Clinical Spectrum and Prognostic Factors of Acute Necrotizing Encephalopathy in Children

This study was conducted to investigate the etiology, the clinical characteristics and prognosis of acute necrotizing encephalopathy (ANE) in Korean children. Six children (1 yr to 7 yr) patients with ANE were enrolled. They were diagnosed by clinical and radiological characteristics and their clinical data were retrospectively analyzed. In a search of clinically plausible causes, brain MRI in all patients, mitochondrial DNA studies for mitochondrial encephalomyopathy, lactic acidosis, and stroke-like episodes (MELAS) and myoclonus epilepsy and ragged red fibers (MERRF) in four patients, and genomic typing on HLA DRB1/HLA DQB genes in three patients were performed. All had precedent illnesses and the main initial symptoms included mental change (83%), seizures (50%), and focal deficits (50%). MRI revealed increased T2 signal density in the bilateral thalami and/or the brainstem in all patients. Mitochondrial DNA studies for MELAS and MERRF were negative in those children and HLA-DRB1*1401, HLA-DRB3*0202, and HLA-DQB1*0502 seemed to be significant. A high dose steroid was given to all patients, which seemed to be partly effective except for 2 patients. In conclusion, ANE is relatively rare, but can result in serious neurological complication in children. Early detection and appropriate treatment may lead to a better neurological outcome.

Key Words : Acute Necrotizing Encephalopathy; Mitochondrial; MELAS Syndrome; MERRF Syndrome; HLA-DR Antigens; HLA-DQ Antigens

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INTRODUCTION

Acute necrotizing encephalopathy (ANE) is a unique acute encephalopathy that predominantly affects infants and young children (1). It is characterized by the features of acute encephalopathy such as seizures and rapid alteration of consciousness after a nonspecific viral illness. Since Mizuguchi et al. proposed this condition as a new entity in 1995 (2), it has occasionally been reported in both Asian and Western countries. An increased level of serum aminotransferase activity and cerebrospinal fluid protein, are the most common abnormalities (1-4). Diagnosis is made mainly by the characteristic findings of computed tomography or magnetic resonance imaging (MRI), which typically show symmetric lesions in the thalami, with variable involvement of the white matter, basal ganglia, brainstem, and cerebellum (1, 4, 5). Regardless of the treatment, the prognosis of ANE varies widely from complete recovery to death (1-6). With the widespread use of MRI, this unique condition is becoming more familiar.

However, the etiology, pathogenesis, guidelines of treatments, or prognostic factors still remain unclear. In this report, we described 6 children with ANE to elucidate the genetic propensity, the clinical/neuroradiological characteristics, the response to medical treatment and prognostic factors.

MATERIALS AND METHODS

A total of 6 children with ANE (aged 1 yr to 7 yr at onset, 3 males and 3 females) was evaluated. The patients were admitted to Kyungpook National University Hospital, Daegu, Korea, from 1999 through 2007. Diagnosis was made mainly by clinical and radiological characteristics. Serological tests or polymerase chain reaction for certain viruses such as Herpes simplex virus, cytomegalovirus, Epstein-Barr virus, influenza virus, and mycoplasma were done in 5 out of them. Brain MRIs were taken in all patients at the time of initial presentation. In a search of other clinically plausible causes, mito-
mitochondrial DNA studies for mitochondrial encephalomyopathy and lactic acidosis and stroke-like episode (MELAS) and myoclonus epilepsy and ragged red fibers (MERRF), particularly point mutation at nt 3243, nt 3271, nt 3252, nt 8344, nt 8356, etc. in the tRNA were screened in four patients. In addition, genomic typing was performed on HLA DRB/HLA DQB genes using polymerase chain reaction (PCR)-SSOP/SSP techniques with Gel immunoelectrophoresis in three patients. This clinical research was approved by Internal Review Board (74005-1540).

**RESULTS**

A total of six children was enrolled in the study. Past medical histories were uneventful in all patients and they had normal developmental milestones. The family histories were also unremarkable. No patients were exposed to any drugs or chemical substances known to cause toxic encephalopathies.

**Clinical features of the subjects**

Clinical characteristics of the subjects are summarized in Table 1. All had precedent illnesses and five out of them (83%) had fever. The initial neurological symptoms include mental change in five patients (83%), seizures (50%) and focal neurological signs (50%).

**Laboratory findings**

Table 2 shows laboratory findings of the subjects. Increased level of serum aminotransferase and lactic dehydrogenase activities may indicate hepatic dysfunction, but their levels varied highly from case to case. Four patients (67%) showed elevation of serum aminotransferase, which increased sharply to 7,000 U/L in 2 out of them. Severe cases revealed laboratory findings compatible with disseminated intravascular coagulation but the clinical symptoms were almost absent. In addition, serum ammonia levels were normal. Serological tests or polymerase chain reaction for certain viruses such as *Herpes simplex* virus, cytomegalovirus, Epstein-Barr virus, influenza virus, and mycoplasma were unremarkable. Mitochondrial DNA studies for MELAS and MERRF were done in 4 patients, and none of them showed any mutations. Genomic typing of HLA DRB/HLA DQB genes in three patients (patient 3, 4, and 5) using PCR-SSOP/SSP techniques with gel immunoelectrophoresis showed HLA-DRB1*1401, HLA-DRB3*0202, and HLA-DQB1*0502 which were possibly significant (Table 3).

Cerebrospinal fluid study revealed normal protein levels in all patients and mild pleocytosis in one patient (patient 4). *Herpes* IgM antibody, oligoclonal band and myelin basic protein were unremarkable in all subjects.

**Radiological findings**

As shown in Fig. 1, brain MRI revealed increased signal

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**Table 1. Clinical features of the subjects with ANE**

| Patient No. | Age (month) | Sex | Precedent illnesses | Initial presentation | Sites of involvement |
|-------------|-------------|-----|---------------------|---------------------|---------------------|
| 1           | 93          | F   | NSFI                | Diplopia            | BG, T, P, C         |
| 2           | 19          | M   | URI & diarrhea      | Stupor/Rigidity     | TT                  |
| 3           | 40          | F   | NSFI                | Stupor/Seizure*     | BG, T               |
| 4           | 68          | M   | NSFI                | Drowsiness/Gait disturbance | TT                  |
| 5           | 73          | F   | NSFI                | Stupor/Seizure      | T, P, C             |
| 6           | 88          | M   | URI                 | Stupor/Seizure      | TT                  |

*Status epilepticus.
ANE, acute necrotizing encephalopathy; NSFI, non specific febrile illness; URI, upper respiratory tract infection; BG, Basal ganglia; T, Thalamus; TT, Thalamotegmentum; P, Pons; C, Cerebellum.

**Table 2. Laboratory findings of the subjects with ANE**

| Patient No. | AST/ALT (U/L) | Glucose (mg/dL) | Ammonia (µg/dL) | Cerebrospinal fluid | mtDNA mutations |
|-------------|---------------|-----------------|-----------------|---------------------|----------------|
| 1           | 26/9          | 145             | 14              | 0                   | 29 83 - - - |
| 2           | 114/42        | 85              | 32              | 0                   | 39 76 - - - |
| 3           | 7,000/4354    | 96              | 48              | 0                   | 30 65 ND ND ND |
| 4           | 34/15         | 85              | 24              | 66                  | 40 61 ND ND ND |
| 5           | 128/44        | 135             | 24              | 0                   | 43 90 ND ND ND |
| 6           | 7,225/2,910   | 108             | 56              | 0                   | 46 61 ND ND ND |

ANE, acute necrotizing encephalopathy; AST, aspartate aminotransferase; ALT, alanine aminotransferase; ND, not done; MBP, myelin basic protein; mtDNA, mitochondrial DNA.
density on T2-weighted imaging in the bilateral thalami and brain stem in almost all patients. Two out of them had lesions in cerebellum (Table 1). The findings were consistent with a unique pattern of ANE. Follow-up MRI was done in 2 weeks to 7 months. In one patient, brain findings had completely resolved, However, in 4 patients of 5 followed-up, a certain degree of sequelae were found (Table 4).

Treatment and outcome

With respect to the treatment, steroids have been used in two regimens: intravenous dexamethasone (patient 2, 3, 6) or methylprednisolone pulse therapy (patient 1, 4, 5). In the dexamethasone group, 1 mg/kg/day of dexamethasone was administered in 4 divided doses for at least 5 days. In the methylprednisolone group, 30 mg/kg/day of methylprednisolone was administered for 5 days. As shown in Table 4, a patient (patient 1) showed excellent outcomes without any neurological sequelae at 6 months after the illness. The other three patients (patient 2, 3, 4) showed a relatively good to fair outcome. Even though they had initial weakness, spasticity on extremities or memory disturbance, their symptoms improved remarkably. Two patients developed epilepsy later on and required prophylactic antiepileptic drugs. Even so, they remained seizure free for more than years, and eventually were taken off antiepileptic drugs without any sequelae. A patient (patient 6) died of cardiorespiratory compromise.

Table 3. HLA-DRB and DQB alleles of 3 patients with ANE

| Patient No. | DRB1    | DRB3   | DQB1       |
|-------------|---------|--------|------------|
| 1           | 0701 1401 | 0202   | 0303 0502  |
| 2           | 1101 1401 | 0202   | 0301 0502  |
| 3           | 0101 1302 | 0301   | 0501 0609  |

ANE, acute necrotizing encephalopathy.

Table 4. Treatment and its outcome in the subjects with ANE

| Patient No. | Treatment                      | Follow-up MRI | Neurological outcome at 6 months |
|-------------|--------------------------------|---------------|---------------------------------|
| 1           | Methylprednisolone pulse       | Complete resolution | Excellent                      |
| 2           | Dexamethasone                  | 50-75% resolution | Fair                           |
| 3           | Dexamethasone                  | 50-75% resolution | Fair                           |
| 4           | Methylprednisolone pulse       | >75% resolution  | Good                           |
| 5           | Methylprednisolone pulse       | <50% resolution  | Poor                           |
| 6           | Dexamethasone                  | Not done       | Poor                           |

*Definition of outcome.

EXCELLANT, complete resolution; Good, almost complete resolution or minimal degree of neurological sequelae (mobile, almost no cognitive/social/emotional impairment); Fair, moderate degree of neurological sequelae (mobile with difficulty, mild to moderate cognitive/social/emotional impairment); Poor, severe degree of neurological sequelae (immobile, severe cognitive/social/emotional impairment).

DISCUSSION

ANE was proposed as a novel disease entity by Mizuguchi et al. in 1995 (2). Patients with ANE manifest fulminating neurologic deterioration with preceding non-specific febrile illness and frequently undergo intractable convulsions. Serious neurological signs such as decorticate, decerebrate posturing or long tract signs may appear. Its mortality is considered to reach as high as 30% (1, 2). Mizuguchi et al. proposed the following diagnostic criteria for acute necrotizing encephalopathy: 1) acute encephalopathy following a viral febrile disease and rapid deterioration in the level of consciousness, convulsion; 2) increased cerebrospinal fluid (CSF) protein without CSF pleocytosis; 3) CT or MRI findings for symmetric, multifocal brain lesions involving bilateral thalami, cerebral periventricular white matter, internal capsule, putamen, upper brain stem tegmentum and cerebellar medulla without involvement of other CNS regions; 4) elevation of serum aminotransferase of variable degrees without hyperammonemia; 5) exclu-
tion of other resembling diseases such as overwhelming bacterial and viral infections, fulminant hepatitis, Reye syndrome, Leigh encephalopathy and related mitochondrial cytopathies, acute disseminated encephalomyelitis or other types of encephalitis, vasculitis, cerebral infarction, and so on (1). Despite the fact that Mizuguchi et al. proposed the diagnostic criteria of ANE, atypical or milder cases have been reported. Yoshi-kawa et al. suggested clinical diversity in ANE. They experienced 2 atypical cases that had selective reversible thalamic involvement and milder clinical manifestations, and that ended up in complete recovery without any sequelae (4). Patients with normal cerebrospinal fluid protein and low levels of transaminase, asymmetric thalamic involvement and no lesion in the brain stem tend to recover well (6). Considering the fact that ANE is a rare neurological condition, we experienced a variety of different cases. They met the diagnostic criteria proposed by Mizuguchi et al. (2).

ANE is known to be one of the neurological complications of viral infections, such as influenza A and B, human herpes-virus 6, varicella zoster or mycoplasma infection (7-14). In addition, it is reportedly associated with the measles virus in an animal study (15). Serological tests or polymerase chain reaction for certain viruses, and mycoplasma were unremarkable in our study. As shown in Table 2, the subjects showed various results in laboratory findings. Interestingly CSF protein was within the normal range in all cases, but all had typical thalamic involvement on the images. In an attempt to exclude other resembling encephalopathies, Herpes IgM, oligoclonal band and myelin basic proteins in CSF were examined and shown unremarkable findings in most cases. Mutations of the mitochondrial DNA were screened to rule out mitochondrial encephalopathies in four out of subjects, which all came back negative (16, 17). Considering the fact of their similarity to mitochondrial disorders on brain MRI, we believe that mitochondrial dysfunction is partly related in this condition, whether primary or not.

As is already known, ANE is a relatively uncommon form of acute encephalopathy especially in East Asia, although European and American cases have been reported. It is thought to have some racial or geographic predilection (1, 6). In this study the HLA typing of 3 patients with ANE were evaluated to see genetic propensity. Genomic typing was performed on their HLA DRB/HLA DQB genes and HLA-DRB1*1401, HLA-DRB3*0202, and HLA-DQB1*0502 were found to be significant as compared with known data of HLA alleles in Korean (Table 3) (18). This is the first study using genomic typing of HLA DRB/HLA DQB genes that has never been done so far and the results may provide the immunogenetic background of ANE. However, further studies are needed to elucidate the condition.

A postmortem examination of the central nervous system could not be conducted, but bilateral thalamic necrosis is the histological hallmark of acute necrotizing encephalopathy. Selective vulnerability of the thalami seems to be a determin-

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