Review Article

Hemolytic uremic syndrome: An updated review

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ABSTRACT

Hemolytic uremic syndrome (HUS) is a microangiopathic thrombotic disease, which is classified into atypical, typical, and secondary types. Thrombocytopenia, acute kidney failure, and hemolysis are the main features of HUS regardless of its type. Infection with Shiga toxin-producing *Escherichia coli* causes typical HUS, and gene mutations trigger atypical HUS, while secondary HUS is associated with bone marrow transplantation, autoimmunity, cancer, and other diseases. New insights into the pathogenesis of HUS have emerged over the past decades, suggesting an important role of the complement system in disease pathogenesis, which has been reinforced by the efficacy of plasma exchange and monoclonal antibodies in its treatment. In this review, we performed an updated review of HUS with a focus on understanding its pathogenesis.

Key words: Atypical hemolytic uremic syndrome, Hemolysis, Hemolytic uremic syndrome, Renal impairment, Shiga toxin, Thrombocytopenia, Typical hemolytic uremic syndrome

Intravascular hemolysis, microthrombi, low platelet count, and renal impairment are features of thrombotic microangiopathies [1], which, as described by Wardle in 1988, are roughly classified into thrombotic thrombocytopenic purpura (TTP) and hemolytic uremic syndrome (HUS). Despite the similarity in their clinical presentation, HUS and TTP have been clearly described as distinct diseases after the identification of von Willebrand factor-cleaving metalloprotease (a disintegrin and metalloprotease with thrombospondin type 1 repeats, member 13 (13ADAMTS13) [2].

HUS is classified into typical, atypical, and secondary subtypes (Fig. 1). The recognized features of HUS occur in HELPP syndrome (elevated hemolysis and liver enzymes and low platelet count) [3], disseminated intravascular coagulation [4], malignant hypertension [5], antiphospholipid antibody syndrome [6], and scleroderma renal crisis [7].

Although there are differences in the pathophysiology of HUS subtypes, clinical differentiation between them is challenging. Typical HUS is the most common type and is associated with Shiga toxins produced by infection with *Escherichia coli* (*E. coli*). Atypical HUS is originally due to gene mutations, leading to complement system alternative pathway activation, injuring the

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vascular endothelial layer. The damaged endothelium causes platelet activation and consumption, resulting in multiple thrombus formations, red blood cell (RBC) lysis, schistocyte, and fragmented RBC formation [8]. Secondary HUS is usually linked to calcineurin inhibitor usage, autoimmune conditions, post-transplantation, viral infection (HIV, Barr, and parvovirus), etc. Before diagnosing atypical HUS, it is essential to rule out primary and secondary HUS, since the treatment options depend on the triggering factors [1]. The aim of this review is to discuss advances in our understanding of the pathogenesis of HUS and the therapeutic options for each subtype.

HUS SUBTYPES

Typical HUS

The worldwide incidence of HUS is 0.5–2.1 cases/10⁶/year, with a peak incidence in children aged <5 years (6.1 cases/10⁶/year). However, there is variation in the incidence of typical HUS in different countries: In the UK, it is 0.91 cases/10⁶/year, in Scotland, 1.25 cases/10⁶/year, and 1.44 cases/10⁶/year in Canada. It was reported that out of 274 typical HUS cases, 122 cases were children aged <4 years in 2015 in the USA [9].

Typical HUS usually begins after a week of the evident clinical gastroenteritis due to Shiga toxin-producing *E. coli* infection

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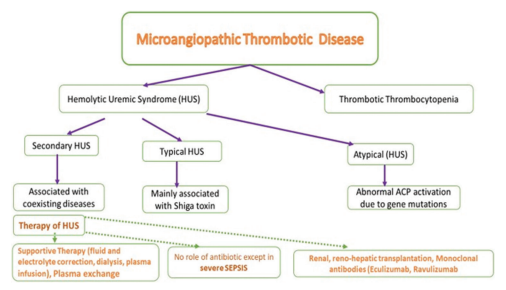


Figure 1: Classifications of microangiopathic thrombotic disease and the therapeutic options of hemolytic uremic syndrome

(mostly serotype O157:H7 or O104:H4). Clinically, typical HUS patients present with prodromal gastroenteritis (83%), fever (56%), bloody diarrhea (50%) for about 2–7 days, generalized weakness, seizures (20%), acute kidney impairment (97%), anuria (55%), hypertension (47%), severe pallor, and fluid overload (69%). Pathogenesis of typical HUS is largely induced by potent Shiga toxins (Shiga-like toxic protein) that combine with glycolipid Gb3 of the cell membrane in the existence of domain B. The combination causes internalization of domain A which inhibits protein synthesis, increasing the injured cell apoptosis [10]. Furthermore, the E. coli toxins enhance functional tissue factor expression, causing vascular endothelium damage [11]. The damaged endothelium promotes microvascular thrombus creation [12], platelet consumption [8], and RBC lysis [13] (Fig. 2). Gb3 expression is predominant in the glomerular endothelium and in the intestinal endothelial layer. The remarkable Gb3 expression has encouraged scientists to postulate the *E. coli* gastroenteritis tissue tropism hypothesis. This hypothesis is supported by the simultaneous damage of intestinal and glomerulus endothelium following E. coli infection, especially that produces type II toxin [14].

Atypical HUS (Figs. 1-3)

The incidence of atypical HUS is 0.23-0.43 cases/10⁶/year. The primary underlying cause of the atypical type of HUS is complement dysregulation [15]. It is alleged that the atypical HUS pathogenesis depends upon the disproportionate complement activation by the alternative pathway. The excessive activation of complement is due to built-in regulators, which result from failure to disable C3b deposition on target cell membrane. Complement product deposition injures vessels' endothelial surface, promoting white blood cell (WBC) and thrombocyte migration and activation, resulting in the creation of thrombit that obstruct blood vessels. The renal impairment that occurs in atypical HUS is primarily due to abnormal complement product depositions in kidney microvasculature, thrombus formation, inducing renal ischemia, and renal impairment [16].

Loss of regulatory protein function is either due to autoantibody formation or genetic mutations. Mutations of some of complement systems factors such as complement factor H (CFH), complement factor I (CFI), membrane cofactor protein (MCP), and complement factor (CF)-3 are found in 40–60% of atypical HUS patients [17], increasing the occurrence rate of atypical HUS. Patients with genetic mutations, infections, autoimmune conditions, drugs, malignancies, or pregnancy are the usual triggers [15,18].

In atypical HUS patients, reported data declared that a mean platelet count usually reduced to 66×10³/μL [19]. Furthermore, significant renal insufficiency may necessitate dialysis in some cases [19]. TTP cases frequently show either minor or major neurological features; however, atypical HUS could exhibit minor neurologic manifestations [20]. Although the complement C3 product is usually low in serum, C4 is commonly normal; however, they are not diagnostic [21,22].

Atypical HUS due to gene mutations of CFH, CFI, CFB, C3, MCP, CDKE, and THBD CFs can be confirmed by gene sequencing; however, it is not essential to be conducted [22]. Gene studying is expensive, and the results take weeks, delaying the required quick, essentially needed therapy. Approximately half of atypical HUS cases have complement gene mutations or formed autoantibodies [22].

The main differences between typical and atypical HUS pathogeneses began when a link was detected between atypical HUS and the existence of gene mutations, altering complement regulator factor H function in the plasma [23]. Other mutations of complement regulators and CFs such as C3, factor B, factor I, and CD46 were detected recently [24]. A few cases often present rare polymorphism or more than one mutation [25]. The clinical severity of both atypical and typical HUS is almost similar, with a death rate of about 5% [26].

Secondary HUS

Secondary HUS is accompanied by coexisting diseases. Streptococcus pneumoniae and influenza virus infection are the

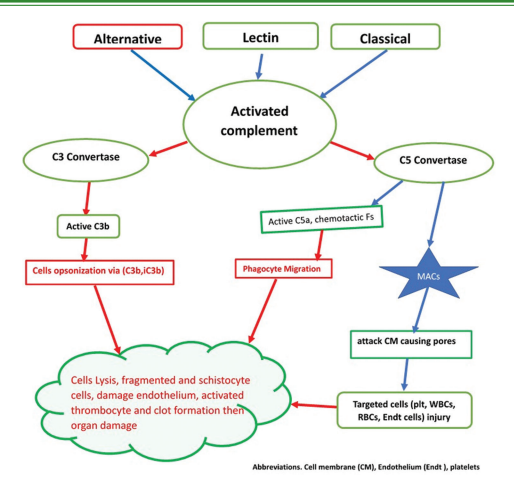


Figure 2: Complement system activation and cell lysis steps

frequently noted organisms in clinically manifested secondary HUS [27,28]. However, these infections may induce HUS, though they are not triggers of secondary HUS. Differentiation between them to be either trigger or cause is not clear. The pneumococci and influenza A contain active neuraminidase enzymes dedicated to removing sialic acid from cell surface [29,30], causing RBC lysis following complement activation by stimulating alternative pathway [31].

PATHOGENESIS OF HUS

Complement Activation Pathways (Figs. 2 and 3)

Complement activation is a body protective mechanism against infection and disease risks by phagocyte activation or direct lysis. Phagocytosis is enhanced by opsonization in the presence of C3b or its fragments. In addition, phagocyte migration is due to chemotactic and peptides such as C5a synthesis and release (Fig. 2) [16]. Direct target cell lysis by complement via creating membrane pores due to membrane assault complexes (MACs) lysis action (Fig. 3) [32,33]. Besides, it is claimed that complement might initiate and promote adaptive immunity [33].

Complement activation is accomplished via an alternative, lectin, and classical pathways. The alternative path is characterized by persistent low-level C3b on surfaces that have contact with plasma [16]. In abnormal complement activation via the alternative pathway, C3 convertase becomes active,

promoting excessive C3b synthesis. The produced C3b initiates cell lysis in the presence of factor B of the cells with no sialic acid glycosaminoglycans on their surface membrane. In addition, the C3 convertase activates the C5 factor (to C5b), promoting WBC sensitization and migration. In addition, C3 convertase promotes MAC formation, initiating cell member pore formation and severe cell damage, direct cell lysis, and cell death [32,33]. Cell opsonization and phagocytosis increase due to C3b and iC3b deposition on the surface of target cells (Fig. 3). In addition, complement initiates and augments adaptive immunity [33].

Endothelial Injury, Leukocyte, and RBC Lysis Mechanism in HUS (Fig. 3)

Complement alternate pathway activation causes deposition of complement assault complex on vascular endothelium cells, RBCs, and leukocytes. Unlike leucocytes and endothelial layer cells, RBCs are probably more prone to fast death due to a lack of cell membrane mending competence, although it is not well documented yet [34]. In atypical HUS, there are two potential mechanisms for RBC destruction, forming RBC fragments and schistocytes [35]. Due to the narrowed vessels, mechanical RBC lysis might occur; however, narrowed capillaries are thought not the cause of RBC lysis or schistocyte formation in HUS in an experimental setting [16]. One of the major weaknesses of this argument is that normal RBCs are usually deformable;

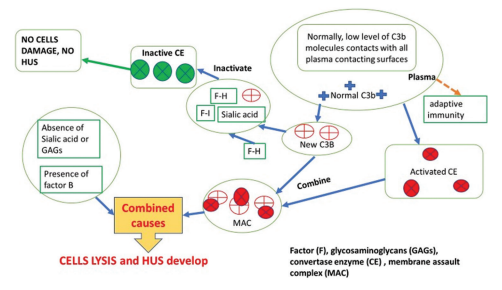


Figure 3: Atypical hemolytic uremic syndrome and alternative pathway

however, RBCs with diameters >3 µm are not easily passed through the microvessels with diameters < 0.3 µm [36]. Secondly, complement-RBCs-mediated lysis is potentially possible; however, the fundamental problem is that the fragmented RBC and schistocyte formation is not a typical feature of RBC destruction in paroxysmal nocturnal hemolytic anemia [37]. Another mechanism involved is possibly due to complement C3d and C3b factor deposition, causing RBC cell membrane stiffness [38]. Therefore, more research is needed to determine what causes the noticeable hemolysis in HUS patients with complement regulatory abnormalities.

Blood vessel endothelial cells and WBCs are more efficient than RBC at removing CF deposits and MACs by internalization or scaling [39]. WBC and endothelial cells possibly have a higher ability to repair their damaged membranes than RBCs; however, a histological study of tissues biopsied from patients with acutestage HUS demonstrated endothelial cell destruction [40]. Endothelial cell membrane injury and death in cases of typical HUS seem due to the Shiga toxins' direct effect. In contrast, the damage of the endothelial cells in atypical HUS is assumed due to lytic or sublytic assault by complement [41]. Other mechanisms via nitric oxide, activated neutrophils and monocytes, released reactive oxygen, free hemoglobin, cytokines, procoagulants, and cellular hypoxia may all induce endothelial cell injuries, sharing in atypical HUS pathogenesis.

Complement Activation and Cell Damage in HUS (Figs. 1 and 2)

However, the damaged endothelium may activate the complement cascade [42], and complement activation can happen via any of the three known mechanisms. In atypical HUS, the alternative pathway is the prim activation pathway [43]. Injuries induced by complement activation increase and can cause more damage to surrounding cells, both those in the damaged vicinity and tissues that the complement products pass through [44]. Endothelial cell injury is assumed the initial step in the pathogenesis of both typical

and atypical HUS [45]; nevertheless, it is unknown which type of cells is injured first. Complement system activation is further promoted by platelet activation [46] and the released hemoglobin from hemolyzed RBCs [47]. These observations had led to a conclusion that in atypical HUS, the damaged RBCs and the other cells may occur simultaneously or even before endothelial cell damage; however, this is unproven yet [48]. Further studies are encouraged to explore this conclusion.

Platelets Effect in Atypical HUS Pathogenesis

Platelet activation mediated by complement has not been extensively investigated in atypical HUS, such as hemolysis and endothelial cell injury. It appears that platelets may have an essential role in atypical HUS pathophysiology, either directly or indirectly, because of low platelet count [49] and the thrombi containing many platelets [48] are hallmarks of this HUS type. In addition to RBCs, platelets are easily prompted by complement attack [50] via the creation of MACs on their membrane [51]; however, platelets have the power to internalize or shed the complement MACs [52]. It is reported that cell membrane sialic acids protect platelet injury via factor H from the destructive action of the complement [48]. Factor-H and membrane regulators' combined action protects thrombocytes against complement effect, but this protection is usually not complete in atypical HUS [16]. New research projects are required in this area to clarify the effect of complement on platelets and thrombus formation.

Coagulation-Associated HUS

Different reports relating complement activation and induction of coagulation have concluded that complement activation causes the initiation and increased coagulation rate. Coagulation can be triggered by complement in two ways. Complement first damages tissue by forming MACs on endothelial layer cells, affecting vessels' endothelium integrity. Second, complement activation

causes active C5 formation and release or creation of C5b-9 complex, activating endothelial cells and causing procoagulative tissue factor expression [53]. It is reported that various molecular interactions between the activated CFs and coagulation proteins [54]. However, the physiological significance of these interactions has not been undoubtedly established. The vital initiative appears due to thrombocytes' activation by MACs [50] or C5a [55]. MACs appear to have a procoagulant impact, mediating platelet prothrombinase rather than causing permanent cell destruction [56,57]. Regardless of the cause of thrombocyte activation, it is widely accepted that activation and aggregation of platelets are intimately ended to coagulation, forming another link between complement activation and coagulation process initiation. Despite the abovementioned possible mechanisms of thrombosis, the real mechanism(s) is (are) not clear yet. Further projects are needed to investigate this topic.

HUS THERAPY (FIG. 1)

Acute HUS management is a collaborative achievement between most medical and surgical specialties, especially the transplant teams. The medical teams usually required in acute HUS therapy are internal medicine, intensive care, nephrology, hematology, and neurology. In the acute stage of HUS, full supportive management is the prim therapy. Supportive therapy includes maintaining good fluid hydration, and normal electrolyte balance is essential. It is also vital to maintain controlled blood pressure values, using renin-angiotensin blockade in typical HUS cases and in those who have chronic kidney disease. In 20-40% of HUS patients, especially those with atypical HUS, neurological manifestations such as seizures occur. To avoid severe trauma and brain damage complications, seizure prevention and medical management of the attacks are essential. Dialysis may be indicated in severe azotemia. Treatment was initially similar for all the HUS types. Although atypical HUS in adult cases is treated ironically by plasma exchange, currently, eculizumab and ravulizumab are approved for adult atypical HUS cases. Plasma exchange principally is the first treatment choice in TTP therapy. In contrast, plasma exchange therapy in cases of atypical HUS is less impressive [15]. Furthermore, when the ADAMTS13 activity is normal as expected in most atypical HUS cases, eculizumab therapy should be tried first, and then plasma exchange is advisable when there is no improvement [18]. Antimicrobial therapy is not indicated, except in severe determined sepsis [58].

Treatment of Typical HUS

There is reasonable evidence that antibiotic coverage is unnecessary unless there is well-documented evidence of sepsis [58]. In an *in vitro* study, initiating antibiotic coverage may inhibit bacterial toxin synthesis to some extent; however, inhibiting DNA synthesis by some antibiotics such as trimethoprim-sulfamethoxazole and ciprofloxacin may increase Shiga toxin formation [59]. In contrast, antibiotics that target organisms' cell walls, DNA transcription, or translation such as azithromycin do not enhance toxin formation.

On the contrary, toxin levels diminished significantly following azithromycin, despite that *E. coli*-O157:H7 type may even remain highly viable. It is not recommended to use plasma exchange in typical HUS management.

In the acute stage of HUS, therapies such as plasma therapy, intravenous globulin, fibrinolysis, antiplatelet, corticosteroids, and even antioxidants have been demonstrated unsuccessful in controlled clinical trials [60]. It was reported that kidney transplantation is advisable in HUS patients, especially children who have a low incidence of end-stage renal failure recurrence rate (0–10%).

Treatment of Atypical HUS

Eculizumab and ravulizumab are approved monoclonal antibodies to treat atypical HUS cases. They prevent complement-mediated microangiopathic thrombus formation. However, it must be emphasized that these two agents are not recommended in patients who had no meningococcal vaccine at least 2 weeks prior to their administration, especially ravulizumab. Plasma exchange must be considered early in atypical HUS therapy (within the earliest 24 h of presentation) and should be continued at least 2 days after complete remission. One or two plasma exchange sessions must be conducted to remove toxins quickly because it reduces mortality significantly. Although plasma infusion may be beneficial in heart or renal failure cases, plasma exchange is preferable because they are more efficient and effective in removing toxins [61]. Intravenous plasma therapy is not indicated HUS induced by S. pneumoniae, while it can aggravate the disease due to the presence of Thomsen-Friedenreich antigen, which stimulates antibody formation by the adults' plasma. Only one case report has described a reasonable response to a high plasma infusion dose (30 mL/kg/week) over 2 years and a half; however, long-term responses are still under evaluation [60].

Treatment of Secondary HUS

Supportive treatment and controlling the coexisting diseases are the mainstay of this type of HUS. Plasma infusion and plasma exchange can be effective, although they are not commonly conducted.

Eculizumab

Eculizumab is a monoclonal antibody that inhibits C5b-9 synthesis to suppress complement terminal component activation. Eculizumab therapy improved kidney parameters, increased platelets, and reduced the need for hemodialysis in cases that had not responded to plasma therapies. A prospective phase II study reported that non-Shiga toxin-induced HUS cases aged ≥12 years noted a significant improvement in glomerular filtration rate (GFR) following early eculizumab administration. Furthermore, it was reported that the insufficient response to plasma infusion or exchange in patients who improved following eculizumab had supported its effectiveness in treating atypical HUS cases [60]. In phase III prospective study, 41 patients with atypical HUS aged ≥18 years, approximately 75% had normal thrombocyte count and

serum lactate dehydrogenase and preserved their kidney function. In addition, life quality improvement, transplant protection, and dialysis withdrawal were reported [22]. However, the number of patients treated with eculizumab was not large enough to draw a firm conclusion; it has been reported that eculizumab has no significant effect in some cases of atypical HUS, particularly those with DGKe mutation [62] or cobalamin C deficiency in adults [63]. However, the number of cases treated was little.

Withdrawal of eculizumab therapy after a mean of 16.5 months of follow-up in 36 atypical HUS cases increased the relapse rate in female patients and in cases with MCP, CFH, and CFI complement gene mutations [64], but the need for dialysis decreased in the cases with recurrence of the atypical HUS. The same study reported that 11 of the relapsed patients had recovered to the basal renal function after resuming eculizumab. Unfortunately, 2 cases had deterioration of their increased basal kidney function values, and one patient developed chronic renal and became dialysis dependent [64]. Fakhouri et al. (2016) studied eculizumab which has been given to 41 cases with atypical HUS; platelet count normalized in 98% at a median of 8 days. After 26 weeks, the same study reported that 88% of the cases had sustained normal thrombocyte count and lactate dehydrogenase values, and the estimated GFR improved by ≥15 mL/min/1.73 m2 in 54% of eculizumab treated patients [22].

Ravulizumab

Ravulizumab is another approved monoclonal antibody that blocks CF C5 activation, preventing its cleavage to C5a and C5b, hindering terminal complement complex creation, thereby preventing RBC hemolysis. Ravulizumab is approved for about 30 months for atypical HUS therapy. In two studies, ravulizumab improved HUS completely in 54% of adult atypical HUS cases within the first 26 weeks. Ravulizumab normalized platelets, LDH in 54% of adult cases of atypical HUS, and improved serum creatinine by 25%. Furthermore, ravulizumab therapy has improved thrombocytopenia in 84%, hemolysis in 77%, and improved kidney parameters in about 60% of adults [65]. The two studies showed a higher response rate to ravulizumab in children with atypical HUS than in adult cases.

Transplantation

Membrane-bound protein is highly present in the kidney tissue, and kidney transplantation improves renal function, improves the outcome, and improves the local level of membrane-bound protein expression that helps in dysfunction improvement and the level of membrane-bound protein expression. Kidney transplantation is not a treatment option of either atypical or secondary HUS, while the failure rate is >90%, and these HUS-type recurrences happen in 50%. The recurrence rate in atypical HUS ranges between 30% and 100%, and it is higher in cases with HF1 mutations than in ones with no mutation.

The liver synthesizes HF1 protein factor; hence, in genetic defect cases of HF1, it was expected that liver transplantation

might improve the outcome. This assumption has encouraged two children to have simultaneous renal and hepatic transplantation, but unfortunately, early liver failure has occurred. This has led to conducting this procedure only for cases that require it as a life-saving measure.

PREVENTION

Informing health authorities about typical HUS as soon as possible is required since it occurs in epidemics, monitoring the probability of index cases, and imposing preventive measures that reduce disease outbreaks. The only available measure to decrease morbidity and mortality is disease prevention. Antibiotics have been proven ineffective in disease prevention and may augment the HUS risk, especially in children who developed *E. coli* O157:H7 infection. Therefore, antibiotics are not advisable in HUS except in severely septic patients [58].

CONCLUSION

The characteristic features of the types of HUS are synchronized small vessels' endothelial cell damage, intravascular RBC lysis, and platelet activation that initiates coagulation, causing small thrombi and tissue ischemic. In atypical HUS pathogenesis, abnormal complement activation via alternative pathway is the main underlying mechanism. Despite the recent evolutions in the HUS pathogenesis, some mechanisms are still not understood clearly and need further research projects.

HUS diagnosis on clinical features is always challenging due to the lack of characteristic universal diagnostic criteria, and it is a disease of exclusion. However, gene mutation determination is helpful in atypical HUS diagnosis, but they are expensive and time-consuming and may delay the urgently required therapy. Prevention of HUS is the best therapy option, especially in non-typical HUS cases. Supportive management and conjunction work between the whole managing teams are highly recommended. Plasma exchange and monoclonal antibodies are the best for anti-FH-positive HUS therapy.

AUTHORS' CONTRIBUTIONS

Habas E aided in conceptualization, methodology, literature review, manuscript preparation, critical review, and revisions of the manuscript. Rayani A, Habas A, Farfar K, Habas E, Elmarghani A, and Elzouki A aided in the literature review, manuscript writing, critical review, and revisions of the manuscript. All the authors reviewed and approved the final version of the manuscript.

CONFLICT OF INTEREST

All authors have no conflict of interest.

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