16. Valves 16 are illustrated in the open position since anterograde blood flow as indicated by arrows 70 and associated blood pressure causes these valves to open and allow the blood to continue to flow in the anterograde direction and into the tumor vasculature.

[0126] FIG. 17 illustrates the initial injection of fluid 76 into side branch artery 64 through catheter 4, channels 38 and open valves 16. When the injection is initiated, the anterograde blood flow carries the injection fluid 76 into the tumor vasculature including tumor arteries 66, and capillaries 68.

[0127] FIG. 18 illustrates injection of fluid 76 at a point when fluid pressure increases within the tumor vasculature and concomitant retrograde arterial blood flow and injection fluid flow in the direction as illustrated by arrows 70. Shown in this figure are catheter 4 of the present disclosure with tumor 60, main artery 62, side branch artery 64, tumor artery 66, and guidewire 74.

35 Injection fluid 76, may contain anti-cancer drugs, radioembolic substances, chemoembolic

substances, embolic agents or the like, which can cause serious complications if delivered to non-target sites. In this case, the retrograde pressure causes valves 16 to close and prevents the reflux of injection fluid into the general circulation, thereby preventing complications associated with delivery of injection fluid to non-target sites.

- [0128] FIG. 19 illustrates a point in time following the completion of fluid injection. At this point, the pressure in the vasculature that is the distal to balloon 8, including side branch 64 and tumor artery 66, is reduced below normal blood pressure due to the gradual uptake of the injected fluid into the tumor vasculature. The blood pressure on the proximal surface of balloon 8 and valves 16 cause them to open allowing anterograde blood flow to be reestablished. When this occurs, the excess fluid 76 distal to balloon 8 and within the side branch artery 64 and tumor vasculature, including tumor artery 66 and tumor capillaries 68, is flushed forward and up into the tumor vasculature, thereby enabling delivery of the entire fluid dose and eliminating fluid reflux and associated complications.
  - [0129] FIG. 20 illustrates the deflation of balloon 8 on catheter 4.
- 15 [0130] FIG. 21 illustrates the withdrawal of catheter 4 into the main artery 62.
  - [0131] FIG. 22 illustrates another embodiment of the present disclosure as described in FIGS. 6, 7 and 8. In this case valve 32 on the distal end of channel 26 of catheter 28 is in the closed position.
  - [0132] FIG 23 illustrates an embodiment of FIG. 22 with valve 32 in the open position.
- [0133] A Method, according to the present disclosure is illustrated by FIGS. 13 through 21; the method applies to both the embodiment illustrated in FIGS. 1B, 2, 3, 4 and 5 and the embodiment illustrated in FIGS. 1C, 6, 7, 8, 22 and 23.
  - [0134] Referring to FIG 24A, a longitudinal cross section of a standard single lumen straight tip catheter 101, having a proximal and distal end, is shown with catheter body 102, and hub 105.
- Hub 105, positioned at the proximal end, further comprises guidewire/injection lumen 110, in fluid communication with a catheter lumen longitudinally oriented and extending from hub 105 and exiting at the distal end of the catheter body 102. The proximal hub connects to a syringe or other means to inject fluids via a luer fitting, thereby allowing injection of a fluid through the longitudinal lumen and exit at the distal end of catheter body 102.
- 30 **[0135]** Referring to FIG. 24B through 24E, longitudinal cross-sections of a sequential assembly of a preferred embodiment of the present disclosure is shown. Referring to FIG 24B, device 103 is shown, having a proximal and a distal end, catheter body 102, catheter extension 104 and hub 106. Hub 106 further comprises handle 109, guidewire/injection luer fitting 110 and balloon fill luer fitting 112. Luer fitting 110 is in fluid communication with a first longitudinal
- 35 guidewire/injection lumen of catheter body 102, extending to the distal end of catheter extension

104, and luer fitting 112, in fluid communication with a second longitudinal balloon fill lumen of catheter body 102, extending to a balloon fill port located near the distal end of catheter body 102. FIG 24C further comprises nose cone 114, positioned on catheter extension 104, forming balloon pocket 116 disposed between the distal end of catheter body 102 and nose cone 114.

5

10

15

20

25

30

35

- Further, a portion of catheter extension 104 can, if desired, extend distal to nose cone 114, thereby forming distal tip 118. FIG 24C, further illustrates occlusion balloon 120 in a radially compressed configuration and FIG 24E illustrating balloon 120 in a radially expanded configuration. Hubs 106 can be constructed from styrene, polyurethane, polypropylene, lipid resistant polycarbonate, polycarbonate, Pebax (polyether block amide), of any durometer, or any convenient material and can have any configuration, including, but not limited to, a solid structure comprising two lumens or tubular extensions of the lumens of catheter body 102, provided that they are in fluid communication as described above. Catheter body 102 can be formed from any plastic or thermoplastic material including polyurethane, PTFE, polyimide,
- formed from any plastic or thermoplastic material including polyurethane, PTFE, polyimide, polypropylene, Pebax or the like, and can comprise a single section or multiple sections of different diameter, durometer, braid or coil reinforcement or any convenient construction with a diameter of between 1Fr and 10 Fr more typically of 2 Fr to 5 Fr. Catheter extension 104 can have a diameter of 0.5 Fr to 5 Fr, more typically of 1Fr to 3 Fr and can be absent or can be of any length, typically 2 mm to 30 mm, more typically from 5 mm to 20 mm. If the catheter extension 104 extends beyond nose cone 114, the section distal to the nose cone forms the distal tip 118.

Distal tip 118 is advantageous when injecting deep into the tumor vasculature is desired and will

- also help tracking of device 103 over a guidewire around sharp corners and through a tortuous vasculature path. Nose cone 114 can be made from any polymer or metal or can be formed from a radiopaque marker band. Balloon pocket 116 can be of any length between 2 mm and 50 mm, more typically between 5 mm and 20 mm. Occlusion balloon 120 has a longitudinal length of 1 mm to 30 mm, more typically of 2 mm to 10 mm and a diameter of 1 mm to 50 mm, typically from 2 mm to 10 mm and can be composed of silicone, polyurethane, polyethylene, PET (polyethylene terephthalate), nylon or the like and can be of any configuration or of any length or shape and can be glued, chemically bonded, heat bonded, RF welded, sonically fused, compressed or crimped under a collar to catheter 102 or catheter extension 104.
- [0136] Referring to FIG 25A, a distal section of a device 128 of a preferred embodiment of the present disclosure is shown and includes catheter body 102, catheter extension 104, nose cone 114, balloon pocket 116, balloon fill lumen 124, guidewire/injection lumen 126 and radially compressed balloon 120. Referring to FIG 25B, balloon 120 is shown in its radially expanded configuration. Balloon fill lumen 124 can be of any convenient shape including but not limited to round, semicircular, or crescent or any shape, typically optimized to provide maximum area

and flow rate. Guidewire/injection lumen 126 is typically round, having a diameter of 0.005" to 0.1", more typically from 0.01" to 0.05"; however, it can be of any desirable shape.

[0137] Referring to FIG 26A, the distal end 133 of an embodiment of the present disclosure is shown with catheter body 102, catheter extension 104, nose cone 114, balloon pocket 116,

- balloon fill lumen 124, guidewire/injection lumen 126 and radially compressed balloon 120. In this instance, the balloon pocket 116 is formed between a proximal collar 130 and a distal collar 114, tapered forward thereby forming a nose cone. The balloon bonding tails 115 can be bonded within the pocket or compressed or bonded under collars 114 and 130. Distal collar 130 can
- comprise a metal, such as a radiopaque marker band or a plastic such as heat shrink tubing and
  can be 1 mm to 20 mm in length, more typically from 2 mm to 10 mm. Balloon fill lumen 124 is
  shown traveling under balloon pocket 116 and ending at its distal end. Guidewire/injection
  lumen 126 is shown traveling longitudinally through catheter 102 and catheter extension 104,
  ending at the distal end of the catheter. Balloon 120 is shown tucked into pocket 116 with outer
- diameter substantially no larger than the outer diameter of catheter body 102. Figure 26B shows the same construction as FIG 26A with balloon 120 in its radially expanded configuration.

  [0138] Referring to FIG 27A, a view of distal section 133 is shown and further illustrates that the

profile of a preferred embodiment of the present disclosure, including the radially compressed balloon 120, the distal collar 114 and the proximal band 130, have an outer diameter equal to or less than that of catheter body 102. FIG 27B shows the same construction as FIG 27A with

balloon 120 expanded from pocket 116 and between collars 114 and 130.

5

25

30

35

[0139] Referring to FIG 28A, a distal section of an alternate embodiment 135 of the present disclosure is shown with outer catheter 119, inner catheter 127, nose cone 125, radially constrained balloon 131 and catheter channels 123. The outer catheter 119 is adapted over inner catheter 121, the catheters configured to provide a radially distributed space between inner and outer catheters extending longitudinally along the length of device 135. Outer and inner catheters can have a length of 10 cm to 250 cm, more typically 50 cm to 150 cm and a diameter of between 0.5 Fr and 10 Fr more typically of 1 Fr to 5 Fr. Inner catheter 127 can have a length less than, equal to, or longer than the outer catheter 119, however in the present figure, inner catheter 127 is shown to be longer than outer catheter 119, its distal end forming the catheter extension 127. Nose cone 125 is disposed along the distal extension of inner catheter 127 at some distance from the distal end of outer catheter 119, the distance being 2 mm to 50 mm, more typically between 5 mm and 20 mm. The balloon pocket is formed between the distal end of catheter 119 and the proximal end of nose cone 125. FIG 28B shows an end view of outer catheter 119, disposed over inner catheter 127, with radially configured channels 123 and stand-

offs 127 disposed between outer catheter 119 and inner catheter 127. Four channels are

illustrated, however device 135 can have 0, 1, 2, 3, 4 or any number of channels and stand-offs, the stand-offs defining the outer edges of the channels 123 and can be formed on either the inner or outer catheter with a height limited only by the diameter of the inner and outer catheters and space there between. Although stand-offs are shown, they are not required, provided that the inner catheter OD is smaller than the outer catheter ID, thereby forming a space between the 5 inner and outer catheters which allow fluid to flow longitudinally along device 135. Device 135 can comprise single layer inner and outer catheters or one or both can have multiple layers. In a preferred embodiment, outer catheter 19 is a three layer construction with an outer Pebax layer, a central polyimide layer including reinforcement such as a coil or braid and an inner Teflon layer. Inner catheter 127 is a single layer of low friction tubing, or tubing of similar construction to that 10 described for the outer catheter 119. FIG 28C shows a longitudinal view of device 135 with outer catheter 119, unexpanded balloon 131, nose cone 125 and a catheter extension of inner catheter 127. Balloon 131 is shown tucked within a pocket formed between the distal end of catheter 119 and the proximal end of nosecone 125. FIG 28D shows a longitudinal cross section of device 135, showing balloon inflation channel 123 disposed between outer catheter 119 and 15 inner catheter 121. In this instance, the distal end of balloon 131 is shown inserted into the proximal end of nose cone 125; however both proximal and distal balloon tails can be bonded directly to inner catheter 127, reflowed into catheter 119 or nose cone 125 or by any means, provided that the balloon tails are positioned approximately below the outer diameter of catheter 119. 20

[0140] Referring to FIG 29A though FIG 29E, examples of balloon configurations that may be used in the device of the present disclosure are shown which can be compliant or noncompliant, dilation or occlusion and can be made from any material including, but not limited to, silicone, polyurethane, polyethylene, PET (polyethylene terephthalate) and nylon.

25 **[0141]** Referring to FIG 30, a surface view of balloon 160 is shown with one-way bypass channels 162 and valves 164, the balloon is described in detail in co-pending patent application number 61/821,058.

30

[0142] Balloon 160 and valves 164 allow flow from the compartment proximal to the proximal surface of balloon 160 to the compartment distal to the distal surface of balloon 160 (anterograde flow) and prevents flow from the compartment distal to the distal surface of balloon 160 to the compartment proximal to the proximal surface of balloon 160 (retrograde flow). Balloon 160 can be disposed on the catheter of the present disclosure and held within a balloon pocket as illustrated in FIGS 24-29 and enable anterograde injection of therapeutic agents from within an artery and into a target while maintaining normal (anterograde) blood flow through channels 162

of balloon 160 and prevent retrograde flow (reflux) of therapeutic agents backward over the catheter, even when pressure distal to balloon 160 is elevated above systolic.

- [0143] Referring to FIG 31, a cross section of balloon 160 is shown with channels 162 and microvalves 164, positioned within channels 162.
- [0144] FIG 32 shows an example of a sequential assembly of an embodiment of the present disclosure. Referring to FIG 32A, catheter 170 of device 168 is shown. FIG 32B illustrates a first step in the construction of device 168 whereby a balloon pocket 172 is formed about a distal section of catheter 170 and a second step, as in FIG 32C, whereby a rounded distal end 174 of catheter 170 is formed and a third step as in FIG 32D whereby a balloon 176, with bonding tails 178 is disposed within pocket 172 of catheter 170 and a fourth optional step whereby catheter 170 or other material is reflowed at position 180 over balloon tails 178. FIG 32F illustrates balloon 176 in its radially expanded configuration with balloon pocket 172 and tails 178 bonded
  - [0145] Referring to FIG 33, three alternate embodiments of the present disclosure are illustrated.
- FIG 10A shows device 173 with catheter 181, balloon pocket 177, radially constrained balloon 178, catheter extension 171 and nose cone 182. FIG 33B shows device 173 with radially expanded balloon 178 and bonding tails 183 bonded within balloon pocket 177. FIG 33C illustrates device 175 with nosecone 182 and proximal bonding tail 191 reflowed into catheter 181 at position 179 and the distal balloon bonding tail 193 reflowed into or under nose cone 182

in pocket 172 without being covered by reflow or other means.

20

30

- to catheter extension 171 at location 184. FIG 33D illustrates device 179 with catheter 181, balloon 178, nose cone 182 and collar 186. Balloon 178 has a proximal tail 191 positioned under collar 186 and distal balloon tail 193 reflowed or bonded to catheter extension 171 and under nose cone 182 or into nose cone 182. FIG 33E shows device 179 with balloon 178 expanded from within the balloon pocket formed between collar 186 and nose cone 182.
- 25 [0146] FIG 34 shows yet another embodiment of the present disclosure with two balloons 188 and 189, catheter 183, balloon pockets 194 and 190, reflow areas 192 and nose cone 189.

  Although the example of FIG 34 shows both balloon 188 and 189 positioned within pockets 194 and 190, only one balloon need be positioned within a pocket.
  - [0147] FIG 35 shows still another embodiment of the present disclosure with balloon 196 containing channels 195 and 197, valve 198 in the closed orientation, valve 199 in the open orientation, collar 183, nose cone 1102 and reflow area 1100. Although valve 198 is shown closed and valve 199 is shown open, they will typically act in unison and all either be simultaneously open or closed.
- [0148] FIG 36 shows still another embodiment of the present disclosure which includes two pressure sensors, positioned distal and proximal to balloon 1110, although a single pressure

sensor positioned either distal or proximal to the balloon can be used. These pressure sensors can be used to monitor and, in conjunction with a syringe, control injection pressure either manually of by an automated means. Alternately pressure proximal or distal to the occlusion balloon can be measured through the catheter using an external pressure gauge (1113), the distal pressure being measured via the guidewire/injection lumen 1105 or any other catheter lumen or other tube. The pressure gauge can be connected to a pump, via a processor, allowing the pump to achieve a defined pressure or be programed to a specific set of pressures, volumes and/or flow rate as a function of time.

5

10

15

20

25

30

35

[0149] Referring to FIG 37, four embodiments of balloon configurations are shown. FIG 37A, shows device 1115 with balloon 1117 and catheter 1116. Balloon 117 in a radially expanded configuration, occupies only part of the circumference of catheter 1116. FIG 37B illustrates device 1118 with catheter 1119 and balloons 1121 whereby the four balloons 1121, in radially expanded configurations are arranged circumferentially about catheter 1119, each occupying a part of the overall outer circumference of catheter 1119. FIG 37C illustrates device 1123, with catheter 1125 and balloon 1127 in a radially expanded configuration, whereby balloon 1127, in a radially constrained configuration is positioned within a pocket of catheter 1125 and the radially outermost part of balloons 1127 is positioned approximately at or below the outer diameter of catheter 1125. FIG 37D illustrates device 1131 with catheter 1135 and balloons 1137 in a radially expanded configuration, whereby balloons 1137, in a radially constrained configuration are positioned within a pocket of catheter 1135 and the radially outermost part of balloons 1137 are approximately positioned at or below the outer diameter of catheter 1135.

[0150] Referring to FIG 38, an anatomical structure 1120 is shown with main artery 1122, right artery 1124, left artery 1126, right capillaries 1128, left capillary 1129, arterial side branch 1136, vein 1130, arteriovenus shunt 1132, tumor 1133, blood flow directional arrows 1134, standard straight tip catheter 1138, and embolization particles 1125. FIG 38A illustrates the beginning of a transarterial embolization (TAE) procedure wherein the embolization particles 1125 are exiting the distal end of catheter 1138 and are carried by forward (antrograde) blood flow into tumor 1133 in a delivery method that is completely mediated by blood flow and normal blood pressure (flow mediated delivery). Capillary beds 1128 and 1129 of tumor 1133 begin to fill with embolic particles 1125 and arteriovenus shunt 1132 carries particles into vein 1130 causing anterograde reflux and non-target embolization. The flow through the areteriovenous shunt 1132 is rapid since the arterial pressure is significantly higher than venous pressure. Referring to FIG 38B, continued injection of particles 1125 from the distal end of standard straight tip catheter 1138 results in the packing of particles and embolization of the distal ends of capillary beds 1128 and 1129. Distal capillary embolization causes the flow through arteriovenous shunt 1132 to stop

5

10

15

20

25

30

35

and pressure to build in left artery 1126. As embolization progresses, the back pressure in artery 1126 continues to rise until embolic particles reflux in the retrograde direction 1142 causing nontarget embolization of the right artery 1124, arterial side branch 1136 and main artery 1122. This situation can cause non-target embolization, loss of an unknown amount of particles, delivery of an unknown and irreproducible dose and non-optimal distribution of embolic particles in the tumor vasculature. In this instance, both anterograde and retrograded reflux can occur. [0151] Referring to FIG 39, anatomical structure 1120 is shown as in FIG 38. In this instance, a balloon 1141, with channels 1143 and one-way valves (FIGS 30 and 31) is positioned about the distal end of catheter 1139. Balloon 1141, so constructed, will allow only anterograde (normal) flow and prohibit retrograde flow. Referring to FIG 39A, balloon 1141 is shown in its radially expanded configuration and blood is flowing through balloon channels 1143 as indicated by blood flow arrow 1134 and into the vasculature of tumor 1133. Embolic particles 1125 are released from the distal end of catheter 1139 and carried forward by blood flow into capillaries 1128 and 1129. Capillary beds 1128 and 1129 of tumor 1133 begin to fill with embolic particles 1125 and arteriovenus shunt 1132 carries particles into vein 1130 causing anterograde reflux and non-target embolization. The flow through the areteriovenous shunt 1132 is rapid since the arterial pressure is significantly higher than venous pressure. Referring to FIG 39B, continued injection of particles 1125 from the distal end of balloon catheter 1139 results in the packing of particles and embolization of the distal ends of capillary beds 1128 and 1129. Distal capillary embolization causes the flow through arteriovenous shunt 1132 to stop and pressure to build in left artery 1126. As embolization progresses, the pressure in artery 1126 continues to rise, however the valves of balloon 1141 close and prohibit retrograde reflux. In this instance, continued injection will increase the packing pressure of particles 1125 and can increase packing density and increase flow into distal locations in the margins of a tumor or other structure thereby improving particle distribution throughout the target vasculature. As part of the present method, pressure distal to the balloon can be regulated between systolic and any pressure above systolic, provided that it is within a range that is safe for the patient. By way of example, injection pressure can be low at the onset of the embolization procedure and increased at some point thereafter to a pressure greater than systolic. Such a point may, for example, be chosen to coincide with the stoppage of flow through arteriovenous shunt 1132. This method may improve particle distribution and packing. Alternately, the injection pressure through catheter 1139 can be high at the onset, thereby forcing particles rapidly into the distal section of capillaries 1128 and 1129 and hasten embolization of arteriovenous shunt 1132, thereby reducing anterograde reflux. Alternately, according to this method, a low to high pressure gradient or a high to low pressure gradient can be used. The aim for the use of a pressure mediated delivery of particles is

to optimize for a low level of anterograde reflux, substantial elimination of retrograde reflux, high particle distribution and high particle density. A pressure sensor as in FIG 36 can be used on the proximal and/or distal side of balloon 1142 to monitor pressure and enable a selection of a procedural end point based on a definitive pressure reading.

[0152] Referring to FIG 40, anatomical structure 1120 is shown as in FIG 38. In this instance, 5 an occlusion balloon 1142 is positioned about the distal end of catheter 1140. Referring to FIG 40A, balloon 1142 of catheter 1140 is shown in a radially expanded configuration. Since expanded balloon 1142 completely occludes artery 1126, all arteries and capillaries distal to the balloon are isolated from the main artery 1122, right artery 1124 and side branch artery 1136 thereby causing blood pressure distal to the balloon to drop from approximately normal arterial 10 pressure of about 80 mmHg to a pressure in the range of 0-50 mmHg. When this happens, blood flow through the arteriovenous shunt 1132 can reverse as shown by blood flow arrow 1135, or the anterograde flow slowed or stopped. Referring to FIG 40B, initial injection of particles 1125 will be against a pressure, with a minimal anterograde flow or into a flow stasis.

15

20

25

30

35

Retrograde pressure flow against the particle injection can result from the flow of venous blood from vein 1130, through arteriovenus shunt 1135 and into the arterial capillary 1129 or from arteriovenous capillary beds associated with capillaries 1128. As particles 1125 are injected, they fill capillaries 1128 and 1129; however, particles cannot easily flow through arteriovenous shunt 1132 because of the reversal or slowing of flow and pressure. Continued injection can result in embolization of the distal portion of capillary 1129 and blockage of arteriovenous shunt 1132 with concomitant reduction or elimination of anterograde reflux. Increasing injection pressure through catheter 1140 following embolization of arteriovenous shunt 1132, can result in a high levels of particle density and distribution. Alternately, according to this method, a gradient can be used. The profile of the pressure gradient can be any function of time and pressure including, but not limited to, a linear or step function from low to high, high to low, alternating high to low and low to high or any other function and can be administered manually, in a semiautomated manner or using a programmable delivery means. Alternately according to this method, a pressure sensor as in FIG 36 can be used on the proximal and/or distal side of balloon 1142 to monitor pressure and select a procedural end point based on a definitive pressure reading. Referring now to FIG 40D, the injection through catheter 1140 into tumor 1133 can be accomplished using an automated pump/pressure monitor system whereby the pressure distal to occlusion balloon 1142 is measured on gauge 1152, the pressure reading transferred through

connection 1154 to pump 1150 which controls the injection of anti-tumor agents from syringe 1156. Pump 1150 can be controlled manually or programed to any function of flow rate, time and/or pressure. The endpoint can be selected at any desirable pressure.

[0153] The aim of the present method is to eliminate retrograde reflux, reduce or eliminate anterograde reflux, control the particle density and distribution, deliver an optimal dose, enable a defined pressure endpoint, improve efficacy and reduce toxicity.

[0154] Referring to FIG. 41A, a longitudinal cross section of a catheter is shown with proximal and distal ends, catheter body 204, distal tip 203 and proximally disposed hub 206. Catheter body 204 has two lumens that are in fluid communication with hub 206, a first lumen extending from port 208 of hub 6 to the distal tip 203 of catheter 204 whereby fluid can be injected from the proximal hub 206 and exit at the distal tip 203 of catheter 204 and a second lumen extending from port 210 of hub 206 to an intermediate location at some distance from the distal tip 203 of catheter 204, the second lumen adapted to communicate with a balloon for inflation and deflation.

5

10

15

20

25

30

35

[0155] Referring to FIG 41B, a longitudinal cross section of a first embodiment of the present disclosure is shown with proximal and distal ends, catheter body 204, distal tip 203, and two layered occlusion balloon 214 with channels 205 and valves 207 and proximally located hub 206. Although balloon 214 is shown with two channels, each with a valve, balloon 214 can have 1, 2, 3, 4 or any number of channels and any number of valves or be without valves. In this instance, the valve configuration allows fluid to flow from the proximal side of balloon 214 to the distal side of balloon 214 and to restrict flow from the distal side of balloon 214 to the proximal side of balloon 214; however, the opposite valve orientation and flow direction is also part of the present disclosure. Catheter body 204 can have a diameter of between 1 Fr and 10 Fr, more typically 2 Fr to 5 Fr and a length of 10 cm to 250 cm, more typically 50 cm to 150 cm. Two layered occlusion balloon 214 can be from 1 mm to 30 mm in diameter, more typically 2 mm to 10 mm in diameter, in its radially expanded configuration.

[0156] Referring to FIG 41C, an alternate embodiment of the device of the present disclosure is shown, having catheter body 218, distal tip 209, hub 206 and umbrella shaped occlusion structure 220. When in its radially expanded configuration, the occlusion structure will completely occlude the flow of the vessel. The umbrella shaped occlusion structure 220 is positioned at some distance from the distal end of catheter 216 and forms an umbrella shaped structure disposed circumferentially about catheter 216 with its outer diameter in contact with the vessel. Umbrella shaped occlusion structure 220 can be from 1 mm to 30 mm in diameter more typically 2 mm to 10 mm in diameter when in its radially expanded configuration and a longitudinal thickness of 0.25 mm to 10 mm, more typically 0.5 mm to 2 mm. Umbrella shaped occlusion structure 220 is shown with its closed end attached to the catheter distal to the open end of the V shape; however, it can be positioned in the opposite orientation or it can be positioned at a 90 degree angle with respect to catheter body 18.

[0157] Referring to FIG 41D, device 222 of the present disclosure is shown having catheter 218, distal tip 209, hub 206, and a unidirectional umbrella occlusion structure 224 with channels 230 and valves 228. Occlusion structure 224 will allow proximal to distal flow and prevent distal to proximal flow.

[0158] Referring to FIG 41E, device 232 is shown with catheter body 234, catheter distal extensions 235 and distal tip 211. Catheter extension 235 can have a diameter of 0.5 Fr to 5 Fr, more typically of 1Fr to 3 Fr and can be absent or can be of any length, typically 2 mm to 30 mm, more typically from 5 mm to 20 mm.

10

15

20

25

30

35

- [0159] Referring to FIG 41F, a preferred embodiment of the present disclosure is shown with catheter body 237, catheter extension 235, distal tip 211, nose-piece 241 and two layered occlusion balloon 243 in its radially expanded configuration. In this instance, two layered occlusion balloon 243 is disposed within a pocket formed on distal catheter extension 235 and between the distal end of catheter body 237 and the proximal end of nose-piece 241. The nose piece can be a tapered nose cone, a distally rounded piece of tubing or catheter, a blunt tube or any structure with a diameter equal to less than the catheter body. When in the radially constrained configuration, the outer diameter of the two layered occlusion balloon 243 has an outside diameter that is about equal to the outer diameter of the catheter body 237.
  - [0160] Referring to FIG 41G, yet another embodiment of device 245 of the present disclosure is shown, having catheter body 247, distal tip 234, nose-piece 241, proximal hub 206 and unidirectional umbrella shaped occlusion structure 224 with channels 230 and valves 228. In this instance, unidirectional umbrella occlusion structure 224 with channels 230 and valves 228 is disposed within a pocket formed on distal catheter extension 234 and between the distal end of catheter body 247 and the proximal end of nose-piece 241. The nose piece 241 can be a tapered nose cone, a radiopaque marker band, a distally rounded piece of tubing or catheter, a blunt tube or any structure of about equal diameter to the catheter body. When in the radially constrained configuration, the unidirectional umbrella shaped occlusion structure 224 has an outside diameter that is about equal to the outer diameter of the catheter body 247.
  - [0161] Referring to FIG 42, four views of a preferred embodiment of the unidirectional occlusion structure of present disclosure is shown. FIG 42A illustrates a two layered unidirectional occlusion structure 236 in its radially expanded configuration (also seen in FIGS 41B and 41F), having a proximal end 238, a distal end 240, balloon 242, balloon sheath 244, channel 246, valve structure 250, outer balloon sheath tail 254, balloon tail 256, flow direction arrow 252 and flow exit 248. When occlusion structure 236 is disposed on a catheter as in FIG 41F, fluid flows in the proximal to distal direction (anterograde) as indicated by arrow 252 through channel 246 and valve 250 and exits out the distal flow exit 248. The anterograde fluid

pressure on the inner surface of balloon sheath 244 at the distal end of channel 246, causes distally directed displacement or deflection of the inner surface of balloon sheath 244 at valve 250, thus allowing fluid to pass through flow exit 248. When flow is reversed, fluid pressure on the outer distal surface of balloon sheath 244 at valve 250 causes the balloon sheath 244 to press against the distal surface of balloon 242, closing valve 250 and preventing retrograde flow. Placing the unidirectional occlusion structure 236 in the opposite direction on the catheter will result in distal to proximal flow and prohibit proximal to distal flow. Although the occlusion structure of FIG 42A is shown with two layers including an inner balloon and an outer sheath, it is understood that the sheath need not be present and a balloon with channels from the proximal surface to the distal surface is considered part of the present disclosure. Balloon 242, including channels 246 can be formed by molding, extruding, vacuum forming or otherwise shaping a material to include the desired number and configuration of channels. Alternately, a standard balloon, including but not limited to, round or oval, can be modified to achieve proximal to distal channels. One method to modify a balloon is by forming longitudinal pleats circumferentially oriented, thereby forming V shaped channels that extend from the proximal surface of the balloon to the distal surface of the balloon. Placing a sheath over such a modified balloon in the same manner as described above would give the same result as the balloon shown in FIGS 42A through 42D.

5

10

15

25

30

35

[0162] Referring to FIG 42B, a side view of the unidirectional occlusion structure 236 of this disclosure is shown with proximal end 238 distal end 240, balloon 242, balloon sheath 244 and flow direction arrow 252.

[0163] Referring to FIG 42C, a proximal view of the unidirectional occlusion structure 236 is shown with proximal end 238, distal end 240 and channels 246.

[0164] Referring to FIG 42D, a distal surface view of the unidirectional occlusion structure 236 is shown with proximal end 238, distal end 240 and flow exit 248. Flow exit 248 is formed as a space between balloon tail 256 and balloon sheath 254. It is also possible to terminate balloon sheath 244 immediately below channels 246 forming a valve 250 that does not include balloon sheath tail 254.

[0165] Referring to FIG 43, an illustration of a prototype of the unidirectional occlusion structure 236 is shown in its radially expanded configuration with proximal end 238, distal end 240, balloon sheath 244, balloon 242 (positioned inside balloon sheath 244), and channel 246. This device was tested and will withstand at least 220 mmHg against its distal surface without retrograde flow.

[0166] Referring to FIG 44, device 60 illustrates the unidirectional occlusion structure 272 in a radially constrained configuration adapted to the distal extension 267 of a catheter 265 with

distal end 262, proximal end 264, proximal collar 266, distal collar 268 (formed into a nose cone) and device pocket 270. Proximal collar 266 and distal collar 268 can comprise a metal, such as a radiopaque marker band, heat shrink tubing or any plastic material such as polyurethane, polyethylene, polystyrene, acetal, PTFE, nylon or the like, and can be 1 mm to 20 mm in length, more typically from 2 mm to 10 mm in length. In this instance, circumferentially oriented occlusion structure 272 is held within pocket 270 of catheter 265, with an outer diameter approximately equal to the outer diameter of catheter 265.

5

10

15

20

25

[0167] Referring to FIG 44B, device 260 is shown with unidirectional occlusion structure 272 in its radially expanded configuration with proximal end 264, distal end 262, balloon 242, balloon sheath 244, valve 250, flow exit 248, channel 246, flow arrow 252, proximal collar 266, distal collar 268, catheter 274, balloon fill lumen 270 and guidewire/injection lumen 276. In this instance, there is no distal balloon sheath tail, the balloon sheath terminating on the balloon surface just below channel 246 and above the perimeter of catheter extension 267, thereby positioning the flow exit between the termination of the balloon sheath and the catheter.

[0168] Referring to FIG 45A, device 280 is shown with catheter 282 and an umbrella shaped structure occlusion structure 2284 in its radially expanded configuration, whereby the umbrella shaped occlusion structure 284 is oriented circumferentially about catheter 282 such that its outer circumference is 360 degrees about catheter 282. When device 280 is placed in an artery or vein and umbrella shaped occlusion structure 284 is placed in its radially expanded configuration, the outer perimeter of occlusion device 284 will be at least in contact with the interior of the vessel wall and substantially occlude flow. FIG 45A shows occlusion structure 284 in forward V orientation and FIG 45B shows device 282 with the umbrella shaped occlusion device 288 in a reverse V configuration. The occlusion structure of the present disclosure can also have a 90 degree orientation with respect to the catheter when in its radially expanded configuration.

[0169] Referring to FIG 46A, device 290 is shown with proximal end 292, distal end 294, catheter 296, two-way occlusion structure 299 in its radially expanded configuration, frame 298 and channels 2100, whereby fluid can flow from proximal to distal or distal to proximal through channels 2100. Although two channels are shown, two-way occlusion structure 298 can have 1, 2, 3 or any number of channels.

30 **[0170]** Referring to FIG 46B, device 2102 is shown with proximal end 2104, distal end 2106, catheter 2108, and unidirectional umbrella shaped occlusion device 2110 comprising, frame 2111, channels 2112 and radial valve 2114, whereby fluid will flow from proximal to distal (anterograde) only, retrograde flow being prohibited by radial valve 2114. Although device 2102 will allow only anterograde flow, if desired, device 2110 of apparatus 2102 can be

configured to allow only retrograde flow and/or have a forward V configuration as shown, or, if desired, a reverse V configuration or an orientation 90 degrees with respect to catheter 2108.

[0171] The frames 298 and 2111 of occlusion structure 2110 can be made of metal, such as shape memory metals nitinol or elgiloy, or plastic such as polyethylene, polyurethane, polystyrene, PTFE, acetal and nylon or elastic materials such as silicone or fabrics such as cotton

5

10

15

35

and rayon and can include a mesh, a wire frame, a diaphragm and can be pleated or otherwise folded or can be any other convenient structure or material provided that it is of sufficient strength and porosity to occlude elevated vascular pressures and capable of integrating channels and valves. Valve 114 can be made from flexible or rigid plastics including polypropylene and polyurethane, elastomeric materials such as silicone and can have a configuration including a flap, sock, cone, duck bill and diaphragm or the like with a thickness of 1 mil to 50 mil, more typically 2 mil to 10 mil.

[0172] Referring to FIG 47A, a distal surface view of a unidirectional occlusion structure 2120 of the present disclosure is illustrated with the catheter 2122 (extending forward), device frame 2124, radial valve 2125 and channels 2126 disposed under radial valve 2125. As shown, radial valve 2125 extends radially outward from catheter 290 and covers all four valves. Four channels are shown in this example; however, any number of channels can be used. This configuration allows flow from the proximal surface to the distal surface of unidirectional umbrella shaped occlusion structure 2120; however, the reverse flow is also possible.

[0173] FIG 47B illustrates another embodiment of the present disclosure comprising unidirectional umbrella shaped occlusion structure 2128 with catheter 2130 (extending forward), device frame 2132, valves 2136 and channels 2134 disposed under valves 2136. In this instance, each channel has a separate valve and although four channels and valves are shown, the device of this disclosure can have any number of channels and valves limited only by the size of the valve and channel and the area of the frames 2124 and 2132. This unidirectional configuration allows flow from the proximal surface to the distal surface of umbrella shaped occlusion structure 2128; however, the opposite flow can be easily achieved by changing the flow direction of the valves or rotating the unidirectional occlusion device 180 degrees on catheters 2122 and 2130.

30 **[0174]** FIG 48 illustrates a prototype micro-valve 2140 configured from 5 mil polyurethane material. This device was tested and will restrain a fluid pressure of at least 220 mmHg applied against its distal surface.

[0175] Referring to FIG 49A, device 2142 is shown with triangular shaped occlusion structure 2145 in its radially expanded configuration and adapted to catheter 2144 whereby the occlusion structure 2145 has frame 2146 oriented circumferentially about catheter 2144 such that its outer

circumference comprises 360 degrees. When device 2142 is placed in an artery or vein, frame 2146 is placed in its radially expanded configuration and the outer perimeter of device frame 2146 will be at least in contact with the interior of the vessel wall and at least substantially occlude flow.

[0176] Referring to FIG 49B, device 2148 is shown with catheter 2150 and a unidirectional 5 triangular shaped occlusion structure 2151 comprising frame 2152, channels 2154 and radial valve 2158. Although reference has been made to a unidirectional occlusion valve with an umbrella shape or a triangular shape, it is understood that any shape including, but not limited to, rectangular, oval, conical, and round can be used. Yet another construction of a unidirectional occlusion structure is a dilation or occlusion balloon or any other medical balloon disposed with 10 channels and valves, the valves extending from a proximal surface to a distal surface. [0177] Referring now to FIG 50A to 50C, a method of deploying an occlusion structure 2191 from a radially constrained configuration to a radially expanded configuration and then returning it to the constrained configuration is shown. FIG 50A shows a longitudinal cross section of device 2180 with a proximal end 2182, a distal end 2184, outer catheter 2186, inner catheter 15 2188, nose cone 2190 and radially constrained unidirectional occlusion structure 2191 with frame 2192, valve 2194, and frame attachment point 2196. Unidirectional occlusion structure 2191 is attached to inner catheter 2188 at attachment point 2196 whereby occlusion device 2191 is preloaded with a force which encourages its distal end to pivot proximally outward at attachment point 2196. In this instance, the outer catheter 2186 constrains occlusion structure 20 2191 against the preloaded force. Device 2180 is first positioned in the vasculature at or in the

[0178] Referring to FIG 50B, outer catheter 2186 is retracted proximally as shown by arrow 2200 while the inner catheter 2188 is held stationary, thereby removing the constraint on occlusion structure 2191, allowing it to pivot outward and in a proximal direction at attachment point 2196 and into its radially expanded configuration. Frame 2192 can be made from a memory metal such as nitinol or elgiloy and pre-formed at attachment point 2196 to the radially expanded configuration thereby pre-loading an outward force on occlusion structure 2191 as it is moved to its radially constrained configuration. If a braided nitinol tube is used, it can be pre-formed into a radially expanded configuration whereby occlusion structure 2191 is oriented circumferentially about catheter 2144 with an outer circumference of 360 degrees. As in this example, the mesh can be coated with polyurethane, PTFE, silicone or the like and channels formed through the mesh and valves placed over the channels.

vicinity of a target structure.

25

30

[0179] Referring to FIG 50C, outer catheter 2186 is retracted distally while holding inner catheter 2188 stationary thereby pivoting frame 2192 distally at attachment point 2198 and placing occlusion structure 2191 in its radially constrained configuration.

[0180] Referring to FIG 51, an anatomical structure 2200 is shown with main artery 2202, right artery 2204, left artery 2206, capillaries 2208, tumor 2209 and blood flow directional arrows 2212. FIG's 51A-51E illustrates a method of the present disclosure wherein a tumor is embolized with drug eluting beads as in Transarterial Chemoembolization (TACE). [0181] In a first step, device 2211, comprising a two lumen catheter 2214 and a radially constrained unidirectional balloon occlusion structure 2216 (also in FIG 42), is advanced over a guidewire 2213 using lumen one (guidewire/injection lumen) of catheter 2214 from an entry point on the surface of the body, usually the femoral artery at the groin, and positioned at, or in the vicinity of, an artery feeding a tumor as in FIG 51A. As indicated by arrows 2212, the blood flows in an anterograde direction over device 2211 and into capillaries 2208 of tumor 2209. [0182] In a second step, the unidirectional balloon occlusion structure 2216 is placed in a radially expanded configuration by inflating the inner balloon of the two layered device of FIG 42 using the second lumen of catheter 2214 (balloon inflation lumen) as seen in anatomical FIG 51B. When placed in a radially expanded configuration, normal blood pressure between about 80 mmHg and 130 mmHg urges valves 2217 of occlusion structure 2216 to the open position, thereby allowing anterograde blood flow through channels 2219 and into the capillaries 2208 of

10

15

20

25

30

tumor 2209.

**[0183]** Figure 51C illustrates a third step whereby chemoembolization particles 2218 are beginning to be injected into left artery 2206 and capillaries 2208 of tumor 209. At this point, valves 2217 of unidirectional occlusion structure 2216 are in the open position and blood is flowing in the anterograde direction through channels 2219 which continues to carry chemoembolization particles 2218 into the vasculature of tumor 2209.

[0184] Referring to FIG 51D, a fourth step is illustrated whereby chemoembolization particles 2218 begin to embolize the distal ends of capillaries 2208, increasing pressure in the proximal section of capillaries 2208 and left artery 2206. This back pressure causes blood flow and chemoembolization particles 2218 to flow in a retrograde direction; however, the back-pressure in left artery 2206 urges valves 2217 to close, thereby maintaining particles 2218 in the vascular compartment distal to occlusion device 2217. Using currently available straight tip catheters, the chemoembolization procedure would be terminated at this point since particles would reflux backward over the catheter and into the general circulation causing non-target embolization and associated complications.

[0185] FIG 51E illustrates a fifth step, not possible using present catheters, whereby embolization particles continue to be injected, without retrograde reflux, and further fill the vasculature of the tumor with particles 2218. This method can both prevent the complications associated with retrograde reflux and allow more particles to enter the tumor.

- [0186] FIG 51F is a final step in the present method whereby occlusion structure 2216 is placed in a radially constrained configuration and device 2211 is withdrawn from the body over guidewire 2213.
- [0187] Although particular mention has been given to a device that is capable of transitioning from a radially constrained configuration to a radially expanded configuration, such a transition is not required. A unidirectional occlusion structure of the present disclosure can be configured in a permanently expanded configuration. In this instance, the occlusion structure may be a highly flexible material such as a low durometer plastic or rubber or a flexible mesh or any material or construction that provides sufficient strength and flexibility to navigate through vasculature and to a target and provide unidirectional occlusion.
- 15 [0188] Referring to FIG 52, distal section 302 of a device is shown with distal end 303, proximal end 304, catheter body 306, distal tip 308, nose cone 310, partial occlusion balloon 312 in a fully expanded configuration, channel 314 and one-way valve 316. In this embodiment, flow is permitted in the proximal to distal direction through channel 314 and restricted, by one-way valve 316 to flow proximally. Partial occlusion balloon 312 can be any shape and diameters

  20 from 1 mm to 30mm more typically from 2 mm to 10 mm.
  - [0189] Referring to FIG 53, a longitudinal section 320 of a distal section of the device of the present disclosure is shown with catheter body 306 distal tip 308, nose cone 310, radially constrained balloon 322, proximal balloon pocket boundary 324 and distal balloon pocket boundary 326. Radially constrained balloon 322 sits within the pocket defined by the distal end of catheter 306 at boundary 324 and the proximal end of nosecone 310 at boundary 326. The outer diameter of the constrained balloon is approximately equal to the outer diameter of catheter body 306. This allows the balloon to sit within the pocket and maintain the catheter at a desirable minimal diameter.

25

30

35

[0190] Referring to FIG 54A, a distal section 330 of the device of the present disclosure is shown with catheter body 306, catheter extension 307, distal tip 308, nose cone 310, balloon pocket 332, guidewire and injection lumen 334 and balloon inflation lumen 336. The catheter body 306 has a diameter of 0.25 mm to 5 mm, more typically from 0.5 mm to 1.5 mm and a length of 10 cm to 240 cm more typically from 75 cm to 150 cm. The catheter extension 307 has a diameter of 0.25 mm to 3 mm more typically from 0.4 mm to 1 mm and a length of 5 mm to 100 mm more typically from 5 mm to 40 mm. The balloon pocket 332 has a depth equal to the

difference in diameter of the catheter body 306 and the catheter extension 307 and a length of 1 mm to 50 mm more typically from 5 mm to 15 mm. The balloon wall thickness and inner diameter are selected, extruded or molded to fit into balloon pocket 332 with minimal balloon extending above of balloon pocket 332.

- 5 [0191] Referring to FIG 54B, distal section 330 includes a balloon 338 in a radially constrained configuration held within balloon pocket 332 and having an outer diameter approximately equal to the outer diameter of catheter body 306.
  - [0192] Referring to FIG 54C, distal section 330 includes balloon 338 in a radially expanded configuration with channels 314 and one-way valves 316 in a closed orientation. Partial occlusion balloon 338 can be any shape and diameters from 1 mm to 30 mm more typically from 2 mm to 10 mm and a length of 1 mm to 50 mm more typically from 5 mm to 15 mm. Channels 314 can be of any shape and configuration and an opening that is calibrated to the desired flow therethrough. In a preferred embodiment, the balloon will have a diameter of 6 mm and a channel diameter of 0.5 mm to 1.5 mm. Balloon 338, including channels 314 can be formed by molding, extruding, vacuum forming or otherwise shaping a material to include the desired number and configuration of channels. Alternately, a standard balloon, including but not limited to, round or oval, can be modified to achieve proximal to distal channels. One method to modify a balloon is by forming longitudinal pleats circumferentially oriented; thereby forming V shaped channels that extend from the proximal end of the balloon to the distal end of the balloon.
- Placing a sheath or film over such a modified balloon results in longitudinal channels and a one way valve as described in co-pending application 61/917,131.
  - [0193] Referring to FIG 55A through 55C, a serial construction of the device of the present disclosure is illustrated.
- [0194] Referring to FIG 55A, a longitudinal view of device construction 350 with catheter body 306, distally located catheter extension 307 and proximally located hub 352 comprising guidewire and injection port 354 and balloon inflation/deflation port 356.
  - [0195] Referring to FIG 55B, a longitudinal view of a device construction 350 is shown with added nose cone 310 and balloon pocket positioned between the distal end of catheter body 306 and the proximal end of nose cone 310.
- 30 [0196] Referring to FIG 55C, a longitudinal view of the device 358 of the present disclosure is shown with catheter body 306, balloon 340 and hub 352. Balloon 340 is shown in a radially expanded configuration with channels 314 and valves 316 in the open position.
  - [0197] Referring to FIG 56, an anatomical structure 360 is shown with tumor 362, main artery 366, distal main artery 367, side branch arteries 370 and 374, tumor capillaries 373, 375, 377 and 379 and blood flow direction arrows 368, 372, 376 and 378. In the case of a tumor in the right

10

15

35

liver lobe, artery 366 is the right hepatic artery and 367 is the distal right hepatic artery which flow toward the tumor as seen by flow direction arrow 368 and 378. In this instance, artery 370 is the gastroduodenal artery and artery 374 is a hepatoenteric artery such as the superduodenal artery, the normal flow of both is away from the hepatic artery, as shown by flow direction arrows 372 and 376, and into arterial networks which supply both the liver and gastrointestinal tract. Blood from the hepatic artery 366 also flows into the tumor capillaries 373, 375, 377, and 379. Normal blood flow through the right hepatic artery is in the range of 4 ml/sec.

[0198] Referring to FIG 57A through 57H, a tumor embolization method according to current medial practice is shown. At least some of the steps shown are used in current catheter based embolization therapy in tumors of the liver

10

15

20

25

30

35

[0199] The first step of the procedure is to advance guidewire 382 from the femoral artery at the groin, through the iliac artery, aorta, celiac artery, hepatic artery and into the right hepatic artery 366 as in anatomical structure 380 of FIG 57A. The diameter of guidewire 382 is typically from about 0.25 mm to 1.25 mm more typically from 0.4 mm to 1 mm.

[0200] In the second step of the procedure, illustrated in FIG 57B, guide-catheter 392 is advanced over guidewire 382 and along the same arterial path as for guidewire 382. Typically, the guide catheter has an outer diameter of about 1.5 mm to 2.5 mm and has a central lumen that can accept a microcatheter with an outer diameter of 0.5 mm to 1.5 mm. The guide catheter is too large to access the vasculature in the vicinity of the tumor and is typically advanced as far along the vascular path toward the tumor as possible. The blood flow follows the same normal pattern as in FIG 56 and flows around the sides of guide catheter 392.

[0201] In the third step shown in FIG 57C, guidewire 382 is optionally removed and replaced with a smaller diameter guidewire 398 that can fit in the central lumen of a microcatheter. Guidewire 398 typically has a diameter of 0.2 mm to 0.75 mm, more typically in the range of 0.25 mm to 0.6 mm.

[0202] The fourth step of the procedure, microcatheter 3104 is advanced over guidewire 398 to a position beyond the distal end of guide catheter 392 and into the vasculature within or in the vicinity of the tumor as shown in anatomical structure 3102 of FIG 57D. Microcatheter 3104 is advanced as close as is practical to the tumor and, if the anatomy allows, into the vasculature of the tumor as in superselective embolization. Microcatheter 3104 typically has a diameter of 0.75 mm to 1.5 mm, more typically at about 1 mm and a total length of 50 cm to 200 cm, more typically from 75 cm to 150 cm. The central lumen microcatheter 3104 is optimized to have an inner diameter as large as possible; however it is usually in the range of about 0.5 mm. [0203] In a fifth step, guidewire 398 is removed from microcatheter 3104 as illustrated in

anatomical structure 3200 of FIG 57E. Removal of guidewire 398 allows the central lumen of

microcatheter 3104 to be used to inject drug and/or embolic materials into the target site within the right hepatic artery and tumor. Blood continues to flow around guide catheter 392 and microcatheter 3104 and into capillaries 373, 375, 377, and 379 of tumor 362, gastroduodenal branch 370 and hepatoenteric branch 374 according to the normal flow pattern shown by arrows 368, 372, 376, and 378.

5

10

15

25

30

35

[0204] In a sixth step illustrated in anatomical structure 3208 of FIG 57F, drug and or embolization agents are injected using a syringe or other means from the proximal end of microcatheter 3104 through guidewire/injection port 354 of hub 52 (FIG 55), longitudinally through guidewire injection lumen 334 (FIG 54), and out the distal end of catheter extension 307. In this instance, embolic particles 3210 are carried by normal blood flow into distal right hepatic artery 3214, tumor capillaries 373, 375, 377 and 379, as illustrated by flow arrow 3212, and into hepatoenteric artery 374 in the direction indicated by flow arrow 376. Drug and/or embolization agents that travel through gastroenteric branch artery 374 or any other arterial branch distal to the distal tip of microcatheter 3104, by normal forward flow, are deposited at non-target sites, including parts of the liver and intestine. This anterograde (to the catheter tip) bypass into distal hepatoenteric arteries can cause serious complications including damage or death to sections of the liver or intestine, gastric ulcers or even the death of the patient. Further, drug and/or embolic agents that travel to non-target sites, fail to enter the tumor; this resulting in a lower than optimal dose to be delivered to the tumor and a lower efficacy than desired.

However, to avoid the aforementioned serious complications, physicians often under-embolize the tumor vasculature.

[0205] Referring to FIG 57G, as forward flow mediated embolization progresses, the distal ends of capillaries 373, 375, 377 and 379 fill with particles 3210 and become embolized. This process dramatically slows the blood flow moving through the tumor and causes a sharp rise in pressure within the tumor vasculature and concomitant retrograde deflection of the high volume blood flow from the hepatic artery. Further, backpressure develops in the distal hepatic artery 3214, resulting in particles to flow in a retrograde direction as in flow direction arrow 3234. This can result in: (1) increased anterograde bypass into hepatoenteric branch 374, reflux over the catheter and into gastrodudenal artery 370 and (3) a signal to the physician that particle injection should stop, even though the tumor is only partially embolized. In this instance, it is possible that larger capillaries become embolized first, due to a larger blood flow while smaller capillaries remain un-embolized. The rapid rise in pressure is in part caused by distal capillary embolization and in part caused by the high volume blood flow from the hepatic artery. Given that embolization is the desired endpoint, it appears that slowing the forward flow of blood from the hepatic artery would allow the tumor to accept the blood and drug and/or embolic agent flow for a longer

period of time and allow more embolization to occur and an improved distribution of particles in the tumor vasculature.

**[0206]** Referring to FIG 57H, injection of drug and/or embolic agents is complete, microcatheter 3104 and guide catheter 392 are removed and the final embolization distribution in tumor capillaries 373, 375, 377 and 379 is shown where, in this example, lager capillary 379 is embolized to the greatest extent, smaller capillary 377 is embolized to a lesser extent and small capillary 373 remains un-embolized.

5

10

15

20

25

30

35

[0207] Referring to FIG 58, a method of tumor embolization, according to the device of the present disclosure is shown. Steps 1 through step 5, shown in FIG's 57A through 57E, are the same for both a standard catheter as shown in FIG 57 and the device of the present disclosure and are not further illustrated.

[0208] Referring to FIG 58A, the device of the present disclosure is positioned in the distal right hepatic artery 3214, with partial occlusion balloon 3256 in its radially expanded configuration comprising channels 3258 and 3260, and one-way valves 3262 and 3264 that are in the open position. FIG 58A shows two channels, however, one, two, three or any number of channels can be used. With or without valves. The maximum channel size is limited by balloon diameter, but can be as small as is practical. Valves 3262 and 3264 can be flap, duck bill, diaphragm, or any type of valve provided that it permits flow only in one direction. Optional pressure sensors 3266, which provides real time pressure measurement in the vascular space distal to partial occlusion balloon 3256 and pressure sensor 3265 which provides real time pressure monitoring in the vascular space proximal to the partial occlusion balloon 3256 are shown. Pressure sensor 3266 which measures pressure in the distal vascular space can be used to signal a procedural endpoint based on a predetermined or non-predetermined pressure reading. This will, for the first time, allow a quantitative and definitive pressure mediated endpoint rather than the present subjective flow mediated endpoint and will enable the procedure to be reproducible and able to be standardized allowing center to center and physician to physician consistency. This is possible only because the vascular space distal to the partial occlusion balloon 3256 is isolated from the vascular space proximal to partial occlusion balloon 3256, thereby allowing that the arterial pressure in the distal space to be closely related to the intra-tumor arterial blood pressure. Blood flow direction through hepatic artery 368, distal hepatic artery 3214 and proximal artery 370 are normal as seen in flow direction arrows 368, 372 and 378 as is the blood flow in tumor capillaries 373, 375, 377 and 379 and illustrated by flow direction arrows 3212. However, partial occlusion balloon 3256 causes a significant reduction in blood flow in distal right hepatic artery 214 and in tumor capillaries 373, 375, 377 and 379. Blood flow can be regulated by the partial occlusion balloon of the present disclosure such that total flow can range from near 100%

(unconcluded flow) to near 0% as in full occlusion. Of most interest is partial occlusion that results in 1% to 25% flow as compared to the un-occluded artery. Therefore, channels 3258 and 3220 allow only a fraction of normal blood flow to pass distally. Blood pressure distal to partial occlusion balloon 3256 is also dramatically reduced by anywhere from about 5 mmHg reduction to 100 mmHg reduction, depending on the nature of the occlusion. This pressure drop causes branch artery 374 to reverse direction as seen by flow direction arrow 376 and now flow toward the distal main artery 3214 and tumor capillaries 373, 374, 377 and 379. The flow reduction and pressure reduction caused by partial occlusion balloon 256 also reduces the flow and pressure within the tumor capillaries 373, 374, 377 and 379. This is of significance because it allows more drug/embolization agents to enter the tumor before backpressure causes flow stasis and retrograde flow.

10

15

20

25

30

35

[0209] Referring now to anatomical structure 3280 of FIG 58B, injection of drug and/or embolic agents is initiated. Blood flow from the proximal main artery 366 is attenuated as it passes through channels 3258 and 3260 and into distal main artery 3214. The anti-cancer agents are carried by the attenuated forward blood flow through distal main artery 3214 and into tumor capillaries 373, 375, 377 and 379. Valves 3262 and 3264 are in the open position as pressure in the distal vascular space is lower than the blood pressure in the vascular space proximal to partial occlusion balloon 3256. Branch artery 374 continues to flow in the reverse direction as indicated by flow direction arrow 376 since blood pressure in the distal right hepatic artery is lower than that of the arterial network connected to the distal end of branch artery 374. In this instance, anterograde drug/embolic agents are prevented from flowing into branch artery 374 and anterograde bypass and non-target delivery does not occur. Optional pressure sensors 3266 and 3265 or pressure measurement through guidewire/injection lumen 334 of catheter 6 (FIG 54A) can be used to monitor real-time pressure.

[0210] Referring to FIG 58C, and looking at anatomical structure 3290, a slower rate of blood flow and lower pressure through distal hepatic artery 3214 allows tumor capillaries 373, 375, 377 and 379 to fill at a slower pace and to a greater distribution than in the current method of full unregulated forward flow. At some point, however, the embolization of tumor capillaries will cause retrograde deflection of blood and anti-tumor agents and a pressure build up in distal hepatic artery 3214 as in FIG 58D. At this point, the increased pressure in distal right hepatic artery 3214 causes valves 3262 and 3264 to close, preventing retrograde flow and non-target embolization through branch artery 370 or any other arteries proximal to partial occlusion balloon 3256. This retrograde deflection and pressure build up will progress at a slower rate as compared to the current standard method of FIG 57. The slower buildup of pressure and retrograde flow allows a larger time window for the physician to terminate the procedure. If

pressure monitoring is done, a defined pressure can be used to terminate the procedure. If the back pressure in distal hepatic artery 3214 exceeds about systolic pressure, branch artery 374 will again follow in the normal flow pattern as illustrated in the flow direction arrow 376 of FIG 56. This situation will allow anterograde drug and/or embolic agents to flow in branch artery 374 and to non-target sites. However, visual observation of contrast movement in branch artery 374 or a defined pressure measurement at or below the flow reversal pressure of branch artery 374 can be used as a procedural endpoint signal.

5

10

15

20

25

35

**[0211]** Referring now to anatomical structure 3300 of FIG 58E, the procedure is complete and catheter 3254, of the present disclosure, and guide catheter 3252 are removed. The distribution and filling of tumor capillaries 373, 375, 377 and 379, using the device of the present disclosure, are improved as compared to the distribution and filling associated with the current microcatheter 3104 of FIG 57H.

[0212] In some embodiments, a pressure feedback loop is implemented to control the rate of fluid introduction (e.g. the rate of embolic bead injection). In such embodiments, a blood pressure sensor may be located at the distal tip of the catheter to monitor the pressure of the vasculature distal to the occlusion or partial occlusion structure. In some embodiments, a predetermined pressure set point may be used. The set point may be an absolute pressure, or a percentage of systolic pressure (such as about 100% of systolic pressure). Such systems can be programmed to maintain the distal vasculature at or below the set point by automatically controlling the infusion rate (e.g. the rate or pressure of an injection pump.) By ensuring that the pressure of the distal vasculature does not exceed systolic pressure, retrograde and/or other undesirable blood flows can be prevented.

[0213] In some embodiments, a rate of pressure change can be used instead of or in addition to a preset pressure in the pressure feedback loop. For example, if the distal vasculature pressure begins to rise faster than a predetermined rate, the rate of embolic substance injection can be slowed, temporarily stopped, or a procedural endpoint may be signaled. The endpoint signal may be an audible, visual, tactile or other signal to persons involved in the procedure, and/or may be a signal that automatically shuts off or changes the state of medical equipment used in the procedure.

30 [0214] FIGS 59A – 59D show the fabrication steps of the distal end of an occlusion catheter according to aspects of the present disclosure.

[0215] FIG 60 shows the completed distal end of the occlusion catheter of FIGS. 59A - 59D being introduced into small branches of a vascular system. The two enlarged contact points of this design allow the catheter to be navigated into smaller vasculature than can be navigated by conventional catheter tips.

[0216] FIG 61A shows an entire occlusion catheter constructed according to principles of the present disclosure. FIG 61B shows details of construction features on the distal end of the occlusion catheter of FIG 61A.

[0217] It is important to note that the device of the present disclosure regulates flow and pressure in the arterial space distal to the partial occlusion balloon, significantly reduces flow rate and pressure in tumor capillaries and causes flow reversal of distal branch arteries. The aforementioned device and method thereby enable substantial elimination of retrograde and anterograde bypass to non-target sites and a more complete filling of the tumor vasculature with drug and or embolic agents and should improve efficacy and reduce complications over standard devices and methods.

## **CLAIMS**

What is claimed is:

5

10

15

20

25

30

35

1. A catheter assembly comprising:

a catheter body having a proximal end, a distal end and a balloon inflation lumen; and an inflatable balloon attachable to the distal end of the catheter body, the balloon having an inner surface that at least partially defines an interior volume, the balloon being configured such that the interior volume can be in fluid communication with the inflation lumen of the catheter body to inflate the balloon, the balloon having a proximal surface and a distal surface,

wherein the balloon is provided with a channel that extends through the balloon, the channel configured to provide fluid communication between the proximal surface of the balloon and the distal surface of the balloon.

- 2. The catheter assembly of claim 1, wherein the catheter assembly comprises a distal tip section that extends beyond the distal end of the catheter body.
- 3. The catheter assembly of claim 1 wherein the balloon comprises a valve configured to affect fluid flow through the channel.
- 4. The catheter assembly of claim 3 wherein the valve is positioned at a distal end of the channel.
- 5. The catheter assembly of claim 3 wherein the valve is positioned within the channel.
- 6. The catheter assembly of claim 3 wherein the valve is configured to allow flow in a distal direction and restrict flow in a proximal direction.
- 7. A device for delivering a therapeutic agent to a target site within a body, the device comprising:

a catheter body having a proximal end, a distal end, a first axial lumen and a second axial lumen, the first axial lumen extending from the proximal end of the catheter body to the distal end of the catheter body and providing fluid communication therebetween, the second axial lumen extending from the proximal end of the catheter body to a more distal location on the catheter body;

a balloon radially disposed near the distal end of the catheter body, the balloon having a proximal balloon surface, a distal balloon surface, a radially constrained configuration and a radially expanded configuration, wherein the balloon is in fluid communication with the second axial lumen and has at least one channel extending from the proximal balloon surface to the distal balloon surface, thereby providing fluid communication therebetween; and

a balloon sheath, disposed over an outer surface of the balloon, the sheath having a proximal side and a distal side, the proximal side having an opening that is in alignment with a

proximal end of the balloon channel, the distal side of the sheath extending over a distal end of the balloon channel, wherein the distal side of the sheath is capable of deflecting distally when a distally directed fluid pressure is applied through the opening in the proximal side, through the balloon channel and against a proximal surface of the distal side of the sheath, thereby allowing fluid to pass through the balloon channel and into a vascular compartment distal to the balloon and the balloon sheath, and wherein the distal side of the sheath is also capable of deflecting proximally onto the distal balloon surface when a pressure is applied to the distal side of the sheath, thereby preventing proximally directed flow in the balloon channel.

5

10

15

20

25

30

35

- 8. The device of claim 7, wherein the interface between the distal side of the sheath and the distal balloon surface forms a unidirectional valve.
  - 9. The device of claim 7, wherein a catheter extension is disposed longitudinally from the distal end of the catheter body.
- 10. The catheter extension as in claim 9, wherein the catheter extension has a diameter less than the catheter body.
- 11. The device of claim 9, wherein the balloon and the balloon sheath are disposed on the catheter extension.
  - 12. The device of claim 9, wherein a nose piece is adapted to the catheter extension at a location distal to the balloon and the balloon sheath.
  - 13. The device of claim 12, wherein the balloon and the balloon sheath are positioned in a pocket between the distal end of the catheter body and a proximal end of the nose piece.
  - 14. The device of claim 9, wherein an outer diameter of the radially constrained balloon and the balloon sheath is less than or equal to an outer diameter of the catheter body.
    - 15. A method of embolizing a tumor, the method comprising:

advancing a device including a catheter body and a partial occlusion structure to a target tumor site within the body;

allowing an anterograde blood flow past the partial occlusion structure, the allowed anterograde blood flow being less than a blood flow that would normally be present if the partial occlusion structure were not in place;

injecting an embolic substance from the device to allow the anterograde blood flow to carry the embolic substance into a vasculature of the tumor target;

monitoring a real time pressure measurement in a vascular space distal to the partial occlusion structure;

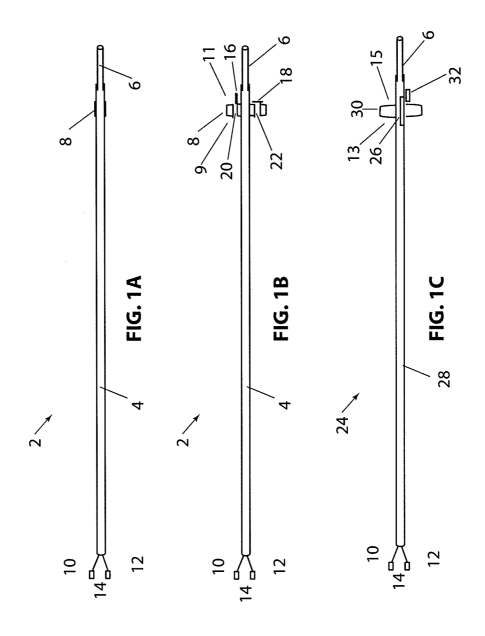
signaling a procedural endpoint based on the monitoring of the pressure measurement; and

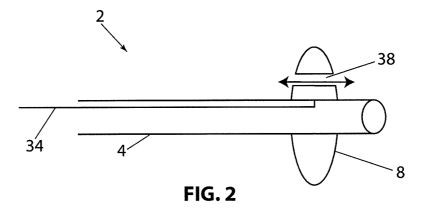
withdrawing the device from the body.

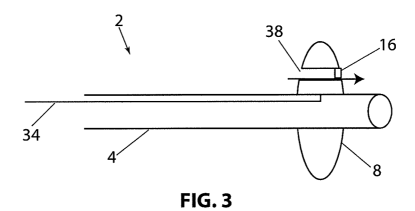
16. The method as in claim 15, wherein the allowed anterograde blood flow is within a range of about 1% to about 25% of a blood flow that would normally be present if the partial occlusion structure were not in place.

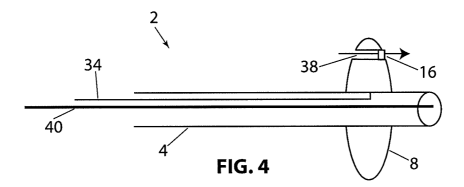
- 17. The method of claim 15, wherein the signaling of the procedural endpoint occurs when a predetermined pressure is reached in the vascular space distal to the partial occlusion structure.
  - 18. The method of claim 17, wherein the predetermined pressure is a predetermined percentage of an un-occluded systolic pressure.
    - 19. The method of claim 17, wherein the predetermined percentage is about 100%.

5









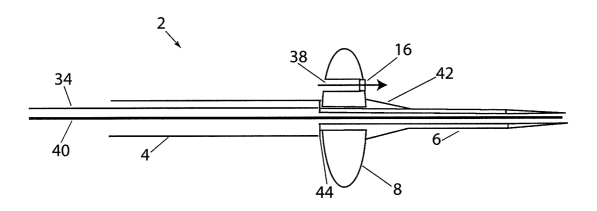
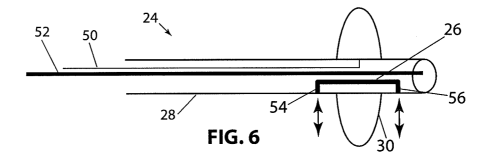
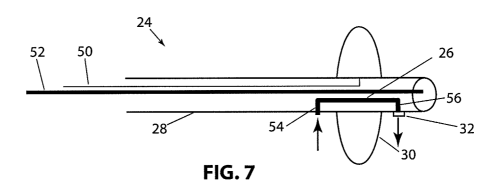
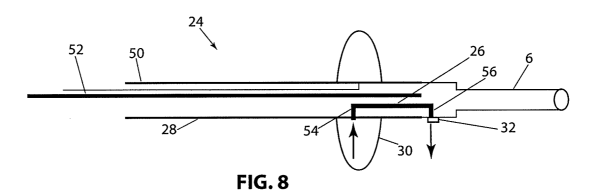
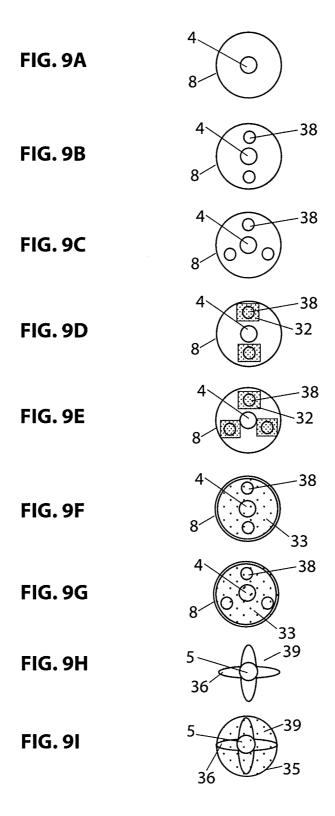


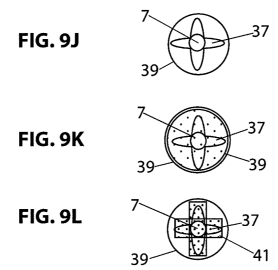
FIG. 5











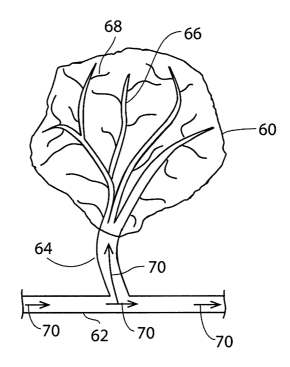


FIG. 10

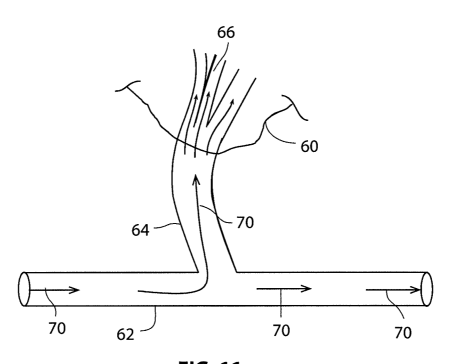


FIG. 11

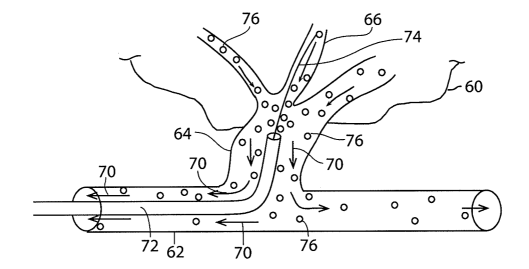


FIG. 12

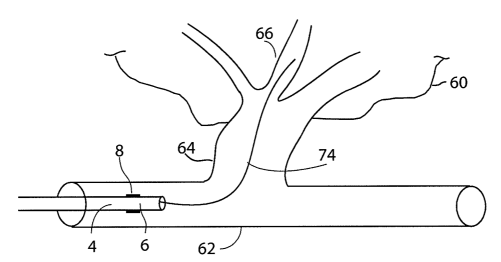


FIG. 13

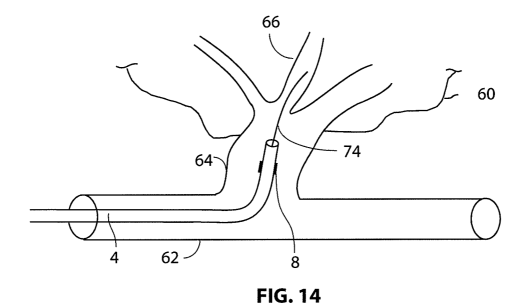


FIG. 15

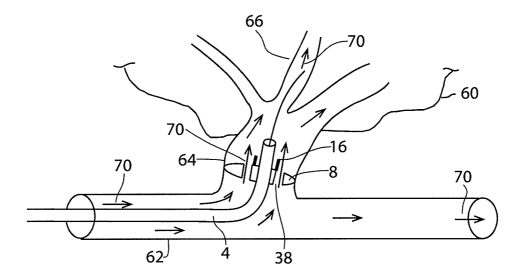


FIG. 16

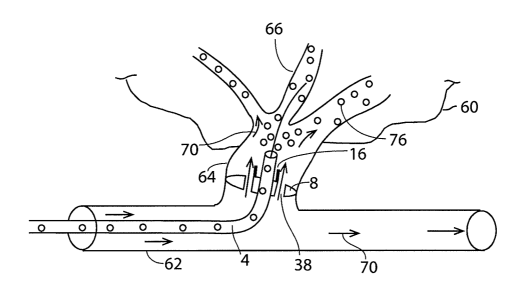
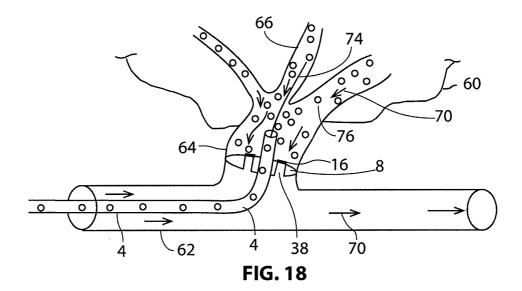
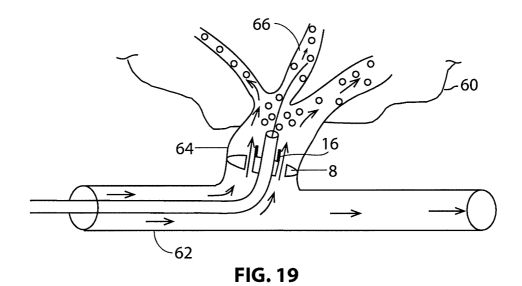
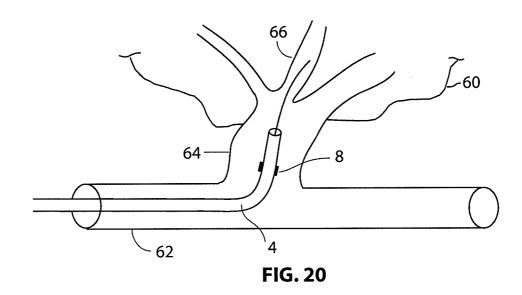
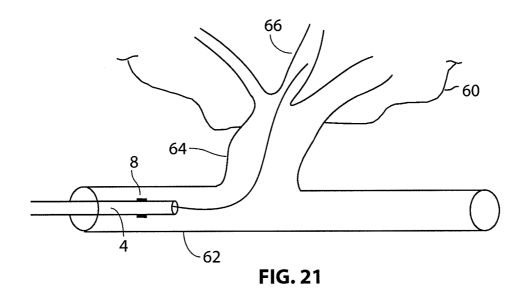


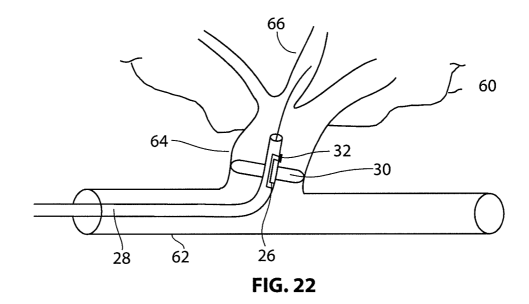
FIG. 17

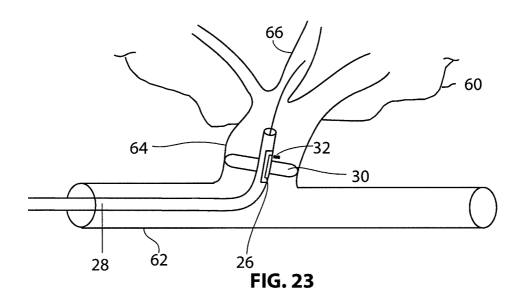


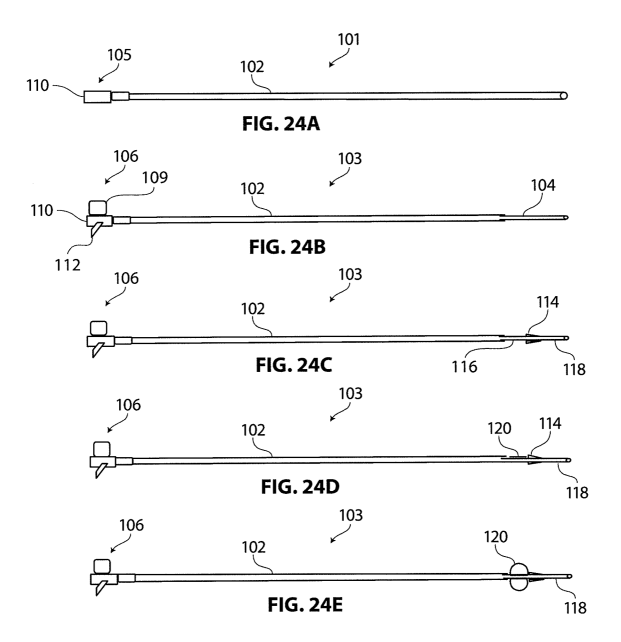












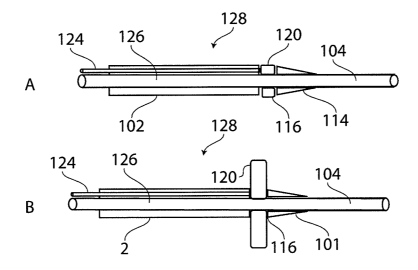


FIG. 25

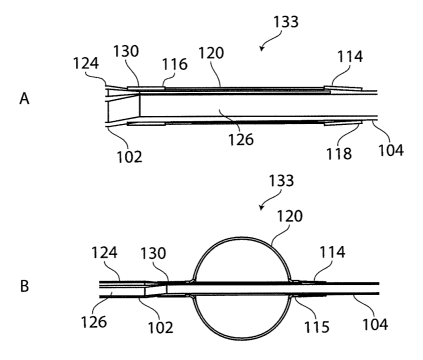


FIG. 26

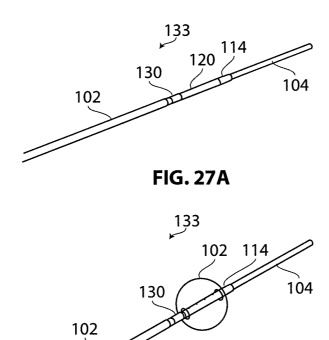


FIG. 27B

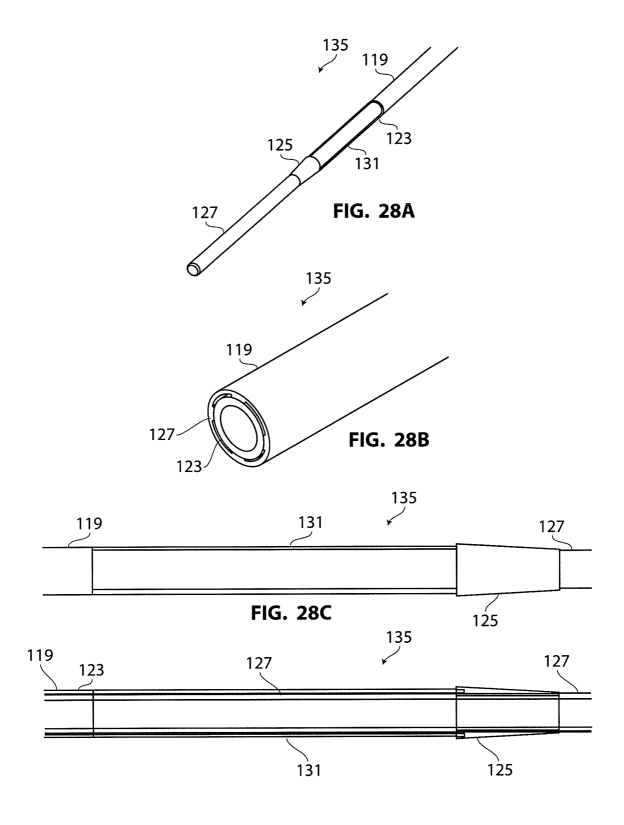


FIG. 28D

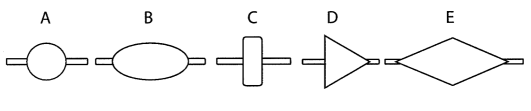


FIG. 29

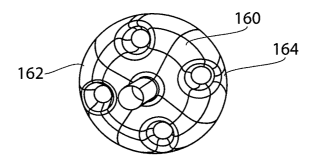


FIG. 30

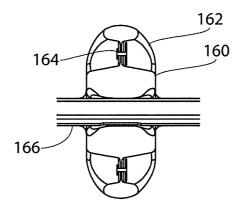
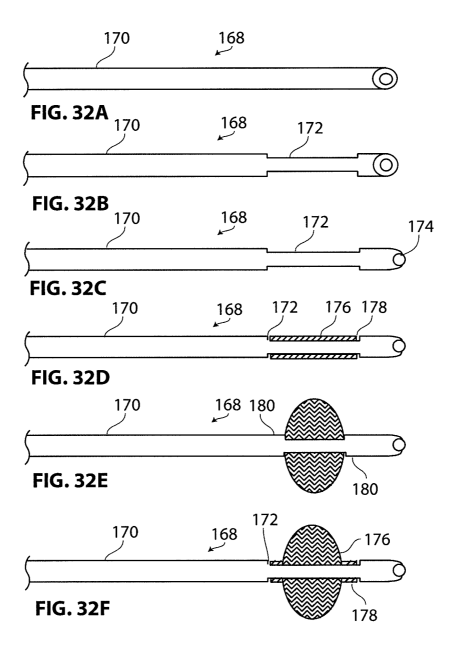
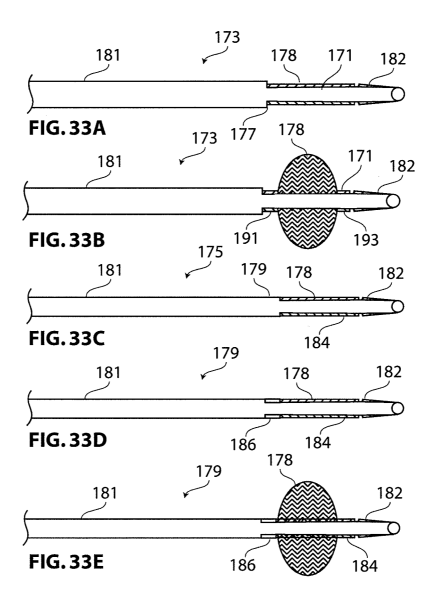
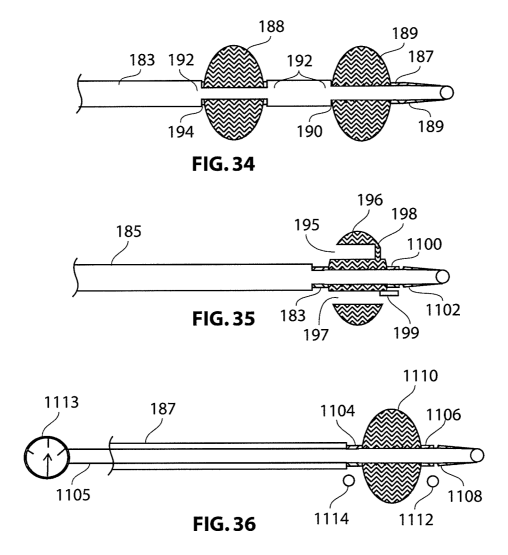
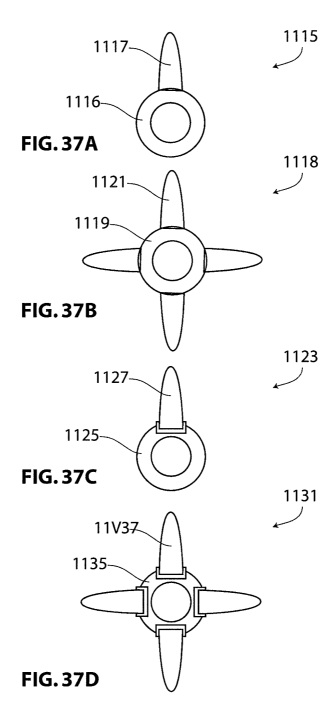


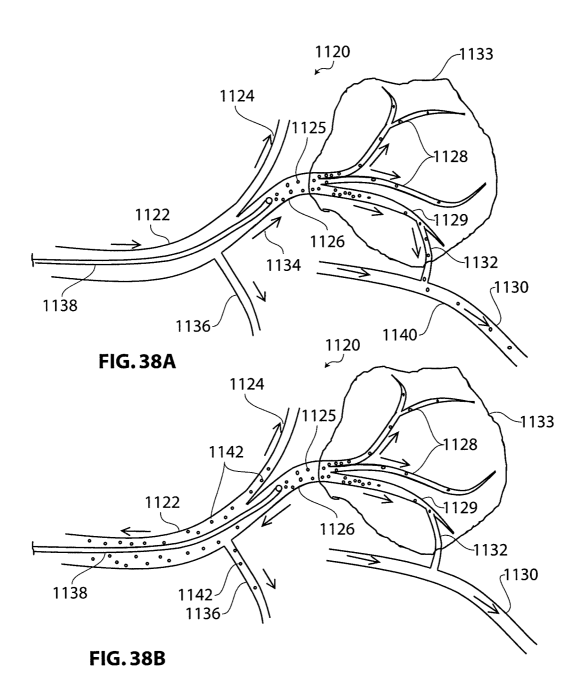
FIG. 31

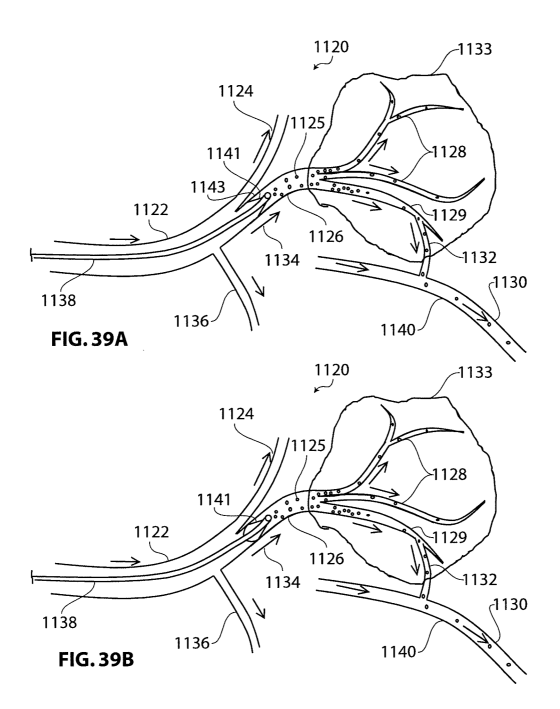


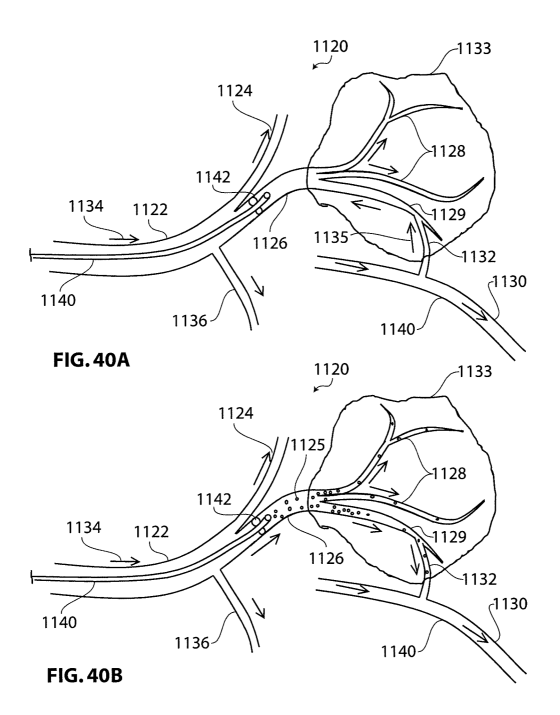




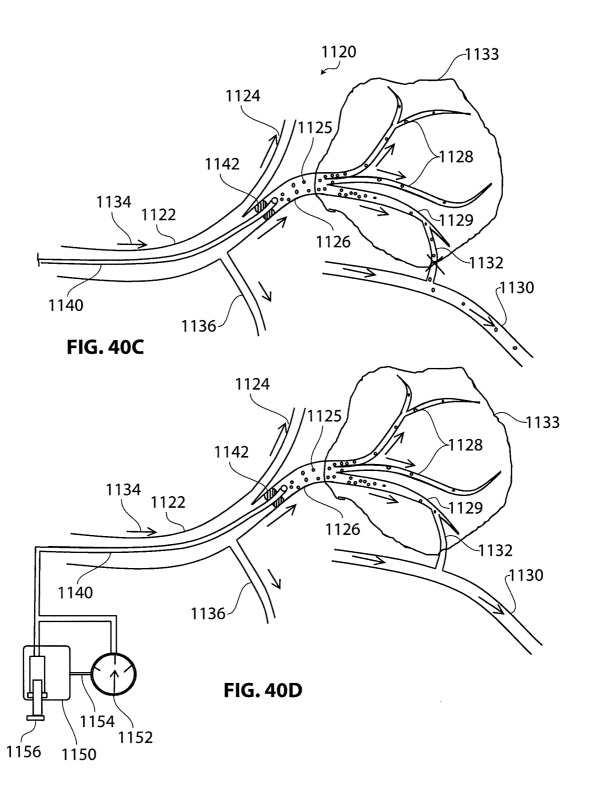


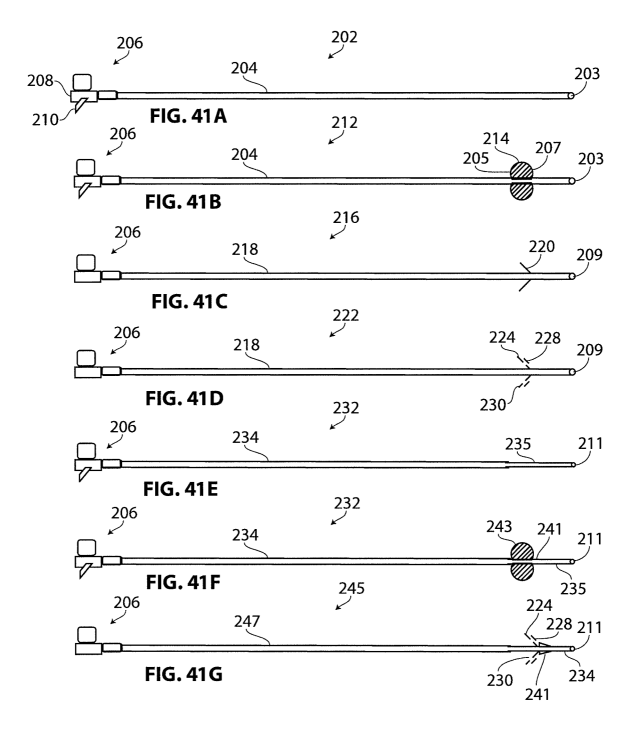


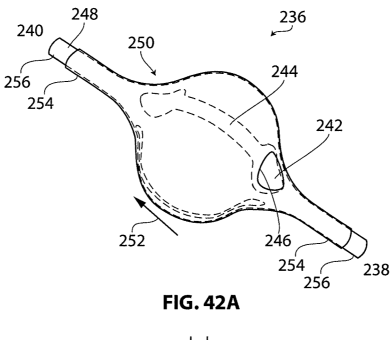




26 / 47







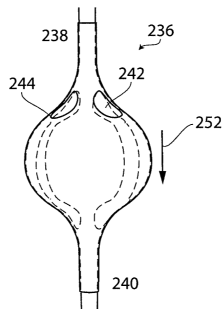
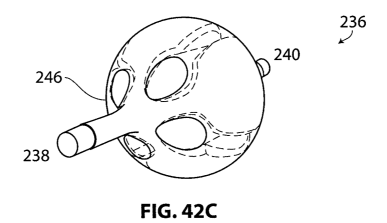
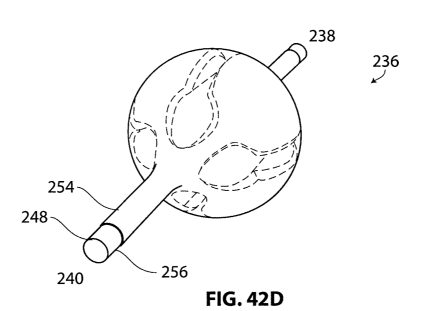


FIG. 42B





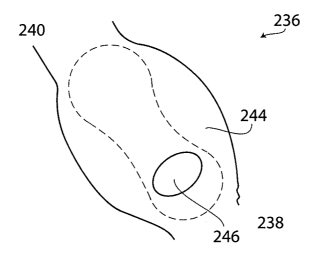


FIG. 43

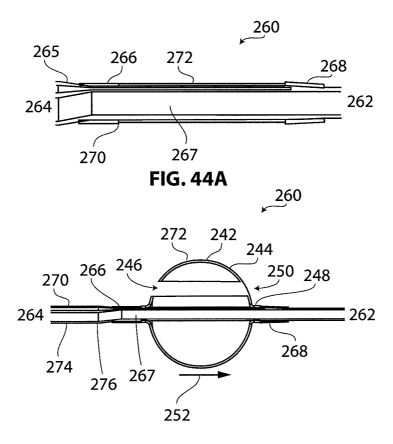
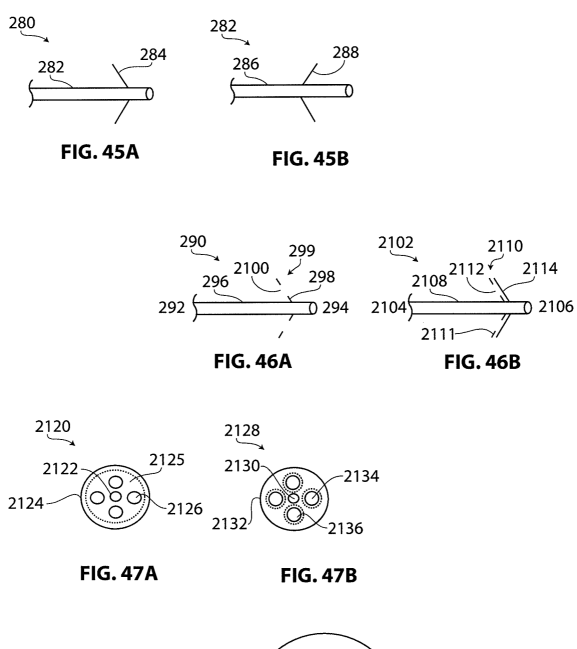
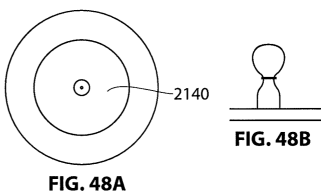
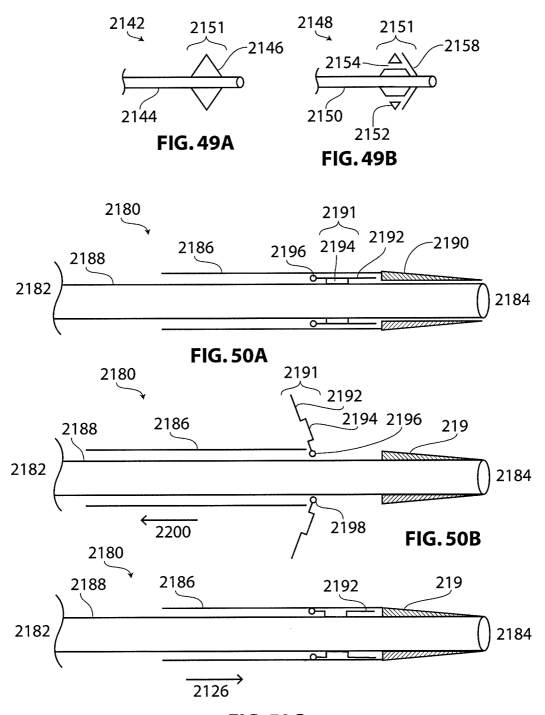


FIG. 44B







**FIG. 50C** 

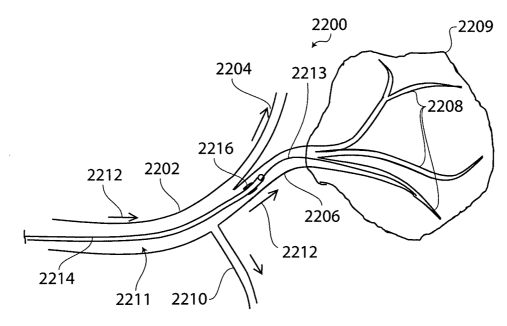
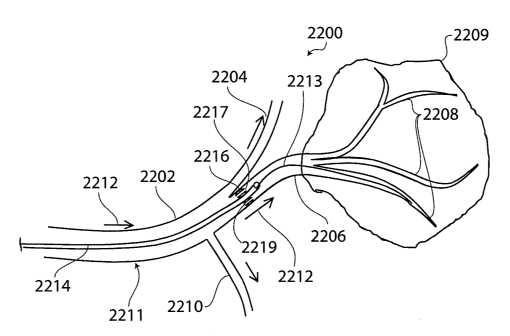


FIG. 51A



**FIG. 51B** 

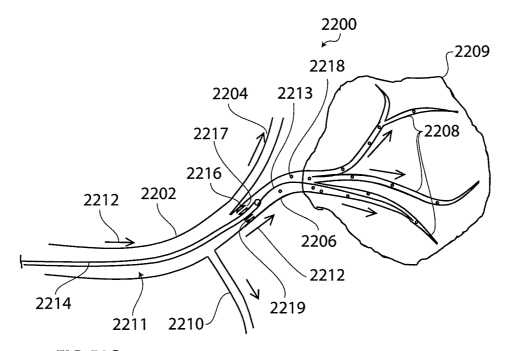


FIG. 51C

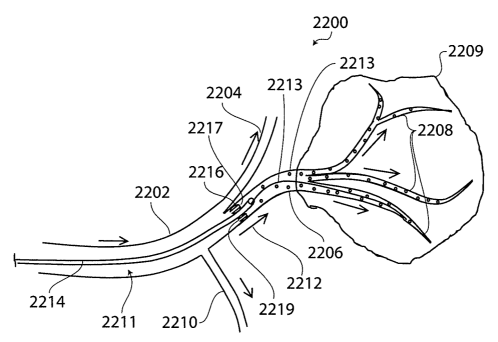


FIG.51D

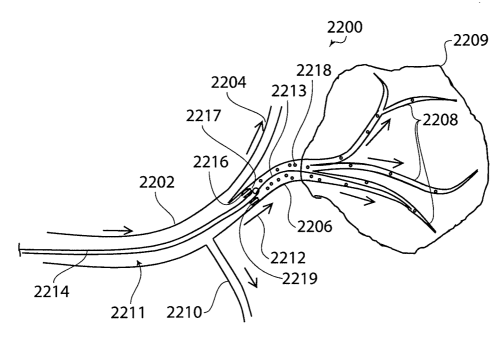


FIG. 51E

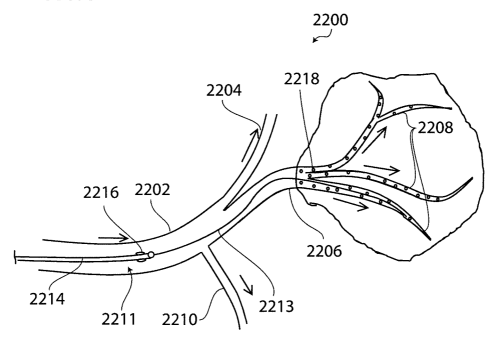


FIG. 51F

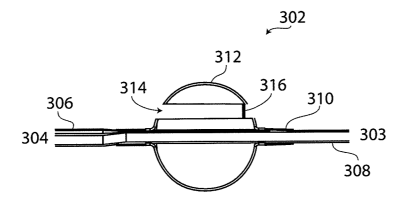


FIG. 52

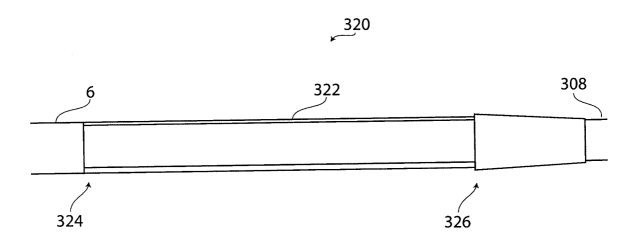


FIG. 53

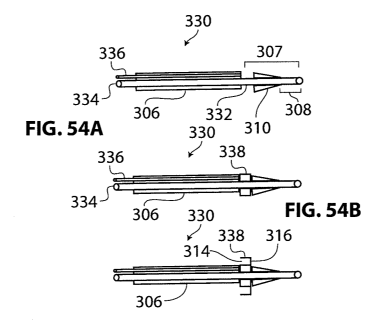
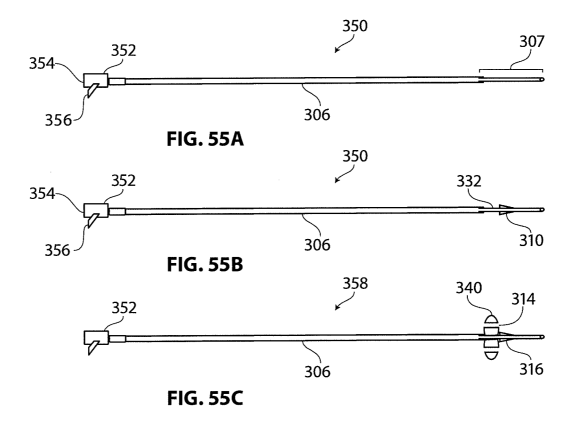
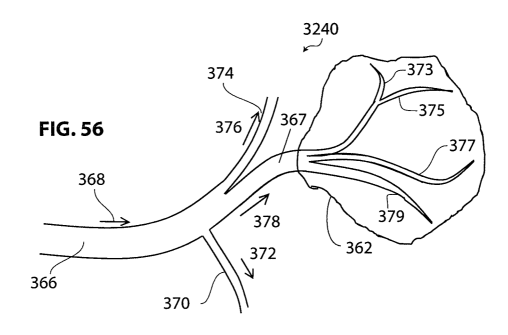
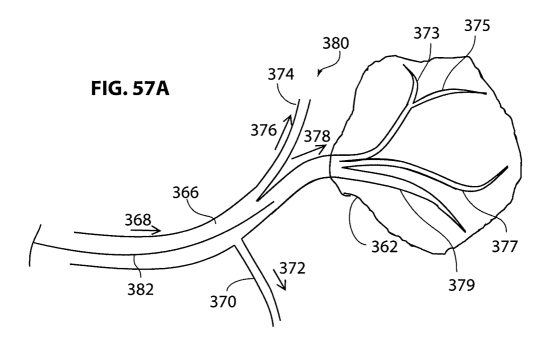
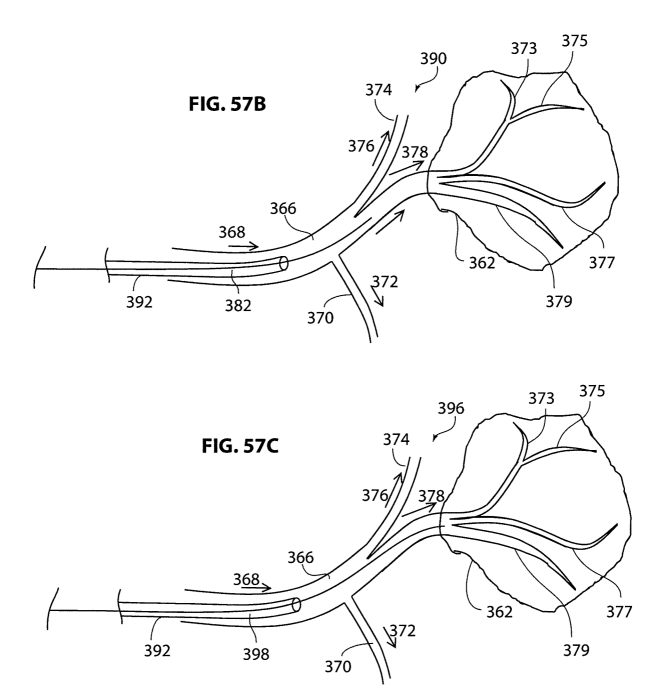


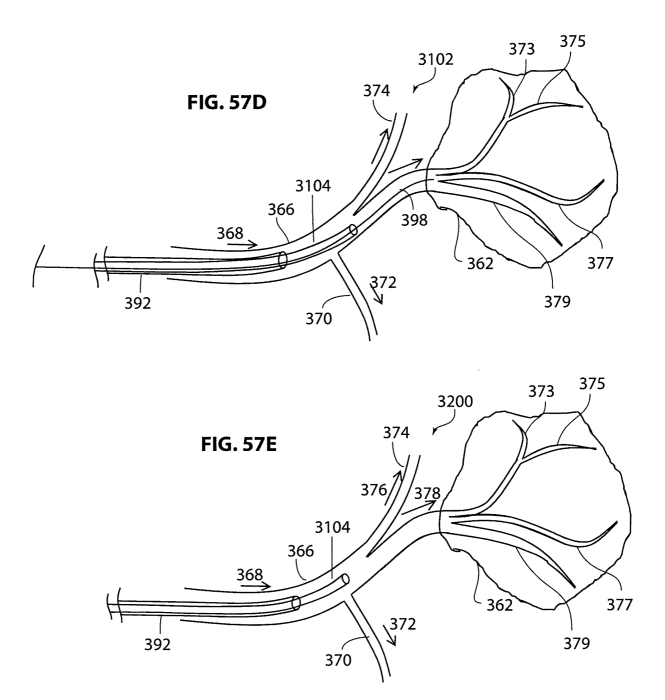
FIG. 54C

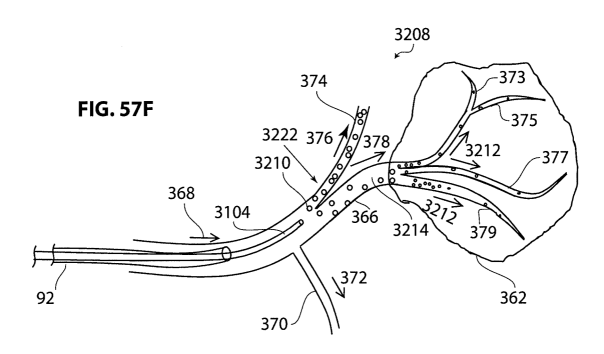


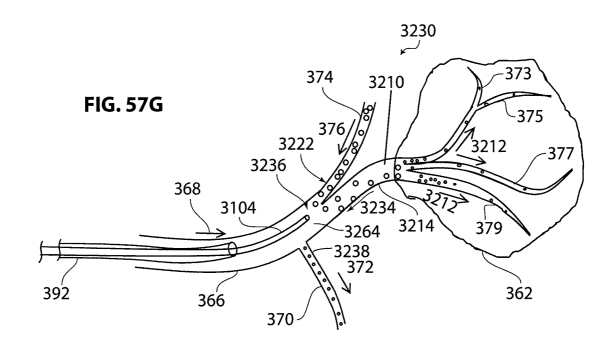












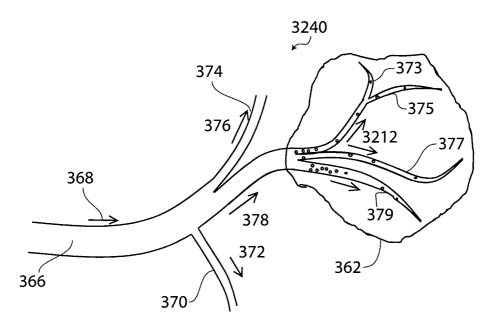
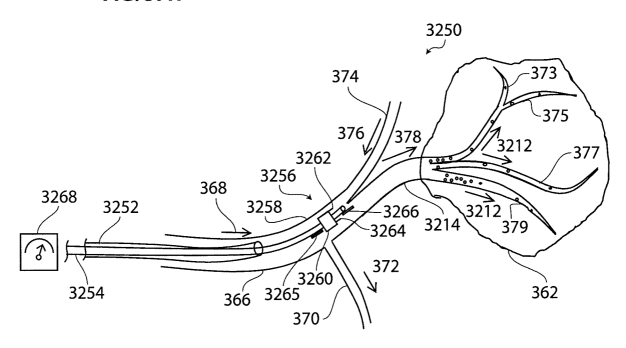
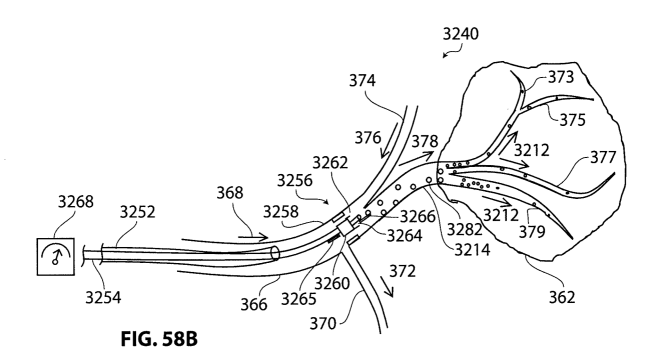
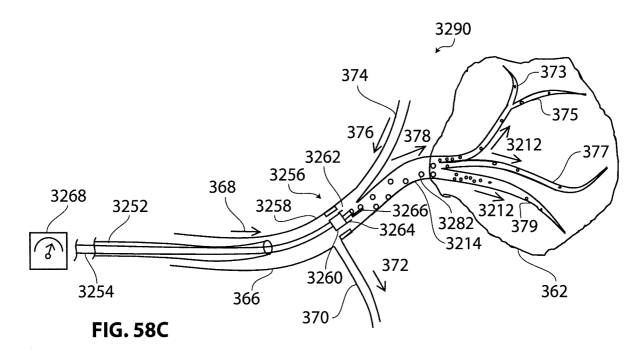


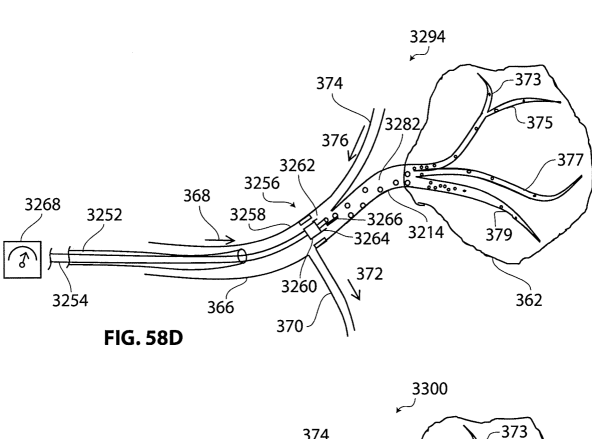
FIG. 57H

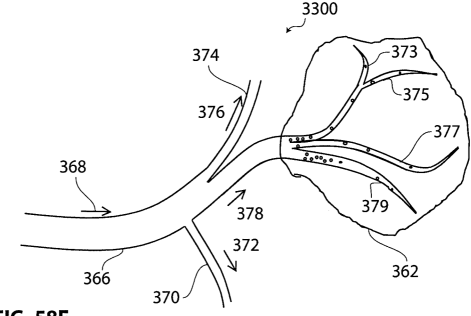


**FIG. 58A** 

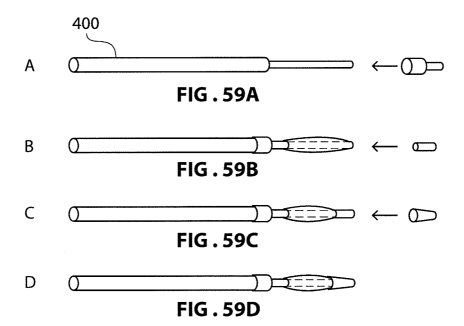








45 / 47



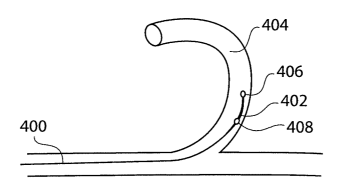
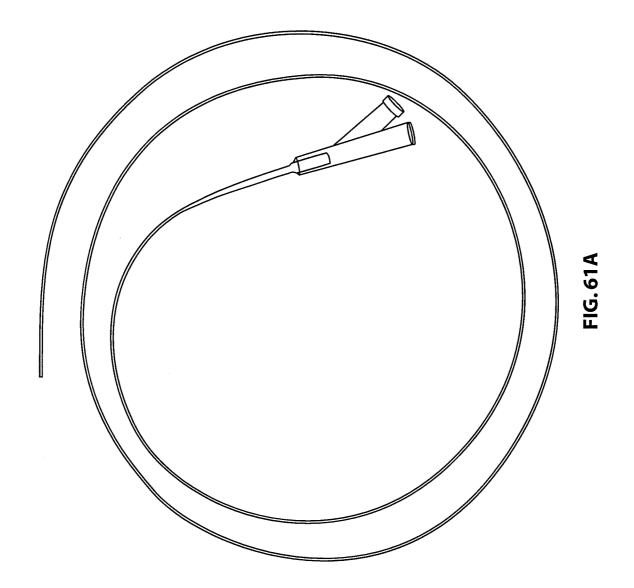
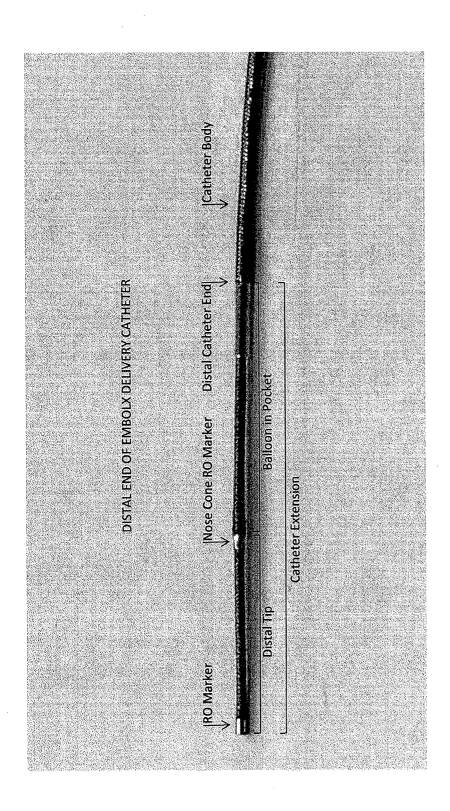


FIG.60





F(G. 6| B

一導管組件可具有一導管體及一可充氣氣球。該導管體具有一近端、一遠端及一氣球充氣腔。該可充氣氣球可附接至該導管體之遠端。該氣球 具有至少部分界定內部體積之一內表面。該氣球被建構以使得該內部體 積可與該導管體之充氣腔形成流體連通中以將該氣球充氣。該氣球亦具 有一近表面及一遠表面。該氣球具有一延伸穿過該氣球之通道。該通道 被建構以提供該氣球之近表面與該氣球之遠表面之間的流體連通。本發 明亦揭示其他導管組件及使用方法。