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Bearingless implantable blood pump

Abstract

Implantable blood pumps and related methods employ a compact rotary motor. The compact rotary motor includes a stator and a rotor. The stator is disposed within a housing circumferentially about a dividing wall such that a blood flow conduit extends through the stator. The stator is disposed circumferentially around at least a portion of the rotor.

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Background/Summary

CROSS REFERENCE TO RELATED APPLICATIONS (1) The present application is a Continuation of U.S. patent application Ser. No. 16/204,015, now U.S. Pat. No. 10,973,967, filed Nov. 29, 2018 (Allowed); which claims the benefit of U.S. application No. 62/615,708 filed Jan.

10, 2018, the full disclosures which are incorporated herein by reference in their entirety for all purposes.

BACKGROUND

(1) Ventricular assist devices, known as VADs, often include an implantable blood pump and are used for both short-term (i.e., days, months) and long-term applications (i.e., years or a lifetime) where a patient's heart is incapable of providing adequate circulation, commonly referred to as heart failure or congestive heart failure. According to the American Heart Association, more than five million Americans are living with heart failure, with about 670,000 new cases diagnosed every year. People with heart failure often have shortness of breath and fatigue. Years of living with blocked arteries and/or high blood pressure can leave a heart too weak to pump enough blood to the body. As symptoms worsen, advanced heart failure develops.

(2) A patient suffering from heart failure may use a VAD while awaiting a heart transplant or as a long term destination therapy. A patient may also use a VAD while recovering from heart surgery. Thus, a VAD can supplement a weak heart (i.e., partial support) or can effectively replace the natural heart's function.

BRIEF SUMMARY

(3) The following presents a simplified summary of some embodiments of the invention in order to provide a basic understanding of the invention. This summary is not an extensive overview of the invention. It is not intended to identify key/critical elements of the invention or to delineate the scope of the invention. Its sole purpose is to present some embodiments of the invention in a simplified form as a prelude to the more detailed description that is presented later.

(4) In many embodiments, an implantable blood pump includes a rotary motor that includes a compact stator assembly. The compact size of the stator assembly is enabled by the stator assembly including a compact stator core, which includes a toroidal portion, and stator coils. Each of the stator coils extend around a respective separated segment of the toroidal portion. In many embodiments, the stator does not extend beyond a disk-shaped volume having a compact thickness (e.g., less than 1.0 inches) in a direction parallel to the axis of rotation of the rotary motor), thereby enabling the stator assembly to have a corresponding compact thickness parallel to the axis of rotation of the rotary motor. In some embodiments, the stator core includes separated stator teeth that extend inwardly from the toroidal portion between adjacent pairs of the stator coils. In some embodiments, the rotary motor includes rotor position sensors (e.g., hall effect sensors). Each of the rotor position sensors can be disposed in or adjacent to a respective gap between adjacent pairs of the stator coils. The compact size of the stator assembly parallel to the axis of rotation of the rotary motor enables the implantable blood pump to have a compact size parallel to the axis of rotation of the rotary motor, thereby requiring less space within the thoracic cavity.

(5) Thus, in one aspect, a first implantable blood pump includes a housing and a rotary motor. The housing defines an inlet opening, an outlet opening, and a dividing wall within the housing defining a blood flow conduit. The blood flow conduit extends between the inlet opening and the outlet opening. The rotary motor includes a stator and a rotor. The stator includes a stator core and stator coils. The stator core includes a toroidal portion and stator teeth. Each of the stator teeth extend toward the rotor from the toroidal portion. Each of the stator teeth is separated from each of an adjacent two of the stator teeth by a respective adjacent intervening segment of the toroidal portion. Each of the stator coils extends around one of the intervening segments of the toroidal portion. The stator is disposed within the housing circumferentially about the dividing wall such that the blood flow conduit extends through the stator core. The stator core is disposed circumferentially around at least a portion of the rotor. The rotor has a rotor axis of rotation and includes a rotor magnet for driving the rotor. The stator teeth axially overlap the rotor magnet with respect to the rotor axis of rotation. In many embodiments, the stator does not extend beyond a disk-shaped volume having a compact thickness (e.g., less than 1.0 inches) in a direction parallel to the rotor axis of rotation.

- (6) In many embodiments, the first implantable blood pump is configured to pump blood from a heart ventricle to the aorta. In some embodiments, the outlet opening is oriented at an angle relative to the input opening. The inlet opening can be oriented to receive blood directly from a heart ventricle and the output opening oriented to output blood in a direction transverse to the orientation of the inlet opening so as to reduce the length of a blood flow cannula used to transfer the blood flow from the output opening to the aorta. The rotor can include centrifugal pump impeller blades.
- (7) In many embodiments of the first implantable blood pump, the rotor defines a rotor blood flow conduit that extends through the stator. For example, in many embodiments, the rotor defines a rotor blood flow conduit that extends through the rotor, thereby extending through the stator.
- (8) The rotor can have any suitable number of magnetic moments. In some embodiments, the rotor has only one magnetic moment.
- (9) In some embodiments, the first implantable blood pump includes one or more rotor position sensors that generate output indicative of the orientation of the rotor for use in electronic commutation of the rotary motor. In some embodiments, the output of the one or more rotor position sensors is indicative of the position of the rotor within the blood flow conduit transverse to the rotor axis of rotation (e.g., in two different directions transverse to the rotor axis of rotation). In some embodiments, the position of the rotor within the blood flow conduit transverse to the rotor axis of rotation is used to control operation of the stator to control magnetic levitation of the rotor within the blood flow conduit. In some embodiments, the one or more rotor position sensors includes hall effect sensors. In some embodiments, each of the hall effect sensors is disposed in or adjacent to a respective gap between an adjacent pair of the stator coils. In some embodiments, each of the hall effect sensors is disposed aligned with and above or below a respective gap between an adjacent pair of the stator coils.
- (10) In some embodiments, the first implantable blood pump includes control electronics disposed within the housing. In such embodiments, the control electronics can be configured to control current passing through each of the stator coils to radially levitate the rotor and rotate the rotor within the blood flow conduit.
- (11) In many embodiments of the first implantable blood pump, an axial position of the rotor along the blood flow conduit is restrained via passive magnetic interaction between the rotor and the stator such that the stator functions as a passive magnetic bearing that controls the axial position of the rotor parallel to the rotor axis of rotation. In such embodiments, the first implantable blood pump can be configured without dedicated magnetic axial bearings that restrain the axial position of the rotor along the blood flow conduit.
- (12) In many embodiments of the first implantable blood pump, the rotor is separated from the dividing wall so as to accommodate flow of blood around the rotor. For example, in some embodiments of the first implantable blood pump, a gap between the rotor and the dividing wall is between about 0.2 mm to about 2 mm with the rotor centered relative to the stator core. A gap between the rotor and at least one of the stator teeth can be between about 0.3 mm to about 2.4 mm with the rotor centered relative to the stator core.
- (13) In another aspect, a second implantable blood pump includes a housing and a rotary motor. The housing defines an inlet opening, an outlet opening, and a dividing wall defining a blood flow conduit extending from the inlet opening to the outlet opening. The rotary motor includes a stator, hall effect sensors, and a rotor. The stator includes a stator core and stator coils. The stator core includes a toroidal portion. Each of the stator coils extends around one of separated segments of the toroidal portion. The stator is disposed within the housing circumferentially about the dividing wall such that the blood flow conduit extends through the stator core. The stator core is disposed circumferentially around at least a portion of the rotor. Each of the hall effect sensors is disposed in a respective gap between an adjacent pair of the stator coils. The rotor has a rotor axis of rotation and includes a rotor magnet for driving the rotor. The stator core axially overlaps with the rotor magnet with respect to the rotor axis of rotation. In many embodiments, the stator does not extend

beyond a disk-shaped volume having a compact thickness (e.g., less than 1.0 inches) in a direction parallel to the rotor axis of rotation.

(14) In many embodiments, the second implantable blood pump is configured to pump blood from a heart ventricle to the aorta. In some embodiments, the outlet opening is oriented at an angle relative to the input opening. The inlet opening can be oriented to receive blood directly from a heart ventricle and the output opening oriented to output blood in a direction transverse to the orientation of the inlet opening so as to reduce the length of a blood flow cannula used to transfer the blood flow from the output opening to the aorta. The rotor can include centrifugal pump impeller blades.

(15) In many embodiments of the second implantable blood pump, the rotor defines a rotor blood flow conduit that extends through the stator. For example, in many embodiments, the rotor defines a rotor blood flow conduit that extends through the rotor, thereby extending through the stator.

(16) The rotor can have any suitable number of magnetic moments. In some embodiments, the rotor has only one magnetic moment.

(17) In some embodiments, the second implantable blood pump includes control electronics disposed within the housing. In such embodiments, the control electronics can be configured to control current passing through each of the stator coils to radially levitate the rotor and rotate the rotor within the blood flow conduit.

(18) In many embodiments of the second implantable blood pump, an axial position of the rotor along the blood flow conduit is restrained via passive magnetic interaction between the rotor and the stator. In such embodiments, the second implantable blood pump can be configured without dedicated magnetic axial bearings that restrain the axial position of the rotor along the blood flow conduit.

(19) In many embodiments of the second implantable blood pump, the rotor is separated from the dividing wall so as to accommodate flow of blood around the rotor. For example, in some embodiments of the second implantable blood pump, a gap between the rotor and the dividing wall is between about 0.2 mm to about 2 mm with the rotor centered relative to the stator core. A gap between the rotor and at least one of the stator coils can be between about 0.3 mm to about 2.4 mm with the rotor centered relative to the stator core.

(20) In another aspect, a method of assisting blood circulation in a patient is provided. The method includes drawing a flow of blood from a patient's heart into a blood flow channel formed by a housing via rotation of a rotor comprising impeller blades. The flow of blood is passed through a toroidal portion of a motor stator core. Delivery of current to each of a plurality of stator coils is controlled to control a radial position of the rotor within the blood flow channel and to control rotation of the rotor within the blood flow channel. The rotor is rotated around a rotor axis of rotation. Each of the stator coils extends around one of separated segments of the toroidal portion. The rotor has permanent magnetic poles for magnetic levitation and rotation of the rotor. The flow of blood is output from the blood flow channel to the patient.

(21) In many embodiments, the method further includes processing output from a plurality of hall sensors to determine an orientation of the rotor. Each of the hall effect sensors can be disposed in a respective gap between an adjacent pair of the stator coils.

(22) In many embodiments, the method further includes supporting control electronics within the housing and between the stator core and the patient's heart. The control electronics can control the delivery of current to each of the stator coils.

(23) In many embodiments, the method further includes flowing blood through and around the rotor. For example, the method can include (a) passing a first portion of the flow of blood through a central aperture formed through the rotor and (b) passing a second portion of the flow of blood through a gap formed between the rotor and the housing.

(24) In many embodiments, the method further includes magnetically levitating the rotor within the blood flow channel. For example, the rotor can be levitated within the blood flow channel such that

the rotor is separated from the housing by a gap between about 0.2 mm to about 2 mm. The rotor can be levitated within the blood flow channel such that the rotor is separated from at least one of the stator coils by a gap between about 0.3 mm to about 2.4 mm.

(25) For a fuller understanding of the nature and advantages of the present invention, reference should be made to the ensuing detailed description and accompanying drawings.

Description

BRIEF DESCRIPTION OF THE DRAWINGS

(1) FIG. 1 is an illustration of a mechanical circulatory support system implanted in a patient's body, in accordance with many embodiments.

(2) FIG. 2 is a cross-sectional view illustration of an implantable blood pump that includes a compact rotary motor that includes a toroidal stator core with teeth, in accordance with some embodiments.

(3) FIG. 3 is an isometric view illustration of the stator core and a rotor component of the compact rotary motor of FIG. 2.

(4) FIG. 4 is an axial view illustration of the stator core and a rotor component of the compact rotary motor of FIG. 2.

(5) FIG. 5 is a cross-sectional view illustration of the stator core, the rotor component, and rotor position sensors of the compact rotary motor of FIG. 2.

(6) FIG. 6 is a cross-sectional view illustration of an implantable blood pump that includes a compact rotary motor that includes a toroidal stator core without teeth, in accordance with some embodiments.

(7) FIG. 7 is an axial view illustration of the stator core and a rotor component of the compact rotary motor of FIG. 6.

(8) FIG. 8 is an isometric view illustration of the stator core, the rotor component, and hall effect sensors of the compact rotary motor of FIG. 6.

(9) FIG. 9 is a cross-sectional view illustration of the stator core, the rotor component, and hall effect sensors of the compact rotary motor of FIG. 6.

(10) FIG. 10 is a simplified schematic diagram illustration of a method of assisting blood circulation in a patient, in accordance with many embodiments.

(11) FIG. 11 is a simplified schematic diagram illustration of additional acts that can be accomplished in the method of FIG. 10.

DETAILED DESCRIPTION

(12) In the following description, various embodiments of the present invention will be described. For purposes of explanation, specific configurations and details are set forth in order to provide a thorough understanding of the embodiments. However, it will also be apparent to one skilled in the art that the present invention may be practiced without the specific details. Furthermore, well-known features may be omitted or simplified in order not to obscure the embodiment being described.

(13) Referring now to the drawings, in which like reference numerals represent like parts throughout the several views, FIG. 1 shows a mechanical circulatory support system 10 implanted in a patient's body 12. The mechanical circulatory support system 10 includes an implantable blood pump assembly 14, a ventricular cuff 16, an outflow cannula 18, an external system controller 20, and power sources 22. The implantable blood pump assembly 14 can include a VAD that is attached to an apex of the left ventricle, as illustrated, or the right ventricle. A respective VAD can be attached to each of the ventricles of the heart 24. The VAD can include a centrifugal pump (as shown) that is capable of pumping the entire output delivered to the left ventricle from the pulmonary circulation (i.e., up to 10 liters per minute). Related blood pumps applicable to the

present invention are described in greater detail below and in U.S. Pat. Nos. 5,695,471, 6,071,093, 6,116,862, 6,186,665, 6,234,772, 6,264,635, 6,688,861, 7,699,586, 7,976,271, 7,976,271, 7,997,854, 8,007,254, 8,152,493, 8,419,609, 8,852,072, 8,652,024, 8,668,473, 8,864,643, 8,882,744, 9,068,572, 9,091,271, 9,265,870, and 9,382,908, all of which are incorporated herein by reference for all purposes in their entirety. The blood pump assembly **14** can be attached to the heart **24** via the ventricular cuff **16**, which can be sewn to the heart **24** and coupled to the blood pump **14**. The other end of the blood pump **14** connects to the ascending aorta (or the pulmonary artery when the VAD is coupled with the right ventricle of the heart) via the outflow cannula **18** so that the VAD effectively diverts blood from the weakened ventricle and propels it for circulation through the patient's vascular system.

(14) FIG. **1** illustrates the mechanical circulatory support system **10** during battery **22** powered operation. A driveline **26** that exits through the patient's abdomen **28** connects the implanted blood pump assembly **14** to the external system controller **20**, which monitors system **10** operation. Related controller systems applicable to the present invention are described in greater detail below and in U.S. Pat. Nos. 5,888,242, 6,991,595, 8,323,174, 8,449,444, 8,506,471, 8,597,350, and 8,657,733, EP 1812094, and U.S. Patent Publication Nos. 2005/0071001 and 2013/0314047, all of which are incorporated herein by reference for all purposes in their entirety. The system **10** can be powered by either one, two, or more batteries **22**. It will be appreciated that although the system controller **20** and power source **22** are illustrated outside/external to the patient body, the driveline **26**, the system controller **20** and/or the power source **22** can be partially or fully implantable within the patient, as separate components or integrated with the blood pump assembly **14**. Examples of such modifications are further described in U.S. Pat. Nos. 8,562,508 and 9,079,043, all of which are incorporated herein by reference for all purposes in their entirety.

(15) FIG. **2** is a cross-sectional view illustration of an implantable blood pump assembly **30**, in accordance with some embodiments. The blood pump assembly **30** can be used in place of the blood pump assembly **14** in the mechanical circulatory support system **10**. The blood pump assembly **30** includes a housing **32** and a compact rotary motor **34**. The compact rotary motor **34** includes a stator **36** and a rotor assembly **38**. The housing **32** defines an inlet opening **40**, an outlet opening **42**, and a blood flow conduit **44** in fluid communication with the inlet opening **40** and the outlet opening **42**. The housing **32** includes a dividing wall **46** that defines the blood flow conduit **44**. The dividing wall **46** also partially defines a compartment in which the stator **36** is disposed and isolates the stator from blood flowing through the blood flow conduit **44**. The rotor assembly **38** includes a rotor magnetic assembly **48** and an impeller blade assembly **50** attached to the rotor magnetic assembly **48**. The rotor magnetic assembly **48** can include any suitable number of permanent magnets (e.g., 1 or more). In operation, the stator **36** generates magnetic fields that interact with the rotor magnetic assembly **48** to levitate the rotor magnetic assembly **48** radially within the blood flow conduit **44**, rotate the rotor magnetic assembly **48** within the blood flow conduit **44** around a rotor axis of rotation **52**, and react axial thrust applied to the rotor assembly **38** parallel to the rotor axis of rotation **52** during pumping of blood through the blood flow conduit **44** via rotation of the rotor assembly **38**.

(16) The housing **32** has a circular shape and is implanted in a patient's body with a first face **54** of the housing **32** facing the patient's heart **24** and a second face **56** of the housing **32** facing away from the heart **24**. The housing **32** includes an inlet cannula **58** that couples with the ventricular cuff **16** and extends into a ventricle of the heart **24**. The second face **56** of the housing **32** has a chamfered edge **60** to avoid irritating other tissue that may come into contact with the blood pump assembly **30**, such as the patient's diaphragm. To construct the illustrated shape of the puck-shaped housing **32** in a compact form, the stator **36** and electronics **62** of the pump assembly **30** are positioned on the inflow side of the housing **32** toward first face **54**, and the rotor assembly **38** is positioned along the second face **56**. This positioning of the stator **36**, electronics **62**, and the rotor assembly **38** permits the edge **60** to be chamfered along the contour of the impeller blade assembly

50.

(17) The blood flow conduit **44** extends from the inlet opening **40** of the inlet cannula **58** through the stator **36** to the outlet opening **42**. The rotor assembly **38** is positioned within the blood flow conduit **44**. The stator **36** is disposed circumferentially around the rotor magnetic assembly **48**. The stator **36** is also positioned relative to the rotor assembly **38** such that, in use, blood flows within the blood flow conduit **44** through the stator **36** and the rotor magnetic assembly **48** before reaching the impeller blade assembly **50**. In some embodiments, the rotor magnetic assembly **48** has a permanent magnetic north pole (N) and a permanent magnetic south pole (S) for combined active and passive magnetic levitation of the rotor magnetic assembly **48** and for rotation of the rotor assembly **38**. In some embodiments, the rotor magnetic assembly **48** has more than one pair of magnetic poles (e.g., 2, 3, 4, 5, or more). The impeller blade assembly **50** includes impeller blades **64**. The impeller blades **64** are located within a volute **66** of the blood flow conduit **44** such that the impeller blades **64** are located proximate to the second face **56** of the housing **32**.

(18) The puck-shaped housing **32** further includes a peripheral wall **68** that extends between the first face **54** and a removable cap **70**. As illustrated, the peripheral wall **68** is formed as a hollow circular cylinder having a width (W) between opposing portions of the peripheral wall **68**. The housing **32** also has a thickness (T) between the first face **54** and the second face **56** that is less than the width (W). The thickness (T) is from about 0.5 inches to about 1.5 inches, and the width (W) is from about 1 inch to about 4 inches. For example, the width (W) can be approximately 2 inches, and the thickness (T) can be approximately 1 inch.

(19) The peripheral wall **68** encloses an internal compartment **72** that surrounds the dividing wall **46** and the blood flow conduit **44**, with the stator **36** and the electronics **62** disposed in the internal compartment **72** about the dividing wall **46**. The removable cap **70** includes the second face **56**, the chamfered edge **60**, and defines the outlet opening **42**. The cap **70** has an inner surface that defines the volute **66** that is in fluid communication with the outlet opening **42**.

(20) Within the internal compartment **72**, the electronics **62** are positioned adjacent to the first face **54** and the stator **36** is positioned adjacent to the electronics **62** on an opposite side of the electronics **62** from the first face **54**. The electronics **62** can include one or more circuit boards and various components carried on the circuit boards to control the operation of the blood pump assembly **30** (e.g., magnetic levitation and/or drive of the rotor assembly **38**) by controlling currents applied to the stator **36**. The housing **32** is configured to receive the electronics **62** within the internal compartment **72** generally parallel to the first face **54** for efficient use of the space within the internal compartment **72**. The electronics **62** also extend radially-inward towards the dividing wall **46** and radially-outward towards the peripheral wall **68**. For example, the internal compartment **72** is generally sized no larger than necessary to accommodate the stator **36** and the electronics **62**, and space for heat dissipation, material expansion, potting materials, and/or other elements used in installing the stator **36** and the electronics **62**. Thus, the external shape of the housing **32** proximate the first face **54** generally fits the shape of the electronics **62** closely to provide external dimensions that are not much greater than the dimensions of the electronics **62**. In the illustrated embodiment, the electronics **62** include Hall effect sensors **74** that generate output indicative of the angular orientation of the rotor magnetic assembly **48** and the transverse position of the rotor magnetic assembly **48** transverse to the rotor axis of rotation **52** in two directions. The output from the Hall effect sensors **74** is used by the electronics **62** to control operation of the stator **36** to levitate and rotate the rotor assembly **38**.

(21) The rotor assembly **38** is arranged within the housing **32** such that the rotor magnetic assembly **48** is located upstream of the impeller blade assembly **50**. The rotor magnetic assembly **48** is disposed within the blood flow conduit **44** proximate the stator **36**. The rotor magnetic assembly **48** and the dividing wall **44** form a gap **76** between the rotor magnetic assembly **48** and the dividing wall **44** when the rotor magnetic assembly **48** is centered within the blood flow conduit **44**. In many embodiments, the gap **76** is from about 0.2 millimeters to about 2 millimeters. In some

embodiments, the gap **76** is approximately 1 millimeter. The north permanent magnetic pole N and the south permanent magnetic pole S of the rotor magnetic assembly **48** provide a permanent magnetic attractive force between the rotor magnetic assembly **48** and the stator **36** that acts as a passive axial force that tends to maintain the rotor magnetic assembly **48** generally axially aligned with the stator **36** relative to the rotor axis of rotation **52** thereby resisting movement of the rotor magnetic assembly **48** towards the first face **54** or towards the second face **56**.

(22) As blood flows through the blood flow conduit **44**, blood flows through a central aperture **78** formed through the rotor magnetic assembly **48**. Blood also flows through the gap **76** between the rotor magnetic assembly **48** and the dividing wall **46** and through a gap **80** between the impeller blade assembly **50** and the inner surface of the cap **70**. The gaps **76** and **80** are large enough to allow adequate blood flow to limit clot formation that may occur if the blood is allowed to become stagnant. The gaps **76** and **80** are also large enough to limit shear forces on the blood cells such that the blood is not damaged when flowing through the blood pump assembly **30**. As a result of the size of the gaps **76** and **80** limiting shear forces on the blood cells, the gaps **76** and **80** are too large to provide a meaningful hydrodynamic suspension effect. That is to say, the blood does not act as a bearing within the gaps **76** and **80**, and the rotor magnetic assembly **48** is only magnetically-levitated.

(23) Because the rotor assembly **38** is radially suspended by active control of the stator **36**, and because the rotor assembly **38** is axially suspended by passive interaction between the stator **36** and the rotor magnetic assembly **48**, no rotor levitation components other than the stator **36** and related components used to control operation of the stator **36** are needed (e.g., proximate the second face **56**) to levitate the rotor assembly **38** transverse to the rotor axis of rotation **52** and to control the position of the rotor assembly **38** parallel to the rotor axis of rotation **52**. By levitating the rotor assembly **38** via the stator **36**, the cap **70** can be contoured to the shape of the impeller blade assembly **50** and the volute **66**. Additionally, levitating the rotor assembly **38** via the stator **36** eliminates the need for electrical connectors extending from the compartment **72** to the cap **70**, which allows the cap **70** to be easily installed and/or removed and eliminates potential sources of pump failure.

(24) FIG. 3 and FIG. 4 show the stator **36** and the rotor magnetic assembly **48**. The stator **36** includes an integral stator core **82** and stator coils **84**. The integral stator core **82** includes a toroidal portion **86** and stator teeth **88**. Each of the stator teeth **88** extends toward the rotor magnetic assembly **48** from the toroidal portion **86**. Each of the stator teeth **88** is separated from each of an adjacent two of the stator teeth **88** by a respective adjacent intervening segment of the toroidal portion **86**. Each of the stator coils **84** extends around one of the intervening segments of the toroidal portion **86**. The stator **36** is disposed within the housing **32** circumferentially around the dividing wall **46** such that the blood flow conduit **44** extends through the stator core **82**. The stator core **82** is disposed circumferentially around the rotor magnetic assembly **48**. In many embodiments, the stator **36** does not extend beyond a disk-shaped volume having a compact thickness (e.g., (H) shown in FIG. 2 less than 1.0 inches) in a direction parallel to the rotor axis of rotation **52**.

(25) FIG. 5 is a cross-sectional view illustration of the stator **36**, the rotor magnetic assembly **48**, and the electronics **62**. In the illustrated embodiment, the electronics **62** include Hall-effect sensors **74**, each of which is disposed adjacent to a respective one of the stator teeth **88**. By positioning the Hall-effect sensors **74** aligned with the stator teeth **88**, the signals generated by the Hall-effect sensors **74** can be processed to track the orientation of the rotor magnetic assembly **48** relative to the stator teeth **88** without adjusting for an orientation difference between the Hall-effect sensors **74** and the stator teeth **88**.

(26) FIG. 6 is a cross-sectional view illustration of an implantable blood pump assembly **130**, in accordance with some embodiments. The blood pump assembly **130** can be used in place of the blood pump assembly **14** in the mechanical circulatory support system **10**. The blood pump

assembly **130** is configured similar to the blood pump assembly **30** except for differences with respect to the stator core **82** and the location of the Hall-effect sensors **74** as described herein. Accordingly, components of the blood pump assembly **130** that are the same or similar to the components of the blood pump assembly **30** are identified using the same or similar reference identifiers in the drawing figures. As illustrated in FIG. **6** and FIG. **7**, the stator core **82** of the blood pump assembly **130** includes the toroidal portion **86** and does not include the stator teeth **88** of the stator core **82** of the blood pump assembly **30**. As illustrated in FIG. **6**, FIG. **8**, and FIG. **9**, each of the Hall-effect sensors **74** in the blood pump assembly **130** is located in a respective gap between adjacent stator coils **86** that corresponds to a space that is occupied by a respective stator tooth **88** in the blood pump assembly **30**.

(27) FIG. **10** is a simplified schematic diagram illustration of a method **200** of assisting blood circulation in a patient, in accordance with many embodiments. Any suitable blood pump assembly, such as the blood pump assemblies **14**, **30**, **130** described herein, can be used to practice the method **200**.

(28) The method **200** includes drawing a flow of blood from a patient's heart into a blood flow channel formed by a housing via rotation of a rotor comprising impeller blades (act **202**). For example, with reference to FIG. **2**, the rotor assembly **38** can be levitated and rotated via application of drive currents to the stator **36**, thereby drawing blood from the patient's ventricle into the inlet cannula **58** and pumping the blood through the blood flow conduit **44**.

(29) The method **200** includes passing the flow of blood through a toroidal portion of a motor stator core (act **204**). For example, with reference to FIG. **2** and FIG. **4**, the flow of blood passes through the toroidal portion **86** of the motor stator core **82** as the blood flows through the blood flow conduit **44**.

(30) The method **200** includes controlling delivery of current to each of a plurality of stator coils to control a radial position of the rotor within the blood flow channel and to control rotation of the rotor within the blood flow channel, the rotor being rotated around a rotor axis of rotation, each of the stator coils extending around one of separated segments of the toroidal portion, the rotor having permanent magnetic poles for magnetic levitation and rotation of the rotor (act **206**). For example, with reference to FIG. **2** through FIG. **4**, delivery of current to each of the stator coils **84** is controlled (e.g., via the electronics **62**) to control a radial position of the rotor magnetic assembly **48** within the blood flow conduit **44** (i.e., transverse to the rotor axis of rotation **52**) and to control rotation of the rotor magnetic assembly **48** within the blood flow conduit **44**. The rotor magnetic assembly **48** is rotated around the rotor axis of rotation **52**. Each of the stator coils **84** extends around one of separated segments of the toroidal portion **86**. The rotor magnetic assembly **48** has permanent magnetic poles for magnetic levitation and rotation of the rotor magnetic assembly **48**.

(31) The method **200** includes outputting the flow of blood from the blood flow channel to the patient (act **208**). For example, referring to FIG. **2**, the blood flowing through the blood flow conduit **44** is output via the outlet opening **42** and to the ascending aorta via the outflow cannula **18**.

(32) FIG. **11** is a simplified schematic diagram illustration of additional acts that can be accomplished in the method **200**. For example, the method **200** can further include processing output from a plurality of Hall-effect sensors to determine the angular orientation of the rotor and the position of the rotor transverse to the rotor axis of rotation in two directions (act **210**). Each of the Hall-effect sensors can be aligned with a respective gap between an adjacent pair of the stator coils (e.g., above the respective gap, below the respective gap, in the respective gap). For example, referring to FIG. **6**, FIG. **8**, and FIG. **9**, output from the Hall-effect sensors **74** is processed (e.g., via the electronics **62**) to determine the orientation of the rotor magnetic assembly **48** for use in controlling supply of current to each of the stator coils **84** to control levitation and rotation of the rotor magnetic assembly **48**. In the blood pump assembly **130**, each of the Hall-effect sensors **74** is disposed in a respective gap between an adjacent pair of the stator coils **84**.

(33) Method **200** can further include supporting control electronics within the housing and between the stator core and the patient's heart, the control electronics controlling the delivery of current to each of the stator coils (act **212**). For example, referring to FIG. 2, the electronics **62** are supported within the housing **32** and control delivery of current to each of the stator coils **84**.

(34) Method **200** can further include passing a first portion of the flow of blood through a central aperture formed through the rotor and passing a second portion of the flow of blood through a gap formed between the rotor and the housing (act **214**). For example, referring to FIG. 2, a first portion of the blood flowing through the blood flow conduit **44** passes through a central aperture formed through the rotor magnetic assembly **48** and a second portion of the blood flowing through the blood flow conduit **44** recirculates back upstream through the gaps **76, 80** formed between the rotor assembly **38** and the housing **32**.

(35) Other variations are within the spirit of the present invention. Thus, while the invention is susceptible to various modifications and alternative constructions, certain illustrated embodiments thereof are shown in the drawings and have been described above in detail. It should be understood, however, that there is no intention to limit the invention to the specific form or forms disclosed, but on the contrary, the intention is to cover all modifications, alternative constructions, and equivalents falling within the spirit and scope of the invention, as defined in the appended claims.

(36) The use of the terms “a” and “an” and “the” and similar referents in the context of describing the invention (especially in the context of the following claims) are to be construed to cover both the singular and the plural, unless otherwise indicated herein or clearly contradicted by context. The terms “comprising,” “having,” “including,” and “containing” are to be construed as open-ended terms (i.e., meaning “including, but not limited to,”) unless otherwise noted. The term “connected” is to be construed as partly or wholly contained within, attached to, or joined together, even if there is something intervening. Recitation of ranges of values herein are merely intended to serve as a shorthand method of referring individually to each separate value falling within the range, unless otherwise indicated herein, and each separate value is incorporated into the specification as if it were individually recited herein. All methods described herein can be performed in any suitable order unless otherwise indicated herein or otherwise clearly contradicted by context. The use of any and all examples, or exemplary language (e.g., “such as”) provided herein, is intended merely to better illuminate embodiments of the invention and does not pose a limitation on the scope of the invention unless otherwise claimed. No language in the specification should be construed as indicating any non-claimed element as essential to the practice of the invention.

(37) Preferred embodiments of this invention are described herein, including the best mode known to the inventors for carrying out the invention. Variations of those preferred embodiments may become apparent to those of ordinary skill in the art upon reading the foregoing description. The inventors expect skilled artisans to employ such variations as appropriate, and the inventors intend for the invention to be practiced otherwise than as specifically described herein. Accordingly, this invention includes all modifications and equivalents of the subject matter recited in the claims appended hereto as permitted by applicable law. Moreover, any combination of the above-described elements in all possible variations thereof is encompassed by the invention unless otherwise indicated herein or otherwise clearly contradicted by context.

(38) All references, including publications, patent applications, and patents, cited herein are hereby incorporated by reference to the same extent as if each reference were individually and specifically indicated to be incorporated by reference and were set forth in its entirety herein.

Claims

1. An implantable blood pump comprising: a housing defining an inlet opening, an outlet opening, and a dividing wall within the housing defining a blood flow passage that extends between the inlet

opening and the outlet opening; a rotary motor including a stator and a rotor; wherein the stator comprises a stator core and stator coils, wherein the stator core has a toroidally-shaped external surface that extends circumferentially and continuously around an axis of rotation of the rotor, wherein each of the stator coils is wound around and encloses a respective circumferentially extending segment of the toroidally-shaped external surface, wherein each of the stator coils is separated from each of two adjacent instances of the stator coils by an intervening gap that corresponds to a respective exposed circumferentially extending segment of the toroidally-shaped external surface, wherein the stator is disposed within the housing circumferentially about the dividing wall such that the blood flow passage extends through the stator core, wherein the stator core is disposed circumferentially around at least a portion of the rotor, wherein the rotor includes a rotor magnet for driving the rotor, and wherein the stator core overlaps the rotor magnet with respect to the axis of rotation of the rotor; and control electronics disposed within the housing and configured to control current passing through each of the stator coils to radially levitate the rotor and rotate the rotor within the blood flow passage.

2. The implantable blood pump of claim 1, wherein the outlet opening is oriented at an angle relative to the inlet opening.

3. The implantable blood pump of claim 1, wherein the rotor comprises centrifugal pump impeller blades.

4. The implantable blood pump of claim 3, wherein the rotor defines a rotor blood flow passage extending through the rotor.

5. The implantable blood pump of claim 1, wherein the rotor defines a rotor blood flow passage extending through the rotor.

6. The implantable blood pump of claim 1, wherein the rotor has only one magnetic moment.

7. The implantable blood pump of claim 1, wherein an axial position of the rotor along the blood flow passage is restrained via passive magnetic interaction between the rotor and the stator.

8. The implantable blood pump of claim 1, wherein the rotor and the dividing wall are separated by a distance in a range from 0.2 mm to 2 mm with the rotor centered relative to the stator core.

9. The implantable blood pump of claim 1, wherein the rotor and at least one of the stator coils are separated by a distance in a range from 0.3 mm to 2.4 mm with the rotor centered relative to the stator core.

10. The implantable blood pump of claim 1, further comprising hall effect sensors for monitoring an orientation and one or more positions of the rotor relative to the stator.

11. A ventricular assist device comprising: a housing defining an inlet opening, an outlet opening, and a dividing wall within the housing defining a blood flow passage that extends between the inlet opening and the outlet opening; a rotary motor including a stator and a rotor; wherein the stator is operable to rotate the rotor around a rotor axis of rotation, wherein the stator comprises a stator core and stator coils, wherein the stator core has a toroidally-shaped external surface that extends circumferentially and continuously around the rotor axis of rotation, wherein each of the stator coils is wound around and encloses a respective circumferentially extending segment of the toroidally-shaped external surface, wherein each of the stator coils is separated from each of two adjacent instances of the stator coils by an intervening gap that corresponds to a respective exposed circumferentially extending segment of the toroidally-shaped external surface, wherein the stator does not extend beyond a disk-shaped volume having a thickness in a direction parallel to the rotor axis of rotation of less than 1.0 inches, wherein the stator is disposed within the housing circumferentially about the dividing wall such that the blood flow passage extends through the stator core, wherein the stator core is disposed circumferentially around at least a portion of the rotor, wherein the rotor includes a rotor magnet for driving the rotor, and wherein the stator core axially overlaps the rotor magnet with respect to the rotor axis of rotation; and control electronics disposed within the housing and configured to control current supplied to the stator to radially levitate the rotor and rotate the rotor within the blood flow passage.

12. The ventricular assist device of claim 11, wherein: a housing comprising an inlet cannula and a first side face from which the inlet cannula extends; the inlet cannula is configured to couple with a ventricular cuff attached to a heart and extend into a ventricle of the heart; and the housing extends by a maximum distance of 1.5 inches from the first side face in a direction away from the inlet cannula.
 13. The ventricular assist device of claim 11, wherein the outlet opening is oriented at an angle relative to the inlet opening.
 14. The ventricular assist device of claim 11, wherein the rotor comprises centrifugal pump impeller blades.
 15. The ventricular assist device of claim 14, wherein the rotor defines a rotor blood flow passage extending through the rotor.
 16. The ventricular assist device of claim 11, wherein the rotor defines a rotor blood flow passage extending through the rotor.
 17. The ventricular assist device of claim 11, wherein the rotor has only one magnetic moment.
 18. The ventricular assist device of claim 11, wherein an axial position of the rotor along the blood flow passage is restrained via passive magnetic interaction between the rotor and the stator.
 19. The ventricular assist device of claim 11, wherein the rotor and the dividing wall are separated by a distance in a range from 0.2 mm to 2 mm with the rotor centered relative to the stator core.
 20. The ventricular assist device of claim 11, wherein the rotor and the stator are separated by a distance in a range from 0.3 mm to 2.4 mm with the rotor centered relative to the stator core.
 21. The ventricular assist device of claim 11, further comprising hall effect sensors for monitoring an orientation and one or more positions of the rotor relative to the stator.
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