Mimics of Autoimmune Encephalitis
Validation of the 2016 Clinical Autoimmune Encephalitis Criteria

Robin W. Van Steenhoven, MD,* Juna M. de Vries, MD, PhD,* Arlette L. Bruijstens, MD, PhD, Manuela Paunovic, PhD, Mariska M. Nagtzaaam, Suzanne C. Franken, MSC, Anna E. Bastiaansen, MD, Marienke A. De Bruijn, MD, PhD, Agnes Van Sonderen, MD, PhD, Marco W.J. Schreurs, PhD, Mayke Gardeniers, MD, Robert M. Verdijk, MD, PhD, Rutger K. Balvers, MD, PhD, Peter A. Sillevis Smitt, MD, PhD, Rinze F. Neuteboom, MD, PhD, and Maarten J. Titulaer, MD, PhD

Neurol Neuroimmunol Neuroinflam 2023;10:e200148. doi:10.1212/NXI.0000000000200148

Correspondence
Dr. Titulaer
m.titulaer@erasmusmc.nl

Abstract

Background and Objectives
The clinical criteria for autoimmune encephalitis (AE) were proposed by Graus et al. in 2016. In this study, the AE criteria were validated in the real world, and common AE mimics were described. In addition, criteria for probable anti-LGI1 encephalitis were proposed and validated.

Methods
In this retrospective cohort study, patients referred to our national referral center with suspicion of AE and specific neuroinflammatory disorders with similar clinical presentations were included from July 2016 to December 2019. Exclusion criteria were pure cerebellar or peripheral nerve system disorders. All patients were evaluated according to the AE criteria.

Results
In total, 239 patients were included (56% female; median age 42 years, range 1–85). AE was diagnosed in 104 patients (44%) and AE mimics in 109 patients (46%). The most common AE mimics and misdiagnoses were neuroinflammatory CNS disorders (26%), psychiatric disorders (19%), epilepsy with a noninflammatory cause (13%), CNS infections (7%), neurodegenerative diseases (7%), and CNS neoplasms (6%). Common confounding factors were mesiotemporal lesions on brain MRI (17%) and false-positive antibodies in serum (12%). Additional mesiotemporal features (involvement extralimbic structures, enhancement, diffusion restriction) were observed more frequently in AE mimics compared with AE (61% vs 24%; p = 0.005). AE criteria showed the following sensitivity and specificity: possible AE, 83% (95% CI 74–89) and 27% (95% CI 20–36); definite autoimmune limbic encephalitis (LE), 10% (95% CI 5–17) and 98% (95% CI 94–100); and probable anti-NMDAR encephalitis, 50% (95% CI 26–74) and 96% (95% CI 92–98), respectively. Specificity of the criteria for probable seronegative AE was 99% (95% CI 96–100). The newly proposed criteria for probable anti-LGI1 encephalitis showed a sensitivity of 66% (95% CI 47–81) and specificity of 96% (95% CI 93–98).

Discussion
AE mimics occur frequently. Common pitfalls in AE misdiagnosis are mesiotemporal lesions (predominantly with atypical features) and false-positive serum antibodies. As expected, the specificity of the criteria for possible AE is low because these criteria represent the minimal requirements for entry in the diagnostic algorithm for AE. Criteria for probable AE (-LGI1, -NMDAR, seronegative) and definite autoimmune LE are applicable for decisions on immunotherapy in early disease stage, as specificity is high.

*These authors contributed equally to this work.

From the Department of Neurology (R.W.V.S., J.M.V., A.L.B., M.P., M.M.N., S.C.F., A.E.B., M.A.D.B., P.A.S.S., M.I.T.), Erasmus MC University Medical Center, Rotterdam; Department of Neurology (A.V.S.), Haaglanden Medical Center, The Hague; Departments of Immunology (M.W.J.S.), Radiology (M.G.), Neuropathology (R.M.V.), and Neurosurgery (R.K.B.), Erasmus MC University Medical Center; and Department of Pediatric Neurology (R.F.N.), Sophia Childrens Hospital, Erasmus MC University Medical Center, Rotterdam, The Netherlands.

Go to Neurology.org/NN for full disclosures. Funding information is provided at the end of the article.

The Article Processing Charge was funded by NIH, Cleveland Clinic, Genzyme.

This is an open access article distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivatives License 4.0 (CC BY-NC-ND), which permits downloading and sharing the work provided it is properly cited. The work cannot be changed in any way or used commercially without permission from the journal.
Glossary

ADEM = acute disseminated encephalomyelitis; AE = autoimmune encephalitis; CBA = cell-based assay; FBDS = faciobrachial dystonic seizures; HE = Hashimoto encephalopathy; IHC = immunohistochemistry; IQR = interquartile range; PNID = probable neuroinflammatory disorder; PNS = paraneoplastic neurologic syndromes; SN-AE = seronegative AE.

Introduction

The discovery of anti-NMDAR encephalitis (anti-NMDARE) in 2007 is regarded as a major breakthrough by introducing autoimmune encephalitis (AE) as a new disease entity with unique characteristics.1,2 In contrast to classical paraneoplastic neurologic syndromes (PNSs),3 AE is associated with neuronal autoantibodies against extracellular antigens, which are directly pathogenic.4 Prompt diagnosis is essential in AE because early administration of immunotherapy improves outcome in most patients with AE.5-7 The diagnosis of AE strongly relies on the identification of neuronal autoantibodies in serum and CSF.4,5 However, it has been stated that former clinical criteria for AE were too reliant on neuronal autoantibody status8 because comprehensive antibody testing can be time consuming and can result in diagnostic and therapeutic delay.5 In addition, the absence of neuronal autoantibodies does not exclude AE9,10 while on the other hand false-positive results may produce an incorrect diagnosis of AE.5,10-12 An important improvement in the clinical approach of patients with suspicion of AE was the publication of the clinical AE criteria in 2016 by Graus et al.5 based on conventional clinical neurologic assessment and standard diagnostic tests (MRI, EEG, and CSF studies). The 2016 AE criteria allow preliminary treatment with immunotherapy by establishment of an early diagnosis of probable or definite AE awaiting neuronal autoantibody status.5 In addition, a novel diagnosis of autoantibody-negative but probable AE and criteria for probable anti-NMDAR encephalitis (anti-NMDARE) were introduced (Figure 1).5 However, because many diseases can resemble AE and immunotherapy may induce serious adverse events or delay of alternative diagnoses,11,13 the diagnostic accuracy of the 2016 AE criteria, particularly specificity, is highly relevant for clinical practice. In this study, we validate the 2016 clinical AE criteria and describe frequently recognized mimics of AE and the red flags to prevent misdiagnosis. In addition, we propose criteria for probable anti-LGI1 encephalitis, to improve outcome by early diagnosis and treatment in this relatively common subtype of AE.7

Methods

Patient Selection and Diagnostics

In this retrospective cohort study, we included children and adults referred to the Erasmus MC University Medical Center with suspicion of AE in the period of July 2016 until December 2019. This study was performed according to the Strengthening the Reporting of Observational Studies in Epidemiology reporting guideline for observational research. The Erasmus MC University Medical Center is the Dutch national referral center for neuroinflammation and accredited European Reference Network site (ERA-RITA). All disorders presented in the diagnostic algorithm for AE proposed by Graus et al.5 were included. Patients with autoimmune cerebellopathies, opsoclonus-myoclonus syndrome, and disorders exclusively affecting the peripheral nerve system were excluded because the 2016 clinical AE criteria focus on patients presenting with a subacute onset of memory deficits or altered mental status.5 Ancillary testing included blood analysis, lumbar puncture, EEG, MRI, and cerebral biopsy, if necessary. All patients underwent extensive neuronal autoantibody testing in serum and CSF, if available, using a combination of tests, including in-house immunohistochemistry (IHC) on rat brain slices.14 Specific cell surface autoantibodies were tested using commercial cell-based assays (CBAs; Euroimmun, Lübeck, Germany) or in-house CBAs. Only those samples with positive CBAs were considered positive that could be confirmed by IHC or, if necessary, live hippocampal cell cultures (LN).14-16 GlyR,17 KLHL-11,18 GFAP,19 IgLON5,20 mGluR1, and mGluR5 were tested by in-house CBAs.21,22 Anti-MOG antibodies were tested using CBA, as described elsewhere.23 Anti-GAD65 was tested by ELISA (Medizym anti-GAD 96, Medipan, Berlin, Germany) and considered clinically relevant if serum concentration was >10,000 IU/mL or CSF concentration was >100 IU/mL (high titer) and IHC showed a compatible staining pattern.24 Antibodies against paraneoplastic neurologic (‘onconeural’) antigens amphiphysin, CV2, Ma1/2, Ri, Yo, Hu, and Tr (DNER) were detected by the combined use of line immunoassay (EUROLINE Paraneoplastic neurological Syndrome 12 Ag (IgG), Euroimmun, Lübeck, Germany), and when positive, it is confirmed using a second antigen-specific line immunoassay (PNS-Blot, Ravo Diagnostika, Freiburg, Germany) and indirect immunofluorescence (Cerebellum (Primary) Slide, Inova Diagnostics, San Diego, CA).

Definitions

All patients were physically seen by the authors (R.W.V.S, A.L.B., Y.S.C., A.E.M.B., J.M.D.V., R.F.N., and M.J.T.). Medical records were reviewed, and definite diagnoses were made by consensus. Patients with positive neuronal autoantibody status and a compatible clinical syndrome, including PNS, were classified as definite AE. An AE mimic was defined as a patient with initial strong suspicion of AE and an alternative final diagnosis. All patients were evaluated according to the 2016 AE criteria,5 including our proposed criteria for probable anti-LGI1 encephalitis (Figure 2). Patients were classified as
probable seronegative AE (SN-AE), if the 2016 criteria were satisfied.\(^5\) Established criteria were used to define acute disseminated encephalomyelitis (ADEM) and Hashimoto encephalopathy (HE).\(^5\) ADEM and HE were classified as separate inflammatory categories (i.e., not as inflammatory AE mimic) because these disorders were also separately included in the 2016 AE criteria.\(^5\) Patients with strong evidence of a neuroinflammatory disorder, who did not fulfill the criteria for probable SN-AE or any other specific inflammatory CNS disorder, were classified in this study as probable neuroinflammatory disorder (PNID), which was considered as an inflammatory subcategory of the mimics. In this study, strong evidence of a neuroinflammatory disorder was defined as the presence of ≥2 of the following characteristics: brain MRI suggestive of AE, CSF pleocytosis, specific oligoclonal bands in CSF, repeated steroid responsiveness, or similar staining

---

### 1.1 Possible autoimmune encephalitis

1. Subacute onset (rapid progression of less than 3 months) of working memory deficits (short-term memory loss), altered mental status, or psychiatric symptoms
2. At least one of the following:
   - New focal CNS findings
   - Seizures not explained by a previously known seizure disorder
   - CSF pleocytosis (while blood cell count >5 cells per mm\(^3\))
3. Reasonable exclusion of alternative causes

### 1.2 Definite autoimmune limbic encephalitis

1. Subacute onset (rapid progression of less than 3 months) of working memory deficits, seizures, or psychiatric symptoms suggesting involvement of the limbic system
2. Bilateral brain abnormalities on T2-weighted fluid-attenuated inversion recovery MRI highly restricted to the medial temporal lobes
3. At least one of the following:
   - CSF pleocytosis
   - EEG with epileptic or slow-wave activity involving the temporal lobes
4. Reasonable exclusion of alternative causes

### 1.3 Probable seronegative autoimmune encephalitis

1. Subacute onset (rapid progression of less than 3 months) of working memory deficits, (short-term memory loss), altered mental status, or psychiatric symptoms
2. Exclusion of well-defined syndromes of autoimmune encephalitis (e.g., typical limbic encephalitis, Bickerstaff brainstem encephalitis, acute disseminated encephalomyelitis)
3. Absence of well characterized autoantibodies in serum and CSF, and at least two of the following criteria:
   - MRI abnormalities suggestive of autoimmune encephalitis
   - CSF pleocytosis, CSF-specific oligoclonal bands, or an elevated CSF IgG index, or both
   - Brain biopsy showing inflammatory infiltrates and excluding other disorder (e.g., tumor)
4. Reasonable exclusion of alternative causes

### 1.4 Probable anti-NMDA receptor encephalitis

Diagnosis can be made when all 3 of the following criteria have been met
1. Rapid onset (less than three months) of at least four of the six following major groups of symptoms:
   - Abnormal (psychiatric) behavior or cognitive dysfunction
   - Speech dysfunction ( pressured speech, verbal reduction, mutism)
   - Seizures
   - Movement disorder, dyskinesias, or rigidity/abnormal postures
   - Decreased level of consciousness
   - Autonomic dysfunction or central hypoventilation
2. At least one of the following laboratory study results:
   - Abnormal EEG (focal or diffuse slow or disorganized activity, epileptic activity, or extreme delta brush)
   - CSF with pleocytosis or oligoclonal bands
3. Reasonable exclusion of alternative causes

Diagnosis can also be made in the presence of three of the above groups of symptoms accompanied by a systemic teratoma

\(^\text{†}\)Antibody testing should include testing of CSF. If only serum is available, confirmatory tests should be included (e.g., live neurons or tissue immunohistochemistry, in addition to cell-based assay)

---

Reprinted with permission from Elsevier.
pattern on IHC in serum and CSF in the absence of a known neuronal autoantibody. Patients exclusively demonstrating a pleocytosis and an altered mental status, new-onset seizures, or focal deficits without other specific evidence of an infectious or inflammatory cause were labeled as encephalitis with unknown cause.

**Standard Protocol Approvals, Registrations, and Patient Consents**

IRB approval was waivered, but informed consent for usage of medical information for research purposes was obtained from all patients or proxies that could be reached.

**Statistics**

We used IBM SPSS 25.0 (SPSS Inc) and Prism 8.4.3 (GraphPad) for statistical analysis. The Pearson χ² test or the Fisher-Freeman-Halton test, when appropriate, was used for patient characteristics analysis and group comparisons, encompassing categorical data. p-values were two-sided and considered statistically significant when below 0.05. Sensitivity and specificity of the 2016 clinical AE criteria were calculated. Sensitivity was defined as the percentage of definite AE and probable SN-AE of all patients fulfilling a specific category of the clinical AE criteria (i.e., true positive), whereas specificity was defined as the percentage of diagnoses other than AE, including other neuroinflammatory disorders, of all patients not fulfilling these criteria (i.e., true negative). No sensitivity was determined for probable SN-AE, in view of the low expected number of patients and absence of a gold standard for this particular diagnosis. Criteria for probable anti-LGI1 encephalitis were also applied to a previous nonoverlapping cohort from our center, described earlier.²⁵ Similarly, criteria for probable anti-NMDARE were calculated in the whole national cohort (that includes the patients from this cohort with anti-NMDARE) to account for potential bias.²⁶

**Data Availability**

The data of this study, coded to adhere to legal privacy regulations, are available on request from any qualified investigator.

**Results**

**Patient Characteristics**

Over a three-and-a-half year period, 310 patients were assessed for eligibility (eFigure 1, links.lww.com/NXI/A889). A total of 239 patients with a suspicion of AE were included, of whom 134 (56%) were female. The median follow-up was 11.0 months (interquartile range [IQR] 1–24.5, range 0–277). Sixty patients (25%) were children (younger than 18 years) at the onset of symptoms. The median age was 42 years (IQR 18–65, range 1–85). Definite AE was diagnosed in 96 patients (40%) and probable SN-AE in 8 patients (3%; Figure 3). Other inflammatory categories included ADEM (9%) and HE (2%). A total of 109 patients (46%) were ultimately classified as AE mimic. In adult patients (age at onset of symptoms 18 years or older), definite AE was diagnosed more frequently (49% vs 13%; p < 0.001) while ADEM was observed more often in children (32% vs 1%; p < 0.001). Patient characteristics and comparison of AE (definite AE and probable SN-AE) vs AE mimics are summarized in Table 1, eTable 1, and eFigure 2. New-onset seizures were observed more frequently in AE than in AE mimics (73% vs 39%; p < 0.001). In addition, patients with AE presented more often (49% vs 21%, p < 0.001) with ≥3 of the following symptoms: working memory deficits, new-onset seizures, behavioral disorders, psychiatric symptoms, and sleeping disorders. Regarding ancillary testing, bilateral mesiotemporal hyperintensities on brain MRI (36% vs 12%; p < 0.001) and epileptic abnormalities on EEG (47% vs 21%; p < 0.001) were described more frequently in AE. A newly diagnosed malignancy was observed more often in patients with AE than in AE mimics (17% vs 1%; p < 0.001).

**AE Mimics and Confounders**

The most frequent AE mimics were CNS inflammatory disorders (26%; PNID 14% and other CNS inflammatory diseases 12%), primary psychiatric disorders (19%), epilepsy with a noninflammatory cause (13%), CNS infectious diseases (7%), encephalitis with unknown cause (7%), neurodegenerative diseases (7%), and primary CNS neoplasms (6%) (Figure 3, specific diagnoses per subcategory are provided in eTable 2, links.lww.com/NXI/A889). In children, primary psychiatric disorders were observed more frequently (36% vs 12%; p = 0.016). Overall, the most frequent confounding factor for AE misdiagnosis was MRI T2/FLAIR hyperintensities involving the mesiotemporal lobe(s), observed in 18 of 109 (17%) AE mimics (Table 1). The presence of ≥1 atypical radiologic feature (involvement of extralimbic structures, enhancement, diffusion restriction), in addition to mesiotemporal T2/FLAIR hyperintensities, was observed
more frequently in AE mimics compared with AE (61% vs 24%; $p = 0.005$). Brain biopsy was performed in 5 of 18 patients with mesiotemporal lobe abnormalities and provided a definite diagnosis in all 5 patients, including GBM (n = 3), B-cell lymphoma, and CNS Whipple disease. The second most common confounder was false-positive or clinically irrelevant antibodies in serum, which was observed in 13 of 109 (12%) AE mimics, including thyroid peroxidase antibodies (anti-TPO; n = 4), NMDAR antibodies (n = 5), a positive VGKC antibody test in the absence of LGI1 and CASPR2 antibodies (n = 2), and Hu and Ma2 antibodies (both n = 1). Sixty-one AE mimics (56%) were treated with immunotherapy, of whom 13 had an CNS inflammatory disease and 15 were classified as PNID (eTable 3, links.lww.com/NXI/A889).

**Validation Clinical AE Criteria**

An evaluation using the diagnostic algorithm for AE is provided for all patients in Figure 4 and for the adult population.
Table 1 Comparison of Patients Characteristics: AE (Definite AE and Probable Seronegative AE) vs AE Mimics<sup>a</sup>

<table>
<thead>
<tr>
<th></th>
<th>AE (n = 104)</th>
<th>AE mimic (n = 109)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female sex, n (%)</td>
<td>59 (57)</td>
<td>62 (57)</td>
<td>0.98</td>
</tr>
<tr>
<td>Age at onset in years (median; IQR; range)</td>
<td>55; 32–67; 9–84</td>
<td>39; 15–65; 1–81</td>
<td>0.017</td>
</tr>
<tr>
<td>Age at onset &lt;18 y, n (%)</td>
<td>8 (8)</td>
<td>33 (30)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Medical history, n (%)</td>
<td>8 (8)</td>
<td>24 (22)</td>
<td>0.74</td>
</tr>
<tr>
<td>Autoimmune disease</td>
<td>21 (20)</td>
<td>6 (6)</td>
<td>0.30</td>
</tr>
<tr>
<td>Active malignancy, n (%)</td>
<td>23 (22)</td>
<td>5 (5)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Unknown at first visit</td>
<td>18 (17)</td>
<td>1 (1)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Subacute presentation (&lt;3 mo), n (%)</td>
<td>90 (87)</td>
<td>88 (81)</td>
<td>0.25</td>
</tr>
<tr>
<td>Symptoms, n (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>New-onset seizures</td>
<td>76 (73)</td>
<td>43 (39)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Working memory deficits</td>
<td>73 (70)</td>
<td>62 (59)</td>
<td>0.077</td>
</tr>
<tr>
<td>Behavioral disorders</td>
<td>65 (63)</td>
<td>61 (58)</td>
<td>0.46</td>
</tr>
<tr>
<td>Sleeping disorders</td>
<td>38 (37)</td>
<td>28 (26)</td>
<td>0.11</td>
</tr>
<tr>
<td>Psychiatric symptoms</td>
<td>35 (34)</td>
<td>48 (45)</td>
<td>0.085</td>
</tr>
<tr>
<td>Autonomous disorders</td>
<td>25 (24)</td>
<td>17 (16)</td>
<td>0.15</td>
</tr>
<tr>
<td>Movement disorder</td>
<td>20 (19)</td>
<td>28 (26)</td>
<td>0.23</td>
</tr>
<tr>
<td>Focal deficits</td>
<td>15 (14)</td>
<td>27 (25)</td>
<td>0.053</td>
</tr>
<tr>
<td>Decreased level of consciousness</td>
<td>12 (12)</td>
<td>18 (17)</td>
<td>0.30</td>
</tr>
<tr>
<td>≥3 symptoms&lt;sup&gt;b&lt;/sup&gt;, n (%)</td>
<td>51 (49)</td>
<td>23 (21)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>MRI of the brain performed, n (%)</td>
<td>101 (97)</td>
<td>105 (96)</td>
<td>0.75</td>
</tr>
<tr>
<td>Mesiotemporal hyperintensities</td>
<td>46 (46)</td>
<td>18 (17)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Bilateral</td>
<td>36 (36)</td>
<td>13 (12)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Unilateral</td>
<td>12 (12)</td>
<td>5 (5)</td>
<td>0.078</td>
</tr>
<tr>
<td>Involvement of extralimbic structures</td>
<td>15 (15)</td>
<td>24 (23)</td>
<td>0.14</td>
</tr>
<tr>
<td>Enhancement</td>
<td>12/72 (17)</td>
<td>16/77 (21)</td>
<td>0.52</td>
</tr>
<tr>
<td>Diffusion restriction</td>
<td>3/101 (3)</td>
<td>7/105 (7)</td>
<td>0.22</td>
</tr>
<tr>
<td>Mesiotemporal hyperintensities with ≥1 atypical MRI feature&lt;sup&gt;c&lt;/sup&gt;</td>
<td>11/46 (24)</td>
<td>11/18 (61)</td>
<td>0.005</td>
</tr>
<tr>
<td>CSF performed, n (%)</td>
<td>101 (97)</td>
<td>98 (90)</td>
<td>0.034</td>
</tr>
<tr>
<td>WBC&gt;5/μL</td>
<td>39/98 (40)</td>
<td>43/96 (45)</td>
<td>0.48</td>
</tr>
<tr>
<td>Oligoclonal bands</td>
<td>14/39 (36)</td>
<td>18/56 (32)</td>
<td>0.70</td>
</tr>
<tr>
<td>EEG performed, n (%)</td>
<td>82 (79)</td>
<td>85 (78)</td>
<td>0.88</td>
</tr>
</tbody>
</table>

Table 1 Comparison of Patients Characteristics: AE (Definite AE and Probable Seronegative AE) vs AE Mimics<sup>a</sup> (continued)

<table>
<thead>
<tr>
<th></th>
<th>AE (n = 104)</th>
<th>AE mimic (n = 109)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Epileptic abnormalities</td>
<td>38 (47)</td>
<td>18 (21)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Focal abnormalities</td>
<td>58 (73)</td>
<td>55 (65)</td>
<td>0.23</td>
</tr>
<tr>
<td>Immunotherapy, n (%)</td>
<td>100 (96)</td>
<td>61&lt;sup&gt;a&lt;/sup&gt; (56)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>First line</td>
<td>99 (95)</td>
<td>61 (56)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Second line</td>
<td>42 (40)</td>
<td>13 (12)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Chronic</td>
<td>70 (67)</td>
<td>22 (20)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

Abbreviations: AE = autoimmune encephalitis; IQR = interquartile range; SN-AE = seronegative autoimmune encephalitis; WBC = white blood cell.
Bold p-values refer to those that meet the predefined cut-off value below 0.05, showing statistical significance.
<sup>a</sup>eTable 1 (links.lww.com/NXI/A889) provides a comparison between all inflammatory CNS disorders vs AE mimics.
<sup>b</sup>Five most frequent symptoms in AE: working memory problems, new-onset seizures, behavioral disorders, sleeping disorders, and psychiatric symptoms.
<sup>c</sup>Involvement of extralimbic structures, enhancement, or diffusion restriction.
<sup>d</sup>28/61 (46%): CNS inflammatory disorders.

(operative age 18 years and older) in eFigure 3 (links.lww.com/NXI/A889). Fifty-five of all 239 patients (23%) with suspicion of AE did not meet the criteria for possible AE, of whom 35 of 55 (63%) were AE mimics and 18 of 55 (33%) definite AE. AE mimics not fulfilling criteria for possible AE included predominantly primary psychiatric disorders (19/35; 54%). Patients with probable and definite AE not fulfilling criteria for possible AE had more frequently high titer anti-GAD65 antibodies (15/18, 83%; \( p < 0.001 \)), were female (17/18, 94%; \( p < 0.001 \)), and had lower median age at symptom onset (31 years, \( p < 0.001 \), eTable 4). A total of 184 of all 239 patients (77%) fulfilled the criteria for possible AE, and 78/184 (42%) of patients satisfying these criteria were diagnosed with definite AE and 8/184 (4%) with probable SN-AE. Thirteen of 184 patients (7%) fulfilling the criteria for possible AE also met the criteria for definite LE, of whom 8 patients (62%) had a diagnosis of definite AE and 2 patients (15%) were finally diagnosed as probable SN-AE. Three patients fulfilling the criteria for definite LE (23%) were classified as AE mimic (Figure 5). Nine of 18 (50%) patients meeting the criteria for probable anti-NMDARE were diagnosed as definite anti-NMDARE while 3 of 9 remaining patients had an alternative neuroinflammatory disorder and 3 patients were classified as PNID. Nine of 184 patients with possible AE (5%) fulfilled the criteria for probable SN-AE. One of these patients was ultimately diagnosed with a GBM (Figure 5). Twenty four of 184 (13%) patients had another specific CNS inflammatory disorder (ADEM 11%, HE 2%). In total, 74 of 184 (40%) patients with possible AE were ultimately classified as AE mimic. The accuracy of the 2016 AE criteria is provided in Table 2, eTable 5, and eTable 6.
Figure 4 Evaluation of Patients According to Diagnostic Algorithm for Autoimmune Encephalitis

Adapted from Graus et al.,5 reprinted with permission from Elsevier. ♦ Anti-MOG was tested in all (n = 21) patients with ADEM; in 10/21 (48%) antibodies were present. *Two patients ultimately diagnosed as probable SN-AE. Three patients fulfilling criteria for definite autoimmune limbic encephalitis were diagnosed as AE mimic after applying diagnostic AE algorithm. † Anti-NMDARE (1), PACNS (1), PML (1), MS (1), and neuro-Sjögren (1). ¥ Encephalitis with unknown cause (3), PNID (3), definite AE (2), and HE (1). Ψ Caspr2 (6), Ma2 (1), and NMDAR (1). Abs = antibodies; ADEM = acute disseminated encephalomyelitis; AE = autoimmune encephalitis; BBE = Bickerstaff brainstem encephalitis; HE = Hashimoto encephalopathy; MOG = myelin oligodendrocyte glycoprotein; NMDARE = anti-NMDA receptor encephalitis; PNID = probable neuroinflammatory disorder; SN-AE = seronegative autoimmune encephalitis.
Criteria for Probable Anti-LGI1 Encephalitis

In total, 32 patients were diagnosed as definite anti-LGI1 encephalitis, of whom 13 (41%) demonstrated faciobrachial dystonic seizures (FBDS) and 12 (38%) demonstrated frequent (>5 per day) stereotypical focal seizures. In all patients, focal seizures had a nonmotor onset, predominantly dyscognitive and autonomous. In the diagnostic algorithm for AE, one patient presented with isolated faciobrachial dystonic and did not fulfill the criteria for possible AE (Figure 4, eFigure 4, links.lww.com/NXI/A889). Three patients with anti-LGI1 encephalitis met the criteria for definite autoimmune LE. Twenty five of 171 (15%) remaining patients with possible AE not fulfilling criteria for definite autoimmune LE met the proposed criteria for probable anti-LGI1 encephalitis. Eighteen of these 25 (72%) patients could be confirmed by antibody testing, whereas the other 7 patients were diagnosed as definite AE with another antibody (eFigure 4). FBDS was exclusively observed in anti-LGI1 encephalitis. Overall, the criteria for probable anti-LGI1 encephalitis showed a sensitivity of 66% (95% CI 47–81) and specificity of 96% (95% CI 93–98; Table 2). It allowed earlier treatment in 25 of 171 (15%) without treating noninflammatory AE mimics erroneously. The criteria for probable anti-LGI1 encephalitis were validated on an earlier described cohort, of whom one overlapping patient was excluded (n = 37), showing a comparable sensitivity (65%; 95% CI 47–80; p = 0.9).

Discussion

In this retrospective cohort study, we describe common AE mimics and validate the 2016 AE criteria using real-world data. In addition, we propose criteria for probable anti-LGI1 encephalitis. We demonstrate that the specificity for probable AE (NMDAR, SN-AE, and LGI1) and definite autoimmune LE criteria is reassuringly high (>95%). Furthermore, we show that AE mimics occur frequently and are diverse. The most common diagnostic categories are primary psychiatric disorders, CNS inflammatory disorders, epilepsy with a noninflammatory cause, CNS infections, neurodegenerative diseases, and primary CNS neoplasms. The sensitivity of the criteria for possible AE was relatively high (83%), which was comparable with previous studies by Li et al.27 and Costa et al.28 This implicates that most patients with AE can be identified by these criteria. However, a substantial part of AE did not fulfill the criteria for possible AE, of whom the majority had anti-GAD65 antibodies with a chronic course. This demonstrates that these criteria focus on patients with a subacute presentation, and sensitivity for neuronal autoantibodies associated with a chronic course (i.e., Caspr2, IgLONS,
GAD65) is only moderate. Ninety percent of patients with a primary psychiatric disorder did not meet the criteria for possible AE, indicating high specificity in this category. However, the overall specificity of the criteria for possible AE was markedly lower (27%), indicating a relatively high rate of false-positive cases and potentially erroneous treatment with immunotherapy. Previous studies reported higher specificities (72%–94%),11,27,28 which is probably explained by differences in AE mimic population. Compared with an earlier study performed by Flanagan et al.,11 we observed a higher frequency of CNS inflammatory and CNS infectious disorders in our study, of whom the majority fulfilled the criteria for possible AE, whereas the occurrence of neurodegenerative diseases was lower. Our findings emphasize that the criteria for possible AE are useful as entry criteria for the diagnostic algorithm of AE. However, possible AE should not be regarded as an established diagnosis and requires ancillary testing because specificity is (too) low. The criteria for probable AE (NMDAR, SN-AE) and definite autoimmune LE were highly specific (>95%), indicating a very low risk of false-positive cases. Li et al.27 reported comparable specificities for probable anti-NMDARE and definite autoimmune LE. We deliberately chose to include other inflammatory CNS disorders in the control group (i.e., patients with a diagnosis other than AE) to obtain optimal specificity for AE. The sensitivity of probable anti-NMDARE criteria in this study (50%) was lower compared with earlier research (81%–90%),29,31 which is probably explained by an underrepresentation of severely affected anti-NMDARE patients in our study. This is supported by a higher sensitivity (58%) if we applied these criteria to the

### Table 2 Accuracy of the 2016 AE Criteriaa

<table>
<thead>
<tr>
<th></th>
<th>Probable SN-AE and definite AE</th>
<th>No AE</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Possible AE+</td>
<td>86</td>
<td>98</td>
<td>184</td>
</tr>
<tr>
<td>Possible AE−</td>
<td>18</td>
<td>37</td>
<td>55</td>
</tr>
<tr>
<td></td>
<td>104</td>
<td>135</td>
<td>239</td>
</tr>
<tr>
<td></td>
<td>Sens 83% [74–89]</td>
<td>Spec 27% [20–36]</td>
<td></td>
</tr>
<tr>
<td>Definite autoimmune LE+</td>
<td>10</td>
<td>3</td>
<td>13</td>
</tr>
<tr>
<td>Definite autoimmune LE−</td>
<td>94</td>
<td>132</td>
<td>226</td>
</tr>
<tr>
<td></td>
<td>104</td>
<td>135</td>
<td>239</td>
</tr>
<tr>
<td></td>
<td>Sens 10% [5–17]</td>
<td>Spec 98% [94–100]</td>
<td></td>
</tr>
<tr>
<td>Probable SN-AE+</td>
<td>8</td>
<td>1</td>
<td>9</td>
</tr>
<tr>
<td>Probable SN-AE−</td>
<td>NA</td>
<td>134</td>
<td>230</td>
</tr>
<tr>
<td></td>
<td>NA</td>
<td>135</td>
<td>239</td>
</tr>
<tr>
<td></td>
<td>Sens: ND</td>
<td>Spec 99% [96–100]</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Anti-NMDARE</td>
<td>No anti-NMDARE</td>
<td></td>
</tr>
<tr>
<td>Probable anti-NMDARE+</td>
<td>9</td>
<td>9</td>
<td>18</td>
</tr>
<tr>
<td>Probable anti-NMDARE−</td>
<td>9</td>
<td>212</td>
<td>221</td>
</tr>
<tr>
<td></td>
<td>18</td>
<td>221</td>
<td>239</td>
</tr>
<tr>
<td></td>
<td>Sens 50% [26–74]</td>
<td>Spec 96% [92–98]</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Anti-LGI1 AE</td>
<td>No anti-LGI1 AE</td>
<td></td>
</tr>
<tr>
<td>Probable anti-LGI1 AE+</td>
<td>21</td>
<td>8</td>
<td>29</td>
</tr>
<tr>
<td>Probable anti-LGI1 AE−</td>
<td>11</td>
<td>199</td>
<td>210</td>
</tr>
<tr>
<td></td>
<td>32</td>
<td>207</td>
<td>239</td>
</tr>
<tr>
<td></td>
<td>Sens 66% [47–81]</td>
<td>Spec 96% [93–98]</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: AE = autoimmune encephalitis; Anti-NMDARE = anti-NMDAR encephalitis; Anti-LGI1 AE = anti-LGI1 autoimmune encephalitis; LE = limbic encephalitis; ND = not determined; NPV = negative predictive value; PPV = positive predictive value; Sens = sensitivity; SN-AE = seronegative autoimmune encephalitis; Spec = specificity.

a eTable 5 (links.lww.com/NXI/A889) provides accuracy of the 2016 AE criteria in the adult population (age at onset ≥18 y), eTable 6 provides accuracy of the 2016 AE criteria including all CNS inflammatory disorders in the disease group.
cortex. The criteria for probable anti-LGI1 encephalitis should be applied in addition to the 2016 clinical AE criteria (i.e., as part of the diagnostic algorithm) and that these criteria are only applicable for patients who also fulfill the criteria for possible AE. FBDS was exclusively observed in anti-LGI1 encephalitis and can be regarded as pathognomonic for this disorder. Therefore, when recognizing FBDS, patients should always be tested for LGI1 antibodies, also in those without cognitive or additional symptoms and not fulfilling criteria for possible AE. Although focal epilepsy has a broad differential diagnosis, we found that frequent stereotypical focal seizures (>5 seizures per day) were also specific for anti-LGI1 encephalitis. All anti-LGI1 encephalitis patients with focal seizures had a nonmotor onset in this study. Particularly, high frequent focal dyscognitive and autonomic seizures should raise suspicion for anti-LGI1 encephalitis, as described earlier. In this study, the most important differential diagnosis of subacute cognitive decline and frequent focal seizures was anti-Caspr2 encephalitis, requiring similar treatment regimens. Three of 32 (10%) patients with anti-LGI1 encephalitis fulfilled the criteria for definite autoimmune LE because bilateral mesiotemporal hyperintensities were observed only in a minority of patients, as described earlier. By including the criteria for probable anti-LGI1 encephalitis to the diagnostic algorithm for AE proposed by Graus et al., a substantial part of anti-LGI1 encephalitis patients can be identified and treated earlier (e.g., prior to antibody test results), which improves outcome. In this study, the criteria for probable anti-LGI1 encephalitis were validated in another cohort from our center without overlap of patients. The LGI1 criteria would profit from validation in cohorts from other countries. A considerable part (46%) of patients were classified as AE mimics. We found that more than half of AE mimics were treated with steroids, supporting the importance of early AE mimic identification because steroids may induce various adverse effects, including a reduction of the diagnostic yield of brain biopsy in CNS lymphoma, deterioration of symptoms in psychiatric disorders, or exacerbation of CNS infections. We described various diagnostic pitfalls that had an important contribution to AE misdiagnosis. First, bilateral mesiotemporal hyperintensities on brain MRI were highly specific for AE, as reported earlier. However, mesiotemporal lesions were also the most important pitfall in AE misdiagnosis (17%) and should therefore be interpreted with caution and rigor. Notably, most AE mimics with mesiotemporal lesions demonstrated additional radiologic features, including enhancements, diffusion restriction, or involvement of extralimbic regions. These features were also observed in a minority of patients with AE but were usually mild and transient, as described earlier. Earlier research showed that diffusion restriction can distinguish HSV1 encephalitis from AE in early stages, whereas mass effect, involvement of extralimbic regions, and enhancement can be seen in gliomas. Furthermore, bilateral mesiotemporal enhancement has been reported in neurosyphilis and CNS Whipple disease. We suggest that mesiotemporal lesions on brain MRI with pronounced additional radiologic features should raise suspicion of an AE mimic, necessitating ancillary testing, including follow-up MRI and brain biopsy in selected cases, particularly if a CNS tumor is suspected. The second most common confounding factor in AE misdiagnosis was false-positive or clinically irrelevant antibody test results (12%). This percentage was notably lower compared with an earlier study by Flanagan et al., reporting positive serum antibodies in 50% of AE misdiagnoses. The testing of extensive antibody panels, with the adjoining risks of false-positive or clinically irrelevant results, which is not advocated nor commonly used within our country, might explain this difference. These findings emphasize the importance of adequate patient selection for antibody studies, as stated earlier, and the relevance of adequate neuronal antibody test methodologies, by using confirmatory test modalities and inclusion of CSF in antibody studies. In addition to diagnostic characteristics, we identified various clinical characteristics that may aid to discriminate between AE and AE mimics. First, the occurrence of seizures was higher in AE and should raise suspicion for AE in patients presenting with subacute cognitive impairment. In particular anti-LGI1, anti-NMDAR, anti-GABAAR, anti-GABAAR, and anti-GAD antibodies are associated with seizures. Second, we show that patients with AE had more frequently a polysymptomatic presentation, as described earlier in various AE subtypes, probably reflecting diffuse or multifocal brain inflammation. Third, a systemic tumor was more common in AE and might suggest a PNS. However, tumor status is frequently unknown at the onset of neurologic symptoms (78% in our study) and therefore less useful. In addition, it is essential to establish causality between type and neurologic syndrome by using the updated PNS-Care Score because comorbid tumors may be
detected. 46 In this study, 8 patients (3%) were diagnosed with SN-AE, which was markedly lower compared with earlier studies. 9 This might be partially explained by extensive testing for relevant antibodies, also in research setting, as well as a very rigorous application of the criteria for SN-AE in our study. However, we observed that a substantial part of patients (n = 15, 8%) could not be classified as SN-AE nor a specific neuroinflammatory disorder, despite a high suspicion of an inflammatory etiology (e.g., suggestive brain MRI or inflammatory CSF profile). In this study, we classified these patients as probable neuroinflammatory disorder (PNID), after thorough exclusion of other diseases, particularly CNS infections and malignancies. This category should be interpreted with caution because heterogeneity is high, and some patients may be diagnosed with another disease at a later stage. Therefore, PNID was classified as AE mimic, to prevent over-interpretation. However, it should be noted that AE mimics also include inflammatory disorders and some patients with PNID might have an inflammatory disorder requiring immunotherapy, although formal criteria are not satisfied. Further research is needed to characterize this heterogeneous patient category, clarify underlying pathogenic mechanisms, and identify new biomarkers. This study has some limitations. First, selection bias probably occurred in this cohort because it was a single-center study from a specialized institution with a relatively high occurrence of neuroinflammatory disorders. However, because we are a national referral center for AE and related disorders, patients from many other (e.g., nonspecialized) institutions were included. Similarly, selection bias mostly influences positive value and negative predictive values, whereas sensitivity and specificity should remain the same. Second, only a small number of pediatric patients with AE were included in our study, in line with the low incidence of AE in children. 32 Consequently, results for this specific patient category should be interpreted carefully. Although the 2016 AE criteria were also considered to allow for inclusion of children, additional pediatric AE criteria were proposed in 2020, that require validation in future research. 47 In summary, criteria for probable and definite AE, including newly proposed criteria for anti-LGI1 encephalitis, are applicable for early decisions on immunotherapy because specificity is high. Specificity of possible AE criteria is low and should, therefore, be regarded as an entry criterion for more extensive investigations, instead of established diagnosis. Various disorders can present as an AE mimic and cause misdiagnosis. Particularly, early identification of CNS infections and CNS tumors is essential because treatment strategies differ substantially.

Acknowledgment
The authors thank all patients for their participation and all referring physicians. M.W.J. Schreurs, P.A. Sillevis Smitt, J.M. de Vries, R.F. Neuteboom, and M.J. Titulaer of this publication are members of the European Reference Network for Rare Immunodeficiency, Autoinflammatory and Autoimmune Diseases-Project ID No 739543 (ERN-RITA; HCP Erasmus MC).

Study Funding
Dr. Titulaer was supported by an Erasmus MC fellowship, has received funding from the Netherlands Organization for Scientific Research (NWO, Veni incentive), ZonMw (Memorial program), the Dutch Epilepsy Foundation (NEF 14-19 & 19-08), Diorapthe (2001 0403), and E-RARE JTC 2018 (UltraAIE, 9003037605).

Disclosure
P.A. Sillevis Smitt holds a patent for the detection of anti-DNER, he received research support from Euroimmun. R.F. Neuteboom reports participates in pediatric MS studies with Novartis, Roche, and Sanofi-Genzyme; he received consultancy fees from Novartis, Sanofi-Genzyme, and Zogenix; he received research grants from the Dutch MS research foundation, DreaMS foundation, Postcode Loterij, Vrienden Loterij, Stichting Vrienden van het Sophia. M.J. Titulaer has filed a patent, on behalf of the Erasmus MC, for methods for typing neurologic disorders and cancer, and devices for use therein; has received research funds for serving on a scientific advisory board of Horizon Therapeutics, for consultation at Guidedpoint Global LLC, for consultation at UCB, for teaching colleagues at Novartis; and has received an unrestricted research grant from Euroimmun AG and from CSL Behring. The other authors report no relevant disclosures. Go to Neurology.org/NN for full disclosures.

Publication History
Received by Neurology: Neuroimmunology & Neuroinflammation May 8, 2023. Accepted in final form June 27, 2023. Submitted and externally peer reviewed. The handling editor was Editor Josep O. Dalmau, MD, PhD, FAAN.

Appendix Authors

<table>
<thead>
<tr>
<th>Name</th>
<th>Location</th>
<th>Contribution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Robin W. Van Steenhoven, MD</td>
<td>Department of Neurology, Erasmus MC University Medical Center, Rotterdam, The Netherlands</td>
<td>Major role in the acquisition of data; study concept or design; analysis or interpretation of data</td>
</tr>
<tr>
<td>Juna M. de Vries, MD, PhD</td>
<td>Department of Neurology, Erasmus MC University Medical Center, Rotterdam, The Netherlands</td>
<td>Major role in the acquisition of data; study concept or design; analysis or interpretation of data</td>
</tr>
<tr>
<td>Arlette L. Bruijstens, MD, PhD</td>
<td>Department of Neurology, Erasmus MC University Medical Center, Rotterdam, The Netherlands</td>
<td>Major role in the acquisition of data</td>
</tr>
<tr>
<td>Manuela Paunovic, PhD</td>
<td>Department of Neurology, Erasmus MC University Medical Center, Rotterdam, The Netherlands</td>
<td>Major role in the acquisition of data</td>
</tr>
</tbody>
</table>

Continued
# References


Mimics of Autoimmune Encephalitis: Validation of the 2016 Clinical Autoimmune Encephalitis Criteria
Robin W. Van Steenhoven, Juna M. de Vries, Arlette L. Bruijstens, et al.
*Neurol Neuroimmunol Neuroinflamm* 2023;10;
DOI 10.1212/NXI.0000000000200148

This information is current as of August 15, 2023

Updated Information & Services
including high resolution figures, can be found at:
http://nn.neurology.org/content/10/6/e200148.full.html

References
This article cites 50 articles, 6 of which you can access for free at:
http://nn.neurology.org/content/10/6/e200148.full.html##ref-list-1

Subspecialty Collections
This article, along with others on similar topics, appears in the following collection(s):
*All Epilepsy/Seizures*
http://nn.neurology.org/cgi/collection/all_epilepsy_seizures
*Autoimmune diseases*
http://nn.neurology.org/cgi/collection/autoimmune_diseases
*Cohort studies*
http://nn.neurology.org/cgi/collection/cohort_studies
*MRI*
http://nn.neurology.org/cgi/collection/mri

Permissions & Licensing
Information about reproducing this article in parts (figures, tables) or in its entirety can be found online at:
http://nn.neurology.org/misc/about.xhtml#permissions

Reprints
Information about ordering reprints can be found online:
http://nn.neurology.org/misc/addir.xhtml#reprintsus