Susceptibility-weighted imaging (SWI) is a new imaging method that maximizes sensitivity to magnetic susceptibility effects and is clinically useful for evaluating trauma and vascular malformations. However, the role of SWI in other neuropathology has not been extensively evaluated. SWI is more sensitive than conventional gradient-echo (GE) techniques for the detection of blood products. Because GE imaging is a well-recognized method of detecting changes indicative of cerebral amyloid angiopathy (CAA), SWI should be as sensitive as well. This is the first reported evaluation of SWI in a patient with suspected CAA showing marked increased sensitivity to the detection of microhemorrhages compared with conventional GE sequences.

CAA is a vascular disease often associated with stroke, especially recurrent hemorrhagic stroke. However, its presentation is varied; depending upon severity, it may not be diagnosed until autopsy. It is not a rare entity; autopsy studies show moderate or severe CAA with an age-dependent prevalence of 8% (for ages 75–84) to 12% (for ages >84), and an even higher prevalence in patients with Alzheimer disease.

Case Report
A 70-year-old man with a past medical history notable for diabetes, hypertension, and ischemic heart disease was referred for evaluation of recurrent cerebral hemorrhages. The patient had experienced a left temporoparietal cerebral hemorrhage approximately 10 years before presentation. Six months before arrival at our hospital, the patient presented with speech difficulty; subsequent brain MR showed a 3.1 × 2.7-cm left temporooccipital lobe hemorrhage with a small zone of surrounding edema. Two months before presentation, the patient underwent coronary artery stent placement and was placed on aspirin and clopidogrel. Within a week, he presented with altered mental status, and a CT without contrast showed a new 2.5-cm intraparenchymal hemorrhage in the left posterior temporoparietal lobe. Because of recurrent hemorrhages in the same location, diagnostic considerations included vascular malformation and amyloid angiopathy. Before MR imaging, the patient underwent cerebral angiography, which demonstrated no evidence of arteriovenous malformation or any pathologic vascular condition. MR angiography (MRA) showed similar negative findings. Brain MR demonstrated ependymomalacia within the left temporoparietal lobe, consistent with the patient’s history of prior repeated hemorrhages. GE and SWI images showed low signal intensity within the left temporoparietal lobe region, consistent with hemosiderin deposition. Both the T2*-weighted GE (Fig 1, left) and SWI image (Fig 1, right) show multiple low-signal intensity areas in a cortical-subcortical distribution, consistent with hemosiderin deposition. However, SWI shows the susceptible foci to be both more numerous and with improved apparent contrast compared with the corresponding conventional GE image. Using the same criteria for counting the small rounded lesions, the SWI projected image shows 201 lesions, whereas the GE-projected image shows 74 lesions. Aggregate imaging findings are consistent with old infarct and highly suggestive of CAA.

Discussion
CAA is well known to manifest as amyloid protein deposits in small arteries. Although CAA is unrelated to systemic amyloidosis, it is associated with increasing age, dementia, Alzheimer disease, dementia pugilistica, postradiation necrosis, and spongiform encephalopathies. No CT- or MR-based imaging technique currently exists to directly visualize or quantify amyloid deposits; diagnosis can only be made by biopsy or autopsy. Pathologic specimens show fibrinoid degeneration and microaneurysms; amyloid stains with Congo red dye exhibit yellow-green birefringence when exposed to polarized light. As the β-amyloid protein builds up within the elastic lamina of vessel walls, vessels lose elasticity and become fragile. Fragile vessels are more easily damaged, and tiny hemorrhages (cerebral microbleeds) occur in and around the arterial vessel wall. Microbleed-induced damage can cause a further loss of vessel wall elasticity, thinning of the vessel wall, or vessel wall dilation. With regard to morbidity and mortality, the most concerning clinical effect of CAA is spontaneous intracranial hemorrhage, which can be recurrent. Cognitive impairment is also a common feature of CAA; 40% of patients with CAA-related intracranial hemorrhage have dementia, and more than 80% of patients with Alzheimer disease have CAA.

Certain patterns of lobar hemorrhage, which can be detected on CT and conventional MR images, are suggestive of CAA. In addition, CAA should be suggested if lobar hemor-
Hemorrhage is identified with a superficial location and cortical involvement. Hemorrhages that involve the cortex and subcortical white matter within the frontal and parietal lobes are most common. Evidence of multiple hemorrhages confined to lobar regions may also be present, and subarachnoid extension of hematomata is more likely in CAA. CAA is a vascular disease that is insensitive to detection by MR angiography, conventional angiography, and digital subtraction angiography. Neither CT nor conventional MR can detect microhemorrhages. Although T2* GE sequences can detect microhemorrhages, 25% of patients ultimately diagnosed with CAA have no microhemorrhages on conventional T2* GE images.

SWI is a 3D, velocity-compensated, GE sequence that combines both magnitude information (used in conventional MR images, including T2* GE) with phase information to accentuate the visibility of susceptible foci (such as small veins and hemorrhage). Microhemorrhages contain hemosiderin, which is paramagnetic relative to normal tissue and leads to large variations in local magnetic fields and a local reduction in T2*. The signal intensity loss is proportional to the amount of hemosiderin present. SWI, with its unique sensitivity to blood products and hemorrhage, is well-suited to detect imaging changes consistent with CAA as evidenced in Fig 1.

SWI imaging, in this case, detected far more microhemorrhages than the conventional T2* GE technique. Conventional T2* techniques underestimate microhemorrhages compared with autopsy, and only 75% of patients with amyloid and lobar hemorrhages show microhemorrhages on T2*-weighted sequences. SWI is a much more sensitive technique than conventional T2* imaging, making it an important method of improving initial diagnosis. Increased sensitivity may allow assessment of the rate of microhemorrhage development or regression, allowing more precise analysis of the natural history of the disease, or assessing response to therapy. Early recognition can be advantageous to patients on anticoagulant or aspirin therapy in that they are at increased risk for subsequent and possibly fatal hemorrhage (reported to be as high as 38%, with a 44% mortality rate). In our case, antithrombotic agents were discontinued because the risk of hemorrhage was determined to be greater for this patient than the risk of thromboembolic disease. Strict blood pressure control may also be of utility in patients with CAA-related hemorrhage.

Conclusions
Imaging the brain with SWI requires only 5 to 7 minutes, and can easily be included in routine neuroimaging protocols. SWI identified many more microhemorrhages than conventional T2*-weighted GE magnitude techniques and may lead to earlier diagnosis of patients with CAA. Improved detection of microhemorrhage may become increasingly important in the diagnosis, management, and monitoring of the therapeutic response of patients with CAA, especially as new therapeutic options, such as low-molecular-weight proteins that reduce amyloid fibril formation, become available.

References