Abstract: An integrated catheter placement system for accurately placing a catheter within a patient’s vasculature is disclosed. In one embodiment, the integrated system comprises a system console, a tip location sensor for temporary placement on the patient’s chest, and an ultrasound probe. The tip location sensor senses a magnetic field of a stylet disposed in a lumen of the catheter when the catheter is disposed in the vasculature. The ultrasound probe ultrasonically images a portion of the vasculature prior to introduction of the catheter. ECG signal-based catheter tip guidance is included in the integrated system to enable guidance of the catheter tip to a desired position with respect to a node of the patient’s heart. Various means for establishing a conductive pathway between a sterile field of the patient and a non-sterile field to enable passage of ECG signals from the catheter to the tip location sensor are also disclosed.

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SYSTEMS AND METHODS FOR BREACHING A STERILE FIELD FOR INTRAVASCULAR PLACEMENT OF A CATHETER

CROSS-REFERENCE TO RELATED APPLICATIONS


BRIEF SUMMARY

[0002] Briefly summarized, embodiments of the present invention are directed to an integrated catheter placement system configured for accurately placing a catheter within the vasculature of a patient. The integrated system employs at least two modalities for improving catheter placement accuracy: 1) ultrasound-assisted guidance for introducing the catheter into the patient's vasculature; and 2) a tip location system ("TLS"), or magnetically-based (e.g., via permanent magnet(s) or electromagnet(s)) tracking of the catheter tip during its advancement through the vasculature to detect and facilitate correction of any tip malposition during such advancement.
In one embodiment, the integrated system comprises a system console including a control processor, a tip location sensor for temporary placement on a portion of a body of the patient, and an ultrasound probe. The tip location sensor senses a magnetic field of a stylet disposed in a lumen of the catheter when the catheter is disposed in the vasculature. The ultrasound probe ultrasonically images a portion of the vasculature prior to introduction of the catheter into the vasculature. In addition, the ultrasound probe includes user input controls for controlling use of the ultrasound probe in an ultrasound mode and use of the tip location sensor in a tip location mode.

In another embodiment, a third modality, i.e., ECG signal-based catheter tip guidance, is included in the system to enable guidance of the catheter tip to a desired position with respect to a node of the patient's heart from which the ECG signals originate. Various means for establishing a conductive pathway between a sterile field of the patient and a non-sterile field to enable passage of ECG signals from the catheter to the tip location sensor are also disclosed. Such means include, for example, connector schemes that establish the conductive pathway through a perforation defined in a sterile barrier, such as a surgical drape, wherein the perforation is isolated by the connector scheme so as to prevent contamination or compromise of the sterile field of the patient.

These and other features of embodiments of the present invention will become more fully apparent from the following description and appended claims, or may be learned by the practice of embodiments of the invention as set forth hereinafter.

**BRIEF DESCRIPTION OF THE DRAWINGS**

A more particular description of the present disclosure will be rendered by reference to specific embodiments thereof that are illustrated in the appended drawings. It is appreciated that these drawings depict only typical embodiments of the invention and are therefore not to be considered limiting of its scope. Example embodiments of the invention will be described and explained with additional specificity and detail through the use of the accompanying drawings in which:

FIG. 1 is a block diagram depicting various elements of an integrated system for intravascular placement of a catheter, according to one example embodiment of the present invention;
FIG. 2 is a simplified view of a patient and a catheter being inserted therein with assistance of the integrated system of FIG. 1;

FIGS. 3A and 3B are views of a probe of the integrated system of FIG. 1;

FIG. 4 is a screenshot of an ultrasound image as depicted on a display of the integrated system of FIG. 1;

FIG. 5 is a perspective view of a stylet employed in connection with the system of FIG. 1 in placing a catheter within a patient vasculature;

FIG. 6 is an icon as depicted on a display of the integrated system of FIG. 1, indicating a position of a distal end of the stylet of FIG. 5 during catheter tip placement procedures;

FIGS. 7A-7E depict various example icons that can be depicted on the display of the integrated system of FIG. 1 during catheter tip placement procedures;

FIGS. 8A-8C are screenshots of images depicted on a display of the integrated system of FIG. 1 during catheter tip placement procedures;

FIG. 9 is a block diagram depicting various elements of an integrated system for intravascular placement of a catheter, according to another example embodiment of the present invention;

FIG. 10 is a simplified view of a patient and a catheter being inserted therein with assistance of the integrated system of FIG. 9;

FIG. 11 is a perspective view of a stylet employed in connection with the integrated system of FIG. 9 in placing a catheter within a patient vasculature;

FIGS. 12A-12E are various views of portions of the stylet of FIG. 11;

FIGS. 13A-13D are various views of a fin connector assembly for use with the integrated system of FIG. 9;

FIGS. 13E-13F are various views of a tether connector for use with the fin connector assembly shown in FIGS. 13A-13D;
FIGS. 14A-14C are views showing the connection of a stylet tether and fin connector to a sensor of the integrated system of FIG. 9;

FIG. 15 is a cross sectional view of the connection of the stylet tether, fin connector, and sensor shown in FIG. 14C;

FIG. 16 is simplified view of an ECG trace of a patient;

FIG. 17 is a screenshot of an image depicted on a display of the integrated system of FIG. 9 during catheter tip placement procedures;

FIG. 18 is a cross sectional view of a fin connector including electrical contacts configured in accordance with one embodiment;

FIGS. 19A and 19B are simplified views of an electrical contact retention system for engagement of a tether connector with a fin connector, in accordance with one embodiment;

FIGS. 20A-20C are various views of one embodiment of a fin connector and a tether connector for establishing a signal pathway through a sterile barrier in connection with use of the integrated system described herein;

FIGS. 21A and 21B are various views of a connector for electrically connecting ECG electrodes to a sensor of the integrated system, according to one embodiment;

FIGS. 22A-22C are various views of one embodiment of a fin connector and a tether connector for establishing a signal pathway through a sterile barrier;

FIGS. 23A and 23B are cross sectional views of a connector system for establishing a signal pathway through a sterile barrier, according to one embodiment;

FIG. 24 is a simplified side view of a connector system for establishing a signal pathway through a sterile barrier, according to one embodiment;

FIGS. 25A and 25B are simplified side views of a connector system for establishing a signal pathway through a sterile barrier, according to one embodiment;
FIGS. 26A and 26B are cross-sectional views of a connector system for establishing a signal pathway through a sterile barrier, according to one embodiment;

FIG. 27 is a simplified view of a connector system for establishing a signal pathway through a sterile barrier, according to one embodiment;

FIG. 28 is a perspective view of stylet including a sterile shield for use with the connector system shown in FIG. 28, according to one embodiment;

FIGS. 29A and 29B are simplified views of the ECG module of FIG. 27, including a connector system for establishing a signal pathway through a sterile barrier, according to one embodiment;

FIG. 30 is a simplified view of a connector system for establishing a signal pathway through a sterile barrier, according to one embodiment;

FIG. 31 is a simplified view of a connector system for establishing a signal pathway through a sterile barrier, according to one embodiment; and

FIG. 32 is a simplified view of elements of a connector system for establishing a signal pathway through a sterile barrier, according to one embodiment.

DETAILED DESCRIPTION OF SELECTED EMBODIMENTS

Reference will now be made to figures wherein like structures will be provided with like reference designations. It is understood that the drawings are diagrammatic and schematic representations of exemplary embodiments of the present invention, and are neither limiting nor necessarily drawn to scale.

FIGS. 1-32 depict various features of embodiments of the present invention, which is generally directed to a catheter placement system configured for accurately placing a catheter within the vasculature of a patient. In one embodiment, the catheter placement system employs at least two modalities for improving catheter placement accuracy: 1) ultrasound-assisted guidance for introducing the catheter into the patient's vasculature; and 2) a tip location/navigation system ("TLS"), or magnetically-based tracking of the catheter tip during its advancement through the tortuous vasculature path to detect and facilitate correction of any tip malposition during such advancement. The ultrasound guidance and tip location features of the present system according to one
embodiment are integrated into a single device for use by a clinician placing the catheter. Integration of these two modalities into a single device simplifies the catheter placement process and results in relatively faster catheter placements. For instance, the integrated catheter placement system enables ultrasound and TLS activities to be viewed from a single display of the integrated system. Also, controls located on an ultrasound probe of the integrated device, which probe is maintained within the sterile field of the patient during catheter placement, can be used to control functionality of the system, thus precluding the need for a clinician to reach out of the sterile field in order to control the system.

[00042] In another embodiment, a third modality, i.e., ECG signal-based catheter tip guidance, is included in the integrated system to enable guidance of the catheter tip to a desired position with respect to a node of the patient's heart from which the ECG signals originate. Such ECG-based positional assistance is also referred to herein as "tip confirmation."

[00043] Combination of the three modalities above according to one embodiment enables the catheter placement system to facilitate catheter placement within the patient's vasculature with a relatively high level of accuracy, i.e., placement of the distal tip of the catheter in a predetermined and desired position. Moreover, because of the ECG-based guidance of the catheter tip, correct tip placement may be confirmed without the need for a confirmatory X-ray. This, in turn, reduces the patient's exposure to potentially harmful x-rays, the cost and time involved in transporting the patient to and from the x-ray department, costly and inconvenient catheter repositioning procedures, etc.

[00044] As the ECG signal-based modality includes a need for passing ECG signals from a catheter assembly disposed in a sterile field of a patient to a data-receiving component of the system disposed in a non-sterile field, embodiments of the present invention are further concerned with various connector systems for establishing a conductive pathway through a sterile barrier separating the sterile and non-sterile fields.

[00045] For clarity it is to be understood that the word "proximal" as used herein refers to a direction relatively closer to a clinician, while the word "distal" refers to a direction relatively further from the clinician. For example, the end of a catheter placed within the body of a patient is considered a distal end of the catheter, while the catheter end remaining outside the body is a proximal end of the catheter. Also, the words
"including," "has," and "having," as used herein, including the claims, shall have the same meaning as the word "comprising."

[00046] Reference is first made to FIGS. 1 and 2 which depict various components of a catheter placement system ("system"), generally designated at 10, configured in accordance with one example embodiment of the present invention. As shown, the system 10 generally includes a console 20, display 30, probe 40, and sensor 50, each of which is described in further detail below.

[00047] FIG. 2 shows the general relation of these components to a patient 70 during a procedure to place a catheter 72 into the patient vasculature through a skin insertion site 73. FIG. 2 shows that the catheter 72 generally includes a proximal portion 74 that remains exterior to the patient and a distal portion 76 that resides within the patient vasculature after placement is complete. The system 10 is employed to ultimately position a distal tip 76A of the catheter 72 in a desired position within the patient vasculature. In one embodiment, the desired position for the catheter distal tip 76A is proximate the patient’s heart, such as in the lower one-third (1/3rd) portion of the Superior Vena Cava ("SVC"). Of course, the system 10 can be employed to place the catheter distal tip in other locations. The catheter proximal portion 74 further includes a hub 74A that provides fluid communication between the one or more lumens of the catheter 72 and one or more extension legs 74B extending proximally from the hub.

[00048] An example implementation of the console 20 is shown in FIG. 8C, though it is appreciated that the console can take one of a variety of forms. A processor 22, including non-volatile memory such as EEPROM for instance, is included in the console 20 for controlling system function during operation of the system 10, thus acting as a control processor. A digital controller/analog interface 24 is also included with the console 20 and is in communication with both the processor 22 and other system components to govern interfacing between the probe 40, sensor 50, and other system components.

[00049] The system 10 further includes ports 52 for connection with the sensor 50 and optional components 54 including a printer, storage media, keyboard, etc. The ports in one embodiment are USB ports, though other port types or a combination of port types can be used for this and the other interfaces connections described herein. A power connection 56 is included with the console 20 to enable operable connection to an
external power supply 58. An internal battery 60 can also be employed, either with or exclusive of an external power supply. Power management circuitry 59 is included with the digital controller/analog interface 24 of the console to regulate power use and distribution.

[00050] The display 30 in the present embodiment is integrated into the console 20 and is used to display information to the clinician during the catheter placement procedure. In another embodiment, the display may be separate from the console. As will be seen, the content depicted by the display 30 changes according to which mode the catheter placement system is in: US, TLS, or in other embodiments, ECG tip confirmation. In one embodiment, a console button interface 32 (see FIGS. 1, 8C) and buttons included on the probe 40 can be used to immediately call up a desired mode to the display 30 by the clinician to assist in the placement procedure. In one embodiment, information from multiple modes, such as TLS and ECG, may be displayed simultaneously, such as in FIG. 17. Thus, the single display 30 of the system console 20 can be employed for ultrasound guidance in accessing a patient's vasculature, TLS guidance during catheter advancement through the vasculature, and (as in later embodiments) ECG-based confirmation of catheter distal tip placement with respect to a node of the patient's heart. In one embodiment, the display 30 is an LCD device.

[00051] FIGS. 3A and 3B depict features of the probe 40 according to one embodiment. The probe 40 is employed in connection with the first modality mentioned above, i.e., ultrasound ("US")-based visualization of a vessel, such as a vein, in preparation for insertion of the catheter 72 into the vasculature. Such visualization gives real time ultrasound guidance for introducing the catheter into the vasculature of the patient and assists in reducing complications typically associated with such introduction, including inadvertent arterial puncture, hematoma, pneumothorax, etc.

[00052] The handheld probe 40 includes a head 80 that houses a piezoelectric array for producing ultrasonic pulses and for receiving echoes thereof after reflection by the patient's body when the head is placed against the patient's skin proximate the prospective insertion site 73 (FIG. x). The probe 40 further includes a plurality of control buttons 84, which can be included on a button pad 82. In the present embodiment, the modality of the system 10 can be controlled by the control buttons 84, thus eliminating the need for the clinician to reach out of the sterile field, which is
established about the patient insertion site prior to catheter placement, to change modes
via use of the console button interface 32.

[00053] As such, in one embodiment a clinician employs the first (US) modality to
determine a suitable insertion site and establish vascular access, such as with a needle or
introducer, then with the catheter. The clinician can then seamlessly switch, via button
pushes on the probe button pad 82, to the second (TLS) modality without having to reach
out of the sterile field. The TLS mode can then be used to assist in advancement of the
catheter 72 through the vasculature toward an intended destination.

[00054] FIG. 1 shows that the probe 40 further includes button and memory controller
42 for governing button and probe operation. The button and memory controller 42 can
include non-volatile memory, such as EEPROM, in one embodiment. The button and
memory controller 42 is in operable communication with a probe interface 44 of the
console 20, which includes a piezo input/output component 44A for interfacing with the
probe piezoelectric array and a button and memory input/output component 44B for
interfacing with the button and memory controller 42.

[00055] FIG. 4 shows an example screenshot 88 as depicted on the display 30 while
the system 10 is in its first ultrasound modality. An image 90 of a subcutaneous region
of the patient 70 is shown, depicting a cross section of a vein 92. The image 90 is
produced by operation of the piezoelectric array of the probe 40. also included on the
display screenshot 88 is a depth scale indicator 94, providing information regarding the
depth of the image 90 below the patient's skin, a lumen size scale 96 that provides
information as to the size of the vein 92 relative to standard catheter lumen sizes, and
other indicia 98 that provide information regarding status of the system 10 or possible
actions to be taken, e.g., freeze frame, image templates, data save, image print, power
status, image brightness, etc.

[00056] Note that while a vein is depicted in the image 90, other body lumens or
portions can be imaged in other embodiments. Note that the US mode shown in FIG. 4
can be simultaneously depicted on the display 30 with other modes, such as the TLS
mode, if desired. In addition to the visual display 30, aural information, such as beeps,
tones, etc., can also be employed by the system 10 to assist the clinician during catheter
placement. Moreover, the buttons included on the probe 40 and the console button
interface 32 can be configured in a variety of ways, including the use of user input
controls in addition to buttons, such as slide switches, toggle switches, electronic or touch-sensitive pads, etc. Additionally, both US and TLS activities can occur simultaneously or exclusively during use of the system 10.

[00057] As just described, the handheld ultrasound probe 40 is employed as part of the integrated catheter placement system 10 to enable US visualization of the peripheral vasculature of a patient in preparation for transcutaneous introduction of the catheter. In the present example embodiment, however, the probe is also employed to control functionality of the TLS portion, or second modality, of the system 10 when navigating the catheter toward its desired destination within the vasculature as described below. Again, as the probe 40 is used within the sterile field of the patient, this feature enables TLS functionality to be controlled entirely from within the sterile field. Thus the probe 40 is a dual-purpose device, enabling convenient control of both US and TLS functionality of the system 10 from the sterile field. In one embodiment, the probe can also be employed to control some or all ECG-related functionality, or third modality, of the catheter placement system 10, as described further below.

[00058] The catheter placement system 10 further includes the second modality mentioned above, i.e., the magnetically-based catheter TLS, or tip location system. The TLS enables the clinician to quickly locate and confirm the position and/or orientation of the catheter 72, such as a peripherally-inserted central catheter ("PICC"), central venous catheter ("CVC"), or other suitable catheter, during initial placement into and advancement through the vasculature of the patient 70. Specifically, the TLS modality detects a magnetic field generated by a magnetic element-equipped tip location styler, which is pre-loaded in one embodiment into a longitudinally defined lumen of the catheter 72, thus enabling the clinician to ascertain the general location and orientation of the catheter tip within the patient body. In one embodiment, the magnetic assembly can be tracked using the teachings of one or more of the following U.S. patents: 5,775,322; 5,879,297; 6,129,668; 6,216,028; and 6,263,230. The contents of the afore-mentioned U.S. patents are incorporated herein by reference in their entireties. The TLS also displays the direction in which the catheter tip is pointing, thus further assisting accurate catheter placement. The TLS further assists the clinician in determining when a malposition of the catheter tip has occurred, such as in the case where the tip has deviated from a desired venous path into another vein.
As mentioned, the TLS utilizes a stylet to enable the distal end of the catheter to be tracked during its advancement through the vasculature. FIG. 5 gives an example of such a stylet 100, which includes a proximal end IOOA and a distal end IOOB. A handle is included at the stylet proximal end IOOA, with a core wire 104 extending distally therefrom. A magnetic assembly is disposed distally of the core wire 104. The magnetic assembly includes one or more magnetic elements 106 disposed adjacent one another proximate the stylet distal end IOOB and encapsulated by tubing 108. In the present embodiment, a plurality of magnetic elements 106 is included, each element including a solid, cylindrically shaped ferromagnetic stacked end-to-end with the other magnetic elements. An adhesive tip 110 can fill the distal tip of the tubing 108, distally to the magnetic elements 106.

Note that in other embodiments, the magnetic elements may vary from the design in not only shape, but also composition, number, size, magnetic type, and position in the stylet distal segment. For example, in one embodiment, the plurality of ferromagnetic magnetic elements is replaced with an electromagnetic assembly, such as an electromagnetic coil, which produces a magnetic field for detection by the sensor. Another example of an assembly usable here can be found in U.S. Patent No. 5,099,845 entitled "Medical Instrument Location Means," which is incorporated herein by reference in its entirety. Yet other examples of stylets including magnetic elements that can be employed with the TLS modality can be found in U.S. Application No. 11/466,602, filed August 23, 2006, and entitled "Stylet Apparatuses and Methods of Manufacture," which is incorporated herein by reference in its entirety. These and other variations are therefore contemplated by embodiments of the present invention. It should be appreciated herein that "stylet" as used herein can include any one of a variety of devices configured for removable placement within a lumen of the catheter to assist in placing a distal end of the catheter in a desired location within the patient's vasculature.

FIG. 2 shows disposal of the stylet 100 substantially within a lumen in the catheter 72 such that the proximal portion thereof extends proximally from the catheter lumen, through the hub 74A and out through a selected one of the extension legs 74B. So disposed within a lumen of the catheter, the distal end IOOB of the stylet 100 is substantially co-terminal with the distal catheter end 76A such that detection by the TLS of the stylet distal end correspondingly indicates the location of the catheter distal end.
The TLS sensor 50 is employed by the system 10 during TLS operation to detect a magnetic field produced by the magnetic elements 106 of the stylet 100. As seen in FIG. 2, the TLS sensor 50 is placed on the chest of the patient during catheter insertion. The TLS sensor 50 is placed on the chest of the patient in a predetermined location, such as through the use of external body landmarks, to enable the magnetic field of the stylet magnetic elements 106, disposed in the catheter 72 as described above, to be detected during catheter transit through the patient vasculature. Again, as the magnetic elements 106 of the stylet magnetic assembly are co-terminal with the distal end 76A of the catheter 72 (FIG. 2), detection by the TLS sensor 50 of the magnetic field of the magnetic elements provides information to the clinician as to the position and orientation of the catheter distal end during its transit.

In greater detail, the TLS sensor 50 is operably connected to the console 20 of the system 10 via one or more of the ports 52, as shown in FIG. 1. Note that other connection schemes between the TLS sensor and the system console can also be used without limitation. As just described, the magnetic elements 106 are employed in the stylet 100 to enable the position of the catheter distal end 76A (FIG. 2) to be observable relative to the TLS sensor 50 placed on the patient's chest. Detection by the TLS sensor 50 of the stylet magnetic elements 106 is graphically displayed on the display 30 of the console 20 during TLS mode. In this way, a clinician placing the catheter is able to generally determine the location of the catheter distal end 76A within the patient vasculature relative to the TLS sensor 50 and detect when catheter malposition, such as advancement of the catheter along an undesired vein, is occurring.

FIGS. 6 and 7A-7E show examples of icons that can be used by the console display 30 to depict detection of the stylet magnetic elements 106 by the TLS sensor 50. In particular, FIG. 6 shows an icon 114 that depicts the distal portion of the stylet 100, including the magnetic elements 106 as detected by the TLS sensor 50 when the magnetic elements are positioned under the TLS sensor. As the stylet distal end 100B is substantially co-terminal with the distal end 76A of the catheter 72, the icon indicates the position and orientation of the catheter distal end. FIGS. 7A-7E show various icons that can be depicted on the on the console display 30 when the magnetic elements 106 of the stylet 100 are not positioned directly under a portion of the TLS sensor 50, but are nonetheless detected nearby. The icons can include half-icons 114A and quarter-icons...
that are displayed according to the position of the stylet magnetic assembly, i.e.,
the magnetic elements 106 in the present embodiment, relative to the TLS sensor 50.

FIGS. 8A-8C depict screenshots taken from the display 30 of the system 10
while in TLS mode, showing how the magnetic assembly of the stylet 100 is depicted.
The screenshot 118 of FIG. 8A shows a representative image 120 of the TLS sensor 50.
Other information is provided on the display screenshot 118, including a depth scale
indicator 124, status/action indicia 126, and icons 128 corresponding to the button
interface 32 included on the console 20 (FIG. 8C). Though the icons 128 in the present
embodiment are simply indicators to guide the user in identifying the purpose of the
Corresponding buttons of the button interface 32, in another embodiment the display can
be made touch-sensitive so that the icons themselves can function as button interfaces
and can change according to the mode the system is in.

During initial stages of catheter advancement through the patient's
vasculature after insertion therein, the distal end 76A of the catheter 72, having the stylet
distal end 100B substantially co-terminal therewith, is relatively distant from the TLS
sensor 50. As such, the display screenshot will indicate "no signal," indicating that the
magnetic field from the stylet magnetic assembly has not been detected. In FIG. 8B, the
magnetic assembly proximate the stylet distal end 10OB has advanced sufficiently close
to the TLS sensor 50 to be detected thereby, though it is not yet under the sensor. This is
indicated by the half-icon 114A shown to the left of the sensor image 120, representing
the stylet magnetic assembly being positioned to the right of the TLS sensor 50 from the
Perspective of the patient.

In FIG. 8C, the magnetic assembly proximate the stylet distal end 100B has
advanced under the TLS sensor 50 such that its position and orientation relative thereto
is detected by the TLS sensor. This is indicated by the icon 114 on the sensor image
120. Note that the button icons 128 provide indications of the actions that can be
performed by pressing the corresponding buttons of the console button interface 32. As
such, the button icons 128 can change according to which modality the system 10 is in,
thus providing flexibility of use for the button interface 32. Note further that, as the
button pad 82 of the probe 40 (FIG. 3A, 3B) includes buttons 84 that mimic several of
the buttons of the button interface 32, the button icons 128 on the display 30 provide a
guide to the clinician for controlling the system 10 with the probe buttons 84 while
remaining in the sterile field. For instance, if the clinician has need to leave TLS mode and return to US (ultrasound) mode, the appropriate control button 84 on the probe button pad 82 can be depressed, and the US mode can be immediately called up, with the display 30 refreshing to accommodate the visual information needed for US functionality, such as that shown in FIG. 4. This is accomplished without a need for the clinician to reach out of the sterile field.

[00068] Reference is now made to FIGS. 9 and 10 in describing the integrated catheter placement system 10 according to another example embodiment. As before, the integrated system 10 includes the console 20, display 30, probe 40 for US functionality, and the TLS sensor 50 for tip location functionality as described above. Note that the system 10 depicted in FIGS. 9 and 10 is similar in many respects to the system shown in FIGS. 1 and 2. As such, only selected differences will be discussed below. The system 10 of FIGS. 9 and 10 includes additional functionality wherein determination of the proximity of the catheter distal tip 76A relative to a sino-atrial ("SA") or other electrical impulse-emitting node of the heart of the patient 70 can be determined, thus providing enhanced ability to accurately place the catheter distal tip in a desired location proximate the node. Also referred to herein as "ECG" or "ECG-based tip confirmation," this third modality of the system 10 enables detection of ECG signals from the SA node in order to place the catheter distal tip in a desired location within the patient vasculature. Note that the US, TLS, and ECG modalities are seamlessly combined in the present system 10, but can be employed in concert or individually to assist in catheter placement. In one embodiment, it is understood that the ECG modality as described herein can be included in a stand-alone system without the inclusion of the US and TLS modalities. Thus, the environments in which the embodiments herein are described are understood as merely example environments and are not considered limiting of the present disclosure.

[00069] FIGS. 9 and 10 show the addition to the system 10 of a stylet 130 configured in accordance with the present embodiment. As an overview, the catheter stylet 130 is removably predisposed within the lumen of the catheter 72 being inserted into the patient 70 via the insertion site 73. The stylet 130, in addition to including a magnetic assembly for the magnetically-based TLS modality, includes a sensing component, *i.e.*, an ECG sensor assembly, proximate its distal end and including a portion that is co-terminal with the distal end of the catheter tip for sensing ECG signals produced by the SA node. In contrast to the previous embodiment, the stylet 130 includes a tether 134 extending from
its proximal end that operably connects to the TLS sensor 50. As will be described in further detail, the stylet tether 134 permits ECG signals detected by the ECG sensor assembly included on a distal portion of the stylet 130 to be conveyed to the TLS sensor 50 during confirmation of the catheter tip location as part of the ECG signal-based tip confirmation modality. Reference and ground ECG lead/electrode pairs 158 attach to the body of the body of the patient 70 and are operably attached to the TLS sensor 50 to enable the system to filter out high level electrical activity unrelated to the electrical activity of the SA node of the heart, thus enabling the ECG-based tip confirmation functionality. Together with the reference and ground signals received from the ECG lead/electrode pairs 158 placed on the patient's skin, the ECG signals sensed by the stylet ECG sensor assembly are received by the TLS sensor 50 positioned on the patient's chest (FIG. 10) or other designated component of the system 10. The TLS sensor 50 and/or console processor 22 can process the ECG signal data to produce an electrocardiogram waveform on the display 30, as will be described. In the case where the TLS sensor 50 processes the ECG signal data, a processor is included therein to perform the intended functionality. If the console 20 processes the ECG signal data, the processor 22, controller 24, or other processor can be utilized in the console to process the data.

[00070] Thus, as it is advanced through the patient vasculature, the catheter 72 equipped with the stylet 130 as described above can advance under the TLS sensor 50, which is positioned on the chest of the patient as shown in FIG. 10. This enables the TLS sensor 50 to detect the position of the magnetic assembly of the stylet 130, which is substantially co-terminal with the distal tip 76A of the catheter as located within the patient's vasculature. The detection by the TLS sensor 50 of the stylet magnetic assembly is depicted on the display 30 during ECG mode. The display 30 further depicts during ECG mode an ECG electrocardiogram waveform produced as a result of patient heart's electrical activity as detected by the ECG sensor assembly of the stylet 130. In greater detail, the ECG electrical activity of the SA node, including the P-wave of the waveform, is detected by the ECG sensor assembly of the stylet (described below) and forwarded to the TLS sensor 50 and console 20. The ECG electrical activity is then processed for depiction on the display 30. A clinician placing the catheter can then observe the ECG data to determine optimum placement of the distal tip 76A of the catheter 72, such as proximate the SA node in one embodiment. In one embodiment, the console 20 includes the electronic components, such as the processor 22 (FIG. 9),
necessary to receive and process the signals detected by the stylet ECG sensor assembly. In another embodiment, the TLS sensor 50 can include the necessary electronic components processing the ECG signals.

[00071] As already discussed, the display 30 is used to display information to the clinician during the catheter placement procedure. The content of the display 30 changes according to which mode the catheter placement system is in: US, TLS, or ECG. Any of the three modes can be immediately called up to the display 30 by the clinician, and in some cases information from multiple modes, such as TLS and ECG, may be displayed simultaneously. In one embodiment, as before, the mode the system is in may be controlled by the control buttons 84 included on the handheld probe 40, thus eliminating the need for the clinician to reach out of the sterile field (such as touching the button interface 32 of the console 20) to change modes. Thus, in the present embodiment the probe 40 is employed to also control some or all ECG-related functionality of the system 10. Note that the button interface 32 or other input configurations can also be used to control system functionality. Also, in addition to the visual display 30, aural information, such as beeps, tones, etc., can also be employed by the system to assist the clinician during catheter placement.

[00072] Reference is now made to FIGS. 11-12E in describing various details of one embodiment of the stylet 130 that is removably loaded into the catheter 72 and employed during insertion to position the distal tip 76A of the catheter in a desired location within the patient vasculature. As shown, the stylet 130 as removed from the catheter defines a proximal end 130A and a distal end 130B. A connector 132 is included at the proximal stylet end 130A, and a tether 134 extends distally from the connector and attaches to a handle 136. A core wire 138 extends distally from the handle 136. The stylet 130 is pre-loaded within a lumen of the catheter 72 in one embodiment such that the distal end 130B is substantially flush, or co-terminal, with the catheter opening at the distal end 76A thereof (FIG. 10), and such that a proximal portion of the core wire 138, the handle 136, and the tether 134 extend proximally from a selected one of the extension tubes 74B. Note that, though described herein as a stylet, in other embodiments a guidewire or other catheter guiding apparatus could include the principles of the embodiment described herein.
The core wire 138 defines an elongate shape and is composed of a suitable stylet material including stainless steel or a memory material such as, in one embodiment, a nickel and titanium-containing alloy commonly known by the acronym "nitinol." Though not shown here, manufacture of the core wire 138 from nitinol in one embodiment enables the portion of the core wire corresponding to a distal segment of the stylet to have a pre-shaped bent configuration so as to urge the distal portion of the catheter 72 into a similar bent configuration. In other embodiments, the core wire includes no pre-shaping. Further, the nitinol construction lends torqueability to the core wire 138 to enable a distal segment of the stylet 130 to be manipulated while disposed within the lumen of the catheter 72, which in turn enables the distal portion of the catheter to be navigated through the vasculature during catheter insertion.

The handle 136 is provided to enable insertion/removal of the stylet from the catheter 72. In embodiments where the stylet core wire 138 is torqueable, the handle 136 further enables the core wire to be rotated within the lumen of the catheter 72, to assist in navigating the catheter distal portion through the vasculature of the patient 70.

The handle 136 attaches to a distal end of the tether 134. In the present embodiment, the tether 134 is a flexible, shielded cable housing one or more conductive wires electrically connected both to the core wire 138, which acts as the ECG sensor assembly referred to above, and the tether connector 132. As such, the tether 134 provides a conductive pathway from the distal portion of the core wire 138 through to the tether connector 132 at proximal end 130A of the stylet 130. As will be explained, the tether connector 132 is configured for operable connection to the TLS sensor 50 on the patient's chest for assisting in navigation of the catheter distal tip 76A to a desired location within the patient vasculature.

As seen in FIGS. 12B-12D, a distal portion of the core wire 138 is gradually tapered, or reduced in diameter, distally from a junction point 142. A sleeve 140 is slid over the reduced-diameter core wire portion. Though of relatively greater diameter here, the sleeve in another embodiment can be sized to substantially match the diameter of the proximal portion of the stylet core wire. The stylet 130 further includes a magnetic assembly disposed proximate the distal end 130B thereof for use during TLS mode. The magnetic assembly in the illustrated embodiment includes a plurality of magnetic elements 144 interposed between an outer surface of the reduced-diameter core wire 138
and an inner surface of the sleeve 140 proximate the stylet distal end 130B. In the present embodiment, the magnetic elements 144 include 20 ferromagnetic magnets of a solid cylindrical shape stacked end-to-end in a manner similar to the stylet 100 of FIG. 2. In other embodiments, however, the magnetic element(s) may vary from this design in not only shape, but also composition, number, size, magnetic type, and position in the stylet. For example, in one embodiment the plurality of magnets of the magnetic assembly is replaced with an electromagnetic coil that produces a magnetic field for detection by the TLS sensor. These and other variations are therefore contemplated by embodiments of the present invention.

[00077] The magnetic elements 144 are employed in the stylet 130 distal portion to enable the position of the stylet distal end 130B to be observable relative to the TLS sensor 50 placed on the patient's chest. As has been mentioned, the TLS sensor 50 is configured to detect the magnetic field of the magnetic elements 144 as the stylet advances with the catheter 72 through the patient vasculature. In this way, a clinician placing the catheter 72 is able to generally determine the location of the catheter distal end 76A within the patient vasculature and detect when catheter malposition is occurring, such as advancement of the catheter along an undesired vein, for instance.

[00078] The stylet 130 further includes the afore-mentioned ECG sensor assembly, according to one embodiment. The ECG sensor assembly enables the stylet 130, disposed in a lumen of the catheter 72 during insertion, to be employed in detecting an intra-atrial ECG signal produced by an SA or other node of the patient's heart, thereby allowing for navigation of the distal tip 76A of the catheter 72 to a predetermined location within the vasculature proximate the patient's heart. Thus, the ECG sensor assembly serves as an aide in confirming proper placement of the catheter distal tip 76A.

[00079] In the embodiment illustrated in FIGS. 11-12E, the ECG sensor assembly includes a distal portion of the core wire 138 disposed proximate the stylet distal end 130B. The core wire 138, being electrically conductive, enables ECG signals to be detected by the distal end thereof and transmitted proximally along the core wire. A conductive material 146, such as a conductive epoxy, fills a distal portion of the sleeve 140 adjacent the distal termination of the core wire 138 so as to be in conductive communication with the distal end of the core wire. This in turn increases the
conductive surface of the distal end 130B of the stylet 130 so as to improve its ability to
detect ECG signals.

[00080] Before catheter placement, the stylet 130 is loaded into a lumen of the
catheter 72. Note that the stylet 130 can come preloaded in the catheter lumen from the
manufacturer, or loaded into the catheter by the clinician prior to catheter insertion. The
stylet 130 is disposed within the catheter lumen such that the distal end 130B of the
stylet 130 is substantially co-terminal with the distal tip 76A of the catheter 72, thus
placing the distal tips of both the stylet and the catheter in substantial alignment with one
another. The co-terminality of the catheter 72 and stylet 130 enables the magnetic
assembly to function with the TLS sensor 50 in TLS mode to track the position of the
catheter distal tip 76A as it advances within the patient vasculature, as has been
described. Note, however, that for the tip confirmation functionality of the system 10,
the distal end 130B of the stylet 130 need not be co-terminal with the catheter distal end
76A. Rather, all that is required is that a conductive path between the vasculature and
the ECG sensor assembly, in this case the core wire 138, be established such that
electrical impulses of the SA node or other node of the patient's heart can be detected.
This conductive path in one embodiment can include various components including
saline solution, blood, etc.

[00081] In one embodiment, once the catheter 72 has been introduced into the patient
vasculature via the insertion site 73 (FIG. 10) the TLS mode of the system 10 can be
employed as already described to advance the catheter distal tip 76A toward its intended
destination proximate the SA node. Upon approaching the region of the heart, the
system 10 can be switched to ECG mode to enable ECG signals emitted by the SA node
to be detected. As the stylet-loaded catheter is advanced toward the patient's heart, the
electrically conductive ECG sensor assembly, including the distal end of the core wire
138 and the conductive material 146, begins to detect the electrical impulses produced by
the SA node. As such, the ECG sensor assembly serves as an electrode for detecting
the ECG signals. The elongate core wire 138 proximal to the core wire distal end serves
as a conductive pathway to convey the electrical impulses produced by the SA node and
received by the ECG sensor assembly to the tether 134.

[00082] The tether 134 conveys the ECG signals to the TLS sensor 50 temporarily
placed on the patient's chest. The tether 134 is operably connected to the TLS sensor 50
via the tether connector 132 or other suitable direct or indirect connective configuration. As described, the ECG signal can then be processed and depicted on the system display 30 (FIG. 9, 10). Monitoring of the ECG signal received by the TLS sensor 50 and displayed by the display 30 enables a clinician to observe and analyze changes in the signal as the catheter distal tip 76A advances toward the SA node. When the received ECG signal matches a desired profile, the clinician can determine that the catheter distal tip 76A has reached a desired position with respect to the SA node. As mentioned, in one embodiment this desired position lies within the lower one-third (1/3rd) portion of the SVC.

[00083] The ECG sensor assembly and magnetic assembly can work in concert in assisting a clinician in placing a catheter within the vasculature. Generally, the magnetic assembly of the stylet 130 assists the clinician in generally navigating the vasculature from initial catheter insertion so as to place the distal end 76A of the catheter 72 in the general region of the patient's heart. The ECG sensor assembly can then be employed to guide the catheter distal end 76A to the desired location within the SVC by enabling the clinician to observe changes in the ECG signals produced by the heart as the stylet ECG sensor assembly approaches the SA node. Again, once a suitable ECG signal profile is observed, the clinician can determine that the distal ends of both the stylet 130 and the catheter 72 have arrived at the desired location with respect to the patient's heart. Once it has been positioned as desired, the catheter 72 may be secured in place and the stylet 130 removed from the catheter lumen. It is noted here that the stylet may include one of a variety of configurations in addition to what is explicitly described herein. In one embodiment, the stylet can attach directly to the console instead of an indirect attachment via the TLS sensor. In another embodiment, the structure of the stylet 130 that enables its TLS and ECG-related functionalities can be integrated into the catheter structure itself. For instance, the magnetic assembly and/or ECG sensor assembly can, in one embodiment, be incorporated into the wall of the catheter.

[00084] FIGS. 13A-15 describe various details relating to the passage of ECG signal data from the stylet tether 134 to the TLS sensor 50 positioned on the patient's chest, according the present embodiment. In particular, this embodiment is concerned with passage of ECG signal data from a sterile field surrounding the catheter 72 and insertion site 73, which includes the stylet 130 and tether 134, and a non-sterile field, such as the patient's chest on which the TLS sensor is positioned. Such passage should not disrupt
the sterile field so that the sterility thereof is compromised. A sterile drape that is positioned over the patient 70 during the catheter insertion procedure defines the majority of the sterile field: areas above the drape are sterile, while areas below (excluding the insertion site and immediately surrounding region) are non-sterile. As will be seen, the discussion below includes at least a first communication node associated with the stylet 130, and a second communication node associated with the TLS sensor 50 that operably connect with one another to enable ECG signal data transfer therebetween.

[00085] One embodiment addressing the passage of ECG signal data from the sterile field to the non-sterile field without compromising the sterility of the former is depicted in FIGS. 13A-15, which depict a "through-drape" implementation also referred to as a "shark fin" implementation. In particular, FIG. 14A shows the TLS sensor 50 as described above for placement on the chest of the patient during a catheter insertion procedure. The TLS sensor 50 includes on a top surface thereof a connector base 152 defining a channel 152A in which are disposed three electrical base contacts 154. A fin connector 156, also shown in FIGS. 13A-13D, is sized to be slidingly received by the channel 152A of the connector base 152, as shown in FIG. 14B and 15. Two ECG lead/electrode pairs 158 extend from the fin connector 156 for placement on the shoulder and torso or other suitable external locations on the patient body. The drape-piercing tether connector 132 is configured to slidingly mate with a portion of the fin connector 156, as will be described further below, to complete a conductive pathway from the stylet 120, through the sterile field to the TLS sensor 50.

[00086] FIGS. 13A-13D show further aspects of the fin connector 156. In particular, the fin connector 156 defines a lower barrel portion 160 that is sized to be received in the channel 152A of the connector base 152 (FIGS. 14B, 15). A hole 162 surrounded by a centering cone 164 is included on a back end of an upper barrel portion 166. The upper barrel portion 166 is sized to receive the tether connector 132 of the stylet 130 (FIGS. 14C, 15) such that a pin contact 170 extending into a channel 172 of the tether connector 132 (FIG. 15) is guided by the centering hole until it seats within the hole 162 of the fin connector 156, thus interconnecting the tether connector with the fin connector. An engagement feature, such as the engagement feature 169 shown in FIGS. 13C and 13D, can be included on either side of the fin connector 156 to engage with corresponding detents 173 (FIG. 13F) on the tether connector 132 to assist with maintaining a mating between the two components. If disengagement between the two components is desired,
a sufficient reverse pull force is applied to the tether connector 132 while holding or securing the fin connector 156 to prevent its removal from the channel 152A of the connector base 152.

[00087] FIG. 13D shows that the fin connector 156 includes a plurality of electrical contacts 168. In the present embodiment, three contacts 168 are included: the two forward-most contact each electrically connecting with a terminal end of one of the ECG leads 158, and the rear contact extending into axial proximity of the hole 162 so as to electrically connect with the pin contact 170 of the tether connector 132 when the latter is mated with the fin connector 156 (FIG. 15). A bottom portion of each contact 168 of the fin connector 156 is positioned to electrically connect with a corresponding one of the base contacts 154 of the TLS sensor connector base 152. In one embodiment, the bottom portion of each contact 168 includes a retention feature, such as an indentation 168A. So configured, each contact 168 can resiliently engage a respective one of the base contacts 154 when the fin connector 156 is received by the TLS sensor connector base 152 such that a tip of each base contact is received in the respective indentation 168A. This configuration provides an additional securement (FIG. 15) to assist in preventing premature separation of the fin connector 156 from the connector base 152. Note that many different retention features between the base contacts 154 and the fin contacts 168 can be included in addition to what is shown and described herein.

[00088] FIGS. 13E and 13F depict various details of the tether connector 132 according to one embodiment, including the tether connector channel 172, the pin contact 170 disposed in the channel, and detents 173 for removably engaging the engagement features 169 of the fin connector 156 (FIGS. 13A-13D), as described above. FIG. 13E further shows a plurality of gripping features 171 as an example of structure that can be included to assist the clinician in grasping the tether connector 132.

[00089] FIG. 14B shows a first connection stage for interconnecting the above described components, wherein the fin connector 156 is removably mated with the TLS sensor connector base 152 by the sliding engagement of the lower barrel portion 160 of the fin connector with the connector base channel 152A. This engagement electrically connects the connector base contacts 154 with the corresponding fin contacts 168 (FIG. 15).
FIG. 14C shows a second connection stage, wherein the tether connector 132 is removably mated with the fin connector 156 by the sliding engagement of the tether connector channel 172 with the upper barrel portion 166 of the fin connector. This engagement electrically connects the tether connector pin contact 170 with the back contact 168 of the fin connector 156, as best seen in FIG. 15. In the present embodiment, the horizontal sliding movement of the tether connector 132 with respect to the fin connector 156 is in the same engagement direction as when the fin connector is slidably mated to the sensor connector base channel 152A (FIG. 14B). In one embodiment, one or both of the stylet 130/tether connector 132 and the fin connector 156 are disposable. Also, the tether connector in one embodiment can be mated to the fin connector after the fin connector has been mated to the TLS sensor, while in another embodiment the tether connector can be first mated to the fin connector through the surgical drape before the fin connector is mated to the TLS sensor.

In the connection scheme shown in FIG. 14C, the stylet 130 is operably connected to the TLS sensor 50 via the tether connector 132, thus enabling the ECG sensor assembly of the stylet to communicate ECG signals to the TLS sensor. In addition, the ECG lead/electrode pairs 158 are operably connected to the TLS sensor 50. In one embodiment, therefore, the tether connector 132 is referred to as a first communication node for the stylet 130, while the fin connector 156 is referred to as a second communication node for the TLS sensor 50. As will be seen, various other first and second communication nodes can be employed to enable the establishment of a conductive pathway between the ECG sensor assembly and the TLS sensor or other system component.

Note that various other connective schemes and structures can be employed to establish operable communication between the stylet and the TLS sensor. For instance, the tether connector can use a slicing contact instead of a pin contact to pierce the drape. Or, the fin connector can be integrally formed with the TLS sensor. These and other configurations are therefore embraced within the scope of embodiments of the present disclosure.

As mentioned, a drape 174 is often placed over the patient 70 and employed as a barrier to separate a sterile field of the patient, e.g., areas and components above the drape and proximate to the insertion site 73 (including the catheter 72, the stylet 130, and
tether 134 (FIG. 10)) from non-sterile areas outside of the sterile field, e.g., areas and components below the drape, including the patient's chest, the sensor 50 (FIG. 10) placed on the chest, and regions immediately surrounding the patient 70, also referred to herein as a non-sterile field. As seen in FIG. 15, the sterile drape 174 used during catheter placement to establish the sterile field is interposed between the interconnection of the tether connector 132 with the fin connector 156. As just described, the tether connector 132 includes the pin contact 170 that is configured to pierce the drape 174 when the two components are mated. This piercing forms a small hole, or perforation 175, in the sterile drape 174 that is occupied by the pin contact 170, thus minimizing the size of the drape perforation by the pin contact. Moreover, the fit between the tether connector 132 and the fin connector 156 is such that the perforation in sterile drape made by piercing of the pin contact 170 is enclosed by the tether connector channel 172, thus preserving the sterility of the drape and preventing a breach in the drape that could compromise the sterile barrier established thereby. The tether connector channel 172 is shaped and configured so as to fold the sterile drape 174 down prior to piercing by the pin contact 170 such that the pin contact does not pierce the drape until it is disposed proximate the hole 162 of the fin connector 156 and such that the drape does not bunch up within the channel. It is noted here that the tether connector 132 and fin connector 156 are configured so as to facilitate alignment therebetween blindly through the opaque sterile drape 174, i.e., via palpation absent visualization by the clinician of both components.

[00094] As already mentioned, note further that the fin contacts 168 of the fin connector 156 as shown in FIG. 15 include the indentations 168A, which are configured to mate with the sensor base contacts 154 in such a way as to assist in retaining the fin connector in engagement with the sensor base channel 152A. This in turn reduces the need for additional apparatus to secure the fin connector 156 to the TLS sensor 50. In other embodiments, retention features that are separate from the electrical contacts can be employed to assist in retaining the fin connector in engagement with the sensor base channel. In one embodiment, the base contacts 154 can be configured as pogo pins such that they are vertically displaceable to assist in retaining the fin connector 156.

[00095] FIG. 16 shows a typical ECG waveform 176 of a patient, including a P-wave and a QRS complex. Generally, and with respect to the present system 10, the amplitude of the P-wave varies as a function of distance of the ECG sensor assembly from the SA
node, which produces the P-wave of the waveform 176. A clinician can use this relationship in determining when the catheter tip is properly positioned proximate the heart. For instance, in one implementation the catheter tip is desirably placed within the lower one-third (1/3rd) of the superior vena cava, as has been discussed. The ECG data detected by the ECG sensor assembly of the stylet 130 is used to reproduce waveforms such as the waveform 176, for depiction on the display 30 of the system 10 during ECG mode.

[00096] Reference is now made to FIG. 17 in describing display aspects of ECG signal data on the display 30 when the system 10 is in ECG mode, the third modality described further above, according to one embodiment. The screenshot 178 of the display 30 includes elements of the TLS modality, including a representative image 120 of the TLS sensor 50, with the icon 114 corresponding to the position of the distal end of the stylet 130 during transit through the patient vasculature. The screenshot 178 further includes a window 180 in which the current ECG waveform captured by the ECG sensor assembly of the stylet 130 and processed by the system 10 is displayed. The window 180 is continually refreshed as new waveforms are detected.

[00097] Window 182 includes a successive depiction of the most recent detected ECG waveforms, and includes a refresh bar 182A, which moves laterally to refresh the waveforms as they are detected. Window 184A is used to display a baseline ECG waveform, captured before the ECG sensor assembly is brought into proximity with the SA node, for comparison purposes to assist the clinician in determining when the desired catheter tip location has been achieved. Windows 184B and 184C can be filled by user-selected detected ECG waveforms when the user pushes a predetermined button on the probe 40 or the console button interface 32. The waveforms in the windows 184B and 184C remain until overwritten by new waveforms as a result of user selection via button pushes or other input. As in previous modes, the depth scale 124, status/action indicia 126, and button icons 128 are included on the display 30. An integrity indicator 186 is also included on the display 30 to give an indication of whether the ECG lead/electrode pairs 158 are operably connected to the TLS sensor 50 and the patient 70.

[00098] As seen above, therefore, the display 30 depicts in one embodiment elements of both the TLS and ECG modalities simultaneously on a single screen, thus offering the clinician ample data to assist in placing the catheter distal tip in a desired position. Note
further that in one embodiment a printout of the screenshot or selected ECG or TLS data can be saved, printed, or otherwise preserved by the system 10 to enable documentation of proper catheter placement.

[00099] Although the embodiments described herein relate to a particular configuration of a catheter, such as a PICC or CVC, such embodiments are merely exemplary. Accordingly, the principles of the present invention can be extended to catheters of many different configurations and designs.

[00100] FIGS. 18-19B depict examples of contact engagement configurations for the tether connector 132 and fin connector 156. Specifically, FIG. 18 depicts the fin contacts 168 of the fin connector 156 according to one embodiment, wherein the rear contact includes a spring clip configuration 168B for receiving the pin contact 170 (FIG. 15) of the tether connector 132 via the centering cone 164 or other aperture defined in the fin connector. FIGS. 19A and 19B depict an engagement scheme according to another embodiment, wherein the pin contact 170 of the tether connector 132 includes a barbed feature 170A that, when inserted into the centering cone 164 or other aperture of the fin connector 156, engages a shoulder 168C defined on the rear fin contact 168 of the fin connector so as to help prevent premature removal of the pin contact from the fin connector. These embodiments thus serve as non-limiting examples of a variety of contact configurations that can be included with the fin connector 156, the sensor connector base 152, and the tether connector 132. Note that unless referred to as otherwise, the contacts described herein are understood to include electrical contacts used in establishing a conductive pathway.

[00101] The embodiments to be described below in connection with FIGS. 20A-32 each depict an example connection scheme as a means for establishing a conductive or other communication pathway between a patient's sterile field and a non-sterile field, i.e., areas outside of the sterile field. Thus, the embodiments described herein serve as examples of structure, material, and/or compositions corresponding to the means for establishing a conductive or other communication pathway. In particular, various embodiments described herein disclose examples for breaching or otherwise circumventing a sterile barrier separating the sterile field from the non-sterile field so as to provide at least a portion of the conductive pathway for the passage of ECG signals from a sensing component such as the ECG sensor assembly of the stylet 130 to the
sensor 50, also referred to herein as a TLS sensor or chest sensor, or other suitable data-receiving component of the system 10. Note that these embodiments are merely examples of a variety of means for establishing such a conductive or other communication pathway, and are not to be considered limiting of the scope of the present disclosure. It is therefore appreciated that the means for establishing a conductive or other communication pathway can be employed for transferring ECG signals or other information, electrical signals, optical signals, etc.

[000102] As will be seen, many of the embodiments to be described include a tether connector, also referred to herein as a first communication node, which is operably connected to the stylet 130 and included in the sterile field, the tether connector is configured to operably attach to a connector included on the sensor 50 or other suitable component of the system 10, also referred to herein as a second communications node, which is disposed outside of the sterile field. Note, however, that the first communication node and second communication node are contemplated as generally referring to various connector interfaces that provide a conductive pathway from the sterile field to the non-sterile field to enable the passage of ECG signals as described above. It is appreciated that the conductive pathway is a communication pathway and includes an electrical pathway, an optical pathway, etc. Further, the communication node connection schemes described and contemplated herein can be employed with systems involving the use of modalities exclusive of ECG signals for navigation or placement of a catheter or other medical device.

[000103] Note further that the embodiments to follow that describe configurations for breaching a drape or other non-transparent sterile barrier are configured such that location of a communication node disposed out-of-sight under the drape/barrier is facilitated by palpation of the clinician, thus easing location and connection of the first and second communication nodes. Also, many of the connector configurations described herein can be configured as one-use, disposable components so as to minimize concerns with infection.

[000104] Reference is now made to FIGS. 20A-20C, which depict a connection scheme as a means for establishing a conductive pathway between sterile and non-sterile fields, according to one embodiment. In particular, FIGS. 20A-20C depict a tether connector 232 that includes an outer housing 234 and a blade holder 236 that attaches to the outer
housing. A blade contact 238 is secured by the blade holder 236 such that the blade
contact extends into a channel 240 of the tether connector. The blade contact 238 serves
to create a slice perforation in a drape that is interposed between the tether connector and
the fin connector 256 when the tether connector 232 is slid on to engage the fin
connector in the manner described in previous embodiments. As before, the outer
housing 234 of the tether connector envelopes and protects the perforation so as to prevent
contamination and compromise of the sterile field.

[000105] FIG. 20C shows that a fin connector 256 includes a fin contact 268 that is
colocated to physically interconnect with the blade contact 238 when the tether
connector is slid on to the fin connector 256, thus establishing a conductive pathway
through the sheath so as to enable ECG signals from an ECG sensing component, i.e., the
ECG sensor assembly described above for instance, to pass to the sensor 50 via the blade
contact 238/fin contact 268 engagement. Note that the particular configuration of the
blade and fin contacts can be varied from what is described herein. For instance, the
tether connector can include two or more blades or contacts for engagement with
the corresponding fin contacts to enable multiple conductive pathways to be established, if
desired. The engagement surfaces of the tether connector and the fin connector can also
vary from what is shown and described. In one embodiment, a light source can be
included with the fin connector or other connectors as described herein so as to provide
illumination through the drape 174 and provide visual assistance in locating the fin
connector for interconnection with the tether connector.

[000106] As seen in FIGS. 14A and 14B, in one embodiment the ECG leads 158 are
permanently connected to the fin connector 156. FIG. 21A depicts another possible
embodiment, wherein the ECG leads are removably attached to the fin connector 156 via
a connector, such as a horseshoe connector 270, best seen in FIG. 21B. FIG. 21A further
shows that the fin connector 156 is permanently attached to the sensor 50. These and
other variations in the connective schemes of the various components of the system 10
are therefore contemplated as falling within the scope of the present disclosure. In
another embodiment, the electrode of each lead is removably attachable from the lead,
such as via a snap connection, for instance.

[000107] Reference is now made to FIGS. 22A-22C, which depict a connection scheme
as a means for establishing a conductive pathway between sterile and non-sterile fields,
according to one embodiment. In particular, FIGS. 22A-22C depict a tether connector 332 that includes a channel 372 for slidably engaging an upper barrel 166 of a fin connector 356 disposed on the sensor 50, in a manner similar to previous embodiments. The tether connector 332 includes a bi-positional top cap 374 to which is attached a pin contact 370 or other piercing contact.

[000108] The top cap 374 is positioned in an un-actuated first position, shown in phantom in FIG. 22B, when the tether connector 332 is first slid on to the fin connector 356. The drape, removed for clarity, is interposed between the upper barrel 166 of the fin connector 356 and the tether connector channel 372, similar to earlier embodiments. After the tether connector 332 is positioned on the fin connector 356, the top cap 374 can then be depressed by the clinician into an actuated second position shown in FIG. 22B, wherein the pin contact 370 is pressed downward through the drape and into operable engagement with a corresponding contact disposed in the fin connector 356. The tether connector 332 is thus positioned as shown in FIG. 22C. In addition to establishing a conductive path through the drape 174, this engagement of the pin contact 370 locks the tether connector 332 on to the fin connector 356 so as to prevent premature separation of the components.

[000109] Reference is now made to FIG. 23A and 23B, which depict a connection scheme as a means for establishing a conductive pathway between sterile and non-sterile fields, according to one embodiment. In particular, FIG. 23A depicts a tether connector 432 including a pin contact 440 or other suitable contact attached to an actuation assembly 442. The actuation assembly 442 includes lever arms for selectively lowering the pin contact 440 through an opening defined by a male end 448 of a housing 446 in which the actuation assembly is disposed. The male end 448 of the housing is configured to be received by a sensor connector receptacle 450 disposed on the sensor 50 or other suitable component of the system, such as a remote module operably connected to the sensor, for instance.

[000110] To interconnect the tether connector 432 to the sensor connector receptacle 450, the male end 448 of the tether connector 432 is brought, above the drape 174, into proximity with the receptacle 450. The actuation assembly 442 is then actuated by raising the lever arms 444, as shown in FIG. 23B. The pin contact 440 is forced downward through the drape 174, thus defining a perforation therein. The male end 448
can then be fully received into the sensor receptacle 450, wherein the pin contact 440 operably connects with a suitable contact of the sensor connector receptacle. The connector scheme shown in FIG. 23A and 23B is useful for imposing a minimal downward force on the body of the patient during connector interconnection. Further, the actuation assembly 442 provides a predetermined force in connecting the first communication node (the tether connector 432) with the second communication node (the sensor connector receptacle 450), and thus does not rely on a clinician's estimation of force to establish the node connection. In another embodiment, the housing 446 and the sensor receptacle 450 can be aligned and mated before the actuation assembly 442 is actuated to pierce the contact 440 through the drape.

[000111] Reference is now made to FIG. 24, which depicts a connection scheme as a means for establishing a conductive pathway between sterile and non-sterile fields, according to one embodiment. As in the embodiment shown in FIGS. 23A and 23B, the present interconnection scheme minimizes downward pressure on the body of the patient during interconnection of the nodes. As shown, a tether connector 532 includes a pin contact 540 or other suitable contact included with a threaded cap 542, which defines threads on an inside surface thereof. The threaded cap 542 is configured to threadingly receive a threaded base 544 disposed on the sensor 50 or other suitable component of the system, such as a remote module operably connected to the sensor, for instance. As before, the drape 174 is interposed therebetween.

[000112] To interconnect the tether connector 532 to the sensor 50, the threaded cap 542 of the tether connector is brought, above the drape 174, into proximity with the threaded base 544 and threaded on to the base. This causes the pin contact 540 to penetrate the drape 174, thus defining a perforation therein. Further threading of the cap 542 on to the base 544 causes the pin contact 540 to engage a contact receptacle 546 included in the base 544, thus operably interconnecting the two nodes. In one embodiment, the tether 134 is rotatably attached to the threaded cap 542 so as to prevent twisting of the tether during threading. The connector scheme shown in FIG. 24 is useful for imposing a minimal downward force on the body of the patient during connector interconnection as the force to join the two connectors is directed laterally with respect to the patient via the threading operation. Note further that a variety of thread configurations and locations, as well as different cap and base configurations, are contemplated by the present disclosure.
Reference is now made to FIGS. 25A and 25B, which depict a connection scheme as a means for establishing a conductive pathway between sterile and non-sterile fields, according to one embodiment. As in the previous embodiment, the present interconnection scheme minimizes downward pressure on the body of the patient during interconnection of the nodes. As depicted in FIGS. 25A and 25B, a tether connector 632 includes one or more piercing contacts, such as pin contacts 640A and 640B that are respectively included on slide arms 642A and 642B. One or more contact receptacles, such as contact receptacles 644A and 644B, are included on a portion of the sensor 50, such as a sensor fin 646, or other suitable system component. As before, the drape 174 is interposed between the tether connector 632 and the sensor fin 646 to serve as a sterile barrier.

To interconnect the tether connector 632 to the sensor fin 646, the tether connector is brought, above the drape 174, into proximity with the sensor fin such that the slide arms 642A and 642B straddle the sensor fin and such that the pin contacts 640A and 640B are aligned with corresponding contact receptacles 644A and 644B, as shown in FIG. 25A. The slide arms 642A and 642B are then slid toward one another such that the pin contacts 640A and 640B penetrate the drape 174, each defining a perforation therein. The slide arms 642A and 642B are slid inward until the pin contacts 640A and 640B seat within and operably connect with the corresponding contact receptacles 644A and 644B, as seen in FIG. 25B, thus interconnecting the two nodes. The connector scheme shown in FIGS. 25A and 25B is useful for imposing a minimal downward force on the body of the patient during connector interconnection as the force to join the two connectors is directed laterally with respect to the patient. Note that the particular configuration of the tether connector, the sensor fin, and the contacts can vary from what is explicitly described herein. For instance, in one embodiment the slide arms can be configured as bi-positional rocker arms that are connected in a see-saw configuration with respect to one another. Also, one, two, or more contacts can be included on the slide arms.

Reference is now made to FIG. 26A and 26B, which depict a connection scheme as a means for establishing a conductive pathway between sterile and non-sterile fields, according to one embodiment. As shown, an integrated connector 730 is incorporated into the drape 174 so as to enable operable interconnection therethrough. In the illustrated embodiment, the integrated connector 730 includes a conductive base.
portion 734 from which extend mechanical connectors, such as snap balls 736A and 736B.

As shown in FIG. 26B, the integrated connector 730 is positioned in the drape 174 as to be connectable with both a suitable receptacle 738 of a tether connector 732 and a suitable receptacle 740 of the sensor 50 or other suitable component of the system 10. In particular, the tether connector 732 can be snap-attached to the integrated connector 730, after which the integrated connector can be attached to the sensor 50, thus providing a suitable pathway for signals from the ECG sensor assembly in the sterile field to be transmitted through the sterile barrier of the drape 174 to the sensor in the non-sterile field. It is appreciated that, in other embodiments, the integrated connector can include other configurations, such as different mechanical connectors, e.g., friction connectors, male/female connectors, etc., and as such the receptacles on the tether connector and sensor can likewise be modified to accommodate the different mechanical connectors. Also, the connective scheme described above can be reversed such that the receptacles are included on the integrated connector and the snap balls on the respective tether connector and sensor. Further, though presently depicted as a unitary component, the integrated connector in other embodiments can include two or more pieces that are attached to each other through a previously defined hole in the drape during manufacture thereof. These and other variations are therefore contemplated.

Reference is now made to FIG. 27, which depicts a connection scheme as a means for establishing a conductive pathway between sterile and non-sterile fields, according to one embodiment. In detail, FIG. 27 depicts an intermediate module, i.e., ECG module 750, disposed outside of the sterile field of the patient, which is operably connected to the sensor 50 of the system 10 via a sensor cable 752. The ECG module 750 is also operably connected to the ECG leads 158. In one embodiment, the ECG module 750 includes the circuitry and other components necessary for receipt and analysis of the ECG signal detected by the ECG sensor assembly of the stylet 130. As such, a conductive pathway is established between the stylet 130 and the ECG module 750 by traversing the sterile field of the patient. In the present embodiment, this is accomplished by a tether connector 762 of the tether 134.

As depicted in FIG. 27, the tether connector 762 operably attaches to a receptacle 764 of the ECG module 750. As shown, the tether connector 762 can include
a sufficiently long handle that enables the clinician to attach the sterile tether connector
to the receptacle 764 of the non-sterile ECG module 750 without touching the ECG
module itself, thus preventing any compromise of the sterile field. In one embodiment,
the handle of the tether connector 762 can include an extendable J-hook contact, for
instance, that can operably connect to a suitable contact of the ECG module.

[000119] FIG. 28 shows another example of a tether connector that can be employed
with the ECG module 750 of FIG. 27 or other suitable component of the system 10 as
part of a connection scheme as a means for establishing a conductive pathway between
sterile and non-sterile fields, according to one embodiment. In particular, FIG. 28
depicts a tether connector 832, which includes a handle and a barbed contact 836 or other
suitable contact at a proximal end thereof. A sterile shield 838 is interposed between the
handle 834 and the contact 836. The sterile shield 838 assists in protecting the hand of
the clinician while inserting the contact 836 into the receptacle 764 of the ECG module
750 in a manner similar to what is shown in FIG. 27. Thus, the sterile shield 838 serves
as an additional barrier to prevent inadvertent contact by the clinician with a component
outside of the sterile field, such as the ECG module 750. Note that the size, shape, and
particular configuration of the sterile shield and/or tether connector can vary from what
is explicitly described in the present embodiment.

[000120] FIGS. 29A and 29B show yet another example of a connection scheme that
can be employed with the ECG module 750 of FIG. 27 or other suitable component of
the system 10 as a means for establishing a conductive pathway between sterile and non-
sterile fields, according to one embodiment. In particular, FIG. 29A shows that the ECG
module 750 can be enveloped by a sterile bag 850. A connector, such as the integrated
connector 730 described above in connection with FIGS. 26A and 26B, can be
incorporated into the bag. As shown in FIG. 29B, an inner snap ball or other mechanical
connector of the integrated connector 730 can be received by the suitably corresponding
receptacle 764 of the ECG module 750. The tether connector of the system 10 can then
be operably connected with the outer snap ball or other connector of the integrated
connector 730, thus establishing a conductive pathway between the sterile field and the
non-sterile field without compromising sterility. Note that the sterile bag 850 can
include any one or more of a variety of suitable materials, including plastic. Note also
that the integrated connector can include other connector configurations in addition to
what is explicitly described herein. In one embodiment, the sterile bag includes no
integrated connector, but rather is pierced by a pin contact of the tether connector, such as the barbed contact 836 included on the tether connector 832 of FIG. 28.

[000121] Reference is now made to FIG. 30, which depicts a connection scheme as a means for establishing a conductive pathway between sterile and non-sterile fields, according to one embodiment. Specifically, the stylet 130 includes a tether connector 862 as a first communication node, as in previous embodiments. A remote sensor connector 864 is also included as a second communications node, and is operably connected to the sensor 50 of the system 10 via a remote sensor connector cable 866. The tether connector 862 and remote sensor connector 864 operably connect to one another along a connection interface 868. The drape 174 that serves as a sterile barrier is interposed between the tether connector 862 and remote sensor connector 864 at the connection interface 868, and a suitable drape piercing configuration is included with the tether connector and the remote sensor connector to establish a conductive pathway through the drape. The present embodiment thus discloses one embodiment wherein the second communication node is located remotely with respect to the sensor 50.

[000122] Reference is now made to FIG. 31, which depicts a connection scheme as a means for establishing a conductive pathway between sterile and non-sterile fields, according to one embodiment. Specifically, the present embodiment includes the tether connector 862 and the remote sensor connector 864 that operably connect to one another along the connection interface 868, as described in connection with FIG. 30, above. The remote sensor connector 864 in the present embodiment is placed proximate the catheter insertion site 73 in a region over which a fenestration 880 defined in the drape 174 (portions of the drape omitted for clarity) is positioned to enable clinician access to the insertion site during catheter placement. The remote sensor connector 864 is adhered to the patient's skin proximate the catheter insertion site 73 with the use of an adhesive, tape, etc., before the region surrounding the insertion site is sterilized in preparation for catheter insertion. Thus, when the insertion site is sterilized, the remote sensor connector 864 is also sterilized. Later, when connection of the tether connector 862 to the remote sensor connector 864 is made, the clinician can handle the latter component without compromising the sterile field of the patient. It is appreciated that the particular configurations of the tether connector and the remote sensor connector can vary while still residing within the scope of the present embodiment.
Reference is now made to FIG. 32, which depicts a connection scheme as a means for establishing a conductive pathway between sterile and non-sterile fields, according to one embodiment. Specifically, FIG. 32 shows the probe 40 employed by the system 10 for US functionality, as described above in connection with FIGS. 3A and 3B. A sterile sheath 900 is placed over the probe 40 so as to bring the probe into the sterile field of the patient. A connection interface, such as a receptacle 910, is included on the probe 900 and is configured so as to be operable connectable with a tether connector 920. In one embodiment, for example, the tether connector 920 includes a pin contact that penetrates the sterile sheath 900 to mate with the receptacle 910 in such a way as to prevent contamination of the sterile field. In this way, the tether connector 920, as a first communication node, operably connects with the probe 40, as a second communications node. In turn, the probe 40 is operably connected to the system console 20, as seen in FIG. 31 for example, so as to enable ECG signals received by the ECG sensor assembly of the stylet 130 via the tether connector 920 to be forwarded to the console, the sensor 50, or other system component for processing, as described above. In another embodiment, the receptacle 910 or other suitable connection interface can be included on the cable connecting the probe 40 to the system console 20. The particular contact configuration of the receptacle 910 and tether connector 920 can be varied according to the understanding of one skilled in the art. For instance, an integrated connector such as that shown in FIGS. 26A and 26B can be incorporated into the sterile sheath in one embodiment. Note further that, though including plastic in the present embodiment, the sterile sheath as described herein can include other suitable materials for providing sterility.

Reference is now made to FIG. 33 in describing means for establishing a conductive pathway between sterile and non-sterile fields, according to one embodiment. As shown, the tether 134 includes a wireless module 950, included within the sterile field, which serves as a first communication node for wirelessly transmitting (via RF or other suitable frequency or frequency range) ECG data received from the ECG sensor assembly of the stylet 130 to a data-receiving component as a second communication node, such as the sensor 50 or other suitable component of the system 10. A wireless module ground electrode 952 is operably connected with the wireless module 950 for placement in the sterile field proximate the catheter insertion site 73. A system ground electrode 158A extends from the sensor 50 for placement outside of the sterile field but
proximate both the catheter insertion site 73 and the location of the wireless module ground electrode 952. One possible placement location for the system ground electrode 158A is beneath the patient arm, as depicted in FIG. 33. The system reference electrode 158B is placed on the lower torso of the patient 70 or other suitable location, as in previous embodiments. Note that the wireless module and system console as discussed herein can be configured in one or more of a variety of ways and include components for wireless signal transmission and reception not specifically detailed herein, such as patch or other antennas, signal transducers, etc.

[000125] With the system configured as shown in FIG. 33, the system ground electrode 158A can be electrically driven such that it produces a voltage that is sensed by the passive wireless module ground electrode 952, given its proximate location with respect to the system ground electrode. This enables both ground electrodes to be at substantially equal electric potentials, thus enabling the wireless module 950 to utilize the wireless module ground electrode 952 and the ECG signals from the ECG sensor assembly of the stylet 130, e.g., the core wire 138 (FIGS. 12C-12E) in one embodiment, to detect and wirelessly transmit the ECG data to the sensor 50 for comparison with the data sensed by the system reference electrode 158B in order to obtain the desired P-wave waveform (e.g., FIG. 16). The data comparison in one embodiment is a differential comparison between the ECG data as obtained by the ECG sensor assembly of the stylet 130, the wireless module ground electrode 952, and the system reference electrode 158B. In one embodiment, the system ground electrode 158A, like the wireless module ground electrode 952, can be passive and not electrically driven. Note also that the analog ECG data can be digitized or otherwise processed by the wireless module 950 before transmission to the sensor 50 or other system component, such as the console 20.

[000126] FIG. 34 describes yet another wireless configuration as a means for establishing a conductive pathway between sterile and non-sterile fields, according to one embodiment. As shown, a positive electrode 954A at a location A and a negative electrode 954B at a location B are included with the sensor 50 and positioned on the torso of the patient 70, while a positive wireless module electrode 956 is included with the wireless node 950, as indicated at location C, positioned on or in the patient proximate the catheter insertion site 73. The ECG sensor assembly of the stylet 130, e.g., the core wire 138 in one embodiment, serves as a negative electrode for the wireless portion of the depicted configuration, indicated at D in FIG. 34 at its final position. Note
that in one embodiment the locations A and B of the electrodes 954A and 954B, respectively, can be altered on the patient body to tune the system 10 for best ECG signal reception.

[000127] In the present embodiment, the electrodes 954A and 954B serve as a first independent source for sampling bipolar ECG signals. The ECG data from these electrodes are digitized and forwarded to the console 20 or other suitable system component via the cable interconnecting the sensor 50 and the console (path 1) outside of the sterile field. The wireless module electrode 956 and the ECG sensor assembly serve as a second independent source for sampling bipolar ECG signals. The ECG data from these electrodes are digitized and forwarded wirelessly to the console 20 via the wireless module 950 (path 2) within the sterile field. Thus, in the present embodiment the wireless module 950 serves as a first communication node, and a wireless receiver of the console 20 as a second communication node for the transfer of ECG signals between the two nodes. Note that the polarities of the afore-mentioned electrodes can be reversed in other embodiments.

[000128] The ECG signals received along both paths 1 and 2 are baseline corrected by appropriate circuitry of the console 20 to adjust for DC offset and drift. After such correction, a non-changing reference, or baseline, P-wave waveform 176A from path 1 can be produced, as seen in FIG. 35A, for example. Similarly, a P-wave waveform 176B as seen in FIG. 35B is produced from path 2, which waveform changes as the stylet 130 within the catheter 72 is advanced toward the heart of the patient. During such advancement, the waveform 176B from path 2 is subtracted from the P-wave waveform 176A from path 1, employing a digital differential amplifier, for instance. This subtraction removes all common components of the waveforms represented by each of the signals, and enables the console 20 to depict via its display 30 only the differences in the two signals, as seen for example by the waveform 176C shown in FIG. 35C. The change in P-wave of the waveform from path 2 can then be easily observed during catheter advancement. Thus the present embodiment enables an easily observable digital display of ECG data to be represented while preventing a physical breaching of a sterile barrier, such as a surgical drape, for the passage of such data.

[000129] Note that in other embodiments the wireless module electrode 956 can include other configurations, including a conductive element imbedded into an
introducer sheath, in contact with the bloodstream of the patient, which is commonly disposed through the insertion site 73 during catheter placement. The introducer can include a connector on a proximal portion thereof to enable a connection with the wireless node 950 to be made, in one embodiment.

[000130] Note further that one or more of a variety of wireless protocols can be employed in transmitting wireless signals in accordance with the embodiments described herein, including one or more of the IEEE 802.11 family of specifications, etc. Also note that in one embodiment the wireless module can be included in a sterile sheath, as described in previous embodiments, to bring the module within the sterile field, together with connectors for operably connecting the wireless module electrode through the sheath or included in the sheath itself. Of course, other methods for maintaining the wireless module within the sterile field can also be employed. In one embodiment, the wireless module can include buttons that further enable control of the system 10 from within the sterile field.

[000131] FIG. 36 shows that in one embodiment the sensor 50 can be retro-fitted with a wireless module 960 to enable signals received by the sensor to be wirelessly transmitted to the console 20 or other suitable component of the system 10. For instance, ECG data received by the ground and reference electrodes 158A, 158B (FIG. 34) can be received by the sensor 50 then wirelessly transmitted to the system console via the wireless module 960. The wireless module 960 can include an antenna or other transmitting component and can operably connect to the sensor 50 via a sensor cable 962 or other suitable interface. Note that the wireless module 960 can be employed in connection with other embodiments described herein, including those depicted in FIGS. 10 and 33, for instance.

[000132] FIG. 37 shows a retention feature for preventing inadvertent separation of the fin connector 156 from the sensor connector base 152 or other receptacle with which the fin connector operably connects, according to one embodiment. As shown, the fin connector 156 includes a retention arm 970 that is resiliently attached to the fin connector body. The retention arm 970 includes a tab 972 that slides over and engages a lip 974 included with the connector base 152 of the sensor 50 when the fin connector 156 is slidably received in the sensor channel 152A (FIG. 14A). The engagement of the tab 972 with the lip 974 prevents inadvertent removal of the fin connector 156 during use.
When removal of the fin connector 156 from the sensor connector base 152 is desired, the retention arm 970 is lifted so as to disengage the tab 972 from the lip 974, after which the fin connector can be slid our of engagement with the sensor channel 152A. This configuration can be employed either with or independent of other retention features, such as the indentations 168A (FIG. 13D). Note that in other embodiments a variety of modifications and configurations can be employed in assisting to maintain engagement between the fin connector and the connector. For instance, the retention arm in one embodiment can be operably attached to one or more of the fin contacts 168 (FIG. 13D) such that displacement, e.g., lifting laterally moving, pinching, etc., of the retention arm or other suitable fin connector component disengages the fin contact(s) from the base contacts (FIG. 15), thus reducing the overall retention force provided by the engagement of the fin contacts with the base contacts. Note further that these principles can be applied to the other connector schemes disclosed or contemplated in addition to the fin connector described here.

[000133] In addition to the above embodiments depicting various connection schemes as means for establishing a conductive pathway between sterile and non-sterile fields, other configurations can be employed, as appreciated by one skilled in the art, for performing the same functionality. Such other configurations can include, for example, wireless transmission of ECG signals from the stylet to the sensor or the system component, the inclusion of electrically conductive thread in the drape, the inclusion of an electrically conductive window (e.g., composed of an electrically conductive plastic or foil) in the sterile drape, etc. In yet another embodiment, a proximal end of the stylet/guidewire itself can be used to pierce the drape for receipt into a connector on the sensor. In this case, no tether is included on the proximal end of the stylet, and the stylet itself serves as the conductive pathway for transmitting ECG signals from the stylet sensor assembly to the sensor on the patient's chest. Such a configuration can allow for over-the-wire placement of the catheter using a stylet/guidewire as described here. As such, the above embodiments should not be construed as being limiting of the present invention in any way.

[000134] Embodiments of the invention may be embodied in other specific forms without departing from the spirit of the present disclosure. The described embodiments are to be considered in all respects only as illustrative, not restrictive. The scope of the embodiments is, therefore, indicated by the appended claims rather than by the foregoing
description. All changes that come within the meaning and range of equivalency of the claims are to be embraced within their scope.

What is claimed is:
CLAIMS

1. A medical device placement system, comprising:
   a medical device assembly including a medical device for placement within a body of a patient and a sensing component, the medical device assembly at least partially disposed in a sterile field;
   a data-receiving component at least partially disposed in a non-sterile field; and
   means for establishing a conductive pathway between the sterile field and the non-sterile field so as to operably connect the sensing component to the data-receiving component.

2. The placement system as defined in claim 1, wherein the medical device includes a catheter, wherein the sensing component is an ECG sensing component for detecting ECG signal data relating to a heart of a patient after insertion of the catheter into the vasculature of the patient, and wherein the data-receiving component is a chest sensor that receives the ECG signal data from the ECG sensing component via the conductive pathway.

3. The placement system as defined in claim 1, wherein the means for establishing the conductive pathway breaches or circumvents a sterile barrier that at least partially separates the sterile field from the non-sterile field.

4. The placement system as defined in claim 3, wherein the sterile barrier includes a drape and wherein the means for establishing the conductive pathway breaches or circumvents the drape in such a way as to prevent contamination of sterile field.

5. The placement system as defined in claim 4, wherein the means establishing the conductive pathway provides a perforation in the drape while also preventing access to the perforation.
6. The placement system as defined in claim 1, wherein the means for establishing the conductive pathway includes:
   a first connector operably connected to the sensing component, the first connector including a first contact; and
   a second connector operably connected to the data-receiving component, the second connector including a second contact that operably connects with the first contact when the first and second connectors connect with one another.

7. The placement system as defined in claim 6, wherein the first and second contacts physically connect to one another through a sterile drape to provide the conductive pathway between the sterile field and the non-sterile field.

8. The placement system as defined in claim 7, wherein the first connector is a tether connector that is operably connected to the sensing component via a tether, and wherein the second connector is directly or remotely attached to the data-receiving component.

9. The placement system as defined in claim 8, wherein the first contact includes a blade for creating a perforation in the drape when the first and second connectors connect with one another.

10. The placement system as defined in claim 8, wherein the first contact includes a pin contact for piercing the drape so as to operably connect with the second contact.

11. The placement system as defined in claim 10, wherein engagement of the pin contact with the second contact prevents relative movement between the tether connector and the second connector.

12. The placement system as defined in claim 8, wherein the tether connector includes a male end that is received into a receptacle included on the data-receiving component.

13. The placement system as defined in claim 8, wherein the tether connector is threadably engageable with the second connector.
14. The placement system as defined in claim 8, wherein the tether connector includes at least one slide arm on which the first contact is included, the slide arm being selectively movable so as to operably connect the first contact with a second contact of the data-receiving component.

15. The placement system as defined in claim 8, wherein the tether connector is selectively connectable to an intermediate module, the intermediate module including the second connector and being operably connected to the data-receiving component.

16. The placement system as defined in claim 15, wherein the tether connector includes a sterile shield disposed proximate the first contact to protect sterility of a hand of a clinician when operably connecting the tether connector to the intermediate module.

17. The placement system as defined in claim 15, wherein the intermediate module is disposed within a sterile sheath so that the intermediate module is disposed within the sterile field, the tether connector able to operably connect with the intermediate module through the sterile sheath.

18. The placement system as defined in claim 17, wherein the intermediate module includes an ultrasound probe of the medical device placement system.

19. The placement system as defined in claim 8, wherein the second connector is remotely disposed from the data-receiving component and wherein the first connector and the second connector operably connect to one another via a fenestration defined in the drape, the fenestration disposed proximate an insertion site for the medical device on the body of the patient.

20. The placement system as defined in claim 1, wherein the means for establishing the conductive pathway includes an integrated connector included in a sterile drape that at least partially separates the sterile field from the non-sterile field.

21. The placement system as defined in claim 20, wherein the integrated connector includes a snap connector for operably connecting with a first connector associated with the sensing component and a second connector associated with the data-receiving component.
22. The placement system as defined in claim 1, wherein the means for establishing the conductive pathway includes a wireless link between the sensing component and the data-receiving component.

23. A method for operably connecting an ECG sensing component included with a medical device at least partially disposed in a sterile field of a patient with a data-receiving component at least partially disposed in a non-sterile field, the method comprising:

    providing the medical device and the ECG sensing component, the ECG sensing component configured to detect ECG signals of the patient after the medical device is at least partially inserted into a body of the patient, the ECG sensing component operably connected to a first communication node disposed in the sterile field; and

    operably connecting a second communication node of the data-receiving component to the first communication node without compromising the sterile field.

24. The method for operably connecting as defined in claim 23, wherein operably connecting the first and second communication nodes enables ECG signals to be passed from the ECG sensing component of the medical device to the data-receiving component, and wherein the second communication node is at least partially disposed in the non-sterile field.

25. The method for operably connecting as defined in claim 23, wherein the sterile and the non-sterile fields are separated by a sterile drape, and wherein operably connecting the second communication node with the first communication node further includes:

    physically connecting the first communication node with the second communication node via a perforation in the drape, a physical connection of the first and second communication nodes covering the breach so as to not compromise the sterile field.

26. The method for operably connecting as defined in claim 25, wherein physically connecting the first communication node with the second communication node produces the perforation in the drape.
27. The method for operably connecting as defined in claim 23, wherein operably connecting the second communication node further includes:

operably connecting the second communication node of the data-receiving component to the first communication node in manner using a force that is substantially lateral with respect to the body of the patient.

28. The method for operably connecting as defined in claim 23, wherein operably connecting the second communication node further comprises:

wirelessly connecting the second communication node of the data-receiving component to the first communication node.

29. A catheter system for providing a conductive pathway between a sterile field of a patient and a non-sterile field, the system comprising:

a stylet positionable in a lumen of a catheter, the stylet including an ECG sensing component that detects ECG signals of the patient, the stylet further including a first connector operably connected to the ECG sensing component and disposed within the sterile field; and

a data-receiving component including a second connector, the second connector at least partially disposed in the non-sterile field and operably connectable to the first connector through a sterile drape without compromising the sterile field such that the ECG signals can be received by the data-receiving component.

30. The catheter system as defined in claim 29, wherein the data-receiving component is a chest sensor of a catheter placement system, and wherein the second connector is removably attachable to the chest sensor.

31. The catheter system as defined in claim 30, wherein the first connector includes a body defining a channel and a pin contact extending into the channel, the pin contact being operably connected to the ECG sensing component via a tether, wherein the first connector slidably receives in the channel thereof an upper portion of the second connector such that the pin contact pierces the drape to define a perforation and operably connects with a contact of the second connector, and wherein the perforation is confined within the channel of the first connector.
32. The catheter system as defined in claim 31, wherein the chest sensor includes a channel that slidably receives a lower portion of the second connector such that at least one contact of the chest sensor is placed in operable communication with at least one contact of the second connector.

33. The catheter system as defined in claim 32, wherein the at least one contact of the second connector includes an indented retention feature that physically engages with the at least one contact of the chest sensor to prevent inadvertent separation of the second connector from the chest sensor.

34. The catheter system as defined in claim 33, wherein the second connector further includes at least one engagement tab that is received by at least one detent defined on the body of the first connector so as to prevent inadvertent separation of the first connector from the second connector.

35. A method for creating a connection through a sterile barrier, the method comprising:
   positioning a sensor on a patient;
   placing the sterile barrier over the sensor; and
   establishing a connection through the sterile barrier between the sensor and a distal end of a catheter.

36. The method for creating a connection as defined in claim 35, further comprising:
   positioning a catheter into the patient using the connection.

37. A method for positioning a catheter within a vasculature of a patient, the method comprising:
   positioning a sensor on the patient;
   placing a sterile barrier over the sensor;
   establishing a connection through the sterile barrier between the sensor and a distal end of the catheter; and
   inserting the catheter into the vasculature of the patient using the connection to position the catheter within the vasculature of the patient.
38. The method for positioning as defined in claim 37, wherein inserting the catheter further includes:

inserting the catheter into the vasculature of the patient using ECG signals from a heart of the patient, the ECG signals being provided via the connection.