Sclerochoroidal calcification is an uncommon condition. Metabolic evaluation and clinical examination are important to exclude associated systemic conditions such as the Bartter and Gitelman syndromes. It has been suggested that the lesions seen in sclerochoroidal calcification are calcium pyrophosphate dihydrate crystals. This report describes the first documented case in the UK of sclerochoroidal calcification associated with Gitelman syndrome and calcium pyrophosphate dihydrate deposition.

CASE REPORT

After an annual optometric examination a 49 year old woman with no ocular symptoms was referred with bilateral fundal abnormalities. There was no previous documentation of the fundal lesions. An initial systemic enquiry was unremarkable and she took no ocular or systemic medication. Best corrected visual acuity was 6/6 (right eye) and 6/9 (left eye). Intraocular pressure was 16 mm Hg bilaterally. Anterior segment examination was unremarkable. Fundoscopy showed numerous discrete yellow/white, slightly elevated lesions of the subretinal pigment epithelium (RPE) located around the superior temporal arcade bilaterally (fig 1). Some depigmentation of the RPE overlying the lesions was also noted. Fluorescein angiography showed non-specific changes with RPE atrophy. B scan ultrasound (fig 2) showed that the lesions were highly echogenic with acoustic shadowing.

Blood results revealed hypokalaemia (3.1 mmol/litre; normal range, 3.5–5.0), hypomagnesaemia (0.30 mmol/litre; normal range, 0.7–1.0), and raised globulin (39 g/litre; normal range, 18–35). Serum electrolytes—sodium, chloride, calcium, and phosphate—were normal, as was her parathyroid hormone value. A 24 hour urine analysis showed hyperkaliuria (102 mmol/24 hours; normal range, 25–100), hypocaliuria (1.6 mmol/24 hours; normal range, 2.5–7.5), natriuresis (227 mmol/24 hours; normal range, 50–200), and normal urine magnesium excretion (4.18 mmol/24 hours; normal range, 3.0–5.0). Blood pH was 7.529 confirming a metabolic alkalosis.

On further direct questioning, the patient described symptoms consistent with pseudogout and plain radiographs revealed extensive CPPD deposition in the fibrocartilage of the knee (fig 3) and in the hyaline cartilages of the shoulder, ankle, and elbow joints. A diagnosis of sclerochoroidal calcification secondary to Gitelman syndrome was made.

DISCUSSION

Idiopathic sclerochoroidal calcification requires no treatment, but screening tests for calcium phosphorous metabolism should be performed to exclude an underlying systemic disorder. Biochemically, Gitelman syndrome and Bartter-like syndromes are characterised by hypokalaemia associated metabolic alkalosis and normotension. Low serum magnesium is occasionally found in patients with Bartter syndrome, so that Gitelman syndrome, in which all patients have hypomagnesaemia, needs to be excluded. Differentiation is achieved by measuring urine calcium. In Gitelman syndrome, hypocalciuria is noted usually in the presence of a normal total serum calcium and ionised calcium, whereas in Bartter syndrome normal or high urine calcium excretion is seen. Our patient had a biochemical profile consistent with a diagnosis of Gitelman syndrome; namely, metabolic hypokalaemia alkalosis in the presence of hypomagnesaemia and hypocalciuria. This renal salt wasting disorder is an autosomal recessive condition caused by a defect in the thiazide
sensitive Na–Cl cotransporter in the distal convoluted tubule. It is often diagnosed in asymptomatic adults who present with unexplained hypokalaemia. However, these patients can be at risk of cardiac arrhythmias and other systemic symptoms as a result of the metabolic disturbance. On suspending sclerochoroidal calcification, it is important to test for associated conditions, such as the Bartter and Gitelman syndromes, because both conditions can now be treated, the latter with oral magnesium pyrrolidone carboxylate.

"On suspecting sclerochoroidal calcification, it is important to test for associated conditions, such as the Bartter and Gitelman syndromes, because both conditions can now be treated."

Chondrocalcinosis has been suggested as a feature of Gitelman syndrome. Excess extracellular inorganic pyrophosphate has been recognised as a probable cause of CPPD crystal deposition disease. In patients with hypophosphatasia, a deficiency of tissue non-specific alkaline phosphatase, crystal deposition disease. In patients with hypophosphatasia, a deficiency of tissue non-specific alkaline phosphatase allows an increase in extracellular inorganic pyrophosphate and thus CPPD disease. Magnesium is a cofactor for alkaline phosphatase and hypomagnesaemia associated with Gitelman syndrome may thus theoretically reduce the activity of alkaline phosphatase and predispose to CPPD disease. It has been suggested that the lesions seen in sclerochoroidal calcification are CPPD crystals, and recently a familial case of sclerochoroidal calcification and chondrocalcinosis has been reported. Our case further suggests that the sclerochoroidal lesions seen in Gitelman syndrome may be crystal deposits of CPPD and that the possible underlying molecular mechanism is secondary to hypomagnesaemia.

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Sclerochoroidal calcification associated with Gitelman syndrome and calcium pyrophosphate dihydrate deposition
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