The Hypodense Artery Sign

Thomas C. Lee, Eric S. Bartlett, Allan J. Fox and Sean P. Symons

*AJNR Am J Neuroradiol* 2005, 26 (8) 2027-2029

http://www.ajnr.org/content/26/8/2027

This information is current as of October 28, 2023.
The Hypodense Artery Sign
Thomas C. Lee, Eric S. Bartlett, Allan J. Fox, and Sean P. Symons

Summary: An acute intracranial embolus may be associated with a hyperdense artery sign on CT, related to acute thrombus within the affected artery, a phenomenon that is well known and has been extensively documented. We present an unusual case of a middle cerebral artery territory acute infarct with a hypodense artery on CT. The Hounsfield unit attenuation of the embolic lesion was fat density. CT angiography and MR imaging confirmed the fatty lesion to be within the middle cerebral artery.

CT findings in acute stroke may be subtle or even undetectable on initial examinations, even when a large territory is involved (1). Various signs have been described as markers of early acute stroke on CT. Increased attenuation of the middle cerebral artery due to clot was first described in the early 1980s (2). Since that time, the “hyperdense middle cerebral artery” sign has been described as a marker of thrombus in the middle cerebral artery (3, 4), and several studies have demonstrated that the hyperdense middle cerebral artery sign is a common finding when CT is performed within a few hours from the onset of symptoms (5–8). False-positive hyperdense middle cerebral artery signs have been noted in patients with a high hematocrit level or calcification in the middle cerebral artery, but in these cases, the abnormal attenuation is usually bilateral (5, 8–10). In our patient, a fat embolus caused a hypodense appearance of the middle cerebral artery with an associated acute middle cerebral artery territory infarct.

Case Report
A 78-year-old woman with no significant medical history underwent an abdominoperineal resection for rectal adenocarcinoma in February 2002. At that time, mitral regurgitation was discovered, which progressed clinically to the point where she required surgical treatment. On January 28, 2004, she underwent mitral valve replacement. The surgical anatomy was complicated, but there were no intraoperative complications. In the immediate postoperative period, she was noted to have a decreased level of consciousness, a dense right hemiplegia, and aphasia. CT of the head performed on the first postoperative day demonstrated an early acute large left middle cerebral artery territory infarct. There was a hypodense filling defect in the left M1 proximal trunk, and the Hounsfield unit measurement of it indicated fat attenuation, with the lowest Hounsfield unit measurement of the intraluminal embolus being –25 H (Fig 1). Two subsequent CT scans, obtained during the next few days, demonstrated similar findings. CT angiography performed on the 13th postoperative day confirmed that the fatty lesion was intraluminal (Fig 2). CT angiography also demonstrated patent left common and internal carotid arteries without significant stenosis or atherosclerotic plaque. The fat embolus likely came from the surrounding tissues during the cardiac surgery.

She remained in the cardiovascular intensive care unit with ventricular pacing for nearly 2 months following the operation. MR imaging was performed 49 days after her stroke, once the pacemaker was removed. MR imaging showed interval evolution of the large left middle cerebral artery infarct. The previously seen infarct edema had resolved. There was hemorrhage within the infarcted territory and in the ventricles, which had occurred in the interval since the prior CT scans but was not

Fig 1. A 78-year-old woman who had undergone a mitral valve replacement. A–C, Axial images from the initial CT head scan obtained on the first postoperative day demonstrate an early left middle cerebral artery territory infarct. There is a fat attenuation filling defect (arrow, B) in the region of the proximal left middle cerebral artery trunk.
clinically symptomatic. Imaging through the left proximal mid-

dle cerebral artery, without and with fat saturation, showed

remnants of the fat embolus within the M1 segment (Fig 3).

**Discussion**

Emboli to the intracranial circulation can originate

from numerous sources. Emboli can originate in the

heart, particularly in the setting of atrial fibrillation. Emboli commonly propagate from the carotid arter-

ties as a result of stenosis or complicated atheroscle-

rotic plaque. Thrombus can form at the site of a vessel
dissection with subsequent embolization to the intra-

cranial circulation. In these scenarios, the emboli are

thrombotic in composition. Fat emboli to the intra-

cranial circulation are uncommon.

Aside from mortality due to other causes, stroke is

the most feared complication of cardiac surgery. The

incidence of stroke after coronary artery bypass is

1.4%, whereas the risk of perioperative stroke for

valve surgery is higher, quoted at 2.5% (11). Persis-
tent atrial fibrillation is the most significant risk factor

for stroke after mechanical mitral valve replacement

(12).

Fat emboli that usually originate from long bone

fractures are more commonly described in the trauma

and orthopedic literature. The major clinical criteria

for diagnosing fat embolism include respiratory dis-

tress and petechiae. Cerebral fat embolism has been

described in patients with a venous-to-arterial shunt

(13).

The typical fat emboli syndrome is generally

thought to be a result of microglobules. The high-

resolution chest CT findings of mild fat embolism are

nonspecific, consisting of bilateral ground-glass opac-

ities and thickening of the interlobular septa. Centri-

lobular nodular opacities are present in some patients

(13). Therefore, the clinical situation is the most im-

portant determinant in diagnosing fat embolus (14,

15).

In our patient, the diagnosis of a fat embolus was

suspected on the basis of the negative Hounsfield unit

attenuation in the middle cerebral artery. Presumably

the mitral valve replacement procedure resulted in

FIG 2. CT angiography axial image obtained 13 days postop-
eratively confirms that the fat attenuation lesion (arrow) is within

the proximal left middle cerebral artery trunk.

FIG 3. Axial and coronal T1-weighted

MR images obtained 49 days postopera-

tively without fat saturation (A,B) and with

fat saturation (C,D). The images without fat

saturation demonstrate the T1-hyperin-
tense embolus (arrow) in the proximal left

M1 trunk. After fat saturation, the T1 hy-

perintensity of the embolus is saturated

(arrow), confirming its fatty composition.

High-signal-intensity methemoglobin in

the adjacent infarct is unchanged before

and after fat saturation.
dislodgement of a macroscopic portion of fat from the cardiac or surrounding tissue, which subsequently embolized to the left middle cerebral artery. Alternatively, fat may have been dislodged during the sternotomy and paradoxically embolized to the brain through an intraoperative atrial defect. Fat containing emboli could also originate from fatty atherosclerotic plaque within the carotid artery or aortic arch. However, there was only minor atherosclerotic plaque in the proximal left internal carotid artery of this patient on CT angiography; therefore, a carotid artery source was excluded. Although the hyperdense middle cerebral artery sign has been widely described as a sign of thromboembolism, a hypodense middle cerebral artery sign associated with acute stroke from a fat embolus is not commonly known.

References