**CASE REPORT**

**Successful treatment of vitamin D unresponsive hypoparathyroidism with multipulse subcutaneous infusion of teriparatide**

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**Abstract**

**Objective:** Hypoparathyroidism is usually controlled with calcium and vitamin-D supplements; in very few cases this treatment fails and teriparatide may be an alternative. We report the first case of hypoparathyroidism refractory to vitamin-D therapy requiring multipulse teriparatide treatment.

**Case report:** A 53 year-old woman presented severe hypocalcemia and hypomagnesemia after thyroidectomy. Preoperatively, mild hypercalciuria was detected with parathyroid hormone (PTH) 69 pg/ml (normal 10–45) and 25-OH-vitamin D 9 ng/ml (normal 20–40) and normal levels of magnesium. No response was seen with oral and i.v. calcium and magnesium, or even with 5 μg calcitriol/day, suggesting a vitamin-D resistance status. Calcium sensor and vitamin-D receptor gene mutation studies were negative.

**Interventions and results:** The following treatments were tried: i) s.c. recombinant human PTH (rhPTH) 1–34 plus oral calcitriol, calcium, and magnesium, was partially effective, but symptoms resumed 4 h after the injection of 20 μg rhPTH; stable calcemia was not achieved even with 4–6 injections/day of teriparatide; ii) two trials of heterologous parathyroid transplant were performed but rejection was detected 3 months after; iii) i.v. magnesium decreased rhPTH requirements but i.m. administration was not tolerated and iv) multipulse s.c. infusion of teriparatide achieved complete normalization of serum calcium, phosphate, magnesium, calcitria and magnesuria with relatively low rhPTH doses (25–35 μg/day) for more than a year.

**Conclusions:** Vitamin-D unresponsiveness leads to uncontrolled hypocalcemia when postsurgical hypoparathyroidism occurs; in situations of no response to usual or higher doses of vitamin-D and s.c. injections of rhPTH, treatment with teriparatide multipulse s.c. infusor is an effective and safe alternative.

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**Introduction**

Injury or removal of the parathyroid glands during neck surgery is the most common cause of acute and chronic hypoparathyroidism (1). Its incidence rate is usually related to the surgeon’s experience, the type of parathyroid pathology and the surgical technique performed. Although postsurgical hyperparathyroidism is usually transitory, it may persist in 0.4–33% of cases depending on the series (2, 3). The conventional treatment of postsurgical chronic hypoparathyroidism is based on calcium salts, vitamin D (mainly calcitriol), and drugs that increase renal tubular reabsorption of calcium as thiazides. Over the past few years, the administration of synthetic recombinant human parathyroid hormone (rhPTH) 1–34 once or twice daily in patients with hypoparathyroidism has proved to reduce urinary calcium excretion compared with calcitriol therapy, and to maintain serum calcium in the normal range, thus avoiding chronic hypercalciuria that may lead to renal function impairment, nephrocalcinosis and renal insufficiency in the long term (4–7).

In the present paper, we describe a 53 year-old woman with post-thyroidectomy uncontrolled hypoparathyroidism despite the usual therapy with high doses of calcitriol plus calcium and magnesium salts. Her life-threatening hypocalcemia and hypomagnesemia required different treatments over 12 months, all of them unsuccessful for controlling calcemia; the final and only effective treatment was a multipulse s.c. infusion of teriparatide which achieved a complete clinical and analytical control of the hypoparathyroid state.

**Patients and methods**

**Case presentation**

A 53 year-old woman was referred to our Endocrinology Service after thyroidectomy performed due to multinodular goiter with mild retrosternal extension...
and subclinical hyperthyroidism. Prior to this, she had already visited our hospital for a larva migrans chronic infestation acquired during a trip to Thailand 9 years before, and was diagnosed 3 years previously. During the follow-up she developed asthma, and IgG subclasses 1 and 3 deficiencies were detected. During the 8 years, she required corticosteroid treatment either inhaled or oral, sometimes during several months in order to prevent and treat episodes of bronchospasm crisis. Consequently, bone mass was periodically checked; bone mass densitometry (BMD) revealed an osteopenic state; besides this, low 25-hydroxyvitamin D (25-OH-vit D) of 9 ng/ml (normal 20–40) was also found, together with a moderate increase in circulating PTH level of 69 pg/ml (normal 10–45), while renal function, calcium (ranging 8.8–9.7 mg/dl) and magnesium levels in plasma and urine were normal. These data were interpreted as corticosteroid-induced osteopenia, associated with vitamin D deficiency and some degree of compensatory hyperparathyroidism; she was therefore treated with cholecalciferol over at least 2 years, with normalization of circulating 25-OH-vit D. No bisphosphonate treatment was used at any time. She also suffered two septic episodes during the previous 3 years, one due to enterococcus faecalis and another of polymicrobial etiology. No septic foci were found despite extensive investigations. Treatment for larva migrans eradication with albendazole, praziquantel, and ivermectine for more than 2 years was ineffective and stopped before treatment of her hyperthyroidism; at present the patient still remains with active infestation.

Total thyroidectomy was performed on April 2005 because of her multinodular goiter, followed by immediate postoperative severe and persistent hypocalcemia. Circulating PTH was undetectable from the first postoperative day. Despite oral calcium and calcitriol started at rapidly progressive doses, serum calcium stayed low (average 6 mg/dl; normal 8.5–10.5) and phosphate high (average 5 meq/dl; normal 2.3–4.3) even with 5 mg calcitriol/day and more than 10 g calcium/day. She complained of fatigue, palpitations, paresthesia and tetany almost daily, and required emergency assistance with i.v. calcium infusion several times a week because of life threatening hypocalcemia mostly due to severe tetania and impairment of asthma, with increasing frequency of episodes that were more refractory to usual treatment under hypocalcemic conditions (Fig. 1A). Hypocalcemia was associated with low serum magnesium (average 1.5 mg/dl; normal 1.8–2.6), requiring high oral doses plus parenteral loads of magnesium. Urinary excretion of calcium and magnesium markedly increased after i.v. infusion, achieving intolerable high levels, with calcitriuria up to 900 mg/24 h. Despite increasing doses of calcitriol that led to circulating 1,25-OH-vitamin D in the upper range of normality (65 to 75 ng/l; normal <70 ng/l), serum calcium levels remained very low, rarely above 7 mg/dl, apparently indicating some state of vitamin D resistance. A trial of thiazide treatment was unsuccessful with virtually no amelioration of serum calcium levels. A potential malabsorptive state was ruled out by a thorough investigation, including xylose test and radiological and endoscopic procedures, all of which resulted normal. Furthermore, negative antitransglutaminase antibodies ruled out celiac disease. A calcium sensor mutation study was negative. Complete sequencing of the vitamin D receptor gene was performed in Dr Peter Malloy’s laboratory at Stanford University as previously described (8) and no mutations were found. After 6 months of uncontrolled life-threatening hypocalcemia and hypomagnesemia under high doses of conventional therapy, the treatments described below were tried after obtaining the patient’s informed consent and the approval of the Ethical Committee of the Hospital Clinic Universitari de Barcelona.

**Methods**

S.c. rhPTH 1–34 (Forsteo, 20 μg/80 μl; Eli Lilly) was started at a dose of 20 μg/12 h s.c., according to previously published data (3, 4) plus oral calcitriol 5 μg/day, 10 g calcium po/day and 0.5 g magnesium po/day, achieving partial serum calcium control (Fig. 1B); yet, symptoms tended to resume 4 h after

![Figure 1](https://www.eje-online.org)

**Figure 1** (A) Response to oral calcium up to 10 g po/day and 1,25-OH-vitamin-D up to 5 μg/day after development of hypoparathyroidism following the neck surgery. (B) After 2 months of unresponsiveness rhPTH was added at increasing doses with some initial response but very modest efficacy during 9 months.
commercial preparation and a complete normalization of serum calcium, phosphate and magnesium, as well as calciuria and magnesuria were achieved in the first day of treatment (Fig. 2B). In the following days, teriparatide infusion rate was tapered down until the minimal dose that allowed a low normal range of calcemia; successful treatment was achieved at final doses of teriparatide ranging between 25–35 μg/day. With this treatment regimen serum calcium levels stayed within normal values in most of the follow-up period, until now.

The pump was programmed in order not to give a theoretically continuous delivery of teriparatide, but a multi-micropulse release of the peptide, trying to mimic the physiologic PTH secretion, which has been described to be about 6–7 bursts per hour (10, 11). The MiniMed 508 pump used in our patient is based on Bio-Pulse Delivery technology, allowing a pulse design modality of treatment in which a pulse of teriparatide was automatically delivered at intervals of 10–14 min.

The reservoir of the pump was refilled every 3 days with no evidence of a reduction in therapeutic effectiveness over time because of potential peptide degradation due to room temperature exposure. The injection site was carefully cleaned and catheter/needle and injection site changed every 3 days, coinciding with the refilling of the teriparatide reservoir. No major problems have arisen since initiation of this therapeutic modality and the patient has remained virtually normal in serum and urinary calcium and magnesium levels. After 3 months of treatment some loss of efficacy was observed, a tendency toward hypocalcemia, which coincided with the development of micronodules in the s.c. abdominal wall. This episode was rapidly resolved by changing the injection site and not using the specific area for several weeks, during which the micronodularity disappeared spontaneously. On another occasion the patient forgot to refill the reservoir; rapid sub-tetany developed, and the alarms of the infusion system switched on; the episode was resolved with the injection of a bolus of 20 μg teriparatide after the reservoir was refilled. Beside these
2 episodes, the patient has remained asymptomatic with normal or near-normal biochemical profile for more than 1 year by using an average of 30 μg/day teriparatide; these requirements have remained very stable during this time, indicating a lack of development of anti-teriparatide antibodies or any sort of tachyphylaxia. She did not refer to bone pain and follow-up evaluation of BMD showed a recovery of bone mass, with an increase in 7% in the first year of treatment, as expected from previously published data (12).

Discussion

We describe for the first time a case in which teriparatide multipulse pump infusion was the only way to control hypocalemia in postparathyroidectomy hypoparathyroidism. Fortunately, in most patients with hypoparathyroidism, regardless of the cause, an acceptable control of calcium levels is possible by different dosage regimes of vitamin D and calcium supplements (1). A therapeutic target of low-normal serum calcium, so as to prevent hypercalcemia and long-term detriment of renal function is the ideal. In the clinical setting this is usually feasible, and this is why it is still the treatment of choice of these patients (13). The high cost of rhPTH and the inconvenience of injection treatment are the reasons why vitamin D plus calcium is preferred to this newer, more physiological approach, despite the risk of hypercalcemia. In our patient, we were unable to reestablish circulating normal calcium or at least to avoid symptoms with this usual vitamin D treatment. Even very high doses of calcitriol (5 μg/day, instead of the recommended 0.25 to 1 μg/day), were not effective. A malabsorption state was ruled out by different procedures, and calcitriol oral loads led to normal or higher than normal circulating 1,25-OH-vit D levels together with intense hypercalcemia, indicating that calcium absorption was probably not impaired, although fractional calcium absorption studies were not performed. In order to explain this intriguing lack of effect of very high doses of calcitriol, vitamin D receptor gene was studied (8), but its complete sequencing showed no mutations. As the main regulatory action of vitamin D is to stimulate calcium absorption in the bowel, hypercalcemia in conjunction with high levels of vitamin D probably indicated that vitamin D was in fact normally acting at the bowel, while maybe a selective or preferential defect of vitamin D action on the kidney caused urinary calcium loss leading to hypercalcemia. Alternatively, hypercalcemia may also be due to a specially exquisite dependence of PTH in this patient. Also, no calcium sensor gene mutations were found (14), and although no other potential tubular defects were explored, such as concomitant magnesium, losing disorders or CLDN16 gene mutations (9, 15), these possibilities were considered unlikely.

At this point, we tested the hypothesis that the patient was extremely dependent on normal circulating PTH for regulation of her calcium levels. The life threatening condition of her hypocalemia state, with aggravation of asthmatic crisis, led to transplantation of cross matched parathyroid tissues from end stage renal tertiary hyperparathyroidism patient. This kind of treatment has been used in patients with unresolved hypoparathyroidism despite classic treatment with calcium plus vitamin D (2). The few reported successful cases achieve normal calcium levels about 3 months after the procedure, once the implanted tissue gets vascularized. Rejection may always happen (16), as unfortunately was the case in our patient.

Administration of rhPTH can be an interesting and effective treatment in selective cases such as the present one. In the cohort studied by Winer et al. (4–6), two injections of a mean of 50 μg of rhPTH achieved an adequate level of circulating calcium in almost all cases. In our patient such an easy control was not possible, even though doses were swiftly raised from those proposed by Winer, until they reached very frequent peptide injections. We have no explanation for this unresponsiveness to multi-injections of teriparatide. Surprisingly, its administration by a multipulse pump driven system resulted in both highly effective and safe treatment over more than 1 year. Moreover, the absolute dosage used was markedly lower in comparison with the previous ineffective multi-injections schedule, which reduced the cost of teriparatide and was a less uncomfortable treatment for the patient. In the last year a new rhPTH delivery system consisting of an implantable biodegradable polymer has been described, allowing pulsatile release of the peptide, but clinical experience is still lacking (17).

Administration of PTH to animal models has been linked to the development of certain tumors as osteosarcoma (18). A more physiological administration such as the one we used, as the number of pulses delivered per hour in our patient is about the same of normal parathyroid glands (10, 11), would probably be safer in this respect; moreover, the total dose given per day in our patient is similar to the one given in the study published by Winer, in which the patients were followed over 3 years without safety problems. A careful follow-up in this regard is provided. In our case no evidence of bone problems were detected after 22 months of teriparatide pump treatment, and additionnally, as expected, a recovery of bone mass was observed. We believe that this modality of teriparatide delivery maybe indeed more safe and may allow a potential lower risk of bone problems at long term because of the intermittent nature of the peptide infusion in contrast to a continuous infusion (19, 20), although unfortunately we cannot prove that the plasmatic profile of rhPTH in our patient would in fact reproduce the pulse design of the infusion system, as we did not measured circulating PTH in a way to assess it.
In summary, we report the first case of long term treatment of vitamin D unresponsive hypoparathyroid patient with teriparatide multipulse pump as the only effective therapeutic modality. In our patient this kind of therapy was shown to be safe, as stable profiles of calcium and magnesium were achieved along 1 year, and should therefore be considered in selected cases such as the one described.

Declaration of interest
The authors declare that there is no conflict of interest that would prejudice their impartiality.

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