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Case report: A 10-year-old girl with primary hypoparathyroidism and systemic lupus
 erythematosus

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#### 43 Abstract

Hypoparathyroidism is a rare disease in children that occurs as a result of autoimmune 44 destruction of the parathyroid glands, a defect in parathyroid gland development or secondary 45 to physical parathyroid gland disturbance. Typical symptoms of hypoparathyroidism present as 46 hypocalcaemia and hyperphosphatemia due to decreased parathyroid hormone secretion and 47 may lead to nerve and muscles disturbances resulting in clinical manifestation of tetany, 48 arrhythmias and epilepsy. Currently, there is no conventional hormone replacement treatment 49 for hypoparathyroidism and therapeutic approaches include normalising mineral levels using 50 an oral calcium supplement and active forms of vitamin D. We present the case of a 10-year-51 old girl with primary hypoparathyroidism who had no prior history of autoimmune disorders, 52 53 but who subsequently developed systemic lupus erythematosus.

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Keywords: primary hypoparathyroidism, hypocalcaemia, hyperphosphatemia, parathyroid
hormone, systemic lupus erythematosus, autoimmune polyglandular syndrome.

#### 57 Background

Hypoparathyroidism is a rare endocrine disease in children caused by parathyroid hormone 58 (PTH) deficiency or resistance to PTH [1]. Primary hypoparathyroidism may occur as a result 59 60 of either autoimmune destruction of the parathyroid glands, a congenital defect of parathyroid gland development as in DiGeorge 22q11.2 deletion syndrome, mutations in the calcium-61 sensing receptor gene, damage to the parathyroids during surgery, most commonly in the 62 63 course of thyroidectomy, or following radiation treatment of the neck [2-4]. The typical manifestations of the disease present when all four parathyroid glands are affected and include 64 hypocalcaemia, hyperphosphatemia and low serum PTH concentrations. The serum mineral 65 66 abnormalities affect nerve and muscle functions leading to tetany, arrhythmias and seizures. There is no conventional hormone replacement therapy and the management of 67 hypocalcaemia and hyperphosphatemia involves an oral calcium supplement and active forms 68 of vitamin D [3]. 69 70 Here, we present the case of a 10-year-old girl with primary hypoparathyroidism who had no 71 prior history of autoimmune disorders, but who developed systemic lupus erythematosus (SLE) a few months after her diagnosis of hypoparathyroidism. To date, there are only a few 72

reported cases of the coexistence of hypoparathyroidism and SLE [5].

#### 74 Case presentation

A 10-year-old Polish female initially presented to the Department of Paediatric Neurology 75 and Rehabilitation with convulsions. Following a confirmation of hypocalcaemia she was 76 77 referred to the Department of Paediatrics, Endocrinology, Diabetology with Cardiology with suspected hypoparathyroidism. The patient had no history of mucocutaneous candidiasis and 78 no history suggesting an endocrine or autoimmune disorder in her family. Her mental 79 development was normal. Her weight was 35 kg (6th percentile of 10-year-old females) and 80 her height was 141 cm (50th percentile of 10-year-old females). She was in early puberty at 81 stage 2 on the Tanner scale, based on breast development. 82

83 The patient was admitted to our department severely unwell with drowsiness and confusion.

84 The physical examination revealed a whole body rash that was attributed to an allergic

reaction to oxcarbazepine that was being used for the treatment of convulsions. She had no
features of ectodermal dystrophy or any signs of primary adrenal insufficiency.

87 Hypoparathyroidism was diagnosed based on very low PTH levels at <3 pg/mL (reference

range, 10-69 pg/mL), low total serum calcium levels at 0.80 mmol/L (reference range, 2.19-

2.69 mmol/L) (Figure 1) and high plasma phosphate levels at 11.8 mg/dL (reference range,

90 3.4-6.2 mg/dL) (Figure 1). Alkaline phosphatase levels and serum magnesium concentrations

91 were normal. Total serum vitamin D was low at 35 nmol/L (reference range, 50-150 nmol/L).

92 Urinary calcium levels were lower than normal at admission.

Hormone analysis showed normal thyroid function (thyroid-stimulating hormone at 1.76

94 IU/L, reference range, 0.28-4.3 IU/L; free T4 at 23 pmol/L, reference range, 14-22 pmol/L).

Adrenal function tests were within the normal range (cortisol at 347.3 nmol/L;

adrenocorticotropic hormone at 2.8 pmol/L; adequate peak cortisol level of 749.5 nmol/L

97 after 250 µg of tetracosactin by intravenous injection). Markers for specific for autoimmune

98 diseases including antibodies against interferon (IFN)-ω, 21-hydroxylase, 17-α-hydroxylase,

99 the calcium-sensing receptor, the acetylcholine receptor, and tissue transglutaminase were not 100 detected (Table 1). Autoantibodies associated with autoimmune thyroid disease and type 1 101 diabetes were also undetectable (Table 1). In blood morphology analysis, there were 102 periodical findings indicating anaemia, leucopaenia and hypoalbuminaemia in biochemical 103 tests.

104 Ultrasonography revealed a slightly enlarged spleen without any features of nephrocalcinosis and no parathyroid pathology. Echocardiography (ECG) did not reveal any heart defect. 105 106 However, effusion in the pericardium was observed coinciding with the fever and elevated inflammatory markers. Long QTc (over 0.5 seconds) was recorded in the ECG. A bone 107 108 densitometry scan showed normal values for bone mass density of the spine L2-L4 (Z-score +0.5) and for total body less head (Z-score +0.1). There was no ectopic calcification observed 109 in magnetic resonance imaging (MRI) of the head. A positron emission tomography-magnetic 110 resonance imaging (PET-MRI) scan of the whole body did not reveal any other pathologies. 111 Electroencephalography was abnormal with pathological changes localised in temporal areas 112 113 on both sides during activity and generalised paroxysmal patterns during sleep. However, 114 there were no more epileptic seizures during clinical observation.

115 The patient was started on intravenous calcium gluconate at 5.5 g elemental calcium/day

along with oral calcium carbonate (0.8 g elemental calcium/day), vitamin D (2000 IU/day)

and the synthetic precursor of the active form of vitamin D3, alfacalcidol  $(2 \mu g/day)$ .

118 Hyperphosphatemia was initially treated with a diet low in phosphates. She was also treated

119 with valproic acid for epilepsy, metoprolol for long QTc, and hydrocortisone for the first nine

120 days to reduce her allergy symptoms. The patient's condition improved and her total serum

121 calcium levels increased, although they remained below the normal range (Figure 1).

122 After fourteen days of hospitalisation, the patient developed a fever with pneumonia and

123 pericardial effusion together with increased inflammatory markers in laboratory tests. C-

reactive protein increased significantly from 30.2 to 1497 nmol/L (reference range, 0-47 124 125 nmol/L). She was treated with antibiotics, although the blood cultures returned negative results. There was no evidence of either viral infections including HIV, B19 parvovirus, 126 127 influenza or of zoonoses or tuberculosis. Furthermore, neoplastic disease was excluded following a bone marrow biopsy. During the course of a chest infection, her total serum 128 calcium levels dropped to below 2 mmol/L. After three weeks of hospitalisation and with a 129 130 high plasma phosphate level of 9.20 mg/dL, she was given sevelamer (initial dose 2.4 g/day), a phosphate-binding medication, to reduce hyperphosphatemia (Figure 1). Finally, the patient 131 was discharged on regular oral calcium carbonate (3.6 g elemental calcium/day), sevelamer 132 133 (4.8 g/day) and alphacalcidol (1  $\mu$ g/day).

Taken together, the clinical symptoms, laboratory test results and the criteria of the American 134 College of Rheumatology for SLE, in particular, epilepsy, leucopaenia, pericardial effusion 135 and anti-nuclear antibodies, the patient was diagnosed with SLE. She was treated 136 subsequently with chloroquine phosphate and oral prednisone (5 mg/day). This treatment 137 138 unexpectedly improved the patient's calcium and phosphate levels (Figure 1) and allowed a reduction in the doses of calcium, sevelamer and vitamin D. During monthly follow up visits, 139 the patient's serum calcium and phosphate levels continued to normalise (Figure 1), albeit 140 with episodes of hypercalcaemia. 141

142 At four months follow-up, the patient was re-evaluated for anti-IFN- $\omega$  and anti-21-hydroylase

antibodies. She was found positive for IFN- $\omega$  antibodies (antibody index 21.4, reference range

144 <5.0) (Table 1). To further investigate the cause of the hypoparathyroidism, we evaluated

145 genetic mutations that are typically connected with the disease. Mutational analysis of genes

146 encoding the calcium-sensing receptor (*CASR*), guanine nucleotide-binding protein subunit

alpha-11 (GNA11), PTH (PTH), glial cells missing transcription factor 2 (GCM2),

autoimmune regulator (AIRE) and GATA binding protein 3 (GATA3) were all negative. In

- addition, whole exome sequence variants in established disease genes and candidate variants
- in genes not yet associated with disease were negative. However, it revealed a heterozygous
- microdeletion in chromosome 15q11.2, inherited from the father and not reported previously
- to be in association with hypoparathyroidism.

#### 153 Discussion

The combination of hypoparathyroidism and SLE is extremely rare therefore the underlyingcauses and significance of these diseases occurring in conjunction is intriguing.

156 Hypoparathyroidism can be caused by several factors including autoimmunity affecting the parathyroid glands, damage to parathyroid glands and genetic mutations such as DiGeorge 157 syndrome or calcium-sensing receptor-activating mutations [6, 7]. In this case, secondary 158 hypoparathyroidism could immediately be discounted as the patient had had no surgery or 159 trauma to the neck or parathyroid glands. The patient did not display symptoms associated 160 with DiGeorge syndrome nor with calcium-sensing receptor-activating mutations. This was 161 162 supported by mutational analysis, which did not reveal mutations in the CASR gene. In view of the above and the association with SLE, it was expected that hypoparathyroidism in this 163 164 patient had an underlying autoimmune origin.

An autoimmune aetiology of the hypoparathyroidism in our patient was considered after her 165 166 hypocalcaemia improved with glucocorticoid treatment. Autoimmune hypoparathyroidism is 167 a major component of autoimmune polyglandular syndrome type 1 (APS1) along with chronic mucocutaneous candidiasis and primary adrenal insufficiency (Addison's disease) [8-12]. 168 APS1 commonly manifests in childhood and our patient presents with one of the major 169 components. However, she does not fully adhere to an APS1 diagnosis as she does not display 170 other symptoms currently. In addition, AIRE gene analysis in our patient did not identify 171 typical known genetic mutations that cause APS1. However, there may be AIRE gene 172 mutations not detected by our analysis and further in-depth mutational analysis is planned. 173 IFN- $\omega$  antibodies are highly specific for APS1 [13,14]. Interestingly, our patient initially 174 175 tested negative for IFN-ω antibodies, but proved positive on re-testing several months after her initial hypoparathyroidism diagnosis. This might suggest that she is progressing towards 176 APS1 and may in time develop another major APS1 autoimmunity. Therefore, monitoring for 177

178	chronic mucocutaneous candidiasis and Addison's disease would seem appropriate in the
179	future. Considering an alternative APS diagnosis, there have been occasional reports of
180	hypoparathyroidism occurring in other APS types [10, 11]. The patient does not conform to
181	an APS2 (Addison's disease, thyroid disease and/or type 1 diabetes) or APS3 (thyroid disease
182	plus one other non-Addison's disease autoimmunity) diagnosis. Thus APS4 (two or more
183	autoimmune diseases not in APS types 1, 2 or 3) was considered. Our patient currently has
184	two autoimmune diseases thus fulfils the criteria for APS4, however, as she matures, she may
185	develop additional autoimmune endocrinopathies and her diagnosis might alter accordingly.
186	In summary, our case report describes the presentation and diagnosis of a 10-year-old girl
187	with hypoparathyroidism who subsequently developed SLE. We suggest that an autoimmune
188	mechanism, potentially associated with an underlying genetic predisposition, is responsible
189	for the disease combination seen in our patient.

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## 234 **Figure legend**

236	Figure 1.	The patient	's total serum	a calcium and	plasma j	phosphate	concentrations	over time.
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- 237 Shading indicates the reference ranges (total serum calcium, 2.19-2.69 mmol/L; plasma
- phosphate, 3.4-6.2 mg/dL). The patient's blood calcium levels were initially low
- 239 (hypocalcaemia) and plasma phosphate levels were initially high (hyperphosphatemia), but
- both normalised following treatment with calcium and prednisone. The start-points of
- treatment with sevelamer (4.8 g/day) and prednisone (5 mg/day) are indicated.

# **Table 1.** Autoantibody screening of the patient

Antibody target	Positive/negative	Result	Reference range
Diabetes screening (Zinc transporter 8; glutamic acid decarboxylase; islet antigen IA2)	Negative	6.7 (index)	≤30 (index)
Interferon-ω (07.03.2018)	Negative	<5 (index)	<5 (index)
Interferon-ω (27.06.2018)	Positive	21.4 (index)	<5 (index)
21-hydroxylase	Negative	<0.4 u/mL	≤0.4 u/mL
Calcium-sensing receptor	Negative	0.52 (index)	1.37 (index)
Acetylcholine receptor	Negative	<0.45 nmol/L	<0.45 nmol/L
17-α-hydroxylase	Negative	<1 U/mL	<1 U/mL
Tissue transglutaminase	Negative	0.4 U/mL	<4 U/mL
Thyroid peroxidase	Negative	5.9 IU/mL	<34 IU/mL
Thyroglobulin	Negative	13.6 IU/mL	<115 IU/mL
Thyroid-stimulating hormone receptor	Negative	0.35 IU/L	<1.75 IU/L

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246	1. Hypoparathyroidism in children might be caused by an autoimmune process, damage to the
247	parathyroid glands or a genetic disorder.

248 2. The typical manifestations of the disease are hypocalcaemia and hyperphosphataemia that

affect nerve and muscle functions leading to tetany, arrhythmias and seizures.

250 3. Autoimmune hypoparathyroidism is often associated with other autoimmune diseases to

constitute autoimmune polyglandular syndromes, typically APS1.

- 4. The management of hypocalcaemia and hyperphosphatemia following decreased PTH
- concentrations involves supplementation with calcium and active forms of vitamin D.

254

## 255 What is new?

- The coexistence of hypoparathyroidism and SLE is extremely rare in children and is nottypical for APS1.

- The normalisation of calcium and phosphate blood concentrations after glucocorticoid
treatment for SLE, reversed the symptoms of hypoparathyroidism indicating a possible
autoimmune basis for the disease.

