Isolated left ventricular apical hypoplasia: a new congenital anomaly described with cardiac tomography

M Fernandez-Valls, M B Srichai, A E Stillman, R D White

METHODS

Patient population
Clinical description
The three patients were a 22 year old women of European descent (case 1), a 46 year old woman of Middle Eastern descent (case 2), and 26 year old man of European descent (case 3). Presenting symptoms were relatively mild and non-specific and included fatigue, shortness of breath, or chest discomfort. Personal and family medical histories were non-specific and included fatigue, shortness of breath, or chest discomfort. 

Prior diagnostic evaluation
ECGs showed normal sinus rate and rhythm, right axis deviation (110–130°), and low precordial voltages with poor R wave progression. Transthoracic echocardiograms showed variable degrees of LV systolic dysfunction (case 1: mild; case 2: mild to moderate; case 3: moderate to severe), always with bulging of the interventricular septum towards the right ventricle (RV), and variable degrees of mitral abnormality (case 1: moderate regurgitation; case 2: prolapse; case 3: none). Left heart catheterisation showed increased LV end diastolic pressures in one patient (case 1: 30–35 mm Hg) but normal indices in another (case 2); in both patients, no coronary artery abnormality was seen on selective angiography.

For all three patients, the primary diagnosis after the aforementioned evaluations and before cardiac tomography was idiopathic cardiomyopathy. All assessments of familial, metabolic, infiltrative, infectious, and inflammatory cardiomyopathy were negative.

Cardiac tomography techniques

Magnetic resonance imaging (MRI) is well established for assessing congenital abnormalities of the heart in adults. Recent advances in cardiac computed tomography (CT), especially multidetector CT (MDCT), have increased its ability to define congenitally abnormal structures.

Cardiac tomography, by MRI or MDCT, shows characteristic gross morphological and tissue changes of acquired or congenital diseases of the left ventricle (LV). We describe cardiac tomographic findings of an apparently new, presumably congenital, LV abnormality noted consistently in three patients. The findings suggested isolated LV apical hypoplasia.

Abbreviations: CT, computed tomography; LV, left ventricular; MDCT, multidetector computed tomography; MRI, magnetic resonance imaging; RV, right ventricle
RESULTS
Cardiac tomography findings
MRI findings
MRI showed the same morphological abnormalities in all three patients: a truncated appearance of the spherical (rather than normal conical) LV with bulging of the interventricular septum towards the RV; invagination of fatty material contiguous with epicardial fat into the area normally occupied by the LV apex; origin of a complex papillary muscle network in the flattened anteroapical (rather than anterolateral and posteromedial) region; and elongation of a normally functioning RV wrapped around the deficient LV apex (fig 1).
Visual assessment of the dynamic MRI data showed that the patients had abruptly limited late diastolic filling in a restrictive pattern and decreased contractility (fig 2). Additionally, visualisation of significant mitral regurgitation was noted in only one patient (case 1: moderate).

MDCT findings
A defective LV apex was visualised on MDCT. Better seen on MDCT than on MRI was the extent to which fatty material replaced the deficient apical myocardium, especially towards the distal interventricular septum (fig 1).

DISCUSSION
MRI and MDCT showed findings of this apparently new anomaly, isolated LV apical hypoplasia: (1) a truncated and spherical LV configuration with rightward bulging of the interventricular septum, associated with impaired late diastolic filling and systolic mechanics; (2) replacement of the LV apical myocardium, especially along the distal interventricular septum, with fatty material contiguous with epicardial fat; (3) origin of a complex papillary muscle network in the flattened anterior apex (but with significant regurgitation in only one of three patients); and (4) elongation of the RV wrapping around the deficient LV apex. These findings account for the clinical presentation and initial diagnostic evaluation suggesting cardiomyopathy.

Differential diagnoses: acquired lesions
Acquired processes were considered as potential causes of this cardiac condition. However, as our patients were dissimilar in age, sex, and ethnic group, causes other than congenital seemed unlikely. Additionally, the morphological changes differed from those of myocarditis or ischaemic heart disease. While Chagas’s disease often leads to LV involvement and may preferentially affect the apex, regions of fibrotic wall thinning commonly cause generalised dilatation with apical aneurysms. None of our patients had such changes or evidence of coronary abnormalities.

Differential diagnoses: congenital lesions
To our knowledge, no prior report matches this description of isolated LV apical hypoplasia. Acknowledging similarities to known conditions, we believe that this cardiac condition is unique.

It is clear that this condition is not hypoplastic left heart syndrome, which is characterised by underdevelopment of the aorta, the aortic valve, and the entire LV resulting from interrupted growth due to genetic influences or secondary to diminished flow to the developing LV. While hypoplastic left heart syndrome is associated with an atretic or stenotic
Endocardium, unlike the configuration found in our patients. Characterised by a diffusely dilated LV with a heavily trabeculated interventricular septum. These morphologies featuring an elongated (as opposed to truncated) RV wrapping around its truncated apex as the primitive ventricle is partitioned in the fifth week with formation of the muscular interventricular septum. Defective LV apical development in our patients may be attributable to relatively inadequate LV to RV dilatation during partitioning, leading to the spherical LV with an elongated RV wrapping around its truncated apex as the interventricular septum continues to develop.

**Cardiac development and potential mechanisms**

Heart development begins around the third week of fetal life with formation of a cardiac tube that develops areas of constriction and dilatation to form primitive chambers. The primitive ventricle is partitioned in the fifth week with formation of the muscular interventricular septum. Defective LV apical development in our patients may be attributable to relatively inadequate LV to RV dilatation during partitioning, leading to the spherical LV with an elongated RV wrapping around its truncated apex as the interventricular septum continues to develop.

**Clinical management**

The natural history of this condition is uncertain, as our patients are relatively young and functional. However, standard treatment of their heart failure has been helpful. While none of them has evidence of tachyarrhythmias, concern for malignant arrhythmias in isolated LV apical hypoplasia seems appropriate.

**Conclusion**

Isolated LV apical hypoplasia is a unique, presumably congenital, cardiac anomaly that is an important condition to recognise.

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**REFERENCES**


While fibrofatty myocardial replacement was seen in our patients, it was limited to the apex of the non-dilated dysmorphic LV, suggesting deficient apical development. In addition, in our patients dysfunction was predominantly diastolic, rather than systolic, and none of them had manifested any clinical signs or symptoms of tachyarrhythmias.

Isolated RV hypoplasia, a rare idiopathic condition characterised by RV underdevelopment with absence of its trabeculated apex, has RV features similar to those affecting the LV in our cases. Patients with this condition also have increased RV and RA pressures due to reduced compliance, not dissimilar to the diastolic dysfunction noted in our patients. However, their myocardial abnormalities are limited to the RV (as opposed to LV) and do not appear to share the fibrofatty morphology characterising our cases.

**Figure 2** Comprehensive evaluation of isolated LV apical hypoplasia by cardiac MRI. Anatomical evaluation with dark blood imaging (A) without and then (B) with fat saturation confirms the fatty nature of the material (large arrows) contiguous with epicardial fat (small arrow) and extending into the truncated LV apex. Dynamic evaluation with (C, D) dynamic bright blood and (E, F) tissue tagged imaging shows the spherical shape of the restrictive LV in late diastole and lack of systolic myocardial deformation due to impaired contractility, especially in the interventricular septum.

mitral valve 95% of the time, our patients had mature valves and malformation was limited to the LV apex.

Non-compaction of the LV results from arrested endomyocardial morphogenesis leading to dysfunction. It is characterised by a diffusely dilated LV with a heavily trabeculated endocardium, unlike the configuration found in our patients.

Congenital LV aneurysm, an idiopathic anomaly of the endocardium and myocardium, and LV diverticulum, part of an idiopathic syndrome with multiple defects, also appear to be different entities. As well as resulting in different LV morphologies featuring an elongated (as opposed to truncated) LV and involvement of the papillary muscles and surrounding myocardium (as opposed to isolated involvement of apex), these conditions usually lead to severe, usually fatal, clinical presentations early in life due to associated intracardiac and extracardiac defects (as opposed to relatively healthy adult presentations). Congenital LV dysplasia caused by abnormal endomyocardial morphogenesis in association with RV dysplasia is a generalised myocardial disease with predominantly RV involvement, which none of our patients had. In previously described cases of isolated LV dysplasia, in one case thinning of the posterior LV myocardial wall was notable with transmural fibrofatty replacement and in the other three there was extensive circumferential fibrofatty infiltration of predominantly LV myocardium causing cardiomegaly, systolic heart failure, and tachycardia. While fibrofatty myocardial replacement was seen in our patients, it was limited to the apex of the non-dilated dysmorphic LV, suggesting deficient apical development. In addition, in our patients dysfunction was predominantly diastolic, rather than systolic, and none of them had manifested any clinical signs or symptoms of tachyarrhythmias.

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**REFERENCES**


Left ventricular pseudoaneurysm following myocardial infarction

An asymptomatic 61 year old man with a history of coronary artery disease, including coronary artery bypass surgery, presented for routine examination after a lateral myocardial infarction treated by thrombolysis five months previously. Physical examination was unremarkable. Echocardiography revealed a dyskinetic cavity connected to the posterolateral wall which communicated freely with the left ventricle. Magnetic resonance imaging (below left) confirmed the diagnosis of a false aneurysm (A) of the left ventricle (LV) by demonstrating a large perforation of the free posterolateral wall (arrows). Invasive coronary angiography revealed a patent left internal mammary artery graft to the left anterior descending coronary artery. A venous bypass to the left circumflex coronary artery demonstrated a severe stenosis at the distal anastomosis. Open heart surgery was performed. The aneurysmal sac was excised and the defect (below right, arrows) was closed with a pericardial patch. The postsurgical period was uneventful.

Rupture of the left ventricular free wall is a catastrophic complication of myocardial infarction, occurring in approximately 4% of patients with infarcts and about 23% of those suffering fatal infarcts. It is four to five times more common than septal rupture. Rarely, free wall rupture is contained by overlying, adherent pericardium, producing what has been termed a pseudoaneurysm or false aneurysm of the left ventricle. These pseudoaneurysms are often detected incidentally by echocardiography or other imaging modalities. Because of their propensity to rupture, emergency surgical intervention is recommended. The patient survived because the pericardium adhered to the epicardium as a consequence of bypass surgery six years earlier.

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