Diabetic muscle infarction

Clinical record

A 55-year-old woman with type 2 diabetes of 8 years’ duration experienced, over 2–3 days, the onset of pain, tenderness and swelling of the medial aspect of her right thigh.

She had recently commenced insulin therapy and was displaying good glycaemic control (HbA1c, level of 6.4%). She had diabetic complications of autonomic and peripheral neuropathy, but no retinopathy. Other medical problems included chronic renal impairment, hypertension, polyarticular gout and hydralazine-induced lupus. A renal biopsy had not been performed, but her renal insufficiency was believed to be a result of diabetic nephropathy and hypertension. She was taking twice-daily mixed insulin (16 units in the morning and 10 units at night), felodipine (10 mg/day), paroxetine (10 mg/day), allopurinol (75 mg/day) and prednisolone (7.5 mg/day).

Her serum creatinine level had peaked at 0.31 mmol/L, but stabilised at 0.21 mmol/L after cessation of an angiotensin-converting enzyme inhibitor. A 24-hour urine collection showed a creatinine clearance rate of 0.12 mL/s (normal range [NR], 1.5–2.5 mL/s) and a protein excretion rate of 4.6 g/day. Renal duplex ultrasound showed that her kidney size was well preserved, but there was a suggestion of renal artery stenosis on the right side.

There was no history of recent injury or injection to her thigh. She had not experienced any rigors and was afebrile. The area of the localised, tender swelling on the medial aspect of her right thigh was not erythematous and no local lymphadenopathy was noted. Apart from the thigh swelling, there was generalised wasting and weakness of the lower limbs, loss of ankle reflexes, and loss of sensation in a stocking distribution, consistent with a diagnosis of peripheral neuropathy. Examination of her left foot revealed two small gangrenous areas. Foot pulses were present and the gangrenous areas were thought to be caused by inappropriate footwear.

Investigations showed an elevated white cell count of $16 \times 10^9$/L (NR, $4.0–11.0 \times 10^9$/L), a platelet count of $432 \times 10^9$/L (NR, $150–400 \times 10^9$/L), an erythrocyte sedimentation rate of 110 mm/h (NR, 7–18 mm/h) and a C-reactive protein level of 119.3 mg/L (NR, 1.6–8.7 mg/L). Creatine kinase (174 U/L; NR, < 215 U/L) and lactate (0.6 mmol/L; NR, 0.5–2.0 mmol/L) levels were normal. Separate to the swelling on the medial aspect of the thigh, a presumed thrombus was palpable in the lateral accessory long saphenous vein. A subsequent ultrasound detected this thrombus extending from the mid thigh to the lateral aspect of the knee, but no deep venous thrombosis was found. She was treated with cephalexin and aspirin.

Over the next two weeks, the thigh swelling evolved into a tender, palpable mass measuring 6 x 20 cm. A computed tomography (CT) scan revealed marked swelling of the entire adductor muscle group, but no discrete mass (Figure 1). In the absence of a clear diagnosis, an exploratory operation of her right thigh was performed. This showed no haemorrhage or abscesses, but evidence of extensive oedema and necrosis of the adductor muscles. Samples taken for microbiological analysis were sterile. Histological examination of a biopsy specimen showed necrotic muscle, an inflammatory cell infiltrate (B) and a blood vessel containing a thrombus (C). No evidence of vasculitis was seen.

The CT scan did not reveal a discrete mass, but extensive oedema of one muscle group and sparing of an adjacent muscle group. Histologically, there was evidence of skeletal muscle fibre necrosis, with a variable amount of muscle regeneration and fibrosis. These are the typical features of diabetic muscle infarction. Reports of spontaneous muscle infarction appear to be virtually confined to patients with diabetes.

Spontaneous muscle infarction is a rare diabetic complication. There have been fewer than 100 patients reported since 1965. However, it is becoming more frequently recognised; almost half of the cases have been reported since 1999. It has a predilection for the quadriceps (62%), hip adductors (13%), hamstrings (8%) and hip flexor (2%) muscles. Rarely, the calf and anterior tibial muscles are...
involved. The pathogenesis of diabetic muscle infarction is still unclear, but a diffuse microangiopathic process, possibly associated with hypoxia–reperfusion injury, has been implicated as a cause.\textsuperscript{1,10,12}

Magnetic resonance imaging (MRI) is the preferred diagnostic test, revealing swollen and oedematous muscles (ie, increased signal intensity of T2-weighted images).\textsuperscript{4,7} Abnormal MRI findings have been reported in all patients with diabetic muscle infarction. Although, in retrospect, our patient’s CT findings were consistent with muscle infarction, a CT scan is considered a less sensitive test, as only 83% of patients with muscle infarction have abnormal CT findings.\textsuperscript{1} An MRI scan was not performed in our patient, as diabetic muscle infarction was not initially considered in the differential diagnosis. Consensus opinion suggests that muscle biopsy is not necessary in a patient presenting with the typical clinical features of diabetic muscle infarction — without fever, erythema, or elevated white cell count — if the MRI findings are appropriate.\textsuperscript{1} 

Management consists of the avoidance of weightbearing, and simple analgesia. After resolution of the acute phase, physical therapy and rehabilitation are useful. As with our patient, total recovery over 4–6 weeks can be expected. In about 50% of patients recurrences occur, but not necessarily in the same muscle group.

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