### Incomplete Susac Syndrome Exacerbated After Natalizumab

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A 66-year-old man with hypertension presented with right-sided numbness and slurred speech. MRI revealed a left frontoparietal area of T2-weighted hyperintensity with focus of central enhancement (figure, A and B). The patient’s symptoms self-resolved within weeks. Over the next 2 years, he experienced multiple relapses including encephalopathy characterized by headache with motor aphasia and sudden incomplete hearing loss. All symptoms except hearing loss were transient, improved with IV methylprednisolone, and were accompanied by new T2 lesions involving the periventricular white matter, corpus callosum, and subcortical white matter, with some demonstrating partial enhancement (figure, C and D).

Extensive workup revealed only anti-endothelial cell antibodies (AECA) (1:50, indirect immunofluorescence, Cornell Dermatopathology Laboratory, New York). CSF analysis performed during remission was unrevealing; oligoclonal bands were absent. Fluorescein angiogram and optical coherence tomography did not show any abnormalities.

The patient was presumed to have multiple sclerosis (MS) and started on natalizumab. Within days of the first infusion, he developed motor aphasia, with MRI revealing extensive T2-hyperintense lesions with corresponding leptomeningeal enhancement (figure, G and H). Repeat CSF analysis revealed elevated protein of 116 mg/dL. Natalizumab antibodies were negative. The patient underwent brain biopsy, which revealed predominantly perivascular demyelination (figure, I) with relative preservation of axons, although axons were focally distended and showed evidence of axonal injury (figure, M) suggestive of demyelinating etiology. In addition, there was evidence of focal microangiopathy. Scattered vessels contained perivascular CD3+ CD8+ T lymphocytes whereas other vessels showed endothelial cell detachment and early fibrin deposition suggestive of an acute endothelial injury. CD68+ and CD163+ macrophages were present in the perivascular space (figure, J). Vascular deposits of C3d (figure, K) with lesser but still discernible deposits of C4d and C5b-9 were noted. There was focal red cell extravasation and hemosiderin deposition along with focal basement membrane zone reduplication compatible with antecedent episodes of vascular injury (figure, L). Glial fibrillary acidic protein stain showed massive reactive gliosis. There was no histopathologic evidence of infection or lymphoma. Overall morphologic features were those of perivenular demyelination and complement-induced endothelial injury.

**Discussion.** Atypical age at onset, episodes of encephalopathy and hearing loss, unusual elevated protein in CSF, extensive leptomeningeal involvement, and remarkable disease exacerbation after natalizumab suggest a diagnosis of Susac syndrome rather than MS. Susac syndrome is presumed to be an autoimmune disorder with microvascular occlusions in the brain, retina, and cochlea.1 The triad of encephalopathy, branch retinal artery occlusions, and hearing deficits is seen at some point in the disease in 85% of cases, but only 13% of patients manifest all these symptoms on presentation.2 Our patient exhibited 2 core features of Susac syndrome. Further findings supporting the diagnosis on MRI include preferential involvement of the central portion of the corpus callosum (known as “snowball” lesions; figure, E), central callosal holes (figure, F), and extensive leptomeningeal enhancement (figure, G and H), reported in 30% of patients with Susac syndrome. The low titers of AECA seen in our patient are suggestive of Susac syndrome, but this finding is not specific.1 Treatments for Susac syndrome are based solely on clinical experience; steroids in conjunction with IV immunoglobulin (IVIg) and/or cyclophosphamide are the most common agents used.1 Our patient has been treated with IVIg for the last 14 months and has had no evidence of clinical or radiologic relapses thus far.

Reports of brain biopsy are rare. Pathologic findings include “focal microangiopathic and/or gliotic changes” and T cell perivascular inflammation with CD8 cell predominance, seen in our case as well.1,4 Some investigators reported evidence of antibody-mediated endothelial cell injury syndrome/endothelial cell necrosis in concert with C3d and C5b-9 deposition within the capillaries and venules of the brain, although this may not be specific because it was also found in some controls.2,4 Our patient’s biopsy also showed findings suggestive of microvascular injury (figure, L) and complement deposition (figure, K),
Fluid-attenuated inversion recovery (FLAIR) sequence with a single large T2-hyperintense signal involving the left frontoparietal lobe (A) with gadolinium-enhanced (Gd\textsuperscript{1}) T1-weighted imaging (T1WI) demonstrating a small central area of contrast enhancement (B). FLAIR sequence demonstrating ongoing new T2 lesions involving bilateral hemispheres (C) with evidence of blood-brain barrier opening, with Gd + T1WI again corresponding to the left parietal lobe lesion (D). Preferential involvement of the central portion of the corpus callosum known as “snowball” lesions (E) as well as a central callosal hole (F). FLAIR sequence demonstrating extensive T2-hyperintense lesions (G) with corresponding postcontrast T1WI confirming leptomeningeal enhancement along the left parietal hemisphere (H) days after the patient’s first natalizumab infusion. (I) Low-power Luxol fast blue (LFB) hematoxylin & eosin (H&E) stain demonstrates white matter with patchy myelin loss. Myelin loss was most prominent in the perivascular white matter, with greater perivenular than periarterial involvement. (J) High-power view of perivascular macrophages also contained small particles of LFB\textsuperscript{–} and periodic acid-Schiff-positive material consistent with the myelin accumulation, which can be seen in demyelinating disorders. (K) Vascular deposits of C3d. (L) A venule infiltrated by macrophages and lymphocytes. There is evidence of vascular compromise characterized by marked red cell extravasation. (M) Neurofilament stain, an axonal marker, showed relative preservation of axons; however, axons were focally distended and showed evidence of axonal injury (yellow arrowheads).
consistent with the endothelial cell injury hypothesis. However, we also observed prominent myelin loss (figure, I), which has not been previously reported in Susac syndrome. It is possible that demyelination in our patient was consequent to microvascular injury, or it may be an unrelated process.

Increased relapse frequency after the start of immunomodulatory drugs effective in MS, such as alemtuzumab, interferon β, and natalizumab, in patients with “atypical” inflammatory diseases of the CNS has been described.5–7 In our case, there was a dramatic temporal relationship between the initiation of natalizumab and disease exacerbation. It is possible that natalizumab altered the balance of the immune system by decreasing circulating T regulatory cells with a subsequent increase in the number of proinflammatory cytokine-producing T cells or blocked egress of regulatory natural killer cells from the periphery into the brain, hence precipitating the relapse.5 Another hypothesis is that natalizumab, being a monoclonal antibody, may be causing activation of the complement and thus inducing the relapse, although no supporting evidence was found in the literature.

Our case highlights the difficulty in diagnosing atypical relapsing idiopathic inflammatory disease of the CNS, even with the help of a brain biopsy, and demonstrates the risk of treating these conditions with MS-specific immune-modulating drugs.

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