Spontaneous Conversion of Atrial Fibrillation to Normal Sinus Rhythm Following Recurrent Cerebral Infarctions

Kyungmi Oh, M.D., Ph.D., Jeong-Yoon Choi, M.D., Byung-Jo Kim, M.D., Ph.D.
Department of Neurology, Korea University Medical Center, Korea University College of Medicine, Seoul, Korea

INTRODUCTION

It has been frequently described that acute stroke is associated with increased incidence of arrhythmias. One of the most common arrhythmia following stroke is atrial fibrillation. Although the pathogenesis of these arrhythmias is still obscure, they are obviously associated with impairment of cardiac autonomic balance between the sympathetic and the parasympathetic nervous system.

Arrhythmia immediately after stroke has been reported in patients with both hemispheric and brainstem cerebral infarcts. Most post-stroke arrhythmias appear in patients with right-sided hemispheric infarcts, especially in patients with right insular infarcts. However, conversion of arrhythmia to normal sinus rhythm following acute cerebral infarct has not been reported. Herein, we present a patient who had spontaneous conversion of atrial fibrillation to normal sinus rhythm after recurrent cerebral infarcts.

CASE REPORT

A 88-year-old man was brought to emergency room with right hemiparesis and aphasia. Six years ago, he was admitted to hospital due to transient ischemic attack and diagnosed of atrial fibrillation, hypertension, hyperlipidemia, gout and chronic renal insufficiency. At that time, brain MRI showed old basal ganglia lacunar infarct without any abnormality in intracranial and extracranial vascular structures. Thereafter, he had regularly visited outpatient clinic of internal medicine with medication including antiplatelet agent. He had not been administered any medication for permanent atrial fibrillation which was continued at regular check-up.

On admission, he was mildly drowsy and not communicative. Neurological examinations showed right hemiparesis and aphasia. Diffusion-weighted MRI showed multiple small cortical infarcts in the left middle cerebral artery territory without involvement of insular cortex (Fig. 1A). Electrocardiogram (12 leads) revealed atrial fibrillation with rapid ventricular rhythm (Fig. 2). Echocardiogram revealed moderate aortic valve regurgitation and left atrial enlargement with diameter of 44.7 mm (Aorta, 28.1 mm) on M-mode study. However, ejection fraction was normal and thrombus was not observed in the atrium or ventricle. We started treatment with...
heparin and digoxin for atrial fibrillation with rapid ventricular rhythm, but did not use antiarrhythmic agent. After few days, digoxin was stopped due to decrement of heart rate less than 40/ min. Seven days after admission, antiocoagulation was stopped because of gross hematuria. Eighteen days after admission, intubation was performed due to purulent sputum and severe stridor induced by epiglottitis. His right-sided weakness progressed to hemiplegia of grade III, but other neurologic symptoms did not deteriorate.

Approximate 1 month after admission, his consciousness was suddenly decreased to stuporous state. On neurologic examination, eyeball was deviated to right side, and left side hemiparesis of grade I was newly developed. Emergent diffusion-weighted MRI revealed that right-sided diffuse subcortical infarcts of middle cerebral artery territory (Fig. 1B). However, insular cortex was not directly involved. Immediately after second ischemic attack, his atrial fibrillation converted spontaneously to normal sinus rhythm which detected by intensive care unit monitoring and ECG follow-up (Fig. 3). Cardiac enzymes were normal and follow-up echocardiogram was not different to previous exam. Normal sinus rhythm was sustained until he died due to sepsis on 11 days after second attack.

**DISCUSSION**

Although there have been many reports about acute onset arrhythmias following stroke, spontaneous conversion of arrhythmia to normal sinus rhythm after stroke has not been described. Post-stroke arrhythmia has been reported to develop a few hours to 3 days after stroke and then spontaneously reversed to normal sinus rhythm after a few days to possibly even a few months. Cerebrogenic mechanism of post-stroke arrhythmias was suggested by the temporal relation between stroke and arrhythmia. In our case, conversion of atrial fibrillation to normal sinus rhythm immediately following second stroke attacks may also support the theory of “cerebral arrhythmogenesis”. However, it is obscure whether this conversion to normal sinus rhythm was prolonged effect because our patient died during acute period of second stroke attack.

Several experimental or clinical evidences that acute stroke deranges cardiovascular autonomic regulatory system and results in electrocardiographic changes or arrhythmias have been reported. This finding was shown in patients with subarachnoid hemorrhage, subdural hematoma, brainstem hemorrhages and infarcts, and both hemispheric hemorrhages and infarcts irrespective of cortical or subcortical areas. Many investigations for cortical lateralization indicated that the right-side hemispheric stroke was more arrhythmogenic than left-side lesion. In particular, right-side insular cortex was suggested as cardiovascular regulation center in brain through few case reports with cerebral infarctions. However, post-stroke arrhythmias were found more commonly in patients with subarachnoid hemorrhage or intracranial hematoma without notion of any specific localization or vascular territory. Our case showed bilateral hemispheric infarctions within the middle cerebral artery territories. After right hemispheric infarction, his atrial fibrillation was converted spontaneously to normal sinus rhythm. This finding supports the hypothesis that cardiovascular regulation center in

![Fig. 1. Diffusion-weighted images (b=1000). A : The images performed at the first attack show acute cerebral cortical infarction on left fronto-temporal areas. B : Follow-up study at the second attack reveals diffuse subcortical infarct in right middle cerebral artery territory and left-sided lesion developed at the first attack.](image1)

![Fig. 2. Electrocardiogram (12 leads) which was performed on admission reveals atrial fibrillation with rapid ventricular rhythm.](image2)

![Fig. 3. Electrocardiograms (lead II) which were performed at intensive care unit before (A) and immediately after (B) the second cerebral infarct developed. The sinus rhythm is sustained until 6 days after the normalization (C). Arrows mark on R of QRS rhythms.](image3)
the brain is more lateralized to right hemisphere. A possible mechanism of cardioversion in our patient can be suggested by previous reports. Increased QT interval, which means the prolongation of action potential, is the main mechanism for arrhythmia conversion of class IA anti-arrhythmic agent such as quinidine, procainamide. The prolongation of QT interval following right hemispheric infarction has reported by previous studies. Although exact mechanism of conversion to normal sinus rhythm is not clearly understood, the possible prolongation of QT interval after the second stroke in right hemisphere may contribute to the spontaneous cardioversion in our case.

Many cerebral infarcts are classified as an undetermined mechanism according to TOAST classification, even the cases are thought clinically to be caused by cardioembolism. Arrhythmia such as atrial fibrillation is a major underlying condition for cardioembolism.

**CONCLUSION**

Our case may suggests possibility that physicians could miss the risk factor due to immediate conversion of arrhythmia to normal sinus rhythm after cerebral infarcts on admission. If no risk factor was revealed by intensive investigation, physicians should concern arrhythmia as a possible etiology of stroke even if the patient has normal sinus rhythm.

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