Transient left septal fascicular block in the scenario of ST-segment elevation myocardial infarction

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Introduction

Although the conduction abnormalities of the left anterior and posterior fascicles, left anterior fascicular block (LAFB), and left posterior fascicular block (LPFB) are commonly diagnosed and well-known to physicians, the abnormalities of the left septal fascicle (LSF) and left septal fascicular block (LSFB), are easily overlooked because most physicians are not aware of the existence of the LSF. LSFB exclusively affects precordial leads without modifying the ECG in the frontal plane, causing prominent anterior QRS forces. LSF is irrigated exclusively by the left anterior descending (LAD) coronary artery, and therefore, LAD occlusion can result in LSFB. We report a case with transient LSFB in the scenario of ST-segment elevation myocardial infarction (STEMI) because of acute occlusion of LAD.

Case Report

A 69-year-old female patient with a history of hypertension, diabetes, and hyperglycemia presented to the emergency department with sudden chest pain for 3 hours. Her mother and brother had a history of coronary heart disease. Furthermore, her brother had received percutaneous coronary intervention (PCI) for treating his underlying heart condition. On admission, her physical examination was unremarkable. Her blood pressure was 112/68 mm Hg and heart rate was 79 beats/min. A baseline ECG was performed and is shown in Figure 1. Her cardiac troponin I level was 0.055 ng/mL (normal, <0.02 ng/mL). The patient was sent to the catheterization laboratory. On the basis of the ECG, we aimed to identify the culprit lesion, and determine why the precordial R/S transition from lead V1 to V2 was so early. The baseline ECG (Fig. 1) showed the precordial R wave “in crescendo” from V1 through V3 and decreasing from V4 to V6; and ST-segment elevation from V2 to V4. The “R” wave jump occurred from V1 to V2. The ECG revealed STEMI caused by total occlusion of the LAD coronary artery. The coronary angiography showed total occlusion of LAD (Fig. 2a). After a successful PCI was performed (Fig. 2b), ECG demonstrated an “R” wave jump from V2 to V3 (Fig. 3). An ECG recorded the next day showed ST-segment resolution and T wave inversion of V2 to V4 (Fig. 4).

Discussion

The transient prominent anterior QRS forces, as shown in Figure 1, are the hallmark of LSFB. LSFB is a relatively common ECG manifestation, but very easily overlooked. The criteria for
ECG diagnosis of LSFB are as follows—QRS duration <120 ms—in general, close to 100 ms; more than 15 mm voltage R waves in V2 and V3 or from V1; increasing for all intermediary precordial leads and decreasing from V5 to V6; “R” wave jump may occur from V1 to V2 (“rS” in V1 for R in V2); absence of QRS axis shift in the frontal plane; and T wave polarity negative in the right precordial leads most of the time (1). LSF was first described by Dr Sunao Tawara, just as LAF and LPF (2).

Although LAFB and LPFB are well known, LSFB is easily overlooked by physicians, especially in the case of ischemic heart disease. Although LAF and LPF may be irrigated by the LAD coronary artery and right coronary artery, LSFB is irrigated exclusively by LAD, and therefore, LAD occlusion can result in LSFB. LSFB exclusively affects precordial leads without modifying the ECG in the frontal plane, causing prominent anterior QRS forces. Other possible etiologies of LSFB are chronic Chagas cardiomyopathy, nonobstructive hypertrophic cardiomyopathy, obstructive hypertrophic cardiomyopathy, aortic valve disease, diabetes mellitus, and Kearns–Sayre syndrome (3). The transitory nature of the electrocardiographic changes rules out any other possible cause.

**Conclusion**

We report a case of transient LSFB caused by the acute occlusion of the proximal LAD. Because transient LSFB may indicate critical obstructive lesions of the proximal LAD, but is easily overlooked, physicians should pay more attention to this ECG pattern.

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