MRI Features of Intracerebral Hemorrhage Within 2 Hours From Symptom Onset

Italo Linfante, MD; Rafael H. Llinas, MD; Louis R. Caplan, MD; Steven Warach, MD, PhD

Background and Purpose—MRI has been increasingly used in the evaluation of acute stroke patients. However, MRI must be able to detect early hemorrhage to be the only imaging screen used before treatment such as thrombolysis. Susceptibility-weighted imaging, an echo-planar T2* sequence, can show intracerebral hemorrhage (ICH) in patients imaged between 2.5 and 5 hours from symptom onset. It is unknown whether MRI can detect ICH earlier than 2.5 hours. We describe 5 patients with ICH who had MRI between 23 and 120 minutes from symptom onset and propose diagnostic patterns of evolution of hyperacute ICH on MRI.

Methods—As part of our acute imaging protocol, all patients with acute stroke within 24 hours from symptom onset were imaged with a set of sequences that included susceptibility-weighted imaging, diffusion- and perfusion-weighted imaging, T1- and T2-weighted imaging, fluid-attenuated inversion recovery (FLAIR), and MR angiography using echo-planar techniques. Five patients with ICH had MRI between 23 and 120 minutes from the onset of symptoms.

Results—ICH was identified in all patients. Distinctive patterns of hyperacute ICH and absence of signs of ischemic stroke were the hallmark features of this diagnosis. The hyperacute hematoma appears to be composed of 3 distinct areas: (1) center: isointense to hyperintense heterogeneous signal on susceptibility-weighted and T2-weighted imaging; (2) periphery: hypointense (susceptibility effect) on susceptibility-weighted and T2-weighted imaging; and (3) rim: hypointense on T1-weighted imaging and hyperintense on T2-weighted imaging, representing vasogenic edema encasing the hematoma.

Conclusions—MRI is able to detect hyperacute ICH and show a pattern of evolution of the hematoma within 2 hours from the onset of symptoms. (Stroke. 1999;30:2263-2267.)

Key Words: hemoglobins ■ intracerebral hemorrhage ■ magnetic resonance imaging ■ stroke ■ tissue plasminogen activator
(FLAIR), T1-weighted imaging (T1WI), T2-weighted imaging (T2WI), MR angiography (MRA), and perfusion-weighted imaging (PWI). Over the last 2-year period, the diagnosis of ICH was made in 5 patients imaged in <2.5 hours from symptom onset. The diagnosis was based on clinical and neuroimaging evaluation by the stroke fellow, the stroke attending physician, and the neuroradiologist. Investigations were ordered by the treating physician. CT scans were not routinely performed in patients who had acute MR studies. Moreover, since the present study was not a comparative study of MRI versus CT, CT scans were only obtained in 2 patients 7 and 9 hours from symptom onset (patients 1 and 3).

**Data Acquisition**

MRI studies were performed on a Siemens Vision 1.5 scanner (Siemens Medical System). All equipment was approved by the Food and Drug Administration for use in patients, and scans were obtained as part of the patients’ routine diagnostic workup. The pulse sequences and typical parameters were as follows: (1) turbo fast low-angle shot (FLASH) T1-weighted localized repetition time (TR), 15 ms; echo time (TE), 6 ms; flip angle, 30°; 3 slices; 3 planes; slice thickness, 8 mm; matrix size, 128×256; field of view (FOV), 300 mm; acquisition time, 10 seconds; (2) DWI, EPI spin echo: TR, 6,000 ms; TE, 99 ms; 18 slices; slice thickness, 7 mm; matrix size, 96×128; FOV, 240 mm; b value, 1000 s/mm²; acquisition time, 30 seconds; (3) SWI, EPI gradient echo: TR, 0.8 ms; TE, 60 ms; flip angle, 60°; 20 slices; slice thickness, 7 mm; matrix size, 96×128; FOV, 240; acquisition time, 2 seconds; (4) T1WI: TR, 600 ms; TE, 12; flip angle, 90°; 20 slices; slice thickness, 5 mm; matrix size, 120×256; FOV, 240; acquisition time, 46 seconds; (5) turbo spin-echo T2/proton density (T2WI): TR, 3800 ms; TE, 22/90 ms; flip angle, 180°; 20 slices; slice thickness, 5 mm; matrix size, 120×256; FOV, 240 mm; acquisition time, 1 minute and 42 seconds; (6) EPI FLAIR: TR, 0.8 ms; TE, 53 ms; flip angle, 90°; 18 slices; slice thickness, 7 mm; matrix size, 84×128; FOV, 240 mm; acquisition time, 4 seconds; and (7) PWI, EPI gradient echo: TR, 0.8 ms; TE, 47 ms; flip angle, 60°; 12 slices; slice thickness, 7 mm; matrix size, 96×128; FOV, 240; acquisition time, 40 seconds, 1 second per set (40 sets).

**Patients**

Patient 1 is a 66-year-old man with hypertension and thrombocytopenia secondary to cyclosporin treatment. At 1:45 AM he had acute onset of right-sided weakness and right hemisensory loss. On examination he was awake, alert, and oriented, and he responded to questions appropriately. He had slight lower face weakness and slight weakness and numbness of the right arm and hand. National Institutes of Health Stroke Scale (NIHSS) score was 4. In the emergency department at 4:40 AM, he suddenly vomited, became mute, developed left eye deviation, lower face weakness, right hemiplegia, and was difficult to arouse. NIHSS score was 22. The scan started at 5:03 AM, 23 minutes from the sudden deterioration. MRI showed a left thalamic bleed with extension to the subcortical white matter. CT obtained 7 hours from symptom onset showed a hyperdense area of the same size and location. The patient’s neurological examination was unchanged when the CT was obtained.

Patient 4 is a 36-year-old woman who had had a cesarean section 10 days previously. At 7:45 AM she fell down with slurred speech and left-sided weakness. Blood pressure was 122/58 mm Hg. She had a left facial weakness, right forced gaze, and dense left hemiplegia. NIHSS score was 15. MRI was performed at 9:26 AM, approximately 101 minutes from symptom onset. MRI showed a right putaminal ICH with extension to the internal capsule.

Patient 5 is an 87-year-old woman with hypertension who had sudden onset of dysphagia, fell down, and had decreased level of consciousness at 11:45 AM. Blood pressure was 215/150 mm Hg. She was difficult to arouse, had marked dysarthria, 1- to 2-mm poorly reactive pupils, vertical nystagmus, dysmetria, and weakness of the right face, arm, and leg. NIHSS score was 12. The scan started at 1:45 PM, 120 minutes from symptom onset. MRI showed a left pontomedullary junction ICH.

**Results**

The diagnosis of ICH was made by the stroke fellows (I.L., R.H.L.), the stroke attending physicians (L.R.C., S.W.), the neurology attending physician, and the neuroradiologist on the basis of risk factors, clinical findings, clinical course, and outcome. ICH was readily identified in all patients by the neuroradiologists as part of their clinical routine evaluation. The simultaneous acquisition of conventional MRI sequences (ie, T1WI, T2WI, and FLAIR) ruled out other pathologies such as abscesses, neoplastic lesions, or ischemic stroke.

The results of SWI, T1WI, T2WI, FLAIR, DWI, and CT are shown in Figure 1 for patient 1 and in Figure 2 for patient 3. The SWI results are shown for all patients chronologically in Figure 3. Characteristic patterns on MRI were best defined on SWI, T2WI, and T1WI.

The lesion can be divided into 3 parts: a center, a periphery, and a surrounding rim. The center has isointense to hyperintense signal characteristics on T2WI and SWI, which may appear heterogeneous, particularly in larger lesions (patient 4, Figure 3). The center is isointense to hypointense on T1WI and becomes smaller over time. The periphery shows an enlarging area of signal loss on SWI. This hypointense area is present to a lesser extent on T2WI and is not apparent on T1WI. A surrounding rim of hyperintensity on T2WI and to a lesser extent on SWI circumscribes the periphery of the hematoma. The surrounding rim is hypointense on T1WI. This area most likely reflects vasogenic edema surrounding the hematoma. For large lesions (eg, patient 4 in this series), the hematoma may have a marbled appearance in the center and/or the periphery. The signal loss is more homogeneous in the patient imaged at 120 minutes and is present only at the periphery of the hematoma in the patient imaged at 23 minutes.

On the basis of the history and neurological examination, we posit that patient 1 had 2 hemorrhages. The first hemorrhage most likely occurred 3 hours before the scan, when he noticed the sudden onset of slight hemiparesis. The second hemorrhage most likely occurred 23 minutes before the scan, when the patient had sudden onset of vomiting, altered consciousness, deviation of the eyes, and hemiplegia. SWI showed a 1- to 2-cm area of signal loss in the left thalamus and the basal ganglia. This area most likely represents the hematoma that was responsible for the sudden onset of mild hemiparesis and (based on the clinical presentation), that occurred approximately 3 hours before the MRI. SWI showed...
another area (3 to 4 cm) that is hyperintense to white matter and hypointense to cerebrospinal fluid, with a periphery of signal loss. We posit that the hyperintensity in the center of the hematoma represents the new hemorrhage that caused sudden vomiting, loss of consciousness, and hemiplegia 23 minutes before the MRI.

**Discussion**

In ICH, hemoglobin passes from an environment with high oxygen concentration (arterial blood) to an environment with a lower oxygen concentration (tissue). It has been proposed that cerebral blood flow and pH are decreased in the under-perfused tissue surrounding the hematoma.9 These alterations might promote local shift in the hemoglobin oxygenation curve in favor of the formation of deoxyhemoglobin. Signal loss on long-TR/short-TE, long-TR/long-TE, and gradient-echo images is the earliest published change observed in a rat model of ICH.10 Such signal characteristic is present at the earliest point at which the animals were imaged, ie, 1 hour from the induction of the hematoma. In those animals, histology correlated signal loss to the transformation (within the erythrocytes) of hemoglobin to deoxyhemoglobin, starting from the most peripheral areas of the hematoma.19 In the series of Patel et al.,7 signal loss was present on SWI, within the hematoma, as early as the patient underwent an MRI scan (eg, 2.5 hours). In agreement with these observations, SWI, in our patients, is the sequence that more clearly shows signal loss due to the presence of deoxyhemoglobin. It seems to
progress from the periphery of the hematoma (Figure 3) toward the center. This observation has been described both in animal models\textsuperscript{10} and in patients with ICH.\textsuperscript{7,10} According to the authors, it can be explained by the presence of deoxyhemoglobin being the highest in the periphery of the hematoma.\textsuperscript{10} The meaning of the increased signal intensity seen on SWI in the center of the acute ICH is unknown. This signal intensity is higher than in normal tissue but lower than in edema or cerebrospinal fluid. We hypothesize that fresh blood in the center of a hyperacute ICH has signal characteristics of a proteinaceous solution since proton relaxation times are not yet significantly influenced by the presence of deoxyhemoglobin.\textsuperscript{5,12}

On the basis of our and other data, a model of ICH MRI signal characteristics from 23 minutes to 5 hours and their interpretation is summarized in the Table. The lesion appears to be made of 3 parts: a center, a periphery, and a surrounding rim. These parts evolve over time. The hyperintense center becomes progressively smaller over time, as deoxygenation of extravascular blood progresses from the periphery inward. For smaller (eg, patient 5 in this series) or later lesions, the hematoma becomes completely hypointense earlier. For larger lesions (eg, patient 4 in this series), the hematoma may have a marbled appearance at the center and periphery. In such cases, the evolution to complete hypointensity might be slower. As shown by T2WI, the outer rim of vasogenic edema enlarges over the hyperacute period. However, on SWI the increased signal intensity due to the vasogenic edema might be overwhelmed by the signal loss because of the susceptibility effect of deoxyhemoglobin.

The increased signal intensity present on FLAIR and T2WI and the hypointensity present on T1WI are useful in ruling out ischemic stroke. In fact, these sequences would be expected to be normal within the first 2 hours after the onset of ischemia. DWI does not seem to be as specific for hemorrhage as it is for ischemic stroke, and the diagnosis of hemorrhage should be based on other sequences. Attempts to measure the apparent diffusion coefficient in ICH are inherently flawed and systematically underestimated since any susceptibility effect will cause dephasing (ie, lower signal intensity) at all b values. Therefore, in an acute ICH, because of the susceptibility effect of deoxyhemoglobin, it would be difficult to measure correctly the apparent diffusion coefficient.

Schellinger et al\textsuperscript{8} measured lesion volume in patients with ICH imaged with MRI and CT between 3 and 6 hours from symptom onset. They concluded that T2WI and T2*-weighted images were as sensitive as CT in identifying ICH in the 9 patients imaged. The authors did not present data on patients

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<th>MRI Signal Features of Hyperacute Hemorrhage</th>
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Figure 3. SWI of all 5 patients showing the pattern of temporal evolution of the signal changes in ICH. SWI shows the progressive extension of the signal loss due to the presence of deoxyhemoglobin from the periphery toward the center of the hematoma. The signal loss is present only in the periphery of the hematoma in patient 1 and is completely hypointense in patient 5.
imaged within 3 hours from symptom onset. To our knowledge, this is the first report of MRI signal changes in patients with ICH imaged between 23 and 120 minutes from symptom onset. EPI/T2*-weighted imaging (SWI) seems the most useful image modality in detecting signal changes in the hyperacute phase of ICH. By SWI, data from the entire brain are acquired in 2 seconds of total scan time (20 slices, 7 mm thick). The finding is useful because a set of MRI sequences (DWI, SWI, FLAIR, T1WI, T2WI, PWI, and MRA) can provide multiple and complementary data in a total scan time of 15 minutes. Such data could be employed in the screening and use of thrombolytic and neuroprotective agents.1a In conclusion, before MRI becomes clinical routine evaluation of ICH, larger prospective investigations are needed. However, we suggest that MRI may be diagnostic in the evaluation of hyperacute ICH.

Acknowledgments
This study was supported by the National Institute of Neurological Diseases and Stroke and by the American Heart Association (Dr Warach). The authors wish to thank Mahesh Patel, MD, and the Department of Neuroradiology of Beth Israel Deaconess Medical Center for the advice in the interpretation of the MR images.

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Stroke. 1999;30:2263-2267
doi: 10.1161/01.STR.30.11.2263

The online version of this article, along with updated information and services, is located on the World Wide Web at:
http://stroke.ahajournals.org/content/30/11/2263

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