Calcinosis Cutis Associated with Severe Atherosclerosis and Severe Calcific Aortic Stenosis: Case Report

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Abstract

Calcinosis cutis is a condition of accumulation of calcium salts within the dermis. We report a case in which Calcinosis cutis is combined with severe cardiac involvement. Our patient suffered from Calcinosis cutis for about 40 years. Except dermal symptoms she presents with severe diffuse cardiovascular calcification. These involve calcification of large and middle arteries (such as femoral arteries, superior mesenteric artery), diffuse severe coronary atherosclerosis, pronounced calcification and phlebitis of the leg veins and calcification of cardiac valves including severe aortic stenosis and calcified mitral valve. Calcinosis cutis is a condition of accumulation of calcium salts within the dermis. We are presenting a case of Calcinosis cutis, with special clinical presentations, with normal serum calcium and phosphorus levels.

The patient has severe calcific aortic stenosis, calcified mitral valve, diffuse severe coronary atherosclerosis, and other large vessels such as femoral artery and artery mesenterica superior. And pronounced calcification and phlebitis the leg veins.

Case Report

An 86-year-old woman presented to the emergency department with recurrent syncope. Her medical history was significant for aortic stenosis.

Calcinosis cutis with ulceration on both lower limbs for about 40 years. Multiple surgical therapies including excisions of ulcerated areas on both legs were performed in the last years. In addition she suffered from a sclerosis of the deep leg veins with severe chronic venous insufficiency (CVI). She had chronic renal disease with normal serum calcium and phosphorus levels, arterial hypertension and hypercholesterolemia.

Moreover, it was sonographically a severe insufficiency of the Vena saphena magna and parva on both sides, and massive calcification of the veins along the two lower legs. This year the patient once suffered from bloody secretions of the left ear [1].

The physical examination was notable for white yellowish, hard, painless lump and small fibrinous ulcerations surrounded by erythema left pretibial (Figure 1). And a systolic ejection murmur, with a maximum in the 2nd ICR right parasternal with radiation in the carotids.

Findings didn’t show any evidence of an acute coronary syndrome in the emergency room. The echocardiography revealed left ventricular hypertrophy with normal systolic LV function (ejection fraction 55%) and calcification of aortic and mitral valves. Doppler sonography revealed mean gradient of 67 mmHg on the aortic valve corresponding to sever aortic stenosis (Figure 2). The gradient on the mitral valve was normal.

Figure 1: Small fibrinous ulcerations surrounded by erythema left pretibial.

Doppler sonography showed calcified intima of both femoral arteries. The X-ray image of the left lower leg showed the calcification of the skin layers.

Due to symptomatic aortic valve stenosis indication for aortic valve replacement was established. Interdisciplinary heart-team made the decision for treatment by transcatheter aortic valve implantation (TAVI) because of high perioperative risk (EuroScore II 6.6%, and STS-Score 7.9%). The computer tomography angiography was carried out in preparation for TAVI (Figure 3).
The patient was successful treated by transfemoral aortic valve implantation with Edward-SAPIEN 26 mm in local anesthesia without major complications. Postoperative echocardiography showed correct position of aortic prosthesis without relevant paravalvular leakage and mean gradient of 20 mmHg (Figure 3). Further hospital stay was uneventful. The patient has recovered quickly after TAVI and could be discharged after five days.

Repeated surgical excision was planned electively after discharge.

Discussion

Calcinosis cutis is characterized by the deposition of insoluble calcium salts in the skin and subcutaneous tissue. The syndrome is divided into five subtypes: dystrophic calcification, metastatic calcification, idiopathic calcification, iatrogenic calcification and calciphylaxis [2].

In all cases of Calcinosis cutis, insoluble compounds of calcium are deposited within the skin due to local and/or systemic factors. These calcium salts consist primarily of hydroxyapatite crystals or amorphous calcium phosphate [3]. The pathogenesis of Calcinosis cutis is still not completely understood and a variety of factors allow for different clinical scenarios to occur.

It should be discussed from which subtype the patient is affected and the pathophysiology of cardiac involvement with vascular calcification.

The symptoms and the underlying condition may indicate partially a calciphylaxis. So the ulcerations in a calciphylaxis are painful. Calciphylaxis presents with small vessel calcification mainly affecting blood vessels of the dermis or subcutaneous fat.

Disturbances in calcium and phosphate metabolism and hyperparathyroidism can be observed [4].

Our patient doesn't have painful ulcers, and has a normal serum calcium and phosphorus levels. Therefore calciphylaxis is unlikely in this case.

The patient has most likely a Calcinosis dystrophic. This subtype shows calcification of vessels, thrombosis, phlebitis, painless ulcerations and a normal serum calcium and phosphorus levels.

Recently, great interest has focused on understanding how atherosclerotic pathology is related to a common plaque constituent: calcium mineral deposits. Important studies in the last decade have now spawned a model wherein calcification in atherosclerotic plaque is viewed as an active complex and therefore presumably regulated process that exhibits intriguing similarities to formation, or remodelling from other tissue like skin and bones [5]. There is an association between anti-centromere antibodies and Calcinosis and atherosclerosis [6,7]. The pathology may be propagated by micro vascular insufficiency, inflammation and local tissue damage in the vessels [8].

A cardiac involvement under Calcinosis is not sufficiently described in the literature. We present a rare case with pronounced cardiovascular involvement including diffuse calcification of small and medium-sized arteries, deep leg veins, moderate mitral valve and severe aortic stenosis. It is common to have aortic stenosis and a mild calcified coronary artery in a 86 year old woman, but due to this pronounced combination with dystrophic calcification should be in this dimension considered. Due to extent of valvular calcification it is
likely that Calcinosis cutis was involved in pathogenesis of the cardiac disease in our patient, however exact mechanism is still not understood. A careful clinical examination and eventually echocardiography should be considered in patients with Calcinosis cutis to diagnose possible cardiovascular involvement.

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References