



73 with late gadolinium enhancement sequences confirming  
74 ischemic injury [9].  
75 The novelty of this series lies in highlighting the diag-  
76 nostic journey of an adolescent and a young adult whose  
77 cardiac symptoms were initially overlooked. We empha-  
78 size the use of stress perfusion CMR and LGE sequences  
79 as the essential 'keys' to uncovering functional coronary  
80 disorders in a demographic typically considered low-risk  
81 by conventional standards.

### Case Presentation

82

#### Case 1 the adolescent with an adult's diagnosis

83

84 A 16-year-old male, previously healthy and physically  
85 active presented with sudden episodes of palpitations and  
86 left arm pain for a few hours. These were unprovoked,  
87 occurring at rest and occasionally accompanied by a brief,  
88 sharp chest discomfort. He had no history of hypertension,  
89 diabetes, smoking, obesity, or family history of premature

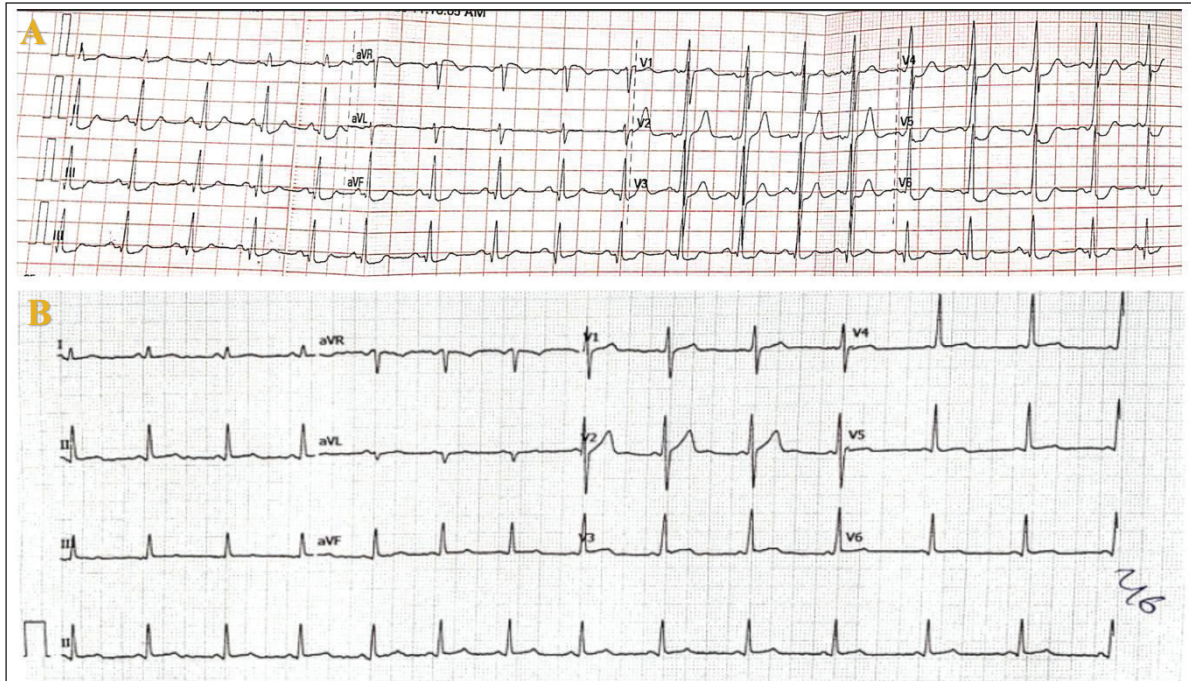


Figure 1. (A) ECG done at the time of symptoms showing sinus tachycardia and ST-T changes. (B) ECG done at Zydus hospital showing normal sinus rhythm and no significant ST-T changes.

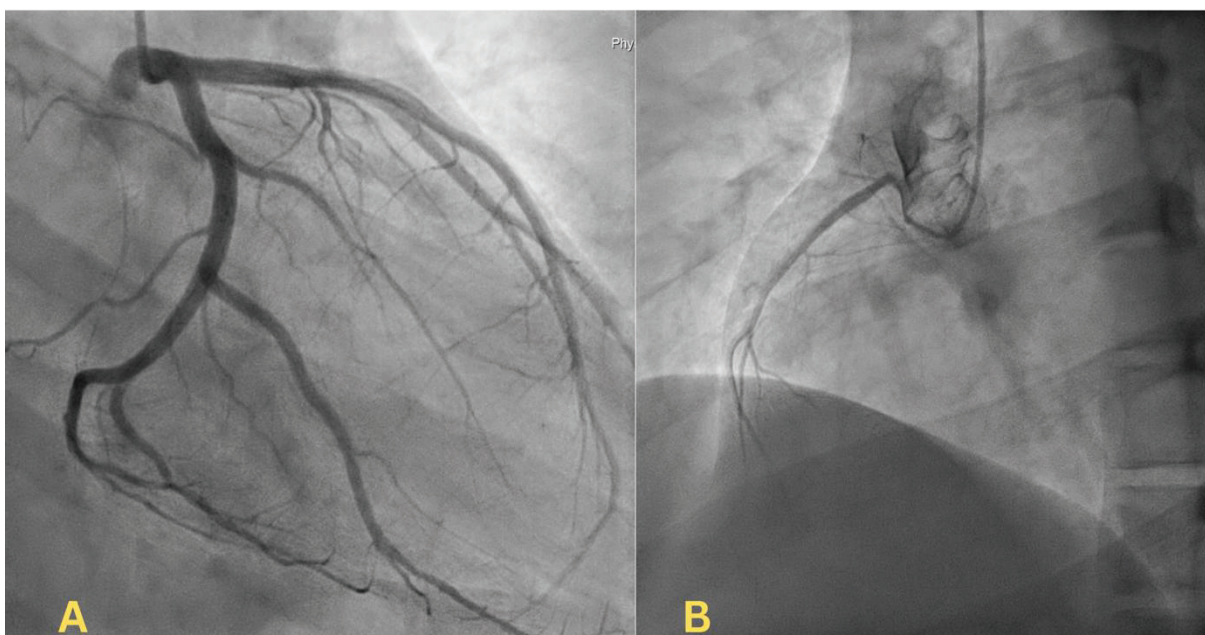
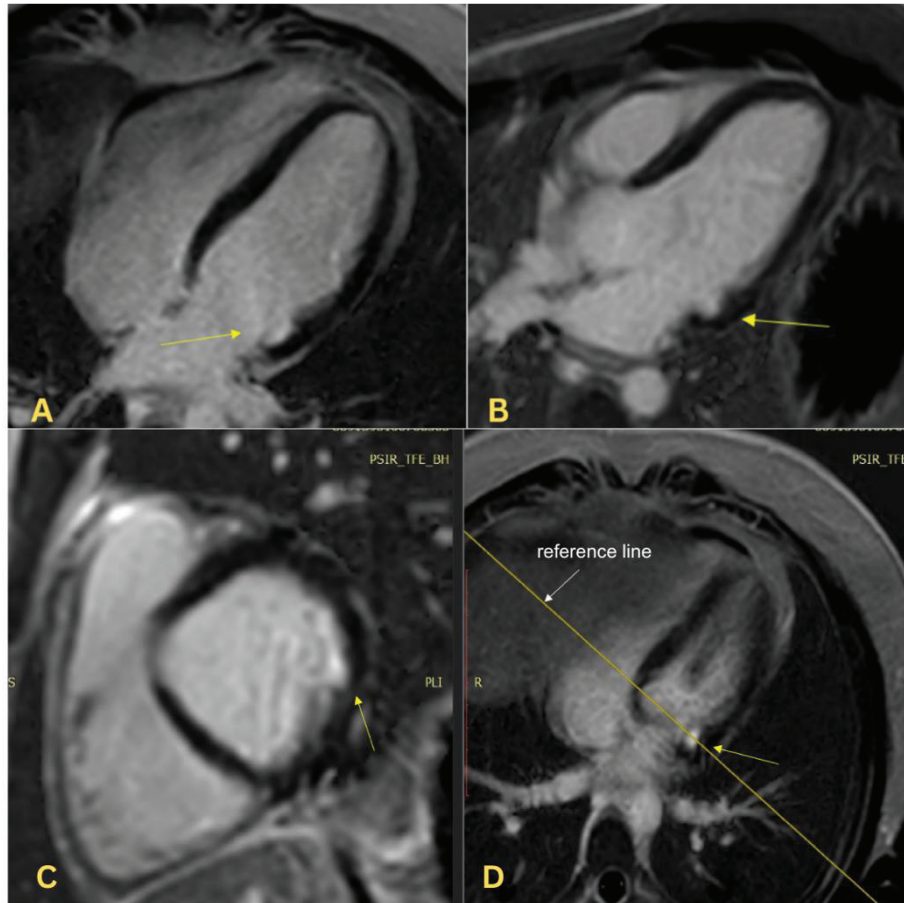
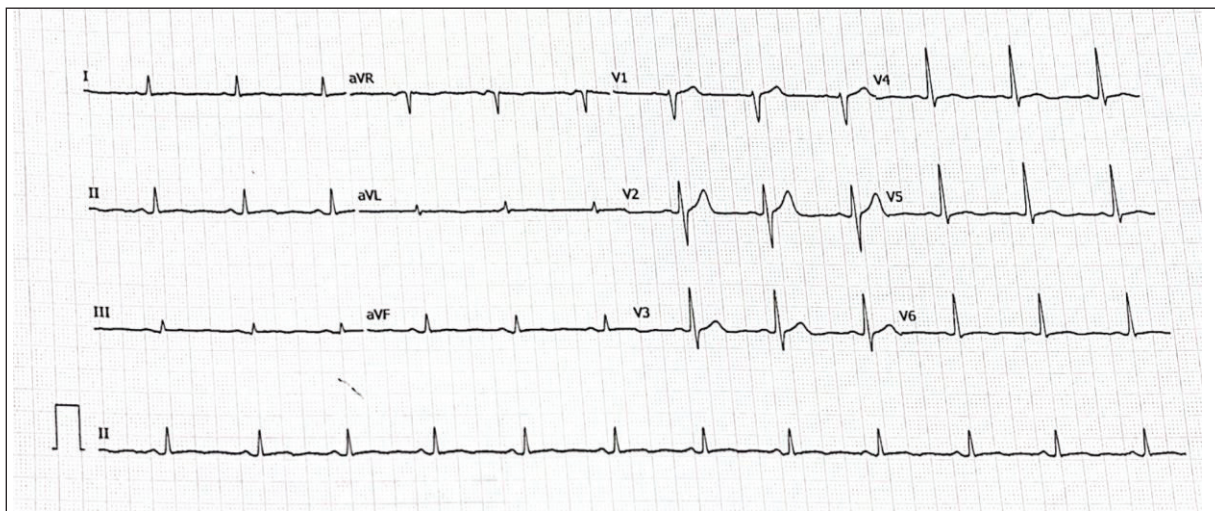


Figure 2. CAG of Case 1 (A) Dominant Left coronary artery system showing normal epicardial vessel (B) Non dominant normal Right coronary artery.



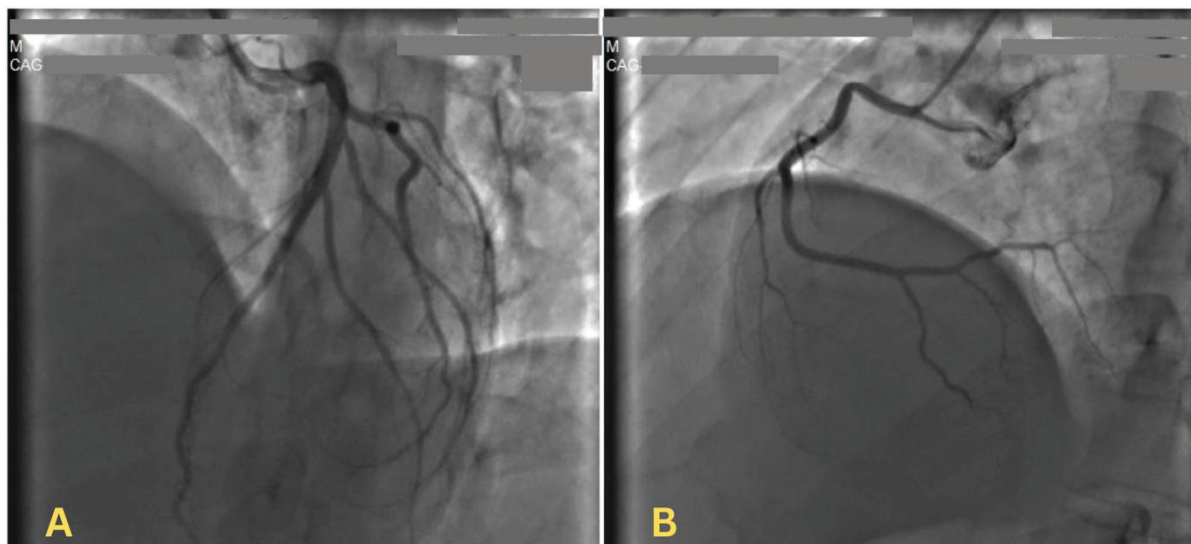
**Figure 3.** CMR-LGE images of Case 1. (A) Four-chamber and (B) three-chamber views demonstrate focal subendocardial enhancement (arrow) in the basal inferolateral wall. (C) Short-axis view showing focal subendocardial enhancement (arrow). (D) Four-chamber view with reference line correlating the short-axis slice location.



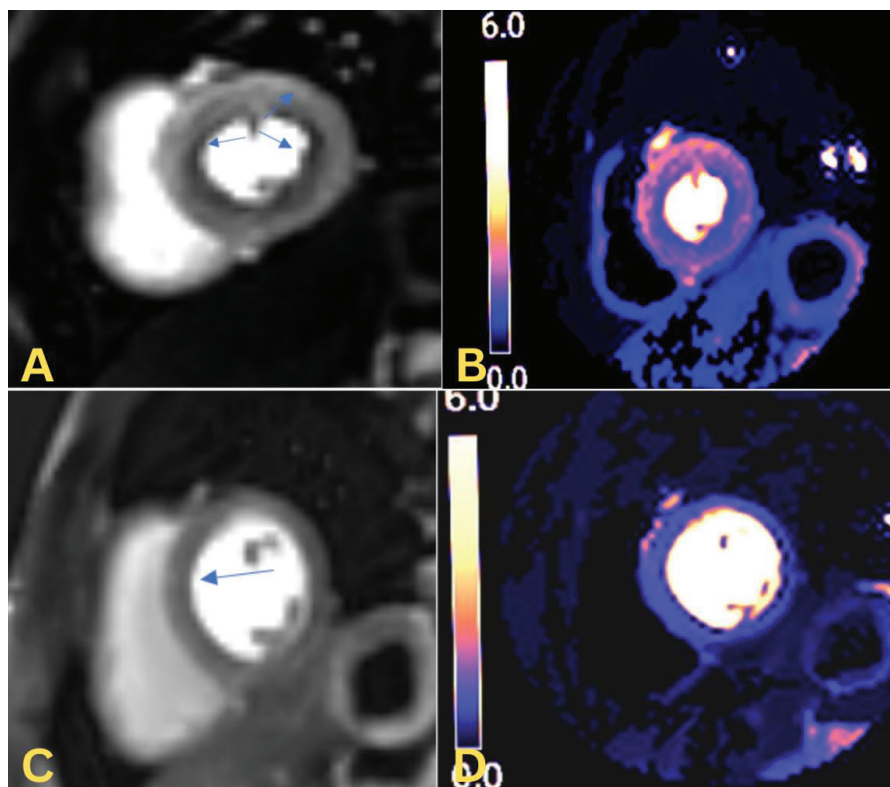
**Figure 4.** Case 2 ECG at the hospital showing NSR and no ST-T changes.

90 coronary disease. His initial evaluations including physical  
91 examination, ECG, ECHO, and cardiac biomarkers  
92 were unremarkable. He was reassured and sent home with  
93 a low-dose beta-blocker.

Nine months later, recurrent paroxysmal palpitations with  
94 left arm discomfort and chest pain at rest prompted readmission.  
95 The first of the serial ECGs obtained at symptom onset  
96 demonstrated sinus tachycardia with transient ST-segment  
97



**Figure 5.** CAG of Case 2 (A) Left coronary system showing normal epicardial vessels (B) Dominant right coronary artery showing normal epicardial vessel.



**Figure 6.** Stress perfusion cardiac magnetic resonance (CMR) in Case 2. (A) Short-axis stress perfusion imaging demonstrating a global subendocardial perfusion defect (arrows). (B) Quantitative perfusion mapping during stress revealed reduced myocardial blood flow (MBF 1.54 ml/min/g) and a mildly reduced myocardial perfusion reserve index (MPRI 2.04). (C) Rest perfusion imaging showed no fixed defects (arrow). (D) Rest perfusion mapping confirmed preserved myocardial blood flow (MBF 0.75 ml/min/g).

98 depression in leads V4–V6 (Figure 1A). A subsequent ECG  
 99 at our hospital showed normal sinus rhythm with no signifi-  
 100 cant ST segment/ T wave changes (Figure 1B).  
 101 Laboratory investigations revealed elevated Troponin  
 102 I of 0.871 ng/ml and High sensitivity troponin I of 614  
 103 ng/l (reference <14 ng/l). Physical examination showed

blood pressure 133/76 mmHg and heart rate 83 beats per 104  
 minute. ECHO confirmed preserved systolic function and 105  
 no structural cardiac abnormalities. 106

Comprehensive laboratory workup excluded systemic 107  
 inflammation (C-Reactive Protein normal), dyslipidemia 108  
 (LDL 101 mg/dl, HDL 37 mg/dl, triglycerides 188 mg/ 109

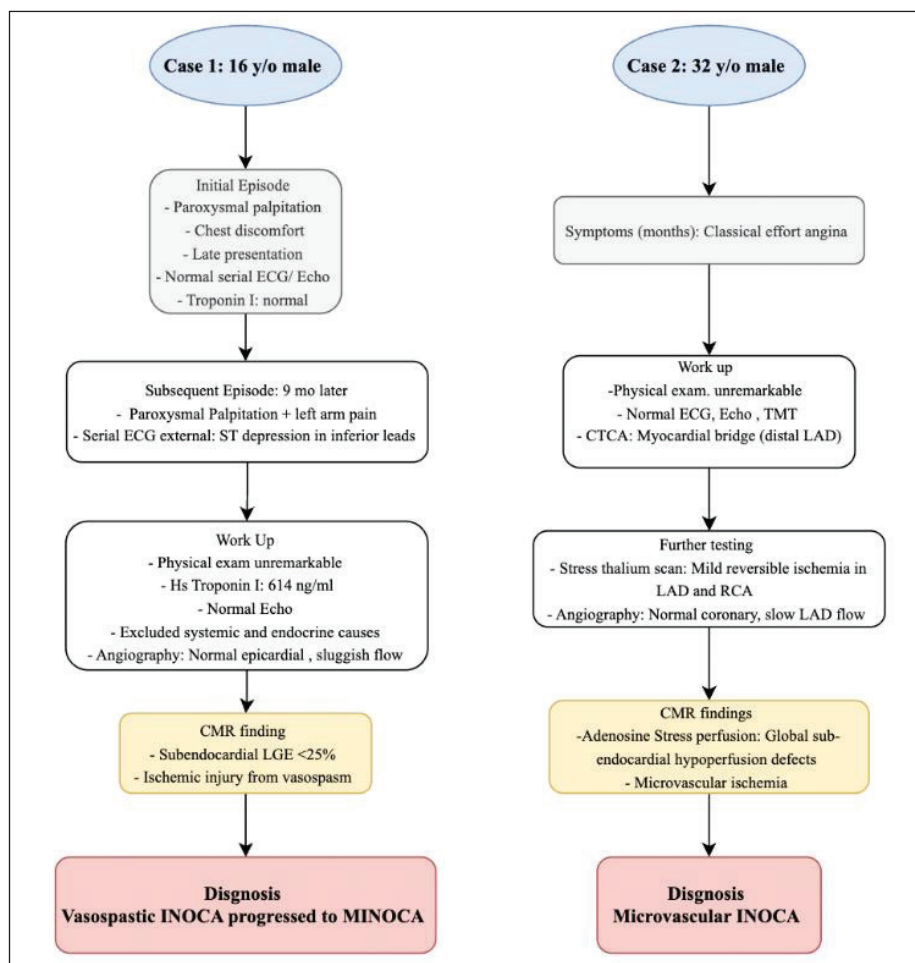


Figure 7. Flow chart summarizing the case series.

110 dl), endocrine disorders, catecholamine-secreting tumors (plasma metanephrines negative) and hematological abnormalities.

113 Coronary angiography was performed, which demonstrated normal epicardial vessels with diffusely sluggish flow (Figure 2).

116 In the absence of anatomical obstruction to explain the myocardial injury, the team moved to CMR for further evaluation, which showed good biventricular systolic function with normal chamber sizes, but revealed a focal area of subendocardial LGE in the basal inferolateral wall affecting less than 25% of the wall's thickness Figure 3. This pattern confirmed a limited subendocardial myocardial infarction in the absence of obstructive coronary artery disease, supporting the diagnosis of MINOCA attributable epicardial or microvascular spasm. The patient was started on a calcium channel blocker, anti-anginal therapy, and antiplatelet agents. At the 6 months follow-up, the patient remains asymptomatic with no further anginal episodes and is doing well clinically.

129 **Case 2 the young adult with invisible yet bothering**  
130 **angina**

131 A 34-year-old physically active male without cardiovascular risk factors or family history of premature coronary  
132

artery disease, presented with a four-month history of classical angina which were characterized by exertional chest pain, dyspnea on exertion, and occasionally a sense of suffocation. Symptoms were reproducible with physical activity and resolved with rest.

138 Six weeks prior, a routine health evaluation included normal resting ECG, ECHO of ejection fraction 60%, no wall motion abnormalities, and normal TMT. At our hospital, a resting ECG again showed no abnormalities (Figure 4).

143 Laboratory investigations and high-sensitive troponin I were normal.

145 A computed tomography coronary angiography (CTCA) was performed which identified a minor myocardial bridge over the distal left anterior descending artery (LAD) with otherwise normal appearing coronary tree. Given the ongoing exertional symptoms, a stress myocardial perfusion imaging (MPI) revealed mild reversible ischemia in both the territories of left anterior descending and right coronary arteries.

153 Invasive coronary angiography demonstrated normal epicardial vessels with no obstructive lesions, but slow coronary flow in the LAD (TIMI 2 flow) (Figure 5).

156 To further investigate for microvascular ischemia, a  
 157 stress cardiac magnetic resonance (CMR) imaging was  
 158 performed (Figure 6) using adenosine infusion (140 mcg/  
 159 kg/min) and gadolinium-based contrast. The study showed  
 160 a mildly reduced LVEF (~53%) without RWMA. During  
 161 stress, a uniform global subendocardial perfusion defect  
 162 was observed in the left ventricle. Quantitative perfusion  
 163 mapping demonstrated reduced stress myocardial blood  
 164 flow (MBF 1.54 ml/min/g; Normal value 2.54 ml/min/g)  
 165 and a mildly reduced myocardial perfusion reserve index  
 166 (MPRI 2.04). LGE sequences showed no evidence of scar  
 167 or fibrosis. Findings are consistent with global microvas-  
 168 cular ischemia in the absence of obstructive epicardial  
 169 disease, leading to a diagnosis of microvascular INOCA.  
 170 The patient was commenced on a diltiazem, nicorandil,  
 171 ranolazine, trimetazidine, and atorvastatin. At 6 months  
 172 follow up he remained asymptomatic.

173 **Discussion**

174 In adolescents, the rarity of atherosclerosis makes vasomo-  
 175 tor dysfunction the leading cause of ischemia. Evidence  
 176 of increased smooth-muscle reactivity and endothelial  
 177 impairment in young patients suggests that coronary  
 178 spasm is a likely mechanism of chest pain occurring at  
 179 rest [5,10]. In chronic coronary syndromes, ischemia  
 180 reflects an imbalance between coronary flow and myo-  
 181 cardial demand.[4] Beyond obstructive disease, impaired  
 182 microvascular dilation or inappropriate vasoconstriction  
 183 can produce this mismatch, underscoring the need for  
 184 mechanism-directed therapy [2,6].

185 These two cases illustrate distinct forms of ischemia  
 186 without obstructive CAD Figure 7. The adolescent  
 187 demonstrates epicardial vasospasm, transient and severe  
 188 vasoconstriction driven by vascular smooth muscle hyper-  
 189 activity, capable of causing myocardial necrosis. The focal  
 190 subendocardial LGE on CMR confirmed limited myocar-  
 191 dial injury from prolong vasospastic episode. In contrast,  
 192 the young adult demonstrates CMD, specifically impaired  
 193 by vasodilatory capacity of microcirculation, revealed by  
 194 stress thallium and stress CMR. The small myocardial  
 195 bridge identified in the distal LAD could not account for  
 196 ischemia in the RCA territory, indicating that CMD was  
 197 the primary driver [11].

198 These pathophysiologic distinctions mandated diver-  
 199 gent management approaches. For the vasospastic  
 200 MINOCA, therapy focused on preventing epicardial  
 201 spasm through calcium channel blockers (diltiazem) as  
 202 first-line treatment, supplemented by nicorandil as an  
 203 additional antispastic agent. Conversely, for the microvas-  
 204 cular INOCA, treatment targeted improving microvascu-  
 205 lar vasodilatory capacity and reducing myocardial oxygen  
 206 demand. The multi-drug regimen included nicorandil  
 207 (direct microvascular vasodilator), ranolazine (particu-  
 208 larly effective in CMD with reduced CFR), and trimetazi-  
 209 dine (metabolic modulator) [4].

While 6-month follow-up confirms sustained symptom  
 resolution, the primary limitation remains the lack of long-  
 term longitudinal data in this young cohort. Additionally,  
 the specialized requirement for stress CMR with quanti-  
 tative perfusion mapping may restrict generalizability in  
 resource-constrained settings.

Nevertheless, the cases show that non-obstructive  
 ischemia can range from microvascular-mediated angina  
 to infarction and underscore the value of precision imag-  
 ing. From a diagnostic perspective the cases emphasize the  
 importance of a stepwise diagnostic approach: excluding  
 obstructive disease, performing structured physiological  
 assessment, and adopting advanced functional imaging to  
 define ischemic mechanism [5]. This prevents unnecessary  
 procedures and therapeutic under-treatment, associated with  
 long term anxiety and poor quality of life [1]. While intra-  
 coronary acetylcholine testing remains the gold standard  
 for diagnosing vasospasm in MINOCA [12], our experi-  
 ence demonstrates that advanced CMR provides a safe and  
 informative alternative when invasive testing is not feasible.

**Conclusion**

These two cases challenge the assumption that myocardial  
 ischemia is the domain of older patients with well-defined  
 risk factors. INOCA can emerge quietly, in individuals at  
 the peak of health, and remain invisible on routine testing  
 until advanced imaging reveals its presence. Both pres-  
 entations underscore a gap in clinical awareness: without  
 considering vasospastic and microvascular mechanisms,  
 patients may be reassured too soon, delaying diagnosis  
 and leaving them vulnerable to ongoing ischemia and its  
 potential complications. Ultimately, these cases demon-  
 strate that the diagnostic ‘key’ for the young, symptomatic  
 patient is the transition from anatomical assessment to  
 functional characterization. By heightened awareness,  
 informed suspicion, and early use of sensitive imaging  
 we move past the common pitfall of ‘atypical’ labels and  
 toward mechanism-specific care that can fundamentally  
 change a patient’s cardiovascular trajectory.

**What is new?**

Young patients presenting with chest pain are often consid-  
 ered as non-cardiac particularly when basic conventional  
 reports are normal. This case series highlights the importance  
 of appropriate clinical history, detailed investigation and  
 advanced imaging in finding out subtle yet clinically relevant  
 and sometimes fatal cardiac conditions in these patients

**List of abbreviations**

ACS	Acute Coronary Syndrome	256
ANOCA	Angina with Non-Obstructive Coronary Arteries	257
CAD	Coronary Artery Disease	258
CAG	Coronary Angiography	259
CAS	Coronary Artery Spasm	260
CBC	Complete Blood Count	261
CMD	Coronary Microvascular Dysfunction	262

263 CMR Cardiac Magnetic Resonance  
 264 CRP C-reactive Protein  
 265 CTCA Computed Tomography Coronary Angiography  
 266 ECG Electrocardiogram  
 267 ECHO Echocardiography  
 268 HDL High-Density Lipoprotein  
 269 INOCA Ischemia with non-obstructive coronary arteries  
 270 LAD Left Anterior Descending (artery)  
 271 LDL Low-Density Lipoprotein  
 272 LGE Late Gadolinium Enhancement  
 273 LVEF Left Ventricular Ejection Fraction  
 274 MBF Myocardial Blood Flow  
 275 MINOCA Myocardial infarction with non-obstructive coronary arteries  
 276  
 277 MPI Myocardial perfusion Imaging  
 278 MPRI Myocardial Perfusion Reserve Index  
 279 MVA Microvascular Angina  
 280 RCA Right Coronary Artery  
 281 RWMA Regional Wall Motion Abnormalities  
 282 TIMI Thrombolysis In Myocardial Infarction (flow grade)  
 283 TMT Treadmill Test  
 284 TSH Thyroid Stimulating Hormone  
 285 VSA Vasospastic angina

286 **Conflict of interest**

287 The authors declare that there is no conflict of interest regarding  
 288 the publication of this article.

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291 **Consent for publication**

292 Written informed consent was obtained from the patient /from  
 293 the parents of the patient.

294 **Ethical Approval**

295 Ethical approval is not required at our institution to publish an  
 296 anonymous case series.

297 **Author details**

298 Naqiya Arsiwala<sup>1</sup>, Abhisheka Tripathi<sup>2</sup>, Binal Raj<sup>3</sup>, Divyesh  
 299 Dadhanian<sup>4</sup>

- 300 1. Medical Student, School of Medicine and Surgery, Università  
 301 Cattolica Del Sacro Cuore, Rome, Italy  
 302 2. Department of Cardiology, Zydus Hospitals and Healthcare  
 303 Research Private Limited, Ahmedabad, India  
 304 3. Clinical Cardiology Associate, Department of Cardiology,  
 305 Ahmedabad, Zydus Hospitals and Healthcare Research  
 306 Private Limited, India.  
 307 4. Cardiovascular Imaging, Department of Interventional  
 308 Radiology, Zydus Hospitals and Healthcare Research Private  
 309 Limited, Ahmedabad, India

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**Summary of the case**

1	<b>PATIENT (GENDER, AGE)</b>	16y/o male and 34 y/o male
2	<b>FINAL DIAGNOSIS</b>	Ischemia with non-obstructive coronary arteries
3	<b>SYMPTOMS</b>	Paroxysmal palpitation and chest Discomfort
4	<b>MEDICATIONS</b>	Symptomatic treatment given
5	<b>CLINICAL PROCEDURE</b>	Echo, TMT, CAG, Stress thallium scan, CMR
6	<b>SPECIALTY</b>	Cardiology