2021

Effect of Presence of Ramus Intermedius Artery on Location of Culprit Lesions in Acute Left Circumflex Coronary Artery Occlusion

Follow this and additional works at: https://www.j-saudi-heart.com/jsha

Part of the Cardiology Commons

This work is licensed under a Creative Commons Attribution-Noncommercial-No Derivative Works 4.0 License.

Recommended Citation
El Zayat, Ahmed; Eldeeb, Mohey; Gad, Marwa; and Ibrahim, Ismail M. (2021) "Effect of Presence of Ramus Intermedius Artery on Location of Culprit Lesions in Acute Left Circumflex Coronary Artery Occlusion," Journal of the Saudi Heart Association: Vol. 33 : Iss. 1 , Article 12.
Available at: https://doi.org/10.37616/2212-5043.1238

This Original Article is brought to you for free and open access by Journal of the Saudi Heart Association. It has been accepted for inclusion in Journal of the Saudi Heart Association by an authorized editor of Journal of the Saudi Heart Association.
Effect of Presence of Ramus Intermedius Artery on Location of Culprit Lesions in Acute Left Circumflex Coronary Artery Occlusion

Ahmed El Zayat*, Mohey Eldeeb, Marwa Gad, Ismail M. Ibrahim

Cardiology Department, Faculty of Medicine, Zagazig University, Egypt

Abstract

Background and aim: Coronary artery anatomy frequently affects location of atherosclerotic plaques and subsequent culprit lesions. We sought to clarify whether presence or absence of Ramus Intermedius coronary artery (RI) would affect location of culprit lesions in acute left circumflex (LCX) coronary artery occlusion.

Methods: The study included 180 patients, 100 with a diagnosis of non-ST elevation myocardial infarction (NSTEMI) and 80 with ST elevation myocardial infarction (STEMI). All culprit lesions were located in the LCX coronary artery. RI group included 45 patients and the No RI group included 135 patients.

Results: Culprit LCX lesions were similarly located at a comparable distance from LCX ostium in both groups and the presence of RI was not associated with significantly more proximally located culprit LCX lesions (34.7 ± 15.2 mm compared to 30.8 ± 17.9 mm respectively, p > 0.05). The frequency distribution of culprit lesions’ distance from LCX ostium showed no significant difference between both groups in any of the segments studied (10 mm each). There was no significant difference between both groups regarding markers of myocardial necrosis size as cardiac biomarkers (peak cardiac troponin-T 1077.4 ± 361.2 pg/dl vs 926 ± 462.2 pg/dl respectively, p = 0.13), (peak creatine kinase-MB 232.2 ± 81 ng/dl vs 194.7 ± 99.2 ng/dl respectively, p = 0.07) or left ventricular ejection fraction (EF 46.3 ± 6.3% vs 48.3 ± 8.3% respectively, p = 0.76).

Conclusion: Presence of RI coronary artery, as an additional flow divider, may not be associated with more proximal culprit lesions, compared to its absence, in cases of acute LCX coronary artery occlusion. Possible underlying pathophysiologic mechanisms remain to be clarified.

Keywords: Left circumflex coronary artery, Ramus intermedius artery, Culprit lesion, STEMI, NSTEMI

1. Introduction

Coronary atherosclerosis and its complications as acute myocardial infarction (AMI) and stroke continue to be a major cause of morbidity and mortality worldwide [1]. It is well-studied that culprit coronary atherosclerotic lesions tend to cluster in the proximal coronary arteries [1,2]. Culprit lesions in AMI are also known to locate close to coronary bifurcations (flow dividers) and major curvatures that prompt atherosclerotic plaques to progress and finally rupture [1,2].

Acute myocardial infarctions secondary to acute left circumflex coronary artery occlusion frequently impose a clinical problem regarding diagnosis and accomplishing reperfusion in a timely fashion. This is mainly because of difficulties in diagnosis on initial electrocardiogram (ECG). This could explain the under-representation of acute left circumflex (LCX) occlusion in acute coronary syndrome cases requiring emergent coronary reperfusion [3–5]. However, it has also been suggested that the
underrepresentation of the LCX as a culprit artery could be actually because of the lower probability of LCX plaques to rupture. It was noticed that proximal LCX segments were even less prone to develop vulnerable plaques that subsequently rupture [6].

In their work, Galbraith et al., found that the presence of Ramus Intermedius (RI) was associated with more proximal left anterior descending coronary artery (LAD) lesions and hence larger anterior infarctions. They suggested that anatomy-induced flow disturbances (by RI as an additional flow divider) have important clinical implications [7]. We sought to study whether same findings of the previous study [7] could be replicated with the LCX as the culprit artery, i.e. whether presence or absence of RI coronary artery would affect location of culprit lesions in acute left circumflex (LCX) coronary artery occlusion.

2. Methods

This prospective cross-sectional study was conducted on patients with acute myocardial infarction (AMI) who were admitted to the coronary care unit of cardiology department, Zagazig University Hospitals, Egypt in the period from January 2017 to February 2020. Included patients had coronary angiography during admission for either primary percutaneous coronary intervention (PPCI) in cases of ST elevation myocardial infarction (STEMI) or non-primary PCI in cases of non-ST elevation myocardial infarction (NSTEMI), with the culprit lesions located in the LCX artery. AMI was diagnosed as per current universal definition for acute myocardial infarction [8].

The study included 180 patients, 100 patients with a diagnosis of NSTEMI and 80 patients with STEMI. Inclusion criteria (1) coronary angiography done during the index hospitalization for AMI. (2) Culprit lesions located in the LCX coronary artery. AMI was diagnosed as per current universal definition for acute myocardial infarction [8].

Exclusion criteria were (1) previous coronary artery bypass grafting (2) acute LCX occlusion with undelaying etiology other than coronary atherosclerotic plaque with superimposed thrombosis (as dissection or embolism) (3) no clearly identifiable culprit lesion (4) previous angioplasty at the site of the culprit lesion.

Study patients were divided into two groups according to the presence or absence of RI coronary artery. Ramus Intermedius coronary artery was said to be present in left main coronary artery trifurcations with the intermediate artery ostial diameter \( \geq 1 \) mm [7]. All RI arteries were identified in the left anterior oblique (LAO) caudal view (spider view) (Fig. 1). RI was present in 45 patients (25%) and absent in 135 (75%) patients.

Written informed consent was obtained from all patients and the study was approved by the local ethical committee and the institutional review board.

Acute LCX occlusion was diagnosed by the absence of anterograde flow i.e. Thrombolysis in myocardial infarction (TIMI) 0/1, and/or presence of LCX thrombus. Cases with TIMI flow grade \( \geq 2 \) were also included, provided that there was ST elevation on the ECG and/or presence of LCX thrombus on the atherosclerotic lesion (9 patients had TIMI = 2 flow).

Quantitative coronary angiography (QCA) analysis of LCX culprit lesion was performed using a standard software package (Q angio XA, Medis, Leiden, the Netherlands). For vessel classification, Coronary Artery Surgery Study (CASS) coronary map was used [9] and to avoid and minimize vessel foreshortening, right anterior oblique view with caudal angulation was used for measurement of distance from LCX ostium to the culprit LCX lesion [9].

Distance from the ostium of the LCX coronary artery to LCX culprit lesion was assessed (in millimeters) and defined as the distance from the ostium to the part of lesion with highest degree of percent stenosis (on QCA) [6].

Coronary angiograms were reported by two independent reviewers who were blinded to the clinical and ECG data.

Markers of myocardial necrosis were measured and compared between both studied groups, which included peak troponin-T, peak creatine kinase-MB and echocardiography-assessed left ventricular ejection fraction (LVEF).

2.1. Statistical analysis

Statistical analyses were performed using SPSS software (version 20.0, SPSS, Chicago, IL).
Fig. 1. Two examples of LCX culprit lesions with RI coronary artery present (A) and absent RI (B) shown in LAO caudal (spider) view.

| Character                          | RI (n = 45)   | No RI (n = 135) | p value |
|----------------------------------|---------------|----------------|---------|
| Age (years)                      | 54.4 ± 11.2   | 57.1 ± 13.7    | 0.41    |
| Sex (male) (n, %)                | 32 (73.3%)    | 105 (77.7%)    | 0.15    |
| Hypertension (n, %)              | 27 (60)       | 74 (54.8)      | 0.55    |
| Diabetes Mellitus (n, %)         | 11 (24.4)     | 27 (20)        | 0.32    |
| Smoking (n, %)                   | 23 (51.1)     | 61 (45.1)      | 0.14    |
| Dyslipidemia (n, %)              | 21 (46.6)     | 54 (40)        | 0.86    |
| Family history of CAD (n, %)     | 12 (26.6)     | 28 (20.7)      | 0.45    |
| Previous MI                      | 11 (24.4)     | 30 (22.2)      | 0.82    |
| Heart rate at presentation (b/m) | 87.4 ± 12.6   | 92.8 ± 18.4    | 0.64    |
| Total ischemic times (symptoms- to-balloon) (min) | 289.5 ± 114.2 | 276.6 ± 141.8 | 0.14 |
| MVD (n, %)                       | 29 (64.4%)    | 92 (68.1)      | 0.85    |

Clinical events (1-month):

Heart failure (1-month) (n, %) 9 (20) 30 (22.2) 0.76
All-cause mortality (1-month) (n, %) 4 (8.9) 11 (8.1) 0.86
Cardiac death (1-month) (n, %) 4 (8.9) 11 (8.1) 0.86

Long-term medications:

Beta Blockers (n, %) 15 (33.3) 53 (39.2) 0.62
ACE/ARB (n, %) 12 (26.7) 30 (22.2) 0.78
Statins (n, %) 8 (17.8) 18 (13.3) 0.65
Aldosterone Receptor Blockers (n, %) 7 (15.6) 18 (13.3) 0.69
Aspirin (n, %) 18 (40) 61 (45.1) 0.31

Medications during hospital admission:

Beta Blockers (n, %) 37 (82.2) 106 (78.5) 0.76
ACE/ARB (n, %) 23 (51.1) 75 (56.1) 0.62
Aldosterone Receptor Blockers (n, %) 27 (60) 73 (54.1) 0.52
Dual anti-platelets (n, %) 45 (100) 135 (100) 1
Statins (n, %) 36 (80) 117 (86.7) 0.48
Dual anti-platelets after discharge 45 (100) 133 (98.5) 0.92

ACE/I: Angiotensin converting enzyme inhibitors. ARB: Angiotensin receptor blockers. MVD (multi-vessel disease, other non-culprit significant plaques, >70% stenosis present in same or other vessels).
Quantitative variables were assessed as mean ± SD, while qualitative ones as numbers and percentages. Chi-square test was used to assess qualitative variables and T test for quantitative variables (study data showed normal distribution). A two tailed P-value < 0.05 was considered significant.

3. Results

There was no significant difference between RI and No-RI groups regarding any of the demographic criteria, total ischemic times, long term medications, medications used during course of hospital admission, use of anti-platelets or 1-month clinical events as shown in Table 1. There was also no significant difference between both groups regarding markers of myocardial necrosis as cardiac biomarkers (cardiac troponin and CK-MB) and LVEF as shown in Table 2. There was also no significant difference between both groups regarding the presence of multivessel disease with significant luminal stenosis >70% in the culprit or other vessels (Table 2).

Table 3 shows that culprit LCX lesions were similarly located at a comparable distance from the LCX ostium in both groups and that the presence of RI was not associated with more proximally located culprit LCX lesions.

The frequency distribution of culprit lesions’ distance from the LCX ostium showed no significant difference between both groups in any of the studied segments (10 mm each). Culprit lesions in both groups were mainly located 20–40 mm from the LCX ostium (75.5% in the RI group compared to 76.9% in the No RI group) as shown in Table 4.

4. Discussion

Plaque rupture is related to intrinsic composition of the individual plaques (i.e. vulnerability) as well as extrinsic forces exerted on plaques (rupture triggers) [10,11]. Intrinsic forces predispose plaques to rupture, whereas extrinsic forces may precipitate disruption if vulnerable plaques are present [12,13]. Pathology studies have shown that >50% of vulnerable plaques were proximally located in coronary tree, one third in midportion, and the rest were in the distal portion of these arteries [14]. Plaque rupture was most frequently encountered in the proximal LAD (66%) followed by RCA (18%) and least found in the LCX (14%) [14]. Many observational studies have validated low shear stress hypothesis of plaque initiation and progression which involved mainly outer side of bifurcations and regions of major curvatures in the arterial tree, where shear stress is lowest [15–19]. On the other hand, plaque rupture and its relation to local shear stress seems to be more complex. In general, plaque rupture is most frequently encountered at area of cap shoulders where shear stress is relatively higher enough to cause endothelial damage followed by rupture of the already thin cap fibroatheroma [20].

We found no significant difference regarding distribution of culprit lesions in LCX coronary artery in presence or absence of RI coronary artery. It was rather expected that culprit LCX lesions would be more proximally located in cases of RI presence, as was the case in LAD culprit lesions in STEMI in the presence of RI [7]. This difference might be peculiar.
to proximal LCX segments, that have been possibly attributed to LCX coronary artery having lowest longitudinal strain compared to other epicardial coronary segments [6]. Insignificant difference regarding location of LCX culprit lesions in both groups also explains the almost similar extent of myocardial necrosis markers as reflected by non-significant differences in peak cardiac biomarkers and left ventricular ejection fraction. This was also reflected on non-significant difference in 1-month hard events as heart failure and mortality. Many confounders that could affect plaques composition, location and vulnerability (as co-morbidities like hypertension, diabetes mellitus, dyslipidemia, etc.) were present in both groups albeit with no significant difference.

Epicardial coronaries are cyclically subjected to both circumferential and longitudinal deformations that might contribute to destabilization and eventual rupture of vulnerable plaques [21]. In their work to study LCX as the culprit artery in STEMI and to define the distribution of such culprit lesions in relation to other two major epicardial coronaries [6], D. Ghanem et al. concluded that LCX plaques seemed less prone to rupture and that its proximal segment was even less so. They also studied LCX culprit lesions in NSTEMI and stable angina and found that culprit and total disease distribution were similar in NSTEMI and chronic stable angina. They suggested that lower LCX longitudinal strain (and subsequently relatively lower shear stress) might have contributed to reduced plaque rupture in STEMI [6]. Using a coronary computed tomographic angiographic (CCTA) database, they created a dynamic 3-dimensional coronary artery anatomic model [6] with retrospective ECG gating. They found that in the studied cases, the mean LAD systolic longitudinal strain was $9.5\% \pm 2.9\%$ and for RCA was $10.1\% \pm 3.9\%$, compared with $15.5\% \pm 2.4\%$ for the LCX ($P < 0.001$). They suggested that significant difference in shortening (referred to as squeezing theory) has contributed substantially to the lower prevalence of the proximal LCX culprit lesions in STEMI [6].

We found that majority of culprit LCX lesions in both groups were located 20–40 mm from the LCX ostium. This was similar to the findings of Katritsis et al. who found that culprit LCX lesions were located 29.73 ± 14.39 mm from the LCX ostium [22].

Presence of coronary segments where plaques are relatively less liable to rupture is very inviting for both clinical and basic research studies to explore possible mechanisms. Differential shear stress and longitudinal strain along specific segments of coronary tree might be reflected on abundance of local mediators with subsequent effects on plaque progression and vulnerability. This could have potential therapeutic implications in both atherosclerotic plaque progression and acute coronary syndromes management.

5. Limitations of the study

This work has some limitations. First, this was a single center study with small number of patients. Second, the presence of many confounders that could affect plaque location, morphology and vulnerability (age, hypertension, diabetes mellitus, dyslipidemia, etc.). Clustering of many risk factors of CAD in the same patient could have affected plaques’ progression and characteristics. Third, infarct size should have been assessed by more accurate imaging modalities as cardiac magnetic resonance imaging (CMR).

6. Conclusion

Presence of RI coronary artery, as an additional flow divider, may not be associated with more proximal culprit lesions, compared to its absence, in cases of acute LCX coronary artery occlusion.

Author's contribution

Conception and design of Study: Ahmed El Zayat, Mohey Eldeeb. Literature review: Ahmed El Zayat, Mohey Eldeeb, Marwa Gad. Acquisition of data: Mohey Eldeeb, Marwa Gad, Ismail M Ibrahim. Analysis and interpretation of data: Mohey Eldeeb, Marwa Gad, Ismail M Ibrahim. Research investigation and analysis: Ahmed El Zayat, Mohey Eldeeb, Ismail M Ibrahim. Data collection: Mohey Eldeeb, Marwa Gad. Drafting of manuscript: Marwa Gad. Revising and editing the manuscript critically for important intellectual contents: Ahmed El Zayat, Mohey Eldeeb, Marwa Gad. Data preparation and presentation: Mohey Eldeeb, Marwa Gad, Ismail M Ibrahim. Supervision of the research: Ahmed El Zayat, Mohey Eldeeb, Marwa Gad, Ismail M Ibrahim. Research coordination and management: Ahmed El Zayat, Mohey Eldeeb, Marwa Gad.

Conflicts of interest

None to declare.

Funding

None.
References

[1] Wang JC, Normand SL, Mauri L, et al. Coronary artery spatial distribution of acute myocardial infarction occlusions. Circulation 2004;110:278–84. https://doi.org/10.1161/01.CIR.0000135468.67850.F4.

[2] Gibson CM, Kirtane AJ, Murphy SA, et al. Distance from the coronary ostium to the culprit lesion in acute ST-elevation myocardial infarction and its implications regarding the potential prevention of proximal plaque rupture. J. Thorb. Thrombolysis 2003;15:189–96. https://doi.org/10.1023/B:THRO.0000011374.60110.bc.

[3] Huey BL, Beller GA, Kaiser DL, Gibson RS. A comprehensive analysis of myocardial infarction due to left circumflex artery occlusion: comparison with infarction due to right coronary artery and left anterior descending artery occlusion. J. Am. Coll. Cardiol. 1988;12:1156–66. https://doi.org/10.1016/0735-1097(88)92594-6.

[4] Wellens HJJ, Conover MB. The ECG in Emergency Decision Making. St. Louis: Saunders Elsevier; 2006. p. 304. ISBN: 9781416002598.

[5] O’Gara PT, Kushner FG, Ascheim DD, et al. ACCF/AHA guideline for the management of ST elevation myocardial infarction. J. Am. Coll. Cardiol. 2013;61:e78–140. https://doi.org/10.1016/j.jacc.2012.11.019.

[6] Ghanim D, Kusniec F, Kinany W, Qarawani D, Meerkin D, O’Gara PT, Kushner FG, Ascheim DD, et al. ACCF/AHA guideline for the management of ST elevation myocardial infarction. J. Am. Coll. Cardiol. 2013;61:e78–140. https://doi.org/10.1016/j.jacc.2012.11.019.

[7] Galbraith EM, McDaniel MC, Jeroudi AM, Kashlan OR, Taha K, Amir O, Carasso S. Left circumflex coronary artery as the culprit vessel in ST-segment-elevation myocardial infarction. J. Am. Coll. Cardiol. 2013;61:e78–140. https://doi.org/10.1016/j.jacc.2012.11.019.

[8] Thysesen Kristian, Alpert Joseph S, Jaffe Allan S, Chaitman Bernard R, Bax Jeroen J, Morrow David A, White Harvey D. Fourth universal definition of myocardial infarction (2018): ESC Scientific Document Group. Eur. Heart J. 2019;40(3):237–69.

[9] Alderman E, Stadius M. The angiographic definitions of the bypass angioplasty revascularization investigation. Coron. Artery Dis. 1992;3:1189–207. Corpus ID: 78859841.

[10] Mann J, Davies MJ. Mechanisms of progression in native coronary artery disease: role of healed plaque disruption. Heart 1999;82:265–8. https://doi.org/10.1136/hrt.82.3.265.

[11] Alderman EL, Corley SD, Fisher LD, Chaitman BR, Faxon DP, Foster ED, et al. Five-year angiographic follow-up of factors associated with progression of coronary artery disease in the coronary artery surgery study (CASS). CASS participating investigators and staff. J. Am. Coll. Cardiol. 1993;22:1141–54. https://doi.org/10.1016/0735-1097(93)90429-5.

[12] Falk E, Shah PK, Fuster V. Coronary plaque disruption. Circulation 1995;92:657–71. https://doi.org/10.1161/01.CIR. 92.3.657.

[13] Lee RT, Kamm RD. Vascular mechanics for the cardiologist. J. Am. Coll. Cardiol. 1994;23:1289–95. https://doi.org/10.1016/0735-1097(94)90369-7.

[14] Virmani R, Burke AP, Farb A, Kolodgie FD. Pathology of the vulnerable plaque. J. Am. Coll. Cardiol. 2006;47:C13–8. https://doi.org/10.1016/j.jacc.2005.10.065.

[15] Asakura T, Karino T. Flow patterns and spatial distribution of atherosclerotic lesions in human coronary arteries. Circ. Res. 1990;66:1045–66. https://doi.org/10.1161/01.res.66.4.1045.

[16] Bharadavaj BK, Mabon RF, Giddens DP. Steady flow in a model of the human carotid bifurcation. Part I — flow visualization. J. Biomech. 1982;15:349–62. https://doi.org/10.1016/0021-9290(82)90057-4.

[17] Motomiya M, Karino T. Flow patterns in the human carotid artery bifurcation. Stroke 1984;15:50–6. https://doi.org/10.1161/01.str.15.1.50.

[18] Zarinis CK, Giddens DP, Bharadavaj BK, Sottiraii VS, Mabon RF, Glagov S. Carotid bifurcation atherosclerosis. Quantitative correlation of plaque localization with flow velocity profiles and wall shear stress. Circ. Res. 1983;53: 502–14. https://doi.org/10.1161/01.res.53.5.502.

[19] Davies PF. Flow-mediated endothelial mechanotransduction. Physiol. Rev. 1995;75:519–60. https://doi.org/10.1152/phys- rev.1995.75.3.519. Kat 38-42.

[20] Mattsson EJ, Kohler TR, Vergel SM, Clowes AW. Increased blood flow induces regression of intimal hyperplasia. Arterioscler. Thromb. Vasc. Biol. 1997;17:2245–9. https://doi.org/10.1161/01.atv.17.10.2245.

[21] Corban MT, Eshtehardi P, Suo J, McDaniel MC, Timmins LH, Rassoul-Arzrumly E, et al. Combination of plaque burden, wall shear stress, and plaque phenotype has incremental value for prediction of coronary atherosclerotic plaque progression and vulnerability. Atherosclerosis 2014; 232(2):271–6. https://doi.org/10.1016/j.atherosclerosis.2013.11.049.

[22] Katrakis DG, Efstatopoulos EP, Pantos J, Korovesis S, Koulaba G, Kazantzidis S, Marmarelis V, Voridis E. Anatomic characteristics of culprit sites in acute coronary syndromes. J. Intervent. Cardiol. 2008;21:140–50. https://doi.org/10.1111/j.1540-8183.2007.00339.x.