Clinically based score predicting cryptogenic NORSE at the early stage of status epilepticus

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Abstract

Objective
To determine whether a clinically based score predicts cryptogenic new-onset refractory status epilepticus (C-NORSE) at the early stage of status epilepticus (SE) with prominent motor symptoms (SE-M) of unclear etiology.

Methods
The score (range 0–6) included 6 clinical features: highly refractoriness to antiseizure drugs, previously healthy individual, presence of prodromal fever, absence of prodromal psychobehavioral or memory alterations, absence of dyskinesias, and symmetric brain MRI abnormalities (the first 2 mandatory). We retrospectively assessed the usefulness of a high scale score (≥5) in predicting C-NORSE in 83 patients with SE-M of unclear etiology, who underwent testing for neuronal surface antibodies (NS-Abs) between January 2007, and December 2019.

Results
Thirty-one (37.3%) patients had a high score. Patients with a high score had more frequent prodromal fever (28/31 vs 24/52), mechanical ventilatory support (31/31 vs 36/52), and symmetric MRI abnormalities (26/31 vs 12/52), had less frequent involuntary movements (2/31 vs 30/52), and had absent prodromal psychobehavioral alterations (0/31 vs 27/52), CSF oligoclonal band detection (0/27 vs 11/38), tumor association (0/31 vs 13/52), or NS-Abs (0/31 vs 29/52) than those with a low score (<5). Thirty-three patients (median age, 27 years; 18 [54.5%] female) were finally regarded as C-NORSE. The sensitivity and specificity of a high score for predicting C-NORSE were 93.9% (95% CI 0.87–0.94) and 100% (95% CI 0.95–1.00), respectively.

Conclusions
Patients with a high score in the indicated scale are more likely to have C-NORSE, making it a useful diagnostic tool at the early stage of SE-M before antibody test results become available.
Glossary

AE = autoimmune encephalitis; AMPAR = α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor; AQP4 = aquaporin-4; ASD = antiseizure drug; CBA = cell-based assay; C-NORSE = cryptogenic NORSE; DWI = diffusion-weighted image; FC = febrile convolution; FIRES = febrile infection-related epilepsy syndrome; FLAIR = fluid-attenuated inversion recovery; GABAαR = γ-aminobutyric acid A receptor; GABAβR = γ-aminobutyric acid B receptor; IgG = immunoglobulin G; IL-6 = interleukin-6; ILAE = International League Against Epilepsy; LGI1 = leucine-rich glioma-inactivated 1; MOG = myelin oligodendrocyte glycoprotein; NCSE = nonconvulsive SE; NMDAR = NMDA receptor; NORSE = new-onset refractory status epilepticus; NS-Abs = neuronal surface antibodies; OCB = oligoclonal band; PMH = past medical history; SE = status epilepticus; SE-M = SE with prominent motor symptoms; WBC = white blood cell.

New-onset refractory status epilepticus (NORSE) is a severe neurologic emergency condition characterized by refractory status epilepticus (SE) without readily identifiable cause in otherwise healthy individuals. The term NORSE is now defined as a clinical presentation, not a specific diagnosis.

According to the consensus definition, NORSE includes patients with viral, paraneoplastic, or autoimmune etiologies; however, it is crucial in clinical practice to differentiate C-NORSE from secondary NORSE with neuronal surface antibodies (NS-Abs) or classical paraneoplastic antineuronal antibodies because treatment strategy and outcome could be different. A large cohort study reported that a half of 130 patients with NORSE remained cryptogenic, but 37% were immune mediated; among those, the most common etiology was anti-NMDA receptor (NMDAR) encephalitis.

Although antibody tests are important to determine the etiology, in an emergency condition, it is often difficult to get the antibody test results in appropriate time. Therefore, we previously developed a clinically based score (range 0–6) based on 6 clinical features to predict C-NORSE at the early stage of convulsive SE, which is currently classified into SE with prominent motor symptoms (SE-M) according to the 2015 International League Against Epilepsy (ILAE) criteria for SE. However, the scale score has not been validated yet.

Here we report the sensitivity and specificity of the high scale score (≥5) in predicting C-NORSE at the early stage of SE-M of unclear etiology (before NS-Ab test results are known).

Methods

Patients selection and antibody assays (study profile)

We first reviewed the clinical information of 180 patients with seizures of unclear etiology on admission or early stage of seizures, in whom NS-Abs were examined to investigate potential immune-mediated etiologies between January 1, 2007, and December 31, 2019 (figure 1). These patients were admitted to Kitasato University Hospital or other associated hospitals between January 1, 1999, and December 31, 2019; in 7 patients who were admitted before January 1, 2007, archived serum/CSF samples obtained at onset of disease were used for antibody assays.

Then, we selected 129 patients who fulfilled the 2015 ILAE criteria for SE. Of those, 46 patients with nonconvulsive SE (NCSE) were excluded because the scale score was originally developed to estimate antibody status in patients with convulsive SE. In this study, we included all patients who developed SE-M regardless of refractoriness to conventional antiseizure drug (ASD) treatment. We assessed the sensitivity and specificity of the high scale score (≥5) in 83 patients with SE-M of unclear etiology during the early stage.

NS-Abs were measured at the laboratory of Josep Dalmau (University of Barcelona) using both a rat brain immunohistochemistry and cell-based assay (CBA)7–13; they included antibodies against the NMDAR, α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor (AMPA R), γ-aminobutyric acid B receptor (GABAβR), γ-aminobutyric acid A receptor (GABAαR), metabotropic glutamate receptor 5, dipeptidyl peptidase-like protein 6, contactin-associated protein-like 2, leucine-rich glioma-inactivated 1 (LG I1), and neurexin 3. Both serum and CSF were examined in all patients except 4 (only CSF [n = 2] or serum [n = 2] was available). In addition to NS-Abs, myelin oligodendrocyte glycoprotein (MOG) and aquaporin-4 (AQP4) antibodies were examined with CBA in patients with overlapping encephalitis and demyelinating syndrome. Antibodies against classical paraneoplastic intracellular antigens (CV2/CRMP5, Ma2, Ri, Yo, Hu, GAD65, and amphiphysin) were measured in serum at Kitasato University with EUROLINE (Euroimmun AG) in patients when associated tumor was suspected or those with NORSE criteria.

Criteria for C-NORSE

Although C-NORSE is not a specific diagnosis, patients were classified into C-NORSE as a subgroup of cryptogenic epileptic syndrome in this study if those fulfilled the following 3 criteria: (1) new-onset refractory SE in previously healthy individual, (2) refractoriness to conventional ASD treatment, and (3) no etiology identified throughout the course of the disease. If the etiology of SE was identified, patients were diagnosed with etiology-based specific diagnosis (e.g., anti-NMDAR encephalitis and anti-LGI1 encephalitis). SE was
considered as refractory when it continued longer than 60 minutes despite adequate administration of benzodiazepines and adequate loading of standard IV ASDs.2,6,15,16 The etiology of NORSE was extensively investigated with CSF examination, malignancy survey, and serologic testing including autoantibodies against neuronal surface and classical para-neoplastic intracellular antigens.

C-NORSE score
C-NORSE score is a clinically based score (range 0–6) based on the following 6 clinical features5 usually obtained within 14 days after admission in general hospital: (1) NORSE highly resistant to conventional ASD treatment, (2) previously healthy individual before the onset of SE, (3) presence of prodromal high fever of unknown origin before the onset of SE, (4) absence of prodromal psychobehavioral or memory alterations before the onset of SE, (5) absence of sustained orofacial-limb dyskinesias despite a profoundly decreased level of consciousness, and (6) symmetric brain MRI abnormalities (table 1).

In the criteria, we previously defined that each feature represents 1 point, but the first 2 clinical features are mandatory.5 Accordingly, if either the first or second feature is absent, the patient is scored 0. We applied 2015 ILAE criteria for SE6 to include patients with SE-M, and all patients underwent EEG and MRI repeatedly during their hospitalization. However, only patients who had electroencephalographic correlates (such as spikes and waves or periodic discharges that explain prominent motor symptoms) were regarded to meet the first clinical feature of the score. Accordingly, patients without apparent electroencephalographic correlates despite convulsive SE or epilepsia partialis continua were scored 0. Symmetric brain MRI abnormalities imply relatively symmetric increased diffusion-weighted image (DWI) or T2/fluid-attenuated inversion recovery (FLAIR) signals in the hippocampus, fimbria, amygdala, claustrum, insula, or perisylvian opercular cortex; these changes may not be seen at the onset of SE-M but often subsequently develop associated with persistent seizure activity.5

Clinical assessments
We assessed the clinical features between patients with a high scale score (≥5) and those with a low scale score (≤4), including sex, age at onset of SE-M, prodromal fever, prodromal psychobehavioral or memory alterations, involuntary movements, mechanical ventilatory support, CSF and MRI findings, and presence of tumor. We reviewed the final diagnosis of these patients after extensive workup and finally determined the sensitivity and specificity of the indicated high scale score. In this study, to focus on the C-NORSE score, we did not assess the efficacy of treatment, such as immunotherapy, or long-term outcome in these patients.
Standard protocol approvals, registrations, and patient consents
The study was approved by Institutional Review Boards of Kitasato University (B18-193). Written informed consent was obtained from the patients or their family members. Information on symptoms, CSF, MRI, EEG, and treatments was obtained from the authors or referring physicians.

Statistical analysis
The Fisher exact test was performed for comparison of categorical variables, and the Mann-Whitney test was used for continuous variables. The statistical significance was set at $p < 0.05$. The sensitivity and specificity of the high C-NORSE score were determined with 2-way contingency table analysis. We used JMP, version 14 (SAS Institute Inc.), for statistical analyses.

Data availability
Any data not published within the article are available and will be shared anonymously by request from any qualified investigator.

Results
Clinical features in patients with a high score and those with a low score
Of 83 patients, 31 (37.3%) had a high score (5–6); 17 patients (54.8%) were female; median age at symptom onset was 27 years (range 5–73 years) (table 2). The remaining 52 patients (62.7%) had a low score (0–4); 37 patients (71.2%) were female; median age at symptom onset was 25 years (range 10–79 years). Other clinical information is shown in table 2. There was no difference between patients with a high score and low score in female sex and median age at onset. However, patients with a high score had more frequent prodromal fever (28/31 vs 24/52), mechanical ventilatory support (31/31 vs 36/52), and symmetric DWI or T2/FLAIR hyperintensities (26/31 vs 12/52) than those with a low score. By contrast, they had less frequent involuntary movements (2/31 vs 30/52) and absent prodromal psychobehavioral alterations (0/31 vs 27/52), CSF oligoclonal band (OCB) detection (0/27 vs 11/38), tumor association (0/31 vs 13/52), or NS-Abs (0/31 vs 29/52) than those with a low score. There was no difference in prodromal headache before the onset of SE, CSF pleocytosis, white blood cell (WBC) counts in CSF, CSF protein levels, or elevated immunoglobulin G (IgG) index.

Final diagnosis
Of 83 patients with 2015 ILAE criteria for SE-M of unclear etiology on admission or early stage of SE, 29 (34.9%) patients were positive for NS-Abs, NMDAR in 26 patients (1 with concurrent AQP4 and 1 with MOG), LGI1 in 1, GABAAR in 1, and unknown antigens (not characterized yet) in 1. No AMPAR or GABAAR antibodies were identified. All antibody-positive patients had a low C-NORSE score: 24 patients had 0, and 5 patients had 1. The remaining 54 patients (65.1%) were negative for NS-Abs; 21 patients were diagnosed with miscellaneous disorders or syndrome including possible autoimmune encephalitis (AE)17 (n = 11), autoantibody-negative but probable AE17 (n = 5), antibody-negative autoimmune limbic encephalitis17 (n = 1), encephalitis associated with systemic lupus erythematosus (n = 2), and nonautoimmune neurologic disorders (n = 2). The remaining 33 patients were finally regarded as C-NORSE based on the above criteria (figure 1).

Clinical features of C-NORSE
Eighteen of 33 patients (54.5%) were female; median age at onset was 27 years (range 5–73 years). Thirty-one patients

Table 1 Components of the C-NORSE score

<table>
<thead>
<tr>
<th>Clinical feature</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. New-onset refractory SE highly resistant to conventional ASD treatment</td>
<td>1</td>
</tr>
<tr>
<td>2. Previously healthy individual before the onset of SE</td>
<td>1</td>
</tr>
<tr>
<td>3. Presence of prodromal high fever of unknown origin before the onset of SE</td>
<td>1</td>
</tr>
<tr>
<td>4. Absence of prodromal psychobehavioral or memory alterations before the onset of SE</td>
<td>1</td>
</tr>
<tr>
<td>5. Absence of sustained orofacial-limb dyskinesias despite profoundly decreased level of consciousness</td>
<td>1</td>
</tr>
<tr>
<td>6. Symmetric DWI or T2/FLAIR hyperintensities</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>6</td>
</tr>
</tbody>
</table>

Abbreviations: ASD = antiseizure drug; C-NORSE = cryptogenic new-onset refractory status epilepticus; DWI = diffusion-weighted image; FLAIR = fluid-attenuated inversion recovery; SE = status epilepticus.

C-NORSE score is a clinically based score (range 0–6) based on the above 6 clinical features (slightly modified from the original one5). In the criteria, each feature represents 1 point, but the first 2 clinical features are mandatory. If either the first or second feature is absent, the patient is scored 0. The sixth feature means relatively symmetric increased DWI or T2/FLAIR signals in the hippocampus, fimbria, amygdala, claustrum, insula or perisylvian opercular cortex: these changes may not be seen at the onset of SE but often subsequently develop associated with persistent seizure activity. The C-NORSE score should be used only to predict C-NORSE at the early state of SE-M of unclear etiology before antibody test results become available, but it should not be used to make a diagnosis (see Text).

4 Neurology Neuroimmunology & Neuroinflammation | Volume 7, Number 5 | September 2020 Neurology.org/NN
had a high score; 23 patients had 6, and 8 patients had 5, but 2 patients had a low score (both 4). Of interest, 7 of the 33 patients (21.2%) had a past medical history (PMH) of febrile convulsion (FC), family history of FC or epilepsy, or both; 3 patients had a PMH of FC (one of them had a family history of FC); 4 patients had no PMH of FC but had a family history of FC (n = 1) or epilepsy (n = 3).

Prodromal symptoms developed before the onset of SE in 31 of 33 patients (93.9%), fever in 28 of 33 patients (84.8%), and headache in 15 of 31 patients (2 unknown). Only 1 patient (3.0%) developed psychobehavioral alterations before the onset of SE, whereas 3 patients (9.1%) showed involuntary movements during the course of the disease, but only 1 patient developed sustained dyskinesias mimicking orofacial-limb dyskinesias. All patients required mechanical ventilatory support due to refractory SE.

Prodromal symptoms mean symptoms/signs that developed before the onset of status epilepticus.

Abbreviations: DWI = diffusion-weighted image; FLAIR = fluid-attenuated inversion recovery; IgG = immunoglobulin G; NS-Ab = neuronal surface antibody; OCB = oligoclonal band; WBC = white blood cell.

Table 2 Clinical features in patients with a high score and those with a low score

<table>
<thead>
<tr>
<th>Feature</th>
<th>High score (n = 31)</th>
<th>Low score (n = 52)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female sex</td>
<td>17 (54.8%)</td>
<td>37 (71.2%)</td>
<td>0.1574</td>
</tr>
<tr>
<td>Median age at symptom onset (y)</td>
<td>27 (5–73)</td>
<td>25 (10–79)</td>
<td>0.4916</td>
</tr>
<tr>
<td>Prodromal fever of unknown origina</td>
<td>28 (90.3%)</td>
<td>24 (46.2%)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Prodromal headachea</td>
<td>15/29 (51.7%)</td>
<td>23 (44.2%)</td>
<td>0.6431</td>
</tr>
<tr>
<td>Prodromal psychobehavioral or memory alterationsa</td>
<td>0 (0.0%)</td>
<td>27 (51.9%)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Involuntary movements</td>
<td>2 (6.5%)</td>
<td>30 (57.7%)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Mechanical ventilatory support</td>
<td>31 (100.0%)</td>
<td>36 (69.2%)</td>
<td>0.0030</td>
</tr>
<tr>
<td>Symmetric DWI or T2/FLAIR hyperintensities</td>
<td>26 (83.9%)</td>
<td>12 (23.1%)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Tumor association</td>
<td>0 (0.0%)</td>
<td>13 (25.0%)</td>
<td>0.0014</td>
</tr>
<tr>
<td>CSF pleocytosis (&gt;5 WBCs/μL)</td>
<td>22 (71.0%)</td>
<td>40 (76.9%)</td>
<td>0.6063</td>
</tr>
<tr>
<td>Median CSF WBC counts (WBCs/μL)</td>
<td>9 (1–224)</td>
<td>17 (0–279)</td>
<td>0.2188</td>
</tr>
<tr>
<td>Median CSF protein (mg/dL)</td>
<td>41 (13–129)</td>
<td>35 (14–534)</td>
<td>0.2685</td>
</tr>
<tr>
<td>CSF OCB detection</td>
<td>0/27 (0.0%)</td>
<td>11/38 (28.9%)</td>
<td>0.0017</td>
</tr>
<tr>
<td>Elevated IgG index (≥0.74)</td>
<td>2/23 (8.7%)</td>
<td>10/38 (26.3%)</td>
<td>0.1113</td>
</tr>
<tr>
<td>NS-Ab detection</td>
<td>0 (0.0%)</td>
<td>29 (55.8%)</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

*Prodromal symptoms mean symptoms/signs that developed before the onset of status epilepticus.

Abbreviations: DWI = diffusion-weighted image; FLAIR = fluid-attenuated inversion recovery; IgG = immunoglobulin G; NS-Ab = neuronal surface antibody; OCB = oligoclonal band; WBC = white blood cell.

NS-Abs were not detected in either serum or CSF. Classical paraneoplastic antineuronal antibodies measured in serum in 28 patients were negative but not measured in 5 (no serum was available for examination). CSF examination revealed a median of 9 WBCs/μL (range 0–224 WBCs/μL) and a median protein level of 41 mg/dL (range 13–129 mg/dL). No CSF-restricted OCBs were detected in 29 examined patients, whereas the IgG index was elevated in 2 of 25 examined patients (8.0%). Ten patients (30.3%) had no pleocytosis (>5 WBCs/μL). Initial brain MRI was unremarkable in 15 patients (45.5%), but follow-up MRIs showed abnormal findings in 30 patients (90.9%); in 27 patients (81.8%), brain MRIs showed symmetric DWI or T2/FLAIR hyperintensities in the medial temporal lobes, basal ganglia, fimbria, claustrum, or perisylvian opercular cortex (figure 2). None of these patients had a tumor identified during the course of the disease.

The sensitivity and specificity of the high C-NORSE score

The sensitivity and specificity of the high score (≥5) for predicting C-NORSE were 93.9% (95% CI 0.87–0.94) and 100% (95% CI 0.95–1.00), respectively.

Discussion

This study shows that (1) patients with the high score are more likely to have C-NORSE, (2) the clinically based score C-NORSE score has high sensitivity and specificity for predicting the C-NORSE, and (3) patients with C-NORSE had distinctive clinical features.

In clinical practice, it is important to estimate antibody status in patients with SE of unclear etiology and identify patients with C-NORSE as early as possible because patients with C-NORSE are usually less responsive to first-line immunotherapy and more likely to have poor long-term outcome with cognitive deficits and refractory partial seizures.
This scale score was originally developed based on our previous preliminary study\textsuperscript{5} that compared the clinical features of 11 adult patients with C-NORSE (aged ≥17 years) with those of 32 patients with anti-NMDAR encephalitis. We previously reported that the C-NORSE score was higher in patients with C-NORSE than those with anti-NMDAR encephalitis; however, the sensitivity and specificity were not determined. After that, we had recruited additional patients since September 2016. In the meantime, the international consensus definition of NORSE was proposed in 2018\textsuperscript{3}; hence, the concept of C-NORSE was much more clearly defined than before. In this study, we adopted the concept of C-NORSE and included pediatric cases as well as newly identified adult cases. Accordingly, we increased the number of patients with C-NORSE from 11 to 33.

In this study, we assessed the sensitivity and specificity of the high score (≥5) in 83 patients with SE-M. In this cohort, the sensitivity and specificity for predicting C-NORSE were 93.9% and 100%, respectively, making it a useful diagnostic tool at the early stage of SE-M of unclear etiology before antibody test results become available.

C-NORSE is a devastating epileptic syndrome of unknown causes, probably of diverse etiologies\textsuperscript{1–5} including autoimmunity, neuroinflammation, or individual susceptibility to seizure. This study highlighted distinctive clinical features of C-NORSE phenotypically different from antibody-positive AE, such as anti-NMDAR, anti-LGI1, or anti-GABA\(_\alpha\)R encephalitis. Patients with C-NORSE often present with high fever of unknown cause, followed by sudden onset of mainly convulsive seizures, leading to refractory SE (occasionally super-refractory SE) requiring a mechanical ventilatory support and continuous infusion of sedative drugs. Early brain MRI is often normal or may show symmetric DWI or T2/FLAIR hyperintensities in the medial temporal lobes,\textsuperscript{5} mimicking autoimmune limbic encephalitis. CSF examination often shows nonspecific mild pleocytosis; however, none of these patients had CSF-restricted OCBs, and the IgG index was not elevated in most of them. Of interest, prodromal psychobehavioral or memory alterations usually did not develop before the onset of SE or decreased level of consciousness. This is highly contrast to those with anti-NMDAR encephalitis\textsuperscript{5,6,17–19} or autoimmune limbic encephalitis,\textsuperscript{17} in

\textbf{Figure 2 Brain MRIs finding obtained from 3 patients with C-NORSE}

\begin{figure}
\centering
\includegraphics[width=\textwidth]{Brain_MRIs.png}
\caption{Brain MRIs show symmetric increased DWI or FLAIR signals in the amygdala, hippocampus, fimbria, claustrum, insula, and fronto-temporal cortex. Basal ganglia and perisylvian opercular cortex are also involved in patients with C-NORSE (not shown). (A) A 37-year-old man; (B) a 49-year-old woman; (C and D) a 39-year-old woman; (A–C) FLAIR, (D) DWI. C-NORSE = cryptogenic new-onset refractory status epilepticus; DWI = diffusion-weighted image; FLAIR = fluid-attenuated inversion recovery.}
\end{figure}
whom these symptoms usually develop in the early course of the disease, and often predominant. Thus, the lack of prodromal psychobehavioral or memory alterations is an important feature in discrimination of C-NORSE from anti-NMDAR encephalitis or limbic encephalitis. The follow-up brain MRIs often show symmetric brain lesions involving the hippocampus, amygdala, limbia, clausstrum, basal ganglia, insular cortex, and perisylvian opercular cortex presumably associated with persistent seizure activity.\textsuperscript{5,20} \text{Such neuroimaging pattern is quite different from autoimmune limbic encephalitis with highly restricted to bilateral medial temporal lobes\textsuperscript{17} or anti-GABAaR encephalitis with multiple corticostriatal lesions.\textsuperscript{13,23} Involuntary movements may develop in patients with C-NORSE due to secondary basal ganglia lesions, but not like NMDAR-associated orofacial-limb dyskinesias\textsuperscript{18} or movement disorders,\textsuperscript{24} or LGI1-associated faciobrachial dystonic seizures.\textsuperscript{25}}

The etiology of C-NORSE remains unknown.\textsuperscript{1–5} It is also controversial whether it is of autoimmune origin.\textsuperscript{5} One might argue that C-NORSE is an epileptic syndrome and should not be confused with AE; randomized controlled trial with immunotherapy has not been conducted yet; therefore, little information is available on the adequate dosage of other immune treatments to formulate any recommendation.\textsuperscript{3} However, it is not easy in clinical practice to exclude a possibility of C-NORSE or antibody-positive AE particularly at the early stage of SE before antibody test results become available; therefore, many patients with NORSE may have been treated with immunotherapy,\textsuperscript{5,26} although the first-line immunotherapy is presumed to be less effective. However, if the C-NORSE score is high (≥5) on referral from other hospital, it is suggested that the patient is more likely to be negative for neuronal antibodies, thus more likely to be less responsive to first-line immunotherapy and have poor outcome. This scoring strategy might help physicians to identify patients with C-NORSE and their decision making in a patient with the high score.

Although the underlying mechanism of C-NORSE is entirely unknown, inflammation-mediated epileptogenesis has been proposed,\textsuperscript{27} in which a vicious cycle that involves inflammation and seizure activity is assumed to lead to cell death and network reorganization, ultimately causing refractory seizure. One previous study reported high levels of cytokines (interleukin-6 [IL-6]) or chemokines (CXCL10 and IL-8) in serum and CSF in pediatric cases of febrile infection-related epilepsy syndrome (FIRES),\textsuperscript{28} which is currently regarded as a subcategory of NORSE.\textsuperscript{3} Among those, proinflammatory cytokines, such as IL-1β and IL-6, have received attention as potential key molecules in C-NORSE. IL-1β has been implicated in seizure-induced neuronal cell death,\textsuperscript{29} SE,\textsuperscript{30} and posttraumatic epilepsy.\textsuperscript{31} Anakinra, IL-1 receptor antagonist, has been reported to be effective in patients with FIRES.\textsuperscript{32,33} IL-6 secreted from macrophages is also important mediator of fever and its deregulated expression is responsible for development of a variety of autoimmune inflammatory diseases.\textsuperscript{34} The efficacy of tocilizumab, IL-6 receptor antagonist, has also been reported in patients with NORSE.\textsuperscript{55} Therefore, elevated CSF levels of proinflammatory cytokines may play an important role in neuroinflammation, leading to development of refractory partial seizures in NORSE or FIRES. In our cohort of patients with C-NORSE, none of them had autoantibodies binding to the neuronal surface membrane with a rat brain immunohistochemistry in either CSF or serum, indicating that autoantibodies may not play an important role in C-NORSE or FIRES, but rather innate immunity may be more important than adaptive immunity as previously described.\textsuperscript{5}

Of interest, 21.2% of patients with C-NORSE had a PMH of FC, family history of FC, or both. In a small group of patients, some genetic predisposition to epileptic seizure might contribute to development of NORSE following fever. Further research is required to determine a role of genomic susceptibility to NORSE. This study has limitations of being retrospective studies and based on the small number of patients included. Genomic studies have not been performed yet in our cohort. Classical paraneoplastic antineuronal antibodies were not examined in all patients. Cytokine or chemokines were not examined in either case. In an emergency situation, some of the components of the score may be difficult to assess historically due to a variety of individual factors. A brain MRI is often difficult to obtain in a ventilated patient with SE-M or cannot be performed on a patient with contraindication (e.g., implanted pacemakers, intracranial aneurysm clips, and iron-based metal implants). When early brain MRI is unremarkable, repeated studies are required to see symmetric MRI abnormalities. However, a brain MRI within the first 24 hours is currently included in the diagnostic checklist for etiology of NORSE,\textsuperscript{36} and follow-up MRI is also important in exclusion of alternative diagnosis (multifocal corticostriatal lesions may appear in the course of the disease in anti-GABAaR encephalitis). It is important to keep in mind that this score was developed in patients with SE-M of unclear etiology. Thus, the results should not be generalized for patients with NCSE.

Despite these limitations, this study demonstrated that the clinically based score is useful for early identification of patients with C-NORSE. However, this score should not be used to make the diagnosis of C-NORSE because NORSE is not a specific diagnosis and exclusion of alternative diagnosis is mandatory. In patients with C-NORSE, irreversible brain damage is expected to occur quickly; thus, early recognition of C-NORSE is crucial. In addition to ASD treatment, we hope that this scoring strategy improves their functional outcome through facilitating early intervention with potential effective drugs that break a vicious cycle of neuroinflammation-induced neuronal damage that consequently increases seizure susceptibility.
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Disclosure

Publication history

Appendix Authors

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<tr>
<th>Name</th>
<th>Location</th>
<th>Contribution</th>
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<td>Atsuko Yanagida, MD</td>
<td>Kitasato University School of Medicine, Sagamihara, Japan</td>
<td>Designed and conceptualized the study; acquisition of data; analyzed and interpreted the data; and drafted and revised the manuscript for intellectual content</td>
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Appendix (continued)

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<tr>
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References
Clinically based score predicting cryptogenic NORSE at the early stage of status epilepticus
Atsuko Yanagida, Naomi Kanazawa, Juntaro Kaneko, et al.
*Neurol Neuroimmunol Neuroinflamm* 2020;7;
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