The ingestible event marker data framework provides a uniform, comprehensive framework to enable various functions and utilities related to ingestible event marker data (IEM data). The functions and utilities include data and/or information having an aspect of data derived from, collected by, aggregated by, or otherwise associated with, an ingestion event.
FIG. 2
FIG. 3

- IEM Data Environment 104
  - IEM Data Source Devices 300
    - Ingestible Devices 300a
    - IEM Data Receivers 300b
    - Health Devices 300c
  - Products 302
    - Ingestible Device / Pharmaceutical Product 302a
  - Events 304
    - Individualized Patient Data 306a
    - Multiple Patient Data 306b
  - Patient Specific Parameters 306
  - IEM Data Algorithms 308
  - Storage Repositories 310
    - Multisensor Lead 312a
  - Other Sources 312
FIG. 5
FIG. 6

PATENT ABSTRACT

The present disclosure relates to a patient-centered medical record system and method that provides a user interface for entry and viewing of patient information. The system includes a server for storing patient data, a database for storing patient information, a feedback loop system for collecting patient feedback, and a data system for accessing and updating patient information. The system also includes a graphical interface for navigating and interacting with the patient information. The disclosure further includes methods for updating patient information and displaying patient information in a user-friendly manner.
INGESTIBLE EVENT MARKER DATA FRAMEWORK

CROSS-REFERENCE TO RELATED APPLICATIONS

Pursuant to 35 U.S.C. §119 (e), this application claims priority to the filing date of U.S. Provisional Patent Application Ser. No. 61/079,082 filed on Jul. 8, 2008; the disclosure of which application is herein incorporated by reference.

FIELD OF THE INVENTION

The present invention relates generally to the technical fields of ingestible devices and communications. More specifically, and in various example embodiments, the present invention relates to a method, article, and system of generating, collecting, managing, distributing, and otherwise utilizing information associated with ingestible events and responses to the ingestible events.

BACKGROUND

Information related to personal events is widely needed in various pursuits. A personal event is an event that is specific to an individual. Examples of personal events include onset of a physiologic parameter of interest, ingestion of a therapeutic agent, etc.

There are many instances where one may want to note a personal event. Examples of such instances include onset of one or more physiologic parameters of interest, including appearance of disease symptoms, administration of medication, ingestion of certain types of foods, commencement of an exercise regimen, ingestion of certain substance, etc.

A variety of different methods and technologies have been developed to note a personal event. For example, techniques have been developed in which individuals can manually record data in a log or physically enter data via a computer device.

The accuracy of such notations may be dependent on the accuracy of data input, the accuracy of proxies used as actual data substitutions, etc. As a result, inaccuracies may occur.

In one example, an individual may suffer from one or multiple health conditions that require therapy with multiple medications. The multiple medications may be prescribed according to an intricate dosing schedule. The complexities associated with multiple health conditions, multiple medication therapies, and intricate dosing schedules may confuse the patient, resulting in inaccurate data capture.

In one example, the individual may have physical or cognitive deficits which may result in difficulties inputting and capturing data. The individual may forget to enter the data, or may enter the data incorrectly.

In one example, the individual may not wish to be inconvenienced and thus may intentionally refuse to enter the data. Conversely, the individual may unintentionally or intentionally enter/record data which is completely inaccurate. For example, the individual may receive periodic, prescheduled reminders to take some medication. The reminders are unable to take into account actual ingestion of the medication. If the individual has already taken the medication, the reminder is both moot and likely to inconvenience the individual. If the medication has not been taken, an inconvenient or unneeded reminder or alert may prompt the user to enter data or send a message advising that the medication has been taken just to quell the alarm while not actually taking the medication. The individual may intentionally leave out portions of the data.

In one example, proxies for data and information may also be inaccurate. For example, “intelligent” medication containers may contain microchips that sense opening of the medication container. From the sensed act of opening the container, an inference may be drawn that medication associated with the medication container has been ingested. The inference may be inaccurate, however, as medication is not necessarily ingested by virtue of opening a medication container.

The above-instances may ripen into further issues if particular parties besides the individual wish to use the individual’s personal event data. Examples of users and potential users (sometimes collectively referred to herein as “party” or “parties”) of personal event data include family and professional caregivers; communication companies; government agencies, e.g., agencies associated with government provided healthcare coverage; private insurance providers; Food and Drug Administration (FDA); Drug Enforcement Administration (DEA); US Bureau of Alcohol, Tobacco, and Firearms (ATF); care providers; medical device manufacturers; patients; clinicians; pharmaceutical manufacturers; pharmacies; web communities; software providers; marketing and financial analysts; and insurance companies.

Competing interests may exist between an individual’s privacy interests in personal event data and the acquisition and appropriation of the personal event data by third parties.

Further, various parties may have a compelling interest in receipt of accurate and comprehensive data, e.g., useful data, either in isolated form (data germane to a particular individual) or empirical form (aggregated data from various sources, various individuals, various personal events of an individual, etc.)

In many circumstances, however, accurate personal event data are not available. The party may have access to faulty data or a crude approximation of the information sought, as discussed above. Thus, the party must rely on such crude proxies to formulate a conclusion. It follows, then, that such conclusions may themselves be skewed or inaccurate. Actions taken in reliance on such conclusions may prove misguided, error-prone, and/or harmful.

To illustrate, a healthcare provider or family member may receive a message from a patient indicating that the patient has taken the medication when, in fact, the patient is merely providing the message without having actually ingested the medication. If the healthcare provider notices changes in the patient’s symptoms in close temporal proximity to receipt of the flawed information suggesting medication ingestion, the healthcare provider may mistakenly conclude that the patient’s symptoms are a result of the medication ingestion. Based on the mistaken conclusion, the healthcare provider may adjust the medication dosage in an attempt to alleviate the symptoms, perhaps to the patient’s detriment.

Of note, the more widely propagated and aggregated the inaccurate data, the more prolific the spread of and reliance on error-associated data and conclusions drawn therefrom.

In addition, recipients of the personal event data may wish to timely receive and utilize such information via a user-friendly, reliable and sophisticated means. The recipi-
ents may wish to receive and/or utilize information in discrete areas, integrate the personal event information with other data, and use the personal event information for various purposes.

[0018] Examples of various purposes include refining and optimizing data such as patient population data; incentivizing individuals or groups based on personal event data, e.g., ingestible event marker data ("IEM data"); corroborating and advancing decisions; supporting stakeholders’ decisions; using IEM data in personalized products and services, e.g., user applications on a mobile telephone; auto refilling prescription medications; managing pharmaceutical life cycle systems and controlled substances; compiling and delivering IP news and information feeds; accessing open sources of anonymized patient population data; determining eligibility and approval for refills, insurance coverage, etc.; using patient tools; participating in social network systems; analyzing aggregated data to derive and/or generate predictive information; supporting and enabling financial transactions; identifying direct and indirect causal failure points in treatment and predict corrective action; and providing dynamic, accurate calendaring/scheduling functions.

[0019] Finally, parties may also wish to access personal event data in conjunction with existing systems, e.g., commercial systems such as automated pharmacy systems, banking and financial systems, etc.

[0020] As can be seen, methods and systems are needed to seamlessly collect, manage, and distribute personal event data to various parties and systems.

[0021] Therefore, there is a need for controlled collection, management, and delivery of accurate personal event data to multi-profile parties for various purposes.

BRIEF SUMMARY OF THE INVENTION

[0022] The ingestible event marker data framework provides a uniform, comprehensive framework to enable various functions and utilities related to ingestible event marker data (IEM data). The functions and utilities include data and/or information having an aspect of data derived from, collected by, aggregated by, or otherwise associated with, an ingestion event. In one example, the IEM data are generated via an ingested device. The term “ingested device” includes any device, mechanism, structure, combined structure, or object capable of ingestion by a human subject or a non-human subject.

[0023] The IEM data framework is highly scalable and integratable with various existing systems, e.g., systems having computer-related component(s). Specific examples of such systems include pharmacy systems, communication systems, financial and banking systems, school systems, medical systems, government agencies, web communities, and personal computer systems. Such existing systems are herein collectively referred to as "commercial systems".

[0024] The IEM data framework enables multiple and various types of implementations. The implementations include various configurations of hardware, software, communication components, and/or data. For example, in one aspect, the IEM data framework is implemented with a basic complement of core components; namely, ingestible event marker data; a hub to receive the ingestible event marker data; and at least one ingestible event marker data system to receive, directly or indirectly, the ingestible event marker data from the hub.

BRIEF DESCRIPTION OF THE FIGURES

[0025] FIG. 1 provides a diagrammatic representation of a communication environment including an IEM data framework, according to one embodiment.

[0026] FIG. 2 provides a diagrammatic representation of the IEM data framework of FIG. 1, according to one embodiment.

[0027] FIG. 3 illustrates IEM data and an IEM data environment associated with the IEM data framework of FIG. 2, according to one embodiment.

[0028] FIG. 4 illustrates a hub associated with the IEM data framework of FIG. 2, according to one embodiment.

[0029] FIG. 5 illustrates exemplary IEM data systems associated with the IEM data framework of FIG. 2, according to one embodiment.

[0030] FIG. 6 illustrates an exemplary IEM data framework having a feedback loop system, according to one embodiment.

[0031] FIG. 7 illustrates an exemplary IEM data framework having a decision support system, according to one embodiment.

[0032] FIG. 8 illustrates an exemplary IEM data framework having an auto refill system, according to one embodiment.

[0033] FIG. 9 illustrates an exemplary IEM data framework having patient tools, according to one embodiment.

[0034] FIG. 10 illustrates an exemplary IEM data framework having a behavioral medicine system, according to one embodiment.

[0035] FIG. 11 illustrates an exemplary IEM data framework having an incentive system, according to one embodiment.

[0036] FIG. 12 illustrates an exemplary IEM data framework having a personalized commercial products/services system, according to one embodiment.

[0037] FIG. 13 illustrates an exemplary IEM data framework having an auto billing system, according to one embodiment.

[0038] FIG. 14 illustrates an exemplary IEM data framework having a tracking system, according to one embodiment.

[0039] FIG. 15 illustrates an exemplary IEM data framework having an interdiction system, according to one embodiment.

[0040] FIG. 16 illustrates an exemplary IEM data framework having a subscription system, according to one embodiment.

[0041] FIG. 17 illustrates an exemplary IEM data framework having an ingestible event marker data collection system, according to one embodiment.

[0042] FIG. 18 illustrates an exemplary IEM data framework having an approval system, according to one embodiment.

[0043] FIG. 19 illustrates an exemplary IEM data framework having a forecasting system, according to one embodiment.

[0044] FIG. 20 illustrates an exemplary IEM data framework having a financial system, according to one embodiment.
FIG. 21 illustrates an exemplary IEM data framework having an ingestible event marker data phone system, according to one embodiment.

FIG. 22 illustrates an exemplary IEM data framework having a social network system, according to one embodiment.

DETAILED DESCRIPTION

1.0 Overview

2.0 Ingestible Event Marker (IEM) Data Framework

2.1 IEM Data

2.1.1 IEM Data Environment

2.1.1.1 IEM Data Source Devices

2.1.1.2 Products

2.1.1.3 Events

2.1.1.4 Patient Specific Parameters

2.1.1.5 IEM Data Algorithms

2.1.1.6 Storage Repositories

2.1.1.7 Other IEM Data Sources

2.2 Hub

2.3 IEM Data Systems

2.3.1 Feedback Loops

2.3.2 Decision Support Systems

2.3.3 Auto Refill Systems

2.3.4 Patient Tools

2.3.5 Behavioral Medicine Systems

2.3.6 Incentive Systems

2.3.7 Personalized Commercial Products/Services

2.3.8 Auto Billing Systems

2.3.9 Tracking Systems

2.3.10 Interdiction Systems

2.3.11 Subscription Systems

2.3.12 IEM Data Collection Systems

2.3.13 Approval Systems

2.3.14 Forecasting Systems

2.3.15 Financial Systems

2.3.16 IEM Data Phone

2.3.17 Social Network System

3.0 IEM Data Framework Method

4.0 IEM Data Framework Article

5.0 IEM Data Framework System

1.0 Overview

The ingestible event marker (IEM) data framework provides an integrated, seamless solution to enable the collection, management, distribution, and utilization of IEM data. The versatile IEM data framework facilitates integration and implementation of the IEM data with existing data and utilization of the IEM data with existing systems, i.e., commercial systems. The information and communication systems include discrete systems, cross-configured systems, and hybrid systems.

Broadly, various aspects of the IEM data framework include a basic complement of core components, e.g., IEM data; a hub; and at least one IEM data system. Any one or a combination of these core components is capable of interoperating, communication, and/or integration with various components of other information/communication systems. The terms “data” and “information” are used interchangeably herein.

The IEM data include information about an ingestion event, information about a response to the ingestion event, or both. The information about an ingestion event may include, for example, information about the ingestion event of a medication or set of medications. The information about a response to the ingestion event may include, for example, physiologic parameter(s) such as a physiologic status or physiologic change event based on the ingestion event. A physiologic status may be, for example, a heart rate, blood pressure measure, etc., ascertained in close temporal proximity to the time of ingestion of medication (and, therefore, likely to be influenced by or as a result of ingestion of the medication.)

Examples of IEM data include data ingestion time (s) of medication, identification of the type(s) of medication ingested at a particular time, the dosage amounts of medication ingested at a particular time, etc.

Typically, the IEM data may be generated and/or communicated via an ingestible device such as an ingestible event marker (IEM), which generates and communicates data associated with the ingestion event. The IEM may be associated, for example, with a receiver, i.e., a device capable of receiving the IEM data on ingestion and further capable of measuring additional IEM data on response to the ingestion event(s). The IEM and the receiver are discussed in detail hereinafter. In various aspects, the ingestible event data may originate from multiple ingestible event markers. In various aspects, the IEM data may be communicated directly from the IEM to a device other than the receiver, e.g., an IEM business system adapted to receive the IEM data directly from the IEM via a communication channel.

In various aspects, the IEM data may be associated with other data, e.g., combined with data related to events other than an ingestion event or response(s) to an ingestion event. Some examples of other data are data associated with various medical devices and data associated with consumer and personal devices such as intelligent devices/appliances. All are discussed in greater detail hereinafter.

In various aspects, the IEM data may be associated with an IEM data environment and/or commercial systems.

In various aspects, the IEM data may be associated with a unique identifier, e.g., sample data reflective of physiologic patterns associated with a particular individual such as heart rate variability, breathing rate, and/or heart rate (ECG) patterns. For example, a portion or all of the IEM data may be compared with a unique identifier generated by or stored on the receiver.

The hub includes any hardware device, software, and/or communications component(s), as well as systems, subsystems, and combinations of the same which generally function to communicate the IEM data. Communication of the IEM data includes receiving, storing, manipulating, displaying, processing, and/or transmitting the IEM data.

In various aspects, the hub also functions to communicate, e.g., receive and transmit, non-IEM data. Non-IEM data includes non-IEM physiologic data. One example is cardiac data generated by a separate cardiac-related device such as an implanted pacemaker and communicated to the hub directly or indirectly, e.g., via the receiver.

Broad categories of hubs include, for example, base stations, personal communication devices, and mobile telephones.

For example, the hub includes a software application associated with a mobile telephone of a patient. The
application and mobile telephone function to receive IEM data from a receiver, which, in turn, receives the IEM data from an ingestible device ingested by the patient. The hub stores, manipulates, and/or forwards the IEM data, alone or in combination with other data, to an IEM data system.

[0087] The IEM data systems include any hardware device, software, and/or communications component, as well as systems and subsystems of the same, which generally function to provide a service or activity related to the IEM data. The IEM data systems, for example, collect, manipulate, calculate, transmit, receive, store, and/or communicate at least a portion of the IEM data.

[0088] Each IEM data system may be built around pre-defined function(s) or service(s) and may be enabled via the IEM data framework.

[0089] One or more IEM data systems may be integrated, interoperate, intercommunicate or otherwise share or further the collection, management, distribution/dissemination, billing or other activities related to IEM data. One example of an IEM data system is a feedback loop system to refine and optimize IEM data and other data, e.g., medical database data.

[0090] Various aspects of the IEM data framework provide on-demand, accurate and efficient services with respect to provision and utilization of IEM data, while reducing redundancies, errors, and inaccuracies associated with personal event data that are sometimes found in the prior art. Various aspects of the IEM data framework further ensure generation and communication of accurate IEM data in a timely manner.

[0091] Further, the IEM data framework is applicable to any communication environment. Communication environments include any environment having therein, or associated with, data or communication of data.

[0092] Various aspects of the IEM data framework utilize the IEM data, the hub, and one or more IEM data systems to enable useful, secure, and efficient use of the IEM data among multi-profile parties in one or various communication environments.

[0093] FIG. 1 provides a diagrammatic representation of communication environment 100 including an IEM data framework 102, according to one embodiment. The communication environment 100 may further include, for example, an IEM data environment 104 and one or more commercial systems 106.

[0094] Communication environment 100 includes any environment having therein, or associated with, data or communication of data. Communication includes any method, act, or vehicle of communication, and/or combinations thereof. For example, communication methods include manual, wired, and wireless, etc. Wireless technologies include radio signals, such as X-rays, ultraviolet light, the visible spectrum, infrared, microwaves, and radio waves, etc. Wireless services include voice and messaging, handheld and other Internet-enabled devices, data networking, etc.

[0095] Vehicles of communication include the Internet, wired channels, wireless channels, communication devices including telephones, computers, wire, radio, optical or other electromagnetic channels, and combinations thereof, including other devices and/or components capable of associated with communicating data. For example, the communication environments include in-body communications; various devices; various modes of communications such as wireless communications, wired communications, and combinations of the same, etc.

[0096] In-body communications include any communication of data or information via the body, i.e., communication via or associated with inter-body aspects, intra-body aspects, and a combination of the same. For example, inter-body aspect include communications associated with devices designed to attach to a body surface. Intra-body aspects include communications associated with data generated from within the body, e.g., by the body itself or by a device implanted, ingested, or otherwise locatable in, or partially in, the body.

[0097] Communications include and/or may be associated with software, hardware, circuitry, various devices, and combinations thereof.

[0098] The devices include devices associated with IEM data generation, transmission, reception, communication, etc. The devices further include various implantable, ingestible, insertable, and/or attachable devices associated with the human body or other living organisms. The devices further include multimedia devices such as telephones, audio players, and multimedia players.

[0099] Wireless communication modes include any mode of communication between points that utilizes, at least in part, wireless technology including various protocols and combinations of protocols associated with wireless transmission, data, and devices. The points include, for example, wireless devices such as wireless headsets; audio and multimedia devices and equipment, such as audio players and multimedia players; telephones, including mobile telephones and cordless telephones; and computers and computer-related devices and components, such as printers.

[0100] Wired communication modes include any mode of communication between points that utilizes wired technology including various protocols and combinations of protocols associated with wired transmission, data, and devices. The points include, for example, devices such as audio and multimedia devices and equipment, such as audio players and multimedia players; telephones, including mobile telephones and cordless telephones; and computers and computer-related devices and components, such as printers.

[0101] The IEM data framework 102 enables exchange, transmission, receipt, manipulation, management, storage, and other activities and events related to IEM data. Such activities and events may be contained within the IEM data framework 102, partially integrated with the IEM data framework 102, or associated with externalities, e.g., activities, systems, components, and the like which are external to the IEM data framework 102. Externalities include, for example, the IEM data environment 104 and commercial systems 106, either or both of which may also be integral to, or partially integrated with, the IEM data framework 102.

[0102] The IEM data environment 104 includes any source of information or data, including remote computer systems, local computer devices, etc. The information or data may comprise IEM data in whole or in part. The information or data may also be independent of the IEM data, e.g., may be capable of aggregation and/or integration with the IEM data.

[0103] The commercial systems 106 include various existing systems that utilize one or various types of data to accomplish a particular purpose. One example of a commercial system is a computerized pharmacy system utilized in a pharmacy. The computerized pharmacy system may function to automatically, e.g., electronically, receive prescriptions,
verify patient and prescription information, verify insurance coverage, process the prescription order, and generate an invoice.

The IEM data framework, the IEM data environment, and the commercial systems are discussed in greater detail hereinafter.

2.0 IEM Data Framework

FIG. 2 provides a diagrammatic representation of the IEM data framework of FIG. 1, according to one embodiment. The IEM data framework includes an ingestion event, i.e., an act of ingestion. Additionally, the IEM data may include, be included in, or be combined with data from other systems or sources, e.g., medical devices, local or remote computer devices and systems, etc. An example of the IEM data is data having an identification of the type of an ingested medication and the time at which the medication was ingested.

The hub includes any hardware, software, and/or communications component(s) in any combination/configuration, which generally function to communicate the IEM data. One example includes communicating the IEM data to the IEM data systems. For example, the hub receives the IEM data from an ingested device and forwards the IEM data, alone or in combination with other data from other sources, to an IEM data system.

The IEM data systems provide discrete services and/or activities related to the IEM data. The discrete services and/or activities include, for example, propagation of information, data, etc., to a particular user, or group of users, via various system component configurations, etc.

In one example, an auto refill system receives IEM data from the hub. The IEM data include an indication that the last remaining pill of a prescription has been ingested. The auto refill system uses this information to contact a local or remote data resource having refill information, verify the refill information, and automatically transmit a request to a pharmacy system (commercial system) for refill of the prescription.

2.1 IEM Data

The ingestible event marker (IEM) data are associated with at least one of an ingestion event and a response to the ingestion event. The ingestion event may be associated with, for example, data related to and/or gathered during transit through the alimentary system, e.g., oral cavity, pharynx, esophagus, stomach, small intestine, large intestine, anus, etc. Examples of IEM data include an ingestion time, identification of ingested substance, expiration date of an associated medication, dosage of an ingested substance, etc. The information about an ingestion event may include, for example, information about the ingestion event of a medication or set of medications. The information about a response to the ingestion event may include, for example, physiologic parameter(s) such as a physiologic status or physiologic change event based on the ingestion event. A physiologic status may be, for example, a heart rate, blood pressure measure, etc., ascertained in close temporal proximity to the time of ingestion.

In various aspects, the IEM data typically may be generated via one or more ingestible event markers (IEMs), discussed hereinafter in detail. The generation of IEM data via multiple IEMs ensures comprehensive data reporting, e.g., data generated from multiple ingestion events of multiple IEMs over a time interval, data generated from multiple IEMs ingested at approximately the same time, etc. In this manner, comprehensive IEM data may be provided.

In various aspects, the IEM data may be communicated to, i.e., received by, a receiver. The receiver may be embodied in various ways, including an implantable device, a semi-implantable device such as a subcutaneous device, and an externally-applied device such as a personal signal receiver. One example of a personal signal receiver is a “patch” receiver which may be removably affixed to the individual’s person, apparel, etc.

In various aspects, the IEM data can be associated with other data, e.g., a personal event not associated with an ingestion event or a response to an ingestion event. A personal event includes any parameter or circumstance associated with a person, e.g., any event associated with ingestion, inhalation, injection, implantation, insertion, and/or imbibing of a device, substance, liquid, etc. A personal event further includes any event associated with personal data, e.g., a physiologic parameter such weight.

In various aspects, the IEM data may be associated with a unique identifier, e.g., heart rate variability, breathing rate, and/or heart rate (ECG) patterns associated with a particular individual. The unique identifier may be variously embodied. One example is a personal identifier assigned to an individual, e.g., an alphanumeric code, etc. Another example is a unique identifier reflective of an individual trait, such as a physiologic pattern.

To illustrate, a patient may ingest an IEM (discussed hereinafter) integrated with medication. The IEM may communicate IEM data to a receiver such as a patch receiver (discussed hereinafter). The data may include, for example, a unique identifier which may be compared to data associated with the receiver for validation purposes.

In one scenario, the IEMs associated with medication prescribed for a particular patient may each be encoded and deployed with corresponding unique identifiers. The unique identifier may be, for example, a predetermined physiologic data sample associated the particular patient. Various physiologic data samples include a data sample reflective of the particular patient’s heart rate variability, a data sample reflective of the particular patient’s breathing rate, a data sample reflective of the particular patients heart rate (ECG) patterns, etc.

When the receiver is affixed or otherwise associated with an individual, programming logic associated with the receiver may receive actual data samples of the individual, e.g., from data sources such as heart devices, etc. The receiver may communicate the actual data samples received from the data sources and the unique identifier(s) received from the IEM(s) to a computer-related device, e.g., a server, which may compare the actual data samples of the individual with the unique identifier to verify that the medication was actually ingested by the particular patient for whom it was prescribed. In various aspects, predetermined actions based on the verification outcome may be taken, e.g., alerts may be sent to a device associated with the prescribing physician, etc.

2.1.1 IEM Data Environment

In various embodiments, IEM data are generated, received, gathered, etc., from one or a variety of sources and comprise various structures, content, types, etc. The IEM data environment includes at least one of an IEM data source device, products, events, patient specific parameters, IEM
data algorithms, and storage repositories. The sources include, for example, various devices, storage repositories, and systems capable of generating, identifying, gathering or otherwise producing data related to ingestion, the ingestion environment, e.g., the alimentary system of a human subject or non-human subject and/or other personal events. The types include, for example, raw data, processed data, aggregated data, combined data, data from various sources, etc. The processed data include, for example, data processed according to a variety of methods, e.g., algorithms such as IEM data algorithms discussed below.

**[0121]** FIG. 3 illustrates IEM data environment 104 associated with the IEM data framework 102 of FIG. 2, according to one embodiment. The IEM data environment 104 includes, for example, IEM data source devices 300, products 302, events 304, patient specific parameters 306, IEM data algorithms 308, storage repositories 310, and other sources 312.

**[0122]** 2.1.1.1 IEM Data Source Devices

**[0123]** The ingestible event marker (IEM) data source devices 300 include, for example, devices capable of gathering, collecting, generating, receiving, storing and/or transmitting, etc., IEM data. One example of such a device is a microchip capable of or otherwise enabling or facilitating the collection, generation, receipt, transmission, etc., of data. Such a microchip may be integrated or associated with the IEM data source devices 300. The IEM data source devices 300 may be embodied, for example, as ingestible devices 300a, receivers 300b, and/or health devices 300c.

**[0124]** In various aspects, IEM data may be related to various devices. For example, a device may be an ingestible device, an inhalable device, an injectable device, an implantable device, an implantable device, and an imbedding device. The foregoing may be embodied, for example, as a microchip alone or in combination with other structural components, each capable of at least one of ingestion, inhalation, injection, implantation, insertion, and imbedment by a human body or a non-human body.

**[0125]** The ingestible device may comprise, for example, a microchip. The microchip may be independently deployed. The microchip may also be attached to, embedded in, or otherwise integrated with a device. The microchip may also be attached to, embedded in, or otherwise integrated with a device. The ingestible device is capable of ascertaining parameter(s) associated with ingestion, e.g., measuring or tallying doses of an ingestible. The ingestible device may also comprise, for example, an ingestible microchip used to ascertain parameter(s), e.g., ingestion time, identify an ingested substance, etc.

**[0126]** The injectable device may comprise, for example, a microchip. The microchip may be independently deployed. The microchip may also be attached to, embedded in, or otherwise integrated with a device. The injectable device is capable of ascertaining parameter(s) associated with injection, e.g., time of injection, identification of an injected substance, etc. In various aspects, the injectable device is capable of injection into a human body or a non-human body, e.g., injection into the alimentary system of a human body.

**[0127]** The implantable device may comprise, for example, a microchip. The microchip may be independently deployed. The microchip may also be attached to, embedded in, or otherwise integrated with a device. The implantable device is capable of ascertaining parameter(s) associated with implantation, e.g., time of implantation, physiologic parameters such as heart rate, EKG data, activity management data, temperature, galvanic skin response data, respiratory data, fluid status data, heart rate variability, etc.

**[0129]** In one aspect, the implantable device is embodied as an implantable receiver, supra, for receiving various data. The implantable receiver may also process, store, transmit, etc. the data. Various other implantable devices include, for example, heart monitors and the like having a microchip to ascertain parameter(s), e.g., heart rate, heart pressure, etc.

**[0130]** The ingestible device may comprise, for example, a microchip. The microchip may be independently deployed. The microchip may also be attached to, embedded in, or otherwise integrated with a device. The ingestible device is capable of ascertaining parameter(s) associated with ingestion, e.g., time of ingestion, physiologic parameters such as environmental content/fluid identification, etc. In one aspect, the ingestible device is embodied as a microchip mechanically associated with a suppository for rectal insertion, vaginal insertion, etc.

**[0131]** The imbibable device may comprise, for example, a microchip. The microchip may be independently deployed. The microchip may also be attached to, embedded in, or otherwise integrated with a substance, e.g., a potable solution or fluid such as a beverage, etc. The imbibable device is capable of ascertaining parameter(s) associated with imbibing, e.g., time of drinking, physiologic parameters such as environmental content/fluid identification, etc. In one aspect, the imbibable device is embodied as a microchip and imbibed together with a beverage. The beverage may aid in swallowing, may be used as a medication, etc.

**[0132]** Further, the IEM data may be associated with administration of a therapeutic agent, etc. For example, administration includes, but is not limited to, parenteral administration, i.e., administration in a manner other than through the alimentary system, such as by intravenous or intramuscular injection or inhalation.

**[0133]** In some aspects, the devices are capable of ingestion, i.e., entry into the alimentary system of a human body or a non-human; inhalation (either the device or a substance associated with the device, e.g., a nasal inhalant). In various aspects the devices are capable of injection, insertion, implantation and/or imbibing, etc., into/within a human body or a non-human body.

**[0134]** The ingestible devices 300g gather/collect/generate IEM data via various methods, e.g., ingestion timing, contact with alimentary system substances, sampling, etc. Further, various ingestible event marker data source devices 300 communicate the IEM data via various methods, e.g., wireless methods, conductive methods via body tissue, etc. The following are examples of the ingestible devices 300g.

**[0135]** A pharma-informatics system described in PCT/US2006/016370, filed Apr. 28, 2006, includes compositions, systems and methods that allow for the detection of the actual physical delivery of a pharmaceutical agent to a body are provided. Embodiments of the compositions include an identifier and an active agent.

**[0136]** An IEM system described in PCT/US2008/52845, filed Feb. 1, 2008, includes an ingestible event marker (IEM) and a personal signal receiver. Aspects of the IEM include an identifier, which may or may not be present in a physiologically acceptable carrier. The identifier is characterized by being activated upon contact with a target internalphysiolog-
cal site of a body, such as digestive tract internal target site. The personal signal receiver is configured to be associated with a physiological location, e.g., inside of or on the body, and to receive a signal of the IEM. During use, the IEM broadcasts a signal which is received by the personal signal receiver.

[0137] The IEM data associated with the IEM system include personal data, e.g., physiologic data generated by the IEM. Examples are derived metrics, e.g., processed physical data to derive various metrics such as time of ingestion data; combined metrics, e.g., derived metrics combined with other derived metric data such as time of ingestion data combined with data identifying the ingested substance; and IEM data, e.g., derived metrics and/or combined metrics aggregated with various physiologic data such as time of ingestion data combined with data identifying the ingested substance and physiologic data such as ECG data, temperature, etc.

[0138] A controlled activation ingestible identifier described in PCT/US07/82563, filed Oct. 17, 2007, includes ingestible compositions such as pharma-informatics enabled compositions. The controlled activation ingestible identifiers include a controlled activation element that provides for activation of the identifier in response to the presence of a predetermined stimulus at a target site of interest.

[0139] A life cycle pharma-informatics system described in U.S. Patent Application Ser. No. 61/034,085, filed Mar. 5, 2008 includes RFID and conductive communications technology combined with medication and/or medication packaging such that the medication can be tracked for the duration of its transmissions while addressing the potential privacy and signal degradation concerns associated with RFID technology.

[0140] The IEM data receivers 300b include devices capable of receipt of IEM data 200. Receipt may be, for example, via wireless or wired channels, etc. The IEM data receiver 300b may also transmit or otherwise forward data. In various aspects, the IEM data receiver 300b may perform, facilitate, or enable various other functionalities related to the IEM data 200 and/or other data. In various aspects, the IEM data receiver 300b may be attachable, implantable, semi-implantable or otherwise associated with a human body or a non-human body.

[0141] The IEM data receiver 300b include personal signal receivers such as patch receivers, e.g., removable attachable externally to a human body or a non-human body; subcutaneous devices; implantable devices external devices, i.e., devices which are not designed for attachment on other permanent or semi-permanent contact with the body, e.g., a mobile telephone. The following are examples of the IEM data receiver 300b.

[0142] The IEM system, PCT/US2008/52845, supra, includes an ingestible event marker (IEM) and/or a personal signal receiver.

[0143] An active signal processing personal health signal receiver described in PCT/US07/24225, filed Nov. 19, 2007, includes a receiver associated with a body, e.g., located inside or within close proximity to a body, configured to receive and decode a signal from an in vivo transmitter which is located inside the body.

[0144] The health devices 300c include multiple devices (and methods associated with the devices) associated with the IEM data 200. The health devices 300c, for example, may gather, collect, aggregate, store, transmit, receive, or otherwise communicate data, including the IEM data 200.

[0145] Communication may be, for example, via wireless or wired channels, etc. The IEM data receiver may also transmit or otherwise forward data. In various aspects, the IEM data receiver 300b may perform, facilitate, or enable various other functionalities related to the IEM data 200 and/or other data. Examples include functions to store data, process data, etc.

[0146] In various aspects, the health device 300c may be attachable, implantable, semi-implantable or otherwise associated with a human body or a non-human body. For example, "intelligent" devices such as intelligent scales, intelligent blood pressure cuffs, intelligent refrigerators, etc., may be integrated in various configurations. As used herein, the term "intelligent devices" refers to one or more devices capable of generating and/or communicating data, e.g., wirelessly transmitted data, via a communication channel to a destination.

[0147] 2.1.1.2 Products

[0148] IEM data 200 also includes IEM data related to products 302. The products 302 include, for example, an ingestible device/pharmaceutical product 302a. One example of an ingestible device/pharmaceutical product 302a is an IEM mechanically associated with medication. The IEM may be mechanically associated with the medication in various ways, including external to the medication, partially integrated with the medication, and wholly integrated with the medication.

[0149] The IEM may be affixed via various means, e.g., with various adhesive or formulated substances. The IEM may be associated with the medication at various phases, e.g., during a medication manufacturing process, at various points in time after a medication manufacturing process, etc.

[0150] 2.1.1.3 Events

[0151] IEM data 200 further includes data related to events 304, e.g., personal events, event parameters, etc. Further examples include time of ingestion of a medication, dosage and identity of medication taken at time of ingestion, etc. Events may include physiologic events, e.g., respiration rate; environmental events, e.g., time of day; usage events, e.g., ingestion of a medication, use of a cardiac resuscitation device, etc.

[0152] 2.1.1.4 Patient Specific Parameters

[0153] IEM data 200 still further includes data related to patient specific parameters 306, e.g., individualized patient data 306a pertaining to an individual patient and multiple patient data 306b pertaining to multiple patients. Examples of patient specific parameters include physiologic data, etc. Multiple patient data include aggregated patient data, patient population data, e.g., combined patient data which includes various predetermined aspects of data regarding at least one patient and excludes data tending to identify a particular patient or an aspect in which the patient has a privacy interest, e.g., name, age, diagnosis and/or other data which the patient wishes to retain as confidential and/or undisclosed to the public.

[0154] 2.1.1.5 IEM Data Algorithms

[0155] IEM data 200 also includes data related to IEM data algorithms 308, e.g., raw data, processed data, or a combination of the same, which undergo processing. In one example, the IEM data 200 have one or more algorithms applied thereto, with processed data as an output. The data, for example, includes individualized patient data 306a and multiple patient data 306b, e.g., patient population data.
The IEM data algorithms may be related to aspects such as data processing associated with the IEM data generated by one or more ingestible devices, e.g., an IEM system.

With respect to IEM data processing associated with an ingestible device, aspects include, for example, transmission of the IEM data generated by the IEM data processing associated with a receiver, and IEM data post-processing aspects.

Transmission aspects of IEM data and algorithms may include, for example, modulation schemes, coding, and error code aspects.

The transmission aspects include, for example, analog, digital, spread spectrum, combinatorial, and contention avoidance.

The analog transmission aspects include, for example, amplitude modulation, single sideband modulation, frequency modulation, phase modulation, quadrature amplitude modulation, and space modulation methods, etc.

The digital transmission aspects include on/off keying, frequency-shift keying, amplitude-shift keying, phase-shift keying, e.g., binary phase-shift keying, quadrature phase-shift keying, higher order and differential encoded, quadrature amplitude modulation, minimum shift keying, continuous phase modulation, pulse-position modulation, trellis coded modulation, and orthogonal frequency-division multiplexing.

The spread spectrum transmission aspects include, for example, frequency hopping spread-spectrum and direct-sequence spread spectrum.

The combinatorial transmission aspects include, for example, binary phase shift-keying with carrier frequency modulation.

The contention avoidance transmission aspects include, for example, duty-cycle modulation and carrier frequency modulation.

The coding aspects include, for example, wake-up schemes, preamble schemes, data packet schemes, and error code schemes.

The wake-up schemes include, for example, multitone schemes and chirp schemes.

The preamble schemes include, for example, unique identifier for packet start schemes.

The data packet schemes include, for example, data related to pill type, pill expiration, manufacturer, lot number, amount, prescribing physician, pharmacy, etc.

The error code schemes include, for example, repetition schemes, parity schemes, checksums, cyclic redundancy checks, hamming distance schemes, and forward error correction schemes, e.g., Reed-Solomon codes, binary Golay codes, convolutional codes, turbo codes, etc.

With respect to IEM data processing and the receiver, considerations may be given to, for example, position, energy conservation schemes, carrier identification, decoding and error correcting.

The position of the receiver includes, for example, the stomach, the side and the xiphoид.

The energy conservation schemes include schemes for a periodic wake-up, e.g., to sense IEM wake-up such that energy, e.g., battery resources, is conserved during non-awake periods.

The carrier identification aspects include, for example, Fourier transform analysis, e.g., fast Fourier transform and discrete Fourier transform, phase locked loop, filter bank, match filter, and combinatorial such as use of previous knowledge about frequency to tune-in.

The decoding aspects and error correcting aspects include, for example, the above-iterated aspects.

With respect to IEM data post-processing, aspects include, for example, pill detection, e.g., multiplicity of identification and count in time aspects, adherence metrics, etc.

With respect to IEM data processing associated with physiologic parameter metrics, aspects include, for example, electrocardiogram (EKG or ECG), impedance, acceleration, optical, pressure, temperature, sound, biochemical/biological, weight, position, derived electromyography (EMG), and electroencephalography (EEG).

IEM data processing related to EKGs includes, for example, compression data, e.g., wavelet and ICA/PCA, R-wave detection such as Hamilton-Tompkins, etc., heart-rate variability, e.g., SDNN, standard deviation in a 24 hour period, standard deviation of consecutive five minute periods, foot print heart rate versus standard heart rate, distribution-based histogram, etc., arrhythmia, and respiration, e.g., principal axis modulation.

IEM data processing related to impedance includes, for example, respiration, fluid status, Galvanic skin response, blood flow, etc.

IEM data processing related to acceleration, includes, for example, direct acceleration, which includes total activity and derived acceleration, which further includes activity type.

IEM data processing related to optical includes, for example, hematocrit, O2 saturation, pulse oximetry, etc.

IEM data processing related to temperature includes, for example, body temperature, heat flux, etc.

IEM data processing related to sound includes, for example, heart sounds, valvular events, etc.

IEM data processing related to biochemical/biological includes, for example, lactose, glucose, antibody, biomarker, bacterial, osmolality, etc.

IEM data processing related to derived data includes, for example, sleep, total energy, etc.

2.1.1.6 Storage Repositories

Ingestible event marker data also includes data related to storage repositories i.e., databases and/or other storage implementations that temporarily and/or permanently retain, store, etc., data related to IEM data, including data to be combined or aggregated with ingestible event marker data.

Storage may be in any form or format, as is known or will be known in the future. In various aspects, the storage repositories may be independently embodied and/or may be partially or wholly integrated with computer-related system(s). The storage repositories, for example, may operate or otherwise be associated with various computer systems, software, hardware, communication components, etc. For example, the storage repositories may be part of a medical office computer system and may contain IEM data related to a particular patient's medication regimen. At various times, e.g., scheduled or ad hoc, various IEM data embodied as medical data may be communicated to/from the storage repositories and/or from/to various points/components.

In another illustration, methods, systems and compositions that allow for treating a patient according to a patient customized therapeutic regimen are described in PCT/US2007/1068, filed May 2, 2007, which include obtaining...
dosage administration information from a patient and using the same to tailor a therapeutic regimen for the patient, as well as preparing and forwarding to the patient physical pharmaceutical dosages based on the customized therapeutic regimen. The dosage administration information from the patient may be stored, for example, on the database 306. The IEM data 200 containing information about the ingestion time of a particular medication can be combined with the dosage administration information to customize the therapeutic regimen.

In various aspects, various other IEM data sources 312 are included. Further, it is noted that data and/or IEM data 200 from multiple sources can be aggregated, integrated, refined, etc. via a variety of methods. To illustrate, IEM data 200 such as ingestion data related to ingestion of a medication are generated from an IEM data source device 300 such as the IEM system. The ingestion data are wirelessly transmitted to an IEM receiver.

Concurrently or in an alternative time period, physiologic data such as cardiac parameters are generated by a health device 300c such as the system for monitoring and treating hemodynamic parameters, supra, is generated and wirelessly transmitted to the IEM data receiver 300b. The IEM data 200 and the cardiac physiologic data are aggregated for onward communication to an IEM data system such as an auto refill system.

To illustrate, cardiac data is derived via various methods and systems. One example is continuous field tomography, e.g., electrical tomography (ET). One continuous field tomography method is described in the U.S. Patent Application Ser. No. 60/797,403, filed May 2, 2006. The cardiac data includes cardiac-related parameters, as well as clinical data for clinical applications. Using ET, various cardiac parameters are measured, such as stroke volume, ejection fraction, dP/dt (max), strain rate (max), peak systolic mitral annular velocity, end systolic volume, end diastolic volume, and QRS length, etc. The cardiac measurements may be used to derive or infer various performance and wellness diagnostics/inferences. For example, an ejection fraction parameter may be used as a basis to predict ventricular synchrony performance.

The metrics generated from the continuous field tomography include, for example, velocity, acceleration, and displacement.

The clinical data derived from the metrics include, for example, left ventricle stiffness as well as ET proxies for other physiologic parameters such as ejection fraction (EF) and dP/dt.

In various aspects, the clinical data may be combined with the IEM data to provide additional information. The information may be useful, for example, in various diagnostic and analytical pursuits. Comprehensive patient-related data displayed having clinical data and IEM data are described in the U.S. Patent Application Ser. No. 61/076,577, filed Jun. 27, 2008, wherein various ET physiologic parameters and derivations such as EF and ventricle stiffness are displayed together with IEM data such as medication ingestion time. From such a display, the efficacy of the medication therapy may be gauged.

Hub 202 includes any hardware device, software, and/or communications component(s), as well as systems, subsystems, and combinations of the same which generally function to communicate the IEM data 200, including receiving, storing, manipulating, displaying, processing, and/or transmitting the IEM data 200.

In various aspects, the hub 202 receives, generates, communicates, and/or transmits, the IEM data 200, alone or in combination with other data, i.e., non-IEM data from various sources. Non-IEM data includes non-IEM physiologic data. Examples of non-IEM data include heart rate, heart rate variability, respiration, physical activity level, wake patterns, temperature, etc.

Communication of the IEM data 200 to and from the hub 202 includes any transmission means or carriers, and combinations thereof, including wireless, wired, RF, conductive, etc. as is known in the art or as may become available in the future.

FIG. 4 illustrates the hub 202 associated with the IEM data framework 102 of FIG. 2, according to one embodiment. The hub 202 comprises various categories of devices, e.g., personal communication devices, base stations, and mobile telephones.

Personal communication devices include, for example, devices having communication and computer functionality and typically intended for individual use, e.g., mobile computers, sometimes referred to as "handheld devices".

Base stations comprise any device or appliance capable of receiving data such as IEM data. Examples include computers, such as desktop computers and laptop computers, and intelligent devices/appliances.

Intelligent devices/appliances include consumer and home devices and appliances that are capable of receipt of data such as IEM data. Intelligent devices/appliances may also perform other data-related functions, e.g., transmit, display, store, and/or process data. Examples of intelligent devices/appliances include devices and appliances having refrigerators, weight scales, toilets, televisions, door frame activity monitors, bedside monitors, bed scales. Such devices and appliances may include additional functionality such as sensing or monitoring various physiologic parameters, e.g., weight, heart rate, etc.

Mobile telephones include telephonic communication devices associated with various mobile technologies, e.g., cellular networks.

In one aspect, the hub 202 includes an IEM data receiver embodied, for example, as a receiver such as a patch receiver 400; a personal communication devices such as a handheld device 402; a base station 404; and a mobile telephone 406.

The patch receiver 400 includes, for example, devices capable of at least receiving data, signals, etc. Patch receivers 400 may be attachable, e.g., permanently or removably attachable externally to a human body or a non-human body. For example, the patch receiver 400 may include a receiver and an adhesive layer to provide for attachment to and removal from a region of skin. Alternatively, the patch receiver 400 may be implantable or semi-implantable, e.g., subcutaneous implantation. One such removably attachable patch receiver 400 is the personal signal receiver of the IEM system described in PCT/US2008/52845, supra.

The handheld device 402, also referred to as a "mobile computer", includes, for example, computing devices having computer-related functionality, e.g., typically having a display screen with touch input functionality, a miniature keyboard, etc. Types of handheld devices include, for
example, a personal digital assistant (PDA) having the input and output combined into a touch-screen interface; and enterprise digital assistants offering integrated data capture devices like bar code, radio frequency identification (RFID), and smart card readers, etc.

[0208] In various aspects, the handheld device 402 includes software, e.g., a software agent/application, associated with the IEM data 200. In various embodiments of the handheld device 402, the software is preconfigured, i.e., configurable by the manufacturer/retailer; configurable by the consumer, i.e., downloadable from a website; or a combination of the same.

[0209] One example of software is an auto refill application related to or integrated with an auto refill system to facilitate automated prescription refill functions.

[0210] The base station 404 includes systems, subsystems, devices, and/or components that receive, transmit, and/or relay the IEM data 200. In various aspects, the base station communicates interoperates with a receiver such as the patch receiver 400 and a communications network such as the Internet. Examples of base stations 404 are computers, e.g., servers, personal computers, desktop computers, laptop computers, intelligent devices/appliances, etc., as heretofore discussed.

[0211] In various aspects, the base station 404 may be embodied as an integrated unit or as distributed components, e.g., a desktop computer and a mobile telephone in communication with one another and in communication with a patch receiver and the Internet.

[0212] In some aspects, the base station 404 includes the functionality to wirelessly receive and/or wirelessly transmit data, e.g., IEM data 200 received from and transmitted to the patch receiver 400 and the Internet.

[0213] Further, in various aspects, the base station 404 may incorporate and/or be associated with, e.g., communicate with, various devices. Such devices may receive, generate, receive, and/or communicate data, e.g., IEM data 200. The devices include, for example, clock radios, intelligent pill dispensers, pill managers, e.g., devices capable of receiving various substances and producing a combined substance, dose(s) of substances, etc., pharmaceutical compounding devices, “intelligent” devices such as scales, blood pressure measurement devices, exercise equipment, e.g., tread mills. Further examples include body weight sensors, motion sensors, position sensors, e.g., bed sensors, chair sensors, portals in doorways, refrigerator and food devices, bathroom facilities devices, etc.

[0214] The mobile telephone 406 includes, for example, devices such as a short-range, portable electronic device used for mobile voice or data communication over a network of specialized cell site base stations. The mobile telephone 406 is sometimes known as or referred to as “mobile”, “wireless”, “cellular phone”, “cell phone”, or “hand phone (HP)”.

[0215] In addition to the standard voice function of a telephone, various embodiments of mobile telephones may support many additional services and accessories such as short message service (SMS) for text messaging, email, packet switching for access to the Internet, java gaming, Bluetooth (short range data/voice communications), infrared, camera with video recorder, and MMS for sending and receiving photos and video. Some embodiments of mobile telephones connect to a cellular network of base stations (cell sites), which is, in turn, interconnected to the public switched telephone network (PSTN) or satellite communications in the case of satellite phones. Various embodiments of mobile telephones can connect to the Internet, at least a portion of which can be navigated using the mobile telephones.

[0216] In various aspects, the mobile telephone 406 includes software, e.g., a software agent/application, associated with the IEM data 200. One example is an auto refill application related to or integrated with an auto refill system to facilitate automated prescription refill functions. In various embodiments of the mobile telephone 406, the software is preconfigured, i.e., configurable by the manufacturer/retailer; configurable by the consumer, i.e., downloadable from a website; or a combination of the same.

[0217] Further, various embodiments of the hub ensure privacy requirements via predetermined methods, e.g., an IEM data source device 300 ingested by an individual transmits sensitive IEM data 200 via body tissues to an IEM data receiver 302 embodied in a patch receiver 400 removably attached to the individual’s body. Signals associated with the sensitive IEM data 200 remain undetectable beyond the individual’s body. Once received by the patch receiver 400, various computing components of the patch receiver 400 cleanse and/or encrypt the IEM data 200 for onward secure transmission. In this manner, breaches of sensitive data transmissions and/or unauthorized access to the sensitive data are avoided.

[0218] Further, various aspects of the hub include combinations of devices. One such combination is an IEM data receiver 300b such as the patch receiver 400 in communication with the handheld device 402 or the mobile telephone 406. Thus, for example, the patch receiver 400 wirelessly transmits IEM data 200 to the mobile telephone 406 having a receiver and a software agent available thereon. The receiver of the mobile telephone 406 receives the IEM data 200. A software agent, e.g., an application, processes the ingested reported data 200 and displays various information related to the IEM data 200 via, for example, a customized graphical user interface (GUI). In some aspects, the software agent generates displays with a predetermined “look and feel”, i.e., recognizable to a user as belonging to a predetermined group of software programs, GUIs, source devices, communities, etc.

[0219] To illustrate the foregoing, the IEM data 200 may include data about an ingested medication. Once received by the mobile telephone 406, the software agent may compare the data about the medication to a predetermined medication regimen. Upon verification that the proper medication has been ingested at the proper time, the software disables an audible alarm scheduled to alert the individual to take the (already ingested) medication, thus averting an unnecessary reminder and removing the annoyance associated therewith. The software agent, via the GUI, displays a standard message to the individual notifying of the medication ingested and the time of the next dosage.

[0220] Additionally, the software agent may include functionality to generate of facilitate a financial transaction. In one example, upon occurrence of a certain event, such as verification that the proper medication has been ingested at the proper time, the software agent generates a predetermined charge for the ingested medication, the verification service, or both. The charge is transmitted to a financial system, e.g., the patient’s cell phone transmits the charge via an IEM data system to a computer system associated with the patient’s financial institution where the charge is automatically applied against a financial account of the patient.
In various other aspects, the transaction model may be based on various parameters. In one example, a transaction is associated with a time based model wherein use of a product or service is charged according to the length of time the product or service is used. In another example, a transaction is associated with a measured value delivery, wherein the value of the product or service is metered, measured, or otherwise valued and charged according to the ascertained value at predetermined time intervals. In still other example, a transaction is associated with therapy delivery, i.e., delivery of a therapeutic substance, event, service, etc. Examples of therapeutic substances include medication. Examples of therapeutic events include cardiac defibrillation acts and cardiac resynchronization acts. Examples of therapeutic services include administration of therapeutics, therapeutic consultations, etc.

The IEM data systems include any hardware component, software component, and/or communications component, as well as networks, systems, and subsystems of the same, which generally function to provide a service, function, activity, etc. related to the IEM data. The IEM data systems, for example, collect, manipulate, calculate, transmit, receive, store, and/or otherwise communicate at least a portion of the IEM data.

Each IEM data system is built around a predefined business function or service and is enabled via the IEM data framework. One or more IEM data systems may be integrated, interoperate, intercommunicate or otherwise share or further the collection, management, distribution, dissemination, billing and/or other activities related to IEM data.

Further, one or more IEM data systems may be associated with one or more commercial systems. For example, one or more IEM data systems may be integrated with, interoperate with, and/or intercommunicate with one or more commercial systems. One or more IEM data systems may otherwise share or further the IEM data related activities with one or more commercial systems.

The IEM data systems include at least one component, e.g., hardware device, software, and/or communications component, which generally function to provide a service or activity related to the IEM data, e.g., a computer to receive IEM data from the hub and display the IEM data in conjunction with other information.

Examples of components include a computer, a receiver, a transmitter, an application, a software module, a data storage medium, a processor, a memory component, a personal communication device, software, a communication link, and a handheld device. It is noted that two or more IEM data systems can cooperatively or independently use one or more of the same components. For example, an auto refill system and an approval system can each access a data storage medium having IEM data related to patients and prescriptions and can each utilize the IEM data for predetermined purposes.

FIG. 5 illustrates exemplary IEM data systems associated with the IEM data framework of FIG. 2, according to one embodiment. The exemplary IEM data systems include, for example, feedback loop systems, decision support systems, auto refill systems, patient tools, behavioral medicine systems, incentive systems, personalized commercial products/services, auto billing systems, tracking systems, interdiction systems, subscription systems, IEM data collections, approval systems, forecasting systems, financial systems, an IEM data phone system, and social networks.

Feedback loop systems aggregate various sources of data, e.g., IEM data, analyze the aggregated data, and/or provide feedback information to multiple profile recipients based on the aggregation/analysis.

FIG. 6 illustrates an exemplary IEM data framework including a feedback loop system, according to one embodiment. The feedback loop system includes, for example, server having application and database. The IEM data framework further includes IEM data and the hub, embodied here as the mobile phone. In various aspects, the feedback loop system may interoperate, or be otherwise associated with, one or more IEM data systems and/or one or more commercial systems.

In one scenario, a patient ingests medication having an ingestible device integrated therein. The ingestible device generates IEM data in the form of medication identification and time of ingestion information. The ingestible device transmits the information to a receiver. The receiver, in turn, communicates the information to the hub embodied as a mobile telephone associated with the patient.

A software agent resident on the mobile telephone aggregates the received medication identification and time of ingestion information with the blood pressure measurement information and forwards the aggregated data to the feedback loop system. The feedback loop system having server, software, database, receives the aggregated data from the mobile telephone and, via the software, compares the aggregated data to patient information in the database to determine if the patient took the most recent dose of medication in a timely manner, and if the patient has consistently taken the medication in a timely manner, and if the blood pressure measurement coincides with an acceptable range of blood pressure measurements.

Based on an analysis of the data, the feedback loop system generates additional IEM data in the form of a decision on patient adherence and a decision on treatment efficacy. The IEM data decisions are stored in database for future reference and forwarded to a commercial system such as a healthcare system associated with a medical center computer system and having patient data such as physician’s medication instructions, etc.

The healthcare system facilitates automatic processing and feedback, enables accessibility to the IEM data, e.g., by a healthcare provider, enables data input, e.g., healthcare instructions by the healthcare provider, etc.

For example, the healthcare system compares the decision data received from the feedback loop system with stored healthcare providers instructions, e.g., medication regimen adherence is satisfactory and no action is needed at this time; medication regimen adherence is not satisfactory and action is needed at this time; medication regimen is satisfactory but action is needed at this time, e.g., titration is needed, etc., and generates the comparison result data for review by the healthcare provider.

The healthcare provider utilizes the information to advantageously adjust patient treatment parameters, e.g., prescription and dosage requirements. The healthcare provider
inputs data based on the comparison results, e.g., the adjusted treatment parameters. The input data are processed by the healthcare system 106a and forwarded to the feedback loop system 204a. The feedback loop system 204a receives the feedback loop data, reconciles the feedback loop data with the patient information resident in the database 504, and forwards the notification to the mobile telephone 406 of the patient 506.

[0238] In various aspects, the feedback loop system 204a and/or the healthcare system 106a cooperate, e.g., communicate with at least one other IEM data system 204 and/or commercial system 106.

[0239] To continue the foregoing illustration, in addition to forwarding the adjusted medication regimen instructions to the patient’s mobile telephone 406, either the feedback loop system 204a or the healthcare system 106a forwards the adjusted medication regimen in the form of a prescription to a commercial system such as a pharmacy system 106b for refill. The pharmacy system 106b fills the prescription and communicates a message to the feedback loop system 204a notifying of the same. The feedback loop system 204a updates the patient’s data in database 504 to reflect the new prescription and fulfillment of the prescription, and communicates the notification to the patient’s mobile telephone 406.

[0240] 2.3.2 Decision Support Systems

[0241] Decision support systems, e.g., personal wellness systems, may generate, store, provide data, e.g., IEM data, which may be used to inform and support decisions, e.g., stakeholders’ decisions. In one example, multiple instances of individualized ingestible event marker data and physiologic data are gathered and combined into anonymized patient population data. Pharmaceutical research and development groups, universities, etc., utilize the data for various purposes, e.g., information to formulate new product lines, adjust existing therapies, etc. The data may be accessed, for example, by subscription to population data feeds, access to the database, etc.

[0242] FIG. 7 illustrates an exemplary IEM data framework 102 having a decision support system 204b, according to one embodiment. The IEM data framework 102 further includes IEM data 200 and the hub 202, shown here embodied as the mobile telephone 406. In various aspects, the feedback loop system 204a may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106.

[0243] In one scenario, IEM data, e.g., IEM data 200a and IEM data 200b, related to multiple individuals, e.g., patient 506a and patient 506b, respectively, are communicated via the hubs, e.g., mobile telephone 406a and mobile telephone 406b, respectively, to the decision support system 204b comprising, for example, server 500, software 502, and database 504. The IEM data 200a and 200b may be encrypted. The decision support system 204b processes and stores the received data. For example, software 502 anonymizes the patient data, i.e., removes all aspects of the data tending to identify an individual and removes, according to a predetermined scheme, all aspects of the data designated as private, sensitive, confidential in nature, etc. The software 502 may provide various other functions such as integrating the anonymized patient data with existing patient population data in the database 504.

[0244] The integrated data in database 504 may be accessed by, delivered to, or otherwise utilized by multiple systems and parties. Such systems include, for example, commercial systems 104 such as pharmaceutical systems 106c and university systems 106d. Parties associated with the pharmaceutical systems 106c may utilize the patient population data, for example, for statistical analysis and projective capabilities such as determining the efficacy, cost efficiency, profit, etc. of a particular medication and projecting from the determination new product line concepts/therapies, etc. Parties associated with universities may utilize the patient population data to research symptomatology, analyze medication risks, etc.

[0245] In various aspects, the decision support system 204b, IEM data system(s), and/or commercial system(s) interoperate, e.g., communicate, therewith.

[0246] To continue the foregoing illustration, in addition to the provision of decision support data such as patient population data, the decision support system 204b communicates patient population data to the feedback loop system 204a. The feedback loop system 204a communicates the patient population data to mobile telephone 406a of patient 506a.

[0247] In one scenario, the decision data derived from a patient population such as medication efficacy may be correlated with an individual’s medication therapy, and communicated via marketing system specifically targeted for that individual.

[0248] 2.3.3 Auto Refill Systems

[0249] Auto refill systems automatically fill or refill prescriptions. In one example, IEM data identifying an ingested medication are gathered and reconciled with current prescription information to identify depleted prescription supplies. If the supply is depleted, a refill order is automatically triggered to the appropriate pharmacy. The pharmacy automatically refills the order, generates a bill, and charges the appropriate account, e.g., via a real-time, online financial transaction.

[0250] FIG. 8 illustrates an exemplary IEM data framework 102 having an auto refill system 204c, according to one embodiment. The IEM data framework 102 further includes IEM data 200 and the hub 202, shown here embodied as the base station 404. In various aspects, the auto refill system 204c may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106.

[0251] In one scenario, the patient 506 ingests prescription medication in conjunction with an ingestible device. The ingestible device identifies the medication type and dosage, and transmits the IEM data 200 via, for example, conductive transmission to the patch receiver 400, which may be removably attached to the patient 506. The patch receiver 400 transmits the IEM data 200 to base station 404. The base station 400 forwards the IEM data 200 to the auto refill system 204c. The software 502 of the auto refill system 204c compares the medication type and dosage of the IEM data 200 against prescription information stored in the database 504. The prescription information, for example, may include the number of tablets in the prescription at time of fill, the dosage instructions, and a running total of the ingested tablets as per previously received information. If the comparison indicates depletion of the prescription medication, database 504 is checked for the number of remaining refills. If refills are remaining, any sensitive data of the IEM data 200 are cleansed, i.e., removed, and a prescription refill request with pertinent information is compiled and transmitted according to predetermined security protocol and via predetermined channel(s) to a commercial system 106 such as the pharmacy system 106b. Upon receipt by the pharmacy system 106b, the refill request is parsed and verified, and the prescription is refilled.
Payment for refill can be effected, for example, via a real-time, online transaction between the pharmacy system 106b and an IEM data system 204 and/or commercial system, e.g., financial transaction system 106c. The financial transaction system 106c, for example, may receive the financial transaction, e.g., prescription refill charge, via a predetermined communication channel. The financial transaction system 106c verifies the patient account information and completes the transaction, notifying the pharmacy system 106b.

Notification of status of refill and payment for refill can be provided via predetermined communication channel(s) to the base station 300, e.g., an email for display on the laptop computer, a text message to the patient’s mobile telephone, etc.

2.3.4 Patient Tools

Patient tools include any data, information, software, websites, etc. that provide information or assist a particular patient focus, e.g., tracking tools to assist a patient in cardiac health management, patient personalization of their own data, etc. Various users may be associated with the patient tools. Examples include various users within a patient community, e.g., patients, family caregivers, and professional caregivers such as physicians.

FIG. 9 illustrates an exemplary IEM data framework 102 having a tools 204d, according to one embodiment. The IEM data framework 102 further includes IEM data 200a, and the hubs, shown here embodied as the base station 404, the mobile telephone 406, and the handheld device 402. In various aspects, the patient tools 204d may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106.

In one scenario, multiple parties such as patients 506a-c access the patient tools 204d, which may be embodied as the server 500 having the software 502 and the database 504 having IEM data 200 in the form of at least patient tools. Patients 506a-c may access the patient tools 204d. For example, via the base station 404, the mobile telephone 406, and the handheld device 402, respectively.

Patient 506b may search the database 504 for patient tools related to mental illness management. The patient tools, for example, may be provided in the form of downloadable data/applications to assist in tracking, monitoring, diagnosing, and notifying a patient of a relevant health issue, e.g., medication dosage schedule, etc. Patient 506b may download the application onto, for example, the mobile telephone 406. Patient 506b may further communicate via, for example, the mobile telephone 406 with at least one commercial system such as the healthcare system 106a, which may provide further medical data, instruction, etc., relevant to the patient 506b’s mental illness management pursuit.

In various aspects the patient tools 204d may be configured for and utilized by for various parties besides the patient, e.g., a patient community, family caregivers, and professional caregivers.

2.3.5 Behavioral Medicine Systems

Behavioral medicine systems may collect, track, and analyze behavior-related data to identify causal failure points in treatment and to predict corrective action by prescribing specific behavior modifications. In various aspects, the behavioral medicine systems may assist patients via questionnaires and patient profile assessment on symptomatology or therapeutic subjects, e.g., in various decision processes by display a menu-guided series of questions and receiving answer(s) from the patient.

FIG. 10 illustrates an exemplary IEM data framework 102 having a behavioral medicine system 204e, according to one embodiment. The IEM data framework 102 further includes IEM data 200 and the hub, shown here embodied as the base station 404 and the mobile telephone 406. In various aspects, the behavioral medicine system 204e may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106.

In one scenario, the behavioral medicine system 204e, e.g., a software agent, may be located in whole or in part on a patient-related device such as the mobile telephone 406. The software agent may assist the patient in various endeavors, e.g., diet choices, smoking cessation, etc. The assistance may be provided, by example, by generating for display on the mobile telephone 406 question sets related to diet and smoking cessation. The patient may answer the questions, e.g., select from various answer options. Based on the patient’s answers to the questions, the software agent may categorize the patient according to predetermined categories. The software agent may provide language and menu choices based on the patient categorization.

In another scenario, patient behavior is tracked with respect to various IEM data, e.g., patient parameters, sometimes referred to herein as “sentinel for wellness”. Examples of sentinels for wellness include medication therapy adherence, weight, blood pressure, etc. The sentinels for wellness may be derived, for example, from various health devices 300: such as intelligent scales, cardiac-related devices, etc.

To illustrate, patient 506 ingests medication according to physician instructions. The IEM data 200 in the form of ingestion information identifying the ingested medication and the time of ingestion are captured via an ingestion device and communicated to the patient’s mobile telephone 406. Also captured via health device(s) 300: at the time of medication ingestion are the patient’s blood pressure and weight. The timing of the foregoing data captures may be synchronized via, for example, software utilizing a reminder system to alert the patient to take the medication at a particular time. Upon receiving the ingestion information, e.g., confirmation of ingestion, the software associated with the mobile telephone 406 communicably triggers health device(s) 300: to determine blood pressure and weight, and forwards such data to the mobile telephone 406 for aggregation with the IEM data 200 in the form of the ingestion information.

The aggregated data may be forwarded to behavioral medicine system 204e, which may be configured, for example, as the mobile telephone and software 406, the server 500 including the software 502 and the database 504, and/or other configurations. Upon receipt of the aggregated data, various processing may take place.

One example of processing is analysis of the IEM data 200 to determine degree of patient adherence to medication regimen, i.e., determine if the patient ingested the prescribed medication in the right dosage at the prescribed time interval(s).

Another example of processing is analysis of the IEM data 200 to determine if the blood pressure measurement is in line with physician expectations. Thus, the notification of patient adherence to the medication regimen and the blood pressure measurement may be communicated to a physician system 106f for review by the patient’s physician. The physician, in turn, may update the IEM data 200, e.g., determine
an adjustment in the medication regimen is needed and communicate, via the behavioral medicine system, the updated medication regimen to the patient's mobile telephone 406 and to the pharmacy system 106b for filling the updated prescription.

[0269] In cases of a nonadherence determination, the physician may alert the patient, via the behavioral medicine system 204c, to make an appointment for a physical review. In various aspects, the behavioral medicine system 204c may generate and/or forward a reminder to the hub, e.g., mobile telephone 406 of the patient 506. The reminder, for example, may include the dosing schedule, a reminder for the upcoming dose, instructions to follow in case of a missed dose, etc.

[0270] In cases of underdosage/overdosage, the behavioral medicine system 204c may interoperate with an alert system, e.g., the IEM data phone system, infra, and compare current dosage information to predetermined thresholds to determine if a critical status dosing event exists, e.g., the patient is critically underdosed or critically overdosed. If such a determination is made, the appropriate system may generate an alert to appropriate parties, e.g., generate a 911 emergency call for medical assistance, generate an emergency alert to the physician system 106f, and generate an alert to a family caregiver system 106g, e.g., a family member's mobile telephone.

[0271] In still another scenario, analysis of the patient's communication patterns/habits is performed to determine patient parameters, indicated actions, etc. To illustrate, an application such as software 502 resident on the mobile telephone 406 tracks the patient's phone usage to determine communication patterns. For example, the family caregivers, physician, etc., may selectively configure tracking parameters of the application to determine various patient communication thresholds, patterns, etc. The software monitors communication from/to the selected device, e.g., the patient's mobile telephone 406. In various aspects, the application mines mobile telephone records of the associated carrier to determine calling and called parties, heavy volume call time, no call times, etc. and builds a profile against the same. The application monitors use of the mobile telephone 406 and identifies significant, e.g., user selected, deviations from the profile. Upon identification of a deviation, the application initiates predetermined actions, e.g., communicates an alert to the physician and/or family caregiver via the healthcare system 106a, the physician system 106f, and/or the family caregiver system 106g.

[0272] Another example of processing is analysis of the IEM data 200 together with data from another source, e.g., aggregated data. The aggregated data may be collected from various sources, aggregated at various and/or multiple points, and/or communicated via various channels to/from various devices.

[0273] To illustrate, cardiovascular data is derived via electrical tomography, as heretofore discussed. The cardiovascular data is communicated directly or indirectly, e.g., via the patch receiver 400, to a software application on the hub, e.g., the mobile telephone 406. The software application on the mobile telephone 406 aggregates the cardiovascular data with the IEM data, e.g., pill ingestion-related data, and displays the various data via a graphical user interface (GUI).

[0274] Subsequent to enrollment, the behavioral medicine system ascertains that the patient has neglected to take the medication at the appropriate times. Reminder alerts for upcoming medication dosing time(s) are sent to the patient via the mobile telephone. Upon receiving the alerts, the patient timely ingests the medication, resulting in a change in the sentinel for wellness.

[0275] 23.6 Incentive Systems

[0276] Incentive systems provide incentives and rebates through various programs. The incentives and rebates are based on, or otherwise associated with, the IEM data. The IEM data may be analyzed via, for example, an IEM data system 204 to determine if certain criteria/thresholds/goals are evident. Based on the determination, incentives tied to or associated with the criteria/threshold/goals may be generated.

[0277] FIG. 11 illustrates an exemplary IEM data framework 102 having an incentive system 204f, according to one embodiment. The IEM data framework 102 further includes IEM data 200 and the hub, shown here embodied as the mobile telephone 406. In various aspects, the incentive system 204f may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106.

[0278] In one scenario, patient adherence is tracked with respect to various patient parameters, e.g., medication therapy and adherence. Incentives may be awarded accordingly. For example, patient 506 ingests medication according to physician instructions. The IEM data 200 in the form of ingestion information identifying the ingested medication and the time of ingestion are captured via an ingestion device and communicated to the patient's mobile telephone 406, and to the behavioral medicine system 204c. The behavioral medicine system 204c verifies patient 506 adherence to the prescribed medication regimen, and sends verification to the incentive system 204f. The incentive system 204f, via the software 502 and the database 504, determines the price paid for the medication, and issues a rebate or credit against the cost. For example, the rebate may be issued and a financial transaction in the amount of the rebate posted to the patient's financial account via the financial transaction system 106c.

[0279] In another example, the rebate may be communicated and applied to an account associated with the patient via the pharmacy system 106b, with, for example, a credit against the next refill for the patient's prescription medication.

[0280] In another example, the patient's blood pressure and weight may be captured via health device(s) 300c at time of medication ingestion. The timing of the foregoing data captures may be synchronized via software utilizing a reminder system to alert the patient to take the medication at a particular time. Upon receiving the ingestion information, e.g., confirmation of ingestion, the software associated with the mobile telephone 406 may communicably trigger health device(s) 300c to determine blood pressure and weight, and forward such data to the mobile telephone 406 for aggregation with the IEM data 200 in the form of the ingestion information. The aggregated data may be communicated to the incentive system 204f where the software 502 and/or database 504 may be utilized to determine if the patient's weight and blood pressure meet acceptable predetermined thresholds. If, for example, the weight exceeds an acceptable threshold, the incentive system 204f may generate an incentive in the form of a discount membership offering at a local health club, etc. The offering may be constructed using various data parameters and demographics, e.g., geographical location of the patient, amount of weight to be lost, health assessment scoring based on individualized patient health parameters, lists of participating health clubs, etc.
The incentive may be communicated to the patient 506 via, for example, the patient’s mobile telephone 506.

Personalized Commercial Products/Services

Personalized commercial products/services provide individualized products and services predicated on or related to IEM data.

FIG. 12 illustrates an exemplary IEM data framework 102 having a personalized commercial products/services system 204a, according to one embodiment. The IEM data framework 102 further includes IEM data 200 and the hub 202. In various aspects, the commercial products/services system 204a may be embodied as, for example, an IEM data device, e.g., a patch receiver. In various aspects, the commercial products/services system 204a may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106.

In one scenario, commercial products/services system 204a further comprise branded or “community” associated products and services.

Auto Billing Systems

Auto billing systems receive, process, and/or facilitate payment via a financial account. Auto billing applications associated with the auto billing system and/or with financial institution systems seamlessly interoperate to generate a bill, verify account holder information, charge an account, etc. Statements are updated to reflect payment information. Similar applications may be applied for prescriptions, consumer products, information provision via personal devices, etc.

FIG. 13 illustrates an exemplary IEM data framework 102 having an auto billing system 204b, according to one embodiment. The IEM data framework 102 further includes IEM data 200 and the hub, shown here embodied as the handheld device 402. In various aspects, the auto billing system 204b may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106.

In one scenario, various parties such as patient 506, physicians, pharmaceutical companies, etc., subscribe to information feeds/patient population data of IEM data 200 to further business goals, manage health care, etc. The parties may receive the information feeds/access population data, etc. via a variety of devices. For example, patient 506 may receive an information feed via hub 202 embodied as the handheld device 402, which, via a software agent, may generate a financial transaction in the form of an invoice for the information feed displayed for the patient 506. Payment may be effected via automated methods.

In a patient selection method, for example, the patient selects various payment options via the software agent resident on the handheld device 402. A payment transaction is generated and communicated to the financial transaction system 106e. The financial transaction system 106e automatically charges an account associated with the patient 506. Confirmation of the payment together with digital, e.g., electronic, copies of the invoice are provided to the software agent resident on the handheld device 402 for the patient 506 to view, etc.

In an automated method, for example, a bill and/or financial transaction are automatically generated upon predetermined criteria. The predetermined criteria include, for example, delivery of information associated with an information feed or other source, access to a data collection, e.g., patient population data stored in a database, etc. The patient selects various payment options via the software agent resident on the handheld device 402, and a payment transaction is generated and communicated to the financial transaction system 106e. The financial transaction system 106e automatically charges an account associated with the patient 506. Confirmation of the payment together with digital copies of the invoice are provided to the software agent resident on the handheld device 402 for the patient 506 to view, etc.

For example, a healthcare provider may access patient population data stored in decision support system 204a via the healthcare system 106a. Software of the decision support system 204a may cooperate with the software 502 and the database 504 of the auto billing system 204b to identify the party to be billed for the access. Upon identification, the auto billing system 204b may automatically generate a bill and/or financial transaction for the access via one or more of the aforementioned channels.

Tracking Systems

Tracking systems track and integrate product movement data. In one example, the life cycle of an ingestible device may be tracked from manufacture to shipment, pharmacy inventory, delivery to patient, ingestion and expulsion.

FIG. 14 illustrates an exemplary IEM data framework 102 having a tracking system 204c, according to one embodiment. The IEM data framework 102 further includes the IEM data 200 and the hub, shown here embodied as a scanner 1402. In various aspects, the tracking system 204c may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106.

In one scenario, a pharmaceutical manufacturer produces an ingestible device 302a such as a particular medication having an IEM system device therein. The IEM system device contains various IEM data 200 such as medication identification, batch number, lot number, and manufacturer identification. The scanner 1402 may be utilized at various times/locations to scan the ingestible device 302a and capture the IEM data 200 associated therewith. The IEM data 200 may then be stored, processed, etc., via, for example, the software 502 and the database 504 of the tracking system 204c. For example, the IEM data 200 may be read by the scanner at a shipping point and when received by a pharmacy to ensure inventory control, distribution integrity, and chain of custody for restricted pharmaceuticals, etc.

The tracking information may be used, for example, by regulatory agencies systems 106 to determine regulatory adherence, etc.

Interdiction Systems

Interdiction systems track, reconcile, and support interdiction programs. The interdiction programs include, for example, programs related to drug identification and use detection by sworn personnel, search and seizure activities, etc.

FIG. 15 illustrates an exemplary IEM data framework 102 having an interdiction system 204d, according to
one embodiment. The IEM data framework 102 further includes IEM data 200 and hub, shown here embodied as a scanner 1402. In various aspects, the interdiction system 204 may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106.

[0301] In one scenario, a pharmaceutical manufacturer produces an ingestible device 302a such as a particular medication having an IEM system device therein. The IEM system device contains various IEM data 200 such as medication identification, batch number, lot number, and manufacturer identification. The scanner 1402 may be utilized at various times/locations to scan the ingestible device 302a and capture the IEM data 200 associated therewith. The IEM data 200 may then be communicated to, for example, the software 502 and the database 504 of the interdiction system 204j, where the IEM data 200 may be accessed by and communicated to regulatory agency systems 106j to facilitate various regulatory and enforcement functions, to locate missing controlled substances, to intercept contraband, to identify unknown substances, and to otherwise support agency and regulatory activities.

[0302] In various aspects, the IEM data 200 may be communicated to/from, for example the interdiction system 204j from/to the tracking system 204i, for processing, storage, etc. For example, the IEM data 200 may be read by the scanner at a shipping point and read by a pharmacy to ensure inventory control, distribution integrity, and chain of custody for restricted pharmaceuticals, etc. The scanned (read) IEM data 200 may be reconciled between the interdiction system 204j and the tracking system 204i to ensure complete shipment, to track shipments through various jurisdictions, etc. In one example, the IEM data 200 such as the identifier data, shipment data, patient information, recipient information, and commercial activities are tracked and reconciled to intercept contraband and otherwise support agency and regulatory activities.

[0303] 2.3.11 Subscription Systems

[0304] Subscription systems enable subscription to various IR information feeds and data/knowledge collections, e.g., IEM data collection system. For example, patients subscribe to IEM data information feeds and/or IEM data collections, which aggregate various sources of data and fuse the data into integrated, individualized information based on the subscriber’s requirements. The information fusion may include, for example, personalized medication regimens and alert applications, individual social community information, music, etc. The information may be automatically billed, for example, under a single point of charge model on a recurring basis. The agent may be provided as part of an embedded device, e.g., standard application on a mobile telephone, etc.

[0305] FIG. 16 illustrates an exemplary IEM data framework 102 having a subscription system 204k, according to one embodiment. The IEM data framework 102 further includes IEM data 200 and the hub, shown here embodied as a mobile telephone 406. In various aspects, the subscription system 204k may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106.

[0306] In one scenario, the patient 506 subscribes to various information feed(s) and/or IEM data collections, discussed hereinafter in detail. The information feed(s) include, for example, structured and non-structured information on a variety of topics generated or delivered from various sources, e.g., websites, blogs, etc. The IEM data collections include storage repositories having IEM data. The storage repositories may be associated, e.g., integral to or remote from, the subscription system 204k. For example, an IEM data collection may be resident in part or wholly in database 504 of the subscription system 204k.

[0307] In one scenario, IEM data 200 are communicated from a subscription source to a subscriber, e.g., a subscriber’s device. The subscription source includes, for example, IEM data systems 204, e.g., the database 504 of the subscription system 204k, feedback loop system 204a, patient tools 204d, and decision support system 204b; commercial systems 106b, e.g., online medical and business information/newsfeed sources, healthcare system 106a; and other sources, e.g., devices associated with the patient 506, the hub, etc. The subscriber includes, for example, a person, group, or resource, e.g., a database, a computer system, server, network, etc.

[0308] In various aspects, subscription services may be initiated via, for example, a software agent resident on the hub or communication with a local or remote system such as the healthcare system 106a.

[0309] In various aspects, the subscription services may be billed and paid via, for example, the subscription system 204k and the financial transaction system 106c.

[0310] In various aspects, the subscription newsfeeds/data may be combined or integrated into a single or multiple newsfeeds, e.g., the software 502 and/or the database 504 of the subscription system 204k may enable data aggregation, etc.

[0311] To illustrate, the patient 506 subscribes to a healthcare newsfeed and a pharmacy newsfeed, one or more having IEM data 200, via the subscription system 204k. The patient subscribes by selecting an application, e.g., software agent resident on the hub, illustratively embodied here as the mobile telephone 406. Once the patient has selected the subscription options, the order is communicated to the subscription system 204k, which, via the software 502 and the database 504, confirms, processes, stores, and bills the order. The subscriber’s financial account may be automatically charged, for example, by communicating invoice information to a financial transaction system 106c associated with the subscriber’s account. Confirmation of the charge may be communicated from the financial transaction system 106c to the subscriber via the subscription system 204k and/or the mobile telephone 406.

[0312] Based on the subscription parameters, the subscription system 204k receives the healthcare newsfeed information and the pharmacy newsfeed information. The software 502 of the subscription system compares subscriber data of the patient 506 in the database 504 against subscriber data found in the pharmacy newsfeed, e.g., patients who are prescribed medications for cardiac therapy. Based on the comparison, software 502 separates the data of the pharmacy newsfeeds relevant to the subscriber, combines the relevant data with the healthcare newsfeed information and communicates the combined newsfeed information to the mobile telephone 406 for access and display.

[0313] 2.3.12 IEM Data Collection System

[0314] The IEM data collection system provides/facilitates access to/storage of the IEM data. Examples of the IEM data include patient population data and electronic medical records. In various aspects, IEM data collections may include
functionality related to the collection, management, manipulation, storage, dissemination, and billing of IEM data. [0315] FIG. 17 illustrates an exemplary IEM data framework 102 having an IEM data collection system 204, according to one embodiment. The IEM data framework 102 further includes IEM data 200 and the hub, shown here embodied as a handheld device 402. In various aspects, the IEM data collection system 204, may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106. [0316] In one scenario, patient population data, e.g., anonymized, empirical patient data, is stored in one or more repositories, e.g., the database 504 of the IEM data collection system 204. The patient population data may be received from various sources, e.g., the IEM data 200 associated with one or more patient 506, IEM data systems 204 such as behavioral medicine systems 204a, prescription systems 204b, patient tools 204c, etc., and commercial systems such as healthcare systems 106a, pharmacy systems 106b, university systems 106c, etc. [0317] In various aspects, the IEM data collection system 204 may be consolidated in a single physical and/or logical location, e.g., the database 504 of the server 500 of the IEM data collection system 204, or distributed across two or more systems or locations, e.g., remotely distributed on multiple IEM data systems 204, associated with commercial systems 106, and/or distributed between the IEM data collection system 204 and other systems/locations. [0318] Multiprofile users may access, utilize, and/or contribute to the IEM data collection system 204. Multiprofile users include, for example, individuals or groups using various methods/devices for access, utilization, and/or contribution. Examples of multiprofile users include patient 506, family members and family caregivers, professionals, academics, corporates, etc. The methods/devices include the hub devices such as a mobile telephone, base station, handheld device, etc., as well as system components associated with IEM data systems and commercial systems, e.g., laptop computer associated with a university network, a desktop computer associated with the family caregiver system 106d, etc. [0319] To continue the foregoing illustration, a researcher, using the university system 106d, accesses the IEM data collection system 204 via the Internet, etc., and submits queries against the patient population data, extracts various data, etc. [0320] In various aspects, the IEM data collection system 204 includes privacy assurance, authentication, and validation mechanisms with respect to financial, medical, and other privacy information. For example, the software 502 may authenticate users. The software 502 may cleanse/verify data to ensure predetermined privacy thresholds are met. [0321] 2.3.13 Approval Systems [0322] Approval systems aggregate and/or analyze various data to enable an informed approval decision. [0323] FIG. 18 illustrates an exemplary IEM data framework 102 having an approval system 204n, according to one embodiment. The IEM data framework 102 further includes IEM data 200, the hub, shown here embodied as a handheld device 402, and an associated intelligent pill dispenser 1802. In various aspects, the approval system 204n may interoperate, or be otherwise associated with, one or more IEM systems 204 and/or one or more commercial systems 106. [0324] In one scenario, the patient 506 opens an intelligent pill dispenser 1802, e.g., a pill dispenser having a microchip and communication abilities. The patient 506 removes a pill having an IEM system from the intelligent pill dispenser 1802. The intelligent pill dispenser 1802, via its microchip, senses the removal of the pill, receives a signal from an IEM system that the patient 506 has ingested the pill, and determines the remaining quantity. If the remaining quantity is fewer than a predetermined threshold quantity, the intelligent pill dispenser 1802 communicates a refill request to the approval system 204n. The approval system 204n, for example, the software 502 and the database 504, verify information associated with the patient 506, e.g., patient name, medication identification, medication ingestion verification, refill timing, etc. The approval system 204n may interoperate with, e.g., communicate with, various IEM data systems 204 and/or commercial systems 106 to obtain/validate information. For example, data provided to/resident in the approval system 204n may be reconciled with medical records of healthcare system 106, the refill request approved by approval system 204n, and a refill communicated to the pharmacy system 106b. [0325] 2.3.14 Forecasting Systems [0326] Forecasting systems aggregate data and/or facilitate analysis of the aggregated data/data collections to derive/generate predictive information. [0327] FIG. 19 illustrates an exemplary IEM data framework 102 having a forecasting system 204n, according to one embodiment. The IEM data framework 102 further includes IEM data 200 and the hub, shown here embodied as a base station 404. In various aspects, the forecasting system 204n, may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106. [0328] In one scenario, for example, IEM data 200 are received by the base station 404 from ingestible devices associated with patients 506a-c. The base station 404 communicates the IEM data 200 to the IEM data collection system 204, which anonymizes the IEM data 200 and aggregates the anonymized IEM data 200 with patient population data. [0329] The IEM data collection system 204 communicates all or a portion of the patient population data to the forecasting system 204n, where the software 502, e.g., one or more applications, processes the patient population data to derive various statistics, conclusions, forecasts, etc., according to predetermined requirements, objectives, etc. For example, the software 502 processes the patient population data and correlates various data such as blood pressure readings over a predetermined period of time versus medication taken versus adherence to medication regimen to determine overall efficacy of medication regimen and to forecast titrated patient dosing based on the overall efficacy findings. [0330] Multiple profile parties, e.g., analysts using the pharmacy systems 106a, agents using the regulatory agency systems 106b, and researchers using the university systems 106c, access the forecasting system 204n. The multiple profile parties utilize various tools, e.g., the software 502, to run analytical and forecasting applications against the patient population data and to access various forecasting data available in connection with the forecasting system 204n. [0331] 2.3.15 Financial Systems [0332] Financial systems support and enable financial transactions associated with IEM data. In various aspects, the financial systems are communicably interoperable with existing automated banking systems and networks, etc.
FIG. 20 illustrates an exemplary IEM data framework 102 having a financial system 204o, according to one embodiment. The IEM data framework 102 further includes IEM data 200 and the hub, shown here embodied as a mobile telephone 406. In various aspects, the financial system 204o may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106.

In one scenario, the patient 506, via the mobile telephone 406, places an order for a product/service, e.g., a new fitted service from the subscription system 204k. The subscription system 204k, via its software, interoperates with the financial system 204o. The subscription system 204k, for example, securely communicates encrypted patient financial information such as account number and subscription information. The financial system 204o authenticates the patient information and securely interoperates with the patient’s financial institution, e.g., via a commercial system 106 such as the financial transaction system 106c to charge the patient’s account and provide charge information/confirmation to the patient 506 via, for example, the mobile telephone 406.

The IEM data phone enables IEM data related applications. For example, application(s) include pill regimen scheduling applications, alert reminder applications, auto refill for medication applications, patient tool applications, social networking applications, incentive tracker applications, auto billing applications, subscription applications, approval applications, and financial transaction applications. The applications may be integrated with, associated with, or independent of one another. The applications may further be manufacturer-installable on the IEM data phone, downloadable or otherwise installable by a wholesaler, retailer, user, etc. Installation may be independent or bundled with other software, products, etc. In various aspects, the applications are user-configurable, downloadable, upgradeable, etc.

In various aspects, the IEM data phone and/or its applications may share common features, e.g., a common graphical user interface (GUI); branding, i.e., a collection of images and ideas representing an economic producer such as concrete symbols embodied as a name, logo, slogan, design scheme, etc. The IEM data phone may also include various connectivity schemes, e.g., Internet and cellular; may provide multimedia capabilities; and may embody various hardware and software configurations. The IEM data phone may be embodied in a variety of devices, e.g., the mobile telephone 406, the handheld device 402, etc.

FIG. 21 illustrates an exemplary IEM data framework 102 having an IEM data phone 204p, according to one embodiment. The IEM data phone 204p may serve as the hub, for example, IEM data framework 102 further includes IEM data 200. In various aspects, the IEM data phone 204p may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106.

In one scenario, the IEM data phone 204p includes the software 502, e.g., a portfolio of branded applications such as pill regimen scheduling, alert reminders, auto refills, patient tools, social networking, incentive trackers, auto billing, subscriptions, approvals, and financial applications.

The pill regimen scheduling application may accept, reconcile, calendar, and manage contraindications and interactions of medication regimen(s). For example, the patient 506 may input information related to one or more prescriptions, including the pharmaceutical name and dosage. The pill regimen scheduling application may check the input information against existing information stored on the IEM data phone 204p, e.g., in the database 504, or elsewhere, e.g., the pharmacy 106b. The pill regimen scheduling application may provide information regarding contraindicated medications, side effects, precautionary instructions. The pill regimen scheduling application may calendar the dosing information and generate alerts, e.g., reminders generated at appropriate times alerting the patient to ingest the medication. The alerts may be audible, visual, email, text message, etc. and may be integrated with, or independent of, alert reminder application(s).

The alert reminder application may accept or access various data associated with scheduling, including IEM data 200, and generate alerts at appropriate times. The alerts may be audible, visual, email, text message, etc. and may be integrated with or independent of alert reminder application(s). The alert application may be user-configurable, e.g., type of alert, repetition of alert, interval of repetition, receivers of alert. The alerts may be associated with various devices of the patient, family caregivers, friends, etc. In one example, the patient 506 may schedule reminders to be sent to the user’s device, e.g., the IEM data phone 204p, the handheld device 402, the base station 404, the mobile telephone 406, etc.

The alert reminder application may be integrated with other applications/systems. To illustrate an IEM system associated with the patient 506 that may, for example, detect medication ingestion event(s) and communicate the IEM data 200 associated with the medication ingestion event(s) to the alert reminder application via the IEM data phone 204p. The alert reminder application may interoperate with the pill regimen scheduling application and perform various checks, e.g., the ingested medication was actually prescribed for the person that ingested it; the ingested medication was ingested in the correct dosage; the ingested medication was ingested at the prescribed time interval, etc.

Predetermined criteria may be used to determine if/when the alert reminders application generates an alert, reminder, etc. To continue with the foregoing illustration, upon a determination that the ingested medication was not prescribed for the person ingesting it or the wrong dosage was ingested, the alert reminder system generates alert(s) to a predetermined destination, e.g., alerts in the form of text messages to mobile telephones associated with the family caregiver system 106g, alerts in the form of email/text messages to the healthcare system 106a and the physician system 106f. If the event is deemed critical, e.g., ingestion of non-prescribed medication, overdose, etc., the alert reminder application may generate a call from the IEM data phone 204p to the emergency assistance system, e.g., place a 911 call. The call (prerecorded audio, text message, etc.) may contain information such as the patient’s name, the nature of the emergency, the ingestion details, physician and family caregiver information, and the physical location of the person ingesting the medication.

The auto refill application may facilitate automatic refill of a prescription medication via interoperation with, for example, the pharmacy system 106b, etc.

The patient tool application may be provided on or accessible from the IEM data phone 204p. For example, software tools for tracking dietary and physiologic symptoms may facilitate user entry of dietary intake and symptoms, collection of device-associated physiologic parameters such
as blood pressure, heart rate, etc., correlation/analysis of the data, and feedback based on the correlation/analysis. The patient tool application may provide data, e.g., the feedback, for display on the IEM data phone 204p, the IEM data system (s) 204, and/or the commercial system(s) 106.

[0346] The social networking application may facilitate social networking functionality. For example, the social networking application may retain various links to selected profiles of various social networks, receive data related to the selected profiles, e.g., updates to the profiles, facilitate messaging and other communication, update the user’s profile, etc., communicate with the IEM data systems(s) 204, and/or the commercial system(s) 106, such as the patient tools/social network 204d and the web communities 106h.

[0347] The incentive tracker application may collect, manage, track, update, etc. incentive information. For example, the incentive tracker application may reconcile data associated with the IEM data collection systems 204 and wholesaler/retailer systems 106 to determine incentive eligibility, e.g., a patient rebate. The incentive tracker application may further tally points under various reward systems, notify the patient 506 of milestones, goals, award of incentive, etc.

[0348] The auto billing application may facilitate billing for various transactions. The auto billing application may interoperate with various applications/systems, including the IEM data system(s) 204 and/or the commercial system(s) 106, such as the billing for an auto refill via the pharmacy system, etc.

[0349] The subscription application facilitates ordering, receipt, management, etc. of various subscriptions, e.g., newsfeeds, access to various data collections on a subscription basis, etc. The subscriptions application may interoperate with various applications/systems, including the IEM data system(s) 204 and/or the commercial system(s) 106, such as the subscription system 204k, the IEM data collection system 204f, etc.

[0350] The approval application aggregates and/or analyzes various sources of data to enable an informed approval decision. The approvals application may interoperate with various applications/systems, including the IEM data system(s) 204 and/or the commercial system(s) 106, such as the auto refill system 204c, the subscription system 204f, the financial systems 204o, the pharmacy systems 106b, the wholesaler/retailer systems 106e, etc.

[0351] The financial application supports and enables financial transactions associated with IEM data 200. The financial application may interoperate with various applications/systems, including the IEM data system(s) 204 and/or the commercial system(s) 106, such as the auto refill system 204c, the incentive system 204f, the subscription system 204f, the approval system 204m, the financial systems 204o, the pharmacy system 106b, the wholesaler/retailer systems 106e.

[0352] 3.0 IEM Data Framework Method

[0353] Social networks are a social structure made of one or more nodes, e.g., components such as websites, accessed by individuals or organizations. The social network is typically tied by one or more specific types of interdependency, such as epidemiology, therapeutic regimen, healthcare management, etc., and thus may attract the interest of otherwise unrelated individuals and groups having in common an interest in the interdependencies. Social networks may be built around various communities, e.g., family caregivers, patients, medical conditions, etc.

[0354] One example of a social network is a patient information community that provides information related to a particular medical condition, treatments, medications, regimens, and side effects based on both provider and anecdotal data. The availability of such data may provide benchmark-type services, e.g., facilitate self-assessment of personal progress and adjustment in therapies and behaviors by comparing and contrasting an individual’s progress with the particulars of others having the same condition, similar therapies, etc.

[0355] FIG. 22 illustrates an exemplary IEM data framework 102 having a social network system 204q, according to one embodiment. The IEM data framework 102 further includes IEM data 200, and the hub, shown here embodied as the base station 404. In various aspects, the social network system 204q may interoperate, or be otherwise associated with, one or more IEM data systems 204 and/or one or more commercial systems 106.

[0356] In one scenario, patient 506 suffers from a cardiac condition. The patient 506 accesses the social network system 204q, which may be embodied as the server 500 having the software 502 and the database 504 having IEM data 200. Patient 506 may access the social network system 204q, for example, via the base station 404. The patient 506 searches the database 504 for patient profiles also having cardiac conditions similar to that of patient 506. The social network system 204q provides multiple profiles of patients having similar conditions. The profiles include various data pertinent to each patient such as medication therapies, personal behavior histories, etc. The patient 506 requests a comparison of his medication therapy, medication therapy adherence, and behavior to that listed in the profiled. The social network system 204q provides the requested comparative data in the form of a graphical display. From the display, the patient 506 is able to determine the profiles having the most favorable treatment outcomes. From such profiles, the patient 506 and/or the social network system 204q analyze the differences between his medication, medication therapy adherence, behavior, etc. and the corresponding interdependencies of the profiles having the most favorable treatment outcomes. The analysis may contrast the differences found in various areas, as well as generate prescriptive advice, e.g., in which areas the patient 506 may want to adjust and specific adjustments based on the analysis. The patient 506 may adopt the prescriptive advice, i.e., adjust accordingly, to improve his own personal outcome. Further, the patient 506 may update the social network system 204q with the adjustment data, which may be used in the future for tracking personal improvement as well as benchmarking purposes by other individuals. In various aspects, the social network system 204q may be communicably associated with other web communities 106h, e.g., youth communities, business communities, etc.

[0357] 3.1 IEM Data Framework Article

[0358] One aspect comprises, for example, receiving, via a hub, ingestible event data that originates from multiple ingested event markers; and communicating, via the hub, at least a portion of the ingestible event marker data to at least one ingestible event marker data system.

[0359] 4.0 IEM Data Framework Article

[0360] One aspect comprises, for example, a storage medium having instructions, that when executed by a computing platform, result in execution of a method of utilizing ingestible event marker data, comprising: receiving, via a hub, the ingestible event data that originates from multiple
ingested event markers; and communicating, via the hub, at least a portion of the ingestible event marker data to at least one ingestible event marker data system.

[0361] 5.0 IEM Data Framework System

[0362] One aspect comprises, for example, a receive module to receive, via a hub, ingestible event data that originates from multiple ingested event markers; and a communicate module to communicate, via the hub, at least a portion of the ingestible event marker data to at least one ingestible event marker data system.

[0363] Further, any of the embodiments disclosed herein may be performed in a data processing system. To illustrate, a diagrammatic system comprises, for example, a processor, a main memory, a static memory, a bus, a video display, an alpha-numeric input device, a cursor control device, a drive unit, a signal generation device, a network interface device, a machine readable medium, instructions and a network, according to one embodiment.

[0364] The diagrammatic system may indicate a particular computer and/or a data processing system in which one or more operations disclosed herein may be performed. The processor may be a microprocessor, a state machine, an application-specific integrated circuit, a field programmable gate array, etc. The main memory may be a dynamic random access memory and/or a primary memory of a computer system. The static memory may be a hard drive, a flash drive, and/or other memory information associated with the data processing system.

[0365] The bus may be an interconnection between various circuits and/or structures of the data processing system. The video display may provide graphical representation of information on the data processing system. The alpha-numeric input device may be a keypad, a keyboard and/or any other input device of text, e.g., a special device to aid the physically challenged. The cursor control device may be a pointing device such as a mouse. The drive unit may be a hard drive, a storage system, and/or other longer term storage subsystem.

[0366] The signal generation device may be a biostimulable or functional operating system of the data processing system. The network interface device may be a device that may perform interface functions such as code conversion, protocol conversion and/or buffering required for communication to and from the network. The machine readable medium may provide instructions on which any of the methods disclosed herein may be performed. The instructions may provide source code and/or data code to the processor to enable any one or more operations disclosed herein.

[0367] Although the present embodiments have been described with reference to specific example embodiments, it will be evident that various modifications and changes may be made to these embodiments without departing from the broader spirit and scope of the various embodiments. For example, the various devices, modules, etc. described herein may be enabled and operated using hardware circuitry, e.g., CMOS based logic circuitry, firmware, software and/or any combination of hardware, firmware, and/or software, e.g., embodied in a machine readable medium.

[0368] In addition, it will be appreciated that the various operations, processes, and methods disclosed herein may be embodied in a machine-readable medium and/or a machine accessible medium compatible with a data processing system, e.g., a computer system, and may be performed in any order. Accordingly, the specification and drawings are to be regarded in an illustrative rather than a restrictive sense.

[0369] Any or all data associated with the aforementioned devices and methods, for example, may be used alone or in combination with other data to constitute IEM data, i.e., data having an IEM data aspect.

[0370] In certain embodiments, the system and/or method further includes/utilizes an element for storing data, e.g., a data storage element, where this element is present on an external device, such as a bedside monitor, PDA, smart phone, computer server, etc. Typically, the data storage element is a computer readable medium. The term “computer readable medium” as used herein refers to any storage or transmission medium that participates in providing instructions and/or data to a computer for execution and/or processing. Examples of storage media include floppy disks, magnetic tape, CD-ROM, a hard disk drive, a ROM or integrated circuit, a magneto-optical disk, or a computer readable card such as a PCMCIA card and the like, whether or not such devices are internal or external to the computer. A file containing information may be “stored” on a computer readable medium, where “storing” means recording information such that it is accessible and retrievable at a later date by a computer and/or computer-related component. With respect to computer readable media, “permanent memory” refers to memory that is permanent. Permanent memory is not erased by termination of the electrical supply to a computer processor. Computer hard-drive ROM, i.e., not used as virtual memory, CD-ROM, floppy disk and DVD are all examples of permanent memory. Random Access Memory (RAM) is an example of non-permanent memory. A file in permanent memory may be editable and re-writable.

[0371] Also provided are computer executable instructions, i.e., programming, for performing the above methods, e.g., for programming the IEM, receiver, and other components of the system. The computer executable instructions are present on a computer readable medium. Accordingly, various aspects provide a computer readable medium containing programming for use in providing ingestible event marker data.

[0372] As such, in certain embodiments the systems include one or more of: a data storage element, a data processing element, a data display element, a data transmission element, a notification mechanism, and a user interface. These elements may be present or otherwise associated with at least one of the ingestible event marker data, the hub, and the IEM data systems.

[0373] One of the above-described systems is reviewed in terms of a receive module and a communicate module. The aspects, however, are not so limited. In a broader sense, the systems are comprised of two or more different modules that communicate with each other, e.g., using the hub functionalities as reviewed above, e.g., using the IEM data in the communication, e.g., using the IEM data systems’ functionalities.

[0374] It is to be understood that this invention is not limited to particular embodiments described, and as such may vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodi-
ments only, and is not intended to be limiting, since the scope of the present invention will be limited only by the appended claims.

[0375] Where a range of values is provided, it is understood that each intervening value, to the tenth of the unit of the lower limit unless the context clearly dictates otherwise, between the upper and lower limit of that range and any other stated or intervening value in that stated range, is encompassed within the invention. The upper and lower limits of these smaller ranges may independently be included in the smaller ranges and are also encompassed within the invention, subject to any specifically excluded limit in the stated range. Where the stated range includes one or both of the limits, ranges excluding either or both of those included limits are also included in the invention.

[0376] Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. Although any methods and materials similar or equivalent to those described herein can also be used in the practice or testing of the present invention, representative illustrative methods and materials are now described.

[0377] All publications and patents cited in this specification are herein incorporated by reference as if each individual publication or patent were specifically and individually indicated to be incorporated by reference and are incorporated herein by reference to disclose and describe the methods and/or materials in connection with which the publications are cited. The citation of any publication is for its disclosure prior to the filing date and should not be construed as an admission that the present invention is not entitled to antedate such publication by virtue of prior invention. Further, the dates of publication provided may be different from the actual publication dates which may need to be independently confirmed.

[0378] It is noted that, as used herein and in the appended claims, the singular forms "a," "an," and "the" include plural referents unless the context clearly dictates otherwise. It is further noted that the claims may be drafted to exclude any optional element. As such, this statement is intended to serve as antecedent basis for use of such exclusive terminology as "solely," "only" and the like in connection with the recitation of claim elements, or use of a "negative" limitation.

[0379] As will be apparent to those of skill in the art upon reading this disclosure, each of the individual embodiments described and illustrated herein has discrete components and features which may be readily separated from or combined with the features of any of the other several embodiments without departing from the scope or spirit of the present invention. Any recited method can be carried out in the order of events recited or in any other order which is logically possible.

[0380] Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it is readily apparent to those of ordinary skill in the art in light of the teachings of this invention that certain changes and modifications may be made thereto without departing from the spirit or scope of the appended claims.

[0381] Accordingly, the preceding merely illustrates the principles of the invention. It will be appreciated that those skilled in the art will be able to devise various arrangements which, although not explicitly described or shown herein, embody the principles of the invention and are included within its spirit and scope. Furthermore, all examples and conditional language recited herein are principally intended to aid the reader in understanding the principles of the invention and the concepts contributed by the inventors to furthering the art, and are to be construed as being without limitation to such specifically recited examples and conditions. Moreover, all statements herein reciting principles, aspects, and embodiments of the invention as well as specific examples thereof, are intended to encompass both structural and functional equivalents thereof. Additionally, it is intended that such equivalents include both currently known equivalents and equivalents developed in the future, i.e., any elements developed that perform the same function, regardless of structure. The scope of the present invention, therefore, is not intended to be limited to the exemplary embodiments shown and described herein. Rather, the scope and spirit of present invention is embodied by the appended claims.

What is claimed is:

1. A system comprising:
   ingestible event marker data;
   a hub to receive the ingestible event marker data; and
   at least one ingestible event marker data system to receive,
   directly or indirectly, the ingestible event marker data from the hub.

2. The system of claim 1, wherein the ingestible event marker data comprises information about an ingestion event.

3. The system of claim 1, wherein the ingestible event marker data comprises information about a response to an ingestion event.

4. The system of claim 1, wherein the ingestible event marker data is generated by an ingestible event marker device.

5. The system of claim 1, wherein the ingestible event marker device comprises an ingestible event marker and a receiver.

6. The system of claim 1, wherein the receiver comprises a personal signal receiver.

7. The system of claim 1, wherein the ingestible event marker data is associated with a unique identifier of an individual.

8. The system of claim 1, wherein the hub comprises at least one element selected from a group consisting essentially of a base station, a personal communication device, and a mobile telephone.

9. The system of claim 8, wherein the base station comprises at least one element selected from a group consisting essentially of a desktop computer, a laptop computer, and an intelligent device/appliance.

10. The system of claim 8, wherein the personal communication device comprises a mobile computer.

11. The system of claim 1, wherein the at least one ingestible event marker data system comprises at least one element selected from a group consisting essentially of: a feedback loop system, a decision support system, an auto refill system, patient tools, a behavioral medicine system, an incentive system, personalized commercial products/services, an auto billing system, a tracking system, an interfederation system, a subscription system, an ingestible event marker data collection, an approval system, a forecasting system, a financial system, an ingestible event marker data phone system, and a social network system.

12. The system of claim 1, wherein the at least one ingestible event marker data system comprises at least one element selected from a group consisting essentially of a computer, a
receiver, a transmitter, an application, a data storage medium, a processor, a memory component, a personal communication device, software, and a mobile telephone.

13. The system of claim 1, further comprising:
an ingestible event marker data source to at least one of
generate at least a portion of the ingestible event marker
data and transmit at least a portion of the ingestible event
marker data.

14. The system of claim 13, wherein the ingestible event
marker data source is selected from a group consisting essen-
tially of an ingestible event marker data source device, a
product, an event, a patient specific parameter, an ingestible
event marker data algorithm, and a storage repository.

15. The system of claim 1, further comprising at least one
commercial system associated with at least one of the ingest-
ible event marker data, the hub, and at least one of the ingest-
ible event marker data systems.

16. The system of claim 15, wherein the commercial sys-
tem comprises at least one element selected from a group
consisting essentially of a healthcare system, a pharmacy
system, a pharmaeutical system, a university system, a finan-
cial transaction system, a physician system, a family care-
giver system, a web community, a regulatory agency system,
and a wholesaler/retailer system.

17. A method comprising:
receiving, via a hub, ingestible event marker data; and
communicating, via the hub, at least a portion of the ingest-
ible event marker data to at least one ingestible event
marker data system.

18. The method of claim 17, wherein the ingestible event
marker data comprises information about an ingestion event.

19. The method of claim 17, wherein the ingestible event
marker data comprises information about a response to an
ingestion event.

20. The method of claim 17 in a form of a machine-readable
medium embodying a set of instructions that, when executed
by a machine, causes the machine to perform the
method of claim 17.

21. An article comprising:
a storage medium having instructions, that when executed
by a computing platform, result in execution of a method
communicating ingestible event marker data via an
ingestible event marker framework, comprising:
receiving, via a hub, the ingestible event marker data;
and
communicating, via the hub, at least a portion of the
ingestible event marker data to at least one ingestible
event marker data system.

22. The article of claim 21, wherein the ingestible event
marker data comprises information about an ingestion event.

23. The article of claim 21, wherein the ingestible event
marker data comprises information about a response to an
ingestion event.

24. A system comprising:
a receive module to receive, via a hub, ingestible event
marker data; and
a communicate module to communicate, via the hub, at
least a portion of the ingestible event marker data to at
least one ingestible event marker data system.

25. The system of claim 24, wherein the ingestible event
marker data comprises information about an ingestion event.

26. The system of claim 25, wherein the ingestible event
marker data comprises information about a response to an
ingestion event.