Brain embolic infarction associated with cardiac amyloidosis in a patient with multiple myeloma: a bone marrow-heart-brain crosstalk

Jaeho Kim, Jaehong Park, Sung-Ji Park, Sung Mok Kim, Kihyun Kim, Jong-Won Chung, Oh Young Bang

1Department of Neurology, Samsung Medical Center, Sungkyunkwan University School of Medicine, Seoul, Korea
2Division of Cardiology, Department of Medicine, Samsung Medical Center, Sungkyunkwan University School of Medicine, Seoul, Korea
3Department of Radiology, Samsung Medical Center, Sungkyunkwan University School of Medicine, Seoul, Korea
4Division of Hematology-Oncology, Department of Medicine, Samsung Medical Center, Sungkyunkwan University School of Medicine, Seoul, Korea

ABSTRACT
Cardiac amyloidosis is characterized by the extracellular fibril deposition of amyloid protein in the myocardium. Cerebral embolism caused by cardiac amyloidosis is rare. We report a case of 59-year-old woman with multiple myeloma who suffered from cerebral infarction probably related to cardiac involvement of amyloidosis, and discuss the pathophysiological mechanisms based on kidney biopsy and characteristic echocardiographic and cardiac magnetic resonance imaging features.

Keywords: Amyloidosis; Cardiomyopathies; Cerebral infarction; Magnetic resonance imaging; Multiple myeloma

INTRODUCTION
Immunoglobulin amyloid light chain amyloidosis may occur as a primary disease or in association with multiple myeloma or other plasma cell dyscrasia [1]. The heart is affected in over 50% of cases [2] but cerebral embolism in relation to cardiac amyloidosis has rarely been reported [1,3]. Herein, we report a patient with multiple myeloma who suffered from embolic infarction associated with cardiac amyloidosis diagnosed based on the kidney biopsy, echocardiographic and cardiac magnetic resonance imaging (MRI) results.

CASE REPORT
A 59-year-old woman suddenly developed vertigo type dizziness. Six years ago, she visited our hospital because of back pain. Spine MRI showed multiple spine mass and computed tomog-
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CT guided bone biopsy revealed clusters of squeezed atypical cells. Laboratory test results showed hypercalcemia, elevated serum lambda (λ) concentration (4,980.00 mg/L) and urine protein electrophoresis revealed suggestive of monoclonal proteinuria (M-protein 2,884.2 mg/day). Kidney biopsy, stained with Congo red, demonstrated amyloid deposition. Based on these findings, she was diagnosed as having multiple myeloma with amyloidosis. Thalidomide, cyclophosphamide, and dexamethasone were prescribed.

The patient was non-smoker and non-alcoholics. On neurologic examination, she showed right-sided ataxia in cerebellar function test and other neurologic examination was normal. Routine electrocardiography (ECG) and 24-hour Holter ECG monitoring revealed episodes of 2nd degree atrioventricular block (Mobitz type I) and no atrial fibrillation or sustained atrial flutter. Laboratory test results showed no significant abnormality of thrombotic profile. Brain CT 12 hours after the onset showed no evidence of intracranial bleeding and brain MRI revealed acute infarcts in the right posterior inferior cerebellar artery (PICA) territories (Fig. 1A). Perfusion-weighted MRI showed perfusion delay related to the lesion and magnetic resonance (MR) angiography showed no evidence of vessel abnormality (Fig. 1B).

Transesophageal echocardiography showed symmetrical thickenings of the left ventricular (LV) and left atrial (LA) wall. LA was enlarged and transmitral Doppler showed diastolic dysfunction, and the cardiac involvement of amyloidosis was suspected (Fig. 2). There was spontaneous echo contrast

Fig. 1. (A) Fluid attenuated inversion recovery brain magnetic resonance imaging shows cerebral infarcts in the right posterior inferior cerebellar artery territory. (B) Magnetic resonance angiography reveals no vessel abnormality.

Fig. 2. Echocardiographic findings. (A) Increased left ventricle basal septal wall thickness (arrow), (B) left atrium (LA) wall thickness (arrows), (C) spontaneous echo contrast without thrombi in LA appendage (circle).
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(SEC) without thrombi in LA appendage (Fig. 2C).

In suspicion of cardiac amyloidosis, she underwent cardiac MRI which showed increased LV basal septal wall thickness and increased both atrial wall thickness. In late gadolinium enhancement imaging, myocardium showed suboptimal nulling in generally and LV subendocardial enhancement of LV myocardium and LA wall enhancement was prominent (Fig. 3). These findings were consistent with cardiac involvement of amyloidosis. The patient was treated with warfarin. She recovered completely 2 days after the onset.

DISCUSSION

This case illustrates an example of the bone marrow-heart-brain crosstalk. The patient had a cerebral infarct in the right PICA territory but no vascular risk factors or atherosclerotic changes in MR angiography suggesting that embolization was the etiology of her stroke. Cardiac amyloidosis was diagnosed by characteristic of echocardiographic and cardiac MRI findings and result of the kidney biopsy.

To the best of our knowledge, this is the first case which showed the cardioembolic stroke caused by cardiomyopathy related to amyloid deposition in patients with multiple myeloma with specific findings of cardiac MRI. Typical features of echocardiography of cardiomyopathy related to cardiac amyloidosis include thickened LV walls with normal to small cavity size, biatrial enlargement, thickened valves, and diastolic dysfunction [4,5]. In addition, typical cardiac MRI features of this condition are tissue characterization of the myocardium: transmural late gadolinium enhancement, LV wall thickness and biatrial enlargement, and pericardial effusion [5].

The mechanisms of embolization in amyloid cardiomyopathy are multifactorial [1,3], (1) mural thrombosis due to blood stasis and intracavitary turbulence [6-8], (2) myocardial ischemic, endocardial damage [7], (3) arrhythmia (e.g., atrial fibrillation) may due to LA dilatation and failure with increasing wall stress [9,10], and (4) plasma hypercoagulability state resulted from nephrotic syndrome, thrombocytosis related to hyposplenism, thrombin-antithrombin pathway impairment [11]. In this case, mural thrombosis due to cardiomyopathy was considered as the mechanisms of stroke, and prophylactic anticoagulation was indicated because SEC was observed in LA appendage.

Etiological workups are complex in patients without demonstrable arterial stenosis or potential sources of cardioembolism [12]. The case presented here demonstrated that multidisciplinary approach involving oncologist, cardiologist, and neurologist are important to evaluation and treatment of patients with ischemic stroke.

CONFLICTS OF INTEREST

No potential conflict of interest relevant to this article was reported.

REFERENCES