The Relevance of the MCP Risk Polymorphism to the Outcome of aHUS Associated With C3 Mutations. A Case Report

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Thrombotic microangiopathy (TMA) has different etiological causes, and not all of them are well understood. In atypical hemolytic uremic syndrome (aHUS), the TMA is caused by the complement dysregulation associated with pathogenic mutations in complement components and its regulators. Here, we describe a pediatric patient with aHUS in whom the relatively benign course of the disease confused the initial diagnosis. A previously healthy 8-year-old boy developed jaundice, hematuria, hemolytic anemia, thrombopenia, and mild acute kidney injury (AKI) in the context of a diarrhea without hypertension or oliguria. Spontaneous and complete recovery was observed from the third day of admission. Persistent low C3 plasma levels after recovery raised the suspicion for aHUS, which prompted clinicians to discard the initial diagnosis of Shigatoxin-associated HUS (STEC-HUS). A thorough genetic and molecular study of the complement revealed the presence of an isolated novel pathogenic C3 mutation. The relatively benign clinical course of the disease as well as the finding of a de novo pathogenic C3 mutation are remarkable aspects of this case. The data are discussed to illustrate the benefits of identifying the TMA etiological factor and the relevant contribution of the MCP aHUS risk polymorphism to the disease severity.

Keywords: C3, MCP risk polymorphism, atypical hemolytic uremic syndrome, de novo mutation, case report

INTRODUCTION

Atypical hemolytic uremic syndrome (aHUS) is an ultra-rare disease characterized by acute kidney injury, thrombocytopenia, and microangiopathic hemolytic anemia, which results from an impaired protection of host endothelial cells from complement damage (1). The complement system is a key element of innate immunity with crucial roles in the elimination of pathogens, immune complexes, or cell remains. The complement activates by three pathways, classical (CP), lectin (LP), and alternative (AP), which generates protease complexes, named C3 convertases that cleave C3 to generate C3b. Convertase-generated C3b can form more AP C3 convertase, providing exponential amplification of the initial activation. Clustering of C3b around the surface-bound C3 convertase generates the C5 convertase, which cleaves C5 and initiates formation of the lytic membrane attack complex (MAC) (2). In health, the activation of C3 in plasma is kept
at a very low level, and the deposition of C3b and further activation of complement are limited to the surface of pathogens by multiple regulatory proteins. The loss of complement regulation leads to the generation of proinflammatory components and/or tissue damage. Both situations have pathological consequences (3). Loss-of-function mutations in genes encoding the regulatory proteins factor H (FH), MCP, and factor I (FI), as well as gain-of-function mutations in the complement activating components factor B (FB) and C3, have been associated with aHUS (4–11). Criteria have been established to facilitate the clinical diagnosis of aHUS, but it is often difficult to exclude STEC-HUS and secondary HUS forms (12). Since 50–70% of aHUS patients have an underlying inherited and/or acquired complement abnormality (13, 14), genetic analyses are recommended to characterize the etiological factor, reinforce diagnosis, and assist patient management. We present a case that was initially classified as STEC-HUS but was reclassified to aHUS based on the complement follow-up and genetic analyses. We discuss the implications of the identification of a de novo gain-of-function C3 mutation in this case and the relevance of genotyping for the MCPggaac aHUS risk polymorphism.

**Clinical Case**

In October 2014, a previously healthy 8-year-old boy was evaluated at the pediatric emergency room (ER) in a tertiary care hospital for hematuria, asthenia, and mild jaundice observed in previous hours. Nonbloody diarrhea had been present for 3 days. Physical examination was unremarkable apart from mild jaundice. Nonfocal or generalizable edema was found. He had no relevant personal or family past history. Initial blood test showed hemoglobin of 12.2 g/dl (>11.5 g/dl), platelets of 35,000/µl (>150,000/µl), creatinine of 84 µmol/L (<61 µmol/L), and normal transaminases, sodium, and potassium. Eight hours later, hemoglobin decreased to 10 g/dl and platelets to 28,900/µl; lactate dehydrogenase (LDH) was determined to be 1,657 U/L (<220 U/L) (Figure 1). Blood test was extended with haptoglobin (undetectable), and a blood smear showed 7–9 schistocytes per field. Creatinine increased to 93 µmol/L. Electrolytes, acid–base balance, and plasma proteins were in normal range. Urine protein to creatinine ratio (UPr/UCr) was 1,921 µg/µmol (<20 µg/µmol). Basic coagulation parameters were normal. Blood pressure remained spontaneously in normal range and diuresis preserved, without involvement of other organs or systems. Patient was admitted to the pediatric ward.

Maximum plasma creatinine was attained on the second day of admission (106 µmol/L). Hemolysis markers started to descend from the third day. He was discharged on the sixth day of admission with hemoglobin of 10.8 g/dl, platelets of 201,000/µl, LDH of 639 U/L, creatinine of 65 µmol/L, and UPr/UCr of 34 µg/µmol (Figure 1).

Additional studies performed during admission revealed the following: plasma homocysteine was normal at 11.4 µmol/L (<15 µmol/L), autoantibodies [antinuclear antibody (ANA), antineutrophil cytoplasmic antibody (ANCA), and antiejectable nuclear antigen (anti-ENA)] were negative, C3 was 51 mg/dl (75–135 mg/dl) and C4 was 21 mg/dl (14–60 mg/dl), and plasma ADAMTS13 activity was 81% (>5%). A stool sample was only obtained after 4 days of admission. Because of its completely normal appearance and the satisfactory evolution of HUS at that moment, it was only tested for Shigatoxin. Blood and urine culture were sterile. The patient was under careful observation during admission without needing renal replacement therapy or any drug. The spontaneous and very favorable evolution, the previous history of diarrhea, and the justification of a negative Shigatoxin assay due to a late stool collection suggested STEC-HUS as the most likely etiology.

Successive controls showed a complete recovery of renal function and absence of hemolytic activity, anemia, and thrombopenia. No treatment was needed during follow-up. However, decreased C3 levels (46–62 mg/dl) (Figure 1) persisted, and subsequent analysis of factor B plasma levels revealed that they were in the lower part of the normal range (85–170 µg/ml) (Figure 2). Hypocomplementemia is not unusual during a STEC-SHU episode, but the complement normalizes afterwards in these patients (16). The persistent hypocomplementemia in our patients did not correlate with his favorable evolution, raising the suspicion of an underlying constitutive complement abnormality that prompted us to the realization of a complete complement molecular and genetic analysis.

To search for mutations in complement genes, we used an in-house next generation sequencing (NGS) panel including all the complement genes relevant to aHUS (17). A complementary analysis of copy number variations was performed by multiplex ligation-dependent probe amplification (MLPA) with the P236 A1 ARMD mix 1 (MRC-Holland, Amsterdam, Netherlands). These analyses identified a C3 mutation in heterozygosis (c535T>C; p.S179P) that was confirmed by Sanger sequencing. This genetic variant has been found previously associated with aHUS (18). No other genetic alterations were found in this patient. Interestingly, none of his parents present this C3 variation (Figure 2A). Paternity was supported by the analysis of CFH and MCP polymorphisms. The patient carries the CFH-H3 aHUS risk polymorphism in heterozygosis, inherited from his father (Figure 2A). He does not carry the MCPggaac aHUS risk polymorphism.

Annotation of C3S179P variant with six pathogenicity prediction algorithms (SIFT, POLYPHEN, Mutation Taster, MutAss, FATHMM and CADD) included in the ANNOVAR server (http://annovar.openbioinformatics.org/) indicated that it is most likely a benign C3 variant. However, because the C3 mutations associated with aHUS are gain-of-function mutations that normally are not predicted pathogenic, we purified the C3 protein from the patient’s plasma and performed a complete functional characterization following standard procedures in our laboratory (15). These analyses demonstrated that the mutant C3S179P is present in the patient’s plasma and shows an altered function with the characteristic of the C3 gain-of-function mutations that associate with aHUS (19–21). Briefly, when purified C3 from the patient was incubated with FB and FD, it completely activated to C3b, suggesting that C3S179P is normally activated by the AP C3 convertase (Figure 2B). When the patient C3b was tested for inactivation by FI in the presence of FH or MCP, we found that it was resistant to inactivation by FI in the
The patient has remained completely asymptomatic without clinical or analytical data of disease activity or renal sequelae for 5 years, with an expectant attitude.

**DISCUSSION**

aHUS is a rare, life-threatening renal pathology associated with complement dysregulation. Mutations in genes encoding the regulatory proteins factor H (CFH), factor H-related protein

**FIGURE 1** | Evolution of hematological parameters and renal activity. Graphics show the blood test determinations since the day of admission (10-Oct-2014) to the day of discharge (15-Oct-2014) and then of the successive routine controls in the following years. Dotted lines represent the upper or minimum value of each parameter from which it is considered pathogenic (hemoglobin > 11.5 g/dl; platelets < 150/ml; LDH > 220 U/L; urine protein/creatinine > 20 µg/µmol; creatinine > 61 µmol/l; C3 < 85 mg/dl).
FIGURE 2. C3S179P is a novel de novo gain-of-function mutations that impairs regulation by MCP. (A) Pedigree depicting the segregation of CFH and MCP polymorphisms organized in haplotype blocks. Single nucleotide polymorphisms are identified by their “rs” numbers. In black are the haplotypes inherited by the patient, identified by a solid square. Asterisk identifies the CFH-H3 aHUS risk haplotype. Plasma levels of C3 and factor B (FB) are indicated for each individual. The patient is the only member of the pedigree carrying the C3S179P mutation. (B) C3 purified from the plasma-ethylenediaminetetraacetic acid (EDTA) of the patient activates normally to C3b by factor B (FB) and factor D (FD). Briefly, C3 was purified using a combination of sodium sulfate precipitation, lysine-sepharose chromatography, DEAE-Sepharose anion exchange chromatography, and Mono S HR 5/5 cation exchange chromatography as previously described (15). (C) C3b generated from the patient’s C3 shows a marked resistant to inactivation by factor I in the presence of MCP. Differences between slopes were tested with a general linear model (GLM), with “time” as an integer variable and “strain” as a nominal one. The time × strain interaction was considered as the estimator of the differences between slopes. MCP (p < 10−8), FH (p = 0.002).

1 (CFHR1), MCP (MCP), and factor I (CFI), as well as mutations in the complement components factor B (CFB) and C3 have been found in 50–70% of aHUS patients (13, 22). Importantly, while mutations in the complement regulators are loss-of-function, mutations in complement components like factor B and C3 are gain-of-function (13, 14). For C3, these aHUS-associated gain-of-function mutations result in C3b activated molecules being resistant to regulation by MCP, but not by factor H (19–21). The genetic and functional analyses performed in our patient concluded that he carries a C3 gain-of-function mutation that is prototypical of aHUS. This explains why our patient presents a constitutive alternative pathway activation with persistent consumption of C3. He has no familial history of aHUS because C3S179P is a de novo mutation, and he is the first in his pedigree carrying this genetic predisposition to aHUS. More interesting is the favorable disease outcome in our patient. Previous studies have shown that C3 mutations, like R161W, tend to be associated with severe aHUS presentations leading to end-stage renal disease. Others, like I1157T, associate with aHUS presentations characterized by multiple recurrences and prolonged favorable outcomes. Interestingly, the presence of the MCPPgaac aHUS risk polymorphism influences the aHUS presentation in all carriers of C3 mutations (10, 19–21, 23), which may be justified because this polymorphism determines reduced expression of MCP on the cellular surface (8). Our aHUS registry includes a total of 13 additional patients having a complete clinical record who carry a clearly pathogenic C3 mutation (Table 1). In total, this series comprises five different C3 mutations. Ten of these aHUS patients also carry the MCPPgaac aHUS risk polymorphism (five in heterozygosis and five in homozygosis) (Table 1). This results in an allele frequency for the MCPPgaac polymorphism in this group of patients (n = 14) of 0.54, which is significantly different (p < 0.0037) from that in the control Spanish population (AF = 0.28; n = 107). Notably, the only patient in this series who have had a relative favorable outcome is the only one who does not carry the MCPPgaac risk polymorphism. Eculizumab treatment was initiated early after aHUS onset or to treat a bad evolution in three patients, and therefore, no conclusions can be reached in them regarding natural progression of aHUS. Notably, nine of the remaining 11 patients reached end-stage renal disease (ESRD) or
had multiple recurrences (Table 1). Eight of these patients carry, in addition to the C3 gain-of-function mutation, the MCPggaaac risk polymorphism or additional pathogenic mutations in the CFH, CFI, MCP, and THBD genes. The only patient who, like our current patient, does not carry the MCPggaaac risk polymorphisms or additional pathogenic mutations had a very late onset (63 years old) without recurrences until she was 76 years old. Currently (80 years old), the patient presents chronic kidney disease but does not require hemodialysis. These registry data suggest that the likely explanation for the favorable disease outcome in our patient is that he does not carry additional genetic risk factors, in particular the MCPggaaac risk polymorphism. A relevant question is why our patient had an aHUS episode. It is known that several viral pathogens interact with MCP and that viral infections may lead to a reduction in the pathogen’s receptor. Therefore, one possibility could be that our patient underwent a transient decrease in the cell surface levels of MCP as a consequence of the infection that triggered the aHUS episode. However, this is just a speculation because, when MCP levels were tested, months after the aHUS episode, they were found normal.

Although our patient is currently asymptomatic and presents normal renal function, our functional characterization of the C3S179P variant indicates that it is an important aHUS genetic risk factor. This has important implications. In fact, we cannot exclude that under exposure to a strong environmental trigger (e.g., an infection), our patient will experience a more severe aHUS recurrence. While we strongly recommend normal life to avoid unnecessary anxiety in the patient and its family, we also suggest active surveillance of the patient with specific recommendations. Regular determination of blood pressure, blood count, and measurement of biochemical markers for hemolysis (bilirubin, LDH, haptoglobin), as well as plasma creatinine, proteinuria, and albuminuria are performed. The patient and his parents are instructed to go the ER in case of presenting symptoms suggesting activity of his disease, such as hematuria, oliguria, edema, and significant general malaise in the context of some intercurrent process that can act as a trigger. Ultimately, it is reassuring to know that we have the “magic bullet” of eculizumab, if this patient should need it (24). In conclusion, complete understanding of the etiological factor in the TMA patient is critical to strengthen diagnosis and assist patient management.

### DATA AVAILABILITY STATEMENT

All datasets presented in this study are included in the article/supplementary files.

### ETHICS STATEMENT

Written informed consent was obtained from the participant’s legal guardian/next of kin for the publication of any potentially identifiable images or data included in this article.

### AUTHOR CONTRIBUTIONS

JL, MS, and SR designed the study. JL, MS, EA, and SR performed the experiments, collected, and analyzed the data. JL and SR drafted the manuscript, which was revised and approved by all
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The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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