Persistent hypercalcemia with similar familial hypocalciuric hypercalcemia features: a case report and literature review

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Abstract

Background: Primary hyperparathyroidism (PHPT) and familial hypocalciuric hypercalcemia (FHH) are the most important differential diagnosis of parathyroid hormone (PTH)-dependent hypercalcemia. The clinical features of FHH and PHPT can overlap in some cases. Therefore, these two diseases must be differentiated to prevent unnecessary parathyroidectomy. Here, we present a case that was not entirely matched with any of the known differential diagnoses of hypercalcemia.

Case presentation: A 19-year-old girl with no history of any disease presented with persistent hypercalcemia without any specific musculoskeletal complaint. We found persistent hypercalcemia in her routine laboratory data from 3 years ago; while no data was available during the childhood period. Her dietary calcium intake was normal. She did not mention any history of renal stone, bone fracture as well as family history of hypercalcemia. Biochemical features showed normal values of serum creatinine, high normal serum calcium (range, 10.3–11.3 mg/dL; normal range: 8.8–10.4), and non-suppressed PTH levels (range, 37.2–58.1 pg/mL; normal range: 10–65). Serum 25 OH vitamin D level at the first visit was 16.1 ng/mL that treated by vitamin D supplementation. Since then, all 25 OH vitamin D levels were in the acceptable range. After correction of vitamin D deficiency during the follow-up period the calcium creatinine clearance ratio(s) (CCCR) were calculated in the range of 0.009 to 0.014 (means below 1%). The clinical and laboratory data indicate more FHH rather than PHPT. Genetic studies were negative for the common genes associated with FHH (CASR, GNA11, and AP2S1 genes) and multiple endocrine neoplasia type1 (MEN1). On the other hand, no evidence of autoimmunity was found in her to support an autoimmune FHH-like syndrome. Hence, the case did not match completely to any diagnosis of FHH and PHPT, so we decided to follow her.

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Conclusion: We presented a patient with FHH phenotype whose common genetic tests were negative. Further research is needed to ascertain other causes leading to similar manifestations.

Keywords: Familial Hypocalciuric hypercalcemia, Primary hyperparathyroidism, Case report, Endocrinology and metabolism, Persistent hypercalcemia

Background
Hypercalcemia has an expanded differential diagnosis of approximately over 25 diseases. Mild-to-moderate hypercalcemia with non-suppressed or high parathyroid hormone (PTH) levels present in patients with primary hyperparathyroidism (PHPT), typically [1]. PHPT is a common endocrine disorder among the general population especially in middle age people and the female gender. Factors favoring PHPT include a history of renal calculi, an asymptomatic patient with a stable prolonged mild hypercalcemia, and rarely the induction of hypercalcemia by thiazides [2]. Familial hypocalciuric hypercalcemia (FHH) and the use of certain medications, such as lithium, could mimic PHPT presentations and should be considered in the differential diagnoses of hypercalcemia with non-suppressed PTH [3].

FHH, also called familial benign hypercalcemia, was initially introduced as a variant of PHPT [4]. It is a rare autosomal-dominant disorder with very low urinary calcium excretion and mild hypercalcemia with high normal or slightly elevated PTH [5]. FHH is a life-long condition that is usually caused by one of many heterozygous inactivating mutations in the calcium-sensing receptor (CASR) gene, which could up-regulate the set point of parathyroid cells. When the CASR receptor is inactivated, PTH is not suppressed despite its relatively high calcium, which makes FHH similar to PHPT. In PHPT, although the renal reabsorption of calcium is higher than normal due to the high PTH level, hypercalciuria still occurs [6]. However, in FHH due to CASR inactivation, a general calcium hyposensitivity occurs which results in more tubular calcium reabsorption. So, calcium/creatinine clearance ratio (CCCR = [24 h urine calcium/ total plasma calcium] × [plasma creatinine /24 h urine creatinine]) is lower in FHH compared to PHPT [7].

In addition to familial forms of FHH which existed from the beginning of life, there are other acquired and sporadic forms with unrevealing family screening: including de novo CASR gene mutations, and CASR auto-antibodies [7]. Patients with CASR autoantibodies mimic the hereditary FHH in biochemical phenotype and could develop FHH-like syndrome. Auto-immune hypocalciuric hypercalcemia is different in pathogenesis from the one caused by inactivating mutations of the CASR [4]. So, a combination of clinical suspicion, biochemical testing, and genetic analysis is needed to distinguish FHH from PHPT. This case is challenging because findings in this case report were not entirely consistent with any of FHH, PHPT, and FHH-like syndrome.

Case presentation
A 19-year-old girl was referred to our endocrinology and metabolism department with elevated plasma calcium levels found 3 years ago and we had no documentation about it before that. She denied any abnormal complaints including specific musculoskeletal pain, polyuria, polydipsia, fatigue, anxiety, sleep disturbance, loss of concentration and cognitive dysfunction, decreased memory, headache and irritability. Moreover, she had no anorexia, constipation, findings related to pancreatitis, peptic ulcer and abdominal pain. She has had regular menstrual periods since puberty. She did not mention the history of drug use. She had no history of renal stone, bone fracture and cardiovascular disease as well as family history of parathyroidectomy. Her parents had normal calcium levels. Physical examinations were unremarkable. Vital signs and body mass index (BMI: 21 kg/m2) were normal and she never had high blood pressure or orthostatic changes. She had a perfectly normal face. She was quite alert and did not show any signs of depression. No skin lesions were observed. No evidence of muscle weakness, skeletal structural disorders or chondrocalcinosis were observed.

Her serum calcium levels ranged from 10.3 to 11.3 mg/dL (normal, 8.8–10.4, with serum albumin between 4 and 4.2 mg/dL in all assays). Inorganic phosphorus levels ranged from 2.8 to 3.4 mg/dL (normal, 2.5–4.5). Serum magnesium levels ranged from 1.8 to 2.5 mEq/L (normal, 1.7–2.2). Intact PTH levels ranged from 37.2 to 58.1 pg/mL (normal, 10–65) and urine CCCR ranged from 0.009 to 0.014 during these 3 years (Table 1). Serum 25 OH vitamin D level at the first visit was 16.1 ng/mL that treated by vitamin D supplementation. Since then, all 25 OH vitamin D levels were in the acceptable range. Thyroid function tests did not show sub-clinical hyperthyroidism. Other laboratory findings were unremarkable (Table 2).

Parathyroid SPECT- CT-MIBI revealed no evidence of parathyroid adenoma. Thyroid, neck, and renal ultrasonography evaluation were normal with no nephrolithiasis or other abnormal findings. On the lateral skull X-ray and both hands AP X-ray, there were no abnormal findings associated with chronic inappropriate high PTH.
level; including subperiosteal bone resorption, brown tumors, bone cysts, or sclerosis.

Genomic DNA was extracted from the submitted specimen and the Lon AmpliSeq 114 Exome RDY Library Preparation Kit was used to amplify some targeted regions of gene panel. Among 19,073 known genes sequenced, variant analysis firstly focused on three genes associated with FHH, including \textit{CASR}, \textit{GNA11}, and \textit{AP2S1}. The further variant analysis was carried out on \textit{MEN1}, \textit{RET}, \textit{CDKN1B}, \textit{GCM2}, \textit{CDC73}, and other genes associated with PHPT induced hypercalcemia. None of the genetic studies in our case were informative.

Although bone mineral density (BMD) measurement is not appropriate at a young age, it was eventually done, because it may help to differentiate FHH from PHPT [8, 9]. BMD of the femoral neck, lumbar spines, and wrist showed normal values (Table 3).

### Discussion and conclusion

We have described a young lady with persistent mild hypercalcemia, high normal PTH, CCCR ≤0.01, without any clinical findings of PHPT, and any related family history of hypercalcemia and nephrolithiasis. Genetic studies were negative for known FHH mutations and the \textit{MEN1} gene. No evidence of the presence of autoimmune FHH like syndrome was detected.

FHH and PHPT are the two most important differential diagnoses in this patient. A two-step diagnostic procedure was used for distinguishing between FHH and PHPT. The first step was the CCCR from a 24-h urine

### Table 1 Data of Ca, PTH, Vit D and CCCR in the case. (July 2018–January 2021)

|                | Serum calcium (mg/dL) | Serum phosphorus (mg/dL) | 25(OH) Vitamin D (ng/mL) | PTH levels (pg/mL) | CCCR  |
|----------------|-----------------------|--------------------------|--------------------------|-------------------|-------|
| July 2018      | 10.8                  | 2.8                      | –                        | 58.1              | –     |
| August 2018    | 10.8                  | 3                        | 16.1                     | 50.8              | 0.009 |
| November 2018  | 10.5                  | 3.3                      | 33.5                     | 40.9              | –     |
| December 2018  | 10.7                  | 3.3                      | 24.7                     | 47.7              | 0.014 |
| July 2019      | 10.4                  | 3.2                      | 33.9                     | –                 | –     |
| May 2020       | 10.8                  | 3.4                      | –                        | 37.2              | 0.009 |
| June 2020      | 10.3                  | 3.2                      | 32.5                     | 40.9              | –     |
| June 2020      | 10.6                  | 3.3                      | –                        | 37.4              | 0.011 |
| October 2020   | 11.3                  | 3.3                      | 50.7                     | –                 | –     |
| November 2020  | 11                    | –                        | –                        | –                 | –     |
| January 2021   | –                     | –                        | 45.6                     | –                 | –     |

\textit{PTH} parathyroid hormone, \textit{CCCR} calcium creatinine clearance ratio.

### Table 2 Other laboratory findings

| Variables                | Value  | Normal range     |
|--------------------------|--------|------------------|
| White Blood Cells count (WBC) mCL | 5600   | 4.4–11.3 (*1000) |
| Hemoglobin concentration (Hb) g/dL | 14.5   | 12.3–15.3        |
| Platelet count (Plt) mCL   | 253,000| 336,000 (*1000)  |
| Blood Urea (mg/dL)         | 25     | 13–51            |
| Serum Creatinine (mg/dL)   | 0.64   | 0.7–1.3          |
| Serum Magnesium (Mg) (mEq/L) | 2.1    | 1.7–2.2          |
| Serum Alkaline Phosphatase (U/L) | 50     | 20–140           |
| Serum thyroid-stimulating hormone (TSH) (mIU/L) | 1.16   | 0.3–4.2          |
| Total T4 (µg/dL)           | 7.0    | 5.0–12.0         |
| Serum Albumin level (g/dL) | 4.1    | 3.5–5.2          |
| Rheumatoid Factor (IU/mL)  | 1.4    | < 14             |
| Immunoglobulin G4 (mg/dL)  | 3.65   | 3–135            |
| Anti-tissue transglutaminase (TTG)-IgA Elisa- Ru/ml | 0.1    | < 3              |
| Anti-thyroid peroxidase (TPO) IU/mL | 0.20  | < 35             |
| Immunoglobulin A (mg/dL)   | 135    | 70–400           |
| Anti-Nuclear Antibodies-IgG (Dilute) | Negative |
sample. A ratio of less than 0.01 suggests FHH, and a ratio of 0.02 or higher could indicate PHPT. For the second step, all patients with a CCCR of 0.020 or less are tested for mutations in the CASR gene. The diagnostic sensitivity of these two steps is 98% [7]. The majority (80%) of patients with FHH have a CCCR of < 0.01; nevertheless, almost 20% of patients with PHPT have a CCCR between 0.01 and 0.02 which can overlap with FHH individuals. A low CCCR may be seen in PHPT with renal insufficiency or in those with severe vitamin D deficiency; the patients have a slightly elevated or high normal serum calcium and a low urinary excretion of calcium [2], while our patient had low CCCR without renal failure and vitamin D deficiency. On the other hand, a large retrospective cohort study was done on 1000 patients with definitive diagnosis of PHPT based on surgery, which showed CCCR is valuable only in 37% patients to correctly differentiate FHH from mild PHPT. In 63% patients with surgically verified PHPT, CCCR was below 2% where both FHH and primary hyperparathyroidism are overlapping [10].

Genetic testing of the CASR gene is helpful in situations in which there is an overlap in the phenotype of PHPT and FHH [11, 12]. On the other hand, PHPT is more prevalent among individuals over 50 years, so young patients with PHPT may be suspected of multiple endocrine neoplasia (MEN) [13], which is why we tested the MEN1 gene mutation for our patient.

FHH is not always a familial disease, despite its name; hence, the lack of evidence in family screening does not rule out FHH. It’s important to note that approximately one-third of individuals with FHH have no mutations within the CASR gene [8]. It may be due to mutations located outside the coding exons 2–7 of the CASR or mutations which were not detected by PCR and sequencing [7].

Kifor et al. reported several patients with PTH-dependent hypercalcemia with presentations that seemed to be caused by autoimmunity (e.g., non-tropical sprue or Hashimoto’s thyroiditis) [4]. Another acquired hypocalciuric hypercalcemia has been reported in a woman with a severe autoimmune disorder because of IgG4 autoantibodies [14]. Although the presence of CASR antibodies is needed for confirmation of the diagnosis of FHH-like syndrome, other obvious autoimmune diseases were also found [4]. In this case, it was not available to check CASR antibodies. However, we can claim that FHH-like syndrome was very unlikely, due to a lack of clinical and laboratory evidence of other associated autoimmune diseases, including non-tropical sprue, Hashimoto’s thyroiditis, Ig G4 related disease, systemic lupus erythematosus, and rheumatoid arthritis.

In our case, the first rank diagnosis was FHH because of young age at the time of hypercalcemia detection, lack of history of fractures, nephrocalcinosis, nephrolithiasis, radiologic suspicion for bone resorption, and a normal BMD as well as CCCR≤0.01 [15]; although these presentations may be present in nearly 20% of patients with PHPT [16]. Also, the absence of autoantibody markers and lack of other autoimmune diseases could rule out FHH-like syndrome. On the other hand, FHH rarity, absence of a family history of FHH, and negative results of relevant genetic tests could support PHPT as the second rank in diagnosis.

Finally, we decided to follow the case rather than referring her to parathyroidectomy, focusing on changes in serum calcium, PTH, and CCCR levels. Also, we plan to repeat BMD at an older age because bone complications could occur in PHPT patients over time, unlike most patients with FHH [17, 18].

Our study has some limitations, including the fact that the CASR autoantibody was not checked in the patient. Also, we did not have any related tests before this period. Finally, although we checked her parent’s laboratory data, we could not access to the exact status of the calcium levels in her brother. Of course, the patient’s family mentioned that their son was healthy.

In conclusion, we have described a young lady with FHH phenotype (e.g., persistent mild hypercalcemia, high normal PTH, CCCR ≤0.01), with negative genetic tests. The findings in this case report were not entirely consistent with any of FHH, PHPT, and FHH-like syndrome. Further researches are needed to ascertain other causes leading to similar manifestations.

**Abbreviations**
FHH: Familial Hypocalciuric Hypercalcemia; PHPT: Primary Hyperparathyroidism; CCCR: Calcium creatinine Clearance Ratio; CaSR: Calcium Sensing Receptor

| Table 3 Dual-energy x-ray Absorptiometry (DEXA) |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Region          | Area (cm²)      | BMC (g)         | BMD (g/cm²)     | T-score         | Z-score         |
| Total Lumbar spine | 67.56            | 70.08           | 1.037           | −0.1            | 0.2             |
| Total Hip       | 31.73            | 30.06           | 0.948           | 0.0             | 0.0             |
| Ultra-distal Radius | 15.30            | 8.02            | 0.524           | −1.0            | −1.0            |
| 10-Year Fracture Risk |
| Major Osteoporotic Fracture | 3.0%        |
| Hip Fracture    | 0.1%             |
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Authors’ contributions
MZ and RHA: wrote the manuscript, MR: contributed to the patient genetic studies, AA and FH: reviewed and edited manuscript. All authors read and approved the final manuscript.

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Availability of data and materials
All data used during the current study are available from the corresponding author on reasonable request.

Declarations

Ethics approval and consent to participate
Not Applicable.

Consent for publication
Written informed consent was obtained from the patient for publication of this case report and any accompanying images. A copy of the written consent is available for review by the Series Editor of this journal.

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

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References
1. Insogna KL. Primary hyperparathyroidism. N Engl J Med. 2018;379(11):1050–9.
2. Lafferty FW. Differential diagnosis of hypercalcemia. J Bone Miner Res. 1991; 6(5):551–9.
3. Pallan S, Rahman MO, Khan AA. Diagnosis and management of primary hyperparathyroidism. Brmj. 2012;344:e1013.
4. Kifor O, Moore FD Jr, Delaney M, Garber J, Hendy GN, Butters R, et al. A syndrome of hypocalciuric hypercalcemia caused by autoantibodies directed at the calcium-sensing receptor. J Clin Endocrinol Metab. 2003;88(1):60–72.
5. Khan AA, Hanley DA, Rizzoli R, Bollerslev J, Young JE, Rejmark L, et al. Primary hyperparathyroidism: review and recommendations on evaluation, diagnosis, and management. A Canadian and international consensus. Osteoporois Int. 2017;28(11):1–19.
6. Phelps KR, Stote KS, Mason D. Tubular calcium reabsorption and other aspects of calcium homeostasis in primary and secondary hyperparathyroidism. Clin Nephrol. 2014;82(2):83–91.
7. Christensen SE, Nissen PH, Vestergaard P, Moskelide L. Familial hypocalciuric hypercalcaemia: a review. Curr Opin Endocrinol Diabetes Obesity. 2011; 18(6):359–70.
8. Shinal M, Dahir K, Broome J. Differentiating familial hypocalciuric hypercalcaemia from primary hyperparathyroidism. Endocr Pract. 2013;19(4):697–702.
9. Isakeni T, Nielsen CS, Christensen SE, Nissen PH, Heikendoff L, Moskelide L. Forearm bone mineral density in familial hypocalciuric hypercalcaemia and primary hyperparathyroidism: a comparative study. Calcif Tissue Int. 2011; 89(4):285.
10. Moore EC, Berber E, Jin J, Krishnamurthy V, Shin J, Spertus A. Calcium-creatinine clearance ratio is not helpful in differentiating primary hyperparathyroidism from familial hypercalciemic Hypocalciuria: a study of 1,000 patients. Endocr Pract. 2018;24(11):988–94.
11. Kearns AE, Wermers RA. Familial Hypocalciuric Hypercalcaemia. Hyperparathyroidism edn: Springer. 2016:105–10.
12. Lee JY, Shoback DM. Familial hypocalciuric hypercalcaemia and related disorders. Best Pract Res Clin Endocrinol Metab. 2018;32(5):609–19.
13. Lassen T, Friis-Hansen L, Rasmussen A, Knigge U, Feldt-Rasmussen U. Primary hyperparathyroidism in young people: When should we perform genetic testing for multiple endocrine neoplasia 1 (MEN-1)? J Clin Endocrinol Metab. 2014;99(11):3983–7.
14. Pallais JC, Kifor O, Chen YB, Slovik D, Brown EM. Acquired hypocalciuric hypercalcemia due to autoantibodies against the calcium-sensing receptor. N Engl J Med. 2004;351(4):362–9.
15. Walker MD, Silverberg SJ. Primary hyperparathyroidism. Nat Rev Endocrinol. 2018;14(2):115.
16. Silverberg SJ, Clarke BL, Peacock M, Bandeira F, Boutroy S, Cusano NE, et al. Current issues in the presentation of asymptomatic primary hyperparathyroidism: proceedings of the fourth international workshop. J Clin Endocrinol Metab. 2014;99(10):3580–94.
17. LAW WM Jr, HW WAHNER, HEATH H III. Bone mineral density and skeletal fractures in familial benign hypercalcemia (hypocalciuric hypercalcemia). Mayo Clin Proc. 1984; Elsevier;1984:811–15.
18. Christensen SE, Nissen PH, Vestergaard P, Heikendoff L, Rejmark L, Brixen K, et al. Skeletal consequences of familial hypocalciuric hypercalcaemia vs. primary hyperparathyroidism. Clin Endocrinol. 2009;71(6):798–807.

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