Atypical hemolytic uremic syndrome secondary to lupus nephritis, responsive to eculizumab

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Abstract

Among the spectrum of disease manifestations associated with systemic lupus erythematosus, lupus nephritis is particularly concerning due to the potential for renal failure. This autoimmune attack may not, however, be limited to the kidney and is increasingly being recognized as a trigger for atypical Hemolytic Uremic Syndrome (aHUS). Atypical HUS falls under the spectrum of the thrombotic microangiopathies (TMA) – a group of disorders characterized by microangiopathic hemolytic anemia, thrombocytopenia, and end organ damage. Although plasma exchange is considered first-line therapy for thrombotic thrombocytopenic purpura – a TMA classically associated with autoimmune depletion of ADAMTS-13 – aHUS demonstrates less reliable responsiveness to this modality. Instead, use of the late complement inhibitor Eculizumab has emerged as an effective modality for the management of such patients. Diagnosis of aHUS, however, is largely clinically based, relying heavily upon a multidisciplinary approach. Herein we present the case of a patient with atypical HUS successfully treated with Eculizumab in the setting of Class IV-G (A) lupus nephritis and hypocomplementemia.

Introduction

Complement mediated TMA (also referred to as atypical Hemolytic Uremic Syndrome) is a rare (annual incidence 1,500,000)1 entity belonging to the family of disorders collectively called thrombotic microangiopathies (TMA).2 The pathophysiology of complement-mediated TMA involves unchecked activation of the alternative pathway leading to excessive deposition and deleterious action of complement on the endothelium, often with a predilection for the renal vasculature.4 Failure to control complement activation is due predominantly to inherited or acquired deficiencies of complement regulatory proteins – described in over 50% of individuals with complement mediated TMA.5,6 Known mutations leading to sporadic and recurrent complement mediated TMA include those that affect regulatory proteins that are both soluble (Factor I, which, in cooperation with Factor H, MCP and other proteins cleaves C4b and C3b; Factor H, which binds C3b and disrupts the alternative pathway C3 convertase) and membrane bound (Membrane Cofactor Protein, CD46); a number of other defects have also been described.7 Patients with complement mediated TMA, even those in whom a clear genetic or antibody-mediated predilection is absent, may experience recurrences that may also damage renal allografts following kidney transplantation.5

Typically, complement mediated TMA will present as a microangiopathic hemolytic anemia, with thrombocytopenia, evidence of mechanical hemolysis (presence of schistocytes on peripheral blood smear, increased LDH, decreased haptoglobin), and varying degrees of end-organ dysfunction.2 Other manifestations include severe hypertension, central nervous system features (such as altered mental status, diplopia, or motor deficits), acute coronary syndrome due to cardiac microangiopathy, distal ischemic gangrene, and acute multi-organ failure.2 Responses to plasma-based therapeutics (either plasma infusion or plasma exchange) in complement-mediated TMA appears highest among those with MCP mutations (97%) and lowest among those with CFI mutations (25%) with 3 year outcomes of ESRD or death as high as 77% among those with CFH and lowest among those with MCP mutations (6%).5 Despite treatment responses to individual bouts of TMA, therefore, plasma based therapeutics are not uniformly effective.

Eculizumab, an inhibitor of human C5 complement protein, has emerged as an important therapeutic agent in the treatment of complement mediated TMA. This agent was studied by Legendre and colleagues in 37 patients with complement mediated TMA with or without thrombocytopenia.8 Eculizumab was dosed intravenously within 1–6 hours of most recent plasma therapy at a dose of 900 mg per week for 4 weeks, then 1200 mg on week 5, then 1200 mg every 2 weeks beginning week 6. Patients continuing plasma based therapeutics during eculizumab treatment received 600 mg eculizumab booster before plasma infusion or within 1 hour following completion of each plasma exchange. In the group with thrombocytopenia, median platelet-count improvements of 73×10³/μL from baseline occurred by week 26 of therapy; 45 patients discontinued dialysis. In the group without thrombocytopenia, 80% achieved TMA-free status.

Here we present a unique case of aHUS which developed in a patient with newly diagnosed systemic lupus erythematosus (SLE), failed a course of therapeutic plasma exchange, but that subsequently responded to eculizumab.

Case Report

A 25 year-old Vietnamese female presented with a one month history of arthralgias, daily symmetric arthritis of the metacarpophalangeal (MCP), proximal interphalangeal (PIP), and metatarsophalangeal (MTP) joints accompanied by two hours of morning stiffness. In addition, she reported myalgias, facial swelling, intermittent nausea, vomiting, diarrhea, and generalized headaches associated with photophobia/sensitivity but no visual or focal neurologic deficits. Ten days prior to admission, she developed a mild, productive cough but no fever or chills. She also had dyspnea on exertion, one-pillow orthopnea, and intermittent pleuritic chest discomfort – most pronounced on deep inspiration. On exam she was noted to have a subtle malar rash and diffuse tenderness on abdominal exam.

Pertinent laboratory results include a positive antinuclear antibody (ANA; in an atypical speckled pattern suggestive of SSA/Ro) with a titer to >320; double-stranded DNA with a titer to >5120; positive ANCA but negative MPO and PR3 antibodies. Anti-Smith IgG was elevated at 31 U (0.9-19), SSA (RO) was elevated...
treatment with empiric antibiotics, high-dose corticosteroids, and mycophenolic acid with progression of renal failure to a peak Cr of 2.5 mg/dL by day 16 and proteinuria (by spot urine Pr/Cr ratio) to a peak of 11.55 mg/mg by day 19 (Figure 1). In addition, a significant progression in her microangiopathic hemolytic picture occurred, with development of red cell transfusion dependence and hemolysis to a nadir hemoglobin of 6.4 g/dL by hospital day 10 and nadir platelet count of 21 K/mcL by day 13. This was accompanied by a rise in the LDH to peak of 1689 U/L on day 12. Concern existed for intermittent mycophenolic acid-mediated cytophenias, so this drug was stopped. To address the lupus nephritis, cyclophosphamide was administered on Day 14.

Plasma-based therapy was initially withheld given absence of microthrombi on preliminary (but not final; Figure 2) renal biopsy report. This was felt to be justified with return of ADAMTS-13 activity level at 99%. However, after further multi-disciplinary discussion, it was decided that a therapeutic trial of plasma exchange would proceed following first dose of rituximab (with appropriate window to prevent premature apheresis removal of rituximab, which was added as a salvage agent for lupus nephritis) with plan to convert to eculizumab if unresponsive. An atypical HUS panel, involving sequencing and analysis of the exonic regions of 12 genes: CFH, MCP (CD146), CFI, C3, CFB, CFHR1, CFHR3, CFHR4, CFHR5, thrombomodulin, plasminogen, DGKE – was negative for disease associated mutations.

Hemolysis and red cell transfusion requirement continued in the face of plasma exchange therapy, but platelet counts were responsive. Creatinine declined during plasma exchange but it was unclear what proportion of this decline represented a therapeutically significant renal response and what proportion represented a plasmapheresis removal effect. Rituximab dose 2 (day 30) and cyclophosphamide dose 2 (day 34) were applied with planned pauses in the apheresis regimen to maximize immunosuppressive agent dwell time. These pauses were associated with immediate declines in the platelet count. Additionally, proteinuria did not appreciably respond to the plasma exchange course with a spot urine Pr/Cr ratio of 10.84 mg/mg on hospital day 50, following 18 rounds of plasma exchange.

Plasma exchange was also declared ineffective at controlling the microangiopathic picture. Following meningococcal and pneumococcal vaccination and institution of antibiotic prophylaxis against meningococcal infection, Eculizumab was initiated on day 49. Eculizumab induced immediate reduction in hemolytic pace allowing spontaneous recovery of hemoglobin levels. Platelet counts, which had been rising in response to plasma exchange, demonstrated a pronounced increment after eculizumab administration followed by progressive recovery thereafter. She was discharged following the second eculizumab dose on day 56.

Haptoglobin, which had been absolutely suppressed throughout her hospitalization, again became detectable following 5 weekly doses (900 mg IV 4 then 1200 mg IV week 5) by day 83 signaling complete resolution of the hemolytic process. In addition, significant

![Figure 1 Renal biopsy images demonstrating the presence of Class IV G (active) lupus nephritis. A) Jones Silver stain demonstrating fibrin thrombus (arrow) within a hilar arteriole; B) Hematoxylin & Eosin stain demonstrating diffuse endocapillary hypercellularity (white arrows) and wire loops (black arrow).](image-url)
renal recovery occurred during Eculizumab therapy with improvement in BUN, Cr and urine Pr:Cr to 25 mg/dL, 1.1 mg/dL, and 1.6 g/day also by day 83. Her proteinuria continued to improve with spot urine Pr:Cr ratio of 0.65 mg/mg by day 160. At last follow up, day 226 eculizumab maintenance had continued to repress hemolytic activity with Hb 11.0 g/dL, platelet count 306 K/mcL, Cr 0.9 mg/dL, and spot urine Pr:Cr 0.46 mg/mg.

Discussion

We describe a case of complement mediated TMA with lupus nephritis as the apparent driver in a patient with negative aHUS gene panel and partial (platelets only) response to therapeutic plasma exchange. Important aspects of this case include the continued hemolysis and transfusion requirement in the face of daily plasma exchange as well as the requirement for at least 5 weekly doses of eculizumab prior to complete cessation of hemolysis.

Our patient demonstrated a significant degree of hemolytic involvement with resolution of haptoglobinemia only after multiple (in our case 5) doses of eculizumab. Coppo et al. reiterate this experience in their case report of a child with diffuse proliferative lupus nephritis also associated with significant proteinuria (with spot urine Pr:Cr as high as 10 mg/mg) and atypical HUS whose haptoglobinemia did not improve until after 3 doses of eculizumab. As with our case, theirs demonstrated exquisite platelet responsiveness to eculizumab, a negative aHUS genetic screening panel, and significant improvement in renal function and proteinuria.

We noted a selective response to plasma exchange in our patient – platelet counts, and possibly creatinine improved, but hemolysis was unresponsive. A very similar patient, also with diffuse proliferative lupus nephritis, Class IV G with nephrotic-range proteinuria (4 g/day) and detectable ADAMTS-13 (33%) activity did respond to treatment with therapeutic plasma exchange. Interestingly, response did not appear to accompany the initial 5/12 plasma exchange but both renal and hematologic responses were noted following initiation of eculizumab.

Interestingly, eculizumab may also bear value in the setting of recalcitrant, non-TMA lupus nephritis. Pickering and colleagues report a patient in whom Class IV-G (A/C) lupus nephritis involving hypocomplementemia and nephrotic range proteinuria refractory to multiple rounds of cyclophosphamide, rituximab, mycophenolate mofetil, and tacrolimus responded to eculizumab therapy. Notably, renal function and proteinuria responded dramatically following initiation of eculizumab therapy (4 weekly doses of 1200 mg followed by two 1200 mg doses every 2 weeks).

The connection between complement activation and thrombotic processes involved during microangiopathy has recently been explored. Binding of C3a and C5a to their respective receptors leads to activation of endothelial membranes and enhanced expression of adhesion molecules and secretion of Von Willebrand Factor and P-selectin with consequent platelet binding and activation. Concomitantly, downregulation of surface thrombomodulin expression further enhances the prothrombotic phenotype by impairing the Protein C pathway. Receptors for C3a and C5a also appear on platelets leading to platelet activation upon complement binding.

An aHUS panel, performed in our patient by an outside laboratory specializing in aHUS diagnostics, was negative. The presence of either a known-but-untested or an as-yet undescribed genetic complement abnormality remains possible. In two large series of patients with both familial and sporadic aHUS, the absence of detectable genetic abnormalities was reported in ~25% to 48% of individuals.5,6
The tendency toward unchecked complement activation following sometimes innocuous triggers, such as upper respiratory tract infection, gastroenteritis, or pregnancy, may initiate a potentially fatal microangiopathic process that ultimately leads to end stage renal disease in over 50% and death in over 30% of affected individuals. While an alternative driver — such as infection — was certainly possible in our patient, there was not an appreciable response with empiric antibiotics and the overall clinical presentation appeared most consistent with systemic lupus erythematosus disease activity.

We therefore conclude that the primary driver for complement activation — as evidenced by hypocomplementemia persisting even in the face of daily receipt of donor plasma (a source of donor complement) during therapeutic apheresis — was a renally focused (lupus nephritis) nidus of intense immunologic activity. Unfortunately, the immunofluorescence samples in our case were corrupted thus no immunofixation results exist. Initiation of eculizumab to halt what was a secondary consequence of primary lupus nephritis proved effective.

Once on eculizumab, it is difficult to know when the medication can be discontinued. In many instances, maintenance treatment is continued indefinitely; an important factor given estimated annual drug costs of $350,000 to $645,000. Early reports suggest that maintenance dosing may be modified, and in some cases even suspended, provided complement activity remains suppressed and patients are carefully surveilled (using home urine dipstick for hemoglobinuria and periodic laboratory testing for markers of hemolysis and schistocytosis) to detect relapse, which would require immediate reinitiation of eculizumab. Gatault et al., studied the use of eculizumab trough levels finding that elimination half-life varied significantly based upon weight with an increase from 7.8 days in 100 kg patient to 19.5 days in a 40 kg patient. Preliminary data, therefore, suggested that 1200 mg maintenance doses could be spaced to every 4 weeks in patients <90 kg and every 6 weeks in patients <70 kg but that additional studies are required to more clearly identify optimal eculizumab dosing, treatment schedules, monitoring, and endpoints for therapy.

Results for ADAMTS-13 activity and aHUS gene panels may be non-diagnostic, but hypocomplementemia is an important clue. Renal outcomes in the setting of complement-mediated TMA, even in the setting of intercurrent hematologic responses, are typically poor. Eculizumab, although expensive, is highly effective at inducing both hematologic and renal responses. Although plasma-based therapeutics can be initiated, clinicians should have a low threshold to move on to late-complement inhibition in the face of non-response — particularly if the ADAMTS-13 activity level is detectable. Although experience is growing in terms of recognition of complement-mediated TMA, additional studies are required to more clearly identify optimal eculizumab dosing, treatment schedules, monitoring, and endpoints for therapy.

References