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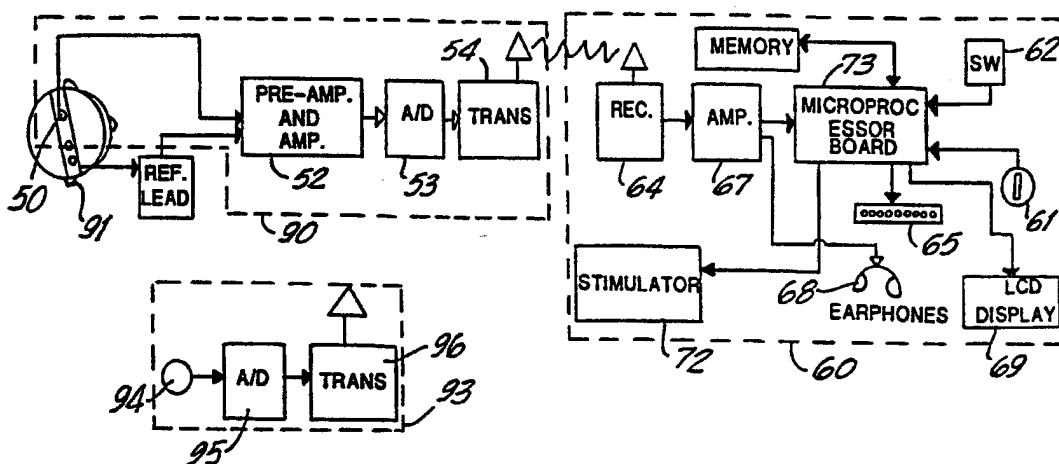
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(54) Title: BRAIN FUNCTION SCAN SYSTEM



(57) Abstract: A portable EEG (electroencephalograph) instrument, especially for use in emergencies and brain assessments in physicians' offices, detects and amplifies brain waves and converts them into digital data for analysis by comparison with data from normal groups. In one embodiment, the EEG electrodes are in a headband which broadcasts the data, by radio or cellular phone, to a local receiver for re-transmission and/or analysis. In another embodiment, the subject is stimulated in two modes, i.e., aural and sensory, at two different frequencies to provide the subject's EPs (Evoked Potentials), assessing transmission through the brainstem and thalamus.

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BRAIN FUNCTION SCAN SYSTEM

BACKGROUND OF THE INVENTION

1. Field Of The Invention

The present invention, called a "brain stethoscope", relates to a medical apparatus and more particularly to a portable EEG (electroencephalograph) device to detect, amplify and analyze brain waves generated by a human and to transmit the results to a remote receiver.

2. Related Art

It has become evident that human brain electrical activity is precisely regulated by a complex brain homeostatic system. Normative values are precisely predictable and have been found to be independent of ethnic factors. Characteristic patterns of deviation from such normative values have been reported for a wide variety of developmental, neurological and psychiatric disorders.

At the present time it is difficult for emergency personnel to determine if a subject has suffered injury to the brain or the spinal cord, cerebrovascular obstruction (stroke) or hemorrhage (bleeding). If these conditions could immediately be identified, patients' lives may be saved through rapid and appropriate treatment, usually determined after a subsequent neurological exam. The causes of abnormal behavior such as violent outbursts are often similarly ambiguous.

It is usual, during a routine medical examination, to evaluate the heart using an EKG (electrocardiogram) device. Usually, there is no attempt to determine if the patient has any brain dysfunction or conditions that may be discoverable using an EEG (electroencephalograph), as generally such devices produce an analog wavy set of waveshape tracings which must be interpreted subjectively by skilled electroencephalographers. Consequently, although the patient may be suffering from brain damage or dysfunction, such as a tumor, it is often not detected in the course of the medical examination. The absence of information about central nervous system (CNS) dysfunction often results in suboptimal treatment.

As an example, a patient arrives at a hospital emergency room (ER) with certain physical symptoms of ischemic stroke, or "brain attack", resulting from blocked blood flow to the brain. Unless the patient is treated promptly, brain cells in the ischemic region would continue to be deprived of oxygen, possibly destroying parts of his cognitive abilities, memory and motor skills and possibly resulting in death. Such adverse effects of ischemic stroke may be halted by immediate and appropriate treatment, for example, injection of tissue plasminogen actuator (tPA), which dissolves clots. However, tPA treatment of a possible stroke victim may be hazardous to initiate, as his physical symptoms may be caused by an intracerebral hemorrhage

which can be worsened by dissolving clots. Quantitative analysis of the EEG (QEEG) may provide a rapid and objective diagnosis between these alternatives.

As another example, a person may be in a coma when emergency ambulance personnel (EMS) arrive. He should not be moved if he has suffered spinal injury. But the ambulance personnel cannot determine if he has suffered spinal injury by simply looking at the comatose patient. Somatosensory evoked potentials (SEPs) provide assessment of the functional integrity of the spinal cord.

Another example of the need for an objective and immediate brain assessment is in situations where there are a number of injured persons who may require medical attention, some of whom may be in a coma. For example, on the battlefield or in the event of a train wreck, it may be necessary to separate comatose patients who are breathing and viable and require immediate treatment, from those who are still breathing but are brain dead. And again, in that situation, it is important to tell if a patient who is comatose but alive has a spinal injury, so that he should not be moved. QEEG, SEPs and brainstem auditory evoked response (BAERs) may provide a rational basis for triage in such situations.

A series of publications and patents in the name of Dr. E. Roy John relate to the field of EEG "neurometrics", which is quantitative electrophysiological measurements (QEEG) evaluated

relative to normative data. Generally, a subject's analog brain waves, at the microvolt level, are amplified, artifacts removed and the amplified brain waves converted to digital data. That data is then analyzed in a computer system to extract numerical descriptors which are compared to a set of norms (reference values), either the subject's own prior data (initial state) or a group of normal subjects of the same age (population norm). Such analyses can quantify the level, if any, of deviation of the activity of any brain region from the reference values.

A computer system based instrument using those principles is the "Spectrum 32" (Cadwell Instruments, Washington). That instrument is large, non-portable and relatively expensive (tens of thousands of dollars). It is generally used by experienced neurologists in a neurology clinic or hospital neurology department. It is not suitable for use in an ambulance, emergency room or a doctor's office for regular medical examinations. Some of the aforementioned patents which relate to neurometrics are U.S. Patents 4,279,258; 4,846,190; 4,913,160; 5,083,571 and 5,287,859, incorporated by reference.

There are a number of patents directed to determine whether a person is alive. For example, Allain U.S. Patent 5,029,590 discloses the use of a pocket-size monitor for life detection. The Allain patent deals primarily with detecting heartbeat via EKG and mentions detecting brain waves using EEG.

In John U.S. Patent 3,706,308 entitled "Life Detecting Medical Instrument" a portable device has EKG and EEG monitors, a stimulator for evoked brain responses (Evoked Potential - EP), an average response computer and a visual display. It determines if a patient is legally dead by comparison of the patient's brain waves with predetermined standards of brain death and does not use comparisons with normal values.

There is an existing need for a portable self-evaluating EEG and EP device which can be monitored by a hand-held control distant from the patient. For example, where an injured person's heartbeat cannot be detected or he is in a coma, he may be taken to a hospital, which has an EEG device and neurologist to detect and evaluate brain waves and to determine whether he is alive and whether his brain is injured. However, in some emergencies, medical personnel need to quickly determine if a patient has had a stroke or if the patient is alive but in a coma, or dead, or if a person has suffered spinal injury. A particular difficulty arises when some patients have spinal injury and are unconscious. In those cases, it would be difficult for medical personnel to ascertain who can safely be moved or should not be moved because of spinal injury. Persons with such conditions may die due to the lack of medical information, for example, a non-spinal injury patient may be in a coma and is not properly and timely transported to a hospital, or may become paralyzed if moved with unrecognized spinal injury.

In general, there are numerous instances in which the ability to make a "brain scan" by a portable EEG/EP device ("Brain Stethoscope") could be valuable in assessing the probability of abnormal brain function rapidly and automatically.

SUMMARY OF THE INVENTION

In accordance with the present invention, there is provided a portable EEG device which can accurately, reliably, continuously and quickly determine if the patient is in a coma, is suffering from concussion or is brain dead; if he is having an ischemic stroke or an intracerebral hemorrhage; if he has a serious spinal injury; and, if his behavior is of concern, whether he has abnormal brain function.

In one embodiment of the present invention, called "Version 1", intended especially for use by emergency personnel and emergency vehicles, in hospital emergency rooms and family physician offices, an EEG device has a limited number of EEG electrodes and may have an EKG electrode, preferably lead 2, and may receive input from a blood pressure device, such as a finger plethysmometer or blood oxygen or saturation meter. Depending upon the particular application, arrays of 1-16 head electrodes may be used, as compared to the International 10/20 system of 19-21 head electrodes generally used in a conventional EEG instrument. The device is small, preferably hand-held, and relatively simple, easy to use and inexpensive. It includes a

software programmed microprocessor having a CPU (Central Processor Unit) which performs the following functions: (i) it steps (polls) through the EEG electrodes if more than one electrode is used, so that each symmetrical pair of electrodes (e.g., P3 & P4) is connected simultaneously (electrode pairs are polled in sequence and the instrument is a two-channel or four-channel device) and evaluates the spontaneous EEG; (ii) it provides a timed sequence of concurrent stimulations in one or two sensory modalities (modes) to the patient, such as an audio tone or click at one repetition rate (F_1) and electrical shocks to peripheral nerves at a second repetition rate (F_2); (iii) based on the responses to these multimodal stimulations, it tests the functional state of the spinal cord (SSEP - Somatosensory Evoked Response) and brain stem (Brain Stem Auditory Evoked Response - BAER); and (iv) it assesses the cardiac rhythm.

Preferably, stimulations are used in two different modes, i.e., auditory clicks and electric pulses to the skin. The stimuli, although concurrent, are at different prime number frequencies to permit separation of different EPs and avoid interference. Such concurrent stimulations for EP permit a more rapid, and less costly, examination and provide the patient's responses more quickly, which is important in emergency situations. Power spectra of spontaneous EEG, waveshapes of Averaged Evoked Potentials, and extracted measures, such as frequency specific power ratios, can be transmitted to a remote

receiver. The latencies of successive EP peaks of the patient may be compared to those of a normal group by use of a normative template.

Preferably, to test for ischemic stroke or intracerebral or subarachnoid hemorrhage, the instrument includes a blood oxygen saturation monitor, using an infra-red or laser source, to alert the user if the patient's blood in the brain or some brain region is deoxygenated.

Another embodiment, called "Version 2", is particularly for use in field conditions in which an immediate indication of brain damage is desired from a number of persons, some of whom may be unconscious. An adhesive patch, or headband, is placed on each subject. It contains one, or more, EEG electrodes, an amplifier, and a local radio transmitter. A stimulus device may optionally be placed on each subject, such as an audio generator in the form of an ear plug, which produces a series of "click" sounds. The subject's brain waves are detected, amplified and modulate the transmitter's carrier wave. A hand-held radio receiver receives the radio waves, demodulates them and converts them into audio tones. The receiver may have an array of LED (Light Emitting Diodes) which blink depending on the power and frequency composition of the brain wave signal. Power ratios in the frequencies of audio or somatosensory stimuli are similarly encoded. With the proper training, brain wave modulated tone signals can be immediately recognized as being generated by an

intact brain or an injured brain. A physician or medical aide who is properly trained to use the Brain Stethoscope may determine, either by reading the LCD screen, listening to the audio tones, or by looking at the blinking LEDs, whether the patient's brain function is abnormal and may evaluate the functional state of various levels of the patient's nervous system.

Another embodiment, called "Version 3", uses a headband (or patch) and a hand-held receiver. The headband has 2-16 EEG electrodes, an amplifier for each electrode, an A/D (Analog/Digital) converter and a local radio transmitter. The transmitter broadcasts an FM or AM carrier which is modulated by the digital data, from the A/D converter, representing the subject's brain waves. The hand-held receiver performs the functions of analyzing the brain waves and stimulating the subject. It includes a display and a microprocessor board. The type of brain wave analysis and stimulation may be the same in Version 1.

In another embodiment, called "Version 4", the EEG device has a single electrode on a headband or an adhesive patch which is placed on a person's head to detect brain waves which are amplified and transmitted by a micro-transmitter to a microprocessor within a hand-held receiver. The microprocessor analyzes the power spectrum of the brain waves by comparison to predetermined norms, or by various ratios of power in different

frequency bands. Version 4 preferably also includes a second patch having an EKG electrode and amplifier. Either or both patches may carry an A/D converter and microtransmitter, the second patch being placed on the skin above the left collarbone.

The hand-held receiver may have LEDs, or a display panel, which displays the results of the analysis, or an audio output.

In "Version 5" a single electrode is placed preferably midway between the ears. A patch containing the electrode also has an amplifier, an A/D converter, microprocessor and a display. The microprocessor analyzes the digital data and indicates if the subject's brain waves are normal or abnormal. This Version 5 may be especially applicable in a battlefield situation.

The invention may be especially useful in cases of an emergency, for example, a wartime or peacetime explosion/disaster situation where a large number of people are injured or dead. The portable EEG device analyzes the power spectrum of brain waves to assist evaluating the degree of injury. By utilizing the portable EEG device, medical personnel may, for example, quickly divide patients into four categories: dead, seriously injured who must be moved to a hospital immediately, injured who can be moved to a hospital later, and injured, who should not be moved without special precautions.

One advantage of the portable EEG device is that it allows personnel at the scene of an emergency to determine almost instantly whether a person is alive but suffering from

concussion; is having an ischemic stroke or is suffering intracerebral bleeding; if the person is dead or in coma; whether the patient has brainstem or spinal injury; or if the patient does not have a heartbeat. The device may be utilized by medical personnel in the field, ambulance medical personnel, firemen and policemen as well as other emergency room personnel, and may be relatively simple to use and low in cost. The instrument, when used in emergency vehicles, such as firetrucks and ambulances, preferably has a built-in cellular telephone which automatically dials-up or otherwise transmits its data to a neurometric computer, for example, at a hospital. Thus, while providing immediate automatic evaluation of patient's brain state at the emergency site, it can transmit a series of brain measurements which are continuously updated to construct a "state trajectory" for remote evaluation by qualified specialists.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a block electronic schematic drawing of the system of the first embodiment;

Figure 2 is a view of the headband and hand-held instrument of the second embodiment of Figure 2;

Figure 3 is a block electronic schematic drawing of the system of the third embodiment;

Figure 4 is a chart showing the timing of two stimulations, namely, auditory clicks and somatosensory skin voltage pulses, which are simultaneous and out-of-phase;

Figure 5 is a block electronic schematic drawing of the system of the fourth embodiment; and

Figure 6 is a block electronic schematic drawing of the entire instrument, including the computer analyzer and display, in a patch or headband; and

Figures 7, 8 and 9 are top plan views of alternative displays for the instruments of Figures 1, 2, 3, 5 or 6.

DETAILED DESCRIPTION

VERSION 1

In the first embodiment, shown in Figure 1, the instrument 1 is a small and inexpensive device which is portable and may be hand-held. It uses a computer system based on a conventional microprocessor, such as an Intel Pentium I (TM) and has a limited internal memory, for example, 100 MB.

The instrument 1 has 1-24 EEG amplifiers, each of which may be connected to a removable EEG electrode shown as electrodes 10a-10p. A suitable electrode uses an adhesive cover, which is removed before applying it to the scalp. The electrode may have multiple small barbs, a needle electrode or a conductive disk, which is removably attached to and may penetrate the patient's skin; the electrode may also use conductive gel, providing rapid

attachment and acceptably low impedance, and may be sterile and disposable. In this, and other embodiments, a self-adhering electrode may be used, for example, the "ZIP-PREP" (TM) electrode having stainless steel micro-barbs in an adhesive gel patch, the patch being applied with finger pressure.

As shown in Figure 1, each of the electrodes 10a-10p is connected to a lead 11a-11p which is connected, within the instrument casing 13, to a low-noise programmable multiplexer or electronic switch 15. The programmable switch 15 may have multiple input lines and one or more output lines and is controlled by the microprocessor (CPU) board 42.

The output lines of switch 15 are connected, to a one-, two- or four-channel low noise preamplifier 12 and to a one-, two- or four-channel amplifier 14. The amplifier 14 is connected to an analog-to-digital converter (A/D) and multiplexer within General Purpose Input Board 43 (GPIB).

Preferably the electrodes are in a stretch hat, elastic flexible band or helmet. The number N of electrodes is generally greater than the number of amplification channels. However, with micro-miniaturization and reduction in cost of amplification channels, each electrode may be connected to its own amplification channel. As many as, but not limited to, 24 electrodes, or as few as one, may be used and as few as one amplification channel, or as many as N , may be used.

The computer system 40 includes a GPIB (General Purpose Input Board) 15 and a microprocessor board 42. The GPIB includes an analog-digital converter (A/D) and multiplexer 43. The computer memory 41 contains a normative or reference EEG and EP database 44. Results stored in the computer memory or other storage means can be transmitted as a digital signal by telephonic or radio transmitter 45 to a remote receiver 46, and may be encoded as audible or visual signals presented by the display 48.

A printer 49 may be used to print out a report on the patient. Preferably the printer is a color printer which is used to generate a topographic "heat scale" color-coded map of the patient's head showing, by its colors, the patient's statistical "normal" and "abnormal" regions.

The analog-to-digital multiplexer (A/D multiplexer) provides a digital output from the analog amplifiers. The A/D multiplexer samples the EEG waves (outputs of amplifiers) at a rate preferably in the range of 200 to 300 times per second (or 5 KHz for BAER + SSEP).

The data from the multiplexer is transmitted to a microprocessor board 42. The microprocessor has been programmed by an external software program means, such as a floppy disk recorder, or other input system, such as read-only memory (ROM). The programmed microprocessor ("firmware") is programmed to

perform the data acquisition and the data analysis described below. The keypad 47 is used to enter the patient's age, name and other information.

The program and its controlled microprocessor condition the input signals and insure that they are valid biological signals, including automatic artifact rejection, and periodic automatic calibration testing and impedance measurements.

The patient is stimulated by the stimulator 16 under control of the microprocessor board 42 of computer system 40. The stimulator 16 may be a lamp 17A which is flashed. In addition, or alternatively, other stimulus devices, such as headphones 17B for auditory stimulation and vibratory device 17C or low level voltage electrical skin stimulation devices may be used.

The computer system 40 automatically provides a timed set of stimuli of various modalities for the stimulator. Stimuli may be delivered separately in each modality or concurrently but at frequencies F_i which are prime numbers which share no common harmonic (see Figure 4). The patient's brain waves will respond to each stimulus with a component at frequency F_i in the EEG power spectrum, providing an "Evoked Potential" (EP). Those brain waves may be averaged, time locked to stimuli onsets to improve the signal/noise ratio, providing an "Average Evoked Response" (AER). Filtering and splitting algorithms may reduce low frequency noise and remove artifact contaminated samples prior to averaging.

The AER is the sum of N samples time locked to the onset of stimuli divided by the number of samples, to provide an updated average. N will vary depending upon the usual signal to noise ratio in each sensory modality. The AER's are held in computer memory 41.

Alternatively, narrow band FFT may be used to compare the power in the EEG at frequencies F_i when the stimulator is on vs. off, using an F-ratio.

The device described herein is capable of evaluating both the spontaneous electrical activity of the resting brain (EEG) and the processing of unimodal or multimodal sensory information after stimulation (EPs).

The switch system 15 is controlled over line 18 by microprocessor board 42. It switches the electrodes 10a-10p to the preamplifiers 12. For example, if 3 preamplifiers are used, one is used for common reference fiducial (vertex lead), such as position C_2 and the other two channels are automatically switched, by the microprocessor 42, between selected electrodes. In this way, one amplifier compresses several channels.

Each of the preamplifiers has an input isolation circuit to protect against current leakage, such as a photo-diode light-emitting diode (LED) isolation coupler. The preamplifiers are protected from electrical interference by a radio-frequency filter and a 60-cycle notch filter. Preferably each preamplifier has a computer-controlled switchable frequency range of 0.5 to

100 Hz, gain of 10,000, or of 100 Hz - 5KHz with gain of 100,000, common mode rejection of 106 dB, and noise of less than 1 microvolt.

EEG recordings may be contaminated by voltages arising from body movements, eye motion or other causes. These artifacts, it is assumed, based on prior studies, generate voltages larger than the brain wave voltage. An updating voltage threshold is computed continuously for each EEG channel, separately, by calculating the root mean squared (r m s) voltage, for example, for a sliding 20-second window and multiplying it by an appropriate constant (r m s voltage is approximately 0.2 standard deviations of amplitude). Segments containing voltages larger than this updated threshold are rejected, unless this option is turned off or the criteria altered by attending personnel. Sampling is suspended after the threshold is exceeded, for example, for one second, to avoid increasing the threshold by incorporating the artifact. It is then resumed. Preferably, those intervals (recording periods on each EEG channel) are rejected in which the voltage (signal) exceeds a multiple of the r m s voltage equal to 6 times (6X) the standard deviation of amplitude. Alternatively, an absolute maximum voltage threshold may be installed. This voltage threshold method provides segments of relatively artifact-free EEG data. The computer system, in

effect, stitches these intervals together to form a continuous artifact-free EEG sample, which is recorded in the computer memory.

Because critical decisions may depend upon the accuracy of the Brain Stethoscope evaluation, ideally odd and even "split half" samples may be constructed by assigning intervals to alternately interloaded but independent samples. For example, the samples from each electrode lead are 2.5 seconds long, the first sample is even-numbered, the second sample is odd-numbered, etc. Then the odd-numbered samples and even-numbered samples are individually compared with the norms. If a true dysfunction exists the measures revealing it must be reliably replicable. An abnormality is defined with significance of $P < 0.05$, for example, "Abnormal" is defined as deviant from normal at the 0.05 level. For example, a "split-half" consists of the first (odd) P1 and second (even) P2 samples. To be "abnormal" the same variable, at the same electrode, must be abnormal (at the 0.05 level) in both split-half samples. The probability that this could occur by chance is $P_1 \times P_2$ (0.05×0.05) or 0.0025. Results from the two split halves may be combined for display, with replicated significant results highlighted.

In addition, and optionally, the instrument may have sensors to determine the heartbeat rate, the blood pressure and the blood oxygen level of the subject. As shown in Figure 1, one or more EKG electrodes 30 are connected to an EKG amplifier 31, with a

bandwidth from 0.5 to 5000 Hz and a gain of 1000, which is connected to an input board 15. As explained below, the heartbeat QRS peaks and the R-R intervals are detected and displayed. A blood pressure (BP) sensor 32 is connected to BP meter (amplifier) 33, which is connected to input board 15. Preferably the BP sensor 32 is a finger tip blood pressure plethysmometer (plethysmograph). An oxygen analyzer 34 (O₂ ANAL), which is an optional device, is connected to input board 15. A suitable oxygen analyzer (blood oxygen saturation monitor) uses an infra-red or laser source to measure oxygen in the blood in body tissues or the brain. For example, the INVOS cerebral oximeter is available from Somanetics Corp., Troy, Michigan.

The Brain Stethoscope of Version 1 can be used in three different modes. Mode 1 is concerned with evaluation of a patient who has possibly suffered traumatic brain/spinal cord injury. Mode 2 is concerned with evaluation of a patient at risk for ischemic stroke (infarct) as against intracerebral hemorrhage. Mode 3 is concerned with the evaluation of overall brain state in patients at risk for a variety of brain dysfunctions, ranging from substance abuse to attention deficit to depression to dementia to psychosis. The same basic hardware and artifact rejection, data acquisition and analysis software in the Brain Stethoscope is used in all three modes of application,

but different lead configurations are required and different software programs are activated in the microprocessor, depending on which application is required.

Mode 1: In evaluation of traumatic brain/spinal cord injury, only three electrodes are required: one electrode on the vertex (Cz), one on the mastoid (left or right) and one on the forehead as ground. One channel of amplification will suffice, with a bandwidth of 0.5 Hz to 5 KHz, recording Cz versus the mastoid reference. The amplifier output is split into a low-pass EEG channel (0.5 Hz to 70 Hz) and a high pass (100 Hz to 1500 Hz) brainstem evoked response channel. The A/D sampling rate should be commensurate with these bandwidths. Auditory clicks, for example, at F1, (40/sec) and 90 dB should be delivered binaurally via stereophonic earphones for 30 seconds every minute. Electrical constant current shocks, for example, at F2, (27/sec), 0.5mS, 20mA, should be delivered to the fingertip of the right index finger or big toe by an electrode pair imbedded in a finger or toe cot, for the 30 seconds every minute when auditory clicks are not being delivered. Preferably a second channel is added for EKG (Lead 2), finger blood pressure plethysmometer, or pulse oximeter, and switched between these inputs to provide information about heart rate, blood pressure and oxygen saturation which is useful to assess shock.

Analysis of Mode 1 Data: The EEG recordings obtained during the evaluation period should be analyzed using very narrow band FFT (Fast Fourier Transform) in narrow, for example, 0.5Hz increments (steps) from 0.5 Hz to 50 Hz. The power in the low delta (0.5 - 1.5 Hz), delta (1.5 - 3.5 Hz), theta (3.5 - 7.5 Hz), alpha (7.5 - 12.5 Hz), beta (12.5 - 20 Hz) and high beta (25 - 50 Hz) frequency bands should be computed and expressed as absolute (μV^2) and relative power (% of power from 1.5 to 20 Hz). EEG segments in which low delta power exceeds 2 times delta power or high beta power exceeds 2 times beta (absolute power) are rejected as contaminated by eye or muscle movements. The approximate age of the patient should be entered via the keypad 47 of the device. Using age-regression equations or normal mean values stored in a table in the normative database 44 (stored in ROM of the microprocessor), the Z score of each of the four bands should be computed, where $Z = [N - P] / \sigma$ and N = normative mean value for a healthy population the same age as the patient, P = mean value computed from patient data and σ = standard deviation of the normal population values. For traumatic brain injury, positive Z-scores are expected for the delta and theta bands. These values are displayed on the screen of display 48 and transmitted to the receiving station 46 and tracked by a frequently updated (1/minute) trajectory. Increasing Z-scores for delta and/or theta suggest increasing intracranial pressure

from edema or intracranial hemorrhage and may require neurosurgical intervention. For ischemia, positive Z-scores are expected in the theta band.

Using very narrow band (VNB) FFT, the power at F1 and F2 is computed every 10 seconds, averaged separately for the 30-second alternating periods of auditory click stimulation at frequency F1 and electric shock stimulation at frequency F2. Preferably the (VNB) FFT is computed at increments (steps) in the range 0.05-0.2 Hz and most preferably at 0.1 Hz (1/10 sec. sample). Taking advantage of the fact that EEG power at a given frequency equals the variance at that frequency, the ratios of power responsive to auditory stimulation F1 (on/off) and somatosensory stimulation F2 (off/on) are calculated. Alternatively, auditory stimuli at F1 and tactile stimuli at F2 are continuous, providing a steady state response. 10-second samples of EEG are collected, FFT computed at 0.1 Hz increments and an average of N samples of the FFT is computed. The power in the F₁ and F₂ windows (Fstim) and the average power in the windows, for example, 40 bins (B) above and below each of the stimulation frequencies, Fav, is used to compute the value of

$$F_{2,B} = \frac{\text{Power Fstim}}{\text{Power Fav}}$$

sensitive. These ratios, treated as F-values, permit statistical assessment of the probability that the auditory stimuli are traversing the brainstem and the somatosensory stimuli are traversing the spinal cord and brainstem to reach the cerebral

cortex. In addition, using trigger pulses at the F1 and F2 frequencies, the microprocessor computes the averaged brainstem auditory evoked responses (BAER) and somatosensory evoked responses (SSER). The averaged BAER and SSER waveshapes, together with the full power spectrum, encoded delta, theta, alpha and beta Z-scores and the F-values for the F1 and F2 power in the EEG, are all transmitted to a remote receiving station 46 for updating a display (compressed spectral and EP waveshape arrays and feature trajectories) and for evaluation relative to normative templates (comparison with normal groups).

For analysis of EKG, QRS peak would be detected and R-R interval calculated for heart rate, displayed on an LCD, shown in Fig. 7, as BEATS/MIN.

Mode 2: The detection of ischemic stroke or intracerebral hemorrhage (bleed) is based upon the basic EEG signs of such cerebrovascular events, arising from the breakage or blockage of a blood vessel. Such events are usually on one side of the brain and give rise to asymmetry in slow brain waves. The electrodes are arranged in symmetrical (homologous) pairs, one electrode of each pair being on the left side of the head and the other electrode being on the right side of the head. In the absence of ischemic stroke, generally the slow waves from both electrodes are equal in amplitude and in phase, i.e., symmetric and synchronous. In this mode, pairs of homologous (L/R) electrodes are polled, preferably in sequence starting from the front

(frontal, temporal, central, parietal, occipital). If any electrode pair shows consistently asymmetric slow waves, especially in the theta band, it is a sign of possible cerebral ischemia. Using increments of 2.5 second artifact-free EEG segments, a sliding window 20 seconds wide is constructed and the average Z-scores of delta, theta, alpha, beta absolute power and relative power and their asymmetries are calculated across the segments within this window. Split-half replication between samples constructed from alternate odd and even segments may be used to validate results by replication, as described for Mode 1. These sequential Z-scores and their Left/Right ratios (asymmetries) are used to construct a state trajectory for each region (electrode). If the slow wave asymmetry rapidly asymptotes to an approximately constant value, or if the absolute or relative power Z-score (especially for theta or delta) in a given region reaches a stable abnormal value ($dz/dt > 0$), the cerebrovascular event is probably occlusion of a vessel (stroke); but if it continues to exacerbate or spread to adjacent regions ($dz/dt > 0$), it is probably a hemorrhage (bleed) which may require rapid intervention. Once the probability of stroke rather than bleed is established, the patient should immediately be treated, for example, with tPA. Successful thrombolytic treatment may restore normal symmetry and Z-values.

Mode 3: In order to obtain a comprehensive QEEG evaluation of brain state, it is advisable to scan the EEG in 8 pairs of homologous electrode placements (FP1/FP2, F3/F4, C3/C4, P3/P4, O1/O2, F7/F8, T3/T4, T5/T6). Each pair of electrodes provides a short period sample, for example, 1-10 seconds and preferably segments of about 2.5 seconds (artifact free). A fiducial electrode, preferably the electrode at position C_z is recorded continuously to confirm stationarity during the full scan. Preferably, linked earlobes (A1+A2) are taken as the reference. The electrodes are scanned in pairs a number of times, for example, 8-48 times and preferably at least 24 times, and ideally 48 times, for a total of 120 seconds/pair. In a two-channel 16-electrode system (8 pairs) at 2.5 seconds per segment per pair, each scan takes 20 seconds. The scans may be performed by an electronic switch, as shown in Figure 1. The ideal 48 scans take 960 seconds, or 16 minutes, and the minimum 8 scans required for high reproducibility would take 160 seconds. In a four-channel system, the corresponding scan times would be 480 seconds and 80 seconds. The trade-off between cost (number of channels) and scan time may involve 2, 4, 8 and 16 channel versions of the Brain Stethoscope for use in situations where rapidity of Mode 3 applications may be important. One additional EEG channel should be dedicated to continuous C_z recording.

For each of the 2.5 second segments during the total scan, the values of the EEG spectral parameters are computed for Cz, which is recorded in every scan, and the mean \bar{M} and standard deviations $\bar{\sigma}$ computed across the set of segments in the scan. Any segment for which a parameter of Cz exceeds $\bar{M} + 2.5 \bar{\sigma}$ should be excluded from the scan. Replication between odd and even split half samples may be used to further establish validity of abnormal findings, as described for Mode 1. Taking advantage of the demonstrated stationarity of the resting EEG, the digital data averaged across the full scan is then compared with norms (population norms) to determine if the patient's brain function is normal or abnormal. A composite 16 channel scan is constructed from the multiple pairwise scans. The averaged spectral parameters for each lead and samples of raw data may be transmitted, by cellular phone or other transmission means, to a receiving PC terminal in a hospital neurology department or neurological center having a suitable computer for its analysis, such as the "Spectrum 32" (Cadwell Instruments) or any computer with neurometric capability. Interpolated statistical probability maps, color coded for significance using a "heat" scale, may be constructed at the remote receiver. Alternatively, the instrument 1 performs its own basic analysis on pairwise data and constructs the composite total scan and may also construct a

topographic map, and preferably provides a simple result, i.e., "normal" or "abnormal", which may be followed by such transmission.

The criteria for "normal" and "abnormal" functions may be a "look-up" table. For example, if the power at any electrode is significantly ($Z \gg 2.0$) below the norm, for the age group of the patient, then the patient is considered "abnormal" and an indication (colored light, digital read-out, buzzer, etc.) will be generated. More specific diagnostic classifications may require transmission of selected clinical observations to a remote center for specialized evaluations.

Further evaluation by the instrument's computer memory may have a set of discriminant functions. Such functions may be empirically derived and installed in the instrument. In one embodiment, the discriminant functions are held as a set in memory as a band or base number for each diagnostic category, see U.S. Patent 5,083,571 at columns 4 and 5. For example, in the diagnostic category of ischemic stroke, the groups may be: normal (no stroke), cerebrovascular compromise (stroke) and intracerebral hemorrhage (bleed).

VERSION 2

The embodiment of Figures 2 and 3 is especially adapted for emergency personnel, such as firemen, military medical corpsmen, etc. It has only one self-adhering electrode 50, or a few such

electrodes in a headband 51 or cap. The electrode 50 is connected to a tiny amplifier and radio-transmitter 54. The amplifier includes a pre-amplifier and high-gain amplifier 52 whose output after A/D conversion 53 modulates the AM transmission of the radio transmitter 54. The entire circuit unit (amplifier and transmitter) is preferably less than 0.25 inches thick and the size of a half-dollar.

The transmitter 54 may be small as it transmits only within a short range, for example, 50-300 feet. As shown in Figure 3, a receiver unit is within a casing 60 which may be handheld and includes a radio receiver 64 tuned to the transmission frequency of the transmitter 54. The unit 60 also includes an amplifier 67 to amplify the demodulated signal and to drive a speaker or earphone (headset) 68. A switch 62 is used to turn the audio on and off and a dial 61 is used to select the band of brain wave frequencies separated by appropriate filters. Position 61a of the dial 61 is the entire spectrum (T-Total) and positions 61b-61e are respectively the delta (1.5 - 3.5 Hz), theta (3.5-7.5 Hz), alpha (7.5 - 12.5 Hz) and beta (12.5 - 20 Hz) bands. Each of these, including the entire spectrum, generates a distinctive warble or series of tones in earphones 68. A person who has been trained can tell, by the intensity of the sequence of such tones, if the subject's brain waves sound normal or abnormal. Split half replication as in Version 1 may be used and significant abnormalities identified by a beep or other appropriate signal.

VERSION 3

As shown in Figure 3, a rapid diagnosis may be obtained by using a hand-held QEEG device.

In Version 3 a headband 90 is applied to a subject. The headband 90 includes a self-attaching electrode set 50, preferably 3-16 electrodes, a pre-amplifier for each electrode, and an amplifier 52 for each pre-amplifier, an A/D converter 53, a micro-transmitter 54 (local battery operated radio transmitters). For example, in a war time battlefield situation the patches 90 may be applied to several (3-6) wounded soldiers within 50-100 feet of a blast site. An ear plug 91 is then inserted into the ear of each subject. The ear plug 91 may be a radio receiver activated by stimulator 72 or, alternatively, it self-generates a sound (click sound) at a selected frequency (F1), for example, at 40 clicks per second.

A patch 93 containing a self-attaching EKG electrode 94, A/D converter 95 and micro-transmitter 96 is placed on the subject's skin above the left collarbone. The amplified and digitized EKG heart waves are transmitted by the local radio transmitter 54 to the hand-held device 60.

The patient is preferably stimulated by audio clicks and vibratory or electrical voltage pulses to the skin to provide EPs by the stimulator 72 which is controlled by the microprocessor board 73. One of the electrodes is connected to the vertex as a

reference (C_2 in 10/20 system). Preferably, to save time, the stimulations are given concurrently, at different frequencies and phase, as shown in Figure 4. For example, a conductive bracelet or ring is adhered to the subject's skin and brief low voltage electrical pulses are transmitted to it. These provide a tingling sensation (somatosensory mode) at frequency F_1 . An earphone plug 91, which is a radio receiver, is inserted into the subject's ear and click sounds (auditory mode) are transmitted to the earphone at a different frequency F_2 , where F_1 and F_2 are different prime numbers.

The brain waves at the vertex, in a normal brain, will reflect both the F_1 and F_2 frequencies. The receiver unit 60, as in the first embodiment (Version 1) produces a Fast Fourier Transform (FFT) of the digital data, based on multiple 2.5 second segments. Separate FFTs are generated for the F_1 and F_2 frequencies, and their harmonics.

An "F ratio" is derived at each of the 4 frequency bands (alpha, beta, delta, theta). The F ratio is the ratio of power, P_{stim} , (mean squared) under stimulation, derived from a narrow band FFT, to a norm (the subject's power, P_{ref} , at that frequency prior to stimulation, e.g., a self-norm). That F ratio may be tracked to determine if there are any changes in the subject's condition. Alternatively, the F ratio may be based on a comparison of the subject's brain waves to a normal group (population norm).

When time and the situation permit, split half replication may be optionally used to validate estimates of abnormality.

An F ratio may be determined in the different ways, as explained above in connection with Version 1. The F ratio is determined for both the F1 and F2 stimulation modes at each of the frequency bands. A trajectory is constructed using a sequence of measurements. If the subject is stable, those F ratios would stay the same and the first derivative of the trajectory would approach zero. The power generated at F1 and F2 includes their harmonics (preferably five harmonics).

An LCD display 69 (Figs. 7 or 9) provides a visual indication of the patient's brain state in each frequency band, relative to normal values, by statistical assessment. The cortical response to auditory stimuli (at F1) and somatosensory stimuli (at F2) is similarly encoded in audio or visual output.

In addition, the receiver unit 60 has a row display 65 which shows amplitude, preferably a row of LCDs. If the brain wave amplitude (total or at each frequency band) is non-existent or low, the LCDs will indicate this appropriately. The frequency band to be displayed is selected by dial 61 or it cycles automatically, or all may be displayed as an array.

The FFT provides information in the frequency domain. The subject's brain waves are compared to a set of normal power spectra held in computer memory and generated using the same stimulus conditions. If the subject's power spectra differ

significantly compared to the range of the predetermined norms, then an "abnormal" display is generated and displayed on alphanumeric display 69.

VERSION 4

In this embodiment, shown in Figure 5, a single EEG electrode 105 is applied to the subject. The electrode is in an adhesive patch 100, about 2 cm in diameter, or in a headband. The patch 100 (or headband) also contains an amplifier 101 (pre-amplifier and amplifier), a battery, an A/D converter 102 and a micro-transmitter 103 (local radio transmitter). The transmitter 103 locally broadcasts the subject's brain waves as digital data, the data having modulated an FM or AM carrier at a selected frequency, or as modulations of the carrier by the analog EEG.

For example, the patches 100 may be applied to several (3-6) subjects within 50-100 of an emergency site. An ear plug 106 is inserted into the ear of each subject. The ear plug self-generates a sound (click sound) at a selected frequency (F1), for example, at 40 clicks per second, and may be a piezo-electric audio generator.

An adhesive patch 93, containing a self-attaching EKG electrode 94, A/D converter 95 and microtransmitter 96, is placed on the subject's skin above the left collarbone.

The hand-held receiver analyzer 110 includes a radio receiver 111, an amplifier 112, a microprocessor board 113, a dial 114, and a display 115, which function the same as the corresponding components in the embodiment of Figure 3.

The microtransmitters 103 and 96 transmit short-range signals which provide the global condition of the subject's cerebral cortex and the brainstem (as evaluated by narrow band fast Fourier transform of the spontaneous EEG and the EEG activated by stimulation from the audio generator plug 106) and the waveshape of the EKG.

Using stored normative data from memory 117, based on age-matched normal controls and a baseline period (about 1 minute) of measurement of the initial state of the subject, the hand-held receiver 110 (Brain Stethoscope) will compute visual or auditory statistical evaluations of the global state of the subject's cortex and brainstem, based upon the narrow band FFT, and the R-R interval of the EKG. These quantitative evaluations (Z-scores relative to population norms) will be presented in visual LED display 115 or as audible signals on earphones 116, separately for each subject. The averaged Brainstem Auditory Evoked Response can be computed and the Peak I-IV measured to assess brainstem integrity. An updating trajectory display 115 will present the evolution of each measure, the slope (first derivative) of which will be automatically evaluated to assess whether the state is stable, improving or deteriorating. When

time and circumstances permit, split-half replication may optionally be used to validate critical findings. Priorities, as to the care of the subjects, can then be assigned in an informed way.

On the trip to the hospital, in an ambulance, the history and updating trajectory, augmented by clinical assessments, would continue to be transmitted by transmitter 121 from the hand-held receiver 110. Samples of EEG Spectra/BAER/EKG waveshapes could be interrogated by the neurometric computer at the hospital (base station).

Assessment of spinal cord injury can be obtained by using the Brain Stem Somatosensory Evoked Potential (BSEP). Acquisition of BSEPs by the Brain Stethoscope can be accomplished by placing a rubber "cot" on the big toe of each foot and stimulating (via a stimulator 120), a pair of electrodes in the toe "cots" at a rate of about 17/sec (F2) - left toe; and 19/sec for (F3) - right toe. The spectral power at a midline forehead electrode, on the subject, at frequencies F2 and F3 will reflect the arrival of information at the cortex via the left and right medical lemniscal pathways. Cortical components and components of the BSEPs, at increasing latencies from 12 ms to 55 ms after each stimulus, will diminish and disappear depending upon the extent of spinal trauma. Finger tip electrodes (median nerve) can similarly be used to assess the level of injury from the 4th

cervical vertebra (C4) and above. This will also estimate the extent of spinal cord injury and the compression caused by intrathecal hemorrhage.

The Brain Stethoscope constructs an "initial state norm" of all extracted EEG/Ep/EKG features during its initial set of measures, and this reference norm includes the variance of each measure. The state trajectory which it constructs utilizes the Z-transform (i.e., $Z = [\text{Present state minus initial state}] / \text{variance of initial state}$), or the F-ratio (present power / initial power). An automatic alarm can be set (auditory or visual) to alert attendants if any change in the state trajectory exceeds 2.56 standard deviation (P 0.01) toward further deterioration. This will provide monitoring of the subject during his transport to the hospital.

VERSION 5

In Version 5, shown in Figure 6, the entire Brain Stethoscope is in a patch 150 or headband. Preferably, as in the prior embodiment of Figure 5, an adhesive patch 150 has a single electrode 151 which preferably is placed midway between the ears, or less preferably on the forehead.

The patch 150 contains an electrode 151, an amplifier 152 (pre-amplifier and amplifier), battery, A/D converter 153, a microprocessor 154 having a computer memory 155, and a display 156. The entire patch is preferably less than 3 cm in diameter.

The microprocessor 154 performs the QEEG analysis, described above, for the receiver of Version 4, and preferably produces a simple normal/abnormal result. Split-half replication may optionally be used to validate critical results, time permitting. That result is shown in the display 156 which is on the front face of the patch 150. The display 156 may be a green LED 157 ("normal") and red LED 158 ("abnormal").

In each of the above-described embodiments a reference lead (reference electrode) "REF/LEAD" should be used in addition to the active EEG electrode(s). Preferably the reference electrode is removably attached at the earlobe or mastoid (temporal bone behind ear). For example, a small 1-3 cm diameter adhesive patch containing a reference electrode may be adhered at the earlobe or mastoid. A wire runs from the reference electrode to the amplifier (pre-amplifier) which may be in a patch with the active electrode.

WHAT IS CLAIMED IS:

1. A medical system to warn of spinal cord, brainstem or brain injury or brain dysfunction of a subject comprising:
 - (a) at least one active EEG (electroencephalograph) electrode means to detect a subject's brain waves;
 - (b) stimulus means to provide concurrent sense stimuli in two or more different stimulus modes (sensory modalities) to the subject, the stimuli in one mode being at frequency F_1 and the stimuli in the other modes(s) being at a different frequency $F_2, F_3, \text{ etc.}$;
 - (c) amplification means to amplify and digitize the detected brain waves;
 - (d) ratio means to receive the digitized brain waves from (c) and produce subject brain wave F ratio data based on the power of brain responses at each of F_1 and F_2 in the presence and absence of stimulation;
 - (e) a receiver including a computer means to compare the subject brain wave F ratio data from (d) with brain wave F ratio data based upon a normal group of patients or in the presence or absence of stimulation; and
 - (f) a warning means in the hand-held receiver to warn if the subject's brain wave F ratio data is abnormal based on the comparison of (e).

2. A medical system as in claim 1 and including FFT means to perform Fast Fourier Transform (FFT) of the digitized brain waves to generate a power spectrum.
3. A medical system as in claim 2 wherein the F ratio data is an F ratio based on the generated power spectrum and the power of brain responses at each of F_1 and F_2 in the presence and absence of stimulation.
4. A medical system as in claim 1 wherein the F ratio data is an F ratio between the power at F_1 or F_2 : $P(F_i) / P(AV)$ where average power $P(AV)$ is across N bins above and below F_1 or F_2 .
5. A medical system as in claim 1 and including means to validate the digitized brain waves by comparing odd and even split-half segments of data from the same electrode, under the same stimulus and about the same time and rejecting segments that are atypical, using a fiducial electrode to assess stationarity.
6. A medical system as in claim 2 wherein the FFT is a very narrow band FFT in increments in the range of 0.05 - 0.5 Hz.
7. A medical system as in claim 1 and, in the receiver, report means to generate a report on the comparison of (e) and transmission means to transmit the report to a base station.

8. A medical system as in claim 1 and means to modulate a carrier wave to modulate the amplified brain waves and to generate an audio signal therefrom.

9. A medical system as in claim 1 wherein the stimulus means includes means to generate an audio click and a somatosensory stimulus produced by applying a constant current or voltage or tactile vibration to sensitive regions of the subject.

10. A medical system as in claim 1 wherein the statistical evaluation of computed measures from the patient (P) is by computing the Z-score, where $Z = (M-P)/\sigma$ and M is the mean value of a normative distribution, P is the current measure from the patient and σ is the standard deviation of the normal age-matched population.

11. A medical system as in claim 1 and including a headband carrying thereon the electrode means and amplification means.

12. A medical system as in claim 1 including a plurality of homologous pairs of electrodes and means to determine asymmetry above a threshold level of the subject's brain waves at the homologous pairs of electrodes.

13. A medical system as in claim 1 and including a headband having thereon the electrode means, amplifier means and a radio broadcast transmitter.

14. A medical system to warn of brain injury comprising:
- (a) at least one active EEG (electroencephalograph) electrode means to detect a subject's analog brain waves, connection means to removably connect the electrode means to a subject's head;
 - (b) amplification means on the connection means to amplify the detected brain waves;
 - (c) radio broadcast means on the connection means to generate a brain wave broadcast signal by modulating a carrier signal with the detected analog brain waves and to broadcast the brain wave signal broadcast;
 - (d) a hand-held radio receiver means to receive and amplify the broadcast brain wave signal; and
 - (e) sound generating means in the hand-held receiver to demodulate the amplified broadcast brain wave signals and to convert demodulated brain waves into tone-like sounds which may be recognized by an operator as reflecting brain damage.
15. A medical system as in claim 14 wherein the connection means is a headband.
16. A medical system as in claim 14 wherein the radio receiver means includes appropriate filter means to select a frequency band from the group of frequency bands of the broadcast brain waves.

17. A medical system as in claim 14 wherein the group of frequency bands comprises delta, theta, alpha and beta bands.

18. A medical system to warn of brain injury to a subject's head, the device comprising:

- (a) an attachment means to removably attach an electrode to the subject's head, at least one EEG (electroencephalograph) electrode means carried on the attachment means to detect a human subject's brain waves;
- (b) amplification means in functional connection with the electrode means to amplify the detected brain waves and analog/digital convertor means to digitize the detected brain waves and to product the subject's digitized brain wave data therefrom;
- (c) a brain wave computer analyzer means on the attachment means to compare the subject's brain wave data with a brain wave data based upon a normal group of subjects stored in the analyzer means; and
- (d) a warning means on the attachment means to warn if the subject's brain wave data is abnormal based on the comparison of (c).

19. A medical system as in claim 18 and including means to validate the digitized brain waves by comparing split-half segments of data from the same electrode and under the same stimulus and about the same time and rejecting segments that do not match.

20. A medical system as in claim 18 wherein the warning means is a plurality of lights or an alphanumeric display panel.

21. A medical system as in claim 18 wherein the attachment means is a patch and the electrode means is a single electrode.

22. A medical system to warn of brain injury to a subject's head, the device comprising:

- (a) an attachment means to removably attach an electrode to the subject's head, at least one EEG (electroencephalograph) electrode means carried on the attachment means to detect a human subject's brain waves;
- (b) amplification means in functional connection with the electrode means to amplify the detected brain waves and analog/digital convertor means to digitize the detected brain waves and to produce the subject's digitized brain wave data therefrom;
- (c) radio transmitter means on the attachment means to broadcast the subject's brain wave data after amplification and modulation;
- (d) a remote radio receiver and demodulator to receive and demodulate the brain wave data;
- (e) a brain wave computer analyzer means to compare the subject's brain wave data with a brain wave data base based upon a normal group of subjects stored in the analyzer means; and

(f) a warning means on the attachment means on the receiver to warn if the subject's brain wave data is abnormal based on the comparison of (e).

23. A medical system as in claim 22 wherein the warning means is a plurality of lights, an audio device or an alphanumeric display panel.

24. A medical system as in claim 22 wherein the attachment means is a patch and the electrode means is a single active electrode and a reference and ground.

25. A medical system as in claim 22 wherein the transmitter means is a radio transmitter or a cellular telephone.

26. A medical system as in claim 22 wherein the transmitted brain wave data includes data from a series of sequential observations.

27. A medical system as in claim 26 wherein the series of sequential observations is divided into an odd and an even split-half and replication of findings between the split halves is used to validate results.

28. A medical system as in claim 22 wherein the series of quantitative measures extracted from the data are used to construct a state trajectory.

29. A medical system as in claim 22 and further including, in the receiver, means to produce an audio output from a broadcast carrier which is modulated by the subject's brain waves.
30. A medical system as in claim 22 and further including an EKG (electrocardiograph) amplifier in the receiver means, an EKG electrode adapted to be removably connected to the subject and connected to the EKG amplifier, and display means in the receiver to display heart beats and the digitized heart rate of the subject.
31. A medical system as in claim 22 and further including a blood pressure monitor in the receiver means, a blood pressure cuff adapted to be removably connected to the subject, and display means in the receiver to display the blood pressure of the subject.
32. A medical system as in claim 22 and further including a blood oxygen meter in the receiving means, an oxygen probe adapted to be removably connected to the subject, and display means in the receiver to display the blood oxygen level of the subject.
33. A medical system as in claim 22 and further including in the receiver, report means to generate a digital data report based on the comparisons of (e), and broadcast means on the receiver to broadcast the report.

34. A medical system as in claim 33 and wherein the receiver is a hand-held local radio receiver, the attachment means includes a radio transmitter to the local receiver, and the receiver includes transmission means to transmit a brain wave data report to a remote receiver.

35. A medical system as in claim 22 wherein the attachment means includes a headband.

36. A medical system as in claim 22 and including at least three electrode means and three amplification means as well as reference and ground electrodes.

37. A medical method to warn of spinal cord, brainstem or brain injury or brain dysfunction of a subject comprising the steps of:

- (a) attaching to a subject at least one EEG (electroencephalograph) electrode means to detect the subject's brain waves;
- (b) providing concurrent sense stimuli in two or more different stimulus modes to the subject, the stimuli in one mode being at frequency F_1 and the stimuli in the other mode(s) being at a different frequency $F_2, F_3, \text{ etc.}$;
- (c) amplifying and digitizing the detected brain waves;
- (d) receiving the digitized brain waves from (c) and producing subject brain wave F ratio data based on the power of brain

responses at each of F_1 and F_2 in the presence and absence of stimulation;

- (e) comparing the subject brain wave F ratio data from (d) with brain wave F ratio data based upon a normal group of patients or in the presence or absence of stimulation of the patient in a receiver including a computer means; and
- (f) generating a warning signal in the hand-held receiver to warn if the subject's brain wave ratio data is abnormal based on the comparison of (e).

38. A medical method as in claim 37 and including performing Fast Fourier Transform (FFT) of the digitized brain waves and generating a power spectrum therefrom.

39. A medical method as in claim 38 wherein the F ratio data is an F ratio based on the generated power spectrum and the power of brain responses at each of F_1 and F_2 in the presence and absence of stimulation.

40. A medical method as in claim 37 wherein the F ratio data is an F ratio between the power at F_1 or F_2 : $P(F_i) / P(AV)$ where average power $P(AV)$ is across N bins above and below F_1 or F_2 .

41. A medical method as in claim 37 and validating the digitized brain waves by comparing odd and even split-half segments of data from the same electrode, under the same

stimulus and about the same time and rejecting segments that do not satisfy criteria for stationarity.

42. A medical method as in claim 38 wherein the FFT is a very narrow band FFT in increments in the range of 0.05 – 0.5 Hz.

43. A medical method as in claim 1 and generating a report, in the hand-held receiver, on the comparison of (e) and transmitting the report to a base station.

44. A medical method to warn of brain injury comprising the steps of:

- (a) removably connecting at least one active EEG (electroencephalograph) electrode to the head of a subject using connection means and detecting the subject's analog brain waves;
- (b) amplifying the detected brain waves using amplification means on the connection means;
- (c) generating a brain wave broadcast signal, using radio broadcast means on the connection means and modulating a carrier signal with the detected analog brain waves and broadcasting the brain wave signal;
- (d) receiving and amplifying the broadcast brain wave signal using a hand-held radio receiver; and
- (e) generating tone-like sounds using the hand-held receiver by demodulating the amplified broadcast brain wave signals and converting the demodulated brain waves into tone-like

sounds which may be recognized by an operator as signifying brain damage.

45. A medical system to transmit data regarding brain injury or dysfunction comprising:

- (a) a headband and at least two EEG (electroencephalograph) electrode means carried therein to detect a human subject's brain waves, the electrode means being located as homologous pairs with one electrode means of each pair on the right side of the brain;
- (b) amplification means to amplify and digitize the detected brain waves and to produce brain wave data therefrom;
- (c) switching means to switch the amplification means, in sequence, to different homologous (L/R) pairs of the electrode means in order to construct a scan;
- (d) means to confirm stationarity of a subject's brain state at a selected fiducial electrode across the set of N scans required to assess the brain state of the subject;
- (e) means to extract measures selected from the group of absolute power, relative power, coherence (synchrony) and symmetry for each sample of EEG from a pair of electrodes and to average each of selected measures across the full set of scans;
- (f) computer means to compare subject's brain wave data of (e)

with brain wave data based upon a normal group of subjects and to produce an almost real-time report based thereon regarding an excess or deficit of energy selected from the group of beta, alpha, theta or delta bands and to assess across sequential scans an increasing shift in local slow wave reflecting injury or dysfunction or, in each pair, asymmetric slow waves indicating cerebral ischemia.

46. A medical system as in claim 45 and print means to print a statistical probability map with interpolated values, using a color-coded heat scale with lines from red to yellow indicating significance of excesses and blue to white indicating deficits relative to the appropriate form.

47. A medical system as in claim 45 and including means to validate the digitized brain waves by comparing odd and even split-half segments of data from the same electrode and under the same stimulus and at about the same time and rejecting segments that do not satisfy criteria of stationarity.

48. A medical system as in claim 45 and display means to construct and display a color-coded topographic map displaying the statistical probability of the values of (f).

49. A medical system as in claim 45 and including computer means to combine the averaged data from all pairs of electrodes into one "virtual simultaneous" recording.

50. A medical system as in claim 45, and stimulus means to apply concurrent stimulation to the subject in different modes, e.g., auditory clicks and electric pulses to the skin, the concurrent stimuli being at two or more different frequencies.

51. A medical system as in claim 46 and including means to validate the digitized brain waves by comparing odd and even split-half segments of data from the same electrode and under the same stimulus and about the same time and rejecting segments that do not satisfy a criterion of stationarity.

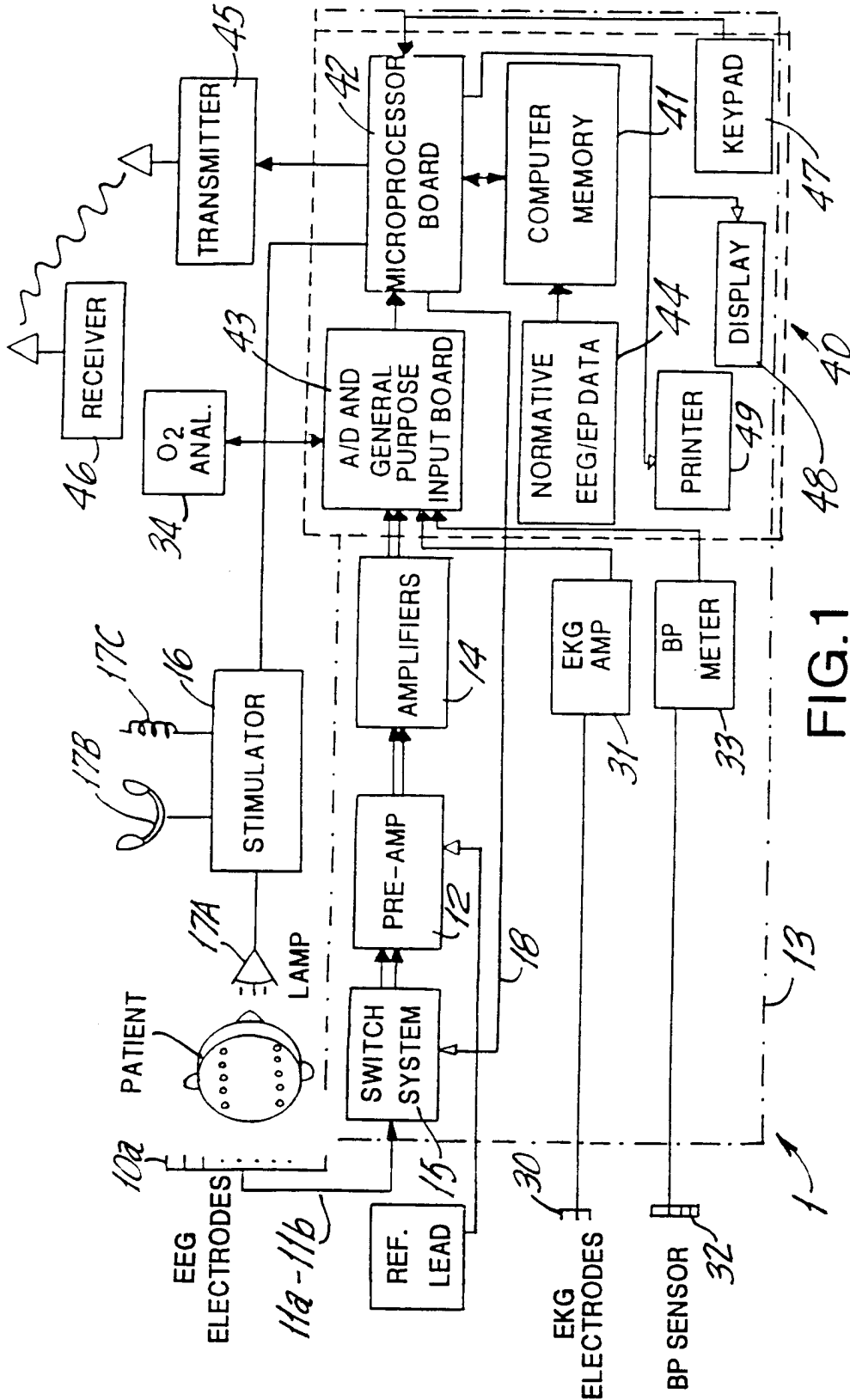


FIG. 1

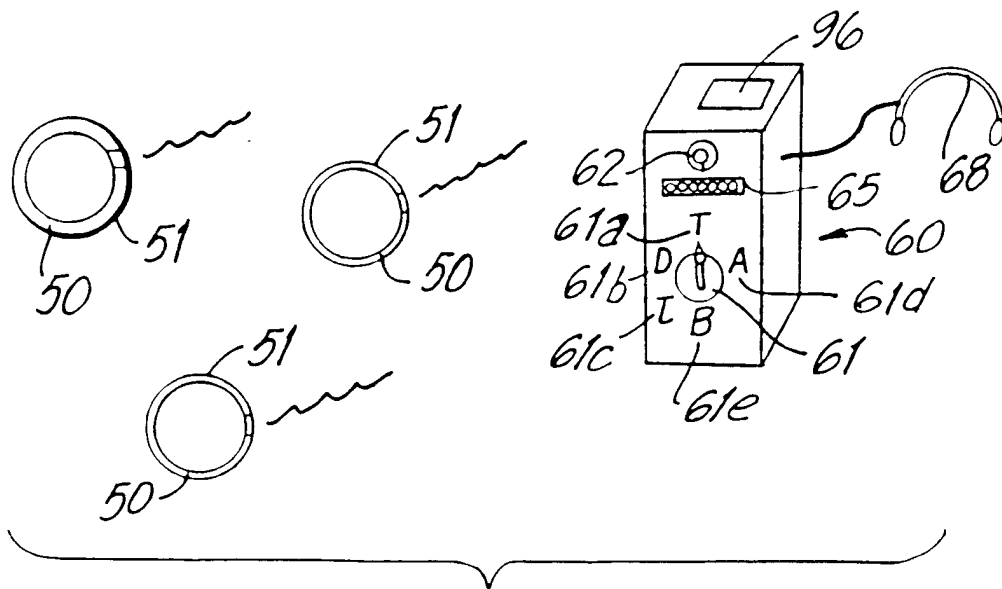


FIG.2

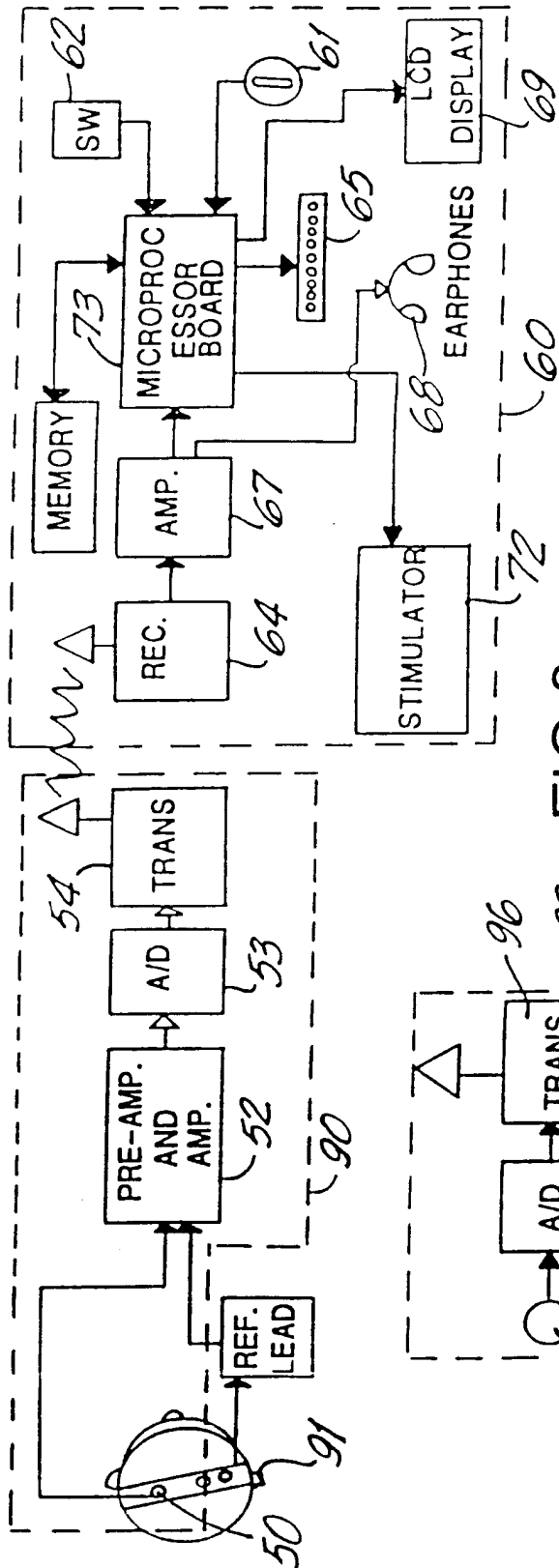


FIG. 3

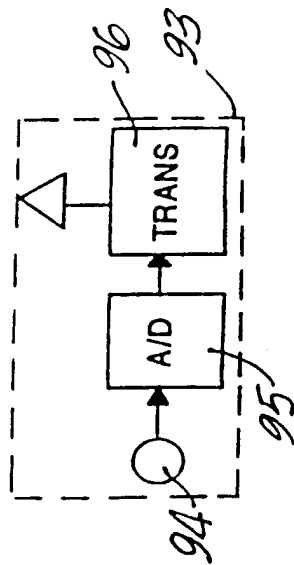
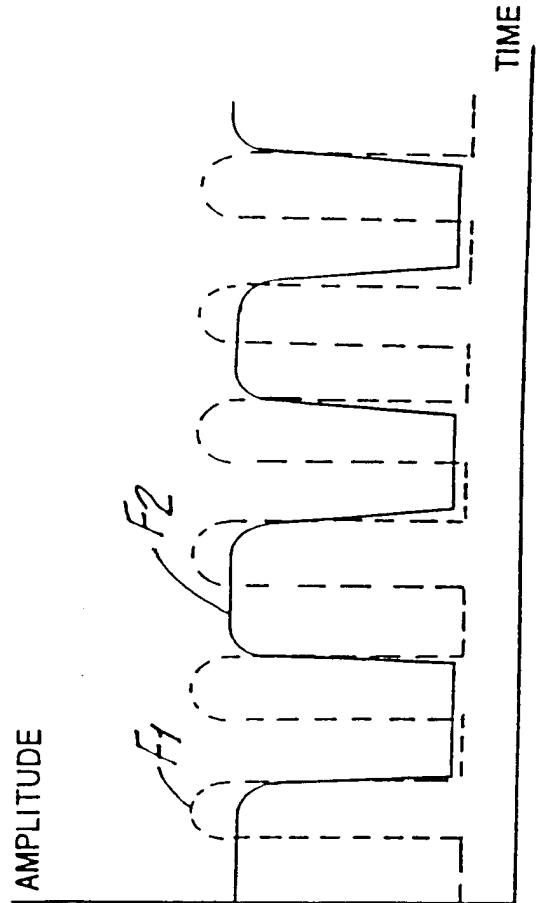


FIG. 4



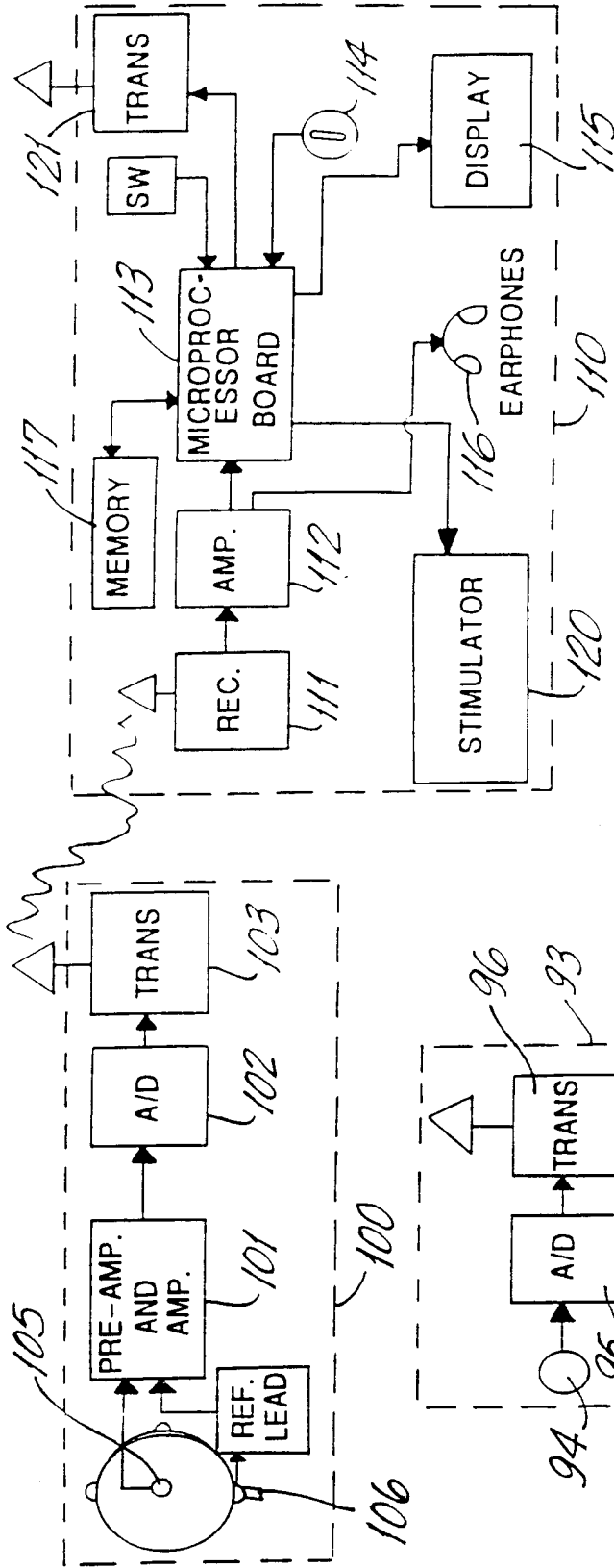


FIG. 5

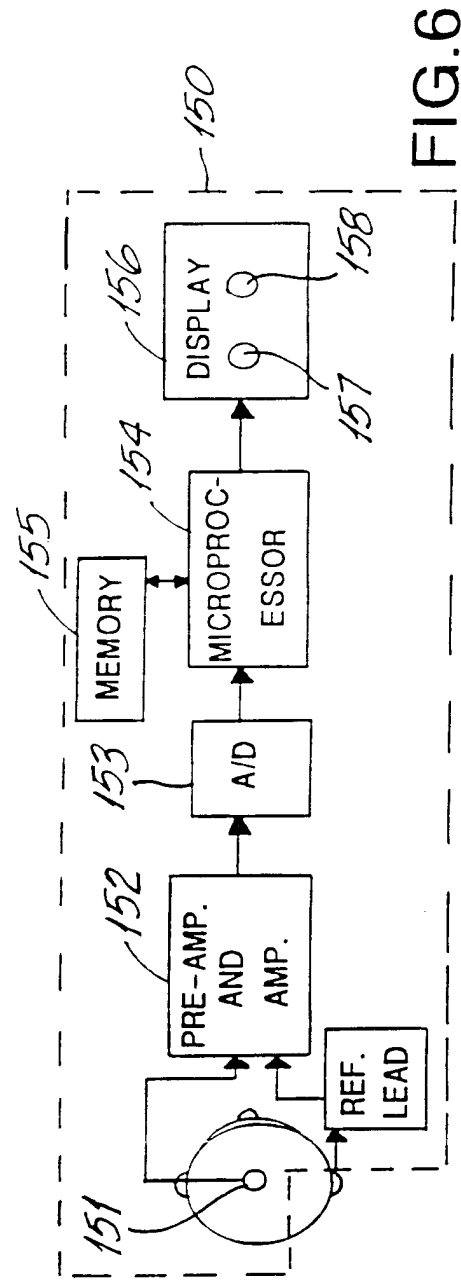


FIG. 6

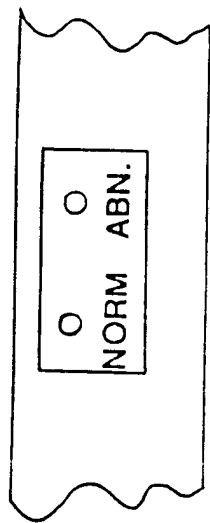


FIG. 7

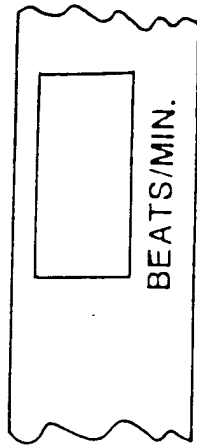


FIG. 8

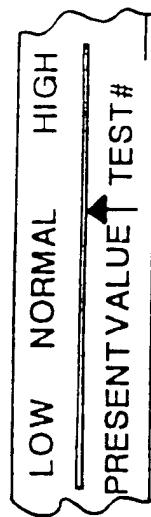


FIG. 9

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US00/10055

A. CLASSIFICATION OF SUBJECT MATTER

IPC(7) :A61B 05/00
US CL :600/544

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 600/544, 545

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

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

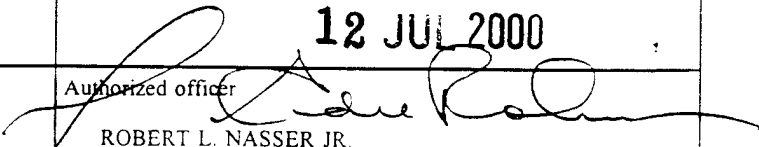
C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	US 5,325,862 A (LEWIS et al.) 05 July 1994, entire document.	1-51
A	US 5,752,922 A (ROSENFELD) 19 May 1998, entire document.	1-51
A	US 5,46,205 A (VIRSU et al.) 05 May 1998, entire document.	1-51

Further documents are listed in the continuation of Box C. See patent family annex.

* Special categories of cited documents	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
"A" document defining the general state of the art which is not considered to be of particular relevance	"N" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
"E" earlier document published on or after the international filing date	"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	"&" document member of the same patent family
"O" document referring to an oral disclosure, use, exhibition or other means	
"P" document published prior to the international filing date but later than the priority date claimed	

Date of the actual completion of the international search 05 JUNE 2000	Date of mailing of the international search report 12 JUL 2000
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Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231 Facsimile No. (703) 305-3230	Authorized officer  ROBERT L. NASSER JR. Telephone No. (703) 308-3251
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专利名称(译)	脑功能扫描系统		
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当前申请(专利权)人(译)	纽约大学		
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发明人	JOHN, ERWIN ROY JOHN, MICHAEL SASHA		
IPC分类号	A61B5/00 A61B5/0476 A61B5/0484		
CPC分类号	A61B5/0476 A61B5/0002 A61B5/0006 A61B5/0484 A61B5/407 A61B5/4076 A61B5/7257		
优先权	09/447369 1999-11-23 US		
其他公开文献	EP1272098B1 EP1272098A4		
外部链接	Espacenet		

摘要(译)

便携式EEG (脑电图) 仪器, 尤其用于医生办公室的紧急情况和大脑评估, 通过与正常组的数据进行比较, 检测和放大脑电波并将其转换为数字数据进行分析。在一个实施例中, EEG电极位于头带中, 该头带通过无线电或蜂窝电话将数据广播到本地接收器以进行重新传输和/或分析。在另一个实施方案中, 以两种不同的频率刺激受试者, 即听觉和感觉, 以提供受试者的EP (诱发电位), 评估通过脑干和丘脑的传播。