Case report

Eye-opening in brain death: A case report and review of the literature

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1. Introduction

Brain death is defined as the complete and permanent absence of all brain functions (Busl and Greer, 2009). Protocols for ascertaining brain death vary widely from country to country (Kondziella, 2020). In Italy, a patient is considered brain dead when he is in a coma, showing apnoea under particular clinical parameters, and having no brain stem reflexes (Legge 29.12.1993 n. 578; Decreto Ministero della Salute 11.04.2008). On the other hand, presence of spinal reflexes, when correctly identified, does not hamper the diagnosis. However, distinguishing between “extracerebral” reflexes and persistent brainstem function at times is challenging, as in the case of eyelid elevation, which is determined by two separate mechanisms. In one case, it can be considered a residual brainstem activity due to the action of the eyelid elevator muscle that is innervated from the fibres of the III cranial nerve. In other cases, it can be considered an “extracebral” reflex because of the action of the Muller’s muscle innervated from sympathetic fibres. We report a case of eyelid elevation following pain stimulation in a patient who met all clinical and instrumental brain-death criteria. Following a review of the literature, we propose an approach to similar cases to allow the timely diagnosis of legal brain death, which is especially important in cases involving potential organ donors.

2. Case report

A 43-year-old woman with a history of craniopharyngioma and pituitary insufficiency was admitted to the emergency room in a deep coma state and acute renal failure following a urinary infection. Fifteen days after admission, her condition rapidly deteriorated, and she suddenly entered cardiopulmonary arrest. An electrocardiogram showed pulseless electrical activity. Cardiopulmonary resuscitation (CPR) was promptly started and 1 mg adrenaline was administrated. She was intubated and moved to the intensive care unit (ICU). CPR continued for about two hours. After resuscitation, she remained comatose with bilateral mydriasis and a Glasgow Coma Scale (GCS) score of 3. Brain computed tomography (CT) showed brainstem and intraventricular haemorrhage with hydrocephalus in the brainstem. On the following day, 24 h after resuscitation, her neurological status had not changed; hence, a neurophysiological evaluation for the long-term neurological prognosis was performed (Scarpino et al., 2018). The electroencephalogram (EEG) showed an isoelectric pattern (Fig. 1), while the somatosensory evoked potentials (SEPs) after stimulation of the median nerves at the wrist revealed the absence of both bulbar and cortical responses in the presence of peripheral and spinal responses (Fig. 2). Since these neurophysiological findings suggested the presence of brain death (Scarpino et al., 2017), her
Brainstem function was tested. Neurological examination showed the absence of the photomotor, oculovestibular, corneal, gag, and cough cranial reflexes. Moreover, the patient, after being hyperventilated (4 acts per minute with a tidal volume of 250 cc) until reaching a blood gas CO₂ value >60 mmHg, was disconnected for one minute from the ventilator, showing the absence of respiratory drive. Even though there was no facial motor response after painful stimulation in the trigeminal territory, the patient presented partial bilateral eyelid elevation, after about two seconds, in response to painful stimulation of the left or right nipple (Fig. 3). Immediately after cessation of the painful stimulation, the eyelid slowly returned to baseline. Inspection of the pupils showed no diometric change. Consequently, intensivists suspended the brain death ruling and scheduled a reevaluation. Two hours later, after re-examination, the same neurological signs were observed. The diagnosis of brain death was again not established and another evaluation was scheduled for the following day. Given the persistence of the same clinical conditions associated with an EEG confirming an isoelectric pattern for the other two days (three days subsequent to cardiac arrest), a cerebral CT angiography was thus arranged, showing no opacification of intracranial arterial circulation and of venous circulation. Therefore, the medical board convened on the same day and established a diagnosis of BD. Eyelid elevation after painful nipple stimulation was no longer present in addition to the absence of other confounding factors for reaching the diagnosis of BD, including episodes of hypotension, hypoxaemia or drug administration. The legal tutor consented to organ donation and no autopsy was performed.

3. Ethical statements

According to Italian law, case reports do not need to be approved by the Ethics Committee; however, the work complies with the ethical guidelines of the Helsinki declaration and the Oviedo convention, as well as the ethical standards of the University of Florence.

4. Discussion

According to Italian law (Legge 29.12.1993 n. 578; Decreto Ministero della Salute 11.04.2008), a first documentation of clinical signs, a repetition of clinical testing and a confirmation of the absence of bioelectrical activity in the brain (EEG) are required in order to establish a diagnosis of brain death (Zappa et al., 2020). The prerequisite for brain death assessment is a body temperature not altering the functions of the brain (usually >35 °C), and a known cause of brain injury. To establish the diagnosis, a Legal Committee must include a medico-legal doctor (delegate of the Head of Hospital or a pathologist), an anaesthetist and a neurophysiologist (either a neurologist or neurosurgeon with expertise in electroencephalography) who must express a unanimous judgment on the time of death if the specific criteria are present.
after two evaluations taken 6 h apart (Decreto Ministero della Salute 11.04.2008). Criteria (Legge 29.12.1993 n. 578; Zappa et al., 2020; Wijdicks et al., 2010; Centro Nazionale Trapianti, 2009) for brain death diagnosis are summarized in Table 1. Spontaneous or triggered spinal activity has no relevance in the diagnosis of brain death (Decreto Ministero della Salute 11.04.2008). During brain death assessment, the request for organ donation is made to whoever legally qualifies.

We presented the case of a patient with a primary acquired brain injury who met all of the clinical criteria for brain death except for the presence of an unknown reflex: bilateral eyelid elevation evoked by painful stimulation on both the right and left nipples, which caused a delay in reaching a diagnosis of brain death.

Eyelid elevation is determined by the action of two muscles with different innervation. This neurological sign might have occurred due to the action of the eyelid elevator muscle that is innervated from the fibres of the III cranial nerve originating in the oculomotor nucleus, located in the mesencephalon. On the other hand, the eyelid elevation might result from Muller’s sympathetic muscle function. The sympathetic fibres, originating from the superior cervical ganglion, are divided into two branches. The first branch travels to where the carotid joins the VI cranial nerve and later the ophthalmic branch of the V cranial nerve, entering the orbit through the optic foramen and supplying the pupil (Parks, et al., 1978). The second branch, instead, which is directed to the Muller’s muscle, leaves the intrapetrous carotid plexus through the petrosal nerve, vidian nerve, and sphenopalatine ganglion reaching the orbit via the inferior orbital fissure (Duke-Elder and Wybar, 1961) (Fig. 4).

In our case, given both the strong clinical and instrumental (SEPs; cranial CT angiography) evidence of lack of any preserved brainstem function, the eyelid opening has been attributed to a residual preserved function of the superior cervical ganglion which was activated by thoracic pain fibres following stimulation to both right and left nipple levels.

Additionally, the patient’s partial and slow eyelid opening was consistent with the activity of a smooth muscle, such as the Muller’s muscle, rather than with the activity of a striated skeletal muscle such as the eyelid elevator muscle, innervated by the III cranial nerve.

Finally, the absence of pupil dilation in the presence of partial eyelid elevation could be attributed to a greater resistance to the loss of intracranial blood flow of the fibres innervating the Muller’s muscle compared to the fibres innervating the pupil (Santamaria et al., 1999).

According to the literature, patients fulfilling all clinical and instrumental criteria of brain death with the exception of neurological signs of eyelid openings are described in only a few previously published papers and all of them differ in some way from our case (Santamaria et al., 1999, Friedman, 1984).

In the paper by Friedman (Friedman, 1984), the author described a monolateral eyelid opening triggered by twisting the ipsilateral nipple. A similar case was also described by Santamaria et al. (Santamaria et al., 1999). Both authors considered the eyelid opening to be an extracranial reflex, although brain death was not declared in either case. Furthermore, the cause of death in the first case (Friedman, 1984) was unknown, whereas the patient in the second case (Santamaria et al., 1999) died from cardiac arrest. To date, we have not found any other similar case reports.

Clinical and instrumental features of these previous patients satisfying clinical and instrumental criteria of brain death except for the presence of an eyelid opening sign are reported in Table 2.

In our case, the patient showed bilateral eyelid elevation evoked by painful stimulation of both right and left nipple levels, regardless of the presence of all other clinical and instrumental (EEG) criteria for a diagnosis of brain death. The brain death diagnosis was halted in order to monitor the clinical conditions. However, based on the hypothesis that this neurological sign was obtained by an extracerebral reflex caused by the presence of a residual preserved function of the superior cervical ganglion activated by thoracic pain fibres following stimulation at both the right and left nipple levels, an angiography was thus performed, revealing the absence of cerebral flow.

In our experience, backed by the scientific literature, a slow partial eyelid opening in response to painful nipple stimulation, with no changes in the pupil diameter, rarely occurs in brain death.

Table 1
Clinical and instrumental criteria for brain death ascertainment.

| Clinical criteria | Instrumental criteria |
|------------------|-----------------------|
| Unconsciousness  | Electroencephalogram for 30 minutes Mandatory to revealing absence of spontaneous or triggered brain electric activity greater than 2 microvolts in all cerebral regions |
| absence of following brain stem reflexes: | cerebral CT angiography or transcranial Doppler only if using of drugs depressant on cerebral functions |
| Oculovestibular reflex | impossibility of evaluating brain stem reflexes or electroencephalogram |
| Corneal reflex | age < 12 months |
| Gag reflex | post-anoxic coma from cardiac arrest that occurred within 24 h |
| Cough reflex | |
| Pupillary inactivity to bright light in both eyes | |
| Motor response after painful stimulation applied to the trigeminal and facial territories: | |
| absence of spontaneous breath for 60 sec without ventilator if pCO₂ > 60 mmHg and pH < 7.40 | |

Fig. 4. Drawing showing the anatomy of the oculo-sympathetic pathway. Sympathetic fibers in the postero-lateral hypothalamus pass through the lateral brain stem and to the cilio-sphincter of Budge and Waller in the intermediolateral gray column of the spinal cord C8–T1. Preganglionic sympathetic nerves exit from the Cilio-sphincter of Budge and pass above the pulmonary apex and ascend up the carotid sheath to the Superior cervical ganglion. The postganglionic sympathetic neurons originate in the superior cervical ganglion and travel up the wall of the internal carotid artery. The sympathetic fibers, which travel with the carotid and supply the pupil, join the sixth nerve and later the ophthamlic branch (1) of the 5th nerve to enter the orbit through the optic foramen, whereas sympathetic fibers to Muller’s muscle leave the intrapetrous carotid plexus through the petrosal nerve, vidian nerve, and the sphenopalatine ganglion reaching the orbit via the inferior orbital fissure. Abbreviations: a, artery; n, nerve.
patients. This exceptional clinical phenomenon should not prevent a diagnosis of BD in a patient who otherwise fulfills the standard criteria for BD, i.e., coma, apnoea, and loss of oculo-vestibular, cervical, gag, cough, and somatomotor reflexes. This unusual response may suggest a residual lower cervical/upper thoracic cord sympathetic function. Notwithstanding the pathophysiological opinion confirming the extracranial origin of eyelid opening, a specific test consisting of the stellate ganglion block could ideally be performed for the confirmation of the clinical suspicion but, this procedure is to date not readily available in the clinical practice. Although the eyelid elevation in our case was considered of reflex nature, the legal committee considered it prudent to also perform cerebral flow evaluation, considering that Italian law requires the observations to proceed in case the BD criteria are met.

Therefore, in this particular clinical condition of slow eyelid opening in response to painful stimulation of the left or right nipple in a patient who otherwise fulfilled all clinical and instrumental BD criteria, we believe that the implementation of cerebral flow evaluation, as well as already performed for other confounding factors, is considered “prudential” as a preliminary assessment prior to determining BD to confirm the absence of any brainstem function.

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### Conflict of Interest

The authors declare that there is no conflict of interest regarding the publication of this article.

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### Table 2

| Source       | Sex | Age | Etiology of Brain Injury | Time to fulfilling of BD criteria | Characteristics of eye opening | EEG | Other test                                      | Cause of death |
|--------------|-----|-----|--------------------------|-----------------------------------|--------------------------------|-----|------------------------------------------------|----------------|
| Friedman, 1984 | M   | 57  | Prolonged Hypotension    | 24 h                              | Monolateral, triggered by twisting ipsilateral nipple, Associated with abdominal movements. |    | Stellate block                                  | Not known      |
| Santamaria et al., 1999 | M   | 47  | Brain-stem Hematoma     | 4 days                            | Monolateral, triggered by twisting ipsilateral nipple, No other motor manifestations |    | Cerebral Angiography                             | Cardiac Arrest  |
| Present Case  | F   | 43  | Intraparenchinal hemorrhage and cardiac arrest | 24 h                              | Bilateral, Triggered by twisting left or right nipple, No associated with other motor manifestations |    | Somatosensory Evoked Potentials, Cerebral Angiography | Brain Death    |

BD = Brain Death; M = Male; F = Female.