A Patient with Proopiomelanocortin Deficiency: An Increasingly Important Diagnosis to Make

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What is already known on this topic?
Proopiomelanocortin (POMC) deficiency is an extremely rare disorder characterized by early-onset obesity, adrenal insufficiency, red hair and decreased skin pigmentation. Hyperphagia, cholestasis, exponential weight gain and adrenal insufficiency are typically observed during the first months of life. In some children, the diagnosis may only be established later.

What this study adds?
This study presents clinical and molecular features of a child with POMC deficiency. We also provide a brief summary of the clinical and genetic features of POMC deficiency based on previously published patient reports and describe how these are providing insight into the role of POMC in the regulation of human metabolism.

Abstract
Proopiomelanocortin (POMC) deficiency is a rare monogenic disorder with early-onset obesity. Investigation of this entity have increased our insight into the important role of the leptin-melanocortin pathway in energy balance. Here, we present a patient with POMC deficiency due to a homozygous c.206delC mutation in the POMC gene. We discuss the pathogenesis of this condition with emphasis on the crosstalk between hypothalamic and peripheral signals in the development of obesity and the POMC-melanocortin 4 receptors system as a target for therapeutic intervention.

Keywords: Obesity, melanocortin 4 receptors, paediatric obesity, proopiomelanocortin deficiency

Introduction
Proopiomelanocortin (POMC) is a 241-amino acid polypeptide that is cleaved via prohormone convertase (PC) to produce the peptides γ, β, α-melanocyte stimulating hormone (MSH), adrenocorticotropin hormone (ACTH), γ, β-lipotrophin and endorphins (1). These peptides stimulate five different melanocortin receptors (MCR) with varying affinity and specificity. Cortisol secretion is regulated through MC2R in the adrenal gland while MC1R regulates skin pigmentation. MC3R and MC4R regulate body weight.

Congenital POMC deficiency develops due to genetic defects in the POMC gene located at Chr.2p23.3. This disorder is characterized by early-onset obesity, adrenal insufficiency, red hair and decreased skin pigmentation (1,2,3,4). Obesity develops as a result of inadequate production of α- and β-MSH, which normally activate the MC3R in the arcuate nucleus and the MC4R in the paraventricular nucleus and...
antagonize the action of agouti-related peptide (AgRP) (4). The hypocortisolaemia and hypopigmentation are due to inadequate stimulation of MC2R and MC1R by POMC-derived peptides in the adrenal gland and skin, respectively. POMC deficiency is rare, but has increased our insight into the important role of the leptin-melanocortin pathway in energy balance.

Here, we present the clinical characteristics of a patient with POMC deficiency due to a mutation in the POMC gene, hoping to contribute to a better understanding of the leptin-melanocortin pathway and to introduce possible treatment options.

**Case Report**

This female patient presented at age 2.5 months with restlessness, cyanosis, and spasms. She was found to be hypoglycaemic with a blood glucose of 31 mg/dL. She was born at 39 weeks gestation with a birth weight of 3000 grams and had no problems during the prenatal or early postnatal period. Her mother and father were not related and she had a healthy brother aged five years. There was no history of relevant disease in the family. Examination findings at presentation revealed growth failure (body weight: 3700 g, 3rd percentile; height: 51 cm, <3rd percentile; head circumference: 35 cm, <3rd percentile) (Figure 1A), red eyebrows and hair and normal female genitalia. Results of further laboratory investigations confirmed the hypoglycaemia (blood glucose: 19 mg/dL) and revealed mild hyponatraemia with a sodium of 132 mmol/L (135-143), accompanied by a potassium of 4.8 mmol/L (3.1-5.5), mildly elevated aspartate transaminase: 123 U/L (<36) and creatinine kinase: 419 U/L (34-204).

Concomitant with the hypoglycaemic state (34 mg/dL), serum and urinary ketones were low and there was no evidence of hyperinsulinaemia or any other metabolic cause for the hypoglycemia (serum insulin: 0.11 U/L, C-peptide: <0.05 nmol/L, lactate: 2.37 mmol/L (0.49-2.19), ammonium: 93.5 μmol/L (13.5-42.8), urinary and blood amino acids and organic acid profile normal). Total/indirect bilirubin levels were 3.7/2.4 mg/dL. However, the child was hypocortisolaemic (cortisol: <5.51 nmol/L) with an undetectable ACTH level (ACTH: <1.1 pmol/L). Other anterior pituitary hormones were as follows: growth hormone 14.8 μg/L; thyroid-stimulating hormone 1.73 U/L; free thyroxine 14.02 pmol/L; prolactin 390 mIU/L (3-24); follicle-stimulating hormone <3 U/L (0.1-3.3); luteinizing hormone <0.07 U/L (0-1.9). A low-dose ACTH stimulation test showed an insufficient cortisol response at 40 minutes (12.1 nmol/L). A magnetic resonance imaging (MRI) scan of the pituitary gland was normal. A diagnosis of isolated central (secondary) ACTH insufficiency, rather than panhypopituitarism was made.

Hydrocortisone treatment was initiated which subsequently enabled successful control of the hypoglycemia. Due to the presence of central adrenal insufficiency together with red hair, a genetic analysis of the POMC gene was undertaken. A homozygous frameshift mutation, c.206delC (p.P69Lfs*2) in the POMC gene was detected (5). This mutation results in a downstream frameshift and premature protein truncation, removing ACTH and other important peptides and most likely completely disrupting POMC function (Figure 1B).

Following this initial presentation, in subsequent months, the child showed rapid increase in growth and developed...
obesity. At 17 months of age her weight was 16.9 kg (>97th percentile), height 80 cm (50th percentile), and head circumference 40 cm (75-90th percentile) (Figure 1A). Her eyebrows and hair were red (Figure 2). The final steroid treatment dose was 8 mg/m²/day. An informed consent form for publication was given by the parents.

Discussion

Congenital isolated ACTH deficiency is a rare condition and the symptoms and signs can be nonspecific. However, it can be life threatening unless appropriate steroid replacement is initiated. POMC is synthesized in the corticotropin cells of the pituitary gland by the action of the transcription factor TBX19/TPT1. POMC is then cleaved to form ACTH by the enzyme PC1 [PC1/3, proprotein convertase subtilisin/kexin (PCSK) type 1] following corticotropin-releasing hormone stimulation (Figure 1B) (6). Isolated ACTH deficiency can result from pathogenic variations of the TBX19 (TPIT), PCSK1 (PC1/3) and POMC genes (6).

Although POMC defects were first reported in 1998, relatively few children with the condition have been reported to date (4). The classic triad of POMC deficiency consists of early-onset obesity, central adrenal insufficiency and red hair. Hyperphagia (80-99%), cholestasis (30-79%, at onset), exponential weight gain (100%) and adrenal insufficiency (30-79%, at onset) are typically observed during the first months of life, but the diagnosis may only be established later in some children. Linear growth is initially normal, as in our patient, and weight gain may not occur initially in a child with uncontrolled adrenal insufficiency. However, weight often increases to above the 90th percentile by the end of the first year. This process likely reflects an insufficiency of hypothalamic POMC. Normally, nutrition and energy hemostasis is balanced by the complex interaction of POMC and AgRP/neuropeptide Y (NPY) with MCR in the hypothalamus (Figure 3) (1,2,3,4). This system is also regulated by peripheral polypeptides such as leptin and ghrelin. Our patient showed a rapid and early-onset weight gain as a result of this process. The weight gain was independent of steroid treatment as only a physiological replacement dose of hydrocortisone was used and subsequent linear growth rate was stabilized on the 50th percentile line, despite ongoing rapid weight gain.

The red hair associated with POMC deficiency is an important sign, especially in children from an ancestral background of dark hair. However, there are a few reports of children with POMC deficiency who do not have red hair or where only the roots of the hair are red (6,7,8,9). Other children may have red hair initially but this turns brown in the first three to four years of life (10).

Other reported features potentially associated with POMC deficiency include pale skin (Fitzpatrick type 1) due to reduced stimulation of MC1R by MSH; central hypothyroidism, possibly due to interactions between POMC and thyrotropin-releasing hormone in the hypothalamus (10,11,12,13); and hypogonadotropic hypogonadism with pubertal growth hormone deficiency reflecting a possible direct interaction between POMC and gonadotropin-
releasing hormone neurons or indirectly via kisspeptin and NPY/AgRP. Our patient did not show hypothyroidism, but did have pale skin and had a low gonadotropin level in early infancy at around the time of the typical “minipuberty”. This finding may reflect impaired gonadotropin release so it is important to monitor the development of puberty in these patients. Transient hyponatraemia has been reported with central ACTH insufficiency, as was seen in our patient (14). This may reflect decreased free water clearance due to hypocortisolæmia, especially with intercurrent infections, or a supportive mineralocorticoid effect of cortisol at times of stress. Detecting hyponatraemia can sometimes lead to a misdiagnosis of primary adrenal insufficiency instead of a secondary or central defect. Finally, developmental delay with abnormal MRI changes has been reported in one child with POMC deficiency. This finding is likely to represent the effects of recurrent hypoglycaemia rather than the underlying condition itself (15).

Genetic analysis was useful to establish the diagnosis of POMC deficiency. Using a custom adrenal array coupled with next generation sequencing we identified a homozygous c.206delC mutation in the child (5). This nucleotide deletion was confirmed by Sanger sequencing and causes a frameshift and premature truncation of the protein at codon 70 (p.Pro69Leufs2*) (Figure 1B). This mutation results in a POMC product that lacks ACTH, α-MSH and other small peptides, and may be subject to non-sense mediated decay. A review of the literature shows that all reported patients with POMC deficiency have homozygous or compound heterozygous mutations in the amino-terminal region of the protein that result in defective ACTH and α-MSH synthesis. The c.206delC change has been reported in two other families in Turkey suggesting a founder effect (7). The only established point mutation in POMC causing a similar phenotype is p.Arg145Cys change that corresponds to a p.Arg8Cys mutation in the ACTH peptide (16). This point mutation results in a bioinactive form of POMC/ACTH with a clinical phenotype of red hair, obesity and central adrenal insufficiency but with elevated ACTH levels on biochemical testing. In addition, isolated obesity has been reported in carriers of POMC mutations or in association with heterozygous point mutations in POMC (especially p.Arg236Gly) (7,17,18,19). Overview of the clinical and molecular features of patients with POMC insufficiency published to date were presented in Table 1 (Table 1). However, the parents of our child had BMIs of 23 kg/m² and 27 kg/m².

Treatment of POMC deficiency can be challenging. Patients require life-long glucocorticoid treatment using replacement doses. Mineralocorticoid replacement is not required. Hypothyroidism should be monitored and treated if present. Early onset obesity can be very difficult to treat beyond standard dietary and lifestyle measures, but the hyperphagic component is especially challenging. Krude et al (10) attempted intranasal ACTH treatment in two index cases with the ACTH peptide fragment identical to α-MSH. However, ACTH treatment at low doses during the first six weeks followed by a high dose (5 mg/day) did not produce a significant response in weight loss. Recently, setmelanotide (Rhythm Pharmaceuticals, Boston, Massachusetts, USA) has been developed as a novel MC4R agonist for the treatment of rare genetic disorders of obesity associated with defects in the MC4 pathway. This novel therapy is currently in Phase II trials in patients with POMC deficiency.
| cDNA | Protein | Ancestry | Age     | ACT H/AI | Red hair          | Obesity        | Other features                                      | Reference |
|------|---------|----------|---------|----------|-------------------|----------------|----------------------------------------------------|-----------|
| c.-11C > A | Alternative translation | Dutch | 4 weeks | ↓ | + Changed to brown at age 2-3 yrs | Early | Conjugated hyperbilirubinemia | 10        |
| c.-11C > A | Alternative translation | German | 5 years | ↓ | + | Early | Subclinical central hypothyroidism | 3         |
| c.-11C > A / c.403_404dupGG | p.Lys136Alafs*23 | Swiss | 6 months | ↓ | + | Early |  | 10 |
| c.64delA | p.Met22Trpfs*49 | Turkish | 3.5 years | ↓ | ND | Early | Developmental delay, ataxia | 15        |
| c.151A > T / c.296delG | p.Lys51* / p.Gly99Alafs*59 | Slovenian | Neonatal | ↓ | + | Early |  | 10 |
| c.202C > T | p.Gln68* | Egyptian | 9 months (transient hypo early) | ↓ | - | Early |  | 9 |
| c.206delC | p.Pro69Leufs*2 | Turkish | 2 years | ↓ | ( + ) Red roots | Early |  | 7 |
| c.206delC | p.Pro69Leufs*2 | Turkish | 2 weeks | ↓ | + Brown later | Early | Central hypothyroidism | 20        |
| c.206delC | p.Pro69Leufs*2 | Turkish | 2.5 months | ↓ | + | Early |  | This report |
| c.223dupC | p.Arg75Profs*44 | North African (Kabilian) | 4 weeks | ↓ | - | Early | GH deficiency hypogonadism | 8         |
| c.231C > A | p.Tyr77* | Hispanic | 9 months | ↓ | - | Early | Apnea, neonatal jaundice; transient hyponatremia | 6         |
| c.256C > T | p.Arg86* | Indian | 1 week | ↓ | ( + ) Skin and hair lighter than expected | Early | Central hypothyroidism | 21        |
| c.296delG | p.Gly99Alafs*59. | Turkish | 3 months | ↓ | + | Early | Transient hypoglycaemia; transient salt-wasting during UTI | 14        |
| c.313G > T / c.433delC | p.Glu105* / p.Arg145Alafs*13 | German | 3 years | ↓ | + | Early | Subclinical central hypothyroidism | 3         |

cdNA: complementary DNA, ND: not described, GH: growth hormone, UTI: urinary tract infection, ACTH: adrenocorticotropic hormone, AI: adrenal insufficiency
In summary, we present a Turkish child with POMC deficiency due to a potential founder POMC mutation. Routine genetic analysis in patients suspected of POMC deficiency is recommended not only to guide long-term prognosis and tailor the personalized management of these patients per se, but also to enable discovery of breakthrough treatments for important public health problems such as obesity.

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Of note, the authors have no links with Rhythm Pharmaceuticals or any company developing novel anti-obesity therapeutics.

Ethics

Informed Consent: The informed consent was taken from the patient’s parents for publication.

Peer-review: Externally peer-reviewed.

Authorship Contributions

Surgical and Medical Practices: Semra Çetinkaya, Erdal Kurnaz, Meliṣkāh Keskin, Elif Sağsak, Şenay Savaş Erdeve, Zehra Aycan, Concept: Semra Çetinkaya, Tülay Güran, John C. Achermann, Design: Semra Çetinkaya, Tülay Güran, John C. Achermann, Data Collection or Processing: Semra Çetinkaya, Erdal Kurnaz, Analysis or Interpretation: Jennifer P. Suntharalingham, Federica Buonocore, Tülay Güran, John C. Achermann, Literature Search: Semra Çetinkaya, Tülay Güran, John C. Achermann, Writing: Semra Çetinkaya, Tülay Güran, John C. Achermann.

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