A 33-year-old man with atypical chest pain

CLINICAL INTRODUCTION
A 33-year-old man with no history of coronary artery disease presented to the rapid access cardiology clinic with an episode of atypical anginal chest pain. He had a 15 pack-year history of smoking and a family history of myocardial infarction under the age of 55. Physical examination and exercise ECG testing were unremarkable. On assessment in the cardiology clinic, blood sampling was notable for an elevated high-sensitivity troponin I of 61 ng/L (99% upper reference level, 34 ng/L). A coronary CT angiogram was performed (figure 1).

QUESTION
Which of the following best explains this presentation?
A. Myocarditis  
B. Myocardial bridging  
C. Kawasaki’s disease  
D. Atherosclerotic plaque rupture  
E. Eosinophilic coronary monoarteritis

Figure 1  CT multiplanar reformatting of the left anterior descending artery with vessel wall analysis of a proximal (blue) and distal (red) segment.
The correct answer is atherosclerotic plaque rupture. Coronary CT angiography demonstrated a minimally stenotic (<25%) non-calciﬁed plaque in the proximal left anterior descending  artery (LAD). The vessel wall is thickened with a region of low CT attenuation surrounding the lumen representing a necrotic core of lipid-rich macrophages. The heterogeneous non-calciﬁed plaque is circumscribed by an outer rim of higher attenuation which is indicative of the napkin-ring sign. Additionally, the plaque has positively remodelled the vessel wall (remodelling index (vessel area in lesion: vessel area in reference segment) of 1.86, positive remodelling deﬁned as remodelling index ≥1.1). In patients with chest pain, the presence of these vulnerable plaque features increases the diagnostic likelihood of acute coronary syndrome.

The presence of coronary risk factors and abnormal coronary artery wall architecture reduces the likelihood of myocarditis. Myocardial bridging frequently occurs in the mid-LAD with plaque development upstream of the bridged segment; however, there is no myocardium surrounding the coronary artery to implicate a ‘tunnelled’ LAD. Kawasaki’s disease often presents in childhood affecting medium-sized arteries and results in luminal dilatation; however, the absence of intraluminal thrombus and coronary artery aneurysm makes this unlikely. Adventitial inﬁltration of eosinophils with spontaneous coronary artery dissection of the tunica media is characteristic of eosinophilic coronary monoarteritis and these features are not present in this instance.

The patient was prescribed aspirin and simvastatin. He presented 4 weeks later with an acute thrombotic occlusion of the proximal LAD that required emergency percutaneous coronary intervention (Figure 2). Identifying vulnerable plaques may help target effective treatment in individuals with active atherosclerosis to prevent rapid plaque progression and rupture.

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Figure 2 (A) Coronary CT angiography of left anterior descending artery (LAD) with non-calciﬁed plaque proximal to ﬁrst diagonal bifurcation. The arrow points to the culprit plaque in the proximal LAD on the Coronary CT angiogram. (B) Invasive coronary angiogram demonstrated complete occlusion of the proximal LAD. (C) Percutaneous coronary intervention at the site of plaque rupture restored coronary blood flow.

REFERENCES