Title: INFUSION THERAPY FLOW RATE ADJUSTMENT SYSTEM AND METHOD

Abstract: Disclosed is a system and method for assisting in verifying the right medication is provided to the right patient, in the right dose, at the right time, and via the right route. The method may include using a scanner to provide first, second, and third signals to a central computer (108) having data defining a first flow rate and first flow rate tolerance. First signal may include data identifying a medication. Second signal may include data identifying a second flow rate. Third signal may include data identifying the volume of medication in a medication container (124). Central computer (108) authorizes the second flow rate if the second flow rate is within the first flow rate tolerance. An infusion pump (120) receives new operating parameters to implement the second flow rate if the central computer (108) authorizes the second flow rate and the central computer (108) documents initiation of the second flow rate using a time source.
INFUSION THERAPY FLOW RATE ADJUSTMENT
SYSTEM AND METHOD

DESCRIPTION

Related Applications


The present application also claims the benefit of U.S. Provision Application Serial No. 60/376,655, entitled “Infusion Therapy System And Method,” filed April 30, 2002.

Technical Field

This invention relates generally to a system and method for medical therapy. More particularly, the present invention relates to a system and method for assisting in verifying that the right medication is efficiently provided to the right patient, in the right dose, at the right time, and via the right route.

Background of the Invention

Patient care systems typically include computer networks, medical devices for treating a patient, and controls for the medical devices. Although patient care systems have been improved through the use of computerized automation systems and methods, patient care systems continue to rely heavily upon manual data management processes for medical devices and controls for medical devices. For example, nursing stations are typically connected to the computer networks in modern hospitals, but it is unusual for the computer network to extend to a patient’s room. Computer networks offer the opportunity for automated data management processing including the operating and monitoring of medical devices and controls for the medical devices at the point-of-care. Despite advances in the field, automated data management technology has been underutilized for point-of-care applications due to a lack of more efficient systems and methods for operating medical devices such as infusion pumps.

Errors can be attributed to a number of things between when a clinician recognizes the need for a treatment and when the treatment is administered to a patient. Traditionally, paper medical administrative records (MARs) have been used to coordinate the treatment decision process and the resulting treatment. However, creating and using paper MARs is a process that
is prone to errors. Paper MARs are generally not verified against system-wide treatment standards. Every clinician may create a MAR in a slightly different manner. Variability in the creation of MARs leads to errors in interpretation of the MARs. Different clinicians may not be aware of what other clinicians are doing in regard to the treatment of the patient. Ultimately, paper MARs result in errors in the treatment administered to patients. One place where these errors are particularly dangerous is in the administration of medical treatment involving medications. It would be beneficial to have an improved system for creating and using MARs to administer medical treatment.

Summary of the Invention

The present invention provides a system and method for assisting in verifying that the right medication is efficiently provided to the right patient, in the right dose, at the right time, and via the right route. The invention also relates to efficiently coordinating infusion therapy with patient care system billing and inventory subsystems.

A first embodiment implemented as a computer program, includes logic for: using a bar code scanner to provide a first signal to a first computer, the first signal including data identifying the medication, the first computer having data defining a first flow rate, the first computer having data defining a first flow rate tolerance, the first computer using a central time source; using the bar code scanner to provide a second signal to the first computer, the second signal including data identifying a second flow rate; using the bar code scanner to provide a third signal to the first computer, the third signal including data identifying the volume of medication in the medication container, where the first computer authorizes the second flow rate if the second flow rate is within the first flow rate tolerance, where the infusion pump receives new operating parameters to implement the second flow rate if the first computer authorizes the second flow rate, and where the first computer documents the initiation of the second flow rate using the central time source.

A second embodiment may be implemented as a method for administering a medication with an infusion pump, the medication being packaged in a plurality of medication containers, the method comprising the steps of: providing a first signal to a first computer, the first signal including data identifying a second flow rate, where the first computer has data defining a first infusion order, the infusion order including a first flow rate, where the plurality of medication containers are prepared according to a first preparation schedule, where the infusion pump receives new operating parameters to implement the second flow rate if the first computer
authorizes the second flow rate; and providing a second signal to the first computer, the second signal triggering a revision of the preparation schedule based on the second flow rate if the first computer authorizes the second flow rate.

A third embodiment may be implemented as a system for creating infusion orders, the system comprising: a first computer screen, the first computer screen offering a plurality of main infusion order types, the main infusion order types including a single dose infusion, a continuous infusion, a sequencing infusion, and an alternating infusion, where the selection of the continuous infusion allows defining of a titrating dose; a second computer screen, where the second computer screen is provided after a main infusion order type is identified in the first computer screen, the second computer screen designed to offer an infusion order subtype, the infusion order subtype being one of the group of infusion order subtypes consisting of TPN, chemotherapy, piggyback, and large volume parental.

Other systems, methods, features, and advantages of the present invention will be, or will become, apparent to one having ordinary skill in the art upon examination of the following drawings and detailed description. It is intended that all such additional systems, methods, features, and advantages included within this description, be within the scope of the present invention, and be protected by the accompanying claims.

Brief Description of the Drawings

The invention can be better understood with reference to the following drawings. The components in the drawings are not necessarily to scale, emphasis instead being placed upon clearly illustrating the principles of the present invention. In the drawings, like reference numerals designate corresponding parts throughout the several views.

FIG. 1 is a graphical representation of a patient care system. The patient care system includes a pharmacy computer, a central system, and a digital assistant at a treatment location;

FIG. 2 is a block diagram of a computer system representative of the pharmacy computer, the central system, and/or the digital assistant of FIG. 1. The system includes an infusion system or a portion of the infusion system;

FIG. 3 is a block diagram showing functional components of the patient care system of FIG. 1.

FIG. 4 is an exemplar computer screen for implementing various functions of the patient care system of FIG. 1;
FIG. 5 is a block diagram showing functional components of the infusion system of FIG. 2. The functional components include blocks for setting infusion system parameters, infusion order creation, infusion order preparation, medication administration, infusion order modifications, and messaging;

FIG. 6 is a block diagram showing functional components for the setting of infusion system parameters of FIG. 5;

FIG. 7 is a block diagram showing functional components for the infusion order creation of FIG. 5;

FIG. 8 is a block diagram showing functional components for the infusion order preparation of FIG. 5;

FIG. 9 is a block diagram showing functional components for the medication administration of FIG. 5; and,

FIG. 10 is a block diagram showing functional components for infusion order documentation 1012, and the infusion order modifications 514 and messaging 520 of FIG. 5.

Detailed Description

While this invention is susceptible of embodiments in many different forms, there is shown in the drawings and will herein be described in detail a preferred embodiment of the invention. The present disclosure is to be considered as an exemplification of the principles of the invention and is not intended to limit the broad aspect of the invention to the embodiment illustrated.

FIG. 1 is a graphical representation of a patient care system 100. Patient care system 100 includes a pharmacy computer 104, a central system 108, and a treatment location 106, linked by a network 102. Patient care system 100 also includes infusion system 210 (FIG. 2). Infusion system 210 is a medication system preferably implemented as a computer program, and in particular an application (i.e., a program or group of programs designed for end users), resident on one or more electronic computing devices within the patient care system 100. As described in detail further herein, the infusion system 210 links clinicians, such as physicians, pharmacists, and nurses, in an interdisciplinary approach to patient care.

Patient care system 100 preferably includes a computerized physician order-entry module (CPOE), an inpatient pharmacy module, a wireless nurse charting system, and an electronic patient medical record. It is desired that patient care system 100 provide a comprehensive patient safety solution for the delivery of medication. Within patient care system 100, software
modules are provided to link together existing patient care systems using interfaces such as HL7 interfaces that are known to those having ordinary skill in the art. Patient care system 100 can operate on a variety of computers and personal digital-assistant products to transmit orders and update patient medical records.

The CPOE enables physicians to enter medication orders, review alerts, reminders, vital signs and results. A pharmacy module checks the prescribed drug against documented patient allergies, and for compatibility with other drugs and food. The pharmacy module also provides real-time data for inventory management. A nurse medication-charting module provides clinical information that is immediately available at the bedside, thus ensuring verification of medication and dosage at the point-of-care.

Patient care system 100 integrates drug delivery products with the information required to ensure safe and effective delivery of medication. The clinical decision supports and accompanying alerts and warnings of the patient care system 100 provide a safety net of support for clinicians as they deliver patient care under increasing time and cost pressures. This information may be supplied through a wireless network that supplies data in a way that improves clinician workflow, making delivery of care easier.

Infusion system 210 provides computerized prescribing and an electronic medical administration record (eMAR). Infusion system 210 puts charting, medication history, and inventory tracking at the clinician's fingertips. Patient care system 100 combines bar-coding and real-time technology to assist in ensuring that the right patient gets the right medication and the right dosage, at the right time, via the right route. Infusion system 210 provides alerts and reminders such as, but not limited to, lab value, out of range, and missed dose.

Patient care system 100 allows medication ordering, dispensing, and administration to take place at the patient's bedside. Physicians can order simple and complex prescriptions, intravenous therapy and total parental nutrition therapy (TPN) using a wireless handheld device. Infusion system 210 checks for drug interactions and other possible errors as well as correct dosage. Infusion system 210 then transmits this data in real-time to the patient care facility or local pharmacy, hospital nursing unit, home care unit, and/or clinic.

The clinician may access a medical records database using a handheld scanning device. The clinician may scan the bar coded medication and the patient's bar coded bracelet to confirm the presence of the right medication, dosage, and time before administering any drugs. Infusion system 210 updates medical and administrative records, thereby eliminating time-consuming
paperwork. Thus infusion system 210 reduces costs and improves efficiency while saving lives. Patient care system 100 may include access-controlled mobile and stationary medication and supply depots, including electronic patient medical records and computerized prescribing, providing complete preparation and inventory management from the point of care to the pharmacy.

As mentioned previously, FIG. 1 is a graphical representation of patient care system 100. The patient care system 100 includes a pharmacy computer 104, a central system 108, and a treatment location 106, linked by a network 102. The pharmacy computer 104 may include a processing unit 104a, a keyboard 104b, a video display 104c, a printer 104d, a bar code reader 104e, and a mouse 104f. Although not shown in FIG. 1, the patient care system 100 may also include subsystems for hospital administration, nursing stations, a clinical information subsystem, a hospital information subsystem, an Admissions Discharge and Transfer (ADT) subsystem, a billing subsystem, and/or other subsystems typically included in patient care systems.

The central system 108 may include a central servicing unit 108a, a database 108b, a video display 108c, input/output components, and many other components known to those having ordinary skill in the art. The network 102 includes a cable communication system 110 portion and a wireless communication system portion. The cable communication system 110 may be, but is not limited to, an Ethernet cabling system, and a thin net system.

The treatment location 106 may include a treatment bed 106a, an infusion pump 120, and medical treatment cart 132. In FIG. 1, a clinician 116 and a patient 112 are shown in the treatment location 106. Medication 124 may be of a type that may be administered using an infusion pump 120. Medication 124 may also be of a type that is administered without using an infusion pump. The medication may be stored in medication storage areas 132a of medical treatment cart 132. The clinician 116 uses a digital assistant 118 to administer medication 124 to the patient 112.

In the course of treating patient 112, the clinician 116 may use the digital assistant 118 to communicate with the cable communication system 110 of the network 102 via a first wireless communication path 126. The infusion pump 120 may also have the ability to communicate with the cable communication system 110 via a second wireless communication path 128. The medication cart 124 may also have the ability to communicate via a wireless communication path (not shown in FIG. 1). A wireless transceiver 114 interfaces with the cable
communication system 110. The wireless communication system portion of the network may employ technology such as, but not limited to, that known to those having ordinary skill in the art as IEEE 802.11b “Wireless Ethernet,” a local area network, wireless local area networks, a network having a tree topography, a network having a ring topography, wireless internet point of presence systems, an Ethernet, the Internet, radio communications, infrared, fiber optic, and telephone. Though shown in FIG. 1 as a wireless communication system, communication paths may be hardwired communication paths.

In the patient care system 100, a physician may order medication 124 for patient 112. The order may also originate with a clinician 116 at the treatment location 106. The physician and/or clinician 116 may use a computerized physician order entry system (CPOE) and/or the medical cart 132 to order the medication 124 for the patient 112. Those having ordinary skill in the art are familiar with basic CPOEs. Despite its name, any clinician 116 may use the CPOE. If the medication 124 is one that is efficient to administer through infusion pump 120, the infusion order includes information for generating operating parameters for the infusion pump 120. The operating parameters are the information and/or instruction set that is necessary to program infusion pump 120 to operate in accordance with the infusion order.

The infusion order may be entered in a variety of locations including the pharmacy, the nursing center, the nursing floor, and treatment location 106. When the order is entered in the pharmacy, it may be entered in the pharmacy computer 104 via input/output devices such as the keyboard 104b, the mouse 104f, a touch screen display, the CPOE system and/or the medical treatment cart 132. Those having ordinary skill in the art are familiar with these and similar input/output devices. The processing unit 104a is able to transform a manually-entered order into computer readable data. Devices such as the CPOE may transform an order into computer readable data prior to introduction to the processing unit 104a. The operating parameters may then be printed in a bar code format by the printer 104d on a medication label 124a. The medication label 124a may then be affixed to a medication 124 container. The medication 124 container is then transported to the treatment location 106. The medication 124 may then be administered to the patient 112 in a variety of ways known in the art including orally and through an infusion pump 120. If the medication 124 is administered orally, the clinician 116 may communicate via the digital assistant 118 and/or the medical cart 132. The medical cart 132 is computerized and generally has a keyboard (not shown), a display 132b, and other input/output devices such as a bar code scanner (not shown).
At the treatment location, the medication 124 may be mounted on the infusion pump 120 and an intravenous (IV) line 130 may be run from the infusion pump 120 to the patient 112. The infusion pump 120 may include a pumping unit 120a, a keypad 120b, a display 120c, an infusion pump ID 120d, and an antenna 120e. Prior art infusion pumps may be provided with a wireless adaptor (not shown) in order to fully implement the system 100. The wireless adaptor may have its own battery if necessary to avoid reducing the battery life of prior art infusion pumps. The wireless adaptor may also use intelligent data management such as, but not limited to, store-and-forward data management and data compression to minimize power consumption. The wireless adaptor may also include the ability to communicate with the digital assistant 118 even when the network 102 is not functioning.

The patient care system 100 may include a variety of identifiers such as, but not limited to, personnel, equipment, and medication identifiers. In FIG. 1, the clinician 116 may have a clinician badge 116a identifier, the patient 112 may have a wristband 112a identifier, the infusion pump 120 may have an infusion pump ID 120d identifier, and the medication 124 may have a medication label 124a identifier. Clinician badge 116a, wristband 112a, infusion pump ID 120d, and medication label 124a include information to identify the personnel, equipment, or medication they are associated with. The identifiers may also have additional information. For example, the medication label 124a may include information regarding the intended recipient of the medication 124, operating parameters for infusion pump 120, and information regarding the lot number and expiration of medication 124. The information included in the identifiers may be printed, but is preferably in a device readable format such as, but not limited to, an optical readable device format such as a bar code, a radio frequency (RF) device readable format such as an RFID, an iButton, a smart card, and a laser readable format. The digital assistant 118 may include a display 118a and may have the ability to read the identifiers including biometric information such as a fingerprint.

The wristband 112a is typically placed on the patient 112 as the patient 112 enters a medical care facility. The wristband 112a includes a patient identifier. The patient identifier may include printed information to identify the patient and additional information such as a treating physician's name(s). The patient identifier for patient 112 may include information such as, but not limited to, the patient's name, age, social security number, the patient's blood type, address, allergies, a hospital ID number, and the name of a patient's relative.

FIG. 2 is a block diagram of a computer 200. Computer 200 may be the pharmacy
computer 104, the central system 108, a CPOE, the digital assistant 118 of FIG. 1, and/or a computer included in any number of other subsystems that communicate via the network 102 such as the medication treatment cart 132. Computer 200 includes an infusion system 210, or a portion of infusion system 210. The invention is described in reference to FIG. 2 as a computer program. However, the invention may be practiced in whole or in part as a method and system other than as a computer program.

A critical concern in the art is that the right medication is administered to the right patient. Therefore, infusion system 210 includes features to assist in assuring that the right medication is administered to the right patient in an efficient manner. Infusion system 210 can be implemented in software, firmware, hardware, or a combination thereof. In one mode, infusion system 210 is implemented in software, as an executable program, and is executed by one or more special or general purpose digital computer(s), such as a personal computer (PC; IBM-compatible, Apple-compatible, or otherwise), personal digital assistant, workstation, minicomputer, or mainframe computer. An example of a general-purpose computer that can implement the infusion system 210 of the present invention is shown in FIG. 2. The infusion system 210 may reside in, or have portions residing in, any computer such as, but not limited to, pharmacy computer 104, central system 108, medication treatment cart 132, and digital assistant 118. Therefore, computer 200 of FIG. 2 may be representative of any computer in which the infusion system 210 resides or partially resides.

Generally, in terms of hardware architecture, as shown in FIG. 2, the computer 200 includes a processor 202, memory 204, and one or more input and/or output (I/O) devices 206 (or peripherals) that are communicatively coupled via a local interface 208. The local interface 208 can be, for example, but not limited to, one or more buses or other wired or wireless connections, as is known in the art. The local interface 208 may have additional elements, which are omitted for simplicity, such as controllers, buffers (caches), drivers, repeaters, and receivers, to enable communications. Further, the local interface may include address, control, and/or data connections to enable appropriate communications among the other computer components.

Processor 202 is a hardware device for executing software, particularly software stored in memory 204. Processor 202 can be any custom made or commercially available processor, a central processing unit (CPU), an auxiliary processor among several processors associated with the computer 200, a semiconductor-based microprocessor (in the form of a microchip or chip
set), a macroprocessor, or generally any device for executing software instructions. Examples of suitable commercially available microprocessors are as follows: a PA-RISC series microprocessor from Hewlett-Packard Company, an 80x86 or Pentium series microprocessor from Intel Corporation, a PowerPC microprocessor from IBM, a Sparc microprocessor from Sun Microsystems, Inc., or a 68xxx series microprocessor from Motorola Corporation. Processor 202 may also represent a distributed processing architecture such as, but not limited to, SQL, Smalltalk, APL, KLisp, Snobol, Developer 200, MUMPS/Magic.

Memory 204 can include any one or a combination of volatile memory elements (e.g., random access memory (RAM, such as DRAM, SRAM, SDRAM, etc.)) and nonvolatile memory elements (e.g., ROM, hard drive, tape, CDROM, etc.). Moreover, memory 204 may incorporate electronic, magnetic, optical, and/or other types of storage media. Memory 204 can have a distributed architecture where various components are situated remote from one another, but are still accessed by processor 202.

The software in memory 204 may include one or more separate programs. The separate programs comprise ordered listings of executable instructions for implementing logical functions. In the example of FIG. 2, the software in memory 204 includes the infusion system 210 in accordance with the present invention and a suitable operating system (O/S) 212. A non-exhaustive list of examples of suitable commercially available operating systems 212 is as follows: (a) a Windows operating system available from Microsoft Corporation; (b) a Netware operating system available from Novell, Inc.; (c) a Macintosh operating system available from Apple Computer, Inc.; (d) a UNIX operating system, which is available for purchase from many vendors, such as the Hewlett-Packard Company, Sun Microsystems, Inc., and AT&T Corporation; (e) a LINUX operating system, which is freeware that is readily available on the Internet; (f) a run time Vxworks operating system from WindRiver Systems, Inc.; or (g) an appliance-based operating system, such as that implemented in handheld computers or personal digital assistants (PDAs) (e.g., PalmOS available from Palm Computing, Inc., and Windows CE available from Microsoft Corporation). Operating system 212 essentially controls the execution of other computer programs, such as infusion system 210, and provides scheduling, input-output control, file and data management, memory management, and communication control and related services.

Infusion system 210 may be a source program, executable program (object code), script, or any other entity comprising a set of instructions to be performed. When a source program,
the program needs to be translated via a compiler, assembler, interpreter, or the like, which may or may not be included within the memory 204, so as to operate properly in connection with the O/S 212. Furthermore, the infusion system 210 can be written as (a) an object-oriented programming language, which has classes of data and methods, or (b) a procedural programming language, which has routines, subroutines, and/or functions, for example, but not limited to, C, C++, Pascal, Basic, Fortran, Cobol, Perl, Java, and Ada. In one embodiment, the system program 210 is written in C++. In other embodiments, the infusion system 210 is created using Power Builder. The I/O devices 206 may include input devices, for example, but not limited to, a keyboard, mouse, scanner, microphone, touch screens, interfaces for various medical devices, bar code readers, stylus, laser readers, radio-frequency device readers, etc. Furthermore, the I/O devices 206 may also include output devices, for example, but not limited to, a printer, bar code printers, displays, etc. Finally, the I/O devices 206 may further include devices that communicate both inputs and outputs, for instance, but not limited to, a modulator/demodulator (modem; for accessing another device, system, or network), a radio frequency (RF) or other transceiver, a telephonic interface, a bridge, a router, etc.

If the computer 200 is a PC, workstation, PDA, or the like, the software in the memory 204 may further include a basic input output system (BIOS) (not shown in FIG. 2). The BIOS is a set of essential software routines that initialize and test hardware at startup, start the O/S 212, and support the transfer of data among the hardware devices. The BIOS is stored in ROM so that the BIOS can be executed when computer 200 is activated.

When computer 200 is in operation, processor 202 is configured to execute software stored within memory 204, to communicate data to and from memory 204, and to generally control operations of computer 200 pursuant to the software. The infusion system 210 and the O/S 212, in whole or in part, but typically the latter, are read by processor 202, perhaps buffered within the processor 202, and then executed.

When the infusion system 210 is implemented in software, as is shown in FIG. 2, it should be noted that the infusion system 210 program can be stored on any computer readable medium for use by or in connection with any computer related system or method. In the context of this document, a computer readable medium is an electronic, magnetic, optical, or other physical device or means that can contain or store a computer program for use by or in connection with a computer related system or method. The infusion system 210 can be embodied in any computer-readable medium for use by or in connection with an instruction
execution system, apparatus, or device, such as a computer-based system, processor-containing system, or other system that can fetch the instructions from the instruction execution system, apparatus, or device and execute the instructions. In the context of this document, a “computer-readable medium” can be any means that can store, communicate, propagate, or transport the program for use by or in connection with the instruction execution system, apparatus, or device. The computer readable medium can be, for example, but not limited to, an electronic, magnetic, optical, electromagnetic, infrared, or semiconductor system, apparatus, device, or propagation medium. More specific examples (a non-exhaustive list) of the computer-readable medium would include the following: an electrical connection (electronic) having one or more wires, a portable computer diskette (magnetic), a random access memory (RAM) (electronic), a read-only memory (ROM) (electronic), an erasable programmable read-only memory (EPROM, EEPROM, or Flash memory) (electronic), an optical fiber (optical), and a portable compact disc read-only memory (CDROM) (optical). Note that the computer-readable medium could even be paper or another suitable medium upon which the program is printed, as the program can be electronically captured, via, for instance, optical scanning of the paper or other medium, then compiled, interpreted or otherwise processed in a suitable manner if necessary, and then stored in a computer memory.

In another embodiment, where the infusion system 210 is implemented in hardware, the infusion system 210 can be implemented with any, or a combination of, the following technologies, which are each well known in the art: a discrete logic circuit(s) having logic gates for implementing logic functions upon data signals, an application specific integrated circuit (ASIC) having appropriate combinational logic gates, a programmable gate array(s) (PGA), a field programmable gate array (FPGA), etc.

Any process descriptions or blocks in figures, such as FIGS. 4-10, should be understood as representing modules, segments, or portions of code which include one or more executable instructions for implementing specific logical functions or steps in the process, and alternate implementations are included within the scope of the embodiments of the present invention in which functions may be executed out of order from that shown or discussed, including substantially concurrently or in reverse order, depending on the functionality involved, as would be understood by those having ordinary skill in the art.

FIG. 3 is a first block diagram 300 showing functional components of the patient care system 100 of FIG. 1. The patient care system 100 may be practiced as a modular system where
the modules represent various functions of the patient care system, including the infusion system. The flexibility of the patient care system and the infusion system may be enhanced when the systems are practiced as modular systems. The modules of the infusion system 210 may be included in various portions of the patient care system 100. The patient care system 100 includes a medication management module 302, a prescription generation module 304, a prescription activation module 306, and a prescription authorization module 308.

The medication management module 302 may coordinate the functions of the other modules in the patient care system 100 that are involved in the administration of medical treatment. The medication management module 302 will generally coordinate with other portions of the patient care system 100. The medication module 302 may include sub-modules for operating and/or interfacing with a CPOE, for operating and/or communicating with point-of-care modules, and for operating and/or communicating with medical treatment comparison modules. In FIG. 3, an admissions, discharge, and transfer (ADT) interface 310, a billing interface 312, a lab interface 314, and a pharmacy interface 316 are shown. ADT interface 310 may be used to capture information such as the patient’s size, weight, and allergies. Pharmacy interface 316 imports orders from the pharmacy. The pharmacy interface 316 may be an HL7 type of interface that interfaces with other systems for entering orders, such as a CPOE. This ability reduces the necessity for entering data into the patient care system 100 more than once. The pharmacy interface 316 may be configured to communicate with commercially available systems such as, but not limited to Cerner, HBOC, Meditech, SMS, and Phamous. Various other interfaces are also known to those having ordinary skill in the art but are not shown in FIG. 3.

The medication management module 302 may have additional features such as the ability to check for adverse reactions due to drug-to-drug incompatibility, duplicate drug administration, drug allergies, drug dosage limitations, drug frequency limitations, drug duration limitations, and drug disease contraindications. Food and alcohol interactions may also be noted. Drug limitations may include limitations such as, but not limited to, limitations associated with adults, children, infants, newborns, premature births, geriatric adults, age groupings, weight groupings, height groupings, and body surface area. Generally, the medication management module 302 will also prevent the entry of the same prescription for the same patient from two different sources within the patient care system 100.
The medication management module 302 may also include the ability to generate reports. The reports include, but are not limited to, end-of-shift, titration information, patient event lists, infusion history, pump performance history, pump location history, and pump maintenance history. The end-of-shift report may include the pump channel, start time, end time, primary infusion, piggyback infusion, medication, dose, rate, pump status, volume infused, volume remaining, time remaining, and the last time cleared. The infusion history report includes medications and volume infused.

The medication management module 302 may also include a medical equipment status database. The medical equipment status database includes data indicating the location of a medical device 332 within the patient care system 100. The medical equipment status database may also include data indicating the past performance of a medical device 332. The medical equipment status database may also include data indicating the maintenance schedule and/or history of a medical device 332.

Infusion prescriptions are entered in prescription entry 324. Prescriptions may include prescriptions such as, but not limited to, single dose infusions, intermittent infusions, continuous infusions, sequencing, titrating, and alternating types. Infusion prescriptions may also include total parenteral nutritional admixtures (TPN), chemotherapy continuous infusion, piggybacks, large volume parenterals, and other infusion prescriptions. The patient care system 100 is capable of functioning without end dates for orders. The patient care system 100 may use a continuous schedule generator that looks ahead a predefined time period and generates a schedule for admixture filling for the time period. The predefined time period may be defined at the patient care system 100 level or at subsystem levels such as the clinical discipline level and an organizational level. The predefined time periods may be adjustable by the clinician entering the order. The schedule may be automatically extendable as long as the order is active in the patient care system 100.

The prescription generation module 304 generates hard prescriptions and electronic (E-copy) prescriptions. Hard prescriptions are generally produced in triplicate in medical facilities. A first hard copy 318 is generally sent to the pharmacy, a second hard copy 320 is generally kept for the patient’s records, and third hard copy 322 is sent to treatment location 106. An electronic prescription is sent to the medication management module 302.

Prescription generation 304 may include confirming operating parameters. The operating parameters may be based on information from prescription entry module 324.
Prescription generation 304 may occur anywhere in the patient care system 100 such as, but not limited to, the pharmacy, the treatment location 106, and a nursing center.

A computerized physician order entry (CPOE) system may be employed to carry out some or all of the functions of the prescription generation module 304. Clinicians 116 may enter data in a variety of manners such as, but not limited to, using a tablet wireless computer, treatment cart 132, and a workstation. The medication management module 302 may interface with more than one prescription generation module 304. The medication management module may receive orders from anywhere within the patient care system 100.

The pharmacy computer 104 is able to access the electronic copy from the medication management module 302. The prescription activation module 306 is a computer-assisted system for coordinating the filling and labeling of prescriptions. The filling of the prescription and the creation or location of medication 124 from stock is handled by the prescription activation module 306.

The patient care system 100 may bypass the prescription activation module 306. This may occur if the ordering clinician 116, such as the patient’s physician, has the authority to immediately activate an order. If the order is immediately activated, the medication management module 302 may go directly to prescription labeling module 326.

In block 326, the patient care system 100 prints the medication label 124. The prescription may be printed remotely and will often be printed by the pharmacy printer 104d. After block 326, the patient care system goes to block 328. In block 328, the medication label 124a is attached to the medication 124. The pharmacist generally provides a visual verification 334 that the medication label 124a matches the first hard copy 318 of the prescription. FIG. 3 shows that a visual verification 334 is also associated with prescription authorization module 308. The medication 124 may then be transported from the pharmacy to the treatment location 106. A portable medical treatment cart 132 may be used for a portion of the route from the pharmacy to the treatment location 106.

The medication label 124a may include information for preparing the infusion bag. If not generated within patient care system 100, medication label 124a may be provided by a bulk medication supplier. If provided by a bulk medication supplier, the patient care system 100 has the capability of gathering the information from the medication label 124a. In addition, the patient care system 100 has the ability to add information, such as a patient identifier, to medication label 124a.
The medication labeling module 328 places the medication label 124 on the medication 124. This may be accomplished manually. This may also be accomplished using an automatic prescription filling and packaging system (not shown). If an automatic filling and packaging system is used, medication labeling module 328 provides data for coordination of the labeling of the medication 124 to the filling and packaging system.

At the treatment location 106, the clinician 116 uses a wireless device 330, such as digital assistant 118 and/or medical treatment cart 132, to verify and administer medication 124 to the patient 112. Wireless device 330 communicates with the medication management module 302 via a communication path, such as first communication path 126.

Clinician 116 generally identifies his/herself by scanning badge 116a, identifies the patient 112 by scanning wristband 112a, identifies the medication 124 by scanning medication label 124a, and identifies the medical device 332, such as infusion pump 120, by scanning label 120d. Clinician 116 may also identify his/herself by providing a fingerprint and/or password. The medical device 332 may be a medical device capable of two-way communication with the medication management module 302. Alternatively, the medical device 332 may only be capable of providing information to the medication management module 302. The infusion program 210 assists the clinician 116 in administering and verifying the medical treatment. The infusion program 210 may include downloading of operating parameters to the medical device 332. Clinician 116 may provide a visual verification to confirm the third copy 322 and/or the MAR matches the labeled medication 124. Scanner 338 may be used to enter machine readable information from the third copy 322 to the wireless device 330 and the medical device 332.

The patient care system 100 includes the ability to make adjustments and modifications to infusion orders. Among other modules that may include the ability to make infusion adjustments are prescription entry 324, prescription activation 306, prescription authorization 308, and prescription modification module 336. Clinician 116 may access prescription modification module 336 in order to make adjustments to an order. The clinician 116 may access the prescription modification module 336 throughout the patient care system 100. However, one very useful location for clinician 116 to access the prescription modification module 336 is at treatment location 106.

In prescription authorization module 308, the patient care system 100 determines whether the clinician 116 has the authority to independently modify an infusion order. The clinician 116 may be recognized by the patient care system 100 as having the authority to
independently modify certain portions of the order. If the clinician 116 does not have the authority to independently modify the order, a pharmacist or physician may be requested to approve the modification entered by the clinician 116.

In one implementation of patient care system 100, an order is entered in pharmacy computer 104. The order includes a first patient identifier and an operating parameter. The pharmacy computer 104 generates a medication label 124a that is affixed to medication 124. The medication 124 is sent to a treatment location 106. At treatment location 106, clinician 116 reads the clinician’s badge 116a, patient’s wristband 112a, and medication label 124a with a digital assistant 118. The digital assistant 118 determines whether medication label 124a and wristband 112a identify the same patient 112. The system 400 then sends the medication identifier to the pharmacy computer 104. The pharmacy computer 104 confirms the medication label 124a identifies the same patient as the order and sends the operating parameter to an infusion pump. The operating parameter may be sent directly to the infusion pump 120. The operating parameter is then used to program the infusion pump to administer the medication 124 to the patient 112.

FIG. 4 is an exemplar computer screen 400 that is useful in implementing various functions of the infusion system 210. In addition to other functions, computer screen 400 may be used to enter new infusion orders, to modify existing infusion orders, and to stop infusion orders. Computer screen 400 includes a processing area 402, search areas 404, a medication information area 406, a titration/tapering criteria area 408, an instruction and note area 410, and a projected solution ingredient area 412. Infusion medication order types include single dose, intermittent, continuous, sequencing, and alternating. Computer screen 400 may be used with digital assistant 118, pharmacy computer 104, infusion pump 120, a CPOE system, and medical treatment cart 132. Computer screen 400 will generally be designed to have the look-and-feel of clinician 116 accessible computer screens throughout the patient care system 100. The functions of computer screen 400 are partially accomplished with database linkage techniques that are familiar to those having ordinary skill in the art such as, but not limited to, hyperlinks, definition boxes, and dropdown menus.

The processing area 402 may include the ability to trigger the creation of an infusion order, a save of an infusion order, and a cancellation of an infusion order. Clinician 116 may customize computer screen 400 to provide the clinician’s 116 preferred order entry procedures. The processing area 402 includes a status indicator for orders. The processing area 402 includes
an area for indicating whether a PRN order (a "when necessary" order) may be placed by clinician 116. The processing area 402 also includes the ability to display and adjust medical device 332 operating parameters, infusion order route, infusion line, infusion administration site, infusion order start time, infusion medication order type, infusion flow rate tolerance, infusion flow rate, infusion duration, and area of preparation (such as pharmacy or a remote site). The processing area 402 may also include an area for linking medical orders to other medical orders such as, linking a physician's infusion order to another medical order that may be entered by another clinician 116. The processing area 402 may include a trigger for displaying data in other areas of the computer screen 400 such as, but not limited to, the projected solutions area 412.

Search areas 404 allow for searching for medications, solutions and/or additives for infusion orders. Default diluents may be provided for orders. If a default dosage for a medication is defined in the patient care system 100, the default dosage may automatically appear with the search result that includes the medication. A search from search area 404 will generally produce the medication name, the route of administration, the cost, the package size, the dosage form, the generic name, whether the medication is a narcotic, whether the medication is controlled, whether formulary, and whether the medication is manufactured.

Medication information area 406 may be used to define infusion order additives and solutions. Medication information area 406 may include separate additive areas and solution areas. The solution area may include a label "Solution/Diluent". The patient care system 100 may use a medication 124 database, a solutions database, and an additive database to populate the medication information area 406 with medications 124, solutions, and additives. Substances identified in one database may also be identified in other databases. The databases may be linked to provide default values for combinations of the medications 124 and solutions.

Titration/tapering criteria area 408 generally applies to continuous infusion orders. Titration defines certain parameters of an order such as dosage and/or flow rate. Dose and flow rate can be entered as an absolute. Also, mathematical symbols such as, but not limited to, greater than ">", less than "<", and equal "=" may be used alone or in combination to enter information in titration/tapering criteria area 408. A calendar may also be used to enter data in titration/tapering criteria area 408. Dosage and flow rate can also be entered as an acceptable range. Titration/tapering criteria area 408 may be hidden when non-continuous infusion orders are entered and/or modified.
Instruction and note area 410 includes the ability to save information such as physician notes regarding a patient 112 and/or an infusion order. The instruction and note area 410 may include a display and lookup area for identifying clinicians 116 that are responsible for the patient 112, such as the patient’s physician.

The projected solutions area 412 displays solution schedules and related ingredients based on the current state of the order being processed for patient 112. The time period projected may be a patient care system 100 default. The time period may also be adjustable by the clinician 116. The projected solutions area 412 may include an adjustable display indicating the time period projected by the patient care system 100. The data displayed in the projected solutions area will generally be saved when an order save is triggered in the processing area 402. The projected solutions area 412 may include the ability to look back over a period of time while modifying a previously entered order. This allows the clinician 116 to view solutions that may have already been prepared according to the unmodified infusion order.

FIG. 5 is a block diagram showing functional components of the infusion system 210 of FIG. 2. The functional components include blocks for setting system parameters 502, infusion order creation 504, infusion order preparation 506, medication administration 512, infusion order modifications 514, and messaging 520. Fig. 5 also includes blocks for pharmacy authorization 508, physician authorization 510, stop orders 516, and inventory and billing 518. FIG. 5 presents one description of the infusion system. However, FIG. 5 does not define a required series of steps for implementing the infusion system. One of the benefits of the infusion system is that clinician’s 116 may access and enter information from a large number of locations, both physical and functional, within the patient care system 100. For example, an infusion order may be created by a physician using a CPOE, by a pharmacist using pharmacy computer 106, by a clinician 116 using digital assistant 118, and by a clinician using medication treatment cart 132.

FIG. 5 may be viewed as first preparing the patient care system 100 for receiving infusion orders – setting system parameters 502; second, creating the infusion order – infusion order creation 504; third, preparing the infusion order – preparation 506; fourth, authorizing the infusion order – pharmacy and physician authorization 508 and 510; fifth, administering the infusion order – medication administration 512; sixth, accounting for the inventory used to prepare the infusion order and billing the patient for the infusion order – inventory and billing 518; seventh, modifying the infusion order – modifications 514; and eight, providing messages
to various personnel and sub-systems regarding the progress of the infusion order – messages
520. Modifications 514 may include stopping the order – stop order 516 – based on information
provided by the ADT interface 310.

Setting system parameters 502 include functional blocks that prepare the infusion system
210 to create and process infusion orders. Setting system parameters 502 includes, but is not
limited to, setting tolerances 542, setting defaults 544, building databases 546, defining
functions 548, and determining system settings 550. Setting system parameters 502 is further
described below in reference to FIG. 6.

Infusion order creation 504 includes functional blocks used to create infusion orders.
Infusion order creation 504 includes functions similar to those described in reference to
prescription generation 304 (FIG. 3). Infusion order creation 504 includes, but is not limited to,
entering information 560, calculations 562, checks 564, and overrides 568. Infusion order
creation is further described below in reference to FIG. 7. The result of infusion order creation
is an infusion order 702 (FIG. 7). Infusion order 702 generally includes an infusion schedule
704 (FIG. 7).

Infusion orders may require authorization as described in reference to block 308 (FIG.
3). In FIG. 5, prescription authorization by the pharmacist and prescription authorization by the
physician are considered separately in functional blocks for pharmacy authorization 508 and
physician authorization 510. Physician authorization 510 is generally not required if the
infusion order is initiated by the physician. The infusion order generally requires pharmacy
authorization 508 and physician authorization 512 if the order is generated by a clinician at the
treatment location 106, other than the pharmacist or physician. However, if medication 124 is
required immediately, the infusion system 210 may allow administering clinicians to bypass
prescription authorization 510 and physician authorization 512. In the case of emergency orders
or non-emergency orders for routine medications, the infusion system 210 may determine there
is no information stored in the patient care system 100 related to the medical treatment the
clinician 116 desires to administer to the patient 112. If the infusion system 100 recognizes the
clinician 116 as having the authority to initiate the desired medical treatment, the system 210
may allow for the administration of the medical treatment without going to blocks 508 and 510.

Infusion order preparation 506 may be accomplished in a number of locations throughout
the medical facility such as, but not limited to, the pharmacy, the nursing center, on the floor,
and the treatment location 106. Preparation 506 includes providing instructions for preparing the medication 124 and minimizing the possibility of errors in medication preparation.

Medication administration 512 takes place at the treatment location 106. The infusion system 210 is designed to make the administration of the order as efficient and accurate as possible. The infusion system 210 provides the administrating clinician with the tools to assist in administering the right medication to the right patient in the right dose at the right time, and via the right route.

Infusion orders are frequently modified. Infusion system 210 provides modifications 514 to account for infusion order modifications. Modification 514 includes modifications to infusion duration, flow rate, infusion site, and stop orders 516. Modification 514 also includes the functional blocks required to implement infusion order modifications.

The infusion system 210 can include patient care system 100 wide defined stop orders 516. Changes in patient status may generate messages 520 for appropriate action. The infusion system 210 coordinates with the ADT interface 310 to automatically stop orders 516 upon discharge or death.

The system 100 includes inventory and billing module 518. Inventory and billing 518 allows the financial transactions associated with patient care to proceed with a minimum of human intervention. The completion of medication administration 512 may trigger patient billing through the billing interface 312. The billing interface may include an HL7 interface. If patients are to be charged based on completion of infusion order preparation 506, the inventory and billing system 210 includes a crediting process. The crediting process may be triggered when infusion bags are returned to the pharmacy for disposal or re-entry into the pharmacy inventory management system.

The infusion system 210 includes a messages module 520 for communicating with real and virtual entities throughout the patient care system 100. For example, when a physician enters a new order, messaging appears in the pharmacy to alert the pharmacists that an infusion order requires authorization. Likewise, when infusion orders are appropriately authorized, the clinician 116 receives messaging on digital assistant 118 to alert the clinician 116 that the infusion order should be administered according to the infusion schedule 704. Overrides 566 may generate messages 520 for the physician and/or the pharmacy. The infusion system 100 may distinguish between system-wide and sub-system overrides in determining whether it is
necessary to generate a message 520. Messaging 520 includes messages received and/or sent to the central system, the pharmacy, the physician, billing, and inventory.

The system may present clinicians 116 with personal computer display views. The personal computer display views summarize outstanding clinical problems for the clinician’s patients. The clinician 116 may quickly retrieve detailed information for the patients. The system 100 may also produce an email or page to digital assistant 118, or other communication device, when certain critical patient conditions prevail.

FIG. 5 also highlights some of the communication paths that occur in patient care system 100. The highlighted communication paths are presented for ease in describing the infusion system 210. Those having ordinary skill in the art recognize that when patient care system 100 is practiced on a network the various functional blocks may communicate with each other via the paths highlighted in FIG. 5 and via paths that are not shown in FIG. 5. Setting system parameters 502 includes communicating data related to the system parameters to infusion order creation 504, via path 522, and/or receiving data from infusion order creation 504 and providing data informing infusion order creation 504 of how the received data relates to the system parameters.

Infusion orders may be passed directly, via path 524, to infusion preparation 506. Infusion orders may also be passed to pharmacy authorization 508, via path 526 and/or to physician authorization, via path 528, before being sent to preparation 506. Path 530 highlights the delivery of the medication 124 from the preparation area to the treatment location 106. Delivery may be accomplished using medication treatment cart 132. Paths 532, 534, 536, and 538 highlight that inventory and billing 518 transactions may be tied to a variety of other functions such as, but not limited to, infusion order creation 504, preparation 506, medication administration 512, and modifications 514. Paths 572, 574, and 576 highlight that a larger number of functions and actors involved in patient care system 100 may generate and receive information via messages 520. Path 582 highlights that system defaults 544 may be created and/or modified by the pharmacist. And, path 580 highlights that information, such as infusion orders, is available to a variety of functional units throughout the system 100.

FIG. 6 is a block diagram showing functional components for the setting of system parameters 502 of FIG. 5. Setting system parameters 502 includes, but is not limited to, setting tolerances 542, setting defaults 544, building databases 546, defining functions 548, and determining system settings 550. Tolerances 542 includes tolerances such as, but not limited
to, net medication tolerances 542a, flow rate tolerances 542b, administration time tolerances 542c, administration system duration 542d, medication duration tolerances 542e, and site change tolerances 542f. The infusion system 210 may also include separate tolerances for order entry and modifications from the ordered tolerances. For example, separate tolerances may be identified such as, but not limited to, an administration system duration 542d, an order entry maximum infusion duration override availability setting, and an administration maximum infusion duration override availability setting.

A net medication tolerance 542a is a maximum concentration of a medication that is safe to administer to a patient. The infusion system 210 associates the net medication tolerances with medications. Net medication tolerances 542a may be defined in medication identification files in a medication database. During infusion order creation 504, the infusion system 210 may determine the flow rate 560e, the number of infusion bags required 562a for a specified period of time, the concentration of the primary ingredient in each infusion bag, the time period over which each infusion bag is to be administered, and the total volume of each infusion bag. Flow rates may be manually entered or adjusted by altering the final concentration or the duration of each infusion bag. In general, the infusion system 210 performs a net concentration check 564a (FIG. 7) to ensure the maximum concentration of the medication is not exceeded. However, if at any time while a clinician 116 is modifying the flow rate by adjusting the final concentration resulting in the final concentration of a solution exceeding the maximum concentration of the medication, the infusion system 210 will send a message 520 to the administering clinician. The administering clinician may be authorized to override the net medication tolerance 542a. The infusion system 210 will usually require the clinician 116 to provide a reason for the override.

Infusion system 210 may include adjustable flow rate tolerances 542b and flow rate adjustment tolerances for administration. Flow rate tolerances 542b are optionally defined for all organizational levels of the patient care system 100. The tolerances 542b may be for the entire patient care system 100, or for sub-systems of the patient care system 100. For example, different flow rate tolerances 542b may apply to sub-systems such as, but not limited to, neonatal, pediatric, psychiatric, specific nursing units, and for specific patients. The flow rate tolerances 542b can be specified relative to the original ordered flow rate or relative to the immediately preceding flow rate. The clinician 116 may also specify a flow rate tolerance specific to a particular order. The infusion system 210 may include a pre-defined indication of whether the administering clinician 116 is permitted to override the flow rate tolerance 542b
without requiring a new order. This indication can apply to the entire patient care system 100, a sub-system, or an individual clinician 116.

The maximum infusion duration 542d may be separately definable for the various portions of the patient care system 100. The maximum infusion duration 542d may also be specific to a particular medication 124. A maximum infusion duration override 568d (FIG. 7) may be provided if it is permissible to override the maximum infusion duration 542d at the time of order entry. An administration maximum infusion duration override may be provided to set whether it is permissible to override the maximum infusion duration 542d at the time of administration and which group of users is allowed to do so. If it is permissible to override during order entry and/or administration, the infusion system 210 may define a subset of the clinicians 116 that have the authority to override the maximum infusion duration 542d.

Defaults 544 include defaults such as, but not limited to, medication diluent defaults 544a, diluent quantity defaults 544b, dose defaults 544c, and units of measure defaults 544d. Units of measurement (UOM) defaults 544d include the ability to specify the units of measurement that are most suitable for different portions of the patient care system 100. For example, medication may be measured in different units by physicians, administering clinicians, pharmacists, financial personnel, and medication screeners. The physician's UOM is generally a measurable value such as “mmol”, “mEq”, “ml”, and/or “mg”, as opposed to “vial” and/or “puff.” The physician's UOM is used for tasks such as ordering and entering information 560.

The Administering clinician's UOM is generally a value that reflects the UOM the medication will be administered in, such as “puff”, “tbsp”, and “tab”. The Administering clinician's UOM is used during medication administration 512. The Administering clinician's UOM may also appear on documentation such as administration reports, admixture fill and manufacturing work orders.

The pharmacy UOM is generally a value that reflects the physical form the medication is dispensed in such as “tab”, “vial”, “inhalator”, and “jar”. The pharmacy UOM is used in preparation 506 and in stocking and dispensing systems. The financial UOM is generally a value that will be used to calculate the financial figures that appear on bills and invoices. The medication screening UOM is generally used when screening the medication.

Units of measurement defaults 544d may be specified using a check-box table where checkmarks are placed in a table correlating the various UOMs with the users of the UOMs.
The infusion system 210 may use the same UOM for more one function. For example, the physician’s UOM may be the same as the pharmacist’s UOM. Setting defaults 544 include data necessary to coordinate the various UOMs. For example, UOM defaults 544d may include the multipliers and dividers necessary to create a one-to-one correspondence between the various UOMs. The UOM defaults 544b may be changed to suit the desires of the individual clinicians. However, the one-to-one correspondence should be maintained by the patient care system 100. The infusion system 210 may be designed to maintain a history of medication unit defaults.

The infusion system 210 may also include a medication measurement suffixes. The medication measurement suffixes may default during order entry. The medication measurement suffixes may be common units of measuring a medication and may include units related to patient characteristics such as body surface area and weight. Medication measurement suffixes may be designated per drug, per order type, per dose, and per UOM.

Building database 546 includes building databases and/or portions of a single database such as, but not limited to, preparation area 546a, additive information 546b, solution 546c, pre-mix definitions 546d, favorites 546e, timing override reasons 546f, flow rate override reasons 546g, translation tables 546h, flow rate description 546i, equipment and routing information 546j, and message trigger 546k.

Timing override reasons 546f include displayable reasons for modifying the timing of infusion orders. For example, timing override reasons 546f may include a stylus selectable reason for digital assistant display 118a for administering an infusion order at a time other than the time specified in the original infusion order. If the clinician 116 administers a medication outside the ordered administration time tolerance 542c, the clinician 116 may be required to choose a reason code for the modification from displayed reasons 1008f (FIG. 10).

Medications 124 and/or infusion orders may have flow rate tolerances, including system flow rate tolerances 542b. The infusion system 210 may include flow rate override reasons table 546g. Flow rate override reasons 546g are notations that the clinician 116 may choose from, and/or supply, if the clinician 116 needs to change the flow rate beyond the bounds defined by the flow rate tolerance 542b. The infusion system 210 may include a defined message trigger 546j indicating whether or not a message should be sent to the patient’s physician if a clinician 116 overrides an order defined flow rate tolerance. The infusion system 210 may also include defined message triggers 546k indicating whether or not a message should be sent, and to whom,
if a clinician 116 overrides a tolerance, such as flow rate tolerances 542b, defined at a level other than the order.

The infusion system 210 may include translation tables 546h such as, but not limited to, a flow rate translation table, a varying ingredient translation table, and varying flow rate translation table. Flow rate translation includes translating an infusion order into a flow rate defined by volume/time where the order is originally specified in any way such as, but not limited to, dosage/time with a particular concentration, volume per unit of weight/time, dosage per unit of body surface area/time, and total dosage and duration.

Varying ingredient translation includes translating a plurality of flow times of infusion orders with varying ingredients in separate infusion bags into the flow rate for the infusion bag currently being administered. Orders with varying ingredients include orders such as, but not limited to, sequencing orders. In sequencing orders, different bags have different ingredients and potentially different flow rates.

Varying flow rate translation includes translation of infusion orders with varying flow rates into the flow rate for the current solution being infused. Varying flow rate orders include orders such as, but not limited to, tapering dose orders and alternating dose orders.

The infusion system 210 may include predefined infusion flow rates 542b. The predefined infusion flow rates 542b may be associated with flow rate descriptions 546i to permit selection from a drop-down list as a shortcut from keying in the flow rate.

Defined functions 548 includes functions such as, but not limited to, preparation area function 548a, bag duration function 548b, verify override requests function 548c, duration to volume function 548d, duration to flow rate function 548e, and flow rate to drip rate function 548f. The infusion system 210 may include a duration-to-volume function 548d to determine the amount to be infused per the infusion order. Flow rate to drip rate function 548f uses information about the medical device 330 to convert flow rates to drip rates.

Determined settings 550 includes settings such as, but not limited to, override authorities 550a, flow rate precision 550b, volume precision 550c, and time precision 550d. The infusion system 210 may determine the total volume of infusions and the flow rate(s) of the infusion order. If these numbers are determined, it is necessary to round the calculated values to flow rate precisions 550b and volume precisions 550c that are comprehensible to clinicians 116 such as the physician, the pharmacist, and the nurse. Flow rate display precision 550b may be set to display the flow rate to a set number of decimal places. Various parts of the patient care system
100 may independently determine the precision for displayed flow rates. For example, the infusion system 210 may display to one decimal place for an adult treatment location, and to three decimal places for a neonatal treatment location. The flow rate precision 550b may reflect the service in which the clinician’s patient(s) are located. The flow rate(s) of the infusion order may be rounded to a system defined precision. The precision may be same for all infusion orders or be dependent on the patient’s service.

Volume display precision 550c may similarly be set to display infusion volumes to a set number of decimal places. Settable time precision 550d may be used to calculate the administration duration period based on flow rate if the infusion is a single dose infusion or an intermittent infusion. The total volume of each infusion bag calculated will be rounded according to the volume precision 550c. The administration time will be rounded by the infusion system 210 according to the set time precision 550d. The time precision 550d may be the same for all infusion orders regardless of the patient’s service or may be service specific.

FIG. 7 is a block diagram showing functional components for infusion order creation 504 of FIG. 5. Infusion order creation 504 includes functional blocks for creating infusion orders. Infusion order creation 504 includes entering information 560, calculations 562, checks 564, and overrides 568. Entering information 560 may include functions such as, but is not limited to, identifying the order type 560a, identifying the medications 560b, identifying the dose 560c, identifying the diluent 560d, identifying the flow rate 560e, and identifying the infusion site 560f.

Infusion order creation 504 is linked to infusion bag preparation 506, and infusion bag delivery (path 530), medication administration 512, and infusion order modifications 514. Infusion order types 560a include order types such as, but not limited to, single dosing, load dosing, intermittent dosing, and continuous. Continuous infusions include alternating infusions, sequencing infusions, tapering infusions, and titrating infusions. Upon selection of the first medication 560b in an infusion order, an infusion order type 560a form for the medication may default. The ordering clinician may have the option of selecting a different order type. The dose 560c and unit of measure 544d may also default. The unit of measure 544d may be correlated with the medication and/or the dose 544c. The infusion system 210 may include a default diluent, or several default diluents, for the medication. One default may be identified as a preferred diluent. A description may be associated with the diluent to assist the ordering
clinician to decide which diluent to select. The diluent description may include a reference avoiding use of a particular diluent if a patient is hypertonic.

The infusion system 210 may also allow additional infusion order types 560a based on the previously mentioned infusion order subtypes. Additional infusion order types 560a include, but are not limited to, TPN infusion orders, chemotherapy continuous infusion orders, piggyback infusion orders, and large volume parenteral infusion orders. The infusion order subtypes may be accessed from different parts of the infusion system 210 allowing sorting and filtering of infusion orders according to the subtypes. A special label format for each infusion order subtype can also be defined to further customize infusion order subtype orders and associated pharmacy workflow.

When searching for a medication 114 during infusion order creation 504, the medication 114 may be flagged as additive and/or a solution to aid the clinician 116 in creating the infusion order. This designation may be made in a medication identification file.

Medication dose 560c may be determined in a number of ways such as, but not limited to, according to body weight, body surface area, and entered according to rate. When the flow rate is not entered, the infusion system 210 will calculate the flow rate according to the dose and time period specified. The ordering clinician may specify the diluent 560d and its quantity. The pharmacy may provide a default for such parameters – see line 582 (FIG. 5). A check 564 may be performed to ensure the net concentration 564a for the medication 560b and the flow rate 564b are appropriate.

The infusion system 210 may identify and/or calculate flow rates 560e based on the patient’s weight, body surface area, and/or a specified frequency and duration of therapy. The ordered flow rate 560e is checked 564b against the flow rate tolerances, such as system flow rate tolerance 542b. The net concentration of the medication 124 may be checked 564a against net concentration tolerances, such as the system net concentration tolerance 542a.

Flow rate 560e may also include displaying descriptions of default flow rates to facilitate the entering of orders. Flow rate 560e may reference flow rate descriptions database 546i.

Calculations 562 may include calculating the dose based on patient weight and/or height (possibly provided by ADT interface 310), the drug amount, diluent volume, concentration, or rate.

Calculations 562 may include, but are not limited to, calculating the flow rate, if not specified in the prescription, the bag quantity 562a or number of infusion bags required for a
specified period of time, the time period over which each infusion bag is to be administered, and the total volume of each infusion and infusion bag based on the concentration of the ingredients in the solution. Flow rates, volume to be infused, and/or duration may be modified. If modified, the infusion system 210 will automatically calculate dependent quantities, based on calculations, if the maximum dosage for the ingredients in the concentration would be exceeded as identified in the ingredient's medication file, the patient care infusion system 210 will alert the pharmacist and/or clinician 116 and may ask for a reason code for the adjustment.

Calculations 562 may include calculations such as, but not limited to, bag quantity calculations 562a, translation calculations 562b, duration to volume calculations 562c, and flow rate to drip rate calculations 562d. Checks 564 include a variety of checks that an infusion order may be subject to. The checks include checks such as, but not limited to, a net concentration check 564a, a flow rate check 564b, an administration time check 564c, a duration check 564c, and an infusion site check 564e. If an infusion order fails a check 564, the clinician 116 may be able to override the check. Overrides 568 may include overrides such as, but not limited to, a net concentration override 566a, a flow rate override 566b, an administration time override 566c, a duration override 566d, and an infusion site override 566e. Overrides 568 may generate messages 520 for the physician and/or the pharmacy. The infusion system 210 may distinguish between system-wide and subsystem overrides in determining whether it is necessary to generate a message 520.

Overrides may include an indication of whether clinicians have the authority to override a tolerance. For example, flow rate override 568b may provide an indication of whether the clinician entering the infusion order has the authority to override the system flow rate tolerance 542b. This indication may apply to the patient care system 100 or a sub-system. Duration override 568d may provide an indication of whether the clinician 116 entering the infusion order has the authority to override the system duration 542d. This indication may apply to the patient care system 100 or a sub-system.

Overrides 566 also include displaying of reasons for the override 568f. Reasons for the overrides 568f may be selected by the clinician 116 from drop-down menus.

The result of the infusion order creation 504 is an infusion order 702. Infusion order 702 may include an infusion schedule 704. The infusion system 210 may look ahead a period of time and generate a the infusion schedule 704 - so long as the infusion order 702 is active - for infusion bag filling for that time period, or longer if specified on demand. The ordering clinician
is not required to specify an end-date for the infusion order. The infusion system 210 may include automatic scheduling of infusion bag delivery based on infusion system 210 defined tolerances 542.

FIG. 8 is a block diagram showing functional components for infusion order preparation 506 of FIG. 5. Infusion preparation 506 includes functional blocks for preparing infusion order 702. Infusion preparation 506 may include, but is not limited to, determining preparation location 506a, scanning ingredients 506b, bag duration checking 506c, and bar code printing 506d for medication labels 124a. Bar code printing 506d may include the functions described above in reference to print label 326 (FIG. 3).

After infusion orders are entered into the infusion system 210, preparation instructions are routed to a preparation location. The preparation location depends upon the infusion system's 100 preparation program 506 and the infusion components. The infusion system 210 may include adjustable databases, such as preparation area database 546a that specify where the infusion order is to be prepared. The infusion order may be prepared in the pharmacy or in a remote location, such as on the floor or at the treatment location 106. The clinician 116 is guided through the preparation process using event management information that may be displayed on digital assistant 118 or another device having a display.

The medication label 124a identifies the ingredients and ingredient concentrations. The medication label 124a may be printed in any location. The medication label 124a generally includes bar code printing 506d. Bar code printing 506b may include printing a bar code label 124a for each infusion bag. The label 124a ensures the correct medication is administered at the correct times and/or in the correct sequence. Alternating and sequencing infusion orders are particularly vulnerable to sequencing and timing errors. Bar code printing 506b may include printing a unique bar code label for every bag in infusion order 702. Bar code printing 506b may also include printing a bar code label 124a that uniquely identifies the combination of ingredients in an infusion bag and the concentration of those ingredients. The bar code for medication 124 may include a prefix, a suffix, and the national drug code (NCD).

FIG. 9 is a block diagram showing functional components for medication administration 512 of FIG. 5. Medication administration 512 includes functional blocks that are used to administer the medication to patient 112. Medication administration 512 may include reading a medication bar code 512a, reading a patient bar code 512b, running an expiration check 512c, providing titrate notification 512d, providing a flow rate to drip rate display 512e, providing "as
needed" infusion initiation 512f, downloading operating parameters 512g, and time monitoring 512h. The infusion system 210 may also translate orders that may have more than one flow rate, such as tapering and alternating orders, into the flow rate for the infusion bag currently being administered. The infusion system 210 may also translate orders having infusion bags with different ingredients, such as sequencing orders, into the flow rate for the infusion bag currently being administered.

Upon administering the medication 124, the clinician 116 scans the medication label 124a. The infusion system 210 includes scanning the bar coded label 24a when initiating the administration of the infusion order, when changing flow rates, changing bags, and/or stopping the infusion order. Infusion system 210 verifies that the infusion bag having the bar coded label should be administered at that time and is for patient 112. The history of the medication administration, including flow rates and volumes administered, may be captured and maintained.

Some infusion orders require hanging of an infusion bag with the intent of only a partial, specific amount of the infusion bag to be administered. The infusion system 210 will allow a clinician 116 to order an amount of an infusion bag to be administered. Most infusion pumps have the ability to define the volume to be administered or the flow rate and time period. Once this time has elapsed, the infusion pump will automatically prevent further administration. Infusion system 210 will, as a reminder to the administering clinician, provide a message on the medication label 114a that it is to be partially administered and the appropriate volume to be administered.

Flow rate to drip rate display 512e uses data generated by flow rate to drip rate functions 548f to provide the administering clinician with drip rates for the current infusion bag. During medication administration 512, the clinician 116 may check on the flow rate and other operating parameters using the digital assistant 118. Flow rate modifications 1002b (FIG. 10) are communicated in real-time.

The infusion system 210 may include PRN or "as needed" infusion initiation 512f. "As needed" infusion initiation 512 causes the creation of a new active order and the preparation of the PRN medication. This option may include prompting the clinician 116 to select a PRN infusion from a list of anticipatory PRN orders placed for the patient and defaulting the requested infusion bags to one. The clinician 116 may have the authority to modify the requested quantity of infusion bags.
Downloading of operating parameters 512g may include determining whether the patient identifier associated with the medical treatment and/or the patient identifier retrieved from the wristband 112a, is the same as the patient identifier associated with the medical treatment at the central location. The determination will often be made by the first computer, for example, the pharmacy computer 104a. If the infusion system 210 determines the various patient identifiers are not the same the system may generate an alarm message 520. If the infusion system 210 determines the various patient identifiers are the same, the infusion system 210 may download the operating parameters directly to the medical device 332. The infusion system 210 may send the operating parameters to a medical device 332, such as infusion pump 120.

One benefit of the system program 210 is that the operating parameters for the medical device 332 do not have to pass through digital assistant 118, or any other computer in the remote location, prior to the operating parameters being available to program the medical device 332. Bypassing computers at the remote location eliminates a potential source of errors in administering medication 124 to a patient 112. The operating parameters for the medical device 332 may be sent “directly” to the medical device 332 assuming the various verifications are achieved. In this context, “directly” meaning that the operating parameters may be sent to the medical device without passing through the digital assistant 118, or any other computer in the remote location.

In another embodiment, the infusion system 210 may include an additional block (not shown) where the central computer accepts a second medication identifier. The clinician 116 at the remote location may enter the second medication identifier. The second medication identifier may be a revised first medication identifier. For example, the second medication identifier may be part of the prescription or electronic physician order entry that is the source for the first patient ID and the operating parameters. The infusion system 210 may then confirm the first and second medication IDs are equivalent prior to sending the operating parameters to the medical device. The second medication ID may be replaced by a revised first medication ID between the time the prescription is entered and the time the medication 124 arrives at the treatment location 106. The infusion system 210 will then sound an alarm if the second medication identifier is not equivalent to the first medication identifier that was included in the medication label 124a. In a further embodiment, the infusion system 210 may include an additional block (not shown) where the operating parameter is used to program the medical device 332.
Various blocks of the infusion system 210, such as block 512, may include displaying treatment information on the digital assistant 118. This may include displaying information that mirrors the information on display 120c of infusion pump 120. The information on display 120c of infusion pump 120 may be supplemented with information about the patient 112, the patient location, and the infusion order. This information may include information regarding multiple channels of infusion pump 120. The displayed information may include information such as, but not limited to, personality, prompt line, status line, operating icons and pump head display. Operating icons include falling drop, stop sign, flow check piggyback, Guardian, and delay start. The pump head display includes information such as the drug label and the infusion rate. Those having ordinary skill in the art are familiar with the displayed information and operating icons described above.

The infusion system 210 time monitoring 512h calculates the time remaining for an order to be completed and the volume of an infusion order that remains to be administered. When the clinician 116 uses the infusion system 210 to administer the infusion order, to make flow rate changes, and to check on the status of an infusion, the infusion system 210 calculates time and volume remaining to be administered and indicates if the calculation indicates a partial bag will be used. For example, on the last bag of an order that is to be stopped before the full volume is administered, and/or on a bag within an order that must be changed before the full volume is administered, the clinician 116 is alerted on digital assistant 118 and/or cart 132. The alert may include a message such as “Please only administer 150 ml.”

Time monitoring 512h includes tracking any modifications made to the flow rate using bar code scanning. The pharmacy is alerted in real time to adjust the preparation 506 of the next required infusion bag according to the modification. Monitoring of preparation 506 and medication administration 512 allows for a just-in-time delivery of medication 124. Just-in-time delivery reduces wastage attributed to discontinued or changed infusion orders. Monitoring also ensures patient 112 safety.

For titrate PRN orders, the clinician 116 is automatically notified of required flow rate changes if the titration conditions in the order indicate that the flow rate must be changed. The infusion system 210 includes defined functions for calculating a conversion of flow rates to drip rates 548f. The infusion system 210 defined values may be adjustable. The infusion system 210 may include automatic translation of flow rate to drip rate 548f to assist the clinician 116 during administration of the treatment.
FIG. 10 is a block diagram showing functional components for infusion order documentation 1012, and the infusion order modifications 514 and messaging 520 of FIG. 5. Modifications 514 include functional blocks used to modify existing infusion orders. Modification 514 may also be viewed as creating new orders to replace existing infusion orders. Modification 514 may include modification changes 1002, generally all ordering options for new orders 1004 are available, rechecks 1006, recheck overrides 1008, and new flow rate to new drip rate display 1010. Infusion order modifications often lead to documentation 1012 and messaging 520. Modifications 514 include the functions described in reference to prescription modification module 336 (FIG. 3). However, modifications 514 are also accessible from other portions of the patient care system 100 such as, but not limited to, prescription entry 324, prescription activation 306, and prescription authorization 308.

Modifications 514 include modifying the duration 1002a, modifying the flow rate 1002b, using a new infusion site 1002c, identifying reasons for modifications 1002d, identifying the column of an infusion bag 1002e, and processing stop orders 1002f. Clinicians 116 may also change an infusion rate without an order if the patient 112 is complaining of discomfort or to facilitate fluid balance, such as when the patient 112 is vomiting.

Modification changes 1002 include identifying a new duration 1002a, identifying a new flow rate 1002b, identifying a new infusion site 1002c, identifying a reason for a modification 1002d, identifying the volume remaining in the infusion bag 1002e, and stop orders 516. The ordering options available during initial infusion order creation 504 are generally available for modifying the infusion order. Ordering options available during initial infusion order creation 504 include those shown in FIG. 7. Rechecks 1006 and recheck overrides 1008 are analogous to checks 564 and overrides 568 that are described in reference to FIG. 7. New flow rate to new flow rate display 1010 assists the clinician and minimizes the possibility of errors during medication administration 512. The modified infusion order may lead to a modified infusion schedule.

Flow rates are frequently modified at the treatment location 106 for reasons such as to catch-up without changing the schedule for preparation when the infusion has been inadvertently stopped for a short time period. Such modifications may not require new infusion schedule 704 to be communicated to the pharmacy. In other cases, the new schedule 704 should be communicated to the pharmacy or other preparation staff. Flow rate modifications 1002b may trigger infusion order scheduling changes and/or messages 520 for appropriate clinicians 116.
When a clinician 116 enters a flow rate modification 1002b into the infusion system 210 at treatment location 106, the clinician 106 may also elect to have the infusion schedule 704 recalculated and sent to the pharmacy. The clinician 116 has the option of requesting new medication labels 124a to be printed by bar code printing 506d module. The new medication labels 124a include data reflecting the new information for any of the previously prepared infusion bags.

The infusion system 210 and/or the clinician may request a modification to the infusion site 1002c. The site may be selected from a list of anatomical representations on a computer screen.

The clinician 116 generally is required to identify a reason for the modification 1002d. Reasons stored in databases such as, but not limited to, override reasons for timing 546f and override reasons for flow rate 546g, may be displayed for easy identification by the clinician 116. There may be a separate hard-coded reason for physician ordered modifications. For physician ordered modifications, the clinician 116 is generally requested to identify the physician.

Prior to implementing the modification, the volume remaining in the current infusion bag is identified 1002e. The clinician 116 may be offered the option of accepting a volume calculated from a displayed value of pre-modification flow rate and/or volume.

If desired, the current infusion may be stopped 1002f. If stopping the order is not required, for example the same infusion bag may be used with a new flow rate and/or a new medication added, the old flow rate may be identified and compared to the modified flow rate.

Any infusion bags that were previously prepared may be checked for expiration based on the new infusion schedule 704. When an infusion order is resumed following either a temporary stop or a hold order, the expiration check may be done regarding expiration of solutions that have already been prepared.

The new infusion schedule 704 is used to control the preparation 506 in the pharmacy or other preparation site. A system default 544 may be set for whether or not any prepared bags should be credited to the patient 112, through the billing interface 312, and whether or not they should be credited to inventory.

Infusion order changes 1002 include all ordering options available 1004 for new orders. The modified flow rate may be rechecked 1006 for rules and tolerances such as, but not limited to, net concentration 1006a, flow rate 1006b, administration time 1006c, duration 1006e, and infusion site 1006f. Overrides 1008 may be available for modifications that are outside of
tolerances. The infusion system 210 may display reasons 1008f for overrides and for administering medications at times other than that specified in the original order. The clinician 116 may be required to identify a reason for the modification.

The infusion system 210 may offer the clinician 116 a display indicating the modified drip rate associated with the modified flow rate 1012. The displayed information may be calculated by the flow rate to drip rate 548f defined function. The infusion system 210 may also be provided with descriptions of typical infusion tubing used within the infusion system 210 for use in calculating drip rates.

A modification results in the infusion system 210 validating the expiration of the infusion bag and providing a message to the clinician 116 if the infusion bag expires prior to the completion of the order. The message may request that the clinician 116 contact the pharmacy. The validation of the expiration of the infusion bag for solutions such as, but not limited to, premixed solutions and solutions manufactured outside of the infusion system 210, may include parsing the scan code.

Flow rate override 1008b may provide an indication of whether the clinician 116 modifying the infusion order has the authority to override the ordered override without requiring approval for a new infusion order. This indication may apply to the patient care system 100 or a sub-system.

Documentation 1012 captures infusion order information in real-time. Documentation includes documenting multiple infusions being administered at the same time and infusion modifications such as, but not limited to, duration changes 1002a, flow rate changes 1002b, volume changes 1012c, and infusion site changes 1002d.

The infusion system 210 may assist the clinician 116 in capturing all changes in flow rate as the changes are occurring. The clinician 116 may change the flow rate as called for in the order, such as to decrease a morphine infusion flow rate from 4 ml to 2 ml. Though the infusion system 210 may recognize the change as a new order, the infusion system 210 may be configured to avoid duplication so that the modified order does not result in the generation of a new bag.

Documentation 1012 includes the ability to document changes such as, but not limited to, an infusion that is stopped temporarily, discontinued, and/or restarted. The clinician 116 may stop infusion for a variety of reasons, such as the infusion site having been compromised, the infusion has been dislodged, and/or the infusion may be heparin/saline locked to facilitate the movement of patient 112. The infusion may be resumed when a new site/infusion has been
reestablished. However the length of time this may take is variable and is generally recorded by the infusion system 210.

Government regulations often require tracking of every step in the process of infusion administration. Infusion system 210 allows the administering clinician 116 to document flow rate modifications on a digital assistant 118, or other computer device, by scanning the medication label 124a and adjusting the flow rate 1002a based on a tolerance, such as a tolerance created by set tolerance 542. A flow rate modification 1002b corresponds in real time with the associated pharmacy’s infusion schedule 704 to ensure just-in-time inventory management of infusion bags to the patient treatment area 106. Documentation 1012 may allow order backdating under some circumstances.

The infusion system 210 includes the ability to document the infusion site 1012d and multiple infusions 1012e for multiple infusion sites. In many situations a patient 112 may have multiple medications 124 and “y-ed” infusions so that the some infusions are running into one site and other infusions are infusing into another site. For example, morphine infusion, antibiotics and normal saline infused into the right arm (site 1) and TPN and 2/3 &1/3 running into a double lumen CVL (site 2). The infusion system 210 allows clinician 116 to document which site the various fluids are infusing through. In treatment locations 106, such as intensive care units, many more than two infusions may be running into one line or one lumen. Clinicians 116 are able to indicate which lumen of a CVL the infusion or medication is running into.

The infusion system 210 includes the ability to document the site location 1012d for infusions and any site location changes. Infusion sites are frequently changed due to occlusions or policy. Therefore, clinicians 116 must document a change in the site location if an infusion becomes dislodged and was subsequently restarted.

The infusion system provides for centralized device configuration. Operating parameters for medical devices 332, such as infusion pump 120, often include defaults and/or tolerances. The defaults and/or tolerances may reside in the infusion system 210, for example flow rate tolerance 542b, and/or in a memory associated with the device 332. For example, infusion pumps 120 may include a database having a table of medications having associated flow rate tolerances. If the clinician 116 enters a flow rate that is beyond the associated flow rate tolerance, the clinician 116 is warned and then may be allowed to proceed - or prohibited from proceeding. Devices 332 such as heart rate monitors may also have configurable tolerances for alerts. In addition to alerts, many other characteristics can typically be configured for devices
332 such as: network name, IP address, polling frequency, and colors. The infusion system 210 includes configuring medical devices 332 individually or in groups from one or more central computers.

System configuration parameters may be defined for a first type of medical device. The system configuration parameters will be sent and accepted by the first type of device unless the particular first type of device has more specific configuration parameters that apply to that particular first type of device. For example, a first plurality of a first type medical device may be located at general care treatment locations. A second plurality of the first type of medical device may be located at an intensive care treatment location. The general care treatment location may not have specific configuration parameters while the intensive care treatment location does have specific treatment parameters. System configuration parameters will apply to all of the first type of medical devices throughout the infusion system 210, i.e. the devices in the general care treatment locations, unless specific configuration parameters apply, e.g. the intensive care treatment location.

For each type of device, specific configuration parameters that apply to all devices of that type across a particular grouping of the devices override the system configuration parameters if a particular device belongs to the group having such a definition, unless the specific configuration parameters are overridden at an even more specific level within the infusion system 210. The groups might be defined as a clinical service, a nursing unit, and/or a combination of service and nursing unit.

For each type of device, the user can define sets of configuration parameters that apply to all devices of that type being used for operations with specified ranges of attributes that override any other definition. In a hospital the operations might consist of infusion orders and the attributes might include patient weight, drug, patient disease state, and patient acuity.

Devices may be identified as part of a general group, a specific group, and/or to be associated with a particular patient by including the device address in a table in a database. General or specific configuration parameters may then be sent to the device according to the identification of the device. The specific configuration parameters may then be read back to the infusion system 210 and compared to the originally sent configuration parameters to verify the original configuration parameters were correctly received by the device 332. If the configuration parameters were not correctly received, the infusion system 210 may provide a message 520 identifying the discrepancies or the communication failure.
The infusion system 210 may detect changes to configuration parameters made at the
device, rather than through a central computer, and send a message and/or alert 520. The
infusion system 210 may also poll the devices to verify their configuration parameters. If system
and/or specific configuration parameters change, the changes may be propagated to all devices
332 identified in the system as belonging to the group according to the groupings identified in
the infusion system 210.

Throughout this document and the related claims, Acentral location® and Aremote
location® are relative terms to each other. Aremote location® is any location where a patient
is receiving treatment through a controlled medical device, such as a patient treatment location
106 where patient 112 is receiving treatment through an infusion pump 120. Acentral
location® is any location, other than the remote location, where parameters for operating the
medical device are accessible such as, but not limited to, the location of the pharmacy computer
104 and the central system 108. In a typical arrangement, several remote locations, such as
treatment location 106, are in communication with a central location.

A method of administering a medication with the infusion system 210 is described
below. The method includes the ability to modify the infusion order. The modifications include
modifications to the flow rate, the infusion site, temporary stops to the infusion, restarting the
infusion, and hanging a new medication 124 container. The method includes: scanning a bar
code associated with the patient 512b; scanning a bar code associated with the medication 512a;
if the infusion is an admixture, validating the expiration 512c; selecting a reason for the
modification 1002d; and recording the remaining volume of the infusion bag or accepting the
value calculated from the previous volume and flow rate 1002e. The validation of the expiration
512c of the infusion bag may include the use of an admixture table and/or a barcode.

The reason for the modification may come from a defined table 546g. The reason for the
modification may also include a hard-coded value for physician-ordered changes. When the
hard-coded value is selected, the clinician 116 is prompted to select the physician from a list of
physicians. The attending physician may be the default in the list of physicians.

There may be a quick select feature to halt the administration of the medication 124, for
element stop order 1002f. If the quick select is not chosen, the following steps may be included:
recording the flow rate and/or accepting the previous value for the flow rate - the previous value
is generally displayed on the digital assistant display 118a, the infusion pump display 120c,
and/or the medical cart 132; comparing the previous flow rate to the ordered flow rate - this
comparison may be accomplished by using infusion system 210 or subsystem rules and tolerances; displaying appropriate messages; conversions between flow rates and drip rates may be displayed 1012 - the conversions may be calculated based on infusion system 210 defined drip-rate conversion tables 548f. The infusion system 210 typically uses descriptions based on the tubing used to make it easy for the clinician 116 to select the correct drip rate conversion.

Changing the flow rate triggers the infusion system 210 to validate the expiration of the infusion bag(s) based on scheduled flow rate. If the solution expires before or during the administration, send a message to the clinician 116, such as “This solution will expire during the scheduled administration period. Please contact the pharmacy.” If it is a premixed infusion bag and/or a customized infusion bag, validate the expiration by parsing the scan code, if possible. Accept the previous infusion site or select a new infusion site location from a list or a graphical anatomical representation. Then recalculate the schedule 704 to implement pharmacy restocking.

Infusion system 210 may include biometrics for identifying patients and clinicians 116. Prior to allowing a clinician 116 to access the infusion system 210, the infusion system 210 accesses information related to the identity of the clinician 116. The infusion system 210 may identify the clinician by using a device, such as a bar code reader, to read the clinicians’ badge 116a. The system may also use biometrics to positively identify the clinician 116, to assure the clinician is an authorized user of the system, and to determine whether the clinician 1176 has authority to access portions of the infusion system 210. The infusion system 210 may require a combination of the clinician badge 116a, or other key, and a verified biometric match in order to grant the clinician access to the infusion system 210. The system may also be configured to terminate access to the infusion system 210 when the clinician badge 116a is removed from the vicinity of the device used to read the clinician badge 116a, or other key.

Biometrics is the technology and science of statistically analyzing measured biological data. One field of biometrics is that of determining unique physical characteristics, such as fingerprints. Biometrics makes it possible to identify individuals to digital systems, such as infusion system 210. A digital persona is created that makes transactions and interactions more convenient and secure. Biometric features for identification include features such as, but not limited to, fingerprint, face, iris and retina scanning, and voice identification. Biometric devices include a scanning or reading device, software to convert the scanned information into a digital format, and a memory to store the biometric information for comparison with a stored record.
Software identifies specific matched points of data that have been processed with an algorithm and compares the data. Unlike passwords, PIN codes, and smartcards, the infusion system 210 biometrics cannot be lost, forgotten, or stolen.

The biometric scanner may be associated with the device for reading the clinician’s badge 116a. For example, the biometric scanner may be a thumb print reader on the handle of a bar code reader. In other embodiments, the biometric scanner and an electronic key reader may be located on the portable medicine cart and/or the medical device. When the clinician 116 places the electronic key within a specified distance of the medical device, a processor will know the specific individual electronic biometric identification file it should expect. The infusion system 210 preferably prompts the clinician 116 to scan his biometric information. The biometric information is entered into the infusion system 210 with some type of biometric reading or scanning device. A one-to-one comparison is made between the scanned biometric information and the previously stored specific individual electronic biometric identification file. This one-to-one identity comparison is more efficient than comparing one-to-many identity files because it does not require searching an entire clinician database for a match. Instead, only one specific comparison is made. If there is a match, then the clinician 116 is granted access to the medical device 332. If there is no match, the clinician 116 is denied access.

In another embodiment, after the infusion system 210 grants access to the clinician 116, the infusion system 210 may terminate that access when the electronic key is removed from the biometric scanner, or the vicinity of the biometric scanner. The vicinity within which the electronic key must be kept may be predetermined and/or may be a variable and programmable infusion system 210 parameter.

In one embodiment, the infusion system 210 includes an encrypted digital fingerprint template, a clinician’s name, a login name, and a password. One technology for implementing the clinician identifier includes “IBUTTON 400” technology from Dallas Semiconductor technology. The infusion system 210 may be activated when the clinician places a finger on a fingerprint scanner. If the infusion system 210 finds a match, the infusion system 210 may request the clinician 116 login to the infusion system 210. If the infusion system 210 does not find a biometric match, the system does not allow the clinician 116 to access the infusion system 210.

In another embodiment, the database storing biometric information may be kept in the central system 108, the pharmacy computer 104, and/or the treatment location 106. At the
treatment location 106, the database may be maintained in the portable cart, the digital assistant 118, and/or the medical device 332. Such distributed databases will allow access to remote devices even if the network 102 is unable to communicate between the various locations. When network 102 communication is reestablished, the remote and central databases may be synchronized with any information modified at the other location so that both infusion system 210 databases are properly updated.

The infusion system 210 provides a closed loop infusion therapy management system. The closed loop begins with a clinician 116 order. Among other methods, the clinician 116 may enter the order through digital assistant 118 and/or medical treatment cart 132. The order is then available in real-time for pharmacy authorization 508 and physician authorization 510. The order is available in real-time as an electronic medication administration record (eMAR). The eMAR is available to the clinician 116 for infusion administration. The infusion system 210 automatically documents medication administration 512 and modifications 514 such as flow rate changes 1002b. Through the process of medication administration 512, the infusion system 210 simultaneously adjusts infusion system 210 and/or sub-system inventory and billing 518. The infusion system 210 also provides event management and decision support data. The infusion system 210 is device independent, meaning that it can be run on workstations, wireless tablets, and handheld digital assistants 100. The infusion system 210 generally runs in real time, however, batch processing and or messaging may be used to coordinate various stages of the infusion system 210 processes.

The closed loop infusion therapy management system includes infusion order entry 560, order preparation 506, and the availability of the status of the infusion. Infusion order entry 560 may be through a number of means such as, but not limited to, the prescription entry module 324, the prescription modification module 336, and the pharmacy interface 316. Computer screen 400 may be employed in entering the infusion order. The status of the infusion provides patient 112 specific usage of infusions and alerts the pharmacy of the need for additional infusion bags.

It should be emphasized that the above-described embodiments of the present invention, particularly, any Apreferred® embodiments, are possible examples of implementations, merely set forth for a clear understanding of the principles of the invention. Many variations and modifications may be made to the above-described embodiment(s) of the invention without substantially departing from the spirit and principles of the invention. All such modifications
are intended to be included herein within the scope of this disclosure and the present invention and protected by the following claims.
Claims

What is claimed is:

1. A system for administering a medication, the medication being in a container, the container having a medication bar code associated therewith, the system comprising:
   a first computer, the first computer having data defining a first flow rate, the first computer having data defining a first flow rate tolerance;
   a central time source;
   an infusion pump;
   a second computer, the second computer designed to accept information from a bar code reader, the second computer designed to provide a first signal to the first computer, the first signal including data identifying the medication, the second computer designed to provide a second signal to the first computer, the second signal including data identifying a second flow rate, the second computer designed to provide a third signal to the first computer, the third signal including data identifying the volume of medication in the medication container,
   where the first computer authorizes the second flow rate if the second flow rate is within the first flow rate tolerance,
   where the infusion pump receives new operating parameters to implement the second flow rate if the first computer authorizes the second flow rate, and
   where the first computer documents the initiation of the second flow rate using the central time source.

2. The system of claim 1,
   where the first computer includes a first flow rate tolerance override, and the first computer includes a database identifying acceptable reasons for overriding the first flow rate tolerance,
   where the second computer is designed to provide a fourth signal to the first computer, the fourth signal including data identifying a first reason for overriding the first flow rate tolerance,
   where the second flow rate is outside of the first flow rate tolerance, and
   where the infusion pump receives new operating parameters if the first reason is an acceptable reason for overriding the first flow rate tolerance.

3. The system of claim 1, where the medication has an expiry, and the first
computer confirms the second flow rate does not exceed the expiry prior to authorizing the second flow rate.

4. The system of claim 1, where the first computer includes data defining a first infusion order, the first infusion order including an order flow rate tolerance, and
   where the first computer authorizes the second flow rate if the second flow rate is within the order flow rate tolerance.

5. The system of claim 1, where a message is sent to a pharmacy if the first computer authorizes the second flow rate.

6. The system of claim 1, where a message is sent to a physician if the first computer authorizes the second flow rate.

7. The system of claim 1, where the first computer includes a database identifying clinician authority levels, where a first plurality of clinicians are identified in the database as having the authority to override the first flow rate tolerance and a second plurality of clinicians are identified in the database as not having the authority to override the first flow rate tolerance,
   where the second computer is designed to provide a fourth signal to the first computer, the fourth signal including data identifying a clinician,
   where the second flow rate is outside of the first flow rate tolerance, and
   where the infusion pump receives new operating parameters if the clinician is in the first plurality.

8. The system of claim 1, where the second computer is designed to provide a display, the display relating the second flow rate to a drip rate.

9. The system of claim 1, where a message is sent to the pharmacy if the second flow rate requires a new infusion bag.

10. The system of claim 1, further comprising one or more computer programs for causing the first computer to authorize the second flow rate if the second flow rate is within the first flow rate tolerance and to document the initiation of the second flow rate using the central time source.
FIG. 4

PROCESSING AREA 402

SEARCH AREA 404

MEDICATION INFORMATION AREA

SOLUTIONS 406

ADDITIVES

SEARCH AREA 404

PROJECTED SOLUTIONS AREA 412

TITRATION/TAPERING AREA 408

INSTRUCTION AND NOTE AREA 410
FIG. 6

- NET MEDICATION (542a)
- FLOW RATE (542b)
- ADMINISTRATION TIME (542c)
- SYSTEM DURATION (542d)
- MEDICATION DURATION (542e)
- SITE CHANGE (542f)
- SET TOLERANCES (542)

- MEDICATION DILUENT (544a)
- DILUENT QUANTITY (544b)
- DOSE (544c)
- UNITS OF MEASURE (544d)
- SET DEFAULTS (544)

- PREPARATION AREA (546a)
- ADDITIVE INFORMATION (546b)
- SOLUTION INFORMATION (546c)
- PRE-MIX DEFINITIONS (546d)
- FAVORITES (546e)
- OVERRIDE REASONS FOR TIMING (546f)
- OVERRIDE REASONS FOR FLOW RATE (546g)
- TRANSLATION TABLES (546h)
- FLOW RATE DESCRIPTIONS (546i)
- EQUIPMENT AND ROUTING INFORMATION (546j)
- MESSAGE TRIGGERS (546k)
- BUILD DATABASE

- PREPARATION AREA (548a)
- BAG DURATION (548b)
- VERIFY OVERRIDE REQUESTS (548c)
- DURATION TO VOLUME (548d)
- DURATION TO FLOW RATE (548e)
- FLOW RATE TO DRIP RATE (548f)
- DEFINE FUNCTIONS (550)
- DETERMINE SETTINGS (550d)

- OVERRIDE AUTHORITIES (550a)
- FLOW RATE PRECISION (550b)
- VOLUME PRECISION (550c)
FIG. 7

ORDER TYPE 560a

MEDICATION 560b

DOSE 560c

DILUENT 560d

FLOW RATE 560e

INFUSION SITE 560f

ENTERING INFORMATION 560

BAG QUANTITY 562a

TRANSLATIONS 562b

DURATION TO VOLUME 562c

FLOW RATE TO DRIP RATE 562d

CALCULATIONS 562

NET CONCENTRATION 564a

FLOW RATE 564b

ADMINISTRATION TIME 564c

DURATION 564d

INFUSION SITE 564e

CHECKS 564

NET CONCENTRATION 566a

FLOW RATE 566b

ADMINISTRATION TIME 566c

DURATION 566d

INFUSION SITE 566e

DISPLAY REASONS 566f

OVERRIDES 566

INFUSION ORDER 702

INFUSION SCHEDULE 704
FIG. 10

NEW DURATION 1002a
NEW FLOW RATE 1002b
NEW INFUSION SITE 1002c
IDENTIFY REASON FOR MODIFICATION 1002d
IDENTIFY VOLUME OF BAG 1002e
STOP ORDER CHANGE 1002f

ALL ORDERING OPTIONS AVAILABLE 1004

NET 1006a CONCENTRATION
FLOW RATE 1006b
ADMINISTRATION TIME 1006c
DURATION 1006d
INFUSION SITE 1006e
RECHECKS 1006

NET 1008a CONCENTRATION
FLOW RATE 1008b
ADMINISTRATION TIME 1008c
DURATION 1008d
INFUSION SITE 1008e
DISPLAY REASONS 1008f
OVERRIDES 1008

NEW FLOW RATE TO NEW DRIP RATE DISPLAY 1012

DURATION 1012a
FLOW RATE 1012b
VOLUME 1012c
INFUSION SITE 1012d
MULTIPLE INFUSIONS 1012e
DOCUMENTATION 1012

SYSTEM 520a
PHARMACY 520b
PHYSICIAN 520c
BILLING 520d
INVENTORY 520e
MESSAGING 520
### INTERNATIONAL SEARCH REPORT

**PCT/US 03/13335**

#### A. CLASSIFICATION OF SUBJECT MATTER

<table>
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<th>IPC</th>
<th>Classification</th>
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<td>7</td>
<td>G06F19/00 A61M5/172</td>
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According to International Patent Classification (IPC) or to both national classification and IPC.

#### B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols):

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

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

**EPO-Internal**

#### C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<tr>
<td>A</td>
<td>US 5 338 157 A (BLOMQUIST )&lt;br&gt;16 August 1994 (1994-08-16)&lt;br&gt;abstract&lt;br&gt;column 4, line 19 - line 29&lt;br&gt;column 6, line 48 - line 55&lt;br&gt;column 15, line 23 - line 40&lt;br&gt;column 16, line 48 - column 17, line 8; figures 1,1A,2</td>
<td>1-10</td>
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<tr>
<td>A</td>
<td>US 5 781 442 A (CHAMBERLAIN ET AL.)&lt;br&gt;14 July 1998 (1998-07-14)&lt;br&gt;abstract&lt;br&gt;column 2, line 23 -column 3, line 20&lt;br&gt;column 9, line 52 - line 62&lt;br&gt;column 11, line 33 - line 55; figures 1-7</td>
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Further documents are listed in the continuation of box C.

Patent family members are listed in annex.

- **A** document described the general state of the art which is not considered to be of particular relevance.
- **E** earlier document published on or after the international filing date.
- **L** document which may throw doubts on the validity of the claimed invention.
- **O** document referring to an oral disclosure, use, exhibition or other means.
- **P** document published prior to the international filing date but later than the priority date claimed.
- **Y** document published after the international filing date or priority date and not in conflict with the invention but cited to understand the principle or theory underlying the invention.
- **X** document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone.
- **V** document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.
- **A** document of the same patent family.

**Date of the actual completion of the international search**

14 October 2003

**Date of mailing of the international search report**

21/10/2003

Name and mailing address of the ISA

European Patent Office, P.B. 5818 Patentlaan 2<br>NL - 2280 HV Rijswijk<br>Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax (+31-70) 340-3016

**Authorized officer**

Michels, N
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| A        | US 5 788 669 A (PETERSON)  
4 August 1998 (1998-08-04)  
column 1, line 51 - column 2, line 26  
column 3, line 2 - line 17  
column 3, line 36 - line 38  
column 5, line 4 - line 64  
column 6, line 49 - line 65; table 50  
column 7, line 34 - line 50; figures 1-3 | 1-10 |
| A        | US 4 756 706 A (RUBALCABA JR. ET AL.)  
12 July 1988 (1988-07-12)  
abstract  
column 6, line 7 - line 51; figures 5,6,12 | 1-10 |
| A        | EP 0 960 627 A (BRAUN MELSUNGEN AG)  
1 December 1999 (1999-12-01) | |
| A        | US 6 135 949 A (JORDAN ET AL.)  
24 October 2000 (2000-10-24) | |
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<tr>
<td>US 4756706 A</td>
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