Hemorrhagic Cardioembolic Stroke Secondary to a Left Ventricular Thrombus: a Therapeutic Dilemma


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Abstract

Cardiogenic embolism is a major cause of stroke and often leads to significant morbidity and mortality. Despite the recent advances in our understanding of the pathophysiology of stroke and its risk factors, diagnosis and therapy; some case scenarios still present a real challenge for the treating physicians. We report a case of a 50 year old male patient presenting with multi-territory cerebral infarctions due to a left ventricular mobile thrombus complicated with hemorrhagic transformation at the time of presentation. Gradual introduction of anticoagulation coupled with a multidisciplinary team approach advocating careful daily clinical assessment of the patient and regular echocardiographic and neuroimaging studies have resulted in a better management and achievement of therapeutic goals.

Keywords: Stroke; Cardioembolism; Mobile thrombus; Hemorrhagic transformation; Echocardiography; Multidisciplinary approach.

Introduction

Embolic stroke is a leading cause of morbidity and mortality. The heart is an important source for the development of embolic strokes. About 15-20% of all ischemic strokes are cardioembolic. Establishing the heart as a source for a patient’s embolic stroke carries important prognostic and therapeutic implications. Patients with such types of stroke often suffer significant morbidity and mortality and have a high risk of recurrence. The presence of a left ventricular thrombus carries additional risks including the risk of hemorrhagic transformation as well as a high risk for recurrent embolic strokes. Managing patients with left ventricular thrombi causing an embolic stroke is challenging especially in the presence of hemorrhagic transformation as we present in this report.

Case Report

A 50-year old right-handed Omani male with a history of smoking and dyslipidemia, taking aspirin and simvastatin, presented with 6 hours history of sudden onset left sided weakness and inability to speak. He had no history of chest pain or palpitations. He did, however, give a history of a recent admission to another hospital with an episode of prolonged chest pain. Details of his management during that admission were not available. Examination showed blood pressure of 115/80 mmHg, regular heart rate of 90 per minute, Glasgow coma scale (GCS) of 12, expressive aphasia, left facial nerve palsy of upper motor neuron type with dense left sided hemiplegia and left homonymous hemianopsia (NIH Stroke Scale score 17). Cardiac examination as well as ECG, cardiac enzymes and basic laboratory investigations was unremarkable. Initial CT brain showed a right occipital infarct with focal areas of hemorrhagic transformation. The patient was admitted to the high dependency unit of the general medical ward with regular assessment of his GCS. A repeat CT brain was performed on the second day and showed a large right middle cerebral artery (MCA) territory infarct with mass effect in addition to the right occipital infarction and the hemorrhagic transformation noted earlier. (Fig. 1)

Echocardiography revealed poor overall systolic function (ejection fraction 45%) with hypokinetic apical and lateral walls with an apical aneurysm and a mobile thrombus measuring about 3.6 cm x 1.7 cm in the apical region of the left ventricle (Fig. 2). The patient was started on 100 ml of Mannitol 20% three times daily with daily monitoring of the electrolytes profile in addition to regular physiotherapy. Forty-eight hours after the onset, he was started on weight-adjusted low molecular weight heparin (LMWH) at a prophylactic dose (enoxaparin 40 mg/day) and clopidogrel 75 mg once a day. He had minimal improvement in his neurological status. His thrombophilia workup (including tests for protein S, protein C and anti-thrombin levels and activities sent before the initiation of LMWH, as well as molecular testing for factor V Leiden and prothrombin gene mutations) were unremarkable. MRI brain was done and confirmed the CT findings.

Daily assessment of the patient by a multi-disciplinary team including a neurologist, a cardiologist and a hematologist in addition to the primary internist was adopted. A weekly echocardiogram was used to assess the thrombus size while weekly neuro-imaging with CT scans was initially used to assess the infarcted areas, especially with regards to hemorrhagic transformation. With no progression...
in the small areas of hemorrhagic transformation on CT scans, the dose of enoxaparin was upgraded to 40 mg twice daily and then after another week, it was increased to 60 mg twice daily with weekly monitoring of anti-Xa levels. Warfarin was introduced with a target INR of 2.0 to 3.0 when the CT showed complete resolution of the cerebral hemorrhage. Follow up echocardiographic studies demonstrated a significant regression in the thrombus size without any evidence of further systemic embolization. The patient was discharged after 7 weeks of hospitalization with NIH Stroke Scale score of 15 and a modified Rankin Scale score of 5.

Discussion

Cardiogenic embolism is recognized to account for one in six strokes.1,2 Common causes of cardiac embolization include atrial fibrillation, ischemic heart disease, rheumatic heart disease and prosthetic cardiac valves.3-6 Left ventricular thrombi (a frequent complication of transmural acute myocardial infarction, likely to be the case in our patient),7,8 are second to atrial fibrillation as a cause of cardiogenic embolization. These are easily detectable by two-dimensional echocardiography, although false-positive studies exist.9 Protruding configuration and free mobility of the thrombus and the presence of adjacent hypokinesia indicate a high embolization potential.10,11 The presence of a potential embolic source does not, by itself, justify the diagnosis of cardiogenic embolism in this patient. However, the abrupt onset of maximal deficit, the presence of multiple brain infarcts involving different vascular territories with hemorrhagic transformation along with the presence of a large left ventricular thrombus indicate that the cause of the infarcts in this patient are indeed cardioembolic.

Hemorrhagic transformation (HT) is not uncommon and patients with cardioembolic infarcts are particularly prone to develop hemorrhagic transformation. HT has been reported in up to 40% of patients with cerebral embolism on CT scans,12 and more than 60% on MRI.13 Older age and large sized infarcts are known to be associated with particularly high risks of hemorrhagic transformation.12-14 In spite of his young age, our patient suffered large cardioembolic infarcts in multiple vascular territories, which led to hemorrhagic transformation.

Patients like the one we are reporting pose a significant challenge. On one hand, he had a major source for recurrent embolic stroke. On the other hand, treating that source with systemic anticoagulation carries the risk of exacerbating the pre-existing cerebral hemorrhage, especially given the size of his initial infarcts. The decision to anticoagulate is generally based on the balance between the relative risk of these two, i.e. the relative risk of embolization without anticoagulation and that of rebleeding or expansion of existing bleeds if anticoagulants are started early. Although the exact risk of each of these is difficult to define, our patient had high risks for both. An extensive literature search to guide our strategy in managing this patient was largely unhelpful and revealed contradicting data. Similar cases are rarely reported in the literature.15 Unfortunately, the available guidelines for the management of stroke do not address this particular issue.16-18 Properly designed clinical trials of the optimal anticoagulation strategy in cardiogenic embolism are not available.19 Several studies have reported no increase in the rate of hemorrhagic transformation, hence supporting the use of immediate anticoagulation in embolic stroke.20-24 However, such studies are either retrospective, including small numbers of patients or, more importantly, have excluded patients with hemorrhagic transformation on initial presentation. Several other studies, on the hand, have shown either an increase in intra-cranial bleeding or no significant reduction in the recurrence of ischemic strokes, death or disability.25-28
As our patient had spontaneous hemorrhagic transformation in a large embolic infarct (both of which are associated with increased risk of further hemorrhage), we opted not to initiate full anticoagulation. This approach is supported by evidence from several reports on patients with high risk of embolization and urgent need for anticoagulation (such as patients with mechanical heart valves) who present with intra-cranial hemorrhage. Such reports indicate that it is safe to withhold anticoagulation for periods extending up to a few weeks. The challenge, however, was that with his large mobile left ventricular thrombus, he had another large embolic infarct within a day of the initial one and without anticoagulation; one risks him getting another major one. We therefore, opted for a graded introduction of anticoagulation initially with prophylactic doses of LMWH going to full dose with close monitoring of anti-Xa levels, as well as regular imaging with CT scans and echocardiography. The patient did not suffer any further emboli or more hemorrhage. He in fact, showed some improvement both clinically and in the size of the left ventricular thrombus.

Conclusion

Left ventricular thrombus complicating ischemic heart disease and causing systemic embolism is a well-recognized clinical situation which requires anticoagulation. This therapy may pose a real challenge in patients with ischemic strokes complicated by hemorrhagic transformation. Multidisciplinary approaches, gradual introduction of anticoagulation guided by the available literature and close monitoring in such patients would result in a better outcome.

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References


