Acute marginal branch occlusion presenting with anterior wall ST segment elevation during percutaneous coronary intervention of the mid right coronary artery

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Case
A 78-year-old male, ex-smoker, with a history of hypertension and hyperlipidemia presented with a one-month duration of worsening angina on exertion. Standard 12 L ECG showed sinus rhythm with early repolarization pattern in the anterolateral leads. Coronary angiography showed minor luminal irregularities in the left coronary vessels (Figure 1 A & B) while the RCA showed high grade non-thrombotic 90% lesion in its mid segment (Figure 2A arrow) followed by another high grade 90% stenosis in the RPDA branch (Figure 2B). A small calibre acute marginal branch (AMB) with an 80% ostial stenosis was noted to take off from the diseased mid RCA segment (Figure 2A arrowhead). Ad hoc percutaneous coronary intervention (PCI) of both the RPDA and mid RCA was carried out.

After predilation, a 2.5x24mm SYNERGY (Boston Scientific) Everolimus drug eluting stent was implanted at the RPDA stenosis followed by another 3.5x24mm SYNERGY (Boston Scientific) Everolimus drug eluting stent at the mid RCA lesion. Following stent postdilation, repeat angiography demonstrated loss of the AMB from the mid RCA. (Figure 3). The patient subsequently developed chest pain and 12L ECG showed ST elevation in leads V1-V4 from baseline, most marked in V2 and V3. (Figure 4)

Attempts to salvage the jailed AMB were unsuccessful due to its angulated origin. IV analgesia with midazolam and fentanyl alleviated the chest pain. The patient remained hemodynamically stable, hence decision was made to stop the procedure and continue optimal medical therapy. At the coronary care unit, bedside transthoracic echocardiogram revealed no anterior wall motion abnormality to suggest LAD occlusion. There was significant increase in serum Troponin I level to 9091ng/mL after 24 hours. Serial 12L ECG monitoring however showed gradual decline of ST elevation in leads V1- V4 (Figures 5 A & B). Resolution of STE was noted nine (9) hours post-procedure (Figure 5C). Repeat transthoracic echocardiogram the following day showed normal LV and RV function with an ejection fraction of 60% with no segmental wall motion abnormalities. Medical therapy was further optimized, and patient was discharged well 3 days after PCI.

Discussion
Our case demonstrated onset of anterior leads (V1-V4) STE with no changes in the limb leads tracing after AMB loss following mid RCA stent implantation. The existing high-grade ostial AMB stenosis predisposed to its closure which resulted in isolated RV infarction (IRVI). RV myocardial infarction most often occurs simultaneously with an inferior LV wall infarction involving an occlusion of the proximal dominant RCA [1]. Left precordial ECG manifestations of RV injury are often absent because they are masked by dominant forces of the accompanying LV inferior wall injury [2,3]. Hence, the predominant ECG presentation of ST elevation (STE) in the inferior limb leads. STE in V4R identifies patients with an acute inferior wall myocardial infarction who are likely to have concomitant RV involvement [1].

However, a larger part of the right ventricle free wall is directed anteriorly and when myocardial injury is isolated only to the right

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Figure 3. A. Mid RCA lesion pre-stenting with acute marginal branch (AMB). B. Mid RCA post-stenting with loss of acute marginal branch (AMB)

Figure 4. A. Baseline 12L ECG. B. 12L ECG showing 2mm ST-elevation of (V1-V3)

Figure 5. (A-D) Serial ECG tracing post-PCI showing gradual resolution of anterior leads STE with no Q wave evolution and absence of concomitant inferior lead STE
ventricle, the presence of minimal inferior wall current and the lack of posterior injury current permits the right ventricle injury current to predominate and manifest STE in leads V1–V4 [4]. Furthermore, the recording of

RV electrical potentials in the left precordial leads depends on the degree of clockwise rotation of the heart in the horizontal plane and on body geometry [5]. Isolated RV infarction (IRVI) is rare and occurs in less than 3% of all patients with myocardial infarction [6]. It may occur by i) proximal occlusion of a dominant right coronary artery (RCA), ii) occlusion of a nondominant RCA [7] and iii) isolated occlusion of an acute marginal branch (AMB) artery after percutaneous coronary intervention as demonstrated in our case.

IRVI should be suspected with the following ECG patterns: [7] (1) dome-like and decreasing ST segment elevation from leads V1 to V3; (2) rapid ST segment normalization and no Q-wave evolution from leads V1 to V3, with or without modest ST segment elevation in lead III (but not aVF) and no resultant Q wave; (3) ST segment elevation in right-sided leads; (4) absence of ST segment depression in aVL; (5) absence of concomitant ST segment elevation in inferior leads (II, III, aVF).

**Isolated anterior lead STE: IRVI versus anterior wall MI**

Anterior acute myocardial infarction (AMI) typically presents with anterior STE with reciprocal changes in the inferior leads [8]. This pattern of concomitant inferior ST segment depression is especially noted among AMI’s involving proximal left anterior descending (LAD) coronary artery occlusion [9,10]. It is however important to note that absence of reciprocal changes in the inferior leads does not automatically rule out anterior wall AMI, as this may distort LAD occlusion often still present with anterior STE with no reciprocal changes [9-11]. In addition, anterior wall AMI due to proximal LAD occlusion often presents with not only inferior ST segment depression but also STE in lead aVR or lateral limb leads, or both [9,10]. For patients with typical symptoms and ECG tracing of anterior lead STE, IRVI should be considered as a differential diagnosis [7,3]. Some authors have suggested a step wise approach in dealing with this finding [7,3]. First is to examine the presence of an incremental or decremental STE pattern from leads V1- V5. A decremental STE pattern favors IRVI [7]. Other investigators have considered the location of lead V1 being directly on top of RV, they have reported that STE primarily in V1 suggests IRVI [12]. Next is to look at the presence of ST depression (STD) in lead aVL. STD in aVL is a sign of reciprocal change in the presence of inferior wall injury, hence its absence points more towards the probability of IRVI [7]. Lastly, the presence of V3R and V4R STE makes the diagnosis IRVI highly probable [7]. No right sided precordial lead tracing was obtained in our case because of the information provided by coronary angiography while the anterior STE took place. Our patient however demonstrated a decremental STE pattern in V2-V4 and the absence of AVL STD. Serial ECG monitoring also did not show STE evolution into Q waves in both the precordial and limb leads (Figure 5 A - D). These findings are consistent with the above mentioned patterns for IRVI.

**Conclusion**

Our case highlighted the importance of recognizing the possibility of IRVI as part of the differential diagnosis when presented with an ECG pattern suggestive of an anterior wall myocardial infarction. This is a rare case demonstrating isolated anterior lead STE mimicking anterior wall MI after loss of a mid RCA AMB during PCI. Consideration of IRVI prevented repeat left coronary circulation angiography and thus spared the patient from additional contrast and radiation. In the acute setting of primary PCI, keeping IRVI in mind along with careful angiographic review and ECG guidance may help correctly identify the culprit vessel when angiography to both left and right coronary circulations are equivocal and when lesions are of equal severity.

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