Abnormal brain metabolism on FDG-PET/CT is a common early finding in autoimmune encephalitis

ABSTRACT

Objective: To compare the rate of abnormal brain metabolism by FDG-PET/CT to other paraclinical findings and to describe brain metabolism patterns in autoimmune encephalitis (AE).

Methods: A retrospective review of clinical data and initial dedicated brain FDG-PET/CT studies for neurology inpatients with AE, per consensus criteria, treated at a single tertiary center over 123 months. Z-score maps of FDG-PET/CT were made using 3-dimensional stereotactic surface projections with comparison to age group-matched controls. Brain region mean Z-scores with magnitudes $\geq 2.00$ were interpreted as significant. Comparisons were made to rates of abnormal initial brain MRI, abnormal initial EEG, and presence of intrathecal inflammation.

Results: Sixty-one patients with AE (32 seropositive) underwent brain FDG-PET/CT at median 4 weeks of symptoms (interquartile range [IQR] 9 weeks) and median 4 days from MRI (IQR 8.5 days). FDG-PET/CT was abnormal in 52 (85%) patients, with 42 (69%) demonstrating only hypometabolism. Isolated hypermetabolism was demonstrated in 2 (3%) patients. Both hypermetabolic and hypometabolic brain regions were noted in 8 (13%) patients. Nine (15%) patients had normal FDG-PET/CT studies. CSF inflammation was evident in 34/55 (62%) patients, whereas initial EEG (17/56, 30%) and MRI (23/57, 40%) were abnormal in fewer. Detection of 2 or more of these paraclinical findings was in weak agreement with abnormal brain FDG-PET/CT ($k = 0.16$, $p = 0.02$).

Conclusions: FDG-PET/CT was more often abnormal than initial EEG, MRI, and CSF studies in neurology inpatients with AE, with brain region hypometabolism the most frequently observed.

Neurol Neuroimmunol Neuroinflamm 2017;4:e352; doi: 10.1212/NXI.0000000000000352

GLOSSARY

AE = autoimmune encephalitis; FDG-PET = 18F-fluorodeoxy-glucose PET; FLAIR = fluid-attenuated inversion recovery; IQR = interquartile range; NMDAR = NMDA receptor; VGKCc = voltage-gated potassium channel-complex.

As early immunotherapy seems to contribute to improved outcomes in autoimmune encephalitis (AE), recent criteria have been proposed to facilitate early diagnosis.\(^1\) 18F-fluorodeoxy-glucose PET (FDG-PET) is only included in criteria for definite autoimmune limbic encephalitis.\(^1\) However, FDG-PET has been recognized as a potentially useful biomarker in suspected AE.\(^2\)–\(^7\) In autoimmune limbic encephalitis, hypermetabolism on FDG-PET in otherwise normal mesial temporal lobe structures by MRI suggests that FDG-PET may be more sensitive than MRI.\(^2\)–\(^4\) Also, particular patterns of metabolism noted by FDG-PET have been identified in certain AE syndromes.\(^3\)–\(^11\) The majority of prior studies of FDG-PET in AE have been limited to qualitative description of FDG-PET findings,\(^2\)\(^,\)\(^3\)\(^,\)\(^5\)\(^,\)\(^6\)\(^,\)\(^10\)\(^,\)\(^12\)–\(^16\) used nondedicated brain FDG-PET studies,\(^8\) have been restricted to specific syndromes,\(^9\)\(^,\)\(^11\)\(^,\)\(^17\)\(^,\)\(^18\) or have made limited comparisons to other diagnostic results incorporated in the current clinical criteria.\(^3\)–\(^6\)

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Funding information and disclosures are provided at the end of the editorial. Go to Neurology.org/nn for full disclosure forms. The Article Processing Charge was funded by the authors.

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We sought to semiquantitatively describe dedicated brain FDG-PET/CT findings for neurology inpatients who met recent AE consensus criteria relative to a database of healthy controls, with comparisons between seronegative and seropositive patients with AE. We also sought to describe the rate of abnormal patterns of brain region metabolism relative to other paraclinical findings in these AE cases as well as prior case series.

METHODS Standard protocol approvals, registrations, and patient consents. This study was approved by the Institutional Review Board of Johns Hopkins University.

Patients. We identified admitted patients with AE who underwent FDG-PET/CT at Johns Hopkins Hospital through the course of admission using the diagnostic terms encephalitis and positron emission tomography (PET) to search the administrative database (December 1, 2005, to March 15, 2016). Patients were cross-referenced with the Johns Hopkins Hospital PET/CT Center database.

Included patients underwent a brain FDG-PET/CT study and had possible or definite AE, including definite limbic encephalitis, per consensus criteria. Diagnostic findings consistent with AE included MRI and EEG abnormalities and the presence of intrathecal inflammation on routine testing. Seropositive patients had a paraneoplastic or cell surface antibody detected in either the serum or the CSF using commercially available assays (Athena Diagnostics, Worcester, MA; Mayo Clinic Laboratories, Rochester, MN).

The electronic medical record was reviewed; data collected were demographic information, clinical history and presentation, diagnostic results, and whether corticosteroids or sedatives were administered within 24 hours preceding FDG-PET/CT study. Clinical MRIs were performed as per the institutional protocol at either 1.5- or 3-T on a Philips (Best, Netherlands), GE Healthcare or Siemens (Erlangen, Germany) scanner. For purposes of this study, T2/fluid-attenuated inversion recovery (FLAIR) signal, diffusion-weighted imaging and apparent diffusion coefficient, and T1 pre- and post-administration of gadolinium sequences were reviewed and rated by each reviewer as consistent or inconsistent with AE, with differences in rating reconciled by discussion between the reviewers.

Statistical methods. The Mann-Whitney U test was used for comparisons of continuous variables. Categorical variables were compared using the χ² test or Fisher exact test, as appropriate. kappa measurement of agreement was performed to assess intermodality agreement of MRI, EEG, and CSF inflammatory markers with brain FDG-PET/CT metabolic patterns. Kappa measurement of agreement was performed for brain FDG-PET/CT metabolic patterns with detection of only 1 or at least 2 diagnostic findings consistent with AE, p < 0.05 was considered significant.

A Friedman test was performed to compare median Z-scores across brain regions for all patients. Serial Wilcoxon rank-sum tests were performed to compare Z-scores between brain regions for all patients with p < 0.008 considered significant after Bonferroni correction. Split-plot analyses of variance with significance of p < 0.008 after Bonferroni correction were performed to compare patterns of FDG-avidity across 6 FDG-PET/CT brain regions within patients and between the seropositive and seronegative AE groups, definite and possible AE groups, as well as for those treated with corticosteroids and those treated with sedatives within 24 hours of brain FDG-PET/CT and those not treated. Comparisons included Z-scores for both left and right hemisphere regions for all patients.

Review of the literature. A PubMed search was performed using positron emission tomography and encephalitis as search terms, updated up to October 27, 2016. Included studies and case series reported brain FDG-PET/CT findings (hypermetabolism and/or hypometabolism) of at least 5 patients with AE or paraneoplastic encephalitis. When provided, reports of abnormalities on brain MRI, EEG, and CSF assays were reviewed.

RESULTS Clinical characteristics of patients with AE undergoing brain FDG-PET/CT. Of the 296 inpatients with the diagnosis of encephalitis, 61 patients met the consensus criteria for AE and underwent brain FDG-PET/CT with studies available for review (table 1). Thirty-two of the 61 (52%) patients had antibodies identified in the serum or CSF, 28 of whom with antibodies with known AE paraneoplastic encephalitis significance, 24/61 (39%) with definite AE antibodies (figure 1). Of the other seropositive patients, 4 were anti–voltage-gated potassium...
channel-complex (VGKCc) seropositive for antibodies different from anti-LGI1 and anti-CASPR2 (3 with supportive CSF, EEG, and/or MRI); 2 were anti-α3 AChR seropositive (1 with supportive EEG); and 2 were anti-striational antibody seropositive (1 with supportive CSF and MRI). Two of the 4 anti-GAD65-seropositive patients had reviewable antibody levels (9,500 and 53,650 U/mL), whereas all had supportive CSF, EEG, and/or MRI.

Seropositive patients were younger than the seronegative patients (median 39 vs 57 years, $p = 0.01$). Durations of symptoms before admission were similar for both groups (median 6 vs 4 weeks, $p = 0.85$). Twelve of the 13 patients were younger than 30 years (5 anti-NMDA receptors [NMDARs]). Fourteen (23%) patients’ CSF were tested for antibodies, with the anti-NMDAR antibody detected in 3 patients, all of whom were negative in the serum. No other

| Table 1  | Clinical characteristics of patients with AE |
|----------|---------------------------------------------|
|          | All (N = 61) | Seropositive (N = 32) | Seronegative (N = 29) | $p$ Value |
| Age, y, median (IQR) | 54 (37) | 39 (44) | 57 (23) | $<0.05^*$ |
| Sex, female, n (%) | 33 (54) | 17 (53) | 16 (55) | 1.00 |
| Race, n (%) | 0.39 |
| White | 37 (61) | 17 (53) | 20 (69) |
| Black | 9 (15) | 5 (16) | 4 (14) |
| Other | 15 (24) | 10 (31) | 5 (17) |
| History of cancer, n (%) | 11 (18) | 3 (9) | 8 (28) | 0.09 |
| Lymphoma | 5 | 0 | 5 |
| Breast | 3 | 2 | 1 |
| Testicular | 1 | 1 | 0 |
| Meningioma | 1 | 0 | 1 |
| Prostate and renal cell | 1 | 0 | 1 |
| Diagnosed with cancer during admission, n (%) | 6 (10) | 4 (13) | 2 (7) | 0.67 |
| Small cell lung | 3 | 3 | 0 |
| Breast | 1 | 0 | 1 |
| Ovarian teratoma | 1 | 0 | 1 |
| Seminoma | 1 | 1 | 0 |
| Duration of neurologic symptoms before admission, wk, median (IQR) | 4 (7.5) | 6 (10) | 4 (8) | 0.85 |
| Neurologic signs and symptoms on admission, n (%) | 0.13 |
| Lethargy | 47 (77) | 22 (69) | 25 (86) |
| Short-term memory impairment | 46 (75) | 25 (78) | 21 (72) | 0.77 |
| Hallucinations | 5 (8) | 4 (13) | 1 (3) | 0.36 |
| Cerebellar signs | 47 (77) | 21 (66) | 26 (90) | $<0.05^*$ |
| Focal weakness | 37 (61) | 17 (53) | 20 (69) | 0.29 |
| Focal numbness | 35 (57) | 15 (47) | 20 (69) | 0.12 |
| Movement disorder | 39 (64) | 19 (59) | 20 (69) | 0.59 |
| Seizures | 25 (41) | 16 (50) | 9 (31) | 0.19 |
| Status epilepticus | 10 (16) | 4 (13) | 6 (21) | 0.50 |
| Cranial neuropathy | 14 (23) | 8 (25) | 6 (21) | 0.77 |
| Aphasia | 25 (41) | 12 (38) | 13 (45) | 0.61 |
| Psychiatric symptoms | 22 (36) | 15 (47) | 7 (24) | 0.11 |
| Focal neurologic findings on admission | 58 (95) | 30 (94) | 28 (97) | 1.00 |
| Multiple focal neurologic findings on admission | 50 (82) | 22 (69) | 28 (97) | $<0.01^*$ |

Abbreviations: AE = autoimmune encephalitis; IQR = interquartile range.
Clinical characteristics of patients with AE who underwent brain FDG-PET/CT through the course of inpatient evaluation.

$^*$Significant.
antibodies were detected in tested CSF samples. Of the 17 patients with a history or subsequent diagnosis of cancer, 7 were found to be seropositive: small cell lung cancer (1 anti-GAD65, 1 anti-Hu, and 1 anti-CV2), breast (1 anti-NMDAR and 1 anti-LGI1), seminoma (1 anti-Ma2), and testicular cancer (1 anti-Ma2).

Other paraclinical findings. Routine CSF studies were consistent with intrathecal inflammation in 34/55 patients (62%, figure e-1 at Neurology.org/nn). Initial EEG for 17/56 (30%) patients demonstrated temporal area slowing, epileptiform discharges, or seizures consistent with the diagnosis of AE. This was more frequently observed among patients younger than 30 years (8/13, 62%) than others (9/43, 21%, \( p = 0.01 \), table e-1). Brain MRI studies for 57 patients were available for blinded review, and 23 (40%) were consistent with the diagnosis of AE. No differences were observed across antibody status, antibody class, or AE classification (table e-1). Fifty-one of the 61 patients (84%) had at least 1 paraclinical finding consistent with AE on routine CSF analysis, brain MRI, or EEG. Thirty-one patients (51%) had only 1 paraclinical finding; 17 patients (28%) had 2 findings; and 3 patients (5%) had 3 findings consistent with AE. Ten patients were included based on clinical criteria, 4 of whom had definite AE based on detected antibodies, and 3 seropositive for other antibodies (table e-2).

Brain FDG-PET/CT findings. Brain FDG-PET/CT was performed a median of 4 weeks after symptom onset (interquartile range [IQR] 9 weeks) and a median of 4 days (IQR 8.5 days) from brain MRI. Brain FDG-PET/CT was abnormal in 52/61 patients (85%, figure e-1) when compared with the healthy control database. FDG-PET/CT demonstrated brain region hypometabolism alone in 42/61 (69%), hypermetabolism alone in 2/61 (3%) patients, and regions of abnormal hypometabolism and abnormal hypermetabolism in 8/61 (13%) of patients (figure e-1, table e-3). No differences were observed across age group, antibody status, antibody class, or AE classification (table e-1). No difference in proportion with abnormal metabolism was noted between those evaluated by FDG-PET/CT within 4 weeks of symptoms (27/31 [87%]) and those evaluated later (25/30 [83%], \( p = 0.73 \)).

Across brain regions in patients with AE, metabolism was greater for the caudate (\( 2.17, \text{IQR} 2.43 \)) relative to the frontal (\( -2.24, \text{IQR} 2.69, p < 0.005 \)), temporal (\( -1.80, \text{IQR} 1.77, p = 0.002 \)), parietal (\( -2.49, \text{IQR} 1.61, p < 0.005 \)), and occipital (\( -2.09, \text{IQR} 2, p < 0.005 \)) brain regions (\( p < 0.005 \), figure 2A). Brain region metabolism patterns did not vary between seropositive and seronegative AE patient groups (\( F(1,120) = 3.18, p = 0.08, \eta^2_p = 0.03, \text{figure 2B} \)) nor definite AE and possible AE (\( F(1,120) = 2.69, p = 0.10, \eta^2_p = 0.02 \)). Similarly, brain region metabolism patterns did not vary between those treated with corticosteroids (\( F(1,120) = 0.200, p = 0.656, \eta^2_p = 0.002 \)) and those treated with
sedatives ($F(1120) = 1.95, p = 0.165, \eta_p^2 = 0.016$) within 24 hours of brain FDG-PET/CT and those not treated.

Concordance between FDG-PET/CT and other paraclinical findings. The finding of an abnormal metabolic pattern on brain FDG-PET/CT was not in agreement with the presence of CSF inflammation on routine assessment (table e-4). By contrast, brain MRI findings consistent with AE were in weak agreement with the finding of abnormal metabolism on brain FDG-PET/CT ($\kappa = 0.17, p < 0.05$), most notably with hypometabolism ($\kappa = 0.25, p < 0.05$; table e-4, figure 3). In general, the presence of any FDG-PET/CT abnormality was not in agreement with the presence of EEG findings consistent with AE (table e-4). However, detection of EEG findings consistent with the diagnosis of AE was in weak agreement with detection of brain region hypermetabolism ($\kappa = 0.16, p < 0.05$) and in fair agreement with having regions of both hypermetabolism and hypometabolism in the same FDG-PET/CT study ($\kappa = 0.26, p < 0.05$, table e-4).

Detection of at least 1 paraclinical finding consistent with AE was not in agreement with detection of abnormal metabolism by FDG-PET/CT ($\kappa = 0.16, p = 0.02$; table e-4).

Literature review. Fourteen studies were identified which met the inclusion criteria (table 2). Of the 139 FDG-PET studies reported, 120 (86%) were abnormal, with 55 (40%) demonstrating both hypometabolism and hypermetabolism, 30 (22%) demonstrating only hypometabolism, and 35 (25%) demonstrating only hypermetabolism. This is compared with the sum report of 38/75 (51%) EEGs, 68/114 (60%) brain MRI, and 45/86 (52%) routine CSF studies consistent with the diagnosis of possible AE.

DISCUSSION Here, we describe dedicated semi-quantitative brain FDG-PET/CT findings among patients meeting the consensus AE criteria. Dedicated brain FDG-PET/CT was abnormal in 85% of patients with AE, and FDG-PET abnormalities were more sensitive for AE compared with EEG, MRI, or routine CSF findings. Although brain region hypometabolism was most commonly noted, some studies demonstrated areas of both hyper- and hypometabolism and a minority demonstrated hypermetabolism alone. The combination of abnormalities in at least 2 of the 3 other paraclinical tests (routine CSF studies, brain MRI, and EEG) was in fair agreement with abnormal findings on dedicated brain
FDG-PET/CT. Our results suggest that brain FDG-PET/CT may be helpful in supporting evidence of brain dysfunction in suspected patients with AE. Brain region hypometabolism in multiple regions likely reflects widespread impairment of neuronal activity in AE. Whether such hypometabolism results from functional changes, structural changes, or a combination of both is not yet clear. Many of the areas of regional hypometabolism did not have correlates on MRI, suggesting the possibility of neuronal dysfunction in the absence of structural disturbance. Longitudinal studies will be needed to clarify whether the observed hypometabolism in AE is reversible. Moreover, although widespread regional hypometabolism was observed across various AE syndromes, there are likely syndrome-specific patterns of brain region metabolism.

Previous series primarily report hypermetabolism in AE. These series contain larger proportions of patients with anti-NMDAR (36/130 reported patients) or patients with anti-LGI1, anti-CASPR2, or anti-VGKCc antibodies (39/130) than our cohort, potentially limiting their generalizability to other seropositive and seronegative AE. We observed brain region hypermetabolism in a subset of patients, many of whom had anti-NMDAR or...
| Reference no. | No. of PET studies/patients in series | Comparison to control population performed? | Serum and/or CSF antibody status (N) | PET demonstrated hypometabolism, N | PET demonstrated hypermetabolism, N | PET demonstrated both hyper-/hypometabolism, N | EEG consistent with AE/EEG performed | MRI consistent with AE/MRI performed | CSF inflammation demonstrated/lumbar punctures performed |
|--------------|-------------------------------------|-----------------------------------------------|-----------------------------------|----------------------------------|----------------------------------|--------------------------------------|---------------------------------|---------------------------------|---------------------------------|
| 2            | 7/7                                 | No                                            | Seronegative (7)                  | 6/7                              | 1                                | 4                                    | 1                               | 2/7                             | 6/7                             |
| 3            | 9/9                                 | No                                            | NMDAR (3), seronegative (6)       | 9/9                              | 3                                | 6                                    |                                 |                                 | 7/9                             |
| 4            | 18/18                               | Yes                                           | Hu (2), Ri (1), GAD65 (1), LGI1 (1), CASPR2 (2), VGKCα (3), NMDAR (2), NMDAR/VGKCα (1), seronegative (5) | 13/18                           | 8                                | 5                                    | 10/18                           | 10/18                           |
| 5            | 13/10                               | No                                            | Hu (2), VGKCα (2), NMDAR (1), "Neuronal Cell Membrane” [2], “Atypical” (1), seronegative (2) | 12/13                           | 1                                | 6                                    | 5                               | 15/17                           |
| 6            | 12/16                               | No                                            | LGI1 (1), VGKCα (1), NMDAR (2), GAD65 (2), Neuropil (1), Ma2 (1), Ma2/Hu (1), nontype (1), seronegative (6) | 11/12                           | 9                                | 1                                    | 1                               | 10/16                           | 10/16                           |
| 8            | 6/6                                 | Yes                                           | NMDAR (6)                         | 6/6                              | 3                                | 3                                    |                                  |                                 | 2/6                             |
| 9            | 10/10                               | Yes                                           | NMDAR (6), LG1 (4)               | 10/10                            | 10                               | 0/10                                 | 2/10                            | 6/10                            |
| 10           | 13/6                                | No                                            | NMDAR (6)                         | 12/13                            | 4                                | 2                                    | 6                               | 6/6                             | 2/6                             | 6/6                             |
| 11           | 18/8                                | Yes                                           | NMDAR (8)                         | 14/18                            | 1                                | 1                                    | 12                              |                                 |                                 |
| 12           | 6/8                                 | No                                            | LGI1 (5), CASPR2 (2), CASPR2/LGI1 (1) | 3/6                              | 1                                | 2                                    | 1/8                             | 6/8                             | 2/8                             |
| 13           | 6/6                                 | No                                            | Seronegative (6)                  | 6/6                              | 1                                | 5                                    |                                 |                                 | 3/5                             |
| 14           | 5/5                                 | No                                            | GABA(B) (5)                       | 3/5                              | 1                                | 2                                    | 3/5                             | 2/5                             | 2/5                             |
| 15           | 10/14                               | No                                            | LGI1 (10)                        | 9/10                             | 8                                | 1                                    | 10/14                           | 10/14                           | 1/13                            |
| 16           | 6/7                                 | No                                            | VGKCα (6), NMDAR (1)             | 6/6                              | 6                                | 6                                    | 7/6                             | 3/7                             | 2/7                             |
| **Total**    | 120/139                             |                                               |                                   | 30                               | 35                               | 55                                   | 38/75                           | 68/114                          | 49/86                           |

Abbreviations: AE — autoimmune encephalitis; CASPR2 — contactin-associated protein-2; GAD65 — 65 KDa glutamic acid decarboxylase; LGI1 — leucine-rich inactivated 1 protein; NMDAR — NMDA receptor; VGKC — voltage-gated potassium channel.

Systematic review of case series reporting FDG-PET findings in AE along with the available EEG, MRI, and CSF study reports.2–6,8–16

*Only anti-VGKC seropositivity reported and potentially includes those seropositive for anti-LGI1, anti-CASPR2, or other VGKC-complex antibodies.
anti-VGKCc encephalitis, compatible with previous literature.5,8–12,15

The current AE consensus criteria only include FDG-PET findings in the criteria for definite autoimmune limbic encephalitis. Bilateral FLAIR/T2 abnormalities of the medial temporal lobes are required, and in the absence of such findings, FDG-PET hypermetabolism in the medial temporal lobes may meet this requirement. Observations provided here suggest that AE may lead to broader metabolic abnormalities detectable by FDG-PET outside the confines of the medial temporal lobes and these may inform future FDG-PET AE criteria.

Concerns raised regarding the incorporation of FDG-PET in the evaluation of patients with AE include availability of FDG-PET imaging modalities in urgent clinical situations. Moreover, as a newer modality, further work is needed to validate it as a method in the diagnosis of AE.24 Worldwide, FDG-PET/CT represents one of the medical imaging modalities with the largest growth in terms of number of scanners.25 In addition, FDG-PET/CT has been found to be diagnostically superior to other conventional imaging modalities in other clinical settings, and it has demonstrated cost-effectiveness in settings such as non–small lung cancer staging.25 FDG-PET also plays an important role in screening for occult malignancy in paraneoplastic syndromes, including encephalitis.26 Thus, FDG-PET is likely to become an increasingly used modality in the evaluation of patients with suspected AE beyond occult malignancy screening. Many institutions use a “vertex to toe” field of view for their whole-body protocols. The addition of a 10-minute dedicated 3D PET acquisition of the brain requires no extra radiopharmaceutical administration, is easily incorporated in conventional clinical workflows, provides increased statistical quality in comparison with “vertex to toe” imaging, and allows for higher-resolution images with more robust quantitation. As the utility of FDG-PET is evaluated further, collaborative evaluation by neurologists and radiologists will be necessary for careful characterization and correlation of syndromes with associated imaging findings, with comparisons to healthy and other neurologic patient populations.

A major limitation of this study is that it is retrospective, involving all patients meeting the criteria for AE who underwent FDG-PET/CT at a single tertiary medical center with associated selection bias. Although performed at a single center, it benefits from the consensus inclusion criteria for AE and uniformity of PET equipment, protocols, and analyses. Also, the observations reported here were limited to those patients admitted to the hospital for onset of symptoms of 3 months or less and do not include findings for those with longer duration of symptoms. Future prospective studies involving serial FDG-PET studies may help clarify the specificity and evolution of patterns of metabolism through the phases of encephalitis, as has been suggested in cases series of specific encephalitides such as anti-NMDAR encephalitis.11 Our study included the initial FDG-PET studies for patients regardless of antibody status, and future larger prospective studies of specific antibody syndromes may further clarify patterns of abnormality, pattern associations with clinical status, and pattern changes in the setting of immune therapy as has been observed in cases of autoimmune dementia.27 Not all patients underwent CSF antibody testing, which may be more sensitive, and thus we may underestimate the number of seropositive patients. However, there was no difference noted in brain region metabolism between seropositive and seronegative groups. Also, 4 patients were anti-VGKCc seropositive without further specification, and although 3 had other findings supportive of AE, the clinical value of such antibodies is unknown and cautious interpretation is advised. One-third of patients studied here were treated with either corticosteroids or sedatives within 24 hours of FDG-PET/CT. Although both corticosteroids and sedatives have been reported to decrease cortical metabolism,19,20 no differences in brain region metabolism were noted between patients exposed and unexposed to these medications before FDG-PET/CT. In addition, FDG-PET/CT metabolism patterns for patients with AE were not compared with other patients with neurologic diseases (such as infectious encephalitis); psychiatric diseases; intoxications; or other syndromes which may also have abnormal FDG-PET findings.21,22,28–31 It will be important for future prospective studies to incorporate patients with other neurologic, psychiatric, and medical diseases to assess the specificity of metabolic findings by FDG-PET described here. Finally, the CortexID control population used for comparison ranges from 30 to 85 years. The 13 patients younger than 30 years studied had a similar rate of abnormal brain region metabolism compared with those older. Ideally, a concurrent age- and sex-matched control population could be used for direct comparison, although such data collection is limited by the radiation exposure to otherwise normal patients.

Here, brain FDG-PET/CT was commonly abnormal in AE, most often demonstrating brain region hypometabolism. The frequency of metabolic abnormalities was greater than that of diagnostic studies currently included in consensus criteria for the diagnosis of AE. Overall, FDG-PET/CT may represent a sensitive and early biomarker for AE and could play a complementary role to currently proposed tests in the diagnosis of AE. Future prospective studies may further clarify the role FDG-PET may play in the
diagnosis and monitoring of AE in general and specific antibody syndromes in particular.

AUTHOR CONTRIBUTIONS
Dr. Probasco: design and conceptualization of the study, analysis and interpretation of the data, and drafting and revising of the manuscript. Dr. Solnes: design and conceptualization of the study, analysis and interpretation of the data, and revising of the manuscript. Mr. Nalluri: analysis and interpretation of data. Mr. Cohen, Dr. Jones, and Dr. Zan: analysis and interpretation of the data and revising of the manuscript. Dr. Javadi and Dr. Venkatesan: design and conceptualization of the study, analysis and interpretation of the data, and revising of the manuscript.

STUDY FUNDING
No targeted funding reported.

DISCLOSURE
J.C. Probasco serves on the editorial board for The Neurohospitalist, is an associate editor for The Neurohospitalist, and is editor-in-chief for NEJM Journal Watch Neurology. L. Solnes, A. Nalluri, J. Cohen, K.M. Jones, E. Zan, and M.S. Javadi report no disclosures. A. Venkatesan received speaker honoraria from Almirall, served as a medical expert for U.S. Journal Watch Neurology, and served as medical editor for Carnival Cruise Lines. Go to Neurology.org/nn for full disclosure forms.

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Received January 23, 2017. Accepted in final form March 27, 2017.

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*Neurol Neuroimmunol Neuroinflamm* 2017;4;
DOI 10.1212/NXI.0000000000000352

This information is current as of May 11, 2017
