Title: PAIN DETECTION SYSTEM AND METHOD UTILIZING NEAR-INFRARED SPECTROSCOPY

Abstract: Disclosed is a pain detection system and method. Preferred methods include capturing imaging data from patient using functional near-infrared spectroscopy. In methods of the invention, pain may be assessed from a non-responsive patient. Preferred methods may further include measuring hemodynamic parameters using the optical imaging data and averaging the measured hemodynamic parameters to reduce background noise. Additionally, a pattern in the averaged hemodynamic parameters is identified and the pattern is compared to a predetermined associated with patient pain. The method further includes detecting, by the processor, a similarity between the pattern and the predetermined pattern and outputting a signal indicating patient pain.
PAIN DETECTION SYSTEM AND METHOD UTILIZING NEAR-INFRARED SPECTROSCOPY

Cross-Reference to Related Applications


The entire teachings of this application are incorporated herein by reference.

Technical Field

In one aspect, systems and methods are provided that detect patient pain during a medical procedure by monitoring the patient throughout the procedure using functional near-infrared spectroscopy.

Background

Many types of medical procedures involve the use of anesthetics, analgesics, and other sedation or partial sedation methods. For example, sedatives are administered before a colonoscopy procedure to reduce anxiety in a patient and reduce discomfort during the procedure. During a colonoscopy procedure many patients report having experienced pain. The pain experienced during the procedure has been observed by practitioners through patient movement or grimacing and may be due to residual effects of the sedative and/or analgesic agent and gender differences. Thus, research implies that nociceptive pathways may be activated during the procedure due to stimuli causing local or central neuronal sensitization and potentially leading to postoperative pain. Routine colonoscopy procedures may be used to evaluate pain signals during a procedure when a patient is under sedation because patient behaviors such as grimacing, moaning, or movement may be an indicator of experienced pain.

Various approaches to minimizing pain during medical procedures have been developed and include the use of carbon dioxide for insufflation, improving anesthetic regimens, providing technical aids, and patient controlled sedation or analgesia. However, precise dose administration of anesthetics may be difficult since dose is determined by a patient's age, medical history, current medications, previous use of anesthetics, current condition (e.g., unstable or critical), and the like. In addition, the
time of administering the anesthetics and commencing the medical procedure may vary
thereby potentially causing insufficient time for the drugs to take effect. As an example,
functional magnetic resonance imaging (fMRI) may capture brain activity associated
with pain. However, fMRI is often difficult to use due to equipment unavailability,
relatively large physical sizes, and relatively high operating costs among other
reasons.

Summary

We now provide new pain recognition systems and methods utilizing a
functional near-infrared spectroscopy system. Preferred systems and methods are
configured to detect pain experienced during a medical procedure while a patient is
sedated or under other forms of anesthesia using near-infrared spectroscopy (NIRS)
system.

In particular systems and methods, a functional near-infrared spectroscopy
(NIRS or fNIRS) system is utilized as the imaging system.

Preferred methods of the invention include recognizing pain in a non-
responsive patient, comprising obtaining functional near-infrared spectroscopy
imaging data from a non-responsive patient to recognize pain in the patient.

In particular systems and methods, the patient is non-responsive, particularly
nonresponsive to commands, such as in a controlled state of anesthesia or an
unconscious sedated state, or is otherwise non-responsive to commands, such as may be
exhibited by a comatose patient, an infant, and the like.

In additional particular aspects, systems and methods are provided that detect
pain based on pain indicia or control data obtained from a subject. This indicia or control
data may be e.g. suitably obtained prior to a patient or subject becoming unresponsive,
such as through sedation. For example, pain can be detected by a responsive patient or
subject through any number of exercises. The patient then may be rendered non-
responsive (e.g. through sedation), and the control data or pain indicia is then
employed to monitor and recognize pain in the unresponsive patient.

In preferred aspects, the NIRS system can probe or detect changes in brain
hemodynamic responses using infrared light wherein changes in the relative
concentration of oxygenated and deoxygenated hemoglobin in the cortex indicates pain experienced by a subject, such as a patient undergoing a medical procedure.

Preferred methods include obtaining or capturing imaging data from subject, e.g. a non-responsive patient, over a period of time (e.g. up to about 0.1, 0.5, 1, 2, 3, 5 or 10 minutes, or more) using near-infrared spectroscopy. Particularly preferred methods include obtaining or capturing optical imaging data from at least one cortical region of the brain of a subject, e.g. a non-responsive patient, over a period of time (e.g. up to about 0.1, 0.5, 1, 2, 3, 5 or 10 minutes, or more) using functional near-infrared spectroscopy. In addition, using the imaging data, preferred methods may include measuring a plurality of hemodynamic parameters, which may include oxygenated hemoglobin, deoxygenated hemoglobin, and/or total hemoglobin, in the at least one cortical region of the brain of the non-responsive patient and preferably averaging the measured plurality of hemodynamic parameters to reduce background noise. A pattern may also be identified in the averaged hemodynamic parameters which may be compared to at least one predetermined pattern associated with patient pain. When a similarity is detected between the pattern and the predetermined pattern, a signal may be output indicating patient pain. In addition, measuring the hemodynamic parameters may include measuring oxygenated and deoxygenated hemoglobin to measure blood flow.

Preferred methods may include performing an analysis exercise such as a finger tapping exercise and a variation of electrical stimulus on a non-responsive patient (e.g. a sedated patient) to inflict a level of pain. In addition, the predetermined pattern that indicated pain may comprise a plurality of peaks in the blood flow over time, wherein the method may include determining whether a time delay between performing the various activities (e.g., finger tapping exercise and the variation of electrical stimulus) and the point at which a peak occurs indicates patient pain. The predetermined pattern associated with pain may be obtained from control data for the hemodynamic parameters, wherein the control data is obtained by acquiring the optical imaging data from the at least one cortical region of the brain of a conscious patient. The optical imaging data may be obtained when the patient is not subject anesthetics and may be acquired from a finger tapping exercise and a variation of electrical stimulus. The method may further include administering an anesthetic regimen based on a particular medical procedure toward the patient after obtaining the control data.
Additionally, preferred systems may include a functional near-infrared spectroscopy configured to capture at least one cortical region of the brain of the patient over a period of time and a processor. The cortical region of the brain may be the somatosensory cortex and the prefrontal cortex. The processor may be configured to measure a plurality of hemodynamic parameters, which may include oxygenated hemoglobin, deoxygenated hemoglobin, and/or total hemoglobin, in the cortical region of the brain of the patient, using the optical imaging data.

Furthermore, the processor may be configured to average the measured hemodynamic parameters to reduce background noise and then may identify a pattern in the averaged hemodynamic parameters. The processor may then compare the pattern to at least one predetermined pattern associated with patient pain and detect a similarity between the pattern and the predetermined pattern. Additionally, the processor may be configured to output a signal indicating patient pain. The processor may be configured e.g. to control an analysis exercise such as a finger tapping exercise and a variation of electrical stimulus on the patient to inflict a level of pain toward the patient from which measurements may be obtained. Further, the processor may be configured to determine whether a time delay between performing the analysis exercise such as a finger tapping exercise and the variations of electrical stimulus and a point when a peak occurs in blood flow indicates pain in the patient.

Methods are also provided that evaluate a treatment and include administering, by a processor, the treatment toward a patient and applying a stimulus toward the patient. Such methods also may include acquiring, with a functional near-infrared spectroscopy, imaging data during the application of the stimulus. Furthermore, from the imaging data, brain activity data representing a hemodynamic parameter in at least one cortical region of the brain of the patient may be determined. The brain activity data may then be compared, by the processor, to control data which may be acquired using the functional near-infrared spectroscopy prior to the imaging data and further outputting an indication of patient pain. A reduction in intensity of the hemodynamic parameter from the control data to the imaging data may indicate a relative efficiency of the treatment as a pain-relieving treatment.

Other aspects of the invention are disclosed infra.
Brief Description of the Drawings

The disclosure will hereinafter be described with reference to the accompanying drawings, wherein like reference numerals denote like elements, and:

FIG. 1 illustrates an exemplary graph showing hemodynamic response functions observed at various cortices according to an exemplary embodiment of the present disclosure;

FIG. 2 illustrates an exemplary graph of hemodynamic response functions according to an exemplary embodiment of the present disclosure;

FIG. 3 illustrates exemplary graphs of hemodynamic response functions observed at right medial prefrontal cortices and somatosensory cortices according to an exemplary embodiment of the present disclosure;

FIG. 4 illustrates exemplary pain pathways associated with the somatosensory cortex and the prefrontal cortex according to an exemplary embodiment of the present disclosure;

FIG. 5 illustrates an exemplary flowchart of the method that recognizes pain in a patient according to an exemplary embodiment of the present disclosure; and

FIG. 6 illustrates an exemplary schematic diagram of the system that recognizes pain in a patient according to an exemplary embodiment of the present disclosure.

Detailed Description

Pain recognition systems and methods are provided that are suitably used to measure pain experienced in a medical procedure, particularly while a patient is unresponsive to commands or communication, such as resulting from sedation. Pain is measured through near-infrared spectroscopy imaging of the patient.

Methods and systems of the invention are particularly useful for a wide variety of medical procedures, including where a patient is sedated to an unconscious or unresponsive state. In the unresponsive state, pain can be assessed and e.g. an anesthetic regimen can be administered based on that pain assessment through the course of the medical procedure. The present pain detection systems and methods can provide
continuous monitoring of neural activity related to pain experienced during medical procedures. The present systems and methods also may provide control of analgesic load to improve patient intraoperative comfort and decrease postoperative pain. In addition, the measure or detection of pain may be used to evaluate the effectiveness of pharmaceuticals for a variety of conditions, including e.g. irritable bowel syndrome.

Preferably functional near-infrared spectroscopy (fNIRS) is employed for imaging of the patient. Functional near-infrared spectroscopy can allow functional imaging of brain activity (or activation) through monitoring of blood oxygenation and blood volume, including in patient’s cortex. Imaging can be performed in the near-infrared range such as between 700-1000 nm.

In particular methods and systems, the patient is in a non-responsive state, such as an unconscious sedated state as may be controlled by administration of anesthesia, and/or may be characterized by partial or complete loss of protective nerve reflexes, including the ability to independently breathe and respond to commands. The patient may be unable to cooperate and may have labile (fluctuating) vital signs. As referred to herein, an unresponsive patient or subject includes a subject exhibiting one or more characteristics of: 1) under the administration of anesthesia and non-responsive to commands, 2) partial or complete loss of protective nerve reflexes, such as loss of ability to i) respond to commands and/or ii) independently breathe. An unresponsive patient also may include patients in a coma or infants (who cannot respond to verbal commands).

Methods and systems of the invention also may be utilized for subject in a conscious state, including a conscious sedated state. As referred to herein, a patient in a conscious state, a conscious sedated state or other similar term is 1) capable of rational responses and/or 2) does not have altered respiratory, cardiac, and/or reflex functions (nerve reflexes from the brain) to the level that requires external support for these vital functions.

In a preferred method for recognizing pain in a non-responsive patient, steps may include obtaining functional near-infrared spectroscopy imaging data from at least one cortical region of the brain of the non-responsive patient; and comparing measured imaging data with patient pain based on control data and thereby recognizing pain in the non-responsive patient. Using the imaging data, one or more hemodynamic parameters may be measured in the at least one cortical region of the brain of the non-responsive patient. Such preferred methods may suitably comprise one or more additional steps of
(i) averaging the measured plurality of hemodynamic parameters e.g. to reduce
background noise; (ii) identifying a pattern in the averaged hemodynamic parameters;
and/or (iii) outputting a signal indicating patient pain.

In a particular preferred method of the invention for recognizing pain in a non-
responsive patient, steps may comprise obtaining functional near-infrared spectroscopy
imaging data from at least one cortical region of the brain of the non-responsive patient;
measuring, by a processor, using the imaging data, a plurality of hemodynamic
parameters in the at least one cortical region of the brain of the non-responsive patient;
averaging, by the processor, the measured plurality of hemodynamic parameters e.g. to
reduce background noise; identifying, by the processor, a pattern in the averaged
hemodynamic parameters; comparing, by the processor, the pattern to at least one
predetermined pattern associated with patient pain based on control data; detecting, by
the processor, a similarity between the pattern and the predetermined pattern; and
outputting, by the processor, a signal indicating patient pain.

In such preferred methods, the hemodynamic parameters include one or more
of oxygenated hemoglobin, deoxygenated hemoglobin, and total hemoglobin.
Measuring at least one hemodynamic parameter may include measuring, by the
processor, oxygenated and deoxygenated hemoglobin to measure blood flow. The at
least one cortical region of the brain may be selected from the somatosensory cortex
and the prefrontal cortex. A predetermined pattern indicating pain may comprise a
plurality of sequential peaks in blood flow over time. The at least one predetermined
pattern associated with pain may be obtained from the control data for the
hemodynamic parameter wherein pain is detected in a conscious patient.

In another particular method for evaluating a medical treatment, steps may
comprise administering a treatment to a patient; applying a stimulus to the patient;
acquiring functional near-infrared spectroscopy imaging data during the application of
the stimulus; determining from the imaging data, brain activity data representing a
hemodynamic parameter in at least one cortical region of the brain of the patient;
comparing the brain activity data representing the hemodynamic parameter to control
data representing the hemodynamic parameter; and outputting an indication of patient
pain. One or more of such steps may be accomplished through use of a processor.

Referring now to the non-limiting exemplary drawings, various pain and
sensory pathways may be measured in the brain using the NIRS system as shown in FIG.
4. The NIRS uses infrared light to probe changes in brain hemodynamic response,
wherein near infrared light penetrates in the tissues and may traverse matter within the skull to the cortex area. A change in the relative concentration of the oxygenated hemoglobin and the deoxygenated hemoglobin in the cortex indicates a change in the number of photons that are escaping the brain.

Specifically, as illustrated in FIG. 6, a functional near-infrared spectroscopy system (NIRS) 625 is a portable, non-invasive inexpensive method of monitoring hemodynamic activity at moderate depths (e.g., surface cortices). The NIRS 625 is configured to characterize relative changes in concentrations of oxygenated hemoglobin (HbO) and deoxygenated hemoglobin (HbR), which combined indicate relate change in total hemoglobin (HbT). The NIRS optodes may be fitted across predetermined cortical regions to thus monitor the hemodynamic functions caused by neuronal activity.

Furthermore, since an NIRS 625 is not influenced by environmental requirements, the system may be used within an operating room. Specifically, the NIRS 625 is configured to deliver photons using fiber optics that are positioned on the head 620 of a patient and are called the source 615. In addition, other fiber cables, detectors 610, are spaced apart from the source 615 to collect the photons that are reflected from the cortical tissue. Furthermore, various detected signals may be measured to detect the intensity, the modulation depth, and the phase delay and a combination thereof may provide information regarding physiological parameters.

Furthermore, as illustrated in FIG. 6, the pain recognition system 600 may be configured to capture optical imaging data from at least one cortical region of the brain over a period of time, using the NIRS system 625. In addition, the system 600 may include a processor 605 that may be configured to measure a plurality of hemodynamic parameters in the at least one cortical region of the brain of the non-responsive patient, using the optical imaging data. The processor 605 may then be configured to average the measure plurality of hemodynamic parameters to reduce background noise in the measurement process. A pattern may then be identified in the averaged hemodynamic parameters. The processor may be configured to compare the pattern to at least one predetermined pattern associated with patient pain and a signal may be output indicating patient pain. Specifically, the predetermined pattern may be associated with control data obtained prior to a stimulus being administered to a patient. The control data may be obtained by fitting the NIRS 625 system to the patient and collecting data at point where the patient indicates experiencing pain, while the patient is conscious (e.g., is not sedated). Thus, when the patient is sedated, the control data may act as an
indication of points where pain is known to exist or be experienced by the patient. Moreover, the hemodynamic parameters may include oxygenated hemoglobin, deoxygenated hemoglobin, and total hemoglobin. In addition, the at least one cortical region of the brain may be selected from the somatosensory cortex and the prefrontal cortex. The processor 605 may be further configured to control a finger tapping exercise and a variation of electrical stimulus on the patient to inflict a level of pain from which measurements may be obtained. Furthermore, the processor 605 may be configured to determine whether a time delay between performing the finger tapping exercise and the variations of electrical stimulus and a point when a peak occurs in blood flow indicates pain in the patient.

Moreover, FIG. 5 illustrates an exemplary flowchart of process 500 of recognizing pain in a patient during a medical procedure. The process 500 may illustratively start at 505 and continue to 510 wherein optical imaging data may be captured from at least one cortical region of the brain of the non-responsive patient over a period of time, using a functional near-infrared spectroscopy. Furthermore, the process includes in 515 measuring, by a processor, a plurality of hemodynamic parameters in the at least one cortical region of the brain of the non-responsive patient, using the optical imaging data. In 520, the process may include averaging, by the processor, the measured plurality of hemodynamic parameters to reduce background noise and in 525, identifying, by the processor, a pattern in the averaged hemodynamic parameters. In addition, in 530, the process may include comparing, by the processor, the pattern to at least one predetermined pattern associated with patient pain. The process may then include in 535, detecting, by the processor, a similarity between the pattern and the predetermined pattern. Lastly, in 540, outputting, by the processor, a signal indicating patient pain, and then the processor may illustratively end in 545.

Furthermore, the hemodynamic parameters may include oxygenated hemoglobin, deoxygenated hemoglobin, and total hemoglobin. The measuring of the hemodynamic parameter may include measuring, by the processor, oxygenated and deoxygenated hemoglobin to measure blood flow. The at least one cortical region of the brain may be selected from the somatosensory cortex and the prefrontal cortex.

Additionally, the process may further include determining, by the processor, whether a time delay between performing the finger tapping exercise and the variation of electrical stimulus and a point when a peak occurs in blood flow indicates pain in the patient. Furthermore, the at least one predetermined pattern associated with pain may be
obtained from control data for the hemodynamic parameter, wherein the control data may be obtained by acquiring the optical imaging data from the at least one cortical region of the brain of a conscious patient. Specifically, the control data may be obtained when a stimulus has not been administered to a patient and includes performing, by the processor, an activity to the patient. The activity performed on the patient may include a finger tapping exercise and a variation of electrical stimulus to inflict pain. In addition, the process may include, administering, by the processor, an anesthetic regimen based on a particular medical procedure toward the patient after obtaining the control data.

In a further embodiment of the present disclosure, a method that evaluates a treatment may include administering, by a processor, the treatment toward a patient and applying a stimulus toward the patient. Furthermore, the process may include acquiring imaging data during the application of the stimulus using a functional near-infrared spectroscopy. In addition, the process may include determining, by the processor, brain activity data representing hemodynamic parameter in at least one cortical region of the brain of the patient, from the imaging data and comparing, by the processor, the brain activity data representing the hemodynamic parameter to control data representing the hemodynamic parameter. The process may further include outputting, by the processor, an indication of patient pain. Additionally, the process may include acquiring, by the processor, control data using the functional near-infrared spectroscopy prior to obtaining the activity imaging data. A determined reduction in the intensity of the hemodynamic parameter from the control data to the imaging data indicates a relative efficiency of the treatment as a pain-relieving treatment.

Methods and systems of the invention may be employed to detect pain and the following non-limiting example is illustrative of the invention.

Example

In this study patients were monitored using NIRS during a routine colonoscopy procedure to analyze brain activity during various procedural events. Specifically, pain was induced by stretching the colon during insufflation and characterizing the signal to determine whether the patient experiences pain.

Methods of Example

Subjects

Seventeen healthy adults were recruited from a population of patients
undergoing screening colonoscopy. Of the seventeen adults, eleven were male and six were female ranging from ages 36-79. Three data sets were eliminated due to no marked stimulus event and two additional data sets were eliminated due to poor signal quality from patient motion. Thus, twelve data sets were used for analysis wherein the subjects included six males.

**Equipment**

The subjects were fitted with a TechEn CW6 NIRS imaging system (TechEn, Milford MA) and data was recorded using the NIRS imaging system during the duration of the medical procedure. The NIRS imaging system was configured to emit near infrared wavelengths of light to measure changes in cortical deoxyhemoglobin and oxyhemoglobin concentration. Auxiliary channels in the data acquisition system were utilized to mark particular activities such as patient grimacing, patient motion, injection of anesthetic regimens, at the like.

**Paradigm**

Finger tapping was delivered manually to each patient's hand which was marked for stimulus delivery. The finger tapping was delivered in substantially the same location using substantially the same pressure each time. Intraoperative recordings were collected and markers were placed in the data set, using an auxiliary data acquisition channel, for the duration of each cardiac ablation event. In particular, three variations of electrical stimulus were applied to the patient.

The patients were all provided with similar anesthetic regimen including midazolam and Demerol. However, the recall of the colonoscopy by some patients is an indication of variation in response to the anesthetic regimen based on difference in the interactions of clinical evaluation, pharmacogemonic, or gender differences.

In addition, nociceptive signals may not be effectively blocked by the anesthetic regimen. Effective blockade of nociceptive signaling is dependent on the intensity, duration, and repetition of afferent signals. Although a patient may not express pain during a medical procedure, nociceptive pathways may still be activated as a result of repeated stimuli which may cause local or central sensitization to occur, thus resulting in pain after the medical procedure.
Data Analysis

Analysis was performed using Homer2 software and the analysis has been previously published Gagnon L, Perdue K, Greve DN, et al. Improved recovery of the hemodynamic response in diffuse optical imaging using short optode separations and state-space modeling. Neuroimage 2011;56:1362-71.

Results

Relative brain activity was measured across the prefrontal and somatosensory cortices as shown in FIG. 2. Specifically, FIG. 2 illustrates that the medial somatosensory cortex experienced the greatest change in oxygenation and hemoglobin. The bi-phasic response, as shown in FIG. 1, shows HbO and HbT peaks occurring about every 4 seconds, observed in two somatosensory channels, thereby indicating patient pain.

In the prefrontal cortex, the bilateral decrease in HbO and HbT prior to time zero, and corresponding rise to a peak value about 3 seconds later indicates patient pain, as shown in FIGS. 1 and 3. In particular, FIG. 3 shows the decrease in oxygenated hemoglobin, at approximately 2 seconds prior to the grimace response, indicating deactivation of medical prefrontal cortex neurons following the initiation of the pain stimulus, based on the approximately 6 second time-to-negative peak observed in studies where pain was applied with known timing.

Discussion

The group average hemodynamic response function observed in the prefrontal and somatosensory cortices is similar to the results obtained in previous near-infrared studies that monitor the brain during controlled pain stimulation. The time offset may be the result of the different methods of marking the stimulus onset. Furthermore, significant delay may be caused by the reaction time of a sedated patient corresponding to a negative-time shift in the hemodynamic response. The increase in signal intensity may indicate increased pain intensity.

The results in the above example show a signal of alterations in brain neuronal activity that corresponds with the timing of insufflation of the colon and facial grimacing thus indicating that the medial procedure produces alterations in brain processing during the elicitation of pain. Furthermore, the NIRS signal observed was similar to the signal produced by a pain stimuli in the same region on a patient not subjected to anesthetics.
Furthermore a reproducible signal over the frontal cortex was observed and although sensory representation of the colon is present in the primary somatosensory cortex, and a potential area to focus on recording a nociceptive signal, the area is substantially small and as a result was only observable in a subset of the patient population. Furthermore, the observed signal was obtained from observation of grimace in the patients, thus indicating that a movement may be observed in conjunction with pain.

The observed signal in frontal cortex is similar to the phasic thermal and electrical stimulation further suggesting that the signal indicates patient pain. Based on the results obtained in this study, afferent pain signals are shown to be processed in the brain of patients undergoing colonoscopy with moderate sedation.

The present pain detector apparatus and methods can provide continuous monitoring of neural activity related to pain experienced during medical procedures. That apparatus may provide control of analgesic load to improve patient intraoperative comfort and decrease postoperative pain. In addition, the measure or detection of pain may be used to evaluate the effectiveness of pharmaceuticals for a variety of conditions.
WHAT IS CLAIMED IS:

1. A method that recognizes pain in a non-responsive patient, comprising:
   obtaining functional near-infrared spectroscopy imaging data from a non-responsive patient to recognize pain in the patient.

2. A method that recognizes pain in a non-responsive patient, comprising:
   obtaining functional near-infrared spectroscopy imaging data from at least one cortical region of the brain of the non-responsive patient;
   comparing measured imaging data with patient pain based on control data and thereby recognizing pain in the non-responsive patient.

3. The method of claim 2 wherein using the imaging data one or more hemodynamic parameters are measured in the at least one cortical region of the brain of the non-responsive patient.

4. The method of claim 3 further comprising one or more of (i) averaging the measured plurality of hemodynamic parameters to reduce background noise; (ii) identifying a pattern in the averaged hemodynamic parameters; and/or (iii) outputting a signal indicating patient pain.

5. The method of any one of claims 3 through 4 wherein the hemodynamic parameters include one or more of oxygenated hemoglobin, deoxygenated hemoglobin, and total hemoglobin.

6. The method of any one of claims 3 through 5 wherein measuring at least one hemodynamic parameter includes measuring oxygenated and deoxygenated hemoglobin to measure blood flow.

7. The method of any one of claims 2 through 6 wherein the at least one cortical region of the brain is selected from the somatosensory cortex and the prefrontal cortex.

8. The method of any one of claims 4 through 7 wherein the predetermined pattern indicating pain comprises a plurality of sequential peaks in blood flow over time.

9. The method of any one of claims 2 through 8 wherein the control data comprises
pain indicia is detected in the patient while a patient is responsive.

10. The method of any one of claims 3 through 9 wherein the pattern associated with pain is obtained from the control data for the hemodynamic parameter wherein pain is detected in a conscious patient.

11. The method of any one of claim 2 through 10 wherein the control data is obtained by acquiring the optical imaging data from the at least one cortical region of the brain of the conscious patient.

12. The method of claim 10 wherein acquiring the optical imaging data when the patient is not subject to anesthetics includes performing an activity to the patient.

13. The method of claim 12 wherein the activity is a finger tapping exercise and a variation of electrical stimulus.

14. The method of claim 12 further comprising administering an anesthetic regimen based on a particular medical procedure toward the patient after obtaining the control data.

15. A system that recognizes pain in a non-responsive patient, comprising:
   a functional near-infrared spectroscopy configured to capture optical imaging data from at least one cortical region of the brain of the non-responsive patient; and
   a processor configured to:
       measure, using the optical imaging data, a plurality of hemodynamic parameters in the at least one cortical region of the brain of the non-responsive patient.

16. The system of claim 15 wherein the processor is further configured to perform one or more of the following (i) through (v):
   (i) average the measured plurality of hemodynamic parameters;
   (ii) identify a pattern in the averaged hemodynamic parameters;
   (iii) compare the pattern to at least one predetermined pattern associated with patient pain;
   (iv) detect a similarity between the pattern and the predetermined pattern; and/or
   (v) output a signal indicating patient pain.
17. The system of claim 15 or 16 wherein the hemodynamic parameters include one or more of oxygenated hemoglobin, deoxygenated hemoglobin, and total hemoglobin.

18. The system of any one of claims 15 through 17 wherein the at least one cortical region of the brain is selected from the somatosensory cortex and the prefrontal cortex.

19. The system of any one of claims 15 through 18 wherein the processor is further configured to control an analysis exercise and a variation of electrical stimulus on the non-responsive patient to inflict a level of pain.

20. The system of claim 15 wherein the processor is further configured to determine whether a time delay between performing an analysis exercise and the variations of electrical stimulus and a point when a peak occurs in blood flow indicates pain in the patient.

21. A method that evaluates a treatment, comprising:
   - administering a treatment to a patient;
   - applying a stimulus to the patient;
   - acquiring functional near-infrared spectroscopy imaging data during the application of the stimulus;
   - determining from the imaging data, brain activity data representing a hemodynamic parameter in at least one cortical region of the brain of the patient;
   - comparing the brain activity data representing the hemodynamic parameter to control data representing the hemodynamic parameter; and
   - outputting an indication of patient pain.

22. The method of claim 21 further comprising acquiring control data using the functional near-infrared spectroscopy prior to obtaining the activity imaging data.

23. The method of claim 21 or 22 wherein determining a reduction in intensity of the hemodynamic parameter from the control data to the imaging data indicates a relative efficiency of the treatment as a pain-relieving treatment.
FIG. 2

Left Prefrontal Cortex

D17  S9  D18

Right Prefrontal Cortex

D20  S10  D19

Right Sensorimotor Cortex

D12  D11  D10  D9  D8  D7  D6  D5  D4

--- Oxygenated hemoglobin (HbO)
---- Deoxygenated hemoglobin (HbR)
------ Total blood volume (HbT)

DXX: Detector Number
SY: Source Number
START

CAPTURE OPTICAL IMAGING DATA FROM AT LEAST ONE CORTICAL REGION OF THE BRAIN

MEASURE HEMODYNAMIC PARAMETERS IN THE CORTICAL REGION OF THE BRAIN

AVERAGE THE MEASURE THE HEMODYNAMIC PARAMETERS

IDENTIFY A PATTERN IN AVERAGED HEMODYNAMIC PARAMETERS

COMPARE PATTERN TO PREDETERMINED PATTERN ASSOCIATED WITH PATIENT PAIN

DETECT SIMILARITY BETWEEN PATTERN AND PREDETERMINED PATTERN

OUTPUT SIGNAL INDICATING PAIN

END

FIG. 5
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER
A61B 5/00(2006.01)i, A61B 5/145(2006.01)i, A61N 1/36(2006.01)i

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)
A61B 5000; A61B 25/00; A61B 50/476; A61N 5/06; A61K 31/55; A61N 1/36; A61M 37/00; A61B 5/145

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched
Korean utility models and applications for utility models
Japanese utility models and applications for utility models

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
eKOMPASS(KIPO internal) & Keywords: recognize, pain, near-infrared spectroscopy, cortical, non-responsive

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<tbody>
<tr>
<td>X</td>
<td>US 2011-0152249 A (DAVID ROGROWSKI et al.) 23 June 2011 See abst ract, paragraphs [0015], [0079], claims 1, 22, 57, 72 and figure 1A.</td>
<td>1-5, 15-17, 20</td>
</tr>
<tr>
<td>A</td>
<td>US 2011-0245708 A (JULIA C. FINKEL et al.) 06 October 2011 See abst ract, paragraphs [0051], [0053] and figures 3A, 3B.</td>
<td>1-5, 15-17, 20</td>
</tr>
<tr>
<td>A</td>
<td>US 2008-0249430 A (ERWIN ROY JOHN et al.) 09 October 2008 See abst ract, paragraph [0018], claim 1 and figure 1.</td>
<td>1-5, 15-17, 20</td>
</tr>
<tr>
<td>A</td>
<td>US 2011-0118661 A (PREMAMIN D. LESS et al.) 19 May 2011 See abst ract, paragraphs [0054], [0072] and figures 1-3.</td>
<td>1-5, 15-17, 20</td>
</tr>
<tr>
<td>A</td>
<td>US 2009-0054955 A (BRIAN H. KOPPEL et al.) 26 February 2009 See abst ract, paragraphs [0077], [0086], claim 18 and figure 1A.</td>
<td>1-5, 15-17, 20</td>
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Further documents are listed in the continuation of Box C. See patent family annex.

Date of the actual completion of the international search
29 May 2014 (29.07.2014)

Date of mailing of the international search report
30 July 2014 (30.07.2014)

Name and mailing address of the ISA/KR
International Application Division
Korean Intellectual Property Office
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Authorized officer
KGM. Tae Hoon

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<table>
<thead>
<tr>
<th>Box No. II</th>
<th>Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)</th>
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<tr>
<td>1. ☒ Claims Nos.: 21-23 because they relate to subject matter not required to be searched by this Authority, namely: Claims 21-23 pertain to a method of treatment of the human body by surgery or by therapy/diagnostic methods, and thus relate to a subject matter which this International Searching Authority is not required, under Article 17(2)(a) of the PCT and Rule 39.1(iv) of the Regulations under the PCT, to search.</td>
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<td>2. ☒ Claims Nos.: 12-14 because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically: Claims 12-14 are unclear because they refer to multiple dependent claim 10 which does not comply with PCT Rule 6.4 (a).</td>
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<td>3. ☒ Claims Nos.: 6-11,18,19 because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).</td>
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<th>Box No. III</th>
<th>Observations where unity of invention is lacking (Continuation of item 3 of first sheet)</th>
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<tr>
<td>1. ☐ As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.</td>
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<td>2. ☒ As all searchable claims could be searched without effort justifying an additional fees, this Authority did not invite payment of any additional fees.</td>
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<td>3. ☐ As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:</td>
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<td>4. ☒ No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:</td>
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**Remark on Protest**
- ☐ The additional search fees were accompanied by the applicant's protest and, where applicable, the payment of a protest fee.
- ☒ The additional search fees were accompanied by the applicant's protest but the applicable protest fee was not paid within the time limit specified in the invitation.
- ☒ No protest accompanied the payment of additional search fees.
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