EXCEPTIONAL CASE

New combined CFH/MCP mutations and a rare clinical course in atypical haemolytic uraemic syndrome

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Abstract

Atypical haemolytic uraemic syndrome (aHUS) is a rare, life-threatening, chronic, genetic disease due to uncontrolled alternative pathway complement activation. In this report, we discuss the case of a heterozygous carrier of a mutation on both factor H and membrane cofactor protein, who persistently presents haemolytic anaemia without need for blood transfusions, normal platelet count, normal renal function and no signs or symptoms of organ injury due to thrombotic microangiopathy 4 years after the diagnosis of aHUS.

Key words: atypical haemolytic uraemic syndrome, membrane cofactor protein, protein factor H

Introduction

Atypical haemolytic uraemic syndrome (aHUS) is a rare, life-threatening, chronic, genetic disease due to uncontrolled alternative pathway complement activation [1]. In most cases, aHUS is associated with mutations or polymorphisms in the genes CFH (encoding complement factor H), accounting for 11–29%, CD46 (MCP, membrane cofactor protein) 3–17% and CFI (encoding complement factor I) 2–17% [2]. Mutations in more than one complement gene have been identified in some patients, and nucleotide polymorphisms in CFH and MCP genes can modulate the risk of aHUS in mutation carriers [3].

Case report

A 31-year-old female presented in our hospital with abdominal pain. The investigations performed showed gallbladder infection and microangiopathic haemolytic anaemia, thrombocytopenia and acute kidney injury. She had a past medical history of kidney stones and one uneventful pregnancy, and no previous history of diarrhoea. She had no family history of kidney disease or aHUS. Initial investigation revealed the following: haemoglobin 8.3 g/L, platelet count 140 × 10⁶/μL and serum creatinine 3.8 mg/dL.

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Table 1. Variations on biochemical parameters during the 4-year follow-up

|                         | Hospital discharge | 1 month | 6 months | 12 months | 18 months | 3 years | 4 years |
|-------------------------|--------------------|---------|----------|-----------|-----------|---------|---------|
| Serum creatinine (mg/dL)| 2.8                | 1.1     | 0.7      | 0.5       | 0.6       | 0.5     | 0.7     |
| Haemoglobin levels (g/dL)| 12.0              | 12.4    | 11.7     | 11.2      | 10.6      | 11.3    | 10.4    |
| Platelet count (cells x 10^5/μL)| 190          | 262     | 253      | 254       | 269       | 195     | 193     |
| LDH (U/L)               | 903                | 1054    | 774      | 942       | 850       | 972     | 1554    |
| Serum haptoglobin (g/L) | <10                | <10     | <10      | <10       | <10       | <10     | <10     |
| UPCR (mg/mg)            | 0.65               | 0.14    | 0.07     | 0.06      | 0.01      | 0.08    | 0.05    |
| Complement C3 (mg/dL)   | 96.8               |         |          |           |           |         |         |
| Complement C4 (mg/dL)   | 21.8               |         |          |           |           |         |         |
| Homocysteine levels (μmol/L) | 10.0          |         |          |           |           |         |         |

UPCR: protein-creatinine ratio on random urine sample.

1054 U/L, serum creatinine 1.1 mg/dL). The study performed at presentation showed normal ADAMTS13 activity and C3 serum levels. The patient was clinically diagnosed with aHUS, possibly triggered by the gallbladder infection. After 4 years, her renal function and platelet number are normal, but the patient has persistent microangiopathic haemolytic anaemia (Table 1). Despite analytical parameters of haemolysis (LDH 1554 U/L, indirect bilirubin 1.54 mg/dL and absent haptoglobin), the patient has no signs or symptoms of organ injury/thrombosis, no hypertension, no proteinuria and the haemoglobin levels remain stable with no need for blood transfusion. During these 4 years, the patient had possible triggering events (infection and cholecystectomy), with no relapse of aHUS. In keeping with this, we performed a genetic analysis to investigate whether the patient carried some genetic abnormalities associated with aHUS. Mutation screening of CFH, MCP and CFI by Sanger sequencing revealed two heterozygous mutations: a CFH (c.3493 C>T, p.His1165Tyr) and an MCP (c.1058 C>T, p.Ala353Val). The first amino acid substitution is not described, and the second is known as a disease mutation [4, 5].

**Discussion**

aHUS is a challenging disease to manage with a relatively poor prognosis, as up to 65% of patients treated with PE will sustain permanent renal damage, have progression to end-stage renal disease (ESRD) or die within 1 year [6]. In the past, it became evident that aHUS is associated with mutations in proteins needed for regulation of the alternative complement pathway [7]. Because of the multiple genetic susceptibility factors involving plasma- and membrane-associated regulators, the rarity of combined-mutated patients and the need for a triggering event, it is difficult to predict the clinical course of aHUS. The combined CFH/MCP mutations reported in our patient (CFH p.His1165Tyr/ MCP p.Ala353Val) have not been previously described. As previously reported, it seems that MCP or CFH mutations confer a predisposition to develop aHUS rather than directly cause the disease, and a second hit is necessary for the full-blown manifestation of the disease, as in all MCP-mutated and 70% of CFH-mutated patients the onset of aHUS was associated with an infectious event [4]. Our patient showed an impaired capacity to protect vascular endothelial cells from complement attack after the activation caused by the gallbladder infection that led to aHUS presentation. After that, the clinical course was clearly distinct from patients with other CFH mutation and was similar to patients with MCP mutations described previously. It has been reported that the most severe prognosis is in the CFH mutation patients, with 70% reaching ESRD, and that patients with MCP mutations have a relapsing course, but none have reached ESRD in the first year [8]. In cases of combined mutations carriers, the concomitant presence of CFH and MCP risk haplotypes significantly increased disease penetration and modified the prognosis [5]. Among patients with CFH mutations, the presence of mutations in other genes did not modify prognosis; however, 50% of patients with combined MCP mutation developed ESRD within 3 years from onset compared with 19% of patients with an isolated MCP mutation [5]. These two mutations (CFH p.His1165Tyr/MCP p.Ala353Val) have a benign phenotype prediction (Polyphen-2), and MCP p.Ala353Val has polymorphic prevalence (1000 Genomes Project). Despite that, their cumulative effect confers a background genetic that seems to be a risk for aHUS; however, their benign character explains the mild clinical course. Indeed, the response to PE would not be expected to be effective for patients with mutations in MCP, a transmembrane protein that is not removed by this treatment, but patients with combined mutations achieved remission with PE similar to patients with single non-MCP mutations [9]. The excellent response of our patient could be related to the replacement of CFH proteins with PE. In our patient, it seems that the mutated expressed proteins cannot fully protect vascular endothelial cells from activated complement translated by the persistently haemolytic microangiopathic anaemia; however, the regulation of the complement attack is enough to protect her from a relapse.

In summary, we describe the first reported case of aHUS with the combination of a previously described MCP mutation with a new CFH mutation associated with a rare clinical outcome (persistent haemolysis without renal failure or any other clinical manifestations of endothelial damage or platelet activation). We may therefore speculate that combination of these frequent mutations will worsen the phenotype and could be a common but under-diagnosed phenomenon. The findings in this patient underscore the influence of different genetic abnormalities on disease presentation, response to therapy and outcome in aHUS patients.

**Conflict of interest statement**

The authors declare that the results presented in this article have not been published previously in whole or part.

(See related article by Sanchez-Nino and Ortiz. Thrombotic microangiopathy: expanding genetic, clinical and therapeutic spectra and the need for worldwide implementation of recent advances. Clin Kidney J 2015; 8: 686–689.)
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