Case Report

Aphasic status epilepticus of frontal origin treated by resective surgery

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Abstract

Aphasic status epilepticus (SE) is a clinical entity of SE, but it has not been well recognized. We report a 43-year-old female with a chronic drug-resistant epilepsy with aphasic SE, treated by resective surgery. The patient showed long-lasting weekly episodes of hypokinesia, slow verbal response, and dysphasia, which were diagnosed as symptoms of aphasic SE. Magnetic resonance imaging showed encephalomalacia in the left frontal lobe with a hemosiderin rim. Intracranial electroencephalography revealed continuous spikes, predominantly on the left superior frontal gyrus with hemosiderin deposit. The aphasic symptoms were seen when ictal discharges gradually spread to the wide area of the left anterior frontal lobe, including the language area. The episodes of recurrent aphasic SE had disappeared by one year after the left anterior frontal resection. We should consider aphasic SE when language impairment is episodic, and consider surgical intervention in cases where it repeatedly occurs despite appropriate medical therapy.

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1. Introduction

A definition had been proposed for non-convulsive status epilepticus (NCSE): a change in behavior and/or mental status associated with 30 min of continuous epileptiform discharges in the electroencephalogram (EEG) [1]. A recent report proposed that emergency treatment should be initiated for focal status epilepticus (SE) with impaired consciousness at 10 min from onset [2]. NCSE is not able to be diagnosed from clinical signs alone. Therefore, emergent EEG is indispensable. Improvement of both clinical and EEG features with the administration of intravenous antiseizure medication may suggest a diagnosis of NCSE [3]. NCSE could be caused by various etiologies, including cerebral stroke [4], infection [5], metabolic disease [6,7], neurological disease [8], or epileptic syndrome as absence epilepsy [9]. It has a higher prevalence in the elderly population and in those with post-stroke epilepsy [4,10]. Aphasia may appear as a focal seizure [11], or as a symptom of SE. Aphasic SE has recently been included in the category of NCSE without coma, and sub-classified in focal SE [2]. Most reported cases with aphasic SE showed de novo onset and improved seizure outcomes with medical therapy [5,7,8,12–18]. However, to the best of our knowledge, no surgical cases via intracranial EEG recording have been reported for repeat aphasic SE. We report the case of a patient with weekly aphasic SE of frontal origin, in which a resective surgery was effective in controlling the aphasic SE.

2. Case presentation

A 43-year-old right-handed female was admitted to our hospital with a 10-year history of drug-resistant seizures. She showed normal mental and physical development. She reported severe headaches during the postpartum period, seven years previous to the seizure onset; although, no neuroimaging examinations were performed. Her seizures began at the age of 33. Her habitual seizures manifested as the following: a focal aware cognitive seizure (aphasia) with abrupt onset; a focal aware clonic seizure, lasting for over 30 min, on the right-side of her face and arm, followed by post-ictal paralysis; and a long-lasting loss of spontaneous speech and hypokinesia with obscure onset, which occurred intermittently for one to two days. During the long-lasting symptom, she often lay in bed throughout the day without eating. This occurred weekly in the last one year before admission to the hospital, despite polytherapy with carbamazepine, levetiracetam, clobazam and perampanel. Neurological examination revealed no abnormality. The Wechsler Adult Intelligence Scale - Third Edition revealed a low intelligence quotient (IQ) (full IQ: 69, verbal IQ: 69, performance IQ: 74). The Standard Language Test of Aphasia disclosed disturbances in word fluency, semantic comprehension, and calculation (Table 1). Long-term scalp video-EEG showed the patient’s indifferent face impression, hypokinesia, decreased spontaneous speech, slow verbal response, and dysphasia with no obvious ictal onset. This occurred approximately when the bilateral frontal dominant continuous slow
waves were recorded superimposed by spikes on Fp1, F3, C3 and F7. She could eat and perform activities of daily living during this time, however, this was done slowly. Her EEG findings and her verbal response markedly improved after an intravenous injection of diazepam. As this altered cognitive function continued for over 30 min, we diagnosed this long-lasting symptom as aphasic SE. Magnetic resonance imaging (MRI) showed encephalomalacia with a hemosiderin rim around the left superior frontal sulcus (Fig. 1 upper right). Interictal [123]-iomazenil single-photon-emission computed tomography showed hypo-accumulation in the left frontal lobe, coincident with the MRI abnormality.

Considering the patient’s seizure frequency and drug-resistance, we subsequently pursued epilepsy surgery localized by intracranial EEG monitoring using subdural electrodes. Intracranial EEG showed frequent spikes on the cortex around the cavity, as well as in the middle and inferior frontal gyri, and the orbitofrontal cortex (apparently without symptoms) (Fig. 2A). Subsequently, continuous spikes in these area were seen to coincide with unpleasant moods (Fig. 2B). The discharges then gradually spread to the posterior middle and inferior frontal gyri, at which she showed indifferent face impression, hypokinesia, decreased spontaneous speech, slow verbal response, and dysphasia (Fig. 2C). The clinical symptoms and EEG findings improved by oral administration of diazepam (Fig. 2D). The anterior language area was detected by direct electrical stimulation (rectangular biphasic 50 Hz, 0.2-millisecond pulse width, 1–5-second duration under 7 mA bipolar stimulation) (Fig. 1, left). With consideration of the extent of sustained spikes and the results of functional mapping, we performed the left anterior frontal resection beyond the MRI abnormality (Fig. 1, upper right). Aphasic SE and the other focal seizures have not been seen as of one year after surgery. However, one week after surgery, the Standard Language Test of Aphasia revealed decreased vocabulary, dysphasia and impaired semantic comprehension with dysgraphia, but she retained normal repetition ability (Table 1). Her speech disturbance was assessed as transcortical sensory aphasia, which improved with speech therapy over one month.

### 3. Discussion

Aphasia may be due to a focal seizure or as a symptom of SE. Epileptic aphasia was defined by Rosenbaum et al. with several criteria as

### Table 1

| SLTA (standard language test of aphasia) | Pre-test | Post-test |
|----------------------------------------|----------|-----------|
| I. Hearing                              |          |           |
| Auditory word recognition               | 10/10    | 10/10     |
| Auditory sentence comprehension         | 10/10    | 10/10     |
| Verbal sequential commands              | 10/10    | 6/10      |
| Speech sound-kana letter choice matching| 10/10    | 10/10     |
| II. Speaking                            |          |           |
| Picture naming                          | 18/20    | 10/20     |
| Word repetition                         | 10/10    | 10/10     |
| Action naming                           | 10/10    | 5/10      |
| Picture story description               | Stage 6  | Stage 4   |
| Sentence repetition                     | 5/5      | 5/5       |
| Animal category fluency                 | 13       | 7         |
| Oral reading of kanji word              | 5/5      | 5/5       |
| Oral reading of kana letter             | 10/10    | 10/10     |
| Oral reading of kana word               | 5/5      | 5/5       |
| Oral reading of sentence                | 5/5      | 5/5       |
| III. Reading                            |          |           |
| Written kanji word-picture choice matching| 10/10    | 10/10     |
| Written kana word-picture choice matching| 10/10    | 10/10     |
| Written sentence-picture choice matching| 9/10     | 9/10      |
| Written sequential commands             | 10/10    | 8/10      |
| IV. Writing                             |          |           |
| Writing kanji names of pictures         | 4/5      | 1/5       |
| Writing kana names of picture           | 5/5      | 4/5       |
| Writing picture story                   | Stage 6  | Stage 4   |
| Writing kana letter to dictation        | 10/10    | 10/10     |
| Writing kanji word to dictation         | 4/5      | 2/5       |
| Writing kana word to dictation          | 5/5      | 5/5       |
| Writing dictated sentences              | 4/5      | 4/5       |
| V. Calculation                          |          |           |
| Written kanji names of pictures         | 13/20    | 10/20     |
| Written kana names of picture           | 5/5      | 4/5       |
| Written picture story                   | Stage 6  | Stage 4   |
| Writing kana letter to dictation        | 10/10    | 10/10     |
| Writing kanji word to dictation         | 4/5      | 2/5       |
| Writing kana word to dictation          | 5/5      | 5/5       |
| Writing dictated sentences              | 4/5      | 4/5       |

Fig. 1. Left anterior frontal resection with preserved language related area. Resection line (dotted line) and results of functional mapping. III: third finger, V: fifth finger, SF: superior frontal, MF: middle frontal, IF: inferior frontal, OF: orbitofrontal, T: temporal. Preoperative and postoperative MRI.
follows: speaking during the seizure, aphasic features, alertness, and simultaneous EEG demonstrating seizure activity [19]. Grimes and Guberman reported a case of **de novo** aphasic SE and proposed a modification of this criteria including that aphasia should be resolved following successful treatment [17]. Recently, aphasic SE was classified in NCSE and sub-classified in NCSE without coma, focal type [2], although the duration of aphasic SE has not been precisely defined. In the present case, we documented the patient’s long-lasting slow verbal response and dysphasia during the scalp Video-EEG monitoring. We diagnosed it as aphasic SE because her language fluency and facial impression recovered, corresponding with EEG improvement, due to an intravenous injection of diazepam.

Regarding aphasic SE, various symptoms have been reported as follows: aphemia/stuttering [12], Wernicke type aphasia [13], severe aphasia [14], and global aphasia [20]. NCSE of frontal origin could manifest as affective indifference [21]. The results of intracranial EEG demonstrated her symptoms (e.g., decreased spontaneous speech, slow verbal response, dysphasia with accompanying indifferent facial impression, and hypokinesia), were of frontal origin. Meanwhile, her preserved consciousness could be explained by the thalamus or brainstem being spared from seizure propagation [22]; although, no depth electrodes were implanted in these deeper brain structures.

The majority of reported cases with aphasic SE showed **de novo** onset with various etiologies, such as a structural lesion, infection, metabolic disease, neurological disease, or upon antiseizure medication taper [5–8,12–14,15–18,23,24]. Most of these cases had improved seizure outcomes with only medical therapies, if the underlying diseases were well treated. With our patient, however, epilepsy surgery was considered necessary because weekly aphasic SE had occurred over the last one year, despite appropriate use of medications. Recently, a child with drug-resistant epilepsy due to focal cortical dysplasia was successfully treated by complete lesionectomy; this was performed soon after the definite diagnosis of NCSE. Rapid cognitive decline of the patient in the last one year was considered to be related to frequent seizures and NCSE [25]. This clinical entity of aphasic SE is noteworthy; it presents the potential benefit from surgical intervention when disabling recurrent episodes are drug-resistant.

We performed intracranial EEG recordings as widespread epileptogenicity was suspected, involving language-related areas. It was challenging to decide the precise extent of the resection with the intracranial EEG findings alone during aphasic SE. Continuous spikes were recorded in the wide area of the left anterior frontal lobe, including the language area; although, precise time correlations between the aphasic symptoms and ictal discharges were at times less apparent. According to prior reports, electro-clinical correlation is sometimes unclear in aphasic SE [26], and aphasia may not correspond directly with the epileptiform activity [27]. Deciding on the border of resection considerably depended on the results of the brain anatomy and functional mapping.

Regarding the etiology of encephalomalacia with hemosiderin deposit, a peripartum intracerebral hemorrhagic stroke was suspected at the previous hospital [28]; however, there was no brain computed tomography evidence to provide radiographic support. NCSE occurs more prevalently in the elderly population [4]. It may also develop in middle-aged adults, similar to our patient, due to cerebral stroke as cerebral strokes are the main cause of adult NCSE [4,10].
4. Conclusion

Our patient presented with unique symptoms due to aphasic SE of frontal origin. This was verified by intracranial EEG. A tailored left anterior frontal resection was effective to control the recurrent episodes of aphasic SE. This case of aphasic SE is remarkable and supports a successful approach from surgical intervention, which may be necessary in cases where aphasic SE is difficult to treat and repetitive resulting in patient disability.

Declaration of competing interest

All the authors have no conflict of interest.

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