Successful Use of Therapeutic Hypothermia for Refractory Nonconvulsive Status Epilepticus

Do-Hyung Kim¹, Hye-hoon Kang¹, Minjung Kim¹, Tae-Won Yang², Oh-Young Kwon¹, Jung Sook Yeom³, Bong Su Kang⁴, Yong Hwan Kim⁵, Young-Soo Kim¹

¹Department of Neurology, Gyeongsang National University Hospital, Gyeongsang National University School of Medicine, Jinju; ²Department of Neurology, Changwon Gyeongsang National University Hospital, Gyeongsang National University School of Medicine, Changwon; ³Department of Pediatrics, Gyeongsang National University Hospital, Gyeongsang National University School of Medicine, Jinju; ⁴Department of Neurology, Yangpyeong Hospital, Yangpyeong; ⁵Department of Emergency Medicine, Samsung Changwon Hospital, Sungkyunkwan University School of Medicine, Changwon, Korea

Therapeutic hypothermia (TH) has rarely been utilized as an adjunct to anticonvulsants in treating patients with refractory convulsive status epilepticus (CSE). However, determining the effectiveness of TH in CSE is difficult due to the unavoidable use of sedative drugs to manage hypothermia. Additionally, the effectiveness of TH has not been studied in patients with refractory non-convulsive status epilepticus (NCSE). Here, we report the successful use of TH without additional sedative drugs in a patient with temporal lobe epilepsy and refractory NCSE. A 46-year-old man was referred to the neurology department because of recurrent seizure attacks. Electroencephalography (EEG) after first-line status treatment showed continuous periodic discharges consistent with NCSE. He was started simultaneously on continuous EEG monitoring and TH, but was not administered any benzodiazepines to control shivering or maintain TH. During TH, EEG abnormalities gradually improved, and the patient regained consciousness in accordance with the improvement in EEG. The patient was alert and his EEG had normalized a few days after starting TH. To the best of our knowledge, this is the first report describing the successful treatment of refractory NCSE with TH. As no sedative drugs were used during the maintenance of hypothermia, NCSE control may have been achieved by TH alone. (2017;7:109-114)

Key words: Hypothermia, induced, Status epilepticus, Electroencephalography

Introduction

Nonconvulsive status epilepticus (NCSE) is defined as a change in mental state from baseline, lasting at least 30-60 minutes, associated with continuous or near-continuous epileptiform discharge on the electroencephalogram (EEG).¹ NCSE is thought to constitute 20-25% of all episodes of status epilepticus (SE).² Refractory SE (RSE) has been associated with high morbidity and mortality rates, and does not always respond to standard pharmacological treatments. When these treatments fail, more active and aggressive alternative treatments are required.

Systemic therapeutic hypothermia (TH) has shown benefits in various clinical settings, including post-cardiac arrest and neonatal hypoxic encephalopathy, and has been found to control intracranial pressure in patients with traumatic brain injury and stroke.³⁻⁵ Experimentally, TH was shown to have anticonvulsant and neuroprotective effects after SE.⁶⁻⁷ Although several case reports have shown promising results following the application of TH in patients with RSE,⁸⁻¹⁰ TH was used as adjuvant therapy in almost all these patients. This report describes the successful use of TH without additional sedative drugs in a patient with temporal lobe epilepsy who experienced refractory NCSE. Findings in this patient suggest that TH can be considered an adjunctive or alternative therapy in patients with medically intractable NCSE.

Case

A 46-year-old man was referred to the neurology department of...
Figure 1. Ten-second EEG trace recordings with a 10-20 electrode system (A) following initial treatment of convulsive status epilepticus with lorazepam and fosphenytoin loading and showing continuous quasi-periodic single or poly-spikes and waves of high amplitude, mainly in both occipitoparietal areas; (B) after phenobarbital loading and just before induction of TH; (C) following induction of hypothermia to the 35°C target temperature; and after (D) 12 hours and (E) 24 hours of hypothermia at 35°C. EEG, electroencephalography.

Figure 2. Brain (A, B) diffusion MRI and (C, D) ADC maps after initial treatment for convulsive status epilepticus. Both showed symmetric restricted diffusion in bilateral occipitoparietal areas. MRI, magnetic resonance imaging; ADC, apparent diffusion coefficient.

Samsung Changwon Hospital because of recurrent generalized tonic-clonic (GTC) seizures. At that time, the patient was undergoing wound care at the Orthopedics Department of our hospital because of necrotizing fasciitis of the right foot. GTC seizures were observed twice by station nurses. The duration of each seizure was slightly over 5 minutes, with the patient not recovering mental status between seizures. After the second GTC seizure, the patient was administered a loading dose of lorazepam, but he remained unconscious, and remnant subtle motor components were observed. The patient’s caregiver informed us that the patient had a previous history of chronic alcoholism and was later diagnosed with diabetes as a post-alcoholic complication of pancreatitis. He had a history of alcohol-related seizures since his late 20s and was diagnosed in his early 30s with temporal lobe epilepsy accompanied by left hippocampal sclerosis. Despite treatment with valproate 600 mg bid, levetiracetam 1,000 mg bid, and vigabatrin 500 mg bid, he was not seizure-free. He was administered 1 mg/kg intravenous lorazepam, followed by a loading dose of 15 mg/kg intravenous fosphenytoin (15 mg/kg). He continued to receive his usual doses of valproate, levetiracetam, and vigabatrin, administered through a nasogastric feeding tube. After lorazepam loading, however, his valproate dose was changed to 450 mg intravenous every 8 hours, and levetiracetam dose to 1,500 mg bid, with no change in vigabatrin dose.

Although his convulsive seizures ceased, the patient did not recov-
Figure 3. Summary of the clinical condition of our patient, including EEG, body temperature, and history of treatment with antiepileptic drugs. CSE, convulsive status epilepticus; NCSE, non-convulsive status epilepticus; EEG, electroencephalography; PHT, phenytoin; LEV, levetiracetam; VPA, valproic acid; VGB, vigabatrin; SE, status epilepticus.

Figure 4. (A) Brain diffusion, (B) ADC, (C) fluid-attenuated inversion recovery (FLAIR), and (D) T1 enhanced MRI after 2 weeks of TH. Follow-up diffusion MRI and the ADC map showed normalization of the areas with initially reduced diffusion. FLAIR and T1 enhanced MRI showed no abnormal findings. ADC, apparent diffusion coefficient; FLAIR, fluid-attenuated inversion recovery; MRI, magnetic resonance imaging; TH, therapeutic hypothermia.

er consciousness. An initial EEG showed continuous periodic discharges in both posterior cerebral hemispheres (Fig. 1A). Diffusion-weighted magnetic resonance imaging (MRI) (Fig. 2A, B) showed symmetric diffusion-restricted lesions in the bilateral thalamus and parieto-occipital cortex with low-signal changes in apparent diffusion coefficient images (Fig. 2C, D). Several follow-up EEGs showed continuous, periodic single or poly-spikes and waves in the occipital areas (Fig. 1B). He was therefore administered loading doses of 600 mg pregabalin and 15 mg/kg phenobarbital on days of hospital stay 2 and 5, respectively (Fig. 3).

On day of hospital stay 6, the patient was started on continuous EEG monitoring and TH, without any additional medication to control seizures. At the time of starting TH, the patient was in a semi-coma and intubated state. TH at a target temperature of 35°C was administered using an Arctic Sun 5000® temperature management system (Medivance Arctic Sun System, Louisville, CO, USA), which uses external gel pads and circulating sterile water. The induction phase lasted 8 hours and the maintenance phase lasted 24 hours at 35°C. The patient was monitored with esophageal core probes, and normothermia was achieved by controlled rewarming over 8 hours. The patient was not administered any benzodiazepine drugs to control shiv-
erring or to maintain hypothermia. Rather, the patient was main-
tained by intravenous infusion of remifentanil hydrochloride, 2,600
mg/day acetaminophen and 11 mg/day buspirone.

EEG after induction of hypothermia to 35°C showed that the peri-
odic epileptiform discharges observed in this patient had decrease in
amplitude without a definite change in pattern (Fig. 1C). During
maintenance of hypothermia, EEG abnormalities gradually improved
and the patient regained consciousness in proportion to the improve-
ment in EEG (Fig. 1D, E). The patient was alert and his EEG normal-
ized on the second day following initiation of the rewarming
procedure. Follow-up MRI two weeks after TH showed normalization
of the areas with initially-reduced diffusion (Fig. 4).

This case report was approved by the institutional review boards
(IRB No. 2016-07-032) of the Sungkyunkwan University Samsung
Changwon Hospital. The patient and his caregivers provided consent
for publication of this report.

Discussion

After controlling generalized convulsive SE in our patient, he was
successfully treated with TH for refractory NCSE. The absence of ad-
titional sedative drugs, except for remifentanil hydrochloride, during
hypothermia maintenance indicates that TH itself may have con-
trolled his refractory NCSE. Although TH has been used as adjunct
therapy for patients with refractory epilepsy and SE,8,11,12 patients
with refractory convulsive SE may require sedative drugs to control
shivering and maintain hypothermia. Simultaneous administration of
sedative drugs limits the ability to evaluate the effects of hypo-
thermia alone. Because our patient did not experience convulsions,
he was not treated with any additional drugs that could have con-
tributed to improvements in EEG or clinical symptoms. Findings from
our patient suggest that mild TH is an appropriate treatment for re-
fractory NCSE and should be considered before administering gen-
eral anesthetics to control seizures.

The use of TH has increased worldwide, as this technique was
found to be advantageous in a variety of clinical settings. Examples
include unconscious adult patients with spontaneous circulation af-
after an out-of-hospital cardiac arrest, near-drowning, traumatic head
injury, hepatic encephalopathy, and stroke.1-5 Experimental studies
showed that TH has anti-epileptic and neuroprotective effects, lead-
ing to its clinical use to treat SE.6,7 The neurophysiological processes
involved in the anticonvulsant effects of hypothermia are still largely
unclear. However, there are theoretical reasons to recommend hypo-
thermia to treat SE. Hypothermia was first reported successful in
treating three children with SE under thiopental anesthesia.8 Case
series and case reports have shown that TH is safe and effective in
patients with RSE.8,13-15 Complications of TH include dysrhythmias,
infected, and primary coagulopathy, but these complications were
reported in patients subjected to severe hypothermia (< 32°C), or
when there were problems controlling temperature.3,4 Because mild
TH has no significant adverse effects, it is regarded as relatively safe
and easy to use.16 As our patient was thought to be less responsive to
conventional AEDs, mild TH was deemed a possible option for man-
aging NCSE prior to applying general anesthesia.

NCSE is defined as a state of ongoing seizures without con-
volusions, usually for longer than 30 minutes, associated with con-
tinuous epileptiform discharges on EEG.1 NCSE constitutes 20% to
25% of episodes of SE17,18 and occurs in approximately 8% of all co-
atose patients without clinical signs of seizure activity.19 NCSE has
been found to persist in 14% of patients after generalized convulsive
SE is controlled.20 Although there is no generally accepted clinical
definition of NCSE, early diagnosis is required to prevent potential
neurological damage. Diagnosis of NCSE requires a combination of
clinical and EEG features and particularly relies on EEG findings.21
Various EEG patterns have been described in NCSE, some of which
clearly denote NCSE. In contrast, other patterns, such as triphasic
waves, generalized periodic epileptiform discharges, and periodic lat-
eralized epileptiform discharges (PLEDs), may not be indicative of
NCSE.22 NCSE has been diagnosed in patients with PLEDs who meet
certain criteria that occur in comatose patients in the aftermath of
generalized tonic-clonic SE.23 Our patient was diagnosed with NCSE
based on these criteria. Additionally, because the EEG and mental
status of our patient gradually improved after hypothermia treat-
ment, this improvement may also be diagnostic of NCSE.

Some NCSE case series have reported high mortality and morbidity
rates,23-25 suggesting that aggressive therapy is warranted. A study of
100 patients with NCSE found that 18 had died,26 with mortality as-
associated with acute medical causes, impairment of mental status,
and development of acute complications.24 However, there is no con-
sensus on robust treatments for NCSE, unlike SE. Therefore, treat-
ment of NCSE depends on the type and cause.27 Because some pa-
tients with NCSE have shown more favorable outcomes than patients
with convulsive SE, caution should be used prior to further treatment
with intravenous anesthetics.28,29 Aggressive pharmacological treat-
ment appears to have a greater risk of morbidity and mortality than
continuing non-convulsive seizure activity.30 Additionally, non-phar-
macological treatments, such as hypothermia, vagal nerve stimulation, and electroconvulsive treatment, are recommended for patients with NCSE, although evidence is insufficient. Therefore, patients, such as ours, treated with various antiepileptic drugs with loading doses for a sufficient time to achieve the maximum effect, and who show no improvement in mental status or EEG may require intravenous anesthetics or other non-pharmacological treatments.

Because SE has heterogeneous causes, which greatly determine prognosis, the response of SE to treatment is difficult to evaluate using consistent and uniform criteria. However, studies have reported that TH has seizure cessation and reduction rates of 62.5% and 15%, respectively, indicating that rate of seizure response in patients with RSE is relatively high (77.5%). No correlation has been observed between target temperature and seizure response, indicating the need for additional studies on the effects, suitable target temperatures, and complications of TH. Furthermore, due to the development of TH tools with few serious adverse effects, the use of mild TH in patients with refractory NCSE should be considered in lieu of intravenous anesthetics.

**Conflicts of interest**

The authors have no relevant conflict of interest to report.

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