



Thrombotic microangiopathy in children

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Abstract

The syndrome of thrombotic microangiopathy (TMA) is a clinical-pathological entity characterized by microangiopathic hemolytic anemia, thrombocytopenia, and end organ involvement. It comprises a spectrum of underlying etiologies that may differ in children and adults. In children, apart from ruling out shigatoxin-associated hemolytic uremic syndrome (HUS) and other infection-associated TMA like *Streptococcus pneumoniae*-HUS, rare inherited causes including complement-associated HUS, cobalamin defects, and mutations in diacylglycerol kinase epsilon gene must be investigated. TMA should also be considered in the setting of solid organ or hematopoietic stem cell transplantation. In this review, acquired and inherited causes of TMA are described with a focus on particularities of the main causes of TMA in children. A pragmatic approach that may help the clinician tailor evaluation and management is provided. The described approach will allow for early initiation of treatment while waiting for the definitive diagnosis of the underlying TMA.

Keywords Thrombotic microangiopathy · Complement · DGKe · Hypertension · Infection · Hemolytic uremic syndrome · Transplant

Introduction

The assessment of an adult patient presenting with thrombotic microangiopathy (TMA) involves work-up for thrombotic thrombocytopenic purpura (TTP) due to deficiency of ADAMTS13 (a disintegrin and metalloproteinase with a thrombospondin motif type 1, member 13), shigatoxin-induced hemolytic uremic syndrome (STEC-HUS), and secondary causes of TMA (infections, drugs, malignant hypertension, autoimmune diseases, and others). If other causes

of TMA have been excluded, then complement-mediated/atypical hemolytic uremic syndrome (CM-HUS or aHUS) should be considered [1–3].

Although there is an overlap of the etiologies causing TMA in both adults and children, some of the diseases are more common in children and other rarer conditions are seen only in children. The main cause of TMA in children is STEC-HUS, followed by CM-TMA and *Streptococcus*-induced TMA (Sp-HUS) [4]. In addition, there are rare conditions such as congenital TTP, vitamin B12 metabolism defects (cobalamin C and G defects), and coagulation disorders (diacylglycerol epsilon mutation or variant *DGKE*) that may manifest as TMA, making the process of differential diagnosis more complex, especially in children younger than 2 years of age. Solid organ and hematopoietic stem cell transplantation (HSCT) may also present with TMA in children and may pose a challenge in differential diagnosis.

Although there are registries of specific diseases such as TTP and aHUS, TMA registries of young children are lacking. In a recent paper from Germany, among 232 patients with TMA, 61% presented with aHUS [5]. In a Brazilian aHUS cohort [6], among the 17 patients who presented with their first manifestation when less than 18 years old, 53% were younger than 2 years of age and one patient who was

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refractory to treatment with eculizumab had a homozygous mutation in *DGKE* [7].

The possibility of various treatment strategies of these conditions underscores the need of a tailored diagnostic approach (Box). This review aims to discuss the causes of TMA—acquired and inherited (Fig. 1)—in children, and propose a practical algorithm to the diagnosis and management of TMA in this population (Fig. 2).

Box—Checklist for diagnosis and evaluation of TMA in children

Identifying the presence of TMA

- Complete blood count
- Peripheral smear for evaluation of schistocytes
- Reticulocyte count
- Bilirubin
- Lactic Dehydrogenase
- Haptoglobin

Evaluating the degree of organ injury

- Complete physical exam including blood pressure and neurologic assessment
- Kidney function: BUN, serum creatinine
- Urinalysis and proteinuria
- Liver enzymes
- Pancreas enzymes: amylase, lipase
- Blood bicarbonate and lactic acid levels
- Troponin

Defining underlying cause (request according to clinical suspicion)

- Shigatoxin in stools
- ADAMTS13 activity and inhibitor
- Blood complement levels
- Cultures: blood, urine, CRL, empyema
- Latex test for *Streptococcus pneumoniae*
- Tests for H1N1 and COVID19
- Other infections according to epidemiology: dengue, leptospirosis
- Homocysteine and vitamin B12 blood levels
- Urinary amino acid chromatography
- Genetic testing: complement panel or Whole Exome Sequencing
- Anti-factor H antibody
- Kidney biopsy when secondary cause suspected or need to evaluate chronicity

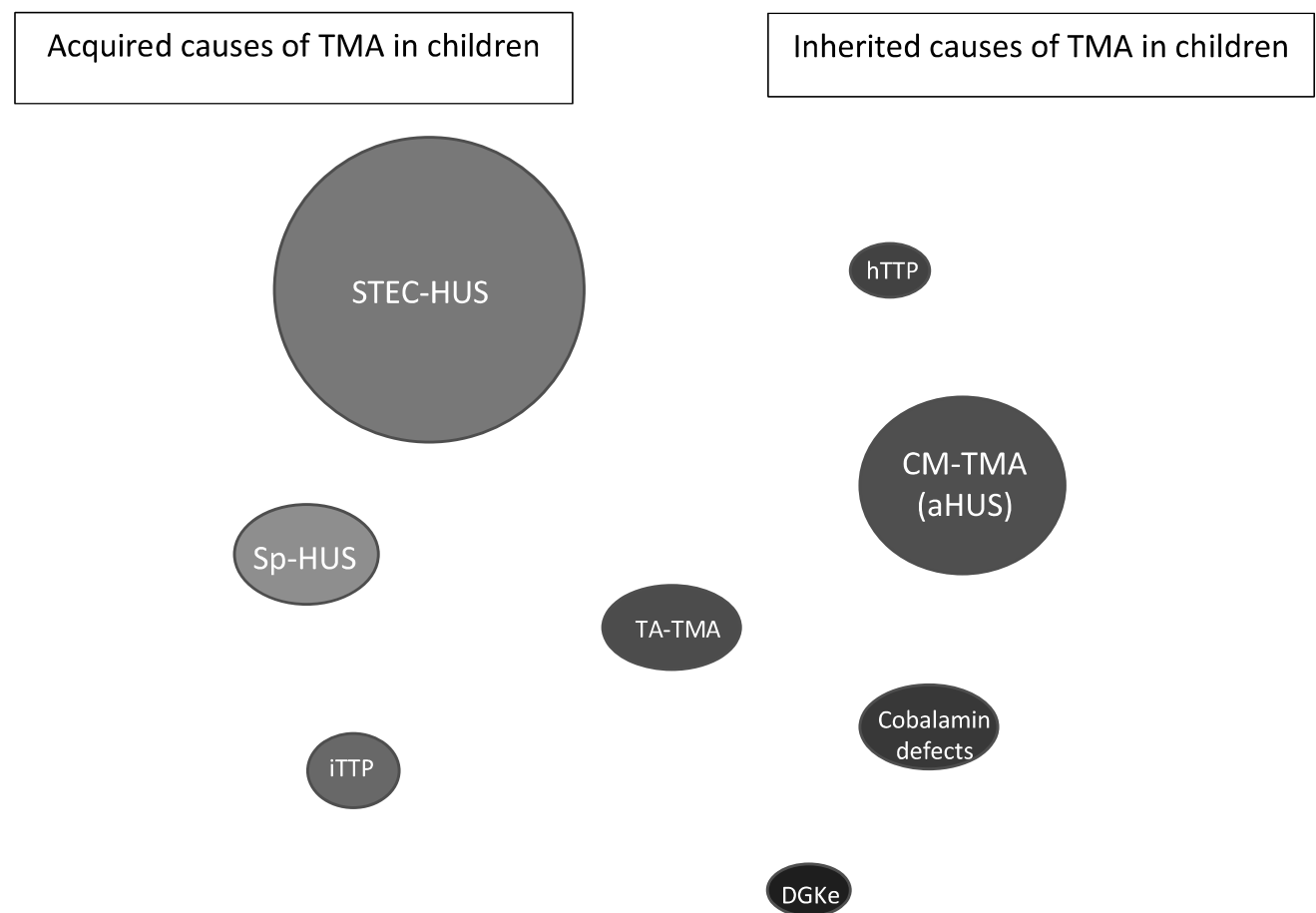


Fig. 1 Acquired and inherited causes of TMA in children. The size of the balloons is proportional to the incidence of each disease. *STEC-HUS* shigatoxin-associated hemolytic uremic syndrome, *Sp-HUS* *Streptococcus pneumoniae* hemolytic uremic syndrome, *DGKe* dia-

cyglycerol kinase epsilon, *CM-HUS* complement-mediated hemolytic uremic syndrome, *iTTP* immune-mediated thrombotic thrombocytopenic purpura, *hTTP* hereditary thrombotic thrombocytopenic purpura, *TA-TMA* transplant-associated TMA

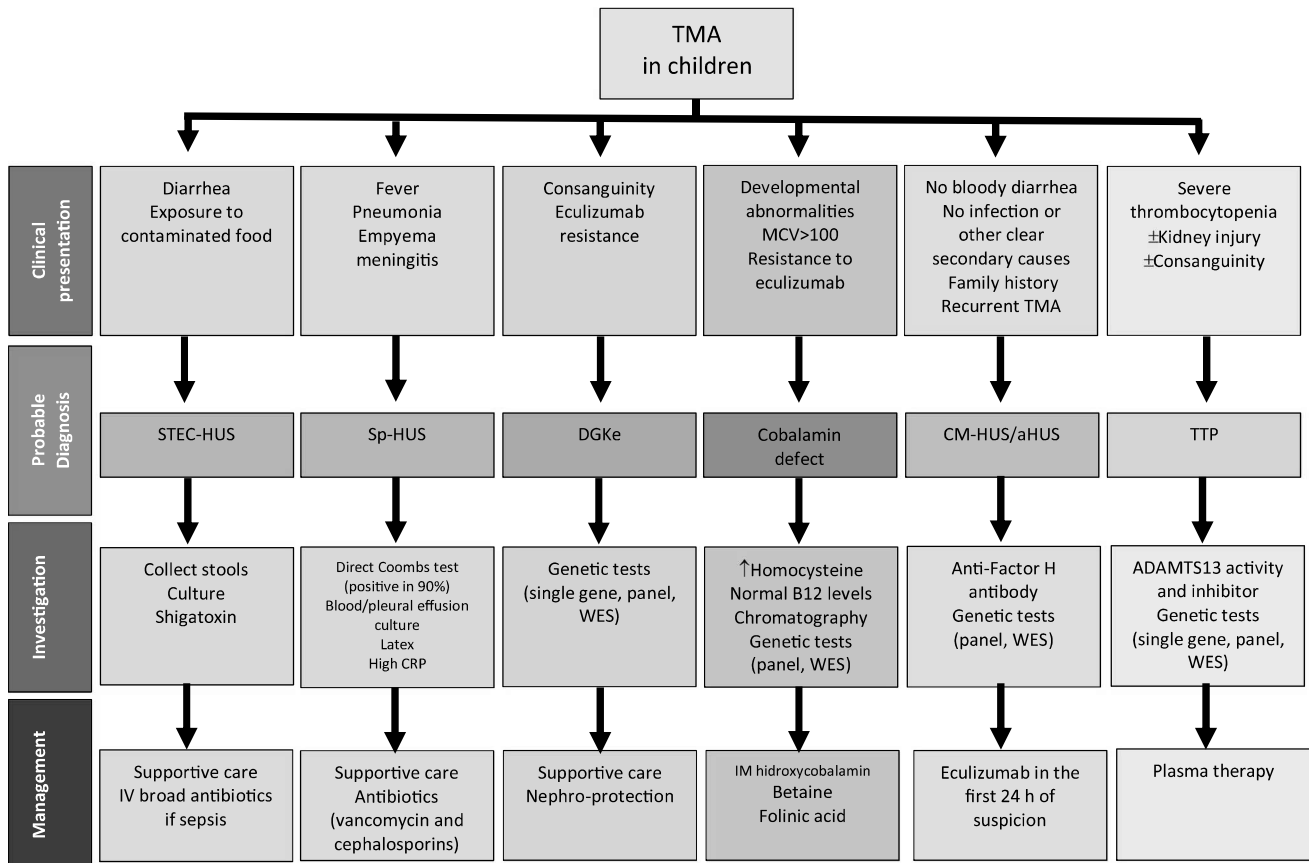


Fig. 2 Differential diagnosis of thrombotic microangiopathy in children: singularities of clinical presentation, tools for investigation and a pragmatic approach to management. *MCV* mean corpuscular volume, *STEC-HUS* shigatoxin-associated hemolytic uremic syndrome, *Sp-HUS* *Streptococcus pneumoniae* hemolytic uremic syndrome,

CRP C-reactive protein, *DGKe* diacylglycerol kinase epsilon, *CM-HUS* complement-mediated hemolytic uremic syndrome, *TTP* thrombotic thrombocytopenic purpura, *WES* whole exome sequencing, *ADAMTS13* a disintegrin and metalloprotease with thrombospondin type-1 repeats, 13th member, *TA-TMA* transplant-associated TMA

BUN blood urea nitrogen; *CRL* cephaloraquidic liquid; *ADAMTS13* a disintegrin and metalloprotease with thrombospondin type-1 repeats, 13th member; *H1N1* quick test to detect influenza strain; *COVID19* antigen or antibody tests to detect current infection by SARS-CoV-2 virus

Shigatoxin-associated hemolytic uremic syndrome

Incidence and epidemiology

STEC-HUS is one of the most common causes of acute kidney injury in children [8, 9]. The disease was first reported in 1955 by Gasser et al. [10]. Diarrhea associated with shigatoxin-producing *Escherichia coli* is estimated to occur in 2.8 million cases/year worldwide and the incidence of STEC-HUS varies from 0.5 cases/100,000 population in Finland [11] to 12 cases/100,000 population in Argentina, where it is endemic [12]. Seasonal variations occur, with a

greater number of cases in the summer. It is more common in children under 5 years of age and *E. coli* O157 is the most common strain producing shigatoxin in this age group in the USA [13]. In a cohort from Finland [11], age under 3 years was a risk factor for HUS in STEC-positive children. In 2011, the largest outbreak to date occurred in Germany with almost 4000 cases of diarrhea due to *E. coli* O104, with 845 cases of STEC-HUS and 54 deaths. There were 90 cases in children with one death and the outcomes were similar to previous case series, although mean age was higher (most children were older than 10 years) [14]. In Europe, non-O157 serotypes of *E. coli* have become more common [15]. In a prospective study in northern Italy, incidence of STEC-HUS among patients with shigatoxin-positive bloody diarrhea stool tests was 15.9% [16].

Clinical presentation and diagnosis

STEC-HUS is characterized by platelet consumption followed by hemolysis and acute kidney injury. Clinically, a

prodrome of profuse, bloody diarrhea occurs about 2 to 5 days after STEC infection. The signs and symptoms of microangiopathy occur a few days after diarrhea, but only in 10 to 20% of patients. Extrarenal manifestations may involve central nervous, cardiovascular, gastrointestinal (including pancreas), and musculoskeletal systems [17]. In addition to the platelet consumption, hemolysis, and kidney injury that characterize TMA, there may be a decrease in the blood level of component 3 (C3) of the complement. In rare cases, a decrease in C4 occurs. This finding is usually present in the most severe cases [18, 19] and is temporary—and may be due to a secondary involvement of the complement system in vascular inflammation [20]. Therefore, the presence of low C3 in patients with TMA does not exclude the possibility of STEC-HUS [21]. Increase in inflammatory markers such as C-reactive protein and D-dimer is frequent in patients with STEC-HUS [22]. It is essential that the collection of fecal material for stool culture or PCR for shigatoxin be done early on admission since positivity decreases after the 10th day of diarrhea, and after 15 days 90% of patients present with negative stool cultures in a prospective cohort from Argentina [23]. We reiterate that it is essential to collect material for shigatoxin in all cases of TMA and to discuss with local health authorities the availability of sending material for the detection of shigatoxin in feces in specialized laboratories. When available, IgA, IgG, and IgM antibodies to *E. coli* lipopolysaccharide (LPS) can be detected later in the course of the disease especially when stool specimens fail to detect the presence of shigatoxin [15]. Specifically for *E. coli* O157 serotype, the Glyco-iELISA to detect IgM antibodies has significantly increased accurate diagnosis of STEC-HUS [24].

When recovery of STEC-HUS extends beyond 2 weeks, the differential diagnosis with atypical HUS should be considered. Clinical and laboratory findings that make STEC-HUS less likely [84]:

- Negative stool culture or PCR-shigatoxin in feces (harvested up to 10 days after diarrhea)
- Persistent thrombocytopenia beyond the first week in the absence of a superimposed infectious condition
- Relapsing pattern of TMA after resolution of initial manifestation
- Kidney injury that persists for more than 4 to 6 weeks in the absence of superimposed septic or hypovolemic shock
- Absence of diarrhea and/or negative shigatoxin test
- Persistently low C3

In these situations, the main differential diagnosis, especially in the pediatric age group, is aHUS [4], which reinforces the importance of early collection of specimens for shigatoxin or stool culture (where shigatoxin is not available) at the admission of ALL patients with TMA.

Genetic variants in alternative complement genes have been reported in patients with STEC-HUS [25] and should be suspected in patients who progress to kidney failure or have recurrent TMA or persistently low C3.

Pathophysiology

Karmali et al. [26] were the first to describe the association between gastrointestinal infections by shigatoxin-producing *E. coli* (Stx; verotoxin) and STEC-HUS. Since then, these strains have been described as a major cause of STEC-HUS in both sporadic cases and in outbreaks in many countries, usually associated with contamination of water, vegetables, beef products, and other foods [9]. HUS–*E. coli* strains produce shigatoxin 1 (Stx1) and/or shigatoxin 2 (stx2). Shigatoxins are 70 kd holotoxins that contain a single A subunit and 5 B subunits. Cell damage results when the B subunit recognizes and binds to globotriaosylceramide (Gb3 or CD77), which resides in the plasma membrane of certain eukaryotic cells including the kidney and brain endothelial cells. Cellular lesions were recently described by Volokhina et al. [27], although the exact mechanism of injury is still unknown. The toxin–Gb3 complex is internalized and enters the cytoplasm, where subunit A is synthesized. Cellular apoptosis results from Gb3 binding with endocytosis, retrograde transport, cytosolic translocation of shigatoxin, and consequent ribosomal inactivation, which leads to endothelial damage, exposure of the subendothelial layer and consequent TMA. The toxin also acts on cell activation, pro-inflammatory and pro-thrombotic pathways, facilitating thrombosis by von Willebrand factor endothelial secretion [8].

Management

There is no specific treatment for STEC-HUS. Clinical care involves supportive measures with a focus on stabilizing the patient to natural resolution and recovery. Early volume expansion has been demonstrated to improve neurological and kidney outcomes of patients with STEC-HUS (weight gain of 12.5% vs. 0%) [28]. A systematic review has shown that no therapy tested to date has shown a greater benefit than supportive treatment [13]. An analysis of the German outbreak—the biggest to date that happened in 2011 with more than 3000 cases of diarrhea and 800 cases of STEC-HUS [29, 30]—showed that the average duration of diarrhea was 9.3 days (median 6 days) and most patients were admitted to the hospital in the first week after the onset of diarrhea. In the German outbreak, plasmapheresis [31] and corticosteroids showed no benefit, as well as the use of the terminal complement blocker eculizumab (Soliris, Alexion Pharma) [32], although results from clinical trials with eculizumab in STEC-HUS are still awaited. For a comprehensive

review of eculizumab in STEC-HUS, we recommend the paper from Walsh and Johnson [33]. More recently, a case report described good outcomes with pulse steroids in a 7-year-old patient with STEC-HUS and late neurologic involvement [22, 34]. Fosfomycin used in the first 2 or 3 days after onset of diarrhea may be beneficial in preventing the evolution to STEC-HUS [35]. In the evolution of STEC-HUS, platelet count was the first to improve (end of the first week), followed by improvement of the hemoglobin and slow drop in lactate dehydrogenase. The acute kidney injury took an average of 2 weeks to recover, starting with a return of diuresis and then a fall in blood urea nitrogen and creatinine. Around 50% of patients required kidney replacement therapy and only three out of 160 patients were dependent on chronic dialysis. However, the kidney injury may be longer in very severe cases (LMPP and MHVG have seen cases that took up to 5 weeks before improvement in kidney function). Approximately 30% of patients with STEC-HUS continue to have variable degrees of neurological or kidney sequelae [13]—proteinuria and systemic arterial hypertension. Therefore, it is recommended that every patient with STEC-HUS be followed by a nephrologist or pediatric nephrologist after hospital discharge. There may be a role for probiotics in the management and prevention of STEC-HUS [36], but further studies are warranted.

Complement-mediated, atypical hemolytic uremic syndrome

Incidence and epidemiology

Atypical hemolytic uremic syndrome (aHUS) is a cause of TMA driven by an alternative complement pathway dysregulation. More recently, the term complement-mediated hemolytic uremic syndrome (CM-HUS) has been advocated since complement overactivation has been found in conditions otherwise considered secondary TMA (as in malignant hypertension, for instance), and this may have important therapeutic implications [37]. For the purposes of this review, aHUS is used for cases of CM-TMA without any other underlying secondary condition (i.e., primary CM-TMA). It is classified as an ultrarare disease with an incidence of less than 1 case per million population per year [38]. Up to 50–60% of patients present with a genetic defect or autoantibody leading to overactivation of alternative complement pathway, but positivity of genetic findings is variable worldwide—from 18% in a cohort from India [39] to 80% in Japan [40]. Progression to kidney failure or death had a significant decrease after introduction of terminal complement inhibitor eculizumab: from 30–50 to 9% in children, and from 56–67 to 6–15% in adults [41].

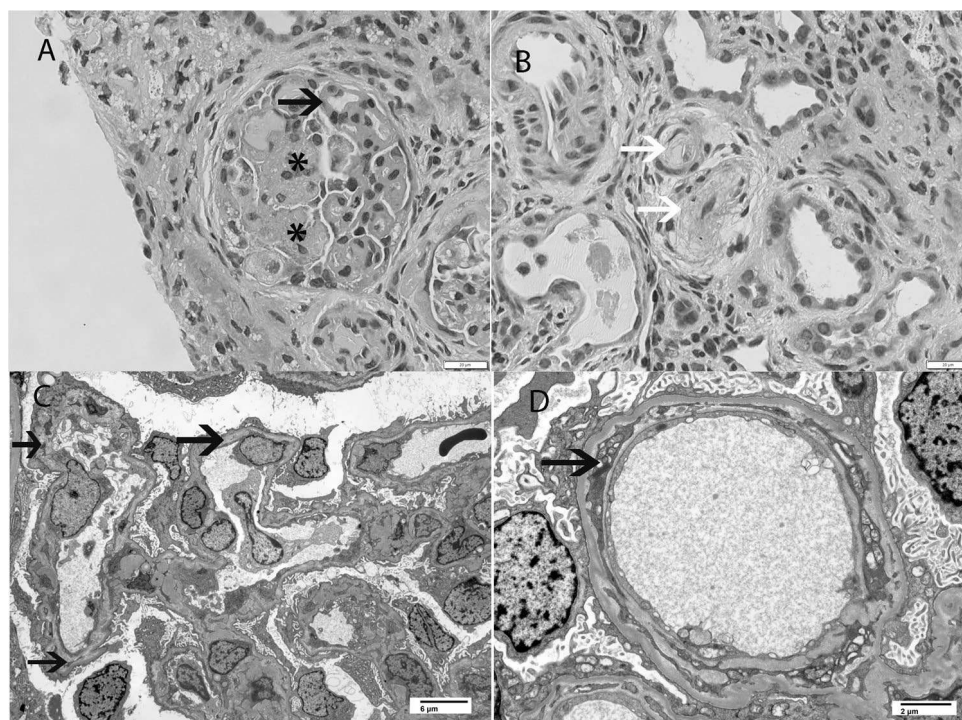
Clinical presentation and diagnosis

aHUS is characterized clinically by the TMA triad: non-immune microangiopathic hemolytic anemia, thrombocytopenia or a significant (> 25%) drop in platelet count, and evidence of organ injury in variable degrees. Since there is no gold standard diagnostic test for aHUS, it is important to rule out other more common causes of TMA [42], which is a challenge in a disease with high morbidity and mortality [43]. One of the key questions in aHUS is the correlation of genetic findings with clinical presentation. In the great majority of patients, genetic variants present in heterozygosity with incomplete penetrance, frequently associated with one or more risk polymorphisms, and an environmental factor may be necessary to result in full blown TMA clinical presentation. In 2007, the French Society of Pediatric Nephrology published data of 46 children with aHUS and found that 52% had a mutation, with *CFH* mutation carrying a worse prognosis and poor transplant outcomes [44]. In 2010, data analysis from 273 patients enrolled in the International Registry of Recurrent and Familial HUS/TTP concluded that adults had a worse prognosis than children, as well as familial cases when compared to sporadic cases. Overall, 75% of patients presented either a mutation or a risk polymorphism, but there was an ethnicity-specific genotype [43]. Bresin et al. [45] found that risk haplotypes in *CFH* and *MCP*, when combined with mutations, carry a worse outcome. More recently, data from The Global aHUS Registry with more than 800 patients enrolled confirmed that adults have worse kidney outcome than children, as well as patients carrying *CFH* pathogenic variants [46].

Pathophysiology

aHUS is a rare disease caused in most cases by a dysregulation of the alternative complement pathway (Fig. 3). The first association between aHUS and the alternative complement defect was described in 1981 [47], when siblings with aHUS and consanguineous parents were found to have decreased blood levels of CFH (Complement Factor H). In 1998, the team lead by T. Goodship [48] unraveled the association between aHUS and chromosome 1q32, which contains the genes for complement regulators. In the following years, defects in other components of complement regulators of the alternative pathway have been successively found—lack of function mutations in CD46 (MCP—membrane cofactor protein), CFI (complement factor I), thrombomodulin (THBD), CFHR1–5 (proteins related to factor H 1 to 5 and hybrid genes), CFP (complement factor properdin), and gain of function mutations in genes encoding C3 convertase components, CFB (complement factor B) and C3. The role of anti-factor H antibodies in the pathogenesis of aHUS, especially in children, was also demonstrated

Fig. 3 Complement-mediated TMA in a child less than 2 years of age. **A** Light microscopy showing glomeruli with mesangiolysis, double contouring of the capillary walls, and lobular accentuation of the glomerular capillary tufts (Periodic acid Schiff stain 60×). **B** Light microscopy showing arterioles with endothelial swelling and near complete occlusion of arteriolar lumen (Periodic acid Schiff stain, 60×). **C, D** Electron microscopy showing capillary wall injury with subendothelial expansion by fluffy granular material, injured cellular elements, and new basement formation resulting in double contours (**C**: 2500×, **D**: 8000×). Asterix indicates mesangiolytic, white arrows indicate arteriolar occlusion by TMA, and black arrows indicate double contour formation



and its detection has therapeutic implications [49]. It is very important to determine the underlying genetic status since it adds to prognosis and management. It is well known that patients with factor H pathogenic variants have worse outcomes and higher rates of post kidney transplant recurrence, and patients with MCP variants or high-risk polymorphisms rarely have aHUS recurrence after transplant, although they might experience recurrent bouts of TMA throughout their life [46, 50, 51].

Management

Besides understanding the pathophysiology of aHUS, determining the underlying defect of the alternative pathway is key to address specific treatment, although it is not necessary to have the result of genetic tests to start treatment. In September 2011, eculizumab (Soliris, Alexion) was approved by the Food and Drug Administration (FDA, USA) as the first terminal complement inhibitor for use in patients with Paroxysmal Nocturnal Hemoglobinuria (PNH) and aHUS. The outcomes of patients with aHUS treated with eculizumab have been published in pivotal trials both in adults [52–54] and children [55], and were extensively reviewed [56]. Almost a decade after the first published use of eculizumab for aHUS [57], the question is whether all patients need lifelong treatment and, if not, who would be eligible to stop. Results of a prospective study on discontinuation of eculizumab [58] in 57 patients from 22 centers in France were published. In this trial, patients were on eculizumab for at least 6 months and in remission

at the time of discontinuation. Importantly, patients should have (1) glomerular filtration rate (GFR) > 60 mL/min/1.73 m² (Schwartz or MDRD formulas for children and adults, respectively) and urinary protein/creatinine ratio (UPC) < 0.05; or (2) stable kidney function in the past six months. The relapse rate was higher in children than adults (31% vs. 19%, respectively) and in patients who carried a pathogenic genetic finding, except for one patient with a negative genetic test—a more detailed genetic analysis revealed a homozygous mutation in the *ADAMTS13* gene and plasma activity of ADAMTS13 was decreased. At the end of the second year of follow-up, patients who had aHUS recurrence evolved with greater proteinuria than those who did not relapse. Although the authors concluded that stopping eculizumab in controlled patients is safe, the risk of relapse needs to be discussed with patients and caregivers and eculizumab should be promptly available if necessary. In addition to eculizumab, a newer formulation of long-acting C5 inhibitor ravulizumab is now available for treatment of aHUS; data from phase 3 trials have demonstrated good efficacy and safety results in adult and pediatric patients [59, 60].

***Streptococcus pneumoniae*-associated hemolytic uremic syndrome**

Incidence and epidemiology

Streptococcus pneumoniae-associated hemolytic uremic syndrome (Sp-HUS) is an important cause of TMA in young children. It accounts for 5–15% of cases of HUS in children (1–3% of all causes of TMA) with increasing prevalence probably due to serotypes not covered by the vaccine [61]. The highest incidence occurs in children under 2 years of age [62] who have Sp-pneumonia/empyema (60%) or meningitis (30%). The incidence of pneumococcal infections that progress with HUS ranges from 0.4 to 0.6% [63]. Prior to 1990, there was a 50% mortality and 67% of the patients who survived progressed to chronic kidney disease (CKD) or hypertension. In 2007, mortality and progression to CKD or hypertension decreased to 12.3% and 16%, respectively, likely due to increased awareness and management in the intensive care unit.

Clinical presentation and diagnosis

The clinical presentation is that of classic TMA triad, occurring usually 7 to 9 days after the onset of pneumococcal infection. The differential diagnosis includes disseminated intravascular coagulation (DIC) secondary to pneumococcal infection, which presents with features of TMA but also bleeding and altered prothrombin time and activated partial thromboplastin time. Evolution of anuria/oliguria and thrombocytopenia is more prolonged than with shigatoxin-associated HUS (STEC-HUS) [61]. The direct Coombs test, unlike all other causes of TMA, is positive in 90% of cases and aids in the diagnosis of Sp-HUS [61].

Pathophysiology

Pneumococci release an enzyme called neuraminidase which removes neuraminic acid and exposes the Thomsen–Friedenreich antigen (T-antigen) present on red blood cells, platelets, and glomerular endothelial cells [64, 65]. The T-antigen is recognized by preformed IgM cold antibodies. More recently, decrease in sialylation of transferrin and IgA1 O-glycans was shown and may add to the pathophysiology of Sp-HUS, which is still not completely understood. In the early 2000s, studies showed that pneumococcal neuroaminidase leads to a disruption in Factor H, making it difficult to bind to C3 convertase effectively, culminating in the activation of the alternative complement pathway and cell injury [66].

Management

Treatment of Sp-HUS is based on supportive therapy with dialysis and initiation of antibiotic therapy. Early detection and rapid onset of antibiotics decrease mortality. In the most severe cases, use of the combination of vancomycin and cephalosporins to broaden the bacterial spectrum should be considered. Based on the cultures and sensitivity tests available, vancomycin should be discontinued, and therapy directed according to the results. The duration of treatment depends on the site of pneumococcal infection; if there is a need for platelet transfusions or red blood cell concentrate, they should be washed to remove IgM antibodies against Thomsen–Friedenreich antigen. The removal of neuroaminidase by plasma exchange is hypothetically indicated. However, plasma exchange is controversial since IgM antibody against the Thomsen–Friedenreich antigen may result in possible increased polyagglutination and aggravation of the microangiopathic phenomenon [62]. There are case reports of the use of complement inhibitor eculizumab in Sp-HUS with good response [67].

Immune-mediated (acquired) thrombotic thrombocytopenic purpura

Incidence and epidemiology

Immune-mediated thrombotic thrombocytopenic purpura (iTTP) is an immunological disorder caused by IgG autoantibodies that inhibit the function of ADAMTS13, which is to cleave the ultralarge multimers of von Willebrand Factor on endothelial cells. Although its incidence is higher in adults than in children, it is a severe condition that needs prompt recognition and treatment.

Clinical presentation and diagnosis

The classic clinical pentad of iTTP comprises hemolytic anemia, thrombocytopenia, neurological symptoms, jaundice and fever, although only a minority present with all symptoms [68]. In order to differentiate TTP from other causes of TMA in the intensive care unit, the Harvard TMA Research Collaborative developed a diagnostic score for TTP called Plasmic Score [69] which includes platelet count, hemolysis, no active cancer, no history of transplant, mean corpuscular volume of red cells <90 fL, normal coagulation, and creatinine <2.0 mg/dL (176 mmol/L). The score can be used on the bedside and has a high predictive value for severe ADAMTS13 deficiency.

Pathophysiology

The basic etiology was discovered in the early 1980s, when giant von Willebrand factor multimers (ULVWF, Ultra Large von Willebrand Factor) were found in the plasma of patients with TTP [70]. ULVWF maintain primary hemostasis, promote platelet adhesion, and aggregation through binding to platelet glycoproteins (GP) 1b–IX and GP IIb/IIIa [71]. In addition, ULVWF binds to free FVIII in the plasma, protecting it against proteolysis. ULVWF is stored mainly in the Weibel Palade bodies of endothelial cells and secreted into plasma in the form of large multimers. ADAMTS13 degrades these ULVWF high-density multimers, cleaving the A2 domain 842Tyr–843Met peptide bond, generating multimers ranging in size between 500 and 20,000 kDa [72]. The action of ADAMTS13 is increased in places or situations where there is greater stress on blood flow, that is, where the ratio between the pressure of blood flow and the diameter of the vessel is high [73]. Severe ADAMTS13 deficiency (defined as less than 10% of activity) results in unusual, giant FVW multimers at risk of platelet thrombus in small vessels causing red blood cell “shear” and occlusion of vessels with tissue ischemia. The HLA variant rs6903608 has been shown to be a risk factor for both immune TTP onset and relapse in an Italian cohort and may serve as a biomarker for worse outcome [74].

Management

Without treatment, iTTP has a 95% mortality rate with most deaths occurring within 24 h after initial presentation [75]. The recognition of iTTP in the first hours (“at bedside”) is crucial to institute treatment. If the collection of ADAMTS13 activity is feasible, it must be done (in a citrated tube) before beginning plasma exchange. It is worth mentioning that collecting or waiting for the result of the ADAMTS13 test *should not delay the start of treatment*, which is a priority. The use of plasma exchange leads to a response rate of 80% and survival greater than 90% when initiated in the first 4 to 8 h of suspected diagnosis and the American Society for Apheresis (ASFA) [76] has classified the procedure as indication level I for iTTP. When unavailable, plasma can be infused in a volume of 25 to 30 mL/kg of fresh plasma until plasmapheresis is available, although a randomized trial showed plasma infusions to be less effective in reducing mortality [77]. In addition, prednisone 1 mg/kg/day or pulse therapy with methylprednisolone should be combined. The current recommendation is daily plasma exchange until platelets are above $150,000/\text{mm}^3$, when plasma exchange can be stopped [78]. Solvent/detergent-treated (inactivates lipid-enveloped viruses) pooled plasma (OCTAPLAS) may be used in children instead of fresh frozen plasma if available [79]. Recently, a new medication for

the treatment of iTTP has been developed. Caplacizumab is a nanobody that binds to von Willebrand factor preventing platelet adhesion. In two prospective studies that enrolled 220 adults, patients who received caplacizumab together with plasma exchange and immunosuppression had lower rates of acute disease exacerbation [80, 81] and no mortality, which led to the drug being FDA and EU-approved. Caplacizumab use together with steroids and rituximab was retrospectively analyzed in 4 children (among 85 patients) in the UK—overall, the bleeding rate was 55%, thrombotic events occurred in 16%, and five patients died (four of whom with the drug starting 48 h after plasma exchange). Since thrombotic events are much more frequent than hemorrhage, platelet transfusions should be avoided except in cases of severe bleeding [76].

Hereditary thrombotic thrombocytopenic purpura

Incidence and epidemiology

Hereditary thrombotic thrombocytopenic purpura (hTTP), also known as Upshaw–Shulman Syndrome [82], is rare (10% of cases of TTP) and caused by homozygous mutations or by compound heterozygous mutations of the *ADAMTS13* gene; therefore, it is more prevalent in offspring of consanguineous parents. Patients with simple heterozygous mutations have no apparent abnormalities. There is a bimodal distribution according to age of diagnosis, with 50% presenting at age 2–5 years and 50% in young women usually triggered by pregnancy [78].

Clinical presentation and diagnosis

The clinical features of the hereditary form are recurrent episodes of microangiopathic hemolytic anemia and thrombocytopenia, usually with neurological manifestations or other signs of organ damage. Its diagnosis requires confirmation of severe ADAMTS13 deficiency (plasma activity < 10%) in the absence of inhibitory antibodies (non-inhibitory antibodies may be present) and confirmed by finding a pathogenic mutation in the *ADAMTS13* gene. There is often a delay in diagnosis of hTTP; the Hereditary TTP Registry [83] has enrolled more than 120 patients with mean age of presentation 4.5 years and mean age of diagnosis 16.7 years, showing the gap in knowledge regarding this disease.

Pathophysiology

Compared to iTTP, rather than decreased ADAMTS13 activity due to antibodies targeting ADAMTS13, in hTTP, ADAMTS13 (antigen) is deficient due to an

underlying genetic abnormality in the *ADAMTS13* gene. When *ADAMTS13* is lacking, ultra-large vWF multimers persist in the circulation resulting in aggregation of platelets, which are activated and form the vWF-rich microthrombi (platelet rich microthrombi are a hallmark of TTP). This process causes microthrombi and occlusion of microvessels leading to organ ischemia. Resultant laboratory features include thrombocytopenia, and evidence of mechanical destruction of red blood cells (schistocytes and hemolytic anemia). Levels of *ADAMTS13* are variable and dependent on the underlying genetic variant; this has an impact on severity and presentation [78].

Management

Current treatment is plasma infusion 10 to 15 mL/kg every 21–28 days.

TMA mediated by cobalamin defects

Incidence and epidemiology

Patients with an inherited vitamin B12 disorder are at risk for developing TMA. It has been mainly observed in neonates or in infants during the first year of life with methylmalonic aciduria and homocystinuria, caused by *cbl* deficiency type C (*cblC*; OMIM #277,400) [84]. *Cbl C* disease mostly manifests as failure to thrive, hypotonia, seizures, microcephaly, and delayed development. Three cases were diagnosed in Brazil, one of them was published in which the whole exome sequencing revealed two novel likely pathogenic heterozygous variants in *MTR* which encodes for methionine synthase, suggesting the diagnosis of methionine synthase deficiency (MSD) *cbl C* disease-mediated TMA [85]. The suspicion was made due to the lack of clinical and laboratory response to the use of eculizumab. This patient had mild neurological impairment that was characterized by developmental delay, mainly his speech. Few adult cases have previously been reported [86, 87].

Clinical presentation and diagnosis

Clinically, patients with a defect in cobalamin metabolism manifest with failure to thrive, hypotonia, seizures, microcephaly, and several degrees of delayed development. Neurological sequelae are common in affected children. Adult-onset disease is rare and may present with neurological manifestations such as cognitive impairment, ataxia, and psychosis. *Cbl* deficiency, or a defect in its metabolism, result in hyperhomocystinemia, decreased plasma methionine level, and methylmalonic aciduria. The reported adults had microangiopathic anemia, thrombocytopenia,

acute kidney injury, and hypertension. Chronic kidney disease with hypertension and proteinuria have been reported in up to 40% of patients.

Pathophysiology

Homozygous or heterozygous mutations in *MMACHC* (chromosome 1p34.1) cause deficiency of the *Cbl-C* enzyme. The *Cbl-C* enzyme is responsible for the metabolism of cobalamin into methylcobalamin and adenosylcobalamin. Furthermore, the lack of *Cbl-C* enzyme leads to the accumulation of homocysteine and methylmalonic acid in endothelial cells, which promote increased platelet aggregation, increased free radicals, increased expression of local procoagulant factors, and induce the binding of tissue plasminogen activator to the endothelium. All of these factors result in endothelial damage and trigger intravascular thrombosis with microangiopathic anemia, thrombocytopenia, and ischemia in organs. The resulting deficiency of methylcobalamin leads to hyperhomocysteinemia in the presence of normal blood levels of vitamin B12 [1].

Management

The main treatment for infants is parenteral hydroxycobalamin [86] (1 mg IM daily), which prevents progression to kidney failure and microangiopathic manifestations. Therefore, we suggest that homocysteine and vitamin B12 in the blood should be performed in all patients with TMA, especially if there is a neurological condition, persistently high levels of lactic dehydrogenase, average corpuscular volume > 100 microns³, or resistance to treatment with complement inhibitor.

TMA-mediated by diacylglycerol kinase epsilon mutation

Incidence and epidemiology

The Mendelian forms of HUS have been associated with complement genes, mostly transmitted as autosomal dominant variants (variable penetrance) and more rarely, autosomal recessive mutations (high penetrance). In 2013, recessive mutations in diacylglycerol kinase epsilon mutation (*DGKE*), a non-complement gene, were described in families with TMA through whole exome sequencing [88]. The incidence is still unknown, but it is an ultrarare cause of TMA.

Clinical presentation and diagnosis

Patients with recessive loss-of-function mutations in *DGKE* gene have phenotypes ranging from membranoproliferative glomerulonephritis, steroid-resistant-nephrotic syndrome [89] to HUS [90]. Usually, patients have hypertension and hematuria and the advance to CKD is slow and progressive. The search for a mutation in the *DGKE* gene is indicated in cases of TMA beginning in the first year of life—especially if there is significant proteinuria and the parents are consanguineous—and in cases where the response to terminal complement inhibition is unsatisfactory.

Pathophysiology

Diacylglycerol kinase (DGK) is a family of enzymes that catalyze the conversion of diacylglycerol (DAG) to phosphatidic acid (genecards.org)—both thought to function as bioactive lipid signaling molecules with important roles in the biosynthesis of complex lipids. Currently, nine members of the DGK family have been identified. DGKE has specificity for arachidonate-containing DAG. A type of TMA due to recessive mutations in the *DGKE* gene has been described [88]—loss of activity of the enzyme DAG, present in endothelial cells, platelets, and podocytes, induces apoptosis in endothelial cells and impairs angiogenic response, leading to a prothrombotic and inflammatory state with consequent TMA. Although the role of complement in kidney disease in patients with *DGKE* mutations was initially neglected, some patients with concomitant mutations in thrombomodulin and C3 have been identified and this possibly explains the spectrum of clinical manifestations in these patients, which normally occur as early as the first year of life. This type of isolated *DGKE* mutation is not associated with post-kidney transplant recurrence.

Management

The benefits of plasma exchange or plasma infusion and immunosuppression have been inconsistent. Recurrences occurred in patients who were receiving eculizumab [42] and recently a Brazilian case was published in which the detection of *DGKE* mutation was made in an adult due to non-response to eculizumab [7]. The treatment so far is restricted to nephroprotective measures (strict control of blood pressure, blockade of the renin–angiotensin–aldosterone system, metabolic control).

Transplant-associated TMA

Transplant-associated TMA (TA-TMA) results from endothelial damage occurring in recipients of solid organ or hematopoietic stem cell transplantation (HSCT). The

pathophysiology of this entity is complex and multifactorial. In solid organ transplant recipients, the main differential diagnoses are ischemia–reperfusion injury, calcineurin-inhibitor toxicity, antibody-mediated rejection, infections, recurrence of aHUS, and STEC-HUS. It is important to rule out these entities and consider complement inhibition early if aHUS recurrence is suspected [91]. Among patients with allogeneic HSCT, TMA may result from multiple mechanisms of endothelial injury that include infections, calcineurin inhibitors, graft versus host disease, and drugs. The cumulative incidence among allogeneic HSCT is 3% [92]. There may be underlying genetic variants in complement genes and the experience of a large cohort published by Jodele et al. [93] showed a significant improvement in 1-year post-HSCT survival from 16.7 to 66% with short course eculizumab use (median 11 infusions). Interestingly, TA-TMA usually presents with hemolysis and platelet consumption that is preceded by hypertension, with only mild and late kidney involvement.

Conclusion

Diagnosis and uncovering the etiology of TMA in children is of utmost importance in management of the disease entity. The causes of TMA in children less than 2 years old is variable and includes the more common entities of STEC-HUS and aHUS; the less common entities such as sp-HUS, iTTP, and hTTP; and rare entities including cblC-TMA and *DGKE*-TMA. TMA in the setting of solid organ or HSC transplantation has a multifactorial pathophysiology and children may have additional underlying complement defects. It is critical to recognize and correctly diagnose the specific TMA entities since the management, outcomes, and prognosis depend on early diagnosis and institution of appropriate treatment.

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Declarations

Conflict of interest LMPP is a speaker for Alexion Brazil, and scientific consultant for Apellis and Novartis.

Disclosure None.

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