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# (54) CHARGER WITH DATA TRANSFER CAPABILITIES

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(51) Int. Cl.

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CPC .... A61B 5/0031; A61B 5/04884; A61B 5/42; A61F 5/0026; A61N 1/36007; A61N 1/362;

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### (57) ABSTRACT

Gastric apparatus (18) is provided, including one or more sensors (70, 72, 74), adapted to generate respective sensor signals responsively to a gastrointestinal (GI) tract physiological parameter of a GI tract of a subject, and an implantable control unit (90), comprising a rechargeable battery and a first set of one or more transducers. The implantable control unit (90) is adapted to receive the sensor signals, using one or more of the transducers of the first set, and transmit data responsively to the sensor signals, using one or more of the transducers of the first set. An external control unit (200), including a power source and a second set of one or more transducers, is adapted to drive the power source to inductively transfer energy via one or more transducers of the second set, so as to recharge the battery, and receive the transmitted data, using one or more transducers of the second set.

## 62 Claims, 12 Drawing Sheets



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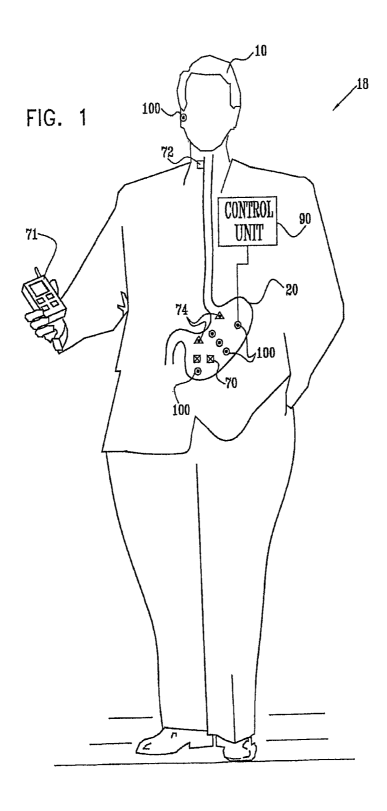
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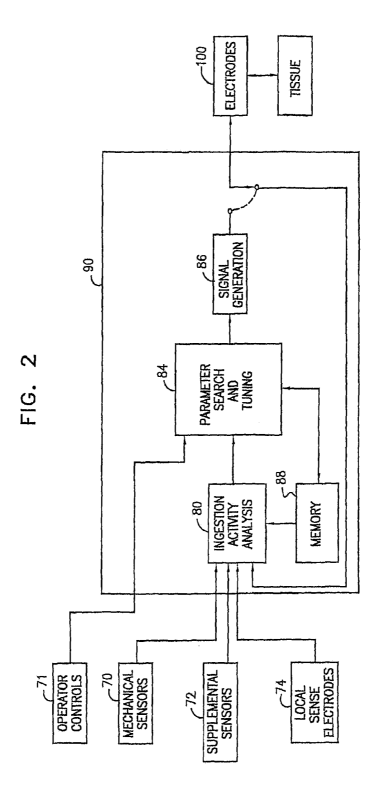
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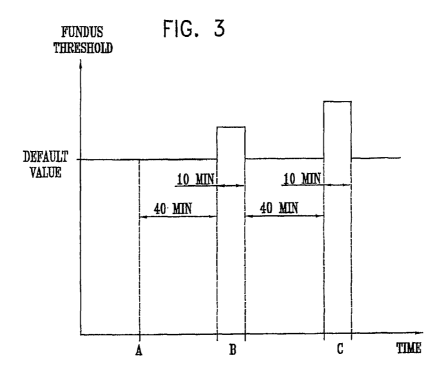
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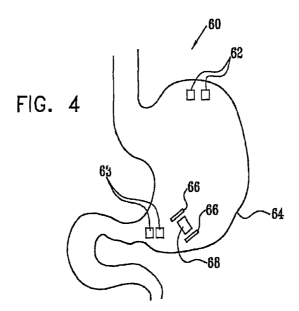
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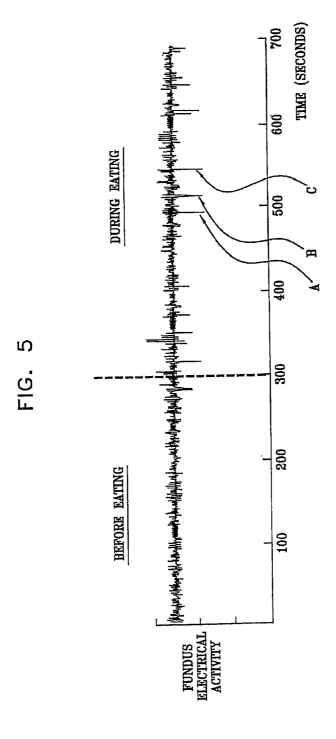


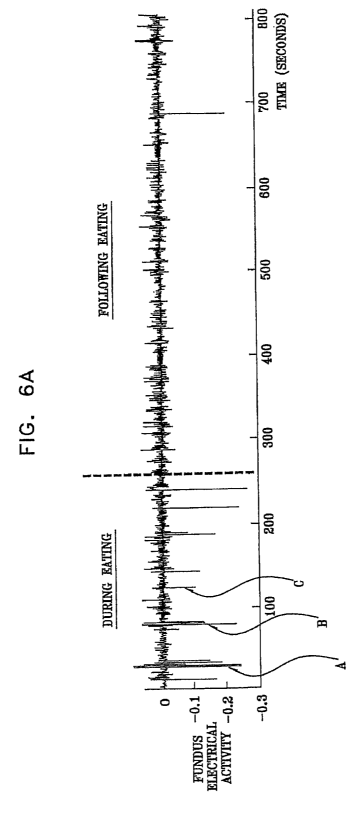


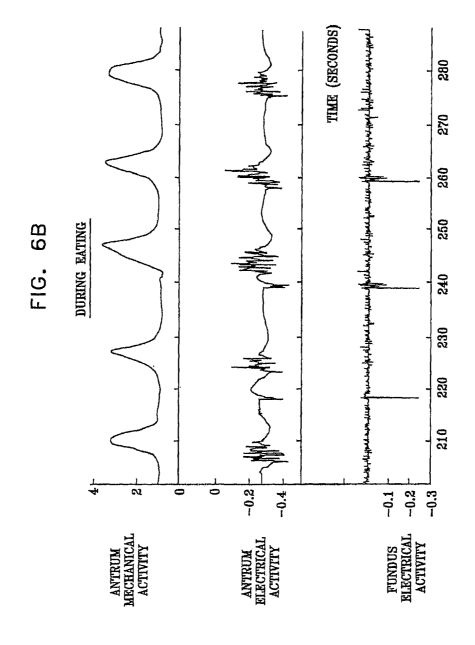
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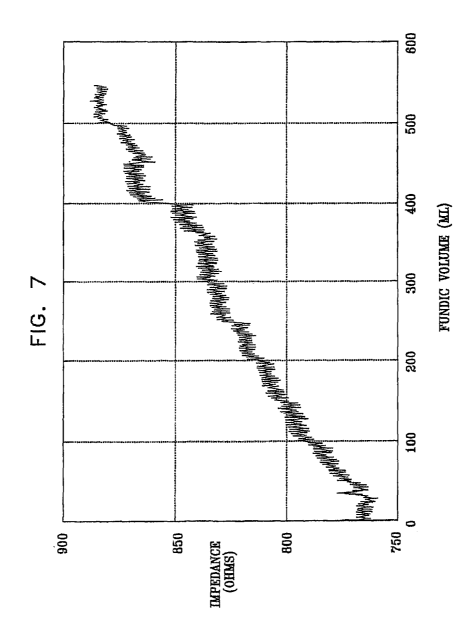


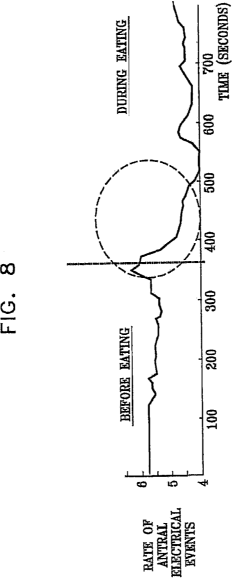




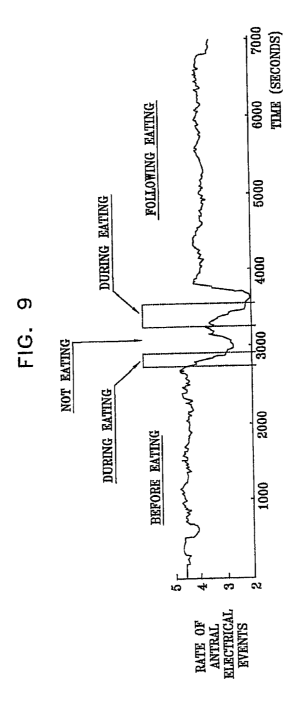


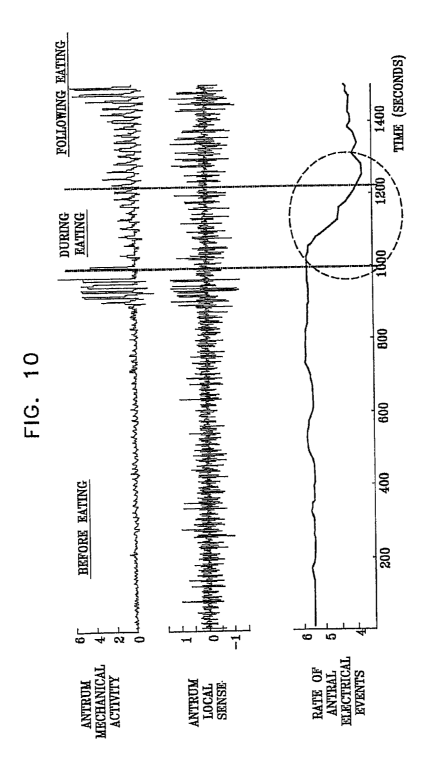






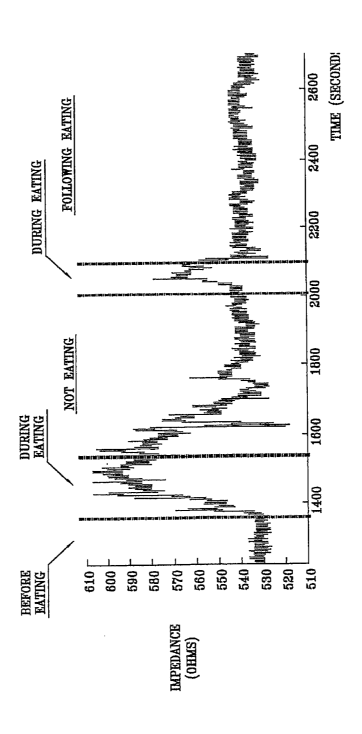
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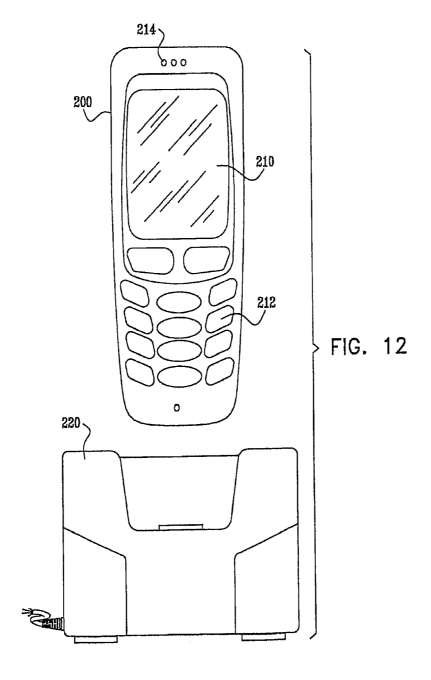




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#### CHARGER WITH DATA TRANSFER CAPABILITIES

# CROSS-REFERENCES TO RELATED APPLICATION

This application claims the benefit of U.S. Provisional Patent Application 60/654,056, filed Feb. 17, 2005, entitled, "Charger with data transfer capabilities," which is assigned to the assignee of the present application and is incorporated <sup>10</sup> herein by reference.

#### FIELD OF THE INVENTION

The present invention relates generally to tracking eating 15 habits, and specifically to invasive techniques and apparatus for detecting and analyzing the swallowing and digesting of food.

#### BACKGROUND OF THE INVENTION

Obesity is a difficult to treat chronic condition defined by a body mass index (BMI=mass/height² [kg/m²]) greater than 30. For obese persons, excessive weight is commonly associated with increased risk of cardiovascular disease, diabetes, 25 degenerative arthritis, endocrine and pulmonary abnormalities, gallbladder disease and hypertension. Additionally, such persons are highly likely to experience psychological difficulties because of lifestyle restrictions such as reduced mobility and physical capacity, due to back pain, joint problems, 30 and shortness of breath. In severe cases, this can contribute to absenteeism and unemployment. Moreover, impairment of body image can lead to significant psychological disturbances. Repeated failures of dieting and exercise to resolve the problem of obesity can result in feelings of despair and the 35 development of clinical depression.

Bariatric surgery is often recommended for persons suffering from morbid obesity. Preferably, the invasive treatment is accompanied by changes in lifestyle, such as improved regulation of eating habits and an appropriate exercise regimen. 40 Such lifestyle changes are dependent upon the self-discipline and cooperation of the patient.

A book entitled, Textbook of Gastroenterology, 3rd edition, edited by Yamada (Lippincott, Williams & Wilkins), which is incorporated herein by reference, has in Chapter 10 45 thereof a description of the physiology of gastric motility and gastric emptying.

An abstract entitled, "Gastric myoelectrical pacing as therapy for morbid obesity: Preliminary results," by Cigaina et al., retrieved on Dec. 24, 2000 from the Web-site http:// 50 www.med-online.com/transneuronix/Product/abstract.htm, which is incorporated herein by reference, describes a method for applying monopolar and bipolar gastric stimulation to achieve weight loss.

An abstract entitled, "Implantable gastric stimulator (IGS) 55 as therapy for morbid obesity: Equipment, surgical technique and stimulation parameters," by Cigaina et al., retrieved on Dec. 24, 2000 from the Web-site http://www.med-online.com/transneuronix/Product/abstract.htm, which is incorporated herein by reference, describes techniques of electrical signal therapy designed to treat obesity.

Stein et al. wrote an article related to providing incentives relating to medical care, entitled, "Carrots and sticks: Impact of an incentive/disincentive employee flexible credit benefit plan on health status and medical costs," American Journal of 65 Health Promotion, May/June 1999, V5, I13, 5, which is incorporated herein by reference.

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Giuffrida wrote an article regarding providing incentives for enhanced patient compliance, entitled, "Should we pay the patient? Review of financial incentives to enhance patient compliance," Biomedical Journal, vol. 315, pp. 703-707, 1997, which is incorporated herein by reference.

U.S. Pat. No. 6,270,455 to Brown, which is incorporated herein by reference, describes a networked system for communicating information to a patient and for remotely monitoring the patient. The system includes a server and a remote interface for entering in the server a set of queries to be answered by the patient. The server may be a web server and the remote interface may be a personal computer or remote terminal connected to the server via the Internet. The system also includes measurement apparatus for providing measurement data related to a patient's condition and treatment, and remotely programmable apparatus connected to the server via a communication network, such as the Internet. The remotely programmable apparatus interacts with the patient in accordance with a script program received from the server. The 20 server includes a script generator for generating the script program from the set of queries entered through the remote interface. The script program is received and executed by the remotely programmable apparatus to communicate the queries to the patient, to receive responses to the queries, and to transmit the responses from the apparatus to the server. The measurement data provided by the measurement apparatus may include physiological condition data and drug delivery measurement data for paperless recordation at a remote loca-

With respect to one embodiment, the Brown patent describes each patient to be monitored being provided with a monitoring device, designed to provide measurements of a physiological condition of the patient, to record the physiological condition measurements, and to transmit the measurements to the patient's remotely programmable apparatus, e.g., through a standard connection cable 30. Examples of suitable types of monitoring devices include blood glucose meters, respiratory flow meters, blood pressure cuffs, electronic weight scales, and pulse rate monitors. The specific type of monitoring device provided to each patient is dependent upon the patient's disease. For example, diabetes patients are provided with a blood glucose meter for measuring blood glucose concentrations, asthma patients are provided with respiratory flow meters for measuring peak flow rates, and obesity patients are provided with weight scales.

U.S. patent application Ser. No. 10/934,155 to Whitehurst et al., filed Sep. 4, 2004, which is incorporated herein by reference, describes an implantable stimulator with at least one electrode and/or leads attached to the stimulator. The stimulator may be implanted adjacent trigeminal, cervical, or other nerves or branches thereof via a surgical procedure in order to electrically stimulate such nerves for the treatment of headache, facial pain, epilepsy, and/or other conditions. In an embodiment, a first external electronic appliance is provided with an electronic interface means for interacting with other computing means, such as by a serial interface cable or infrared link to a personal computer or to a telephone modem or the like. Such interface means may permit a clinician to monitor the status of the implantable stimulator and prescribe new stimulation parameters from a remote location. The first external appliance may also be adapted to (a) transmit electrical power to the implantable stimulator via a second external appliance positioned near the patient, in order to power the implantable stimulator and/or recharge the a power source/ storage device thereof; and/or (b) transmit data via the second external appliance to the implantable stimulator, in order to change operational parameters (e.g., electrical stimulation

parameters) used by the stimulator. The implantable stimulator may also be adapted to (a) transmit sensed data indicating a need for treatment or in response to stimulation from the implantable stimulator (e.g., ENG, EEC, change in neurotransmitter or medication level, or other activity) to the first external appliance via the second external appliance; and/or (b) transmit data indicating the state of the implantable stimulator (e.g., battery level, stimulation settings, etc.) to the first external appliance via the second external appliance.

U.S. Pat. No. 6,129,685 to Howard, which is incorporated 10 herein by reference, describes apparatus and methods for regulating appetite by electrical stimulation of the hypothalamus and by microinfusion of an appropriate quantity of a suitable drug to a distinct site or region within the hypothalamus

U.S. Pat. No. 4,823,808 to Clegg et al., which is incorporated herein by reference, describes a method for treating obesity, including receiving a physiological measurement and generating audio or visual feedback for the patient to hear or see. The feedback is used for purposes of teaching behavior 20 modification.

U.S. Pat. No. 5,868,141 to Ellias, which is incorporated herein by reference, describes an endoscopic stomach insert for reducing a patient's desire to eat.

U.S. Pat. No. 6,067,991 to Forsell, U.S. Pat. No. 5,601,604 25 to Vincent, U.S. Pat. No. 5,234,454 to Bangs, U.S. Pat. No. 4,133,315 to Berman et al., U.S. Pat. No. 4,416,267 to Garren et al., and U.S. Pat. Nos. 4,592,339, 5,449,368, 5,226,429 and 5,074,868 to Kuzmak, which are incorporated herein by reference, describe mechanical instruments for implantation in 30 or around the stomach of an obese patient.

U.S. Pat. No. 5,690,691 to Chen et al., which is incorporated herein by reference, describes a gastric pacemaker for treating obesity and other disorders. The pacemaker includes multiple electrodes which are placed at various positions on 35 the gastrointestinal (GI) tract, and deliver phased electrical stimulation to pace peristaltic movement of material through the GI tract.

U.S. Pat. No. 5,423,872 to Cigaina, which is incorporated herein by reference, describes apparatus for applying electrical pulses to the distal gastric antrum of a patient, so as to reduce the motility of the stomach and to thereby treat obesity or another disorder.

U.S. Pat. Nos. 5,188,104 and 5,263,480 to Wernicke et al., which are incorporated herein by reference, describe a 45 method for stimulating the vagus nerve of a patient so as to alleviate an eating disorder.

U.S. Pat. Nos. 6,104,955, 6,091,992, and 5,836,994 to Bourgeois, U.S. Pat. No. 6,026,326 to Bardy, and U.S. Pat. No. 3,411,507 to Wingrove, which are incorporated herein by 50 reference, describe the application of electrical signals to the GI tract to treat various physiological disorders.

U.S. Pat. No. 5,979,449 to Steer, which is incorporated herein by reference, describes an oral appliance for appetite suppression.

U.S. Pat. No. 4,975,682 to Kerr et al., which is incorporated herein by reference, describes apparatus for food intake regulation which is external to the body and which is based upon the voluntary cooperation of the subject in order to be effective.

U.S. Pat. No. 5,861,014 to Familoni, U.S. Pat. No. 5,716, 385 to Mittal et al., and U.S. Pat. No. 5,995,872 to Bourgeois, are incorporated herein by reference, and describe methods and apparatus for stimulation of tissue, particularly gastrointestinal tract tissue.

PCT Patent Publication WO 98/10830 to Ben-Haim et al., entitled, "Fencing of cardiac muscles," and U.S. patent appli-

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cation Ser. No. 09/254,903 in the national phase thereof, both of which are assigned to the assignee of the present patent application and are incorporated herein by reference, describe various methods for controlling the behavior of muscle tissue, for example by blocking or altering the transmission of signals therethrough.

PCT Patent Publication WO 99/03533 to Ben-Haim et al., entitled, "Smooth muscle controller," and U.S. patent application Ser. No. 09/481,253 in the national phase thereof, both of which are assigned to the assignee of the present patent application and are incorporated herein by reference, describe apparatus and methods for applying signals to smooth muscle so as to modify the behavior thereof. In particular, apparatus for controlling the stomach is described in which a controller applies an electrical field to electrodes on the stomach wall so as to modify the reaction of muscle tissue therein to an activation signal, while not generating a propagating action potential in the tissue. In the context of the present patent application and in the claims, the use of such a non-excitatory signal to modify the response of one or more cells to electrical activation thereof, without inducing action potentials in the cells, is referred to as Excitable-Tissue Control (ETC). Use of an ETC signal is described in the PCT Patent Publication with respect to treating obesity, by applying the ETC signal to the stomach so as to delay or prevent emptying of the stomach. In addition, a method is described for increasing the motility of the gastrointestinal tract, by applying an ETC signal to a portion of the tract in order to increase the contraction force generated in the portion.

PCT Patent Publication WO 02/082968 to Policker et al., entitled, "Analysis of eating habits," which is assigned to the assignee of the present patent application and is incorporated herein by reference, describes apparatus and methods for detecting the occurrence of an eating event by a subject and analyzing the quantity and characteristics of the food ingested.

US Patent Application Publication 2004/0098068 to Carbunaru et al., which is incorporated herein by reference, describes techniques for both recharging and communicating with a stimulator having a rechargeable battery, which stimulator is implanted deeply in the body, in particular where the stimulator is a micro stimulator. The system includes a base station and an external device, e.g., a chair pad. The chair pad may contain an antenna/charging coil and a booster coil. The antenna/charging coil can be used for charging the rechargeable battery and also for communicating with the stimulator using frequency shift keying and on-off keying. The booster coil can be used to recharge a battery depleted to zero volts. The base station connected to the chair pad may be used to power the antenna/charging coil and the booster coil.

U.S. Pat. No. 6,516,227 to Meadows et al., which is incorporated herein by reference, describes a spinal cord stimulation (SCS) system including a replenishable power source, e.g., a rechargeable battery, that requires only an occasional recharge. The SCS system monitors the state of charge of the internal power source and controls the charging process by monitoring the amount of energy used by the SCS system, and hence the state of charge of the power source. A suitable bidirectional telemetry link allows the SCS system to inform the patient or clinician regarding the status of the system, including the state of charge, and makes requests to initiate an external charge process.

US Patent Application Publication 2003/0114899 to Woods et al., which is incorporated herein by reference, describes techniques for detecting the status of a rechargeable battery included within an implantable medical device. The medical device can incorporate a status indicator which sig-

nals the user concerning the battery status, e.g., low battery level. The signal may be audible or it may arise from an electrical stimulation that is perceptually distinguished from the operative, therapeutic stimulation. The external programmer may also incorporate a second battery status indicator that is visual, audible, or physically felt. Battery status data may be conveyed on visual displays on the external programmer by uploading this information from the medical device using a bi-directional telemetry link.

U.S. Pat. No. 6,185,452 to Schulman et al., which is incorporated herein by reference, describes a device configured for implanting beneath a patient's skin for the purpose of tissue stimulation (e.g., nerve or muscle stimulation) and/or parameter monitoring and/or data communication. Alternatively, the device is configurable to monitor a biological parameter or to operate as a transponder to retransmit received command messages.

US Patent Application Publication 2004/0106963 to Tsukamoto et al., which is incorporated herein by reference, describes an implantable integrated power module incorporating a power source (e.g., a battery), a power management circuit, a magnetically inductive coupling system for remote communication and/or inductive charging, and a homing device for locating the implanted inductive charging coil. Communication (one- or two-way) may be carried out using the inductive charging link, a separate inductive pathway, or another pathway such as RF or light waves.

#### SUMMARY OF THE INVENTION

An embodiment of the invention provides apparatus and methods for detecting and tracking the swallowing of solids and liquids.

An embodiment of the invention provides apparatus and methods for detecting, tracking, quantifying and determining 35 the qualitative character of ingested liquids and solids.

An embodiment of the invention provides apparatus and methods for treating obesity.

An embodiment of the invention provides apparatus and methods that enable the implementation of changes in food 40 ingestion habits in a predictable and controlled manner.

An embodiment of the invention provides methods and apparatus for regulating food ingestion.

An embodiment of the invention provides apparatus and methods for bariatric surgery that are less drastic than those 45 currently employed.

In some embodiments of the present invention, apparatus for detecting, tracking, quantifying and determining the qualitative character of ingested liquids and solids comprises a sensor coupled to a patient's gastrointestinal tract. Typi- 50 cally, the sensor generates a signal indicative of the swallowing of food. An analysis module typically determines a quality of the food, for example, whether it is predominantly solid or liquid, and stores this information in an electronic memory. Alternatively or additionally, the analysis module determines 55 other characteristics of the ingested material, for example, the nutritional, chemical, and/or caloric content. "Food," as used in the context of the present patent application and in the claims, is to be understood as including both solid and liquid food. "Swallowing," as used in the context of the present 60 patent application and in the claims, is to be understood as being indicative of the onset of eating.

In some embodiments of the present invention, swallowing is detected by tracking the electrical activity in muscle tissue in the fundic region of the stomach. Typically, the commence-65 ment of enhanced electrical activity is also detected in muscle tissue in the antral region of the stomach. Measurement of the

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time delay between swallowing and the commencement of electrical activity in the antrum is typically used to differentiate between solid and liquid matter, which are generally passed at different rates through the stomach.

Alternatively or additionally, swallowing is detected by at least one sensor placed at a site on the gastrointestinal tract other than the fundic region of the stomach, and the sensor generates a signal indicative of swallowing. Appropriate sites include, but are not limited to, a site on the esophagus, a site on the stomach, and a site on the throat. Whenever detection of swallowing is described in the present patent application with respect to fundic activity, it is to be understood as being by way of example, and not as excluding detection by a sensor located elsewhere on the gastrointestinal tract.

Typically, measurement of the intensity and/or duration of the electrical activity in the antral region is correlated with aspects of fundic electrical activity denoting swallowing, as described hereinbelow, such that ingested matter of differing chemical and nutritional content can be distinguished. Further typically, the amount of food accumulated in the fundus or antrum is estimated by measuring a level of electrical activity at various sites in the stomach.

Typically, electrical activity response criteria of the stomach of an individual patient are determined and calibrated by measuring the response of the patient's stomach to various types of solid and liquid food. To ensure appropriate compliance, calibration is typically performed under the supervision of a healthcare worker. For illustration, a table such as the following may be established for a particular patient. Except with respect to the example of sugarless chewing gum, these illustrative values are shown with respect to a constant volume of food or liquid ingested (e.g., 100 ml of steak, water, or tomato juice).

TABLE I

| Substance                     | Fundic<br>activity<br>level | Antral<br>activity<br>level | Time delay until<br>onset of antral<br>activity |
|-------------------------------|-----------------------------|-----------------------------|---|
| Sugarless chewing gum         | 1                           | 1                           | _   |
| Non-caloric liquid - Water    | 2                           | 1                           | _   |
| Caloric liquid - Tomato juice | 2                           | 2                           | <1 Minute                                       |
| Caloric liquid - Milk         | 2                           | 2                           | <1 Minute                                       |
| Solid - Apple                 | 2                           | 2                           | Minutes   |
| Solid - Meat                  | 2                           | 3                           | Minutes   |
|                               |                             |                             |   |

In this illustration, the measured data are typically analyzed to determine signal characteristics corresponding to the indicated fundic and antral electrical activity levels. For example, calibration of fundic activity during the chewing of sugarless gum typically yields a low level indication of swallowing, while calibration during the swallowing of liquids and solids yields a greater fundic response. Similarly, there is typically no significant antral response to the patient drinking water, while calibration during the digestion of liquids or solids having higher caloric content yields a greater antral response. Measurements are typically made of the delay time between swallowing and the commencement of antral activity, because consumption of liquids is typically characterized by a rapid transition from the fundus to the antrum, while solids typically stay in the fundus for at least about 10 minutes prior to being passed to the antrum. Typically, a large variety of liquids and solids are used to establish a profile of electrical response characteristics for each patient.

In some embodiments of the present invention, eating detection is accomplished by monitoring the impedance of the fundus and the rate of the antral slow waves, whereby an

eating event is indicated when both the impedance of the fundus and the rate of the antral slow waves cross threshold values within a certain time period. Threshold values may be (a) generally predetermined, or (b) determined for each individual patient during a calibration process, in which the 5 patient ingests various types of food while the impedance of the fundus and the rate of the antral slow waves are monitored, along with other relevant physiological data.

The threshold values indicating an eating event are typically updated to ensure accurate detection of eating by the 10 patient. For some applications, the threshold values indicative of eating are modified through the use of a control unit that adapts the threshold values by checking that an indicated eating event corresponds to an actual eating event. Such checking may include relying on the patient to periodically 15 verify or deny an eating event and/or through additional sensor information. For example, a repeated false positive indication of eating due to normal gastric activity would cause one or more of the threshold values used to signify an eating event to be modified.

Alternatively or additionally, the control unit is adapted to change one or more of the threshold values in response to a physiological event that has a tendency to cause false indications of eating activity. For example, a phenomenon known as change in rhythm of antral electrical activity. This change in antral electrical activity is largely unrelated to eating, but can lead to false indications of eating activity. Since MMC activity lasts about 10 minutes and appears in a cyclical manner with a period of about 40 minutes, the control unit is adapted 30 to identify MMC activity and respond, such that false positive identifications of eating activity are reduced. For example, when a change in antral electrical activity is detected, which may be indicative of eating, the control unit examines data on antral electrical activity from 30 to 50 minutes prior thereto, 35 may be induced by pacing the rectus abdominus muscle. searching for similar activity that may be indicative of MMC activity. If the current activity is likely to be related to MMC activity, then the fundic threshold level signifying an eating event is increased during the subsequent times that are between 30 and 50 minutes in the future (i.e., when subse-40 quent MMC activity is expected), thus reducing the likelihood of false positives relating to MMC contractions. As appropriate, other periodic physiological activities of the gastrointestinal system are treated in a similar manner. It is to be understood that the period of the MMC activity is described 45 herein as being between 30 and 50 minutes by way of illustration and not limitation. In some patients, the period of the MMC activity may be higher, e.g., 50 to 90 minutes, or 90 to 120 minutes. For some applications, a calibration period is provided to determine the length of the period for each 50

For some applications, various supplemental sensors are also applied to the gastrointestinal tract or elsewhere on or in the patient's body. These supplemental sensors, which may comprise pH sensors, blood sugar sensors, ultrasound trans- 55 ducers or mechanical sensors, typically convey signals to a control unit of the apparatus indicative of a characteristic of solids or liquids ingested by the patient. For example, an ultrasound transducer may be coupled to indicate whether ingesta are solid or liquid, and a pH sensor may indicate that 60 an acidic drink such as tomato juice was consumed rather than a more basic liquid such as milk.

In some embodiments, the collected data are stored and intermittently uploaded to an external computer, typically by a wireless communications link, for review by the patient's 65 physician, to enable monitoring of the patient's adherence to a dietary regimen.

For some applications, a specific schedule of allowed food ingestion is pre-programmed by the physician into the memory, and a processor is continuously operative to detect whether food consumption is taking place in accordance with the programmed schedule. For some patients, the schedule may be less strict with respect to drinking certain types of liquids, and more strict with respect to eating certain types of solid food. When an exception from the schedule is detected, the processor typically actuates a signal generator to convey an ingestion-control signal to the patient, in order to encourage the patient to adhere to the schedule. Typically, but not necessarily, apparatus and methods described in U.S. Provisional Patent Application 60/259,925, entitled, "Regulation of eating habits," filed Jan. 5, 2001, and in a PCT Patent Application PCT/IL02/00007, entitled, "Regulation of eating habits," filed Jan. 3, 2002, which published as WO 02/053093, both of which are assigned to the assignee of the present patent application and incorporated herein by reference, are utilized in the administration of the ingestion-con-20 trol signal. Alternatively or additionally, the signal generator generates a visual, audio, or other cue or causes another reasonable discomfort to encourage the patient to adhere to

For embodiments in which this form of dietary monitoring the migrating motor complex (MMC) is characterized by a 25 is supplemented by dietary regulation, the apparatus typically compares the indications of actual food and drink consumption with the pre-programmed schedule. In the event of a sufficient level of patient non-compliance, the ingestion-control signal is typically delivered to the patient's stomach via a set of electrodes placed in a vicinity thereof, so as to induce a sensation of discomfort or minor nausea. For example, an unpleasant sensation, such as nausea, may be induced by altering the natural electrical activity of the stomach, thereby inducing gastric dysrhythmia, or, alternatively, discomfort

Alternatively or additionally, the signal is applied to another site on or in the patient's body. For example, the ingestion-control signal may be applied mechanically or electrically in a vicinity of the cochlear nerve, so as to induce vertigo. Alternatively, the signal is applied so as to generate a brief pain sensation anywhere on the patient's body, which only recurs if the patient continues to eat. Further alternatively, the signal is applied to the esophagus or to the lower esophageal sphincter, so as to cause contraction of muscle tissue therein, thereby making any further eating difficult or very uncomfortable.

Alternatively or additionally, the ingestion-control signal is configured so as to induce a feeling of satiation, typically but not necessarily in accordance with methods described in U.S. patent application Ser. No. 09/734,358, entitled, "Acute and chronic electrical signal therapy for obesity," filed on Dec. 21, 2000, which issued as U.S. Pat. No. 6,600,953, and which is assigned to the assignee of the present patent application and is incorporated herein by reference. For example, methods described in that application for engendering a feeling of satiation may be applied in conjunction with embodiments of the present invention, such that muscles in the vicinity of stretch receptors in the stomach are caused to contract, thereby resulting in decreased hunger sensations. Alternatively or additionally, the feeling of satiation is induced by applying electrical signals which enhance the mobility of chyme from the fundus to the antrum of the stomach, where stretch-receptor signals are generally generated to a greater extent for a given quantity of food than in the fundus.

In another embodiment, when an exception from the schedule of allowed food ingestion is detected, the processor typically conveys the exception to an external operator con(

trol unit, which in turn wirelessly communicates the exception in real time to a remote computer system. The remote computer system can be configured to analyze the exception based on predetermined rules and, if necessary, perform an appropriate action, such as notification of a healthcare 5 worker, care provider, or family member of the patient, in order to encourage the patient to adhere to the schedule.

Typically, the schedule of allowed food ingestion can be modified after implantation of the apparatus, typically by means of a wireless communications link. In this manner, the schedule can be adjusted in response to changes in the patient's eating habits and experience with the apparatus.

In an embodiment, antral electrical activity of a subject is monitored, and a signal is applied to a vagus nerve of the subject in temporal coordination with the monitored antral electrical activity. For example, the signal may be applied during a slow wave as indicated by antral electrical or mechanical activity, or by other means. Alternatively or additionally, the signal is applied within 5 or 10 seconds before an anticipated slow wave, or within 5 or 10 seconds after a slow wave. As appropriate, this technique of vagus nerve stimulation may be coordinated, alternatively or additionally, with measurements of fundic impedance.

Typically, bursts of antral electrical activity occur several 25 times a minute. In an embodiment, a signal is applied to the vagus nerve synchronized with each burst (e.g., (a) during the burst, (b) shortly following a defined feature of the burst, (c) prior to an anticipated feature of the burst, or (d) following a slow wave). For some applications, techniques described: (a) 30 herein, (b) in the above-cited U.S. Pat. Nos. 5,188,104 and 5,263,480 to Wernicke et al., and/or (c) in the other references cited in the Background section of this application, are adapted for use in carrying out this embodiment of the present invention.

There is therefore provided, in accordance with an embodiment of the present invention, gastric apparatus, including:

a gastrointestinal sensor, adapted to be coupled to a gastrointestinal site of a subject and to generate a gastrointestinal sensor signal responsive to a property of the gastrointestinal 40 site;

a set of one or more antral sensors, adapted to be coupled to an antral site of an antrum of the stomach and to generate an antral sensor signal responsive to a property of the antrum; and

a control unit, adapted to receive and analyze the gastrointestinal and antral sensor signals, and to determine, responsive thereto, a characteristic of a food ingested by the subject.

Typically, the control unit is adapted to be implanted in the 50 subject.

In an embodiment, the characteristic of the ingested food includes a caloric content of the ingested food, and the control unit is adapted to determine the caloric content. Alternatively or additionally, the characteristic of the ingested food 55 includes a chemical content of the ingested food, and the control unit is adapted to determine the chemical content. Further alternatively or additionally, the characteristic of the ingested food includes a nutritional content of the ingested food, and the control unit is adapted to determine the nutritional content.

For some applications, the apparatus includes an operator unit, which is adapted to be disposed external to the subject and to transmit a control signal to the control unit.

In an embodiment, the gastrointestinal sensor is adapted to 65 generate a swallowing sensor signal responsive to swallowing by the subject. Typically, the gastrointestinal sensor is

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adapted to be placed at an esophageal site of the subject, a site of the stomach of the subject, and/or a site of a throat of the subject.

Typically, the gastrointestinal sensor includes a set of one or more fundic sensors, adapted to be coupled to a fundic site of a fundus of the stomach of the subject and to generate a fundic sensor signal responsive to a property of the fundus, and the control unit is adapted to receive and analyze the fundic and antral sensor signals, and to determine, responsive thereto, the characteristic of the ingested food. In an embodiment, the fundic sensor set includes one or more strain gauges. Alternatively or additionally, the antral sensor set includes one or more strain gauges.

Typically, the fundic sensor set includes a set of fundic electrodes, adapted to generate a fundic electrode signal responsive to a property of the fundus, the antral sensor set includes a set of antral electrodes, adapted to generate an antral electrode signal responsive to a property of the antrum, and the control unit is adapted to receive and analyze the fundic and antral electrode signals, and to determine, responsive thereto, the characteristic of the ingested food. For example, the control unit may be adapted to determine, responsive to an analysis of at least one of the electrode signals, an amount of the ingested food accumulated in a region of the stomach. Alternatively or additionally, the control unit is adapted to count, responsive to an analysis of at least one of the electrode signals, a number of meals ingested by the subject during a period of time.

The antral electrode set typically includes two antral electrodes, adapted to be coupled to two sites of the antrum, and the control unit is adapted to identify a measure of electrical impedance between the two sites of the antrum. In this case, the control unit is typically adapted to determine the characteristic of the ingested food, responsive to a change in the 35 measure of electrical impedance. For some applications, the fundic electrode set includes two fundic electrodes, adapted to be coupled to two sites of the fundus, and the control unit is adapted to identify a measure of electrical impedance between the two sites of the fundus. For example, the control unit may be adapted to determine the characteristic of the ingested food, responsive to a change in the measure of electrical impedance. Alternatively or additionally, the control unit is adapted to identify an increased measure of electrical impedance relative to a baseline value as indicative of eating. Further alternatively or additionally, the control unit is adapted to identify a substantial return towards a baseline value of the measure of electrical impedance as indicative of a termination of eating.

For some applications, the control unit is adapted to identify an increase in the measure of electrical impedance as indicative of an onset of eating. For example, the control unit may be adapted to detect the onset of eating, responsive to the increase in the measure of electrical impedance being greater than 5 ohms per centimeter of distance between the two sites of the fundus.

In an embodiment, the control unit is adapted to perform a calibration including measurement of a response of the fundic and antral electrode signals to ingestion by the subject of one or more test foods. For example, the one or more foods may include one or more solid foods, and the control unit may be adapted to perform the calibration responsive to ingestion of the one or more solid foods. Alternatively or additionally, the one or more foods includes one or more liquid foods, and the control unit is adapted to perform the calibration responsive to ingestion of the one or more liquid foods. Further alternatively or additionally, the one or more foods includes one or more solid foods and one or more liquid foods, and herein the

control unit is adapted to perform the calibration responsive to ingestion of the one or more solid foods and the one or more liquid foods.

In an embodiment, the antral electrode set is adapted to generate the antral electrode signal responsive to an electrical potential change generated responsive to a contraction of a muscle of the antrum. In this case, the control unit is typically adapted to determine, responsive to an amplitude of the antral electrode signal, the characteristic of the ingested food. Alternatively or additionally, the control unit is adapted to determine, responsive to a frequency of the antral electrode signal, the characteristic of the ingested food. Further alternatively or additionally, the control unit is adapted to determine, responsive to a spike energy per antral cycle of electrical activity, the characteristic of the ingested food. Still further alternatively or additionally, the control unit is adapted to determine, responsive to a duration of the antral electrode signal, the characteristic of the ingested food.

In an embodiment, the control unit is adapted to determine, 20 responsive to a change in a rate of antral electrode signal events, the characteristic of the ingested food. The control unit may alternatively or additionally be adapted to identify an increase in an amplitude of the antral electrode signal as indicative of an onset of a cephalic phase occurring in the 25 subject. For some applications, the control unit is adapted to identify an increase in an amplitude of the antral electrode signal as indicative of an onset of antral digestion. Alternatively or additionally, the control unit is adapted to identify a reduction in a rate of antral electrode signal events as indicative of an onset of antral digestion.

For some applications, the control unit is adapted to identify an increased amplitude of the antral electrode signal relative to a baseline value as indicative of antral digestion. Alternatively or additionally, the control unit is adapted to identify a reduced rate of antral electrode signal events relative to a baseline value as indicative of antral digestion. Further alternatively or additionally, the control unit is adapted to identify a substantial return towards a baseline value of an amplitude of the antral electrode signal as indicative of a 40 termination of antral digestion. Still further alternatively or additionally, the control unit is adapted to identify a substantial return towards a baseline value of a rate of antral electrode signal events as indicative of a termination of antral digestion.

The control unit is typically adapted to determine the char- 45 acteristic of the ingested food, responsive to a time delay between an onset of eating and an onset of a decreased rate of electrical events in the antrum. For some applications, the control unit is adapted to determine the characteristic of the ingested food, responsive to the time delay and responsive to 50 a threshold time delay. Alternatively or additionally, the control unit is adapted to determine, responsive to the time delay, an extent to which the ingested food includes solid food matter. In this case, the control unit is typically adapted to determine that the ingested food includes solid food matter, 55 responsive to the time delay being more than about one minute. For some applications, the control unit is adapted to determine that the ingested food includes predominantly solid food matter, responsive to the time delay being more than about 5 minutes.

In an embodiment, the control unit is adapted to determine, responsive to the time delay, an extent to which the ingested food includes liquid food matter. For example, the control unit may be adapted to determine that the ingested food includes liquid food matter, responsive to the time delay being less than about 5 minutes. Alternatively or additionally, the control unit is adapted to determine that the ingested food

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includes predominantly liquid food matter, responsive to the time delay being less than about one minute.

For some applications, the control unit is adapted to determine the characteristic of the ingested food, responsive to a time delay between an onset of eating and an onset of increased electrical activity in the antrum. In this case, the control unit is typically adapted to determine the characteristic of the ingested food, responsive to the time delay and responsive to a threshold time delay. In an embodiment, the control unit is adapted to determine, responsive to the time delay, an extent to which the ingested food includes solid food matter. For example, the control unit may be adapted to determine that the ingested food includes solid food matter, responsive to the time delay being more than about one minute. Alternatively or additionally, the control unit may be adapted to determine that the ingested food includes predominantly solid food matter, responsive to the time delay being more than about 5 minutes. In an embodiment, the control unit is adapted to determine, responsive to the time delay, an extent to which the ingested food includes liquid food matter. For example, the control unit may be adapted to determine that the ingested food includes liquid food matter, responsive to the time delay being less than about 5 minutes. Alternatively or additionally, the control unit is adapted to determine that the ingested food includes predominantly liquid food matter, responsive to the time delay being less than about one

Typically, the fundic electrode set is adapted to generate the fundic electrode signal responsive to an electrical potential change generated responsive to a contraction of a muscle of the fundus. For example, the control unit may be adapted to determine the characteristic of the ingested food responsive to an amplitude of the fundic electrode signal, a frequency of the fundic electrode signal, and/or a change in a rate of fundic electrode signal events of the fundic electrode signal.

In an embodiment, the control unit is adapted to identify an increased amplitude of the fundic electrode signal relative to a baseline value as indicative of eating. Alternatively or additionally, the control unit is adapted to identify an increased frequency of the fundic electrode signal relative to a baseline value as indicative of eating. Further alternatively or additionally, the control unit is adapted to identify a substantial return towards a baseline value of an amplitude of the fundic electrode signal as indicative of a termination of eating. Still further alternatively or additionally, the control unit is adapted to identify a substantial return towards a baseline value of a frequency of the fundic electrode signal as indicative of a termination of eating. For some applications, the control unit is adapted to identify an increase in an amplitude of the fundic electrode signal as indicative of an onset of eating. For example, the control unit may be adapted to detect the onset of eating responsive to the increase in the amplitude of the fundic electrode signal being greater than about 20

In an embodiment, the control unit is adapted to identify an increase in a frequency of the fundic electrode signal as indicative of an onset of eating. For example, the control unit may be adapted to detect the onset of eating, responsive to the increase in the frequency being greater than about 10 percent.

In an embodiment, the control unit includes a memory, adapted to store a result of the analysis performed by the control unit. Typically, the memory is adapted to upload the stored result to an external computer, e.g., by using a wireless communications link.

In an embodiment, the apparatus includes a supplemental sensor adapted to be placed at a site of the subject and to

convey a supplemental sensor signal to the control unit. The control unit is typically adapted to receive and analyze the supplemental sensor signal, and to determine, responsive thereto, the characteristic of the ingested food. Alternatively or additionally, the control unit is adapted to receive and 5 analyze the supplemental sensor signal, and to determine, responsive thereto, an onset of eating by the subject. Further alternatively or additionally, the control unit is adapted to receive and analyze the supplemental sensor signal, and to determine, responsive thereto, eating by the subject. Still 10 further alternatively or additionally, the control unit is adapted to receive and analyze the supplemental sensor signal, and to determine, responsive thereto, a termination of eating by the subject. Typically, the supplemental sensor includes an electrode, a pH sensor, a blood sugar sensor, an 15 ultrasound transducer, and/or a mechanical sensor. In an embodiment, the supplemental sensor is adapted to be placed at a gastrointestinal site of the subject, an esophageal site of the subject, a site of the stomach of the subject, and/or a site of a throat of the subject.

For some applications, the control unit includes a memory, adapted to store a schedule of allowed food ingestion, wherein the apparatus includes an operator unit, which is adapted to be disposed external to the subject, and wherein the operator unit is adapted to generate an external cue when 25 the analysis performed by the control unit is indicative of the subject not eating in accordance with the ingestion schedule. For example, the external cue may include a visual cue, and the operator unit is adapted to generate the visual cue. Alternatively or additionally, the external cue includes an audio 30 cue, and the operator unit is adapted to generate the audio cue. For some applications, the operator unit includes a user override, adapted to be used by the subject and adapted to disable the cue. Alternatively or additionally, the operator unit is adapted to modify the schedule stored in the memory. For 35 example, the operator unit may be adapted to modify the schedule responsive to information obtained by the operator unit, e.g., via a wireless communications link.

In an embodiment, the apparatus includes a set of one or more current-application electrodes, adapted to be coupled to 40 a tissue of the subject, and wherein the control unit is adapted to drive a current, responsive to the analysis, through the set of current-application electrodes into the tissue. For example, the current-application electrode set may be adapted to be placed at an aural site of the subject, at an esophageal site of 45 the subject, and/or at a site of the stomach of the subject. For some applications, the control unit is adapted to drive the current into the tissue responsive to the characteristic of the ingested food. In an embodiment, the control unit is adapted to apply a pacing signal to a rectus abdominus muscle of the 50 subject. For some applications, the control unit is adapted to drive the current into the tissue responsive to a time of the subject eating.

In an embodiment, the control unit is adapted to configure rhythmia. Alternatively or additionally, the control unit is adapted to configure the current such that driving the current disrupts coupling of gastric mechanical activity and gastric electrical activity of the subject. Further alternatively or additionally, the control unit is adapted to configure the current 60 property of the antrum; and such that driving the current induces a sensation of discomfort in the subject. Still further alternatively or additionally, the control unit is adapted to configure the current such that driving the current induces a sensation of nausea in the subject. Yet further alternatively or additionally, the control unit 65 is adapted to configure the current such that driving the current induces a sensation of vertigo in the subject.

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In an embodiment, the control unit is adapted to drive the current-application electrode set to apply an Excitable-Tissue Control (ETC) signal to the tissue. For example, the control unit may be adapted to drive the current-application electrode set to apply a stimulatory pulse at a site of application of the ETC signal. Alternatively or additionally, the control unit is adapted to drive the current-application electrode set to apply a stimulatory pulse to tissue at a site other than a site of application of the ETC signal. Still further alternatively or additionally, the control unit is adapted to drive the currentapplication electrode set to apply the ETC signal in order to increase an aspect of contraction of the tissue. For some applications, the control unit is adapted to drive the currentapplication electrode set to apply the ETC signal in order to cause tissue contraction in a first portion of the stomach of the subject, and stretching of a stretch receptor of the stomach in a second portion of the stomach. Alternatively or additionally, the control unit is adapted to drive the current-application 20 electrode set to apply the ETC signal in order to increase a contraction strength of tissue in a vicinity of a stretch receptor of the stomach of the subject, so as to increase a sensation of satiation of the subject. Further alternatively or additionally, the control unit is adapted to drive the current-application electrode set to apply the ETC signal to the tissue so as to enhance movement of chyme from a fundus to the antrum of the stomach of the subject.

In an embodiment, the control unit includes a memory, adapted to store a schedule of allowed food ingestion, and wherein the control unit is adapted to withhold driving the current when the analysis performed by the control unit is indicative of the subject eating in accordance with the ingestion schedule. Typically, the ingestion schedule includes types of foods and associated amounts permitted during a time period, and the control unit is adapted to withhold driving the current when the analysis is indicative of the subject eating in accordance with the ingestion schedule. Alternatively or additionally, the ingestion schedule includes a number of meals permitted during a time period, and the control is adapted to withhold driving the current when the analysis is indicative of the subject eating in accordance with the ingestion schedule. Further alternatively or additionally, the ingestion schedule includes an amount of food permitted at a certain meal, and the control is adapted to withhold driving the current when the analysis is indicative of the subject eating in accordance with the ingestion schedule.

Typically, the memory is adapted to download a new schedule from an external computer. For some applications, the apparatus includes an operator unit, which is adapted to be disposed external to the subject and to transmit a control signal to the control unit. In an embodiment, the operator unit includes a user override, adapted to be used by the subject and adapted to withhold driving the current.

There is further provided, in accordance with an embodithe current such that driving the current induces gastric dys- 55 ment of the present invention, a method for analyzing gastric function of a stomach of a subject, including:

> sensing a property of a gastrointestinal tract of the stomach; sensing a property of an antrum of the stomach;

analyzing the property of the gastrointestinal tract and the

determining, responsive to the analysis, a characteristic of a food ingested by the subject.

There is also provided, in accordance with an embodiment of the present invention, gastric apparatus, including:

one or more sensors, adapted to generate respective sensor signals responsive to activity of a gastrointestinal tract of a subject; and

a control unit, adapted to:

receive and analyze the sensor signals,

determine that an eating event has occurred, responsive to at least one of the sensor signals and a threshold,

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identify an aspect of at least one of the sensor signals 5 deriving from rhythmic activity of the gastrointestinal tract that is not indicative of current eating by the subject, and

modify the threshold responsive to the aspect of the signals that derives from activity that is not indicative of current eating.

In an embodiment, the control unit is adapted to modify the threshold if the aspect is indicative of a migrating motor complex (MMC).

In an embodiment:

in modifying the threshold, the control unit is adapted to modify the threshold in a threshold-modification direction,

the control unit is adapted, at a later time at least 30 minutes 20 following modifying the threshold, to further modify the threshold in the threshold-modification direction responsive to identifying at the later time the aspect of the signals deriving from activity that is not indicative of current eating by the subject.

In an embodiment, the control unit is adapted, at a later time at least 30 minutes following modifying the threshold, to at least partially restore the threshold towards a previous value thereof, responsive to not identifying at the later time the aspect of the signals deriving from activity that is not 30 indicative of current eating by the subject.

In an embodiment, the control unit is adapted to modify the threshold responsive to a relationship between a previous portion of the sensor signals and a current portion of the sensor signals.

In an embodiment, the control unit is adapted to modify the threshold responsive to the relationship, the previous portion being between about 30 and about 50 minutes prior to the current portion.

In an embodiment, the control unit is adapted to identify 40 the previous portion of the sensor signals as being indicative of a migrating motor complex (MMC) and to identify the current portion of the sensor signals as being indicative of a MMC, and to modify the threshold responsive to identifying the previous and current portions as being indicative of the 45 MMC.

There is additionally provided, in accordance with an embodiment of the present invention, apparatus, including:

a sensor, adapted to generate a signal responsive to antral electrical activity of a subject; and

a vagus nerve stimulator, adapted to receive the signal and, responsive thereto, to stimulate a vagus nerve of the subject in temporal coordination with an aspect of the sensed signal.

There is still additionally provided, in accordance with an embodiment of the present invention, a selection method, 55 including:

non-invasively recording a fasting electrogastrogram of a natient: and

identifying the patient as a candidate for implantation of a medical device responsive to a rate of slow wave activity of 60 the patient being greater than a threshold rate.

In an embodiment, the threshold rate is at least 2.9 cycles

In an embodiment, identifying the patient includes identifying the patient as a candidate for implantation of a device 65 capable of applying an ETC signal, responsive to the rate of slow wave activity being greater than the threshold rate.

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In an embodiment, the method includes rejecting the patient as a candidate for implantation of the medical device responsive to the rate of slow wave activity being less than 2.9 cycles per minute.

There is also provided, in accordance with an embodiment of the present invention, gastric apparatus, including:

one or more sensors, adapted to generate respective sensor signals responsive to activity of a gastrointestinal tract of a subject;

an implantable control unit, comprising a rechargeable battery and at least one first coil, the implantable control unit adapted to receive the sensor signals, and transmit data responsive thereto; and

an external control unit, comprising a power source and at least one second coil, the external control unit adapted to:

drive the power source to inductively transfer energy via the second coil to the first coil, so as to recharge the battery,

receive the transmitted data.

There is further provided, in accordance with an embodiment of the present invention, gastric apparatus, including:

one or more sensors, adapted to generate respective sensor signals responsively to a gastrointestinal (GI) tract physiological parameter of a GI tract of a subject;

an implantable control unit, including a rechargeable battery and a first set of one or more transducers, the implantable control unit adapted to receive the sensor signals, and transmit data responsively thereto, using one or more of the transducers of the first set; and

an external control unit, including a power source and a second set of one or more transducers, the external control unit adapted to:

drive the power source to wirelessly transfer energy via one or more of the transducers of the second set, and receive the transmitted data, using one or more of the transducers of the second set, and

the implantable control unit is adapted to receive the transmitted energy, using one or more of the transducers of the first set, and charge the battery using the energy.

In an embodiment, the GI tract physiological parameter includes a parameter indicative of electrical activity of the GI tract, and the sensors are adapted to generate the signals responsively to the electrical activity parameter. Alternatively or additionally, the GI tract physiological parameter includes a parameter indicative of an impedance of the GI tract, and the sensors are adapted to generate the signals responsively to the impedance parameter. Further alternatively or additionally, the GI tract physiological parameter includes a parameter indicative of stretching of the GI tract, and the sensors are adapted to generate the signals responsively to the stretching parameter.

In an embodiment, the external control unit is adapted to transmit information to the implantable control unit.

For some applications, at least one of the transducers of the first set includes a coil, and at least one of the transducers of the second set includes a coil. Alternatively, at least one of the transducers of the first set includes a piezoelectric transducer, and at least one of the transducers of the second set includes a piezoelectric transducer. For some applications, the implantable control unit is adapted to transmit the data using the same one or more transducers of the first set that the implantable control unit uses to receive the energy. Alternatively or additionally, the external control unit is adapted to transfer the energy using the same one or more transducers of the second set that the external control unit uses to receive the transmitted data.

For some applications, the apparatus includes an external recharging element, and the power source of the external control unit includes a rechargeable battery adapted to be recharged using the recharging element.

In an embodiment, the apparatus includes an implantable 5 stimulator coupled to the implantable control unit, and the implantable control unit is adapted to drive the stimulator to apply stimulation to the GI tract.

For some applications, the implantable control unit is adapted to configure the stimulation to treat diabetes and/or 10 obesity of the subject.

In an embodiment, the external control unit is adapted to monitor, using the received data, information regarding the applied stimulation, the information selected from the group consisting of: an amount of time per day that the implantable control unit drives the stimulator to apply the stimulation to the GI tract, and a number of times per day that the implantable control unit drives the stimulator to apply the stimulation to the GI tract. For some applications, the external control unit includes an output element, which is adapted to output the 20 information regarding the applied stimulation.

In an embodiment, the GI tract includes a stomach of the subject, and the one or more sensors are adapted to generate the respective sensor signals responsively to the GI tract physiological parameter of the stomach. For some applications, the external control unit is adapted to monitor a gastric slow wave rate of the subject responsively to the received data.

In an embodiment, the external control unit is adapted to store information regarding eating habits of the subject. For 30 some applications, the external control unit includes an output element, which is adapted to output the eating habit information. For some applications, the external control unit is adapted to transmit the eating habit information to the implantable control unit, and the implantable control unit is 35 adapted to detect eating of the subject responsively to the sensor signals using at least one eating detection parameter set responsively to the eating habit information. For some applications, the external control unit is adapted to transmit the eating habit information to the implantable control unit, 40 and the implantable control unit is adapted to detect eating of the subject responsively to the sensor signals using an eating detection algorithm having a threshold set responsively to the eating habit information.

For some applications, the external control unit includes at 45 least one input element, and the external control unit is adapted to determine at least a portion of the eating habit information in response to information input by a human via the input element. For some applications, the eating habit information is reflected in a meal log, and the input element is 50 adapted to accept input of the meal log by the human.

For some applications, the external control unit is adapted to determine at least a portion of the eating habit information responsively to the received data. For example, the eating habit information may include information selected from the 55 group consisting of: a number of meals consumed per time period by the subject, a quantity of food consumed by the subject, a quantity of food consumed per time period by the subject, and a number of eating detections per time period, and the external control unit is adapted to store the selected 60 eating habit information.

For some applications, the eating habit information includes an indication that the subject follows a particular eating regimen, and the external control unit is adapted to transmit the indication to the implantable control unit. For 65 some applications, the implantable control unit is adapted to detect eating of the subject responsively to the sensor signals

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using an eating detection algorithm having a threshold set responsively to the indication. For some applications, the apparatus includes an implantable stimulator coupled to the implantable control unit, and the implantable control unit is adapted to drive the stimulator to apply stimulation to the GI tract, and set a parameter of the stimulation at least in part responsively to the indication.

In an embodiment, the external control unit is adapted to receive an indication of a non-GI tract physiological parameter of the subject. For some applications, the external control unit includes an output element, which is adapted to output the indication and eating habit information of the subject. For some applications, the indication of the non-GI tract physiological parameter includes an indication of a body composition of the subject, and the external control unit is adapted to receive the indication of the body composition. For some applications, the indication of the non-GI tract physiological parameter is selected from the group consisting of: a heart rate of the subject, at least one parameter of an ECG of the subject, and a blood pressure of the subject, and the external control unit is adapted to receive the selected indication of the non-GI tract physiological parameter. For some applications, the indication of the non-GI tract physiological parameter includes an indication of a level of exercise of the subject, and the external control unit is adapted to receive the exercise level indication.

For some applications, the indication of the non-GI tract physiological parameter includes an indication of a weight of the subject, and the external control unit is adapted to receive the indication of the weight. For example, the indication of the weight may include an indication selected from the group consisting of: a weight of the subject, and a body mass index of the subject, and the external control unit is adapted to receive the selected indication. For some applications, the external control unit is adapted to store eating habit information selected from the group consisting of: a number of meals consumed per time period by the subject, a quantity of food consumed by the subject, and a quantity of food consumed per time period by the subject, and the external control unit includes an output element, which is adapted to output the weight and the selected eating habit information.

For some applications, the indication of the non-GI tract physiological parameter includes a blood glucose level of the subject, and the external control unit is adapted to receive the blood glucose level. For some applications, the external control unit is adapted to receive an indication of eating of a meal by the subject, and the external control unit includes an output element, which is adapted to output the blood glucose level in conjunction with the indication of meal eating.

For some applications, the apparatus includes an implantable stimulator coupled to the implantable control unit; the external control unit is adapted to transmit, to the implantable control unit, information selected from the group consisting of: the indication, and information derived from an analysis of the indication; and the implantable control unit is adapted to drive the stimulator to apply stimulation to the GI tract. For some applications, the implantable control unit is adapted to modify a parameter of the stimulation at least in part responsively to the information. For example, the stimulation parameter may include an intensity of the stimulation, and the implantable control unit is adapted to modify the intensity of the stimulation at least in part responsively to the information, and/or the stimulation parameter may include a timing parameter of the stimulation, and the implantable control unit is adapted to modify the timing parameter of the stimulation at least in part responsively to the information.

For some applications, the implantable control unit is adapted to change a mode of the stimulation at least in part responsively to the information. For some applications, the non-GI tract physiological parameter includes a blood glucose level of the subject, and the implantable control unit is 5 adapted to change the mode of the stimulation from a glucoselevel-control mode to an early-satiety-induction mode, in response to a determination that the blood glucose level is at a desired level. Alternatively or additionally, the non-GI tract physiological parameter includes a weight of the subject, and the implantable control unit is adapted to increase a relative amount of time that the mode of the stimulation is a glucoselevel-control mode compared to an early-satiety-induction mode, responsively to the weight.

In an embodiment, the external control unit is adapted to be coupled to a remote service provider. For some applications, the apparatus includes an external cradle, and the external control unit is adapted to be removably coupled to the cradle, and to be coupled to the service provider via the cradle. For 20 some applications, the external control unit is adapted to receive information from the service provider. Alternatively or additionally, the external control unit is adapted to send information to the service provider selected from the group consisting of: at least a portion of the received data, and 25 second set, and the implantable control unit includes a information derived from an analysis of at least a portion of the received data.

There is yet further provided, in accordance with an embodiment of the present invention, gastric apparatus for use with a scale, the apparatus including:

one or more sensors, adapted to generate respective sensor signals responsively to a gastrointestinal (GI) tract physiological parameter of a GI tract of a subject;

an implantable control unit, including a first set of one or more transducers, the implantable control unit adapted 35 to transmit data responsively to the sensor signals, using one or more of the transducers of the first set; and

an external control unit, including a second set of one or more transducers, the external control unit adapted to: receive the transmitted data, using one or more of the 40 transducers of the second set, and

be coupled to the scale, and to receive an indication of a weight of the subject from the scale.

In an embodiment, the external control unit includes a power source, and is adapted to drive the power source to 45 wirelessly transfer energy via one or more transducers of the second set, and the implantable control unit includes a rechargeable battery, and is adapted to received the transmitted energy, using one or more transducers of the first set, and charge the battery using the energy.

For some applications, the indication of the weight includes an indication selected from the group consisting of: a weight of the subject, and a body mass index of the subject, and the external control unit is adapted to receive the selected

In an embodiment, the external control unit is adapted to determine eating habit information responsively to the received data, and the external control unit includes an output element, which is adapted to output the weight indication and the eating habit information. For some applications, the eating habit information includes information selected from the group consisting of: a number of meals consumed per time period by the subject, a quantity of food consumed by the subject, a quantity of food consumed per time period by the subject, and a number of eating detections per time period, 65 and the output element is adapted to output the weight indication and the selected eating habit information.

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For some applications, the output element is adapted to output the weight indication in conjunction with the eating habit information. For example, the output element may be adapted to display a graph including both the weight indication and the eating habit information.

There is still further provided, in accordance with an embodiment of the present invention, gastric apparatus for use with a glucose monitor, the apparatus including:

one or more sensors, adapted to generate respective sensor signals responsively to a gastrointestinal (GI) tract physiological parameter of a GI tract of a subject;

an implantable control unit, including a first set of one or more transducers, the implantable control unit adapted to transmit data responsively to the sensor signals, using one or 15 more of the transducers of the first set; and

an external control unit, including a second set of one or more transducers, the external control unit adapted to: receive the transmitted data, using one or more of the transducers of the second set, and

be coupled to the glucose monitor, and receive a blood glucose level of the subject from the glucose monitor.

In an embodiment, the external control unit includes a power source, and is adapted to drive the power source to wirelessly transfer energy via one or more transducers of the rechargeable battery, and is adapted to received the transmitted energy, using one or more transducers of the first set, and charge the battery using the energy.

In an embodiment, the external control unit is adapted to determine eating habit information responsively to the received data, and the external control unit includes an output element, which is adapted to output the glucose level and the eating habit information. For some applications, the eating habit information includes information selected from the group consisting of an occurrence of eating by the subject, a number of meals consumed per time period by the subject, a quantity of food consumed by the subject, a quantity of food consumed per time period by the subject, and a number of eating detections per time period, and the output element is adapted to output the glucose level and the selected eating habit information.

For some applications, the output element is adapted to output the glucose level in conjunction with the eating habit information. For example, the output element may be adapted to display a graph including both the glucose level and the eating habit information.

There is also provided, in accordance with an embodiment of the present invention, gastric apparatus for use in conjunction with a remote service provider, the apparatus including: one or more sensors, adapted to generate respective sensor

signals responsively to a gastrointestinal (GI) tract physiological parameter of a GI tract of a subject;

an implantable control unit, including a first set of one or more transducers, the implantable control unit adapted to transmit data responsively to the sensor signals, using one or more of the transducers of the first set; and

an external control unit, including a second set of one or more transducers, the external control unit adapted to: receive the transmitted data, using one or more of the transducers of the second set, and

send eating-related information to the service provider selected from the group consisting of: at least a portion of the received data, and information derived from an analysis of at least a portion of the received

There is further provided, in accordance with an embodiment of the present invention, apparatus including:

an implantable control unit, including a first set of one or more transducers, the implantable control unit adapted to transmit data using one or more of the transducers of the first set:

an external receiver, including a second set of one or more 5 transducers, the external control unit adapted to receive the data using one or more of the transducers of the second set;

a processing unit, adapted to determine a financial incentive for the subject responsively to at least a portion of the 10 data

In an embodiment, the processing unit is adapted to be located remotely from the subject.

In an embodiment, the processing unit is adapted to be located proximate to the subject.

In an embodiment, the processing unit is adapted to be coupled to a point-of-sale terminal.

There is still further provided, in accordance with an embodiment of the present invention, a method including:

implanting, in a body of a subject, a control unit having a 20 rechargeable battery;

generating one or more signals responsively to a gastrointestinal (GI) tract physiological parameter of a GI tract of the subject, and receiving the signals at the control unit;

transmitting, from the control unit, data responsively to the 25 signals;

transmitting energy from a site outside the body to the control unit, and recharging the battery using the energy; and receiving the transmitted data at the site outside the body.

For some applications, the subject suffers from diabetes, 30 and implanting the control unit includes implanting the control unit in the subject that suffers from the diabetes. Alternatively or additionally, the subject suffers from obesity, and implanting the control unit includes implanting the control unit in the subject that suffers from the obesity. Further alternatively or additionally, the subject suffers from a heart condition, and implanting the control unit includes implanting the control unit in the subject that suffers from the heart condition.

In an embodiment, the method includes providing a financial incentive to the subject responsively to a physiological improvement in the subject determined responsively to the transmitted data. For some applications, providing the financial incentive includes calculating a value of the incentive responsively to a measure of the physiological improvement. 45 Alternatively or additionally, providing the financial incentive includes awarding a coupon to the subject.

There is additionally provided, in accordance with an embodiment of the present invention, a method including:

generating, from within a body of a subject, one or more 50 signals responsively to a physiological parameter of a gastrointestinal (GI) tract of the subject;

transmitting, from within the body, data responsively to the signals;

receiving the data at a site external to the body; and receiving an indication of a weight of the subject.

For some applications, the indication of the weight includes an indication selected from the group consisting of: a weight of the subject, and a body mass index of the subject, and receiving the indication includes receiving the selected 60 indication.

In an embodiment, the method includes providing a financial incentive to the subject based on a measure of successful weight loss, determined responsively to the indication of the weight. For some applications, the measure is selected from 65 the group consisting of: a magnitude of the weight loss, and a rate of the weight loss, and providing the financial incentive

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includes providing the financial inventive based on the selected measure of successful weight loss.

In an embodiment, the method includes: determining eating habit information responsively to the received data; and outputting the weight indication and the eating habit information. For some applications, the eating habit information includes information selected from the group consisting of a number of meals consumed per time period by the subject, a quantity of food consumed by the subject, a quantity of food consumed per time period by the subject, and a number of eating detections per time period, and outputting includes outputting the weight indication and the selected eating habit information. For some applications, outputting includes outputting the weight indication in conjunction with the eating habit information. For example, outputting may include displaying a graph including both the weight indication and the eating habit information.

There is yet additionally provided, in accordance with an embodiment of the present invention, a method including:

generating, from within a body of a subject, one or more signals responsively to a physiological parameter of a gastrointestinal (GI) tract of the subject;

transmitting, from within the body, data responsively to the signals;

receiving the data at a site external to the subject; sensing a blood glucose level of the subject;

determining eating habit information responsively to the received data; and

outputting the sensed glucose level and the eating habit information.

In an embodiment, the method includes providing a financial incentive to the subject based on a measure of blood glucose control, determined responsively to the blood glucose level.

There is also provided, in accordance with an embodiment of the present invention, a method including:

generating, from within a body of a subject, one or more signals responsively to a gastrointestinal (GI) tract physiological parameter of a GI tract of the subject;

transmitting, from within the body, data responsively to the signals;

receiving the data at a site external to the body; and sending eating-related information to a service provider, the information selected from the group consisting of: at least a portion of the received data, and information derived from an analysis of at least a portion of the received data.

For some applications, the eating-related information includes information selected from the group consisting of: an occurrence of eating by the subject, a number of meals consumed per time period by the subject, a quantity of food consumed by the subject, a quantity of food consumed per time period by the subject, and a number of eating detections per time period, and sending the eating-related information includes sending the selected eating-related information.

For some applications, the method includes sending a reminder from the service provider to the site, and communicating the reminder to the subject.

In an embodiment, the method includes analyzing, at the service provider, the eating-related information. For some applications, analyzing the eating-related information includes developing a recommendation based on the eating-related information, and including sending the recommendation from the service provider to the site, and communicating the recommendation to the subject from the site. For example, the recommendation may be selected from the group consisting of: a recommended food, and a recommended recipe, and sending the recommendation includes sending the selected

recommendation. For some applications, the recommendation is commercial in nature, and sending the recommendation includes sending the commercial recommendation.

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There is further provided, in accordance with an embodiment of the present invention, a method including:

implanting a device within a body of a subject; transmitting data from the device;

receiving the data at a site external to the body; and providing a financial incentive to the subject responsively to the data

For some applications, transmitting the data includes wirelessly transmitting the data from the device.

For some applications, the subject suffers from diabetes, and implanting the device includes implanting the device in the subject who suffers from the diabetes. Alternatively or 15 additionally, the subject suffers from obesity, and implanting the device includes implanting the device in the subject who suffers from the obesity. Further alternatively or additionally, the subject suffers from a heart condition, and implanting the device includes implanting the device in the subject who 20 suffers from the heart condition.

For some applications, receiving the data includes logging the data, and providing the financial incentive includes providing the financial incentive responsively to the logged data.

For some applications, providing the financial incentive 25 includes awarding a coupon to the subject.

For some applications, transmitting the data includes transmitting personal data stored in the device.

For some applications, the site includes a point-of-sale site, and receiving the data includes receiving the data at the point- of-sale site, and providing the financial incentive includes providing the financial incentive at the point-of-sale site.

For some applications, providing the financial incentive includes providing a discount on a health-related product or service.

For some applications, providing the financial incentive includes providing an entertainment-related product or service at a reduced price or for free.

In an embodiment, providing the financial incentive includes providing the financial incentive in response to 40 determining, responsively to the data, that the subject has followed a prescribed regimen. For some applications, the regimen includes a diet regimen, and providing the financial incentive includes providing the financial incentive upon determining, responsively to the data, that the subject has 45 adhered to the prescribed diet regimen. Alternatively or additionally, the regimen includes an exercise regimen, and providing the financial incentive upon determining, responsively to the data, that the subject has adhered to the prescribed exercise regimen.

In an embodiment, the site includes a site proximate the subject, and receiving the data at the site includes receiving the data at the proximate site. For some applications, providing the financial incentive includes determining the financial incentive at the proximate site.

In an embodiment, the site includes a site remote from the subject, and receiving the data at the site includes receiving the data at a site proximate the subject, and transmitting the data from the proximate site to the remote site. For some applications, providing the financial incentive includes determining the financial incentive at the remote site.

In an embodiment, providing the financial incentive includes providing the financial incentive upon determining, responsively to the data, that the subject has attainted a health goal. For some applications, the health goal includes a blood 65 glucose control-related goal, and providing the financial incentive includes providing the financial incentive upon

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determining that the subject has achieved the blood glucose control-related goal. For example, the blood glucose control-related goal may include a goal of achieving a peak daily blood glucose level that is lower than a prescribed level, and providing the financial incentive includes providing the financial incentive upon determining that the subject has achieved the peak daily blood glucose level. Alternatively or additionally, the blood glucose control-related goal may include a goal of achieving an average blood glucose level that is lower than a prescribed level, and providing the financial incentive includes providing the financial incentive upon determining that the subject has achieved the average blood glucose level.

In an embodiment, providing the financial incentive includes: analyzing the data to determine a physiological improvement of the subject; and providing the financial incentive responsively to the physiological improvement. For some applications, providing the financial incentive includes calculating a value of the incentive responsively to a measure of the physiological improvement.

In an embodiment, implanting the device includes applying stimulation from the device to tissue of the subject, and transmitting the data includes transmitting the data responsively to a parameter of the applied stimulation. For some applications, applying the stimulation includes applying the stimulation to a gastrointestinal (GI) tract of the subject, a pancreas of the subject, a heart of the subject, and/or nervous tissue of the subject.

In an embodiment, implanting the device includes implanting a device that includes at least one sensor, and transmitting the data includes generating a sensor signal using the sensor, and transmitting the data responsively to the sensor signal. For some applications, generating the sensor signal includes generating the sensor signal responsively to a physiological parameter of a pancreas of the subject. For some applications, generating the sensor signal includes generating the sensor signal responsively to a blood glucose level of the subject. For some applications, generating the sensor signal includes generating the sensor signal responsively to a state of a gastrointestinal (GI) tract of the subject.

For some applications, generating the sensor signal includes generating the sensor signal responsively to a physiological parameter of a gastrointestinal (GI) tract of the subject. For some applications, providing the financial incentive includes: sending eating-related information to a service provider selected from the group consisting of: at least a portion of the data, and information derived from an analysis of at least a portion of the data; and determining the financial incentive at the service provide responsively to the eating-related information.

For some applications, generating the sensor signal includes generating the sensor signal responsively to a physical activity level of the subject. For example, generating the sensor signal may include sensing acceleration of the subject.

For some applications, providing the financial incentive includes providing the financial incentive in response to determining, responsively to the data, that the subject has followed an exercise regimen.

In an embodiment, the method includes receiving an indication of a weight of the subject, and providing the incentive responsively to the weight indication and the data. For example, the indication of the weight may include an indication selected from the group consisting of: a weight of the subject, and a body mass index of the subject, and receiving the indication includes receiving the selected indication.

The present invention will be more fully understood from the following detailed description of embodiments thereof, taken together with the drawings, in which:

#### BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a schematic illustration of apparatus for treating obesity, in accordance with a typical embodiment of the present invention;

FIG. 2 is a schematic block diagram showing a control unit <sup>10</sup> of the apparatus of FIG. 1, in accordance with a typical embodiment of the present invention;

FIG. 3 is a graph showing a default fundus threshold value signifying an eating event, and modifications of the threshold value, in accordance with an embodiment of the present 15 invention:

FIG. 4 is a schematic diagram showing experimental apparatus used to measure electrical responses to eating in the stomach of a normal rabbit, in accordance with a typical embodiment of the present invention;

FIG. 5 is a graph showing electrical activity in the fundus of a normal rabbit before and during eating, and results of analysis thereof, in accordance with a typical embodiment of the present invention;

FIG. **6A** is a graph showing electrical activity in the fundus <sup>25</sup> of a normal rabbit during and following eating, and results of analysis thereof, in accordance with a typical embodiment of the present invention:

FIG. **6**B is a graph showing details of electrical and mechanical activity recorded during the taking of the data <sup>30</sup> shown in FIG. **6**A;

FIG. 7 is a graph showing detail of electrical fundic activity, measured in accordance with a typical embodiment of the present invention;

FIG. **8** is a graph showing the rate of electrical events in the 35 antrum of a normal dog before and during eating, and results of analysis thereof, in accordance with a typical embodiment of the present invention;

FIG. 9 is a graph showing the rate of electrical events in the antrum of a normal dog before, during, and after eating, and 40 results of analysis thereof, in accordance with a typical embodiment of the present invention;

FIG. **10** is a graph showing electrical and mechanical activity and the rate of electrical events in the antrum of a normal dog before, during, and after eating, and results of analysis 45 thereof, in accordance with a typical embodiment of the present invention;

FIG. 11 is a graph showing fundic electrical activity in a normal dog during several periods of eating and non-eating, and results of analysis thereof, in accordance with a typical 50 embodiment of the present invention; and

FIG. 12 is a schematic illustration of a portable control charger, in accordance with an embodiment of the present invention.

#### DETAILED DESCRIPTION OF EMBODIMENTS

FIG. 1 is a schematic illustration of diet evaluation apparatus 18, which detects when a patient 10 swallows, and detects the type and amount of matter ingested, in accordance 60 with a typical embodiment of the present invention. Typically, but not necessarily, apparatus 18 additionally determines, responsive to the detection, whether to apply electrical energy to modify the activity of tissue of patient 10. Apparatus 18 typically comprises mechanical sensors 70, supplemental 65 sensors 72, local sense electrodes 74, operator controls 71, and one or more current-application electrodes 100.

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Electrodes 74 and 100 are typically coupled to the serosal layer of a stomach 20 and/or inserted into the muscular layer of the stomach in the fundic and antral regions. Alternatively or additionally, the electrodes are coupled elsewhere on the stomach, gastrointestinal tract, or to other suitable locations in or on the patient's body. The number of electrodes and sensors, as well as the positions thereof, are shown in FIG. 1 by way of example, and other sites on stomach 20 or in or on the patient's body are appropriate for electrode and sensor placement in other applications of the present invention. Different types of electrodes known in the art are typically selected based on the specific condition of the patient's disorder, and may comprise stitch, coil, screw, patch, basket, needle and/or wire electrodes, or substantially any other electrode known in the art of electrical stimulation or sensing in tissue.

Typically, apparatus 18 is implanted in patient 10 in a manner generally similar to that used to implant gastric pacemakers or other apparatus for stimulating or sensing in the gastrointestinal tract that are known in the art. As appropriate, techniques described in one or more of the references cited in the Background section of the present patent application may be adapted for use with these embodiments of the present invention. Other methods and apparatus useful in carrying out some embodiments of the present invention are described in the above-cited U.S. Provisional Patent Application 60/259, 925, entitled, "Regulation of eating habits," filed on Jan. 5, 2001, and in the above-cited PCT patent application and in the above-cited U.S. patent application Ser. No. 09/734,358, entitled, "Acute and chronic electrical signal therapy for obesity," filed on Dec. 11, 2000, which are assigned to the assignee of the present patent application and are incorporated herein by reference.

FIG. 2 is a schematic block diagram illustrating details of operation of a control unit 90 of apparatus 18, in accordance with a typical embodiment of the present invention. Typically, control unit 90 is implanted in patient 10, and receives signals from mechanical sensors 70, supplemental sensors 72, and local sense electrodes 74, all of which are typically implanted on the gastrointestinal tract of the patient or elsewhere on or in the body of the patient. These sensors and electrodes are typically adapted to provide an "ingestion activity analysis" block 80 of the control unit with information about food ingestion and/or the present state of the stomach.

Typically, using techniques described hereinbelow, analysis block 80 determines each time that the patient swallows, and also the character and amount of the ingested matter. For example, local sense electrodes 74 coupled to the fundus of the stomach may send signals indicative of fundic electrical activity to analysis block 80, and analysis block 80 identifies aspects of these signals that are characteristic of swallowing of food by the patient. Additionally, mechanical sensors 70 and local sensor electrodes 74 coupled to the corpus and antral regions of the stomach typically send signals which 55 analysis block 80 identifies as indicative of the onset, duration, and/or intensity of the digestive process in those regions. Typically, these data are utilized by analysis block 80 to determine a quality of the ingested matter, for example, whether it is predominantly solid or liquid. Alternatively or additionally, these data may be used to determine other characteristics of the ingested material, for example, its nutritional, chemical, and/or caloric content.

In a typical embodiment, analysis block **80** determines the time delay between swallowing (as measured, typically, by local sense electrodes **74** on the fundus) and the commencement of electrical and mechanical activity in the antrum. This delay is typically used to differentiate between the ingestion

of solid and liquid matter, because solids are generally held in the fundus for at least about 10 minutes before being passed to the antrum, while liquids are generally passed to the antrum essentially immediately.

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Alternatively or additionally, the amount of food accumu- 5 lated in the various regions of stomach 20 is estimated by measuring a level of electrical or mechanical activity in a vicinity of those regions. For example, in some embodiments of the present invention, eating detection is accomplished by monitoring the impedance of the fundus and the rate of the antral slow waves, whereby an eating event is indicated when both the impedance of the fundus and the rate of the antral slow waves cross threshold values within a certain time period.

Further alternatively or additionally, analysis block **80** pro- 15 cesses data from supplemental sensors 72 indicative of the blood sugar level of the patient, to enable an evaluation of whether and of what type of food has been ingested.

In order to improve the accuracy of the analyses described hereinabove, analysis block 80 is typically calibrated by mea-20 suring the appropriate electrical response criteria of stomach 20 of patient 10 to various types of solid and liquid food.

For some applications, analysis block 80 stores the results of its analysis in a memory block 88 of control unit 90, and these results are later uploaded to an external computer, typi-25 cally by a wireless communications link, for review by the patient's physician. Alternatively or additionally, analysis block 80 conveys results of its analysis of the inputs from mechanical sensors 70, supplemental sensors 72, and local sense electrodes 74, to a "parameter search and tuning" block 30 84 of control unit 90. The parameter search and tuning block adapts the threshold values indicative of eating by checking that an indicated eating event corresponds to an actual eating event. For example, the parameter search and tuning block may rely on the patient to periodically verify or deny an eating 35 detections/number of actual eating events). event by using operator control 71. Alternatively or additionally, the parameter search and tuning block utilizes additional sensor information such as antrum impedance, which varies during eating while remaining steady in the absence of eating. In an embodiment, a false positive indication of an eating 40 event may cause one or more of the threshold values to be increased, while a false negative may cause one or more of the threshold values to be decreased.

Alternatively or additionally, search and tuning block 84 is adapted to change one or more of the threshold values in 45 response to a physiological event that has a tendency to cause false indications of eating activity. For example, the migrating motor complex (MMC) is characterized by increased antral electrical activity, which can lead to false indications of eating activity. Since MMC activity lasts about 10 minutes 50 and appears in a cyclical manner with a time lag of about 40 minutes between events, the search and tuning block is adapted to identify MMC activity and respond, such that false positive identifications of eating activity are reduced.

In an embodiment, a calibration period is provided in 55 which a record is generated of actual eating events by the subject. As appropriate, the calibration period may be about one day, several days, or longer than a week. The record of actual eating events may comprise, for example, entries made by the subject or another person in an electronic or non- 60 electronic journal, or using other techniques known in the art for detecting swallowing or otherwise detecting eating. In this embodiment, thresholds for fundic impedance and antral electrical activity are set responsive to (a) the record of actual eating events and (b) measurements of fundic impedance and 65 antral electrical activity made during the calibration period. Typically (but not necessarily), some preference is given to

reducing false negatives relative to reducing false positives. In other words, for many applications, it is more important to avoid missing a detection of an eating event than to avoid incorrectly reporting that an eating event occurred. (For other applications, both are of equal importance, or the latter is

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In an embodiment, two parallel matrices are generated in response to the record of actual eating events and the measurements of fundic impedance and antral electrical activity made during the calibration period. The first matrix, a false negative matrix, has a range of thresholds for changes in fundic impedance on the x-axis of the matrix, and a range of antral electrical activity on the y-axis of the matrix. For clarity of description (although not necessarily in practice), the range of fundic impedance extends from normalized values of 1 to 20, and the range of antral electrical activity also extends from 1 to 20. The false negative matrix is then generated as a  $20\times20$ matrix. Each cell in the false negative matrix represents a given combination of possible thresholds of fundic impedance and antral electrical activity. The value stored in a given cell represents a value associated with the extent of false negatives that would have been generated for the given fundic impedance threshold and antral electrical activity threshold represented by that cell. For example, if a normalized fundic impedance threshold of 2 and a normalized antral electrical activity threshold of 7 yielded no false negatives during the calibration period, then the value of the false negative matrix at cell (2, 7) would be zero. Similarly, if a normalized fundic impedance threshold of 20 and a normalized antral electrical activity value of 20 did not result in an identification of any of several actual eating events during the calibration period, then the value of the false negative matrix at cell (20, 20) would be 100%. For some applications, the value of a given cell in the false negative matrix is defined as: 1-(number of correct

A corresponding false positive matrix is generated. The x-axis and y-axis of the false positive matrix are analogous to the corresponding axes of the false negative matrix. Typically, the value in each cell of the false positive matrix reflects the total or average number of false positive indications of eating in a given time period (e.g., one day, or throughout the calibration period).

A calibration period analysis algorithm typically identifies one or more near-minimum values in the false negative matrix. These near-minimum values (NMV's) typically are located in one or more "clouds" on the false negative matrix. Similarly, the calibration period analysis algorithm typically identifies one or more NMV's in the false positive matrix. (The term "near-minimum value" is understood to include actual minimum values, as well.) The near-minimum values are typically located in one or more clouds on the false positive matrix, as well. For some applications, in order to determine a suitable fundic impedance threshold and a suitable antral electrical activity threshold for use during regular operation of an eating detection algorithm, the calibration period analysis algorithm first identifies cells in the false negative matrix that are HMV's, and then determines which of the corresponding cells in the false positive matrix are also

In some cases, only a single set of fundic impedance and antral electrical activity threshold values (x, y) is an NMV in both the false negative and the false positive matrix. In these cases, this set typically defines the thresholds for use in regular operation of the eating detection algorithm.

In other cases, multiple sets (x(i), y(i)) of thresholds are identified that correspond to an NMV in both the false negative matrix and the false positive matrix. In these cases, for

some applications, the calibration period analysis algorithm determines one of the multiple threshold sets that is likely to have a high level of "stability" during regular operation of the eating detection algorithm. To determine stability, the calibration period analysis algorithm typically determines which cell in the false negative matrix having an NMV is not adjacent to or relatively near to one or more cells in the false negative matrix having relatively high values (i.e., indicating many false negatives).

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(a) electrodes implanted on the fundus, for measuring fundic impedance, and (b) electrodes implanted on the antrum, for measuring the rate of antral electrical activity, in accordance with an embodiment of the present invention. During an approximately six hour monitoring period, the patient was free to eat whatever she chose, whenever she chose to eat. During this period, the patient recorded eating three times (pizza bread at 09:45, pasta with cheese at 12:30, and candy at 14:30).

TABLE II

|    |      |      | ;    | Sample | false ne | gative i | natrix |      |      |      |     |
|----|------|------|------|--------|----------|----------|--------|------|------|------|-----|
|    | 18   | 18.5 | 19   | 19.5   | 20       | 20.5     | 21     | 21.5 | 22   | 22.5 | 23  |
| 10 | 33   | 33   | 33   | 33     | 33       | 33       | 33     | 33   | 33   | 67   | 67  |
| 12 | 33   | 33   | 33   | 33     | 33       | 33       | 33     | 33   | 33   | 67   | 67  |
| 14 | 33   | 33   | 33   | 33     | 33       | 33       | 33     | 33   | 33   | 67   | 67  |
| 16 | 33   | 33   | 33   | 33     | 33       | 33       | 33     | 33   | 33   | 67   | 67  |
| 18 | 33   | 33   | 33   | 33     | 33       | 33       | 33     | 33   | 33   | 67   | 67  |
| 20 | 33   | 33   | 33   | 33     | 33       | 33       | 33     | 33   | 33   | 67   | 67  |
| 22 | 33   | 33   | 33   | 33     | 33       | 33       | 33     | 33   | 33   | 67   | 67  |
| 24 | 33   | 33   | 33   | 33     | 33       | 33       | 33     | 33   | 33   | 67   | 67  |
| 26 | 33   | 33   | 33   | 33     | 33       | 33       | 33     | 33   | 33   | 67   | 67  |
| 28 | 33   | 33   | 33   | 33     | 33       | 33       | 33     | 33   | 33   | 67   | 67  |
| 30 | [33] | [33] | [33] | [33]   | [33]     | [33]     | [33]   | [33] | [33] | 67   | 67  |
| 32 | [33] | [33] | [33] | [33]   | *33*     | [33]     | [33]   | [33] | [33] | 67   | 67  |
| 34 | [33] | [33] | [33] | [33]   | [33]     | [33]     | [33]   | [33] | [33] | 67   | 67  |
| 36 | [33] | [33] | [33] | [33]   | [33]     | [33]     | [33]   | [33] | [33] | 67   | 67  |
| 38 | [33] | [33] | [33] | [33]   | [33]     | [33]     | [33]   | [33] | [33] | 67   | 67  |
| 40 | 67   | 67   | 67   | 67     | 67       | 67       | 67     | 67   | 67   | 100  | 100 |
| 42 | 67   | 67   | 67   | 67     | 67       | 67       | 67     | 67   | 67   | 100  | 100 |
| 44 | 100  | 100  | 100  | 100    | 100      | 100      | 100    | 100  | 100  | 100  | 100 |
| 46 | 100  | 100  | 100  | 100    | 100      | 100      | 100    | 100  | 100  | 100  | 100 |

For example, one cell (x1,y1) in the false negative matrix having an NMV of 5% may be relatively near to a second cell (x1-1,y1+2) having an NMV of 30%. Another cell (x2,y2) in the false negative matrix having an NMV of 5% may have no cells within +/-2 on the x-axis or the y-axis having an NMV greater than 25%. In this case, the fundic impedance and antral electrical activity thresholds represented by the second cell would be selected by the calibration period analysis algorithm

In another example, a summing algorithm typically weighted by proximity is used to evaluate the neighborhood (e.g., +/-3 cells) of all cells having an NMV. The cell that is 45 both NMV and having the lowest "false negative sum" for its neighborhood is selected by the calibration period analysis algorithm to represent the fundic impedance and antral electrical activity thresholds during regular operation of the eating detection algorithm.

For some applications, the stability determination described hereinabove is performed with respect to values in the false positive matrix or with respect to values in both the false positive matrix and the false negative matrix.

In an embodiment, if the calibration period analysis algorithm identifies n cells having stable NMV's in both the false negative and false positive matrices, then during regular operation of the eating detection algorithm, an evaluation of each of the corresponding n sets of fundic impedance and antral electrical activity thresholds is performed. A determination of an eating event is made responsive to some or all of the n sets of thresholds (e.g., responsive to the measured fundic impedance and antral electrical activity exceeding the corresponding thresholds of ½ of the n sets).

Table II and Table III (below) display results obtained based on an experiment with an obese human patient having

TABLE III

| _ |    |     |      |     |        |          |          |        |      |     |      |    |
|---|----|-----|------|-----|--------|----------|----------|--------|------|-----|------|----|
| _ |    |     |      | Ş   | Sample | false po | sitive r | natrix |      |     |      |    |
|   |    | 18  | 18.5 | 19  | 19.5   | 20       | 20.5     | 21     | 21.5 | 22  | 22.5 | 23 |
|   | 10 | 7   | 7    | 7   | 6      | 6        | 4        | 4      | 4    | 4   | 4    | 4  |
|   | 12 | 5   | 5    | 5   | 5      | 5        | 4        | 4      | 4    | 4   | 4    | 4  |
|   | 14 | 4   | 4    | 4   | 4      | 4        | 3        | 3      | 3    | 3   | 3    | 2  |
|   | 16 | 3   | 3    | 3   | 3      | 3        | 3        | 3      | 3    | 3   | 3    | 2  |
|   | 18 | 3   | 3    | 3   | 3      | 3        | 3        | 3      | 3    | 3   | 3    | 2  |
|   | 20 | 2   | 2    | 2   | 2      | 2        | 2        | 2      | 2    | 2   | 2    | 1  |
|   | 22 | 2   | 2    | 2   | 2      | 2        | 2        | 2      | 2    | 2   | 1    | 1  |
|   | 24 | 1   | 1    | 1   | 1      | 1        | 1        | 1      | 1    | 1   | 1    | 1  |
|   | 26 | 1   | 1    | 1   | 1      | 1        | 1        | 1      | 1    | 1   | 1    | 1  |
|   | 28 | 1   | 1    | 1   | 1      | 1        | 1        | 1      | 1    | 1   | 1    | 1  |
|   | 30 | [0] | [0]  | [0] | [0]    | [0]      | [0]      | [0]    | [0]  | [0] | 0    | 0  |
|   | 32 | [0] | [0]  | [0] | [0]    | *0*      | [0]      | [0]    | [0]  | [0] | 0    | 0  |
|   | 34 | [0] | [0]  | [0] | [0]    | [0]      | [0]      | [0]    | [0]  | [0] | 0    | 0  |
|   | 36 | [0] | [0]  | [0] | [0]    | [0]      | [0]      | [0]    | [0]  | [0] | 0    | 0  |
|   | 38 | [0] | [0]  | [0] | [0]    | [0]      | [0]      | [0]    | [0]  | [0] | 0    | 0  |
|   | 40 | 0   | 0    | 0   | 0      | 0        | 0        | 0      | 0    | 0   | 0    | 0  |
|   | 42 | 0   | 0    | 0   | 0      | 0        | 0        | 0      | 0    | 0   | 0    | 0  |
|   | 44 | 0   | 0    | 0   | 0      | 0        | 0        | 0      | 0    | 0   | 0    | 0  |
|   | 46 | 0   | 0    | 0   | 0      | 0        | 0        | 0      | 0    | 0   | 0    | 0  |

An experiment-analysis algorithm was set to have a "refractory" period of 30 minutes following each detection of eating, during which a new detection of eating would not be registered. A "hysteresis" period of 8 minutes was set, whereby a detection of eating would only be made if fundic impedance and antral electrical activity passed respective thresholds within 8 minutes of each other. Lastly, an identified eating detection was counted as being a true eating detection if it occurred within 15 minutes prior to or following the time of eating as recorded by the patient.

The x-axis in Table II (and Table III) represents threshold rates of antral electrical activity, in minutes. The y-axis represents fundic impedance, in ohms. Thus, for example, Table

II shows that if the threshold for antral electrical activity were set at 19 seconds (such that only antral electrical activity occurring slower than once every 19 seconds generates an indication of potential eating), and if the threshold for fundic impedance change were set at 16 ohms (such that an indication of potential eating is only generated if the fundic impedance increases by more than 16 ohms), then two actual eating events would have been correctly detected, and one actual eating event would not have been detected. This defines a false negative percentage of 33%.

Similarly, in Table III, the total number of false positive indications of eating are shown in the matrix as a function of particular threshold settings. For example, for a fundic impedance threshold of 19 ohms, and an antral electrical activity threshold rate of 21 seconds, no false positive indications of eating were generated.

Cells in each matrix were automatically analyzed to determine optimal or near-optimal threshold settings for use in regular operation of an eating detection algorithm. Suitable thresholds optimizing both the fundic impedance threshold and the antral electrical activity rate threshold are marked by cells having square brackets surrounding their values. Thus, for this patient, it was found that a fundic impedance threshold ranging from 30 to 38 ohms and an antral electrical activity rate threshold ranging from 18 to 22 seconds is generally optimal. A particular threshold set (32 ohms, 20 seconds) was identified by the automated analysis as being particularly stable (in addition to minimising false positive and false negative indications), and is marked by a "\*\*".

In an embodiment, the following algorithm and/or pseudocode is executed to determine the fundic impedance and antral electrical activity thresholds. Typically, the identified user would be a physician reviewing records of actual eating events ("bookmarks") and measured data.

- 1. Input session
  - 1. User chooses files of data.
  - 2. For each selected file, allow the user to review the bookmarks and have the following options:
    - i. remove/approve each bookmark
    - ii. accept all bookmarks without reviewing
    - iii. add a bookmark (e.g., to represent an actual eating event not previously recorded)
    - iv. impose a fixed time shift on all bookmarks (e.g., in case the "time stamps" of the bookmarks are not aligned with the time stamps of measurements of 45 fundic impedance
  - 3. The user will then select the following detection criteria ranges or approve the default:
    - i. Time before and after a meal bookmark that will be considered as a true detection, if detected (default: 50 +/-15 min)
    - ii. Time before and after a meal that will not be considered as a false detection, if a detection is made (default: from 15 min before bookmark to 45 min after bookmark)
- 2. For all possible rate and impedance thresholds:
  - Find all "eating detection events," i.e., times during which both the fundic impedance and the rate of slow waves cross their corresponding thresholds in a common time frame, typically 3-10 minutes
  - Calculate false negative percentage (FN) (percentage of meals that did not have a detection within the pre-defined "true detection time limits"
  - Calculate false positives per day (FP) (number of detections that did not correspond to an actual eating event, 65 i.e., a bookmark, within the pre-defined "false detection range," divided by the number of days).

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- 3. Identify the "optimal set" of thresholds combination according to the following rule:
  - Find all threshold pairs (fundic impedance/rate of antral electrical activity) that generate the minimal (or near minimal) false negative percentage.
  - 2. In this set of threshold pairs, find the corresponding subset with minimal false positives per day. The results of these two steps are defined as the "optimal set"
- 4. Find closed spaces of points in the optimal set:
  - i. Set cloud number=1
  - ii. Unmark all points in the optimal set
  - iii. Find the unmarked point in the optimal set that has minimal value of F and R values:  $\min_{r,f} \{P(r,f): P \in \text{optimal set}\}$  and mark it
  - iv. Set Rcount=1; Fcount=0;
  - v. If P(r+count,f+Fcount)∈optimal set then
    - 1. mark P(r+Rcount; f+Fcount)
    - 2. Increment Rcount
    - 3. goto (iv)

else

- if there is an unmarked point in the range P(r:r+1, f: f+1) that belongs to the optimal set then
- set Rcount and Fcount so that r+Rcount and f+Fcount will point to that pair
- 2. mark the selected pair
- 3. goto iv

else

- 1. set cloud-number=cloud-number+1
- 2. define a cloud (cloud-number) and assign all marked points from the optimal set to it
- 3. remove all marked points from the optimal set
- 4. goto (iii)
- 5. Find cloud's edges
- 1. set  $P=\min_{r,f}\{P(r,f):P\in cloud\}$
- 2. set count=1
- 3. Unmark all F/R matrix members
- 4. set  $E(count) = min_{Rcount}(P(r+Rcount,f) \notin cloud)$
- 5. mark E(count)
- 6. in the range  $P(R_{E(count)} \pm 1, F_{E(count)} \pm 1)$  find an unmarked point  $P \notin cloud$  for which the set  $N(R_p \pm 1, F_p \pm 1) \notin cloud$  is not empty
- 7. If P exists then
  - a. Set count=count+1
  - b. Set E(count)=P
  - c. Goto 5

Else

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- a. in the range  $P(R_{E(1)}\pm 1, F_{E(1)}\pm 1)$  find an unmarked point  $P\notin \text{cloud}$  for which the set  $N(R_p\pm 1, F_p\pm 1)$   $\notin \text{cloud}$  is not empty
- b. if P exists then
  - 1. Set count=count+1
  - 2. Set E(count)=P
  - 3. Goto 5
- d. Else check if E(1) and E(count) are neighbors
- e. If they are neighbors, then goto 6 else create a set of additional points for the set E according to the following rules:
  - 1. the number of the points will equal max(abs( $R_{E(1)}$   $R_{E(count)}$ ), abs( $F_{E(1)}$   $F_{E(count)}$ ))
  - 2. Each new point will have the FN, FP values of the opposite edge point
- 6. Grade each point in each cloud
- 1. For each cloud
- 2. for each P∈cloud
- 3. grade(P)=0
- 4. for each  $E_{count}$  point in E

- 5.  $\operatorname{grade}(P) = \operatorname{grade}(P) + (\operatorname{FN}(E_{count}) + \operatorname{FP}(E_{count})/4)/\operatorname{dist}(P, P)$
- 7. Choose the 3 points with minimal grade.
- 8. Offer these three combinations to the user, and specify for each of them the values of false negative and false positive 5 it generates.

FIG. 3 is a graph showing an example of a change in a sensing threshold value during suspected periods of MMC activity, in accordance with an embodiment of the present invention. Time period B indicates a time when increased antral electrical activity is detected, which may be indicative of either eating or MMC activity. Search and tuning block 84 then examines data indicating antral electrical activity from 30 to 50 minutes prior to the present activity (i.e., around time A) to determine whether similar activity occurred at time A. If similar activity occurred at time A, the present activity during time period B is of increased likelihood to be related to MMC activity. Therefore, as shown in FIG. 3, the fundus threshold level is increased during time period B, thus reducing the 20 likelihood of false positives relating to MMC activity. In this manner, fundus impedance levels measured prior to time period B generally only need to exceed the default value shown in FIG. 3 in order to produce an indication of eating. Fundus impedance levels during time period B, however, 25 typically need to exceed the elevated threshold in order to produce an indication of eating. Typically, the fundus threshold value is further increased in a subsequent time period C. 30 to 50 minutes after time period B, if antral electrical activity is high during period C, as this is evidence of likely 30 MMC activity. For some applications, other periodic physiological activity of the gastrointestinal tract is treated in a similar manner.

In some other embodiments of the present invention, eating detection is accomplished by monitoring the rate of antral 35 electrical events. Results described hereinbelow show that the rate of antral electrical events typically decreases upon the commencement of eating. For some applications, the reduction of antral electrical events is used in addition to or instead of the techniques described hereinabove for the identification 40 of eating activities. Combining several detection criteria for the onset of eating is typically used to reduce the number of false positives and false negatives.

In an embodiment, the standard deviation of the values of event-to-event time differences are evaluated in a given sliding time window. As appropriate, the events can be detection of electrical activity or mechanical activity in the antrum. The length of the sliding time frame is typically between about 20 seconds and about 2 minutes, but, as appropriate, can be between about 20 seconds and about 10 minutes. In order to detect eating, the measured data are evaluated so as to identify a single event-to-event time difference which is more that 2-3 times the standard deviation of the former events.

With reference to FIG. 2, block 84 evaluates the analysis performed by analysis block 80 with respect to a pre-programmed or variable ingestion schedule stored in memory block 88, so as to determine whether the patient is eating in compliance with a schedule. Typically, the schedule can be modified after implantation of control unit 90, by communication from operator controls 71 using methods described 60 hereinbelow. If it is determined that the patient's eating is not in compliance with the schedule (e.g., the patient has eaten too much at one meal, or has eaten too many meals in a day, or has had too much of a certain type of food or drink), then block 84 typically actuates a signal generator block 86 to 65 generate electrical signals that are applied by current-application electrodes 100 to tissue of patient 10. Block 86 typi-

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cally comprises amplifiers, isolation units, and other standard circuitry known in the art of electrical signal generation.

The signals generated by block **86** are typically configured so as to induce a response appropriate for controlling the patient's eating habits. For example, block **86** may drive current-application electrodes **100** to apply signals to the stomach that induce gastric dysrhythmia and the resultant feeling of discomfort or nausea. Alternatively or additionally, the signals are applied to an aural site of patient **10** (e.g., in a vicinity of the cochlear nerve or the tympanic membrane), and are configured to induce vertigo, or another unpleasant balance-related sensation. Alternatively or additionally, block **86** generates a visual, audio, or other cue to encourage the patient to adhere to the schedule.

For some applications, control unit 90 drives electrodes 100 to apply a modulation signal to muscle in one area of stomach 20, so as to induce a contraction of the stimulated muscle that, in turn, induces satiety when food in an adjacent area of the stomach causes additional stretching of stretchreceptors therein. This signal may be applied in addition to or instead of the signals described hereinabove that produce gastric or other discomfort. The form of contraction-mediated stretching utilized in these applications simulates the normal appetite-reduction action of the stomach's stretch-receptors, without the patient having eaten the quantities of food which would normally be required to trigger this appetite-reduction response. In a typical application, current-application electrodes 100 are placed around the body of the stomach and are driven to induce a generally steady-state contraction of the corpus, which simulates electrically the squeezing of the corpus produced mechanically by implanted gastric bands known in the art.

Typically, the signals applied by current-application electrodes **100** include, as appropriate, an Excitable-Tissue Control (ETC) signal and/or an excitatory signal that induces contraction of muscles of the stomach. Aspects of ETC signal application are typically performed in accordance with techniques described in the above-referenced PCT Publications WO 99/03533 and WO 97/25098 and their corresponding U.S. national phase application Ser. Nos. 09/481,253 and 09/101,723, mutatis mutandis.

Typically, evaluation apparatus 18 includes remote operator controls 71, external to the patient's body. This remote unit is typically configured to enable the patient or his physician to change parameters of the ingestion schedule stored in memory block 88. For example, if the patient has lost weight, the physician may change the ingestion schedule to allow a single mid-afternoon snack. Alternatively or additionally, operator controls 71 comprise an override button, so that the patient may eat outside of the designated meal times, or consume a particular food or drink not in accordance with the schedule, if the need arises. Operator controls 71 typically communicate with control unit 90 using standard methods known in the art, such as magnetic induction or radio frequency signals.

FIG. 4 is a schematic diagram showing experimental apparatus 60 used to measure electrical responses to eating in the stomach 64 of a normal rabbit, in accordance with a typical embodiment of the present invention. Bipolar sense electrodes 62 were coupled to the fundus of stomach 64, and bipolar sense electrodes 63 were coupled to the antrum of the stomach. Additionally, two stitch electrodes 66 with a strain gauge 68 located therebetween were coupled to the antrum.

Reference is now made to FIGS. **5**, **6**A and **6**B, which are graphs showing the results of experiments performed using apparatus **60** in a rabbit, in accordance with a typical embodiment of the present invention. FIG. **5** shows electrical activity

in the fundus, measured during a five minute period before the rabbit was fed solid food, and during a more than six minute period while the rabbit was eating solid food. It can be seen that the second period is distinguished by markedly increased electrical activity. Spikes, typified by those marked "A," "B," 5 and "C" in this graph, are typically identified by a control unit operating in accordance with these embodiments of the present invention, and are interpreted as indications of eating. It is noted that in the case of the rabbit experiment shown in FIG. 5, electrical activity as measured by spikes per unit time 10 increased by a factor of about 8, and is therefore considered to be a good indication of the initiation and continuation of eating.

FIG. 6A is a graph showing the electrical response of the fundus of the rabbit stomach, and the results of analysis 15 thereof, in accordance with an embodiment of the present invention. In this experiment, the measurements were first taken for five minutes while the rabbit was eating solid food, and were continued for almost 10 minutes after the rabbit ceased eating. It is clearly seen that the period after the rabbit 20 ate is characterized by significantly less electrical activity than that which occurred during eating. Spikes, such as those marked "A," "B," and "C" in this graph, occur at a rate at least 15 times higher during eating than thereafter, and are therefore typically used by a control unit to determine both the 25 onset and the termination of eating.

FIG. 6B is an expanded view of some of the data shown in FIG. 6A, additionally showing simultaneous mechanical and electrical activity in the antrum of the rabbit. The top graph shows mechanical activity in the antrum as measured by 30 strain gauge 68 (FIG. 4), and the middle graph shows electrical activity in the antrum, measured by electrodes 63 during the same time period. The repeated co-occurrence of antral mechanical and electrical activity, as seen in FIG. 6B, is indicative of the expected antral mechanical response to 35 antral electrical activity.

The bottom graph of FIG. **6**B shows the measured electrical activity in the fundus during the same period, i.e., while the rabbit was eating. It can be seen that, while there is close correlation between mechanical and electrical activity in the 40 antrum, there is not such a close correlation between fundic electrical activity and either measure of antral activity. Control unit **90** (FIG. **2**) is therefore generally enabled to measure and differentiate between fundic and antral response, and to utilize this information to facilitate the evaluations and determinations described herein.

FIG. 7 is a graph showing electrical impedance measurements made between two stitch electrodes placed in the stomach of a pig, in accordance with an embodiment of the present invention. In this experiment, fundic volume was measured at the same time as the impedance was measured, and the data show a clear dependence of the impedance on the volume. It is hypothesized that as the fundus distends, the fundic wall thickness decreases, producing a corresponding increase in electrical impedance. Alternatively or additionally, the 55 increased distance between the two electrodes produced as a result of the distension causes the electrical impedance to increase. Similar experimental results (not shown) were obtained when impedance and volume measurements were made in the antrum. Moreover, changes in impedance were 60 found to correlate with waves of antral activity.

Reference is now made to FIGS. **8**, **9**, **10**, and **11**, which are graphs showing the results of experiments performed using apparatus (not shown) similar to apparatus **60** in several normal dogs, in accordance with a typical embodiment of the 65 present invention. All of the dogs fasted for approximately 24 hours prior to eating during the experiments.

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FIG. 8 shows the rate of electrical events in the antrum in a dog, measured during a six minute period before the dog was fed solid food and during a more than seven minute period while the dog was eating solid food. Electrical events that were recorded were spikes in the signal of amplitude at least a threshold amount greater than the signal noise. It will be appreciated that detecting changes in other events may be useful for some applications. It will also be appreciated that whereas data shown in the figures reflects measurements of antral electrical events, for some applications the analysis techniques described herein may also be implemented with respect to the rate of fundic electrical events.

It can be seen that the second period is distinguished by a markedly decreased rate of antral electrical events. Such a decrease is typically identified by a control unit operating in accordance with these embodiments of the present invention, and is interpreted as an indication of eating. It is noted that the rate of antral electrical events, as measured by events per unit time, decreased on average by about 20% beginning about one minute after the initiation of eating, and is therefore considered to be a good indication of the initiation and continuation of eating. (Decreases of up to about 50% were seen in other experiments.) Alternatively or additionally, responsive to a calibration procedure, such a decrease in the rate of antral electrical events may be used to determine other characteristics of the ingested material, for example, its nutritional, chemical, and/or caloric content. Similar results were obtained in experiments on two other dogs (not shown).

FIG. 9 is a graph showing the rate of electrical events in the antrum in a second dog, measured during a more than 40 minute period before the dog was fed solid food, during an approximately 13 minute period while the dog was eating solid food (interrupted by an approximately 6 minute period of non-eating), and during an almost 60 minute period after the dog ceased eating. It is clearly seen that the period beginning approximately four minutes after the dog ceased eating is characterized by return to a rate of antral electrical events almost equal to the rate prior to eating, and significantly higher than the reduced rate during eating. The rate of antral electrical events is therefore typically used by a control unit to determine both the onset and the termination of antral activity.

FIG. 10 is a graph showing simultaneous mechanical activity, electrical activity, and rate of electrical events in the antrum of a third dog, measured during a more than 16 minute period before the dog was fed solid food, during an approximately 3.5 minute period while the dog was eating solid food, and during a more than four minute period after the dog ceased eating. The top graph shows mechanical activity in the antrum as measured by a strain gauge, and the middle graph shows electrical activity in the antrum, measured by electrodes during the same time period. It can be seen that co-occurring mechanical and electrical activity began approximately 1.5 minutes prior to the beginning of eating, corresponding with the onset of cephalic phase activity (brain activity reflecting the mental anticipation of eating).

The bottom graph of FIG. 10 shows the rate of electrical events in the antrum of the dog. It can be seen that the second period is distinguished by a markedly decreased rate of antral electrical events, consistent with the results of the first dog experiment described hereinabove. An increase in mechanical and/or electrical antral activity prior to eating as occurred in this experiment is typically identified by a control unit operating in accordance with these embodiments of the present invention, and provides additional information that can be interpreted together with information such as the decreased rate of antral electrical events observed in this

experiment to provide indications of anticipation of eating, eating and/or gastric digestion.

FIG. 11 is a graph showing electrical impedance measurements made between two stitch electrodes in the fundus of a fourth dog, measured during five sequential periods: (1) an 5 approximately 22 minute period before the dog was fed solid food (portion of period not shown), (2) an approximately three minute period while the dog was eating solid food, (3) an approximately 7.5 minute period during which the dog did not eat, (4) an approximately one minute period while the dog was eating solid food, and (5) a greater than 10 minute period after the dog ceased eating.

It can be seen that the eating periods (second and fourth periods) are distinguished by markedly increased fundic electrical impedance. Such increases are typically identified by a control unit operating in accordance with these embodiments of the present invention, and are interpreted as indications of eating. This interpretation is supported by the correlation between impedance and volume measurements in the fundus 20 obtained in the pig experiments described hereinabove. It is noted that in the case of the dog experiment shown in FIG. 11. the fundic electrical impedance, as measured in ohms, increased by more than about 12%, beginning less than about one minute after the initiation of eating during the second 25 trols 71, described hereinabove with reference to FIGS. 1 and period, and by about 5% beginning less than about one minute after the initiation of eating during the fourth period. The fundic electrical impedance is therefore considered to be a good indication of the initiation and continuation of eating. Similar results were obtained in two other experiments on 30 different days on the same dog (not shown).

It is clearly seen in FIG. 11 that the period beginning almost immediately after the dog ceased eating (the fifth period) is characterized by a return of fundic electrical impedance to a value almost equal to that prior to eating, and significantly 35 lower than the increased value observed during eating. Fundic electrical impedance is therefore typically used by a control unit to determine both the onset and the termination of eating.

The inventors have observed that fundic electrical impedance (e.g., as measured in the case of the dog experiment 40 shown in FIG. 11), as an indicator of eating, typically exhibits lower variability than antral electrical impedance, and is less affected by movement and/or change in posture of the subject. Fundic electrical impedance also typically provides more reliable detection of eating than antral activity.

In typical embodiments of the present invention, measurements of antral and/or fundic electrical impedance are used in conjunction with or separately from other indicators of swallowing or digestion, described hereinabove, in order to track a patient's eating habits. Advantageously, impedance mea- 50 surements made between two electrodes located even at mutually remote sites on a portion of the stomach can be accurate indicators of global strain of that portion, while a mechanical strain gauge placed at a particular site on the stomach generally only yields an indication of strain at that 55

It will be recognized by persons skilled in the art that more complex combinations of variations in levels of electrical or mechanical activity in different regions of the stomach may occur than those demonstrated in the experiments described 60 hereinabove.

For example, certain electrical or mechanical activity may lag the eating of certain amounts and types of food. Examples of more complex combinations (not shown) were obtained in additional experiments in other dogs. Analysis block 80, with 65 proper calibration as described hereinabove, can readily be enabled to evaluate such complex combinations.

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Reference is now made to FIG. 12, which is a schematic illustration of a portable control charger 200, in accordance with an embodiment of the present invention. In this embodiment, control unit 90 of apparatus 18 further comprises:

- (a) a replenishable power source, e.g., a rechargeable battery, which requires periodic recharging, e.g., at least once per week;
- (b) recharging circuitry;
- (c) at least one wireless transducer, for wirelessly receiving externally-generated energy (e.g., at least one coil for inductively or electromagnetically receiving externallygenerated energy (e.g., RF energy), or at least one piezoelectric transducer for receiving externally-generated ultrasound energy); and
- (d) at least one wireless transducer for transmitting and/or receiving data (e.g., at least one coil for inductively or electromagnetically transmitting the data (e.g., as RF energy), or at least one piezoelectric transducer for transmitting the data as ultrasound energy).

For some applications, a single set of at least one transducer both receives the externally-generated energy and transmits/receives the data.

Control charger 200 typically functions as operator con-2, receiving data from and sending data to control unit 90. In addition, control charger 200 is adapted to inductively charge the rechargeable battery of control unit 90.

Control charger 200 typically comprises various input/ output elements, such as a display screen 210, a keypad or keyboard 212, and a speaker 214. Control charger 200 also comprises a rechargeable battery, which is typically recharged using standard household AC current, such as via an AC-powered recharging cradle 220, similar to conventional portable telephone recharging cradles. Control charger 200 further comprises at least one transducer (e.g., at least one coil or at least one piezoelectric transducer) for wirelessly transmitting energy to the coil of control unit 90 (e.g., inductively, electromagnetically, or via ultrasound) and at least one transducer (e.g., at least one coil or at least one piezoelectric transducer) for transmitting data to and/or receiving data from control unit 90. For some applications, a single set of at least one transducer both transmits the energy and transmits/receives the data.

In addition to its charging functionality and the functionality of operator controls 71, control charger 200 is typically programmed (in hardware and/or in software) to have functionality related to the patient's eating habits and/or weight loss program. Such functionality optionally utilizes data received from control unit 90, and/or transmits data to control unit 90. Such functionality typically includes one or more of the following:

personal digital assistant (PDA) functionality, e.g., a program for helping the patient keep a record of and/or modify his eating habits. For some applications, data received from control unit 90 are used by the program. For example, the program may record and display the number of meals or quantity of food consumed each day, and give encouraging messages when the number of meals or quantity of food is in compliance with a prescribed eating program. For some applications, the patient maintains a meal log, by keying meal details into keypad 212;

connectivity to a scale, for inputting measurements of the patient's weight. For some applications, a graph of the patient's weight is displayed next to a graph of the number of meals or quantity of food consumed during the

same time period, in order to encourage the patient to be in compliance with a prescribed eating program;

glucose monitoring, such as for diabetic patients. For some applications, monitored glucose levels are displayed in conjunction with detected meals, in order to intermittently make the patient aware of how his eating behavior affects an important parameter of health. The blood glucose monitor may be integrated in the charger or control unit 90, or external thereto;

body composition monitoring, measured by control unit **90** or by an external gauge. An example of body composition monitoring is body fat monitoring;

heart rate, ECG, and/or blood pressure monitoring, which may be integrated in the charger or control unit **90**, or external thereto:

monitoring of time or energy invested in exercise, e.g., via a home treadmill or bicycle (typically but not necessarily externally wired to the charger). Alternatively or additionally, an implanted accelerometer monitors body movement, thereby giving an indication of overall 20 physical activity;

monitoring of gastric slow wave rate (e.g., fasting and fed) and/or gastric contraction amplitude (e.g., fasting and fed). This monitoring is typically performed by control unit 90:

connectivity to a remote service provider, such as over a wireless or wired connection, for sending data to or receiving data from the service provider. This connectivity is optionally obtained via cradle **220**. In an embodiment, the remote service provider:

 a) tracks the patient's eating habits for analysis by a healthcare worker (e.g., dietitian or physician);

b) sends recommendations to control charger 200 for the patient, such as for display on display screen 210. Such recommendations may be patient-specific, such 35 as determined based on analysis of the patient's eating habits, and may include recommendations for specific dietetic foods or recipes. Such recommendations may also be commercial in nature; and/or

c) sends reminders to the patient, e.g., regarding setting 40
or going to healthcare appointments, or attending
weight-related support group meetings;

displaying system parameters such as time from last battery charge, last known battery voltage, estimated current battery voltage level; and/or

displaying average number of eating detections per day and/or a total amount of time each day that control unit **90** applies stimulation to the stomach.

For some applications, glucose monitoring, patient weight data, and/or another external feedback mechanism is used to 50 provide feedback to the patient and/or automatically change a parameter of stimulation. For example, in response to feedback indicating that the patient has attained a desired weight, or is losing weight more quickly than desired, a level of intensity of the stimulation may be lowered, so as to reduce an 55 intensity of the induced sensation or satiety or discomfort.

For some applications, glucose monitoring, patient weight data, and/or another external feedback mechanism is used to automatically change a mode of stimulation. For example, in response to an indication that blood glucose levels are at a 60 desired level, control unit 90 may switch operation of the implanted apparatus from a glucose-level-control mode of operation to an early-satiety-induction (and typically weight-loss-induction) mode of operation. Similarly, if a desired patient weight or rate of weight loss has been attained, then 65 control unit 90 may enter the glucose-level-control mode of operation, or increase the relative amount of time that it is in

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the glucose-level-control mode of operation compared to the early-satiety-induction mode of operation. As appropriate, the glucose-level-control mode of operation may be such as to: (a) induce an acute reduction of glucose level (e.g., post-prandial glucose level), and/or (b) induce a chronic reduction of glucose level (e.g., as indicated by a fasting glucose level of the patient).

For some applications, control unit 90 receives, from a PDA or other input device, an indication of a particular eating behavior of the patient, and sets an eating detection parameter responsive to the indicated eating behavior. For example, the indicated behavior may include an indication by the patient that he is on a particular eating regimen (e.g., a high protein diet, a low fat diet, a low carbohydrate diet, or a popular diet program, such as the Atkins diet). For some patients, control unit 90 is programmed to regulate a threshold of an eating detection algorithm in response to expected eating behaviors of the patient. For example, in response to a particular diet, a patient or a group of patients may demonstrate modulated intensity or timing parameters of fundic or antral activity, and control unit 90 may be programmed to automatically modify corresponding thresholds in its eating detection algorithm responsive to the effect of the particular diet on the fundic or antral activity.

Alternatively or additionally, control unit **90** receives, from a PDA or other input device, an indication of a particular eating behavior of the patient, and sets a stimulation parameter responsive to the indicated eating behavior.

For some applications, a financial incentive is provided to the patient, based on a measurement of success such as a magnitude of weight loss and/or a rate of weight loss. For example, a patient may receive a coupon to a clothing store or a health food store in response to losing a certain amount of weight in a week.

For some applications, some or all of the data types described above, and/or personal data of the patient (such as a standard medical record) are stored in the implantable device, and may be modified and/or read by an external programmer such as charger 200 or by a customized reading device. For some applications, such personal data are used to improve patient care, e.g., for a patient admitted to a hospital, or for a patient being admitted to a hospital. Alternatively or additionally, the personal data are used to facilitate giving financial incentives to the patient. For example, virtual coupons may be stored in charger 200 or in another device, and physiological data or other collected data may be used to calculate a coupon value for the patient based on the patient's achievements and/or improvements. For some applications, techniques described herein are performed using data stored in charger 200 as well as patient records or other records stored on a server of a healthcare provider.

For some applications, functionality described herein with respect to a charger is embodied in an external device that does not charge control unit 90. For example, such an external device may comprise a cash register in a store, which comprises a reader capable of reading at least some of the personal data stored in the implantable device for the purpose of delivering or receiving virtual coupons or delivering another financial incentive.

For some applications, techniques described herein with respect to treatment of obesity are applied, alternatively or additionally, in the treatment or tracking of another medical condition, such as congestive heart failure (CHF), diabetes, hypertension, snoring, sleep apnea, or a disorder related to the nervous system (e.g., epilepsy, pain, motor dysfunction, or a psychiatric disorder). For example, the techniques described herein may be adapted to provide evaluation and/or monitor-

ing of disease progression, by providing suitable data transfer between an implantable device and an external data logger (e.g., embodied in a charger like charger 200).

For example, apparatus comprising an implantable cardiac-treatment device and an external data logger (e.g., embodied in a charger like charger 200) may comprise any of the functionalities described hereinabove with respect to charger 200 and control unit 90, mutatis mutandis, and/or one or more of the following:

patient activity sensors, e.g., sensors for measuring motion, 10 walking distance, coughing, and/or sleeping of the patient. Such sensors may be used, for example, to facilitate generation of a patient-viewable output correlating gradual increases in daily walking with gradual improvement of a parameter related to heart failure; and 15 internal or external sensors and an analysis unit, for identifying changes in one or more heart-related parameters, such as heart rate variability (typically reduced when

such as heart rate variability (typically reduced when heart failure progresses), breathing sounds, ST segment elevation (e.g., as an indirect indication of cardiac 20 ischemia), heart rate elevation, aortic flow (e.g., measured using echo- or impedance-based techniques), dp/dt, or dZ/dt.

As appropriate based on the type of data collected, statistics are typically generated (e.g., hourly, daily, or weekly 25 statistics), and a summary is presented to the patient and/or transmitted to the patient's physician or to a remote data collection center.

For some applications, changes in thoracic electrical impedance are measured and presented to the patient and/or 30 transmitted to the patient's physician or to a remote data collection center. The measurements are used as an indication of changes in thoracic fluid volume, and may be used to facilitate earlier clinical assessment of changes in the patient's state (e.g., worsening CHF), and/or to initiate a 35 behavior-modification program for the patient.

In an embodiment, techniques for data logging, analysis, and transmission, and activities subsequent thereto, are applied with respect to modifying a therapy performed using an intra-body device, such as a pacemaker, ICD, ventricular 40 assist device, or electrical cardiac contractility modulation device (as are known in the art). Alternatively or additionally, techniques for data logging, analysis, and transmission, and activities subsequent thereto, are applied with respect to modifying a therapy performed without using an intra-body 45 device, such a drug therapy.

It is thus to be appreciated that techniques described hereinabove with respect to obesity may be applied, mutatis mutandis, with respect to the treatment of cardiac conditions. For example, an external data logger (e.g., incorporated in a charger) may be used to track overall activity levels of a patient via an implanted accelerometer, and to provide virtual coupons to the patient in response to activity levels that are indicative of the patient following a prescribed exercise regi-

For some applications, techniques described hereinabove with respect to obesity and/or cardiac conditions are applied, mutatis mutandis, to the treatment of diabetes. For example, tracking of changes in the condition of a diabetic patient may be facilitated by communication of data between an external 60 data logger and one or more of the following: (a) an implanted pancreatic stimulator, (b) a sensor implanted on the pancreas, (c) an implanted blood constituent sensor (e.g., a blood glucose sensor), or (d) an implanted sensor that detects changes in state of the GI tract.

In an embodiment, incentives are provided to a diabetic patient for following a prescribed health regimen. The health 42

regimen may include, for example, a diet regimen, an exercise regimen, or a behavior regimen (e.g., measuring blood glucose levels a prescribed number of times per day). Alternatively or additionally, the incentives may be provided based upon attainment of a health goal, or one in a series of interim health goals. Such health goals may include, for example, a peak daily blood glucose level that is lower than a prescribed level, an average blood glucose level that is lower than a prescribed level, or a level of weight loss that is in accordance with a prescribed weight loss plan.

It is noted that techniques described separately with respect to obesity, diabetes, hypertension, cardiovascular disease, snoring, and sleep apnea may be performed together. For example, weight data collected with respect to treating obesity may be used, additionally, in the treatment of diabetes or a cardiovascular disease of the patient.

It is noted that coupons are described herein by way of illustration and not limitation, and that the scope of the present invention includes other forms of financial and non-financial incentives, as well (e.g., tax rebates, reduced health insurance premiums, workplace rewards, eligibility for participation in health-related programs or non-health-related programs, and free entrance to theaters or other places of entertainment).

It will be appreciated by persons skilled in the art that the present invention is not limited to what has been particularly shown and described hereinabove. Rather, the scope of the present invention includes both combinations and subcombinations of the various features described hereinabove, as well as variations and modifications thereof that are not in the prior art, which would occur to persons skilled in the art upon reading the foregoing description.

The invention claimed is:

1. Apparatus, comprising:

one or more sensors, adapted to generate respective sensor signals responsively to a physiological parameter of a subject;

an implantable control unit, comprising a rechargeable battery and a first set of one or more transducers, the implantable control unit adapted to receive the sensor signals, and transmit data responsively thereto, using one or more of the transducers of the first set; and

an external control unit, comprising a power source, a second set of one or more transducers, and a display screen, the external control unit adapted to:

drive the power source to wirelessly transfer energy via one or more of the transducers of the second set,

receive the transmitted data, using one or more of the transducers of the second set,

be in connective interface with a remote service provider over a connection over which the external control unit is recharged,

send to the remote service provider eating-related information including a number of meals consumed per time period by the subject,

receive, from the remote service provider, subject-specific recommendations generated by the remote service provider based on an analysis of received eatingrelated information, and

display the subject-specific recommendations on the display screen.

wherein the implantable control unit is adapted to receive the transmitted energy, using one or more of the transducers of the first set, and charge the battery using the energy.

2. The apparatus according to claim 1, wherein the physiological parameter includes a gastrointestinal (GI) tract

physiological parameter of a GI tract of the subject, and wherein the one or more sensors are adapted to generate the respective sensor signals responsively to the GI tract physiological parameter.

- 3. The apparatus according to claim 2, wherein the GI tract 5 physiological parameter includes a parameter indicative of electrical activity of the GI tract, and wherein the sensors are adapted to generate the signals responsively to the electrical activity parameter.
- **4.** The apparatus according to claim **2**, wherein the GI tract physiological parameter includes a parameter indicative of an impedance of the GI tract, and wherein the sensors are adapted to generate the signals responsively to the impedance parameter.
- 5. The apparatus according to claim 2, comprising an implantable stimulator coupled to the implantable control unit, wherein the implantable control unit is adapted to drive the stimulator to apply stimulation to the GI tract.
- **6**. The apparatus according to claim **5**, wherein the implantable control unit is adapted to configure the stimulation to treat diabetes of the subject.
- 7. The apparatus according to claim 5, wherein the implantable control unit is adapted to configure the stimulation to treat obesity of the subject.
- 8. The apparatus according to claim 5, wherein the external control unit is adapted to monitor, using the received data, information regarding the applied stimulation, the information selected from the group consisting of: an amount of time per day that the implantable control unit drives the stimulator to apply the stimulation to the GI tract, and a number of times per day that the implantable control unit drives the stimulator to apply the stimulation to the GI tract.
- **9**. The apparatus according to claim **8**, wherein the external control unit comprises an output element, which is adapted to 35 output the information regarding the applied stimulation.
- 10. The apparatus according to claim 2, wherein the GI tract includes a stomach of the subject, and wherein the one or more sensors are adapted to generate the respective sensor signals responsively to the GI tract physiological parameter 40 of the stomach.
- 11. The apparatus according to claim 10, wherein the external control unit is adapted to monitor a gastric slow wave rate of the subject responsively to the received data.
- 12. The apparatus according to claim 2, wherein the external control unit is adapted to store information regarding eating habits of the subject.
- 13. The apparatus according to claim 12, wherein the external control unit comprises an output element, which is adapted to output the eating habit information.
- 14. The apparatus according to claim 12, wherein the external control unit is adapted to transmit the eating habit information to the implantable control unit, and wherein the implantable control unit is adapted to detect eating of the subject responsively to the sensor signals using at least one 55 eating detection parameter set responsively to the eating habit information.
- 15. The apparatus according to claim 12, wherein the external control unit is adapted to transmit the eating habit information to the implantable control unit, and wherein the 60 implantable control unit is adapted to detect eating of the subject responsively to the sensor signals using an eating detection algorithm having a threshold set responsively to the eating habit information.
- 16. The apparatus according to claim 12, wherein the external control unit comprises at least one input element, and wherein the external control unit is adapted to determine at

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least a portion of the eating habit information in response to information input by a human via the input element.

- 17. The apparatus according to claim 16, wherein the eating habit information is reflected in a meal log, and wherein the input element is adapted to accept input of the meal log by the human
- 18. The apparatus according to claim 12, wherein the external control unit is adapted to determine at least a portion of the eating habit information responsively to the received data.
- 19. The apparatus according to claim 18, wherein the eating habit information includes information selected from the group consisting of: a number of meals consumed per time period by the subject, a quantity of food consumed by the subject, a quantity of food consumed per time period by the subject, and a number of eating detections per time period, and wherein the external control unit is adapted to store the selected eating habit information.
- 20. The apparatus according to claim 12, wherein the eating habit information includes an indication that the subject follows a particular eating regimen, and wherein the external control unit is adapted to transmit the indication to the implantable control unit.
- 21. The apparatus according to claim 20, wherein the implantable control unit is adapted to detect eating of the subject responsively to the sensor signals using an eating detection algorithm having a threshold set responsively to the indication.
  - 22. The apparatus according to claim 20, comprising an implantable stimulator coupled to the implantable control unit, wherein the implantable control unit is adapted to drive the stimulator to apply stimulation to the GI tract, and set a parameter of the stimulation at least in part responsively to the indication.
  - 23. The apparatus according to claim 1, wherein the external control unit is adapted to transmit information to the implantable control unit.
  - **24**. The apparatus according to claim **1**, wherein at least one of the transducers of the first set comprises a coil, and wherein at least one of the transducers of the second set comprises a coil.
  - 25. The apparatus according to claim 1, wherein at least one of the transducers of the first set comprises a piezoelectric transducer, and wherein at least one of the transducers of the second set comprises a piezoelectric transducer.
  - 26. The apparatus according to claim 1, wherein the implantable control unit is adapted to transmit the data using the same one or more transducers of the first set that the implantable control unit uses to receive the energy.
  - 27. The apparatus according to claim 1, wherein the external control unit is adapted to transfer the energy using the same one or more transducers of the second set that the external control unit uses to receive the transmitted data.
  - 28. The apparatus according to claim 1, comprising an external recharging element, wherein the power source of the external control unit comprises a rechargeable battery adapted to be recharged using the recharging element.
  - 29. The apparatus according to claim 1, comprising an external cradle, wherein the external control unit is adapted to be removably coupled to the cradle, and to be connected to the service provider via the cradle.
  - 30. The apparatus according to claim 1, wherein the physiological parameter includes a heart-related physiological parameter of the subject, and wherein the one or more sensors are adapted to generate the respective sensor signals responsively to the heart-related physiological parameter.
  - 31. The apparatus according to claim 1, and comprising an implantable stimulator coupled to the implantable control

unit, wherein the implantable control unit is adapted to drive the stimulator to apply stimulation to a heart of the subject.

- **32**. The apparatus according to claim 1, wherein the subject-specific recommendation is for a specific food.
- 33. The apparatus according to claim 1, wherein the external control unit is adapted to receive from the remove service provider reminders generated by the remote service provider based on the eating-related information, and display the reminders on the display screen.
- **34**. The apparatus according to claim 1, further comprising  $^{10}$  the remote service provider.
- **35**. The apparatus according to claim **2**, wherein the external control unit is adapted to receive an indication of a non-GI tract physiological parameter of the subject.
- **36**. The apparatus according to claim **35**, wherein the external control unit comprises an output element, which is adapted to output the indication and eating habit information of the subject.
- **37**. The apparatus according to claim **35**, wherein the indication of the non-GI tract physiological parameter includes an indication of a body composition of the subject, and wherein the external control unit is adapted to receive the indication of the body composition.
- **38**. The apparatus according to claim **35**, wherein the indication of the non-GI tract physiological parameter is selected from the group consisting of: a heart rate of the subject, at least one parameter of an ECG of the subject, and a blood pressure of the subject, and wherein the external control unit is adapted to receive the selected indication of the non-GI 30 tract physiological parameter.
- **39**. The apparatus according to claim **35**, wherein the indication of the non-GI tract physiological parameter includes an indication of a level of exercise of the subject, and wherein the external control unit is adapted to receive the exercise 35 level indication.
- **40**. The apparatus according to claim **35**, wherein the indication of the non-GI tract physiological parameter includes an indication of a weight of the subject, and wherein the external control unit is adapted to receive the indication of the 40 weight.
- **41**. The apparatus according to claim **40**, wherein the indication of the weight includes an indication selected from the group consisting of: a weight of the subject, and a body mass index of the subject, and wherein the external control unit is 45 adapted to receive the selected indication.
- **42**. The apparatus according to claim **40**, wherein the external control unit is adapted to store eating habit information selected from the group consisting of: a number of meals consumed per time period by the subject, a quantity of food consumed per time period by the subject, and a quantity of food consumed per time period by the subject, and wherein the external control unit comprises an output element, which is adapted to output the weight and the selected eating habit information.
- 43. The apparatus according to claim 35, wherein the indication of the non-GI tract physiological parameter includes a blood glucose level of the subject, and wherein the external control unit is adapted to receive the blood glucose level.
- **44.** The apparatus according to claim **43**, wherein the external control unit is adapted to receive an indication of eating of 60 a meal by the subject, and wherein the external control unit comprises an output element, which is adapted to output the blood glucose level in conjunction with the indication of meal eating.
- **45**. The apparatus according to claim **35**, comprising an 65 implantable stimulator coupled to the implantable control unit.

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- wherein the external control unit is adapted to transmit, to the implantable control unit, information selected from the group consisting of: the indication, and information derived from an analysis of the indication, and
- wherein the implantable control unit is adapted to drive the stimulator to apply stimulation to the GI tract.
- **46**. The apparatus according to claim **45**, wherein the implantable control unit is adapted to modify a parameter of the stimulation at least in part responsively to the information.
- **47**. The apparatus according to claim **46**, wherein the stimulation parameter includes an intensity of the stimulation, and wherein the implantable control unit is adapted to modify the intensity of the stimulation at least in part responsively to the information.
- **48**. The apparatus according to claim **46**, wherein the stimulation parameter includes a timing parameter of the stimulation, and wherein the implantable control unit is adapted to modify the timing parameter of the stimulation at least in part responsively to the information.
- **49**. The apparatus according to claim **45**, wherein the implantable control unit is adapted to change a mode of the stimulation at least in part responsively to the information.
- **50**. The apparatus according to claim **49**, wherein the non-GI tract physiological parameter includes a blood glucose level of the subject, and wherein the implantable control unit is adapted to change the mode of the stimulation from a glucose-level-control mode to an early-satiety-induction mode, in response to a determination that the blood glucose level is at a desired level.
- **51**. The apparatus according to claim **49**, wherein the non-GI tract physiological parameter includes a weight of the subject, and wherein the implantable control unit is adapted to increase a relative amount of time that the mode of the stimulation is a glucose-level-control mode compared to an early-satiety-induction mode, responsively to the weight.
  - **52**. A method comprising:
  - implanting, in a body of a subject, an implantable control unit having a rechargeable battery;
  - generating, by one or more sensors, respective signals responsively to a physiological parameter of the subject, and receiving the signals at the implantable control unit;
  - transmitting, from the implantable control unit, data responsively to the signals, using one or more transducers of a first set of transducers of the implantable control unit.
  - transmitting energy from an external control unit at a site outside the body to the implantable control unit, by driving, by the external control unit, a power source of the external control unit to wirelessly transfer the energy via one or more transducers of a second set of transducers of the external control unit;
  - receiving the transmitted energy, using one or more of the transducers of the first set, and recharging the battery using the energy;
  - receiving the transmitted data by the external control unit at the site outside the body, using one or more of the transducers of the second set;
  - sending, by the external control unit, eating-related information including a number of meals consumed per time period by the subject, to a remote service provider with which the external control unit is in connective interface over a connection over which the external control unit is recharged;
  - receiving, by the external control unit from the remote service provider, subject-specific recommendations generated by the remote service provider based on an analysis of received eating-related information; and

- displaying, by the external control unit, the subject-specific recommendations on a display screen of the external control unit.
- **53**. The method according to claim **52**, wherein the subject suffers from diabetes, and wherein implanting the implantable control unit comprises implanting the implantable control unit in the subject that suffers from the diabetes.
- **54**. The method according to claim **52**, wherein the subject suffers from obesity, and wherein implanting the implantable control unit comprises implanting the implantable control <sub>10</sub> unit in the subject that suffers from the obesity.
- **55**. The method according to claim **52**, wherein the subject suffers from a heart condition, and wherein implanting the implantable control unit comprises implanting the implantable control unit in the subject that suffers from the heart 15 condition.
- **56**. The method according to claim **52**, comprising providing a financial incentive to the subject responsively to a physiological improvement in the subject determined responsively to the transmitted data.
- **57**. The method according to claim **56**, wherein providing the financial incentive comprises calculating a value of the incentive responsively to a measure of the physiological improvement.

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- **58**. The method according to claim **56**, wherein providing the financial incentive comprises awarding a coupon to the subject.
- **59**. The method according to claim **52**, wherein the physiological parameter includes a gastrointestinal (GI) tract physiological parameter of a GI tract of the subject, and wherein generating comprises generating the one or more signals responsively to the GI tract physiological parameter.
- **60**. The method according to claim **52**, wherein the physiological parameter includes a heart-related physiological parameter of the subject, and wherein generating comprises generating the one or more signals responsively to the heart-related physiological parameter.
- **61**. The method according to claim **52**, wherein the subject-specific recommendation is for a specific food.
  - 62. The method according to claim 52, further comprising: receiving, by the external control unit, from the remote service provider, reminders generated by the remote service provider based on the eating-related information, and

displaying, by the external control unit, the reminders on the display screen.

\* \* \* \* \*



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## 摘要(译)

提供了胃设备(18),包括一个或多个传感器(72,74),适于响应于胃肠(GI)生理道生成相应的传感器信号。受试者胃肠道的参数和可植入控制单元(90),包括可充电电池和第一组一个或多个换能器。可植入控制单元(90)适于使用第一组中的一个或多个换能器接收传感器信号,并且使用一个或多个换能器响应于传感器信号发送数据。第一组。外部控制单元(200),包括电源和第二组一个或多个换能器,适于驱动电源通过第二组的一个或多个换能器感应地传递能量,以便使用第二组的一个或多个换能器对电池再充电,并接收发送的数据。

