POTT’S PUFFY TUMOUR — REPORT OF A GROTESQUE CASE

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Sir Percival Pott in the 18th century described osteomyelitis of the frontal skull bone with associated subperiosteal abscess collection, which became known as Pott’s Puffy Tumour. The incidence of the disease has been reported to have progressively dropped since the advent of antibiotics in the 20th century. Most of the frontal skull infections are thought to result from contiguous spread from the paranasal sinuses, though post traumatic cases were originally described, and staphylococcus is the commonest isolated pathogen. Osteomyelitis of other skull bones has been described with associated abscesses as well, and in some cases, there is an accompanying intracranial empyema. Cranial imaging with Computerized Tomography is the gold standard in evaluation, and abscess drainage with craniectomy, followed by weeks of broad spectrum antibiotic therapy is the mode of treatment. We report a case of grotesque looking Pott’s Puffy Tumour in a 16-year old male student who was admitted into our service and was treated operatively and discharged home without neurological deficits. The size of the lesion was as worrisome as the duration of ill health in this era of antibiotics when the availability of medical services in Nigeria is presumed to have improved significantly.

Key words: • Antibiotics • craniectomy • drainage • incision • osteomyelitis • trauma

Pott’s puffy tumour as originally described by the British surgeon, Sir Percival Pott [1714-1788], was a sub-periosteal abscess collection following an osteomyelitis of the frontal bone as a result of trauma to that bone [1,2]. It is equally known to complicate infections of the paranasal sinuses in majority of cases [3,4,5,6,7,8,9]. But osteomyelitis could also affect the other skull bones, and not the frontal bone in isolation. Also, beyond the extracranial abscess collection, intracranial abscesses, orbital sepsis and periorbital oedema have been reported to complicate this disease [10,11,12,13,3]. Staphylococci have been reported as the most common infective agents, but anaerobes are increasingly being cultured, especially, in the paediatric cases [14,12]. With the advent of antibiotics, Pott’s puffy tumour became increasingly rare as less number of cases became reported, but it is not yet extinct [15,16,17,18,11,12,13]. Imaging studies readily diagnose the lesion in the affected cases, and also reveal contiguous infections of the
paranasal sinuses and the intracranial cavity, and with computed tomography, the full extent of bone infection is better defined. Abscess drainage and craniectomy followed by 4 – 6 weeks of antibiotic therapy usually cures the disease, and in most cases without neurological deficits.

We report the case of a 16-year old male student who was admitted into our service with a grotesque looking Pott’s Puffy Tumour, without any known predisposing factor, and was treated operatively and discharged home without neurological deficits. The size of the lesion was as worrisome as the duration of ill health, in this era of antibiotics when the availability of medical services in Nigeria is presumed to have improved significantly.

Case Report

I.O. is a 16-year male right handed secondary school student who was referred to our service on 5th February, 2009 with complaints of recurrent fever, scalp pains and scalp swellings of 6 months’ duration.

There were associated headaches, neck pains, bilateral eye swellings and purulent eye discharge. The swellings had appeared serially at 2-monthly intervals in the parietal region and were treated with herbs and orthodox drugs, discharging pus but with persistence of scalp pains. The present episode started 1 week earlier with a frontal scalp swelling that rapidly spread over most of the scalp. There was no history of trauma to the head, or URTI, prior to onset and no associated alteration in sensorium, sphincteric dysfunction, hemi-corporeal weakness, seizures or vomiting.

Clinically, he was a young male, fully conscious but with a grotesque C-shaped cystic scalp collection involving the left frontal, left parietal and right parietal regions in coalescence. There was intense bi-periorbital oedema and mechanical ptosis with underlying chemosis, and moderate nuchal rigidity, but equivocal Kernig’s sign. There were no cranial nerve, gross motor, sensory or autonomic deficits, and no lateralizing signs.

We made a clinical diagnosis of Pott’s Puffy Tumour and requested for laboratory workup and skull x-rays because our Centre did not have Computed Tomographic facilities and the closest Centre with the facility was about 100 km away. The skull x-rays showed osteomyelitis of the left frontal and left parietal bones but no paranasal sinus lesions. The HIV I&II screening was non-reactive, genotype was ‘AS’ and total white cell count was 18.5 x 10^9/L with 60% neutrophilia.

We subsequently did a preliminary incision and drainage within six hours of admission to evacuate the abscess and relieve the pressure symptoms on the head and eyes. The procedure yielded frank pus, 1500 ml, and some specimen was sent for microbiology but the culture yielded no growth. He was placed on twice-daily dressings with honey-soaked gauze packing and parenteral antibiotics. By the 12th day post-op we sent him for CT scanning with bone window and contrast which confirmed fronto-parietal osteomyelitis but no intracranial abscess. Craniectomy and delayed primary scalp closure were done and the patient was placed on post-operative antibiotics for another two weeks and subsequently, discharged home on per-oral antibiotics with no neurological deficits.

Discussion

Pott’s Puffy Tumour is thought to be a disappearing disease in the antibiotic era, and most are thought to arise from contiguous paranasal sinus spread mostly from the frontal bones according to many reports [19,20,1,3,5,10,11,12,13,15,16,17,18]. Our case appears, instead, to have been a case of de novo parietal osteomyelitis that was neglected for long and extended to involve the frontal bone over a period of 6 months. There was no history suggestive of trauma prior to the onset of illness but this would not be completely ruled out in an active young male student. There was, however, a definite denial of any history of upper respiratory tract infection in association with the fever, scalp pains and scalp swellings which was quite unlike other reports [3,5]. But periorbital
oedema and orbital cellulitis had in our case have been reported by Lamoreau and Fanciullo, Morley, and Hayek, et al to be associated with this disease [7,8,9]. Despite the classical advocacy of urgent CT scanning and definitive surgical intervention, we modified our protocols in the light of our technical handicaps and did a preliminary incision and drainage to relieve sepsis, reduce pains and save life, before resorting to a classic work-up and definitive treatment; and still got a good outcome. It would have been unwise to insist on a full work-up with CT scanning in the light of the enormous size of the abscess and painful incapacitation of this patient, thus we went ahead with a simple drainage procedure first, and only obtained a proper imaging in a more stable patient, following which we did craniectomy as the definitive procedure.

**Conclusion**

In these modern times, it should be realized that sub-optimal care and mismanagement of clinical cases still pervade our health system and a case as grotesque as ours could still be found in this era of antibiotics. Also, without readily available Computed Tomography, one could resourcefully modify the standard protocols to provide a fairly good surgical care for the patient; so long as the basic principles are strictly adhered to, thereby reducing morbidity and mortality.

**References**

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Comment

The historical description of a sub-periosteal or sub-galeal purulent collection at the cephalic region, by Sir Percival Pott (1714-1788), known classically as Pott’ Puffy Tumor, recently falls in the definition of a subgaleal and / or sub-periosteal abscess, with the classical characteristics of inflammation secondary to an infection, with the presence of flushing, heat, pain and increased volume (tumor).

This purulent collection is usually due to the spread of an infection from the paranasal sinus reached by a cranial fracture, which allows the passage of germs into neighboring tissues; also is due to traumatic brain injury, with open trauma of the scalp, but contaminated, and subsequently due to an inappropriate treatment and control, infecting the skull, establishing an osteomielitis, thus allowing the collection of germs in closed areas at the subperiosteal or sub-galeal spaces; and finally, that could correspond to the case reported by J.K. Emegjulu and I. BC. Iloabachie, to minor trauma, accidental or provoked, some caused by sharp objects, which can cause late osteomyelitis of the calvarium.

It’s necessary to comment, that noting the patient’s clinical conditions reported, the authors performed the best, that is to solve the subgaleal abscess with conventional surgical techniques, primarily to drain the pus and then studying the patient in order to search the origin of the infection, which was observed in the phase 3D Computed Tomography, which corresponded to a location of the osteomyelitis at the parietal bone, left posterior region.

Later, it was the best performed, removing the bone with osteomyelitis and infected splinters, and starting specific antibiotic treatment according to the antibiogram of the evacuated pus. The cranioplasty must be made on a deferred time, once septic parameters have normalized. It is a well-described and documented case by the authors, which was resolved in an appropriate manner, despite not having some immediate accessory diagnostic tools.

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