Simultaneous Thrombosis in a Normal Left Ventricle and Normal Carotid Artery in a Patient With a Stroke Secondary to Iron Deficiency Anemia

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Abstract
Iron deficiency anemia (IDA) is implicated as a cause of stroke, particularly in young patients without cardiovascular disease. In such patients, thrombi sometimes form in carotid arteries or the aorta. We report here a patient with a stroke secondary to IDA with thrombi in the normal left ventricle and normal carotid artery. The patient was a 45-year-old woman with severe IDA who developed cerebral infarction in the right middle cerebral artery. She had no other thrombophilia or cardiovascular diseases. Echocardiography showed a left ventricular thrombus without cardiac disease, and the carotid ultrasound showed a mobile thrombus attached to the right internal carotid artery without atherosclerosis. Antithrombotic therapy with iron supplementation removed both thrombi within 2 weeks. This is the first case of IDA with a ventricular thrombus in a normal heart. This case identifies a new site for thrombosis in IDA and shows that patients with IDA may present with simultaneous thrombosis at separate sites. IDA must therefore be recognized as a cause of stroke or systemic embolism, particularly in patients without detectable cardiovascular disease. In such patients, it is important to conduct careful investigations to detect thrombosis.

Keywords: Carotid artery thrombosis; Echocardiography; Embolism; Iron deficiency anemia; Left ventricular thrombosis; Menorrhagia; Stroke; Cardiac thrombosis; Cerebral infarction

Introduction
Iron deficiency anemia (IDA) is a common disease implicated as an uncommon cause of stroke and is epidemiologically associated with increased risk of stroke [1, 2]. Moreover, several case reports suggest a causative role of IDA in young and otherwise healthy patients with stroke or with systemic embolism where thrombi are sometimes detected on an apparently intact arterial wall such as the carotid artery [3-9] or aorta [10, 11]. Here we describe a patient with thrombi in an otherwise normal carotid artery and left ventricle, which is a previously unreported site for thrombosis in patients with IDA. Our case exemplifies a new site for thrombosis in IDA and indicates that patients with IDA may present with simultaneous thrombosis at separate sites.

Case Report
A 45-year-old woman presented with left hemiparesis. Three years before admission she was diagnosed with IDA, which was attributed to menorrhagia; however, she did not take medication and was otherwise healthy. She experienced headache and left hemiparesis upon awakening and was admitted to the hospital. Her blood pressure was 122/80 mm Hg, pulse rate regular at 90/min and anemic conjunctiva. She was alert but slightly agitated. Neurological examination revealed left hemiparesis and left hemispatial neglect, and magnetic resonance (MR) imaging revealed right frontoparietal infarction. MR angiography revealed occlusion of the distal branches of the right middle cerebral artery, and MR venography revealed normal dural sinuses. The electrocardiogram was normal. Hemoglobin was 6.0 g/dL, mean corpuscular volume 65.3 fL, platelets 573,000/μL, serum iron 19 μg/dL, total iron-binding capacity 497 μg/dL, transferrin saturation 3.8% and ferritin 1.6 ng/mL. Protein S, protein C, antithrombin III and antiphospholipid antibody values were normal, indicating the absence of thrombophilia.

The next day, transthoracic echocardiography revealed a mobile thrombus in the left ventricle apparently attached to the chordae tendineae of the mitral valve (Fig. 1). Chamber
size, wall motion and cardiac valves were normal. Carotid ultrasound revealed a mobile thrombus attached to the wall of the right internal carotid artery (Fig. 2). There was no detectable atherosclerotic lesion with a maximal intima-media thickness of 1.0 mm. She was treated with aspirin, clopidogrel and argatroban (a direct thrombin inhibitor) as well as iron supplementation. Her symptoms resolved after 2 weeks, and no thromboembolism occurred after admission.

On day 4 of hospitalization, ultrasound revealed that the carotid thrombus was reduced in size by 50%, and on day 8, carotid ultrasound did not detect a thrombus. On day 11, transesophageal echocardiography did not detect a cardiac thrombus; no septal defect or patent foramen ovale was present, and the thoracic aorta was normal. On day 14, transthoracic echocardiography revealed no left ventricular thrombus.

Discussion

IDA is implicated as a cause of stroke or systemic thromboembolism, and epidemiological studies [1, 2] revealed that IDA is more prevalent in patients with stroke compared with controls. Moreover, when IDA is associated with stroke or a systemic embolism, large thrombi are sometimes present in an apparently intact carotid artery [3-9] as in our present case or less commonly in the aorta [10, 11]. These patients are young women 20 - 50 years of age with severe IDA (Hb 5.5 - 10 g/dL), which is often caused by menorrhagia. More important, they have no known cardiovascular disease. These common features, taken together with the characteristics of our present patient, suggest that IDA plays a causative role in certain young patients with stroke rather than as a risk factor.

The present case reveals that the left ventricle represents a new site for thrombosis in patients with IDA. In the absence of detectable cardiac disease, left ventricular thrombosis is rare and described only anecdotally. Many of these patients have prothrombotic disorders such as protein C deficiency [12], protein S deficiency [13], essential thrombocytemia [14] and antiphospholipid antibody syndrome [15]; however, there are no reports of patients with left ventricular thrombus associated with IDA. Because our patient had no other detectable prothrombotic disorders, we concluded that IDA was the sole cause of left ventricular thrombosis. Moreover, since left ventricular thrombosis is rare in the absence of cardiac disease, such a thrombus may be overlooked, particularly when it is small or less mobile. Conversely, when the thrombus is large and mobile, it may be indistinguishable from cardiac myxoma [14-17], and this diagnostic uncertainty may lead to surgery. In our present patient, the thrombus was distinguished from myxoma by its relatively small size and its resolution after treatment. Therefore, it is important to recognize IDA as a cause of left ventricular thrombosis to avoid unnecessary surgery as well as to identify the embolic source.

The characteristics of our present patient, taken together
Ventricular and Carotid Thrombosis in Iron Deficiency Anemia


