Electroencephalography for the diagnosis of brain death

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Electroencephalography (EEG) is frequently used to assist the diagnosis of brain death. However, to date there have been no guidelines in terms of EEG criteria for determining brain death in Korea, despite EEG being mandatory. The purpose of this review is to provide an update on the evidence and controversies with regarding to the utilization of EEG for determining brain death and to serve as a cornerstone for the development of future guidelines. To determine brain death, electrocerebral inactivity (ECI) should be demonstrated on EEG at a sensitivity of 2 \( \mu V/mm \) using double-distance electrodes spaced 10 centimeters or more apart from each other for at least 30 minutes, with intense somatosensory or audiovisual stimuli. ECI should be also verified by checking the integrity of the system. Additional monitoring is needed if extracerebral potentials cannot be eliminated. Interpreting EEG at high sensitivities, which is required for the diagnosis of brain death, can pose a diagnostic challenge. Furthermore, EEG is affected by physiologic variables and drugs. However, no consensus exists as to the minimal requirements for blood pressure, oxygen saturation, and body temperature during the EEG recording itself, the minimal time for observation after the brain injury or rewarming from hypothermia, and how to determine brain death when the findings of ECI is equivocal. Therefore, there is a strong need to establish detailed guidelines for performing EEG to determine brain death.

Key words: Electroencephalography; Brain death; Electrocerebral inactivity
Brain death is a term that implies irreversible loss of function of entire brain, including brainstem and hemisphere. The concept of brain death first emerged in the 1950s, driven by progress in critical care medicine such as cardiopulmonary resuscitation and the development of mechanical ventilation.

Since the ad hoc Committee of Harvard Medical School defined the characteristics of irreversible coma as 1) unreceptivity and unresponsiveness, 2) the absence of movement and breathing, 3) the absence of brainstem reflexes, and 4) flat electroencephalogram, a myriad of criteria for determining brain death have been suggested and evolved. However, a recent review reported that there was significant variation among the criteria and protocols to determine brain death according to countries and institutions, although most countries have legal provisions on brain death. Particularly, ancillary tests to confirm brain death are used in variation. The most frequently required test is an electroencephalography (EEG), which is mandatory in 28% and optional in 47% of countries. In Korea, standards for determining brain death were first established in 1983 by the ‘Committee for defining death’ of the Korean Medical Association. On Feb 8, 1999 ‘Laws on organ transplantation’ have been established, defining brain death only for the purpose of organ transplantation and requiring a flat EEG for at least 30 min. The most recent Korean standards for determining brain death have been stated in Article 21 of the enforcement decree of the Organ Transplant Act.

Despite the fact that number of annual cases requiring determination of brain death reached 537 cases in 2015, to date no specific guidelines or consensus with regard to performing and interpreting EEGs for the diagnosis of brain death have been established in Korea. The goal of this review is to provide an update on the role of EEG for determining brain death and the potential confounding factors, for development of future guidelines.

**EEG FINDINGS REPRESENTATIVE OF BRAIN DEATH**

The consensus is that EEG potentials should be absent to declare brain death. A survey by the American EEG Society’s ad hoc Committee on the EEG Criteria for the Determination of Cerebral Death in the late 1960s revealed that, three out of 2,650 adult cases of coma with presumably “isoelectric” EEGs met the committee’s criteria showed any recovery of cerebral function. These three cases were due to massive overdoses of nervous system depressants. Non-physiologic terms such as “electrocerebral silence”, “isoelectric,” “linear,” and “flat” were replaced in the 1970s with the term “electrocerebral inactivity (ECI),” which appeared in the Glossary of the International Federation of Clinical Neurophysiology.

ECI was defined as absence over all regions of head of identifiable electrical activity or cerebral origin, whether spontaneous or induced by physiologic stimuli and pharmacological agents. The widely accepted working definition of ECI is the absence of non-artifactual electrical activity with a peak to peak amplitude above 2 μV when recording from scalp electrode pairs that were 10 cm or further apart.

**CLINICAL SETTING AND CONFOUNDING FACTORS**

ECI is required to confirm cessation of brain function, but this doesn’t ensure either the irreversibility or loss of whole brain function. EEG as well as the clinical examination is affected by physiologic variables and medication. The Korean law lists the following as a prerequisite for determining brain death: A. The absence of any treatable drug toxicities (such as due to anesthetics, hypnotics, sedatives, muscle relaxants, or poisons); B. The absence of any treatable endocrinological disorders (such as hepatic coma, uremic coma, hypoglycemic encephalopathy); C. The absence of hypothermia (defined as a rectal temperature of 32°C or lower); D. The absence of a shock-like state. If sedative medications were used before the diagnosis of brain death, the French guidelines recommend using the techniques based on the study of intracerebral blood flow such as cerebral angiography, which are not influenced by the medications, rather than EEG.

American Clinical Neurophysiology Society (ACNS) guidelines 2016 newly addressed the item that blood pressure and oxygen saturation should be recorded. The French guidelines on EEG recommended maintaining the patient’s body temperature above 34°C and that the EEG should be performed at least 12 hours after cardiorespiratory arrest. Particularly, determining brain death needs caution in the setting of...
therapeutic hypothermia. There had been a case of a patient who had regained cough reflex and spontaneous respiration in the operating room for organ procurement 48 hours after rewarming from therapeutic hypothermia.\textsuperscript{13}

The guidelines as to when the EEG should be recorded after the brain injury, as well as blood pressure, oxygen saturation, body temperature, and the minimal time required after rewarming from hypothermia should be further developed. Moreover, ECI may be seen in cases where subcortical or brainstem function remains intact as EEG represents cortical activity.\textsuperscript{14} This may be seen in hypoxic ischemic encephalopathy as the cerebral cortex is more susceptible to ischemia.\textsuperscript{15}

\section*{REQUIREMENTS FOR EEG ACQUISITION}

\subsection*{1. Electrodes and montage}

A full set of scalp electrodes over all major brain areas using conventional 10-20 system should be used. In certain cases, recording with a full set of conventional 10-20 scalp locations may not be feasible such as due to head trauma or recent surgery. In these cases, the electrode positions may be moved as necessary, as long as careful documentation is made and the inter-electrode distances remain at least 10 centimeters.\textsuperscript{10}

The French guidelines stated that a minimum of 8 electrodes should be placed, whereas the American Academy of Neurology (AAN) recommend at least 16 channels.\textsuperscript{7} French guidelines recommended the placement of FP2, C4, O2, T4, FP1, C3, O1, T3 and Cz electrodes. The ACNS guidelines maintained that midline electrodes (Fz, Cz, Pz) should be included because these electrodes are useful for the detection of residual low-voltage physiologic activity, and are relatively free from artifacts.\textsuperscript{10}

Although a ground electrode should be added in most cases, a ground electrode should not be used if grounds from other electrical equipment are already attached to the patient, which often happens in intensive care units because of safety issue according to ACNS guidelines 2006.\textsuperscript{9} This recommendation was replaced with the requirement of an ‘isolated’ ground electrode in ACNS guidelines 2016.\textsuperscript{10}

Inter-electrode impedance should be balanced between 100 and 10,000 Ohms as a routine.\textsuperscript{10,16} High impedance results in more artifacts and can be decreased by cleansing the scalp. It is also affected by the quality of the electrodes. Impedances below 100 Ohms may be a possible reason for a false-positive ECI recording. It is essential that excess electrode paste does not spread from one electrode to another, creating a shunt or short circuit, which may also attenuate the signal.\textsuperscript{10} Electrode impedance should be rechecked during the recording when any type of artifact appears. normal impedance in an electrode demonstrating noise may still necessitate changing or modifying that electrode.\textsuperscript{16}

The use of needle electrodes and “electrode caps” should be avoided.\textsuperscript{10} Not only are there concerns such as patient discomfort and potential risk of injury to the staff, but these electrodes have higher impedance than cup electrodes, resulting in potentially higher levels of noise.\textsuperscript{16} When using electrode caps, excess jelly spreading from one electrode to another can create a short circuit, which may attenuate the signal.\textsuperscript{9}

\subsection*{2. EEG recording}

A sampling rate of at least 256 Hz is recommended according to the French guidelines.\textsuperscript{12} The sensitivity must be increased to a maximum of 2 μV/mm. The calibration voltage should be either 2 or 5 μV, which differs from routine EEG recording.\textsuperscript{10}

With regard to filter settings, the high-frequency (low pass) filters should not be set below 30 Hz, and the low-frequency (high pass) filters should not be set above 1 Hz.\textsuperscript{10} The French recommendations are more strict; 70 Hz or higher for the low pass filter and 0.53 Hz for the high pass filter.\textsuperscript{12} The 50 or 60-Hz notch filter can be used with care, but there should be segments of EEG that are recorded without this filter for comparison.\textsuperscript{10}

If the EEG suggests ECI, the integrity of the system should be tested by gently touching or shaking each electrode of the montage to create an artifact potential. This verifies that the electrode board is connected to the recording device and that the montage settings match the electrode placement.\textsuperscript{10,12} At least 30 minutes of EEG recording is recommended by Korean law as described in most guidelines,\textsuperscript{10,12} based on the observation that intermittent cortical activity with amplitudes as low as 2 μV may appear at intervals of up to 20 minutes during continuous recording for up to 8 hours.\textsuperscript{17}

Auditory and bilateral somatosensory stimuli (touch and pain) should be repeatedly performed, and be clearly marked
on the recording (i.e., the point where the somatosensory stimulus was applied). The technician must document all the events that occur during the test on the EEG tracings.

The Korean law mandates that EEG should be performed once after the second clinical examination and the apnea test in patients aged one year or above, and twice (before and after the second clinical examination) in infants between ages two months and one year. Although it is stated to perform a repeat clinical examination 48 hours later in those less than 1 year of age, it is unclear as to what the time intervals between the two EEGs should be. The French guidelines recommend a repeat EEG in all age groups. The time interval between the two EEGs was recommended to be at least 4 hours (the starting point being the beginning of each EEG recording) for those more than 1 year of age; 24 hours between the ages of 1 month and 1 year; 48 hours between the ages of 7 days and 1 month. The American guidelines state that one EEG should be sufficient but added the following comments: In the event that technical or other difficulties lead to uncertainty in the evaluation of ECI, the entire procedure should be repeated after an interval. This may be as short as 6 hours in adult patients, but in neonates and children the interval should be at least 24 hours. Consideration should be given to other confirmatory tests if technical limitations are unlikely to be overcome in subsequent recordings, based on the opinions of the treating physicians.

The basis as to why a single EEG is sufficient is described in the American EEG guidelines as the following; in the Collaborative Study of Cerebral Death, no patients survived for more than a short period after an EEG showed ECI, excluding the cases which were due to overdosing with central nervous system (CNS) depressants. This provides evidence that a single EEG demonstrating ECI is a highly reliable tool for determining cortical death.

3. Evaluation of extracerebral potentials

The technician should recognize the common sources of artifact, either internal or external, such as electrical artifact originating from electric beds, IV drips, blood warmers, or other electrical devices. Electric potentials of extracerebral origin often cannot be avoided at high sensitivities in intensive care unit (ICU) and responsible for the disagreement in the interpretation. Therefore, additional monitoring is required to differentiate cerebral from extracerebral potentials. The ACNS guidelines recommend the following (modified):

1. Electrocardiogram monitoring is essential. This may be monitored by using two precordial disc electrodes.
2. Continuous video recording is strongly recommended to help identify any artifact in the EEG recording.
3. Respiration artifact must be documented with specific notations on the record or monitored by a transducer if this cannot be eliminated. Brief disconnection of the respirator should allow definitive identification of the artifact, if clinically feasible.
4. Motion monitors should be placed if intermittent movements are noted in a limb.
5. An additional monitor is often needed for other artifacts emanating from the patient or the local environment. The most convenient for this purpose is a pair of electrodes on the dorsum of the hand separated by about 6 to 7 cm.
6. Machine noise, thermal noise, and electrical interference

Fig. 1. An example of a dummy patient with a three-lead device (Grass Technologies, West Warwick, RI, USA). This can be used to measure machine noise and external interference which enters to the recording system.
entering the recording system from the jack box to the amplifiers may be measured on a “dummy patient” (Fig. 1), i.e., placing a 10,000-Ohm resistor between input terminal 1 (G1) and input terminal 2 (G2) of one of the channels, as long as either G1 or G2 is shorted to the reference electrode.

7) If electromyogram potentials are of such amplitude as to obscure the tracing, it may be necessary to reduce or eliminate them by using a short-term neuromuscular blocking agent. As this may interfere with the neurologic examination and cause other problems, neuromuscular blockade should be performed under the guidance of an experienced physician familiar using medications in critically ill patients.

INTERPRETATION

An estimate must be made of whether the amplitudes exceed 2 μV. The interpretation should be carried out using long distance montages such as longitudinal or transverse bipolar montages with double distance electrodes. Filters should be avoided. It is actually very difficult to acquire EEG without artifacts higher than 2 μV/mm in ICU (Fig. 2). Hence, determining ECI creates diagnostic challenges, and significant inter- and intra-rater disagreement has been reported. In practice, it is necessary to tolerate artifacts, as long as it can be ensured that they are artifacts. Determining whether the signals shown in EEG are cerebral or not can be done cautiously based on the results of the additional monitoring as indicated in III-3. Reviewing an accompanying video recording can also be useful.

When a lack of confidence exists, the EEG report must reflect such uncertainty, and the record cannot be stated as demonstrating ECI. In Korea, ECI is mandatory for declaring brain death by law, and no guidelines exist for determining brain death when it is uncertain whether the EEG truly shows ECI or not. According to statistics from Korea Organ Donation Agency since 2010, only one patient was diagnosed with brain death without the EEG results clearly demonstrating ECI due to difficulties in performing the EEG itself as a result of craniectomy (personal communication). Details of the EEG reports for the diagnosis of brain death which were suggested in French recommendation are as follows (modified):

1. Administrative data: the administrative data for routine EEG as well as the location and exact times indicating the beginning and end of the recording must be specified.
2. Clinical information: the circumstances and origin of the coma, especially the presumed date of the onset of the coma; the time lapse from the event which resulted in brain death; imaging data; clinical criteria for brain death; the presence of interfering factors: toxicological or metabolic; drugs administered in the past 24 hours; body temperature; blood pressure; oxygen saturation; specific recording conditions (e.g. scalp condition, environment).
3. EEG description: read-out on long-distance montages; the duration of the interpretable recording; background activity: indicate the maximum amplitude measured aside from the artifacts; results from auditory, somatosensory tactile and nociceptive stimulations; describe an eventual diffusion of the electrocardiogram and its topography (diffuse, diffusion only on certain derivations).
4. Conclusion: Absence of all brain electrical activity under the stated conditions of the EEG recording.

PERSONNEL

The recording should be performed by a specialized technician, qualified to identify and eliminate artifacts, implement the required settings, perform standard activation procedures and document all relevant data in the recording. The report should only be created by a physician specializing in EEG or Neurophysiology.

CONCLUSIONS

To determine brain death, ECI should be demonstrated on scalp EEG which is recorded as following criteria. 1. The sensitivity of 2 mW/mm; 2. Inter-electrode distances more than 10 centimeters; 3. Covering over all major brain areas including midline area 4. Recording for at least 30 minutes; 5. Giving intense somatosensory or audiovisual stimuli. ECI should be also verified by checking the integrity of the system. Evaluation of extracerebral potentials is required, because such potentials are often unavoidable at high sensitivity in ICU. Physicians should keep in mind the anatomical and physio-
logic limitation of scalp EEG. A detailed guideline is needed for determining ECI and confirming brain death in cases of uncertainty in the evaluation of ECI.

Fig. 2. EEG of a 70-year-old comatose male recorded 3 days after cardiopulmonary arrest. (A) The EEG appeared flat at sensitivity 7 μV/mm without any reactivity to painful stimuli. (B) Low-voltage, mixed-frequency activity was seen at sensitivity 2 μV/mm in the same epoch, which could not be eliminated. The patient never regained consciousness, spontaneous respiration, or brainstem reflexes after the cardiopulmonary arrest. A repeat EEG performed 12 days after the arrest showed the same pattern. EEG, electroencephalography.

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