Tearing while bowling

Pairoj Chattranukulchai,1 Kittichai Luengtaviboon,2 Monravee Tumkosit3

DESCRIPTION

A 38-year-old man with asymptomatic chest wall deformity presented with acute, severe retrosternal pain experienced while he was bowling. The pain radiated to the back and down to abdomen in a tearing fashion. Physical examination revealed tall stature, arachnodactyly, marked funnel chest (figure 1A) and severe thoracic scoliosis as demonstrated

![Figure 1](A) Physical examination reveals marked funnel chest. (B) Chest film shows severe thoracic scoliosis with leftward shift of the heart.

![Figure 2](A) Contrast-enhanced axial CT demonstrate an annuloaortic ectasia of aortic root (red asterix) with intimal flap separates the true from false lumen at descending aorta (black asterisk). Severe pectus excavatum with very short distance between sternum and vertebrae results in complete encompassing of the heart in the left hemithorax (arrows). (B) Sagittal image also shows the severely depressed sternum and the extent of floating dissection flap along descending aorta, consistent with type B aortic dissection.

1Division of Cardiology, Department of Medicine, Faculty of Medicine, Chulalongkorn University, Cardiac Center, King Chulalongkorn Memorial Hospital, Bangkok, Thailand
2Division of Cardiothoracic Surgery, Department of Surgery, Faculty of Medicine, Chulalongkorn University, Bangkok, Thailand
3Department of Radiology, Faculty of Medicine, Chulalongkorn University, Bangkok, Thailand

Correspondence to
Dr Pairoj Chattranukulchai,
pairoj.md@gmail.com

To cite: Chattranukulchai P, Luengtaviboon K, Tumkosit M. BMJ Case Rep Published online: [please include Day Month Year] doi:10.1136/bcr-2013-010389
by chest film (figure 1B). He was diaphoretic with blood pressure in both arms were 120/80 mm Hg where remarkably diminished to 80/60 mm Hg in both legs. Faint diastolic blowing murmur was heard along the left parasternal border; these factors prompted the suspicion of aortic dissection.

An echocardiography revealed annuloaortic ectasia with aortic root dilation to 52 mm associated with mild aortic regurgitation. The dilated descending thoracic aorta with intimal flap was also noted.

A contrast-enhanced axial CT revealed severe pectus excavatum with a very short distance between the sternum and the vertebrae resulting in complete encompassing of the heart in the left hemithorax (figure 2A, arrows). Annuloaortic ectasia (figure 2A, red asterix) with type B aortic dissection were confirmed. The intimal flap separated true from false lumen (figure 2B, black asterisk) starting from the proximal descending thoracic aorta down to the aortic bifurcation. The sagittal image also demonstrated the severely depressed sternum and the extent of the floating dissection flap (figure 2B).

The patient was diagnosed as having Marfan syndrome—the intrinsic aortic wall weakness predisposing to aneurysm formation and dissection. Late diagnosis has a negative influence on the outcome. Early detection includes interval monitoring of aortic enlargement by regular imaging of the aorta along with preventive therapies consist of oral β-blocker along and prophylactic aortic repair when indicated could prevent this catastrophic event. The patient underwent open aortic repair and his postoperative follow-up was uneventful.

**Learning points**

- Marfan syndrome is an inherited disorder of connective tissue.
- Clinical manifestations comprise classic ocular, musculoskeletal and cardiovascular abnormalities which include aneurysm formation and dissection of aorta.
- Early detection including regular imaging of the aorta and timely surgical repair when indicated are crucial to prevent catastrophic aortic dissection.

**Contributors** PC was involved in writing the manuscript. KL and MT reviewed and edited the manuscript.

**Competing interests** None.

**Patient consent** Obtained.

**Provenance and peer review** Not commissioned; externally peer reviewed.

**REFERENCES**
