A 70 year old man, a non-smoker, presented with a painful swelling over the right sternoclavicular joint and low grade continuous fever for a period of two months. The patient had been empirically started on antituberculous therapy on the suspicion of tuberculous osteomyelitis, but his fever and swelling persisted.

On clinical examination, a hard and tender swelling was present over the right sternoclavicular joint. The swelling was fixed to the underlying bone, but not to the overlying skin. The patient did not have any skin lesions. Examination of his respiratory system revealed no abnormality. Examination of other systems was unremarkable.

Blood investigations showed a haemoglobin concentration of 160 g/l. The total white blood cell count was $10.5 \times 10^9$ with 72% polymorphonuclear cells, 26% lymphocytes, and 2% eosinophils. ELISA testing for HIV infection was negative. Other haematological and biochemical parameters were within normal limits.

Fine needle aspiration cytology of the swelling showed a few red blood cells, polymorphonuclear cells, lymphocytes, and a few epithelioid cells in a necrotic background. No granulomas or giant cells were seen.

Computed tomography (plain and contrast) of the thorax revealed a right sternoclavicular and first costoclavicular joints arthropathy with subchondral sclerosis and abnormal periarticular soft tissue mass. A technetium-99 bone scan was done (shown in fig 1) which revealed the diagnosis.

Questions
(1) What does the bone scan show?
(2) What is the diagnosis and name the associated syndrome.
### Answers

**QUESTION 1**
The bone scintigraphy shows an increased radiotracer uptake in the region of the right sternoclavicular joint and the manubrium sterni ("bullhead" sign). Increased uptake was also seen in the region of the left knee joint, the left ankle joint, D9 thoracic vertebrae, and the posterior end of the left eighth rib.

**QUESTION 2**
The diagnosis is sternocostoclavicular hyperostosis (SCCH). The associated syndromes are called SAPHO syndrome (an acronym for synovitis, acne, pustulosis, hyperostosis, and osteitis), CRMO (chronic recurrent multifocal osteomyelitis), ACW syndrome (anterior chest wall), or PAO (pustulotic arthropoesteitis).

### Discussion

SCCH is observed mainly in young and middle aged adults, and is rarely seen above 60 years of age. Bone and joint involvement are the commonest findings. Patients present with pain and swelling of gradual onset, most commonly involving the medial end of the clavicle and the manubrium sterni. Involvement of only one clavicle and the adjacent part of the manubrium sterni are also a frequent event. The ribs are similarly involved, with changes occurring in the anterior costochondral junction and/or in the posterior costal arch, leading to limitation of the thoracic cage mobility. The swelling might also present as a solitary neck mass or as thoracic sinus formation. Flat bones like the ilium, the mandible, long bones, or spine could also be involved in the disease process. The involved bone shows sclerotic changes with periosteal, articular, or periarthritis inflammation.

The skin involvement in SAPHO and PAO is usually in the form of palmpoplantar pustulosis, palmpoplantar psoriasis, or severe acne known as acne conglobata or acne fulminans. Though skin involvement is a common feature, the absence of skin lesions at the time of presentation is only apparent because the skin lesions may have been transitory, or may develop decades after the bony manifestation. The patient may have other manifestations like thoracic outlet syndrome, thrombosis of the subclavian vein, or superior vena cava syndrome.

Diagnosis can be made by radiography. Bone scintigraphy using radiotracer material like technetium-99 is the imaging modality of choice. It reveals hot spots in the areas of increased uptake. “Bullhead” sign refers to increased uptake by the medial end of the clavicles and manubrium sterni, corresponding to a bull’s head. This sign is a typical and a highly specific manifestation of SCCH syndrome and helps to confer the diagnosis. Also a bone scan is able to detect early bone involvement, which would not yet be seen radiographically. Ultrasonography, computed tomography, and magnetic resonance imaging contribute little to the identification and location of the lesion.

### Learning points
- Sternocostoclavicular hyperostosis (SCCH) is a benign condition that involves bones and joints.
- SCCH is seen in the young and middle aged, rarely in old age.
- Associated syndromes include SAPHO, PAO with skin involvement.
- Diagnosis is by bone scan with classical “bullhead sign”.
- Treatment consists of anti-inflammatory drugs, rarely in severe cases with steroids.
- SCCH should be considered in the differential diagnosis of bacterial osteomyelitis, Paget’s disease, Ewing’s sarcoma, and infectious spondylodiscitis.

### Final diagnosis

Sternocostoclavicular hyperostosis.

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Painful chest wall swelling

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