Fatal Hypocalcaemia Due to Hungry Bone Syndrome with Secondary Refractory HyperParathyroidism after Parathyroidectomy: A Case Report

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ABSTRACT

Introduction: Hungry bone syndrome (HBS) refers to the rapid, profound, and prolonged hypocalcaemia associated with hypophosphatemia and hypomagnesaemia, and is exacerbated by suppressed parathyroid hormone (PTH) levels, which follows parathyroidectomy in patients with severe primary hyperparathyroidism (PHPT) and preoperative high bone turnover. [1]

Case report: This report concerns a dialysed patient who underwent surgical treatment for secondary refractory hyperparathyroidism. Haemodialysis was carried out pre-operatively, and subsequently, a total parathyroidectomy with auto-transplantation of parathyroid tissue in the sternocleidomastoid muscle (SCM) was performed. Rapid and progressive hypocalcaemia symptoms developed during the second day postoperatively. Acute cardiac symptoms with tachyarrhythmia, haemodynamic instability and finally asystole occurred, which required cardiopulmonary resuscitation (CPR). The ionic calcium level was 2.2 mg/dL being consistent with a diagnosis of HBS. A second cardiac arrest unresponsive to CPR followed an initial period of normal sinus rhythm. Death ensued shortly after. Before death, the ionic calcium was 3.1 mg/dL.

Conclusion: HBS, after parathyroidectomy in patients with secondary hyperparathyroidism (SHPT), may be severe, prolonged and sometimes fatal. Generally, HBS symptomatology is that of a mild hypocalcaemia. It can, however, include heart rhythm disturbances with haemodynamic alterations requiring intensive care measurements and even cardiopulmonary resuscitation. A close clinical and laboratory post-parathyroidectomy monitoring of dialysed patients is of the utmost importance.

Keywords: hungry bone syndrome, secondary hyperparathyroidism, cardiac complications

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INTRODUCTION

Hungry bone syndrome (HBS) denotes to the rapid, profound, and prolonged hypocalcaemia with attendant hypophosphataemia and hypomagnesaemia, and is aggravated by lowered parathyroid hormone (PTH) levels, which follows parathyroidectomy in patients with severe primary hyperparathyroidism (PHPT) and preoperative high bone turnover [1].

This clinical state usually extends for more than three to four days after a parathyroidectomy. It is argued that HBS is due to the sudden increased calcium influx into the “starved” bone, primed by the decrease of serum parathyroid hormone level (iPTH) post-parathyroidectomy [1]. Hypocalcaemia and HBS are considered either as a postoperative complication or conversely as indicative of a “successful” parathyroidectomy [2]. HBS occurs both after parathyroidectomy performed in primary hyperparathyroidism (PHP) and refractory secondary hyperparathyroidism (SHPT), in the latter case exhibiting more severe and prolonged forms [3]. Several HBS risk factors are described, including el-
evated preoperative levels of serum iPTH or alkaline phosphatase (AlkPhos) levels, advanced age, preoperative bone damages [4]. HBS symptoms are similar with those of a severe hypocalcaemia: from minor parasthesia, carpopedal spasms, positive Chvostek and Trousseau signs to convulsions, laryngospasm or acute cardiac dysrhythmias.

The current case report concern the treatment of a dialysed patient with severe HBS which occurred after a total parathyroidectomy performed for secondary refractory hyperparathyroidism, in whom symptoms of hypocalcaemia developed the day following the operation. Heart rhythm disturbances were followed by cardiac arrest. Resuscitation measures were unsuccessful.

**Case report**

A 65-year-old man, BMI of 33 kg/m², was admitted to the Nephrology and Endocrinology/Internal Medicine Department of Harghita General Hospital (Harghita General Hospital, Miercurea Ciuc, Romania), with generalised weakness, severe pruritus and difficulty walking due to severe left leg pain. His medical history included end-stage renal disease (ESRD) due to adult polycystic disease having been on chronic haemodialysis three times weekly for the past eight and a half years. He had been diagnosed with secondary hyperparathyroidism approximately three years ago, and since then he had been treated with active metabolites of vitamin D and bisphosphonates. Longstanding comorbidities such as arterial hypertension, aortic and mitral stenosis, atrial fibrillation, and dyslipidaemia were also noted. For these, he had been prescribed antihypertensive, statins and antiplatelet drugs. Additionally, a diagnosis of calciphylaxis was included in the patient’s medical papers, without any indication as to who made the initial diagnosis. A few months prior to admission, calcium based phosphate binders had been prescribed; sevelamer carbonate [Renagel, Sanofi, Germany] oral capsules, 800 mg three times daily.

Taking into consideration the refractory secondary hyperparathyroidism with no improvement in the patient condition under the medical treatment stated above, along with the very high levels of iPTH, hypocalcaemia and hyperphosphatemia, the attending nephrologist decided to refer the patient to the Second Surgical Department Targu Mures for parathyroidectomy.

Physical examination on admission to the surgical department was non-specific for other endocrine diseases. Blood chemistry showed serum calcium levels of 11.6 mg/dL, nv: 9-10.5 mg/dL, and serum phosphorus levels of 5.6 mg/dL, nv: 2.7-4.5 mg/dL, very high levels of serum intact parathyroid hormone of 1257 pg/mL, nv: 10-65 pg/mL and high level of alkaline phosphatase of 376 U/L, nv: 40-150 U/L. These data are consistent with a high bone turnover due to uremic osteodystrophy. Table 1 details all laboratory data generated on admission.

Targeted parathyroid imaging, i.e. 99mTc Sestamibi scintigraphy, ordered by the nephrologist in Tg Mures Nuclear Medicine Department, two months earlier, revealed a single image suggestive of parathyroid adenoma, located on the lower part of the right thyroid lobe (Figure 1).

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### Table 1. First admission, preoperative and postoperative day 1 and 2 laboratory data

|                      | First admission values | Preoperative values | Postoperative values (POD 1) | Postoperative values (POD 2) |
|----------------------|-----------------------|--------------------|-----------------------------|-----------------------------|
| WBC (vn: 3.6-10 x 10³) | 8.2 x 10³             | 7.1 x 10³          | 7.3 x 10³                   | -                           |
| Haemoglobin (vn:12-17 g/dL) | 14.6 g/dL           | 14 g/dL            | 13.7 g/dL                   | 13.5 g/dL                   |
| Haematocrit (vn: 36-54 %) | 43%                  | 42%                | 41%                         | 38%                         |
| Total calcium (vn: 9-10.5 mg/dL) | 10.9 mg/dL        | 11.6 mg/dL         | 7.3 mg/dL                   | -                           |
| Ionic calcium (vn: 3.8-4.8 mg/dL) | -                   | -                  | -                           | 2.2-3.1 mg/dL               |
| Phosphate (vn: 2.7-4.5 mg/dL) | 4.9 mg/dL          | 5.6 mg/dL          | 5.3 mg/dL                   | -                           |
| Sodium (vn: 136-145 mmol/L) | 139 mmol/L         | 141 mmol/L         | 136 mmol/L                  | 138 mmol/L                  |
| Potassium (vn: 3.5-5.1 mmol/L) | 4.9 mmol/L      | 5.1 mmol/L         | 6.2 mmol/L                  | 6.2 mmol/L                  |
| Chloride (vn: 98-107 mmol/L) | 104 mmol/L        | 102 mmol/L         | 99 mmol/L                   | -                           |
| AlkPhos (vn: 40-150 U/L) | 364 U/L            | 376 U/L            | 311 U/L                     | -                           |
| iPTH (vn: 10-65 pg/mL) | 1198 pg/mL        | 1257 pg/mL         | 47.5 pg/mL                  | -                           |
| Creatinine (vn: 0.72-1.25 mg/dL) | 5.1 mg/dL       | 5.29 mg/dL         | 6.7 mg/dL                   | -                           |
| Glucose (vn: 80-115 mg/dL) | 81 mg/dL           | 89 mg/dL           | 96 mg/dL                    | 85 mg/dL                    |

# = WBC= white blood cell count, AlkPhos= alkaline phosphatase, iPTH= intact parathyroid hormone
On admission to the surgical department, the initial preoperative protocol included endocrinology reassessment of the case which confirmed the diagnosis and surgical indication. Cardiological and anaesthesia evaluations were also conducted before surgery. Additionally, preoperative dialysis was carried out the day before surgery.

The patient underwent a total parathyroidectomy with autotransplantation of parathyroid tissue (tPtx+AT) in sternocleidomastoid muscle (SCM). The weight of the resected parathyroids was 1.42 g.

The procedure was done under general anaesthesia, and no complications occurred.

Immediately after the operation, the serum iPTH level dropped to 47.5 pg/mL, and the patient was transferred to the ward in a stable condition.

The first postoperative supplementation of calcium in dialysate fluid was performed on the first day after the operation (POD 1). In spite of this a post-dialysis serum total calcium level of 7.3 mg/dL was recorded for which continuous micro-infusion of 10 ml calcium gluconate 94 mg/ml [B Braun Melsungen AG Germany] and then 30 ml of calcium gluconate in continuous micro-infusions four times daily. Within a few hours of the administration of the supplementary calcium, the patient developed atrial fibrillation with tachyarrhythmia (heart rate > 150/min), acutely altered mental status and a rapid drop in blood pressure to 70/40 mm Hg.

Norepinephrine [Noradrenaline, 2mg/ml, Aguettant Ltd, UK] was commenced as intravenous infusion for haemodynamic support and synchronised cardioversion. Shortly after that, the patient developed asystole, for which cardiopulmonary resuscitation (CPR) was performed for thirteen minutes. After an initial period of no more than ten minutes, when there was a return to sinus rhythm and a slightly hemodynamic amelioration, another cardiac arrest ensued for which standard hospital resuscitation protocol was initiated. External thoracic compressions and ventilation, CPR was performed for thirty minutes during which time a total of 10 mg of epinephrine [adrenaline, Terapia SA, Romania] at 1 mg/ml, was given.

The patient was unresponsive to these resuscitation manoeuvres and died. Before the patient’s death, ionic calcium was 3.1 mg/dL.

**DISCUSSIONS**

Hypocalcemia after parathyroidectomy is expected to happen but is generally oligosymptomatic and is solved in the first 3-4 days postoperatively. Extending this period of severe hypocalcaemia, possibly accompanied by other ionic disorders such as hypophosphatemia and hypomagnesaemia, defines HBS. Strictly nosologically, within the same syndrome hypocalcaemia immediately after parathyroidectomy may occur, which is based on the same pathophysiological mechanism i.e. the sudden decrease of postoperative iPTH and the impact that the hormone has on the osteoclastic resorption [5].

The prevalence of HBS is contrarily reported in the literature, with figures ranging between 4-95% depending on several factors, i.e. primary or secondary parathyroid disease, geographic area [6]. The complication occurs more frequently after parathyroidectomy for re-
nal hyperparathyroidism, where the clinical manifestation of HBS is more severe [7]. Such a case, with rapid fatal development shortly after parathyroidectomy, is presented here. The literature reports relatively few such situations, those reported generally concerning post-surgical treatment of SHPT [8,9].

From the pathophysiological point of view, HBS can be defined as an alteration of the bone remodelling process, which usually removes the osteoclastic activity from the affected bone and replaces it with new bone osteoblastic activity. This balance is lost in primary and secondary hyperparathyroidism, where the increased osteoclastic activity is stimulated by increased secretion of iPTH, which explains demineralisation and hypercalcemia in hyperparathyroidism. The sudden reduction of iPTH levels immediately after parathyroidectomy primes a change in ionic fluxes, with rapid remineralisation of the “starved” bone, and sometimes severe and prolonged reduction of serum calcium and other ions [10]. The syndrome should not be confused with postoperative hypocalcaemia that occurs in other situations such as thyroid surgery or bariatric surgery, where the pathophysiology of the complication is different, i.e. an inadvertent resection of parathyroids, intestinal malabsorption [11,12]. Similar ionic disorders including hypocalcaemia, hypokalemia also occur in other postoperative or posttraumatic conditions such as peripheral or abdominal compartment syndrome, rhabdomyolysis, but in these cases, the pathophysiology differs from HBS [13,14].

The literature describes several risk factors for HBS [2]. Among these factors, most authors agree that the presence of preoperative parathyroid-related bone lesions (osteitis fibrosa cystica, brown tumours, pathologic fractures) is significantly associated with the development of postoperative HBS [15,16]. In the case described here, there were no X-rays on admission to objectify these signs, but patient described multiple fractures in the disease's history. High levels of preoperative serum AlkPhos is also considered a predictive factor for parathyroid-related bone disease and therefore for HBS [16]; our patient's preoperative serum level of AlkPhos was 376 U/L (nv: 40-150 U/L). There are also other risk factors for HBS, which were noted in this case: high values of preoperative serum iPTH, older age, the increased volume and weight of resected parathyroid glands. As regards this latter aspect, Zamboni & Folse [17] confirm the positive statistical correlation between the weight of the resected adenomas and the postoperative incidence of hypocalcaemia and HBS.

Surgical interventions indicated in SHPT are subtotal parathyroidectomy or total parathyroidectomy with or without autotransplantation of parathyroid tissue. A total parathyroidectomy with parathyroid autotransplantation in the sternocleidomastoid muscle was performed in the presented case. Regarding the incidence of postoperative HBS, literature does not describe statistical differences between parathyroidectomy techniques. Total parathyroidectomy without autotransplantation was initially credited with a higher risk of postoperative hypocalcaemia, which is why the method was abandoned for many years. In recent years, however, various studies have demonstrated that this technique is feasible without higher postoperative morbidity, including hypocalcaemia and HBS [18].

Clinical manifestations of HBS are similar with those of severe hypocalcaemia: from minor carpopedal spasms, perioral paresthesia, tingling extremities, clinical signs of hypocalcaemia (Chvostek, Trousseau) to more severe symptoms such as convulsions, cardiac symptoms, coma and even fatal outcome. Our patient did not experience symptoms of hypocalcaemia on the first postoperative day; 24 hours after surgery, however, a sudden cardiac tachyarrhythmia occurs, very likely in the context of hypocalcaemia, the ionic calcium determined at the time being very low. Heart rhythm disorders are also a part of classical semiology of hypocalcaemia, in the context of hypocalcemic cardiomyopathy being frequently described QT interval prolongation, tachyarrhythmia or postoperative exacerbation of latent cardiac insufficiency [19,20], as it is probably also the case of the patient presented.

Hungry bone syndrome's treatment is primarily a preventive one. Patients in the risk group may benefit from preoperative administration of vitamin D or its active metabolites. Boyle et al. [21] showed that the preoperative treatment of severe hyperparathyroidism with calcitriol (1,25 (OH),D), started few weeks before parathyroidectomy, significantly reduces the preoperative values of AlkPhos and the need for postoperative calcium administration. Other authors stipulated the beneficial effect of bisphosphonates in reducing preoperative calcium level and postoperative HBS incidence [22]. When HBS diagnosis is established the curative treatment includes administration of elemental calcium, initially consisting of IV calcium gluconate and once the patient can tolerate, oral calcium supplements,
tailoring the dose and the frequency of administration according to serum calcium level [18].

Several electrical cardiac changes seen during severe hypocalcemia episode due to the same reason, as explained above were not objectively demonstrate in the present case.

**CONCLUSION**

HBS is a severe complication of parathyroidectomy in patients with PHPT and SHPT. In the case of SHPT, hungry bone syndrome may by severe, prolonged and sometimes fatal. Generally, HBS symptomatology is that of a mild hypocalcaemia; in some cases, however, the symptomatology includes heart rhythm disturbances with hemodynamic alterations requiring intensive care measurements and even cardiopulmonary resuscitation. Therefore chronic dialysed patients with recent parathyroidectomy should be closely monitored, here including serial serum calcium dosing, for proper substitutive treatment in the hours and days following surgery.

**CONFLICT OF INTEREST**

None to declare.

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