Rapid development of a tophus following ipsilateral hemiparesis

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Abstract
An 85 year old man with a longstanding history of non-tophaceous gout developed a single large tophus, unassociated with inflammation, on his paretic leg over a six week period following an acute hemiplegia. The rapidity of tophus formation, its localisation to the paretic limb, and the apparent blunting of the acute inflammatory response represent a previously unreported interaction between gout and neurological deficit.

Considerable interest exists in the interaction between neurological deficits and the clinical expression of rheumatic diseases. The sparing effect of upper or lower neurological deficit on the subsequent development of arthropathy, including gout, has been well described. Conversely, a few cases have been described in which exacerbation of established joint disease has followed hemiparesis. We report an apparently unique patient with a history of recurrent acute gout, in whom the development of a hemiparesis was followed by the rapid appearance of a non-inflammatory tophus on the heel of his paretic foot.

Case report
A previously fit 85 year old right handed man was admitted with an acute left hemiplegia. Since the age of 50 he had had recurrent acute attacks of gout limited to both feet (including classic podagra), treated with short intermittent courses of non-steroidal anti-inflammatory drugs and allopurinol. He had not taken such drugs for many months. There were no associated risk factors for gout (obesity, excess alcohol, chronic diuretic treatment, family history) and he had never developed tophi. Examination confirmed marked left sided weakness affecting the arm more than the leg, mild sensory signs but no visual field deficit. A computed tomography brain scan showed right frontoparietal infarction.

Over the subsequent six weeks his motor function improved, though he still required the assistance of one person for most activities. At this time he complained of a swelling over his left heel which prevented him from wearing his normal footwear. The swelling, at the level of the pre-Achilles bursa, was 2×3 cm, firm, and non-tender. The overlying skin was healthy and intact with no signs of local inflammation. No other swellings were apparent elsewhere. A lateral radiograph (fig 1) showed soft tissue swelling superficial to the Achilles tendon insertion with an underlying small focus of calcification. Aspiration of the swelling confirmed urate crystals in a typical tophus 'sheet' configuration (compensated polarised light microscopy). After aspiration macroscopically white crystal deposits exuded through the needle track (fig 2), without provoking an inflammatory response.

Further investigations showed: urea 11.7 mmol/l (normal range 1-6.5); creatinine 167 μmol/l (60-120); uric acid 406 μmol/l (100-400); calcium and fasting lipoproteins within normal limits. Radiographs of feet, knees, pelvis, and hands showed only minor changes of osteoarthritis (knees, first metatarsophalangeal joints) and no chondrocalcinosis. Treatment was established with allopurinol 100 mg daily. Because the

Figure 1 Lateral left heel radiograph showing soft tissue swelling and central area of calcification.
Rapid development following unidentified trauma, vascular alteration to our calcific focus effects Discussion was weeks after his developing tophi. The possibility of tophi strongly increased in tissue damage, and lower limbs lesions. The association with symptomatic chronic gout. In our patient the speed of formation of his first ever tophus was rapid. The absence of associated inflammation is also remarkable, particularly as newly forming (compared with older) urate crystals might be expected to expose more ‘naked’, non-protein coated surfaces for interaction with cell surfaces and inflammatory mediators. The morphology of the urate crystals seemed typical of those seen in tophi, though it is possible that such rapidly forming crystals might present less active surfaces with respect to surface charge and ‘roughness’. Diminution of the inflammatory reaction consequent upon altered neurovascular responses would seem a more plausible explanation, possibly mediated through reduction in neurotransmitters.

Mechanisms to explain exacerbation of arthropathy (rheumatoid arthritis, osteoarthritides) following hemiparesis favour neurovascular alteration of blood flow or altered neurotrophic effects on inflammation. The converse sparing effect of upper and lower motor neurone lesions on development of locomotor disease (including gout) has more commonly been attributed to disuse with reduction of damaging mechanical factors and intra-articular pressure. Acute self limiting arthropathy has been seen in hemiplegic limbs of patients with no prior history of arthritis shortly after a cerebrovascular accident, although reaction to underlying infection has been incriminated, the cause remains speculative.

In the present case it is possible that minor tissue damage or unrecognised trauma, consequent upon weakness and sensory deficit, promoted tophus formation in a predisposed patient. Alternatively, altered neurovascular response in a paretic limb following stroke may lead to increased tissue urate concentrations and possibly predispose to the development of tophi. Although there was no overt pre-Achilles trauma, ulceration or bursitis, the calcific focus on the radiograph might support local tissue injury and indicate that a small, unidentified (mixed) crystal deposit had been present at this site for some time. Whether a crystal mass originated before or after the stroke, however, the hemiparesis seems to have encouraged accelerated crystal deposition and tophus development.

The only comparable case is that reported by Cosgrave and Lewkonia of a patient with posttraumatic ‘frozen shoulder’ who developed tophi in the ipsilateral hand six months later. Although there was no neurological deficit, disuse or altered regional blood flow were again both proposed as mechanisms of causation.

Tophus formation is normally regarded as a slow process. Although in vitro formation of monosodium urate monohydrate crystals can be rapid, in vivo formation is estimated to take many months, and tophi usually occur in association with symptomatic chronic gout. In our patient the speed of formation of his first ever tophus was rapid. The absence of associated inflammation is also remarkable, particularly as newly forming (compared with older) urate crystals might be expected to expose more ‘naked’, non-protein coated surfaces for interaction with cell surfaces and inflammatory mediators. The morphology of the urate crystals seemed typical of those seen in tophi, though it is possible that such rapidly forming crystals might present less active surfaces with respect to surface charge and ‘roughness’. Diminution of the inflammatory reaction consequent upon altered neurovascular responses would seem a more plausible explanation, possibly mediated through reduction in neurotransmitters.