Non-traumatic fractures following seizures: two case reports

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Abstract

Introduction: Seizures with or without trauma may cause fractures that occur commonly in epileptic seizures. Fracture risk is less reported in non-epileptic seizures. Some metabolic conditions leading to a decrease in bone mineral density may cause fractures secondary to non-epileptic seizure.

Case presentation: We describe two cases of non-traumatic acetabular and vertebrae fractures following seizures without history of epilepsy. They occurred in two male patients, 18 and 48 years old suffering respectively from hypercorticism and poorly controlled diabetes mellitus. Seizures, occurring inside hospital, were secondary to hypertensive encephalopathy crisis with hypokaliemia in the first case and severe hypoglycaemia in the second one. Fracture was promoted by a decrease in mineral bone density caused respectively by hypercorticism and diabetic chronic renal failure.

Conclusion: These observations emphasize that fracture prevention among patients with decreased mineral bone density should include the avoidance of metabolic causes of seizure.

Introduction

Seizures increase fracture risk [1], occurring in the whole of the skeleton even without trauma [2]. This risk is mainly reported in epileptic seizures because of repeated occurrence of attacks and osteopenia due to the use of anti-epileptic drugs [1]. Literature data have shown that fractures are more related to the seizure itself rather than to the iatrogenic factor [1]. Moreover, some metabolic conditions may lead to a decrease in bone mineral density and hence may be considered as risk factors of fracture when seizures occur. We report here two cases of this unusual situation to resort its eventual prevention.

Case presentation

Case report 1
An Arabic 48-year-old man from North Africa (Tunisia) with a ten year history of diabetes mellitus is treated by insulin twice a day for seven years. His diabetes is poorly controlled mainly due to lack of observance. Additionally, he suffered from depression with suicide ideas. Shortly after an overdose of insulin, he developed generalized tonic-clonic seizures that was not followed by trauma and that regressed after parenteral hypertonic glucose serum perfusion.

Laboratory assessment showed a severe hypoglycemia (capillary glycemia at 1.8 mmol/l) and nephrotic syndrome with renal failure (creatininemia at 241 μmol/l). After seizure, the patient presented a disability of the left lower limb. X-ray examination showed a fracture in the left iliac bone localized in the acetabular floor (Figure 1) while brain CT-scan was normal. The patient was referred to orthopedic center where he was treated conservatively. Six months later, the patient was able to walk. Insulin therapy was adjusted aiming to avoid hypoglycemia.

Case report 2
An Arabic 18-year-old man from North Africa (Tunisia) was referred to our department for suspicion of Cushing syndrome. On physical examination, the patient was obese (BMI at 32 kg/m^2), he had purplish abdominal striae and hypertension (systolic pressure between 15 and 17 mmHg). Neurological examination was normal.

Laboratory examination showed hypokalemia at 2.1 mmol/l confirmed in several samples separately performed while glycemia and calcemia were at normal range. Urinary free cortisol excretion was high (462...
μmol/24 h) and was not suppressed after both low and high doses of dexamethasone confirming the diagnosis of Cushing syndrome. X-ray examination (Figure 2) and bone densitometry revealed decrease of bone mineral density with a Z-score of -2. Brain MRI imaging showed a double pituitary adenoma without any other hemispheric lesion. The diagnosis of corticotropic adenoma was retained.

During an acute hypertensive peak at 220 mm Hg, the patient developed tonic-clonic seizures without any trauma. His glycemia was normal (5.6 mmol/l). Seizures regressed after intravenous load of Loxen* and Diazepam* however the patient suffered from mild low back pain. Radiographies identified L2, L3, L4 and L5 burst fractures (Figure 3).

Cushing disease was surgically treated with regression of hypertension and weight. Bone mineral density was checked two years later and showed an improved Z-score.

Discussion
We reported here two cases of hip and vertebrae fractures following non-epileptic generalized tonic-clonic seizures without any trauma or fall. Several data have reported fractures during seizures. Fracture risk is notably increased mainly among epileptic patients. It occurs in 11% of patients mainly within the first two years of diagnosis. This risk is directly related to seizures in 43% of cases in one of the largest series [3].

Fractures concern the whole of skeleton particularly the skull, members, vertebrae and the hip and may be multiples, bilateral or associated to other lesions such as dislocations of the shoulder [4,5]. They induce an increase of management cost of epileptic disease [6]. Acetabular fracture observed in the first patient has also been reported and may be serious leading to fatal pelvic haemorrhage [7,8].

Non-epileptic seizures reported in our cases are unusual. In our first case, seizures are related to severe hypoglycaemia, which is common cause of seizures. They were reported in 4.7% of all the hospitalisations for hypoglycaemias especially in old and depressive patients [9]. Concerning the second patient, seizures are caused by encephalopathy hypertensive crisis in addition to hypokaliemia due to hypercorticism. These results are in line with database that emphasizes the role of hypertensive encephalopathy [10], and ionic disorders in seizures such as hypokaliemia and hypocalcaemia [5].
Fractures occurring without any trauma suggest previous bone fragility due to other illness [5]. Bone fragility has been confirmed in our second patient by bone mineral densitometry assessment before seizures.

Our cases concern male young patients who are not at risk of primary osteoporosis. Among classic reported causes of male secondary osteoporosis [11], we can incriminate secondary hyperparathyroidism caused by diabetic renal failure in the first patient [12] and hypercorticism in the second one [13].

Hypercorticism is one of the main diseases that cause osteoporosis leading to fractures. Vertebral bodies and ribs are the typical sites of fracture both in Cushing’s syndrome and in patients using glucocorticoid drugs long-term [14]. The frequency of vertebral fractures has ranged from 16 to 20% [14]. Decreased bone mineralization is particularly pronounced in children and in young females with Cushing’s syndrome [14]. Interestingly, earlier studies had shown that young and adolescent patients undergoing glucocorticoid therapy may lose bone mass more rapidly than older patients [15]. Fracture risk is higher at the onset of disease and may reveal it. It may also occur during clinical investigations such as in our patient [13] or when diagnosis is delayed [16] leading to definitive bone sequels.

Nevertheless, diabetes mellitus is an unusual cause of osteoporosis and usually we find associated fracture risk factors [17]. Diabetes could play an indirect role through increasing fall risk-related fractures due to feet injuries [18]. Of interest, some prospective data showed that diabetes mellitus is a statistically risk factor of osteoporosis especially in insulin treated diabetes, long history of diabetes or in elderly [19].

The role of metabolic control in osteopenia is still not well established despite that some data suggest that insulin, growth factors or osmotic factors such as extracellular glucose levels may play a significant role in the osteoblast metabolism [20].

**Conclusion**
These cases demonstrate the possibility of non-traumatic fracture following seizure. Diabetes mellitus and
hypercorticism may cause, even in males, osteoporosis and subsequently are considered as risk factors of seizures and fractures. Importantly, these conditions may be well-managed to prevent osteoporosis. We insist not only on avoiding falls and trauma but also on preventing metabolic causes of seizures such as hypoglycaemia, ionic disturbances and hypertensive encephalopathy.

Consent
Written informed consent was obtained from the patient for publication of this case report and accompanying images. A copy of the written consent is available for review from the journal’s Editor-in-Chief.

Authors’ contributions
KA, IS, STA followed the patients reported here. KA, IS wrote the paper, MCC, AMB, LC reviewed the paper.

Competing interests
The authors declare that they have no competing interests.

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