HA HEAT EXCHANGE BETWEEN THE SUBJECT'S FLOWING BLOOD AND THE HEAT EXCHANGE MEDIUM BEING CIRCULATED THROUGH THE TUBE.
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HEAT EXCHANGE CATHETERS AND THEIR METHODS OF MANUFACTURE AND USE

Related Application

[0001] This patent application claims priority to United States Provisional Patent Application No. 61/542,004 filed September 30, 2011, the entire disclosure of which is expressly incorporated herein by reference.

Field of the Invention

[0002] The present invention relates to medicine and biomedical engineering and more particularly to heat exchange catheter devices and their methods of manufacture and use.

Background of the Invention

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[0004] Hypothermia can be induced in humans and some animals for the purpose of protecting various organs and tissues (e.g., heart, brain, kidneys) against the effects of ischemic, anoxic or toxic insult. For example, animal studies and/or clinical trials suggest that mild hypothermia can have neuroprotective and/or cardioprotective effects in animals or humans who suffer from ischemic cardiac events (e.g., myocardial infarct, acute coronary syndromes, etc.), postanoxic coma after cardiopulmonary resuscitation, traumatic brain injury, stroke, subarachnoid hemorrhage, fever and neurological injury. Also, studies have shown that whole body hypothermia can ameliorate the toxic effects of radiographic contrast media on the kidneys (e.g., radiocontrast nephropathy) of patients with pre-existing renal impairment who undergo angiography procedures.
One method for inducing hypothermia is by endovascular temperature management (ETM) wherein a heat exchange catheter is inserted into a blood vessel and a thermal exchange fluid is then circulated through the heat exchange catheter. This technique can effectively cool blood flowing through the subject's vasculature and, as a result, lower the core body temperature of the subject to some desired target temperature. ETM is also capable of warming the body and/or of controlling body temperature to maintain a monitored body temperature at some selected temperature. If a controlled rate of re-warming or re-cooling from the selected target temperature is desired, that too can be accomplished by carefully controlling the amount of heat added or removed from the body and thereby controlling the temperature change of the patient.

A number of heat exchange catheters are currently available for use in ETM. One such catheter, the Solex™ Catheter available from ZOLL Circulation, Inc. of Sunnyvale, California, generally comprises a flexible catheter shaft having curved loops of tubing protruding from opposite sides of a distal portion of the catheter. In operation, the distal portion of the Solex™ catheter is inserted into the vasculature of a subject and a heated or cooled thermal exchange medium is then circulated through the curved loops of tubing. This results in exchange of heat between the circulating thermal exchange medium and blood flowing through the subject's vasculature, without causing the thermal exchange medium to be infused into the subject's bloodstream.

The current Solex™ catheter is formed of two subassemblies, a proximal portion and a distal portion. The proximal portion is a dual lumen tube that has an inflow lumen through which the thermal exchange fluid flows in the distal direction and an outflow lumen through which the thermal exchange fluid flows in the proximal direction. The distal portion is a coiled shaft with thin walled tubing "sewn" onto it to form the protruding curved loops of tubing. These proximal and distal portions are then joined together to make the final catheter. This two-piece construction requires a jointure or connection midway along the catheter shaft and is relatively labor intensive.

Additionally, the coiled shaft on the distal portion of the SolexTM catheter gives rise to a "bumpy" feel as the catheter is withdrawn through a
vascular introducer, such as during removal of the catheter from the subject's body. The elimination of the coiled distal shaft in favor of a smoother one-piece shaft could lessen or eliminate such "bumpy" feel as the catheter is being removed from the patient.  

[0009] Accordingly, there exists a need in the art for the development of new methods for catheter manufacture that may be used for the manufacture of the Solex™ catheter and/or other catheters having similar construction, thereby eliminating the need for formation of a jointure or connection midway along the catheter shaft and potentially offering other advantages, such as: 1) reducing the number of parts used in manufacturing the catheter and/or 2) reducing the labor and manual endeavor required for manufacture of the catheter, such as eliminating the need for the delicate sewing step to attach the thin-walled tubing to the coiled shaft and/or 3) reducing or eliminating the "bumpy" feel that the catheter has when being removed from the patient.

Summary of the Inventions

[0010] In accordance with the present invention, there is provided a catheter device having a one-piece or unitary catheter shaft with protruding loops of tubing which form a closed-loop recirculation circuit through which thermal exchange fluid or other flowable matter may be circulated.  

[0011] Further in accordance with the present invention, there is provided a catheter device comprising: a) an elongate catheter body having, a proximal end, a distal end, an inflow lumen and an outflow lumen; b) a plurality of bores extending transversely through the catheter body at spaced-apart locations; c) a tube having a first end connected to the inflow lumen and a second end connected to the outflow lumen, said tube passing in alternate transverse directions through adjacent ones of said bores such that tubing loops comprising portions of said tube protrude outwardly from opposite sides of the catheter body.  

[0012] Still further in accordance with the present invention, there is provided a method for manufacturing a catheter device, said method comprising the steps of: a) providing or obtaining an elongate catheter body having, a proximal end, a distal end, an inflow lumen and an outflow lumen; b) forming a plurality of bores transversely through the catheter body at spaced-
apart locations; c) passing a tube in alternate transverse directions through adjacent ones of said bores such that loops of the tube protrude outwardly from opposite sides of the catheter body; d) connecting a first end of the tube to the inflow lumen; and e) connecting a second end of the tube to the outflow lumen. The bores may be formed by any suitable means such as drilling, punching, laser cutting, blade cutting, water jet cutting, or other manual or automated techniques known in the art. In some embodiments, the bores may be formed by advancing a punch through the catheter body at an angle (e.g., substantially a right angle or any other desired angle) relative to a longitudinal axis of the catheter body. If oval or ovoid bores are desired, such punch may be of oval or ovoid shape.

[0013] Still further aspects and details of the present invention will be understood upon reading of the detailed description and examples set forth herebelow.

**Brief Description of the Drawings**

[0014] Figure 1 is a schematic diagram of an endovascular temperature management system of the present invention.

[0015] Figure 2 is a partial perspective view of a portion of a catheter body having a plurality of transverse bores formed therein during manufacture of a catheter device of the present invention.

[0016] Figure 3 is a partial perspective view of the catheter body portion of Figure 2 having a tube passed in alternating directions through the transverse bores such that loops of the tube protrude on opposite sides of the catheter body during manufacture of a catheter device of the present invention.

[0017] Figure 3A is a transverse cross sectional view through line 3A-3A of Figure 3.

[0018] Figure 3B is a transverse cross sectional view through line 3B-3B of Figure 3.

[0019] Figure 3C is a transverse cross sectional view through line 3C-3C of Figure 3.

[0020] Figure 4 is a side view of a fully assembled catheter device of the present invention.

[0021] Figure 4A is a distal end view of the catheter device of Figure 4.
Detailed Description of the Invention

[0022] The following detailed description and the accompanying drawings to which it refers are intended to describe some, but not necessarily all, examples or embodiments of the invention. The described embodiments are to be considered in all respects only as illustrative and not restrictive. The contents of this detailed description and the accompanying drawings do not limit the scope of the invention in any way.

[0023] Figure 1 is a schematic diagram of an endovascular temperature management system 10 which comprises a heat exchange catheter 12 of a type having heat exchange tubing loops 14, an extracorporeal control console C and at least one body temperature sensor TS. In this example, the extracorporeal control console C contains a controller (e.g., a microprocessor controller), a user interface UI for inputting data to the controller, a heater/cooler for adjusting the temperature of a thermal exchange medium (e.g., 0.9% sodium chloride solution) and a pump for pumping the thermal exchange medium.

[0024] The catheter 12 is connected to the extracorporeal control console C by way of an inflow line IL and an outflow line OL so that the pump within the console C will circulate temperature-controlled thermal exchange medium through heat exchange tubing loops 14 which protrude from a distal portion of the heat exchange catheter 12. In operation, the catheter 12 is inserted into the vasculature of a subject so that blood flowing through the subject's vasculature will pass over and in heat exchange proximity with the heat exchange tubing loops 14, thereby allowing heat to be exchanged between the circulating thermal exchange medium and the subject's flowing blood. Warming or cooling of the subject's flowing blood then results in warming or cooling of all or a desired portion of the subject's body. In the particular non-limiting example shown in Figure 1, the distal portion of the catheter 12 is positioned so that the heat exchange tubing loops 14 reside within the subject's inferior vena cava IVC, such catheter positioning being suitable for applications wherein whole body temperature management is desired.

[0025] The temperature sensor(s) TS may be positioned on or in the subject's body to measure the temperature of all or part of the body where it is desired to effect temperature modification or control. The controller within the
console C receives signals from the temperature sensor(s) TS indicating the currently sensed body temperature. A desired target temperature may be input via the user interface UI and the controller will then issue control signals to the heater cooler and/or pump to adjust the temperature and/or the flowate of the heat exchange medium in an effort to attain and/or maintain the target body temperature. A control console of the type shown in Figure 1 and described in this example is commercially available as the Thermogard XP™ Temperature Management System from ZOLL Circulation of Sunnyvale, California. Other potentially useable control consoles are available from other sources and/or described in the following United States patents and published patent applications, the entire disclosures of which are expressly incorporated herein by reference:

[0026] The catheter 12 of this example is constructed and manufactured in the manner shown in Figures 2-4A. A one-piece catheter body 12 is used in this example. As may be appreciated from the cross-sectional views of Figures 3A-3C, the catheter body 12 of this example has an inflow lumen 17, an outflow lumen 15, an optional distal infusion or guidewire lumen 16, an optional medial infusion lumen 18 and an optional proximal infusion lumen 19. The individual lumens 15, 16, 17, 18, 19 may be integrally formed (e.g., extruded) within the catheter body 12 or may comprise one or more separate tube(s) that are passed through a lumen of the catheter body 12. In this particular example, the one-piece catheter body is formed of an extruded multilumen tube. Also, in this example, the catheter body 12 has an optional atraumatic distal tip member 11 attached to its distal end.

[0027] At the time of manufacture, a series of transverse bores 13 are formed through the catheter body 12, as shown in Figure 2. These transverse bores 13 may be formed at any suitable angle relative to the longitudinal axis LA of the catheter body 12. In the particular example shown, the bores 13 are oval or ovoid shaped bores formed by advancing an oval or ovoid shaped punch through the catheter body 12 on a predetermined trajectory and such bores extend through the catheter body 12 substantially at right angles relative to the longitudinal axis LA of the catheter body 12. The trajectory of the bores 13 will avoid obliteration of any lumen(s) that are required to remain in tact to carry apparatus or thermal exchange media through that portion of
the catheter body 12. In this example, optional medial infusion lumen 18 and optional proximal infusion lumen 19 are terminally sealed to proximal and medial infusion lumen outlet openings 44, 46 (seen on Figure 4), which are located proximal to where the bores 13 are formed. Therefore, the medial and proximal infusion lumens 18, 19 are nonfunctional in the region where the bores 13 are formed. Also, in this example, the outflow lumen 15 is terminally sealed to the proximal end of tube 14 at window 20. Thus, the portion of outflow lumen 15 that extends through the region where the bores 13 are formed is non-function. Thus, in this example, the positioning of the catheter body 12 and the trajectory of the punch are controlled so that each bore 13 only obliterates unused portions of the outflow lumen 15, optional medial infusion lumen 18 and optional proximal infusion lumen 19, while leaving the inflow lumen 15 and optional distal infusion or guidewire lumen 16 in tact.

[0028] The catheter body 12 may be appropriately sized and formed of any material(s) suitable for the intended applications of the catheter device. For example, in many applications, it will be desirable for the catheter body 12 to have enough rigidity and wall thickness to contain working pressures of up to about 100 psi while being sufficiently flexible to navigate through the intended blood vessels or other body lumens to the desired location within a subject's body. Typically, this may be accomplished by a catheter body that has an outer diameter of 6 Fr (.080") to 14 Fr (.180") and is formed of a biocompatible polyurethane (e.g., Elastollan™ available from BASF Corporation, Florham Park, NJ or Tecothane™ available from The Lubrizol Corporation, Wickliffe, OH) or polyether block amide (e.g., Pebax™ available from Arkema, Inc., Philadelphia, PA).

[0029] The heat exchange tube 14 may be appropriately sized and formed of any material(s) suitable for the intended applications of the catheter device. For example, in many applications, it will be desirable for the heat exchange tube 14 to a) have a thin wall thickness (typically around .001") to best facilitate heat transfer, b) have sufficient tensile strength to withstand pressures of up to about 100 psi and c) be sufficiently rigid or semi-rigid so as not to expand uncontrollably under pressure. Thus, it will be desirable for the tube 14 to be formed of a material capable of being extruded and/or blown into a tube having such wall thickness and properties. Examples of materials
that may be suitable for forming the tube 14 include polyethylene terephthalates (PETs) available from a variety of sources or polyether block amide (e.g., Pebax™ available from Arkema, Inc., Philadelphia, PA).

[0030] Also, as seen in Figure 2, a window into the inflow lumen 22 is skived or otherwise formed at a first location in a wall of the catheter body 12 and a window into the outflow lumen 20 is skived or otherwise formed at a second location in a wall of the catheter body 12. Thereafter, as illustrated Figure 3, one end of a tube 14 is passed in alternate transverse directions (e.g., back and forth) through adjacent ones of the bores 13 so that generally U shaped loops of the tube 14 protrude outwardly from opposite sides of the catheter body. A first end (in this example the distal end) of the tube 14 is then inserted through window 22 into inflow 17 lumen and secured to the wall of the inflow lumen 17. This forms a sealed connection through which inflowing thermal exchange medium will flow from the inflow lumen 17 into the first (e.g., distal) end of the tube. Sealing attachment of the tube 14 to the luminal wall of the inflow lumen 17 may be accomplished by any suitable means such as heat sealing or by adhesive bonding. Examples of adhesives that are useable for this purpose include but are not necessarily limited to cyanoacrylate adhesives (e.g. Loctite 4011) available from the Henkel Corporate, Westlake, OH, UV curing acrylic adhesives (e.g., e.g. Loctite 3311) available from Henkel Corporate, Westlake, OH, and epoxy adhesives (e.g. Loctite 3981) available from Henkel Corporate, Westlake, OH). A second end of the tube 14 (in this example the proximal end) is inserted through window 20 into outflow lumen 15 and secured to the wall of the outflow lumen 15 in the same manner as described above. This forms a sealed connection through which thermal exchange medium will flow out of the tube 14 and into the outflow lumen 15. By this arrangement, thermal exchange fluid will enter the distal end of the tube 14 and will exit the proximal end of the tube 14. However, in some embodiments, it may be desirable to reverse the connections such that the proximal end of the tube 14 will be connected to the inflow lumen 17 and the distal end of the tube 14 will be connected to the outflow lumen 15. In such alternative embodiments, the trajectory of the bores 13 may obliterate the distal (unused) portion of the inflow lumen 17 while leaving that portion of the outflow lumen 15 in tact.
In some embodiments, it will be desirable to form the protruding U shaped loops of tubing into desired shapes by thermosetting or other suitable forming techniques. As shown in Figures 4 and 4A, in this example, the protruding U shaped loops of tubing 14 are formed so as to circumferentially curve around one side of the catheter body 12, as shown. Also, as shown in Figures 4 and 4A, when fully assembled, the catheter device of this invention includes a hub 30 on its proximal end, with an outflow lumen connector 32 (connected to outflow lumen 17), inflow lumen connector 34 (connected to inflow lumen 15), medial infusion lumen connector 36 (connected to medial infusion lumen 18), distal infusion of guidewire lumen connector 36 (connected to the distal infusion/guidewire lumen 16) and proximal infusion lumen connector 36 (connected to proximal infusion lumen 19) extending therefrom. Optional graduated distance markings 42 are formed on a proximal region of the catheter body 12 to indicate the length of catheter that is indwelling in the body at any particular time. Also, an optional proximal radiographic marker 48 and an optional distal radiographic marker 50 are located on the catheter body to facilitate radiographic determination of the location of the heat exchanging region (e.g., the protruding tube loops 14) within a subject's body.

It is to be appreciated that the invention has been described hereabove with reference to certain examples or embodiments of the invention but that various additions, deletions, alterations and modifications may be made to those examples and embodiments without departing from the intended spirit and scope of the invention. For example, any element or attribute of one embodiment or example may be incorporated into or used with another embodiment or example, unless otherwise specified if to do so would render the embodiment or example unsuitable for its intended use. Also, where the steps of a method or process have been described or listed in a particular order, the order of such steps may be changed unless otherwise specified or unless doing so would render the method or process unworkable for its intended purpose. All reasonable additions, deletions, modifications and alterations are to be considered equivalents of the described examples and embodiments and are to be included within the scope of the following claims.
Claims

What is claimed is:

1. A catheter device comprising:
   
a) an elongate catheter body having, a proximal end, a distal end, an inflow lumen and an outflow lumen;
   
b) a plurality of bores extending transversely through the catheter body at spaced-apart locations;
   
c) a tube having a first end connected to the inflow lumen and a second end connected to the outflow lumen, said tube passing in alternate transverse directions through adjacent ones of said bores such that tubing loops comprising portions of said tube protrude outwardly from opposite sides of the catheter body.

2. A catheter device according to claim 1 wherein the catheter body further comprises a through tube or through lumen that extends through at least a portion of the catheter body and which terminates distally in a distal port.

3. A catheter device according to claim 2 wherein the through tube or through lumen is useable as a guidewire passageway to facilitate advancement of the catheter body over a guidewire.

4. A catheter device according to claim 2 wherein the through tube or through lumen is useable as a delivery passageway to facilitate delivery of a therapeutic substance or device through the catheter device and into the subject's body.

5. A catheter device according to page 1 wherein the tubing loops are formed to curved shapes.
6. A catheter device according to claim 1 wherein a window into the inflow lumen is formed at a first location in a wall of the catheter and the first end of the tube extends through that window into the inflow lumen and is secured to a wall of the inflow lumen.

7. A catheter device according to claim 6 wherein a window into the outflow lumen is formed at a second location in a wall of the catheter and the second end of the tube extends through that window into the outflow lumen and is secured to a wall of the outflow lumen.

8. A catheter device according to claim 7 wherein the window into the outflow lumen is located distal to the window into the inflow lumen such that fluid will progress through the loops of tubing in a distal direction.

9. A catheter device according to claim 7 wherein the window into the outflow lumen is located proximal to the window into the inflow lumen such that fluid will progress through the loops of tubing in a proximal direction.

10. A catheter device according to claim 1 wherein the bores are ovoid or oval in shape.

11. A catheter device according to claim 1 wherein the first end of the tube is connected to the inflow lumen at a location proximal to the transverse bores and the second end of the tube is connected to the outflow lumen at a location distal to the transverse bores.

12. A catheter device according to claim 11 wherein the transverse bores extend through the catheter body in a manner that obliterates adjacent portions of the inflow lumen but leaves adjacent portions of the outflow lumen in tact.

13. A catheter device according to claim 1 wherein the first end of the tube is connected to the inflow lumen at a location distal to the transverse bores and the second end of the tube is connected to the outflow lumen at a location proximal to the transverse bores.
14. A catheter device according to claim 13 wherein the transverse bores extend through the catheter body in a manner that obliterates adjacent portions of the outflow lumen but leaves adjacent portions of the inflow lumen in tact.

15. A system comprising a catheter device according to claim 1 in combination with apparatus for circulating heated or cooled thermal exchange fluid though the catheter device.

16. A system according to claim 15 further comprising a temperature sensor useable for sensing the temperature of all or a portion of a subject's body.

17. A system according to claim 16 further comprising a controller which controls the temperature and/or flowrate of thermal exchange that is circulated through the catheter device.

18. A system according to claim 17 wherein the controller receives a) an input target body temperature and b) signals from the temperature sensor indicative of the sensed body temperature and wherein the controller is programmed to vary the temperature and/or flowrate of the thermal exchange fluid in a manner that causes or attempts to cause the sensed body temperature to be the same as or within an acceptable range of the target body temperature.

19. A method for manufacturing a catheter device, said method comprising the steps of:

A) providing or obtaining an elongate catheter body having, a proximal end, a distal end, an inflow lumen and an outflow lumen;

B) forming a plurality of bores transversely through the catheter body at spaced-apart locations;

C) passing a tube in alternate transverse directions through adjacent ones of said bores such that loops of the tube protrude outwardly from opposite sides of the catheter body;
D) connecting a first end of the tube to the inflow lumen; and

E) connecting a second end of the tube to the outflow lumen.

20. A method according to claim 19 wherein the catheter body further comprises a through tube or through lumen that extends through at least a portion of the catheter body and terminates distally in a distal port.

21. A method according to claim 19 further comprising the step of forming tubing loops to desired shapes.

22. A method according to claim 21 wherein the tubing loops are formed to curved shapes.

23. A method according to claim 21 wherein the tubing loops are thermoset in desired shapes.

24. A method according to claim 19 wherein step D comprises:

    forming a window into the inflow lumen at a first location in a wall of the catheter;

    inserting the first end of the tube through the window into the inflow lumen; and,

    securing the tube to a wall of the inflow lumen.

25. A method according to claim 24 wherein the window is formed by skiving a window into the inflow lumen at a first location in a wall of the catheter.

26. A method according to claim 24 wherein step E comprises:

    forming a second window into the outflow lumen at a second location in a wall of the catheter;

    inserting the second end of the tube through the second window into the outflow lumen; and,
securing the second tube to a wall of the outflow lumen.

27. A method according to claim 26 wherein the second location is distal to the first location.

28. A method according to claim 26 wherein the second location is proximal to the first location.

29. A method according to claim 26 wherein the first location is proximal to the transverse bores and the second location is distal to the transverse bores.

30. A method according to claim 26 wherein the first location is distal to the transverse bores and the second location is proximal to the transverse bores.

31. A method according to claim 29 wherein the transverse bores are formed in a manner that obliterates adjacent portions of the inflow lumen but leaves adjacent portions of the outflow lumen in tact.

32. A method according to claim 30 wherein the transverse bores are formed in a manner that obliterates adjacent portions of the outflow lumen but leaves adjacent portions of the inflow lumen in tact.

33. A method according to claim 19 wherein the bores formed in Step B are oval or ovoid in shape.

34. A method according to claim 19 wherein a longitudinal axis is projectable through the catheter body and wherein the bores are formed in Step B by advancing a punch through the catheter body at an angle relative to the longitudinal axis of the catheter body.

35. A method according to claim 34 wherein the punch is advanced at substantially a right angle to the longitudinal axis of the catheter body.

36. A method according to claim 34 wherein the punch has an oval or ovoid shape which causes the bores to be oval or ovoid in shape.
INTERNATIONAL SEARCH REPORT

International application No.
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A. CLASSIFICATION OF SUBJECT MATTER
IPC(8) - A61 F 7/12 (2012.01)
USPC - 607/105

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)
IPC(8) - A61B 18/01; A61F 7/00, 7/12 (2012.01)
USPC - 606/21; 607/105, 113

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

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

MicroPatent

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
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* Further documents are listed in the continuation of Box C.

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Name and mailing address of the ISA/US
Mail Stop PCT, Attn: ISA/US, Commissioner for Patents
P.O. Box 1450, Alexandria, Virginia 22313-1450

Facsimile No. 571-273-3201

Authorized officer: Blaine R. Copenheaver
PCT Helpdesk: 571-272-4000
PCT OSP: 571-272-7774

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