The thrombotic microangiopathy (TMA) syndromes are extraordinarily diverse. They may be hereditary or acquired. They occur in children and adults. The onset can be sudden or gradual. Despite their diversity, TMA syndromes are united by common, defining clinical and pathological features. The clinical features include microangiopathic hemolytic anemia, thrombocytopenia, and organ injury. The pathological features are vascular damage that is manifested by arteriolar and capillary thrombosis with characteristic abnormalities in the endothelium and vessel wall. We focus on nine disorders that we describe as primary TMA syndromes, for which there is evidence supporting a defined abnormality as the probable cause (Table 1 and Fig. 1; and the interactive graphic, available with the full text of this article at NEJM.org). For clarity of this discussion, the names that have been chosen for these syndromes reflect their cause. However, we retain the common names of thrombotic thrombocytopenic purpura (TTP) for ADAMTS13 deficiency–mediated TMA and the hemolytic–uremic syndrome for Shiga toxin–mediated TMA (ST-HUS) because these names are familiar. We do not use the term “atypical HUS,” which was historically used to distinguish heterogeneous, uncharacterized syndromes from ST-HUS, since the term lacks both specificity and a suggestion of cause. We also do not use the term “idiopathic” with any of the primary TMA syndromes.

The presence of a causal abnormality, such as ADAMTS13 deficiency or a complement mutation, may not be clinically expressed until a condition, such as pregnancy, surgery, or an inflammatory disorder, precipitates an acute TMA episode. The treatment of such patients is focused on the cause of the primary TMA syndrome, not the precipitating condition. These patients are distinct from many other patients who have microangiopathic hemolytic anemia and thrombocytopenia that are manifestations of an underlying disorder (Table 2). The treatment of such patients is focused on the underlying disorder.

Knowledge of the pathogenesis, management, and outcomes of the primary TMA syndromes has accelerated in recent years (Fig. 2; and Table S1 in the Supplementary Appendix, available at NEJM.org). The objective of this review is to provide a unified perspective of these syndromes.

### TTP (ACQUIRED AND HEREDITARY)

**BACKGROUND**

In 1924, Moschcowitz described a 16-year-old girl with weakness, pallor, purpura, and hemiparesis who died after 14 days with cardiac failure. Autopsy revealed hyaline thrombi in terminal arterioles and capillaries throughout most organs, including the kidneys. This report was the first description of TMA, presumably TTP, also called ADAMTS13 deficiency–mediated TMA.

**CAUSE**

In 1982, unusually large multimers of von Willebrand factor were observed in patients with chronic, relapsing (hereditary) TTP. This finding led to the discovery of...
**Table 1. Primary Thrombotic Microangiopathy (TMA) Syndromes.**

<table>
<thead>
<tr>
<th>Name</th>
<th>Cause</th>
<th>Clinical Features</th>
<th>Initial Management</th>
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<tbody>
<tr>
<td><strong>Hereditary disorders</strong></td>
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<tr>
<td>ADAMTS13 deficiency–</td>
<td>Homozygous or compound heterozygous ADAMTS13 mutations</td>
<td>Initial presentation is typically in children but may also be in adults; possible evidence of ischemic organ injury; acute kidney injury is uncommon; patients with heterozygous mutations are asymptomatic.</td>
<td>Plasma infusion</td>
</tr>
<tr>
<td>mediated TMA (also called TTP)</td>
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<tr>
<td>Complement-mediated TMA</td>
<td>Mutations in CFH, CFI, CFB, C3, CD46, and other complement genes causing uncontrolled activation of the alternative pathway of complement</td>
<td>Initial presentation is often in children but may also be in adults; acute kidney injury is common; patients with heterozygous mutations may be asymptomatic.</td>
<td>Plasma infusion or exchange, anticomplement agent</td>
</tr>
<tr>
<td>Metabolism-mediated TMA</td>
<td>Homozygous mutations in MMACHC (encoding methylmalonic aciduria and homocystinuria type C protein)</td>
<td>Initial presentation is typically in children &lt;1 year of age; also reported in one young adult with hypertension and acute kidney injury.</td>
<td>Vitamin B12, betaine, folinic acid</td>
</tr>
<tr>
<td>Coagulation-mediated TMA</td>
<td>Homozygous mutations in DGKE; mutations in PLG and THBD also implicated</td>
<td>Initial presentation with acute kidney injury is typically in children &lt;1 year of age with DGKE mutations; clinical features of disorders associated with other mutations have not been described.</td>
<td>Plasma infusion</td>
</tr>
<tr>
<td><strong>Acquired disorders</strong></td>
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</tr>
<tr>
<td>ADAMTS13 deficiency–</td>
<td>Autoantibody inhibition of ADAMTS13 activity</td>
<td>Initial presentation is uncommon in children; often presents with evidence of ischemic organ injury; acute kidney injury is uncommon.</td>
<td>Plasma exchange, immunosuppression</td>
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<tr>
<td>mediated TMA (also called TTP)</td>
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</tr>
<tr>
<td>Shiga toxin–mediated TMA</td>
<td>Enteric infection with a Shiga toxin–secreting strain of <em>Escherichia coli</em> or <em>Shigella dysenteriae</em></td>
<td>Initial presentation is more common in young children, typically with acute kidney injury; most cases are sporadic; large outbreaks also occur.</td>
<td>Supportive care</td>
</tr>
<tr>
<td>(also called ST-HUS)</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Drug-mediated TMA (immune reaction)</td>
<td>Quinine and possibly other drugs, with multiple cells affected by drug-dependent antibodies</td>
<td>Initial presentation is a sudden onset of severe systemic symptoms with anuric acute kidney injury.</td>
<td>Removal of drug, supportive care</td>
</tr>
<tr>
<td>Drug-mediated TMA (toxic dose–related reaction)</td>
<td>Multiple potential mechanisms (e.g., VEGF inhibition)</td>
<td>Gradual onset of renal failure occurs over weeks or months.</td>
<td>Removal of drug, supportive care</td>
</tr>
<tr>
<td>Complement-mediated TMA</td>
<td>Antibody inhibition of complement factor H activity</td>
<td>Initial presentation is acute kidney injury in children or adults.</td>
<td>Plasma exchange, immunosuppression, anticomplement agent</td>
</tr>
</tbody>
</table>

* The primary TMA syndromes are described by evidence supporting a defined cause. Shiga toxin–mediated TMA (also called Shiga toxin–related hemolytic–uremic syndrome [ST-HUS]) occurs primarily in children and may be the most common of the nine primary TMA syndromes. Among adults, acquired thrombotic thrombocytopenic purpura (TTP) may be the most common primary TMA syndrome; acquired TTP is rare in children, in whom the incidence may be similar to that of hereditary TTP. The frequencies of TMAs that are mediated by complement, metabolism, coagulation, or drugs are unknown. The demonstration of antibodies that can neutralize the activity of complement factor H suggests that acquired TMA mediated by a deficiency in complement factor H may occur. DGKE denotes diacylglycerol kinase ε, PLG plasminogen, THBD thrombomodulin, and VEGF vascular endothelial growth factor.

A von Willebrand factor–cleaving protease that was subsequently characterized as ADAMTS13. ADAMTS13 cleaves von Willebrand factor multimers that are secreted from vascular endothelial cells. ADAMTS13 deficiency results in unusually large von Willebrand factor multimers and the risk of platelet thrombi in small vessels with high shear rates. Hereditary TTP (also called Upshaw–Schullman syndrome) is caused by homozygous or compound heterozygous ADAMTS13 mutations. Patients with heterozygous mutations have no apparent abnormalities. Acquired TTP is an autoimmune disorder caused by autoantibody inhibition of ADAMTS13 activity. The incidence of acquired TTP is much greater in adults (2.9 cases per 1 million per year) than in children (0.1 cases per 1 million per year). Factors that are associ-
ated with an increased frequency of this disorder include an age of 18 to 50 years, black race, and female sex.

**PRESENTATION AND DIAGNOSIS**

Among the primary TMA syndromes, TTP is unique for rarely causing severe acute kidney injury (Fig. 3). The clinical features of hereditary TTP are recurrent episodes of microangiopathic hemolytic anemia and thrombocytopenia, often with neurologic abnormalities or other signs of organ injury. Diagnosis of hereditary TTP requires documentation of ADAMTS13 deficiency and an absence of ADAMTS13 autoantibody inhibitor, and confirmation requires documentation of ADAMTS13 mutations. Hereditary TTP may be apparent at birth, with microangiopathic hemolytic anemia and thrombocytopenia, or not until adulthood, when it may be precipitated by a condition such as pregnancy. Although the severity of the condition may be related to ADAMTS13 mutations, observations of heterogeneity among siblings suggest that clinical manifestations require additional genetic or environmental factors, similar to observations in Adams13-deficient mice.

Presenting clinical features of acquired TTP are diverse; some patients have minimal abnormalities, whereas others are critically ill. Weakness, gastrointestinal symptoms, purpura, and transient focal neurologic abnormalities are common. However, one third of patients have no neurologic abnormalities. Most patients have normal or only transient, mildly elevated creatinine values. Diagnostic criteria are the presence of microangiopathic hemolytic anemia and
thrombocytopenia without another apparent cause. Thus, the exclusion of other primary TMA syndromes may not be possible. An ADAMTS13 level indicating less than 10% of normal activity supports the clinical diagnosis of acquired TTP. It identifies almost all patients at risk for relapse, but this level is neither sufficiently sensitive to identify all patients with TTP nor sufficiently specific to exclude patients with underlying disorders.

TREATMENT

The treatment for hereditary TTP is ADAMTS13 replacement by plasma infusion. Patients with severe plasma allergic reactions have been effectively treated with plasma-derived factor VIII concentrate that contains ADAMTS13. Although many patients require plasma only when thrombocytopenia or symptoms occur, others may require regular prophylactic plasma infusions.

Before the use of plasma exchange, survival from acquired TTP was 10%. In 1991, a randomized, controlled trial documented a survival rate of 78% with plasma exchange. The high mortality without treatment creates urgency to begin plasma exchange, which often results in treatment of patients who do not have TTP. Glucocorticoids are standard treatment; rituximab and other immunosuppressive agents are appropriate when the clinical course is complicated. Dialysis is rarely required.

LONG-TERM OUTCOMES

The long-term outcomes of patients with hereditary TTP are unknown. Experimental data suggest that ADAMTS13 provides protection against atherosclerosis, but it is unknown whether patients with hereditary TTP are at increased risk for cardiovascular disease. Long-term follow-up of patients with acquired TTP has revealed a risk of relapse and an increased prevalence of cognitive impairment, major depression, systemic lupus erythematosus, hypertension, and death.

FUTURE NEEDS

If long-term follow-up shows that hereditary TTP causes increased morbidities, prophylactic treatment will become more important. The development of recombinant ADAMTS13 would make prophylactic treatment simpler and safer. For the treatment of patients with acquired TTP, safer and more accessible alternatives to plasma exchange are needed.

<table>
<thead>
<tr>
<th>Table 2. Common Disorders Associated with Microangiopathic Hemolytic Anemia and Thrombocytopenia.</th>
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<tbody>
<tr>
<td><strong>Systemic infection</strong></td>
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<tr>
<td><strong>Systemic cancer</strong></td>
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<tr>
<td><strong>Severe preeclampsia, eclampsia, HELLP syndrome</strong></td>
</tr>
<tr>
<td><strong>Severe hypertension</strong></td>
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<tr>
<td><strong>Autoimmune disorders (e.g., systemic lupus erythematosus, systemic sclerosis, antiphospholipid syndrome)</strong></td>
</tr>
<tr>
<td><strong>Hematopoietic stem-cell or organ transplantation</strong></td>
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</table>

* Listed are disorders that may initially suggest the diagnosis of a primary TMA syndrome. Many different systemic infections (viral, such as human immunodeficiency virus and cytomegalovirus; fungal, such as aspergillus; and bacterial) and many different systemic cancers may be associated with microangiopathic hemolytic anemia and thrombocytopenia without overt disseminated intravascular coagulation (DIC). Many other disorders, such as any condition associated with DIC, may also present with microangiopathic hemolytic anemia and thrombocytopenia and must also be considered as alternative causes in the evaluation of patients for the possible diagnosis of a primary TMA syndrome. Some of these disorders (e.g., severe hypertension, systemic lupus erythematosus, and systemic sclerosis) may also be associated with the characteristic pathological features of TMA. These disorders may directly cause the clinical and pathological features of TMA, a hypothesis supported by resolution of these features with effective treatment of the disorder. HELLP denotes hemolysis, elevated liver-enzyme levels, and low platelets.

BACKGROUND

TMA that is characterized by predominant renal failure and described as HUS was recognized as a familial disorder in 1975. In 1981, two brothers with TMA were found to have a deficiency of complement factor H. The association between TMA and mutations in the gene encoding complement factor H (CFH) was established in 1998. Subsequently, mutations in multiple other factors facilitating increased complement activation by the alternative pathway have been identified in patients with TMA.

CAUSE

Complement-mediated TMA results from uncontrolled activation of the alternative pathway of complement. Unlike the other two pathways of complement activation, the alternative pathway is constitutively active as a result of spontaneous hydrolysis of C3 to C3b. In the absence of normal regulation, C3b deposition on tissues may increase markedly, resulting in increased formation of the C5b-9 terminal complement complex (also called the membrane-attack complex) and injury of normal cells. The precise role of complement dysregulation in TMA has not been fully...
defined. Endothelial injury as well as complement dysregulation on the platelet surface causing activation may be involved.\(^\text{30}\)

Hereditary complement-mediated TMA may result from either a loss-of-function mutation in a regulatory gene (CFH, CFI, or CD46) or a gain-of-function mutation in an effector gene (CFB or C3).\(^\text{31,32}\) Most complement mutations that are
associated with TMA are heterozygous, even though many family members with heterozygous mutations are asymptomatic. A difference between probands and family members in the presence of additional modifying genes may explain this discrepancy. Other genetic abnormalities have been identified in patients with complement-mediated TMA, including single-nucleotide polymorphisms in \( CFH \) and \( CD46 \), copy-number variations in the \( CFH \)-related 1 and 3 genes (\( CFHR1 \) and \( CFHR3 \)), and fusion genes of the \( CFHR \) region with \( CFH \) caused by nonallelic homologous recombination. These additional genetic abnormalities may contribute to the loss of alternative pathway regulation and increased risk of TMA. In addition to genetic abnormalities, a functional deficiency in complement factor H may result from antibodies to the comple-
ment, resulting in acquired TMA. CFH antibodies account for about 10% of complement-mediated TMA. These antibodies are responsible for defective CFH-dependent cell protection.

**PRESENTATION AND DIAGNOSIS**

Acute kidney injury and hypertension are prominent abnormalities in complement-mediated TMA. Current diagnostic criteria are those that were used in clinical trials involving a total of 37 patients, which supported the approval of eculizumab (a humanized monoclonal antibody that blocks the generation of C5a and C5b) for the treatment of “atypical HUS” in 2011. These criteria include all of the following: a serum creatinine level at or above the upper limit of the normal range, microangiopathic hemolytic anemia, thrombocytopenia, ADAMTS13 activity of 5% or more, and negative stool tests for Shiga toxin–producing infection.33 These criteria are not specific; they may also occur in all other primary TMA syndromes as well as in other patients with microangiopathic hemolytic anemia and thrombocytopenia. Complement
genetic studies, now commercially available with a rapid return of results, may provide a more specific diagnosis. Normal plasma levels of C3, C4, and complement factors H, B, and I do not exclude the diagnosis of complement-mediated TMA.

TREATMENT
Anticomplement therapy can be used to supplement plasma therapy and potentially preempt liver transplantation. Eculizumab is currently the only available anticomplement agent. Its effect may be limited among patients who have C5 mutations. The nonspecific diagnostic criteria and the fact that patients with no identified complement mutation may have a response to anticomplement therapy makes the decision to use an anticomplement agent as initial therapy difficult. Anticomplement therapy is a reasonable initial treatment for patients with antibodies against complement factor H. However, the use of immunosuppression to reduce the antibody titer should be considered. The high cost of eculizumab (wholesale acquisition cost for 1 year of treatment for an adult, $614,736; University of Oklahoma Medical Center Pharmacy, March 5, 2014) and the implication that it should be continued indefinitely are critical issues. The risk of meningococcal infection in connection with eculizumab therapy must be considered.

LONG-TERM OUTCOMES
Before the use of eculizumab, the risks of end-stage renal disease or death and of recurrence after kidney transplantation were primarily dependent on mutational analyses, with CFH mutations causing the greatest risk. It is assumed that anticomplement treatment will improve outcomes.

FUTURE NEEDS
Clinical trials to determine the most appropriate length of anticomplement therapy and to develop surveillance markers to confirm renal remission are critical. Further genetic investigation is required to explain the heterozygosity and low penetrance of currently identified complement mutations and also to determine the cause of disease in patients who may have complement-mediated TMA but in whom no mutation has been identified.

<table>
<thead>
<tr>
<th>SHIGA TOXIN–MEDIATED HEMOLYTIC–UREMIC SYNDROME</th>
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<tr>
<td><strong>BACKGROUND</strong></td>
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| The name “hemolytic–uremic syndrome” was first proposed in 1955. The typical clinical features of ST-HUS were first described in a 1962 report involving five children between the ages of 6 and 10 months who had diarrhea preceding renal failure. An infectious cause was suspected, but an association between ST-HUS and an enteric infection, 
| Shigella dysenteriae type 1, was not recognized until 1975. It was not until 1983, during an investigation of outbreaks of hemorrhagic colitis, that Escherichia coli O157:H7 was identified as a pathogen. The same year, the association between ST-HUS and Shiga toxin–producing E. coli was described. Multiple E. coli strains produce Shiga toxin; E. coli O157:H7 is the most common pathogen associated with ST-HUS in Europe and the Americas. S. dysenteriae type 1 remains an endemic cause of ST-HUS in other countries. Although ST-HUS is popularized by large outbreaks, most occurrences are sporadic. ST-HUS is much more common among children (median age, 2 years), in whom mortality is 3%. ST-HUS in adults is more severe, with higher mortality. |
| **CAUSE** |
| Shiga toxin–producing E. coli are common intestinal bacteria in cattle, consistent with the rural predominance of endemic ST-HUS. Outbreaks result from contaminated water, beef products, vegetables, and other foods. Cell damage results when Shiga toxin binds to globotriaosylceramide (Gb3, also known as CD77 or ceramide trihexoside) on endothelial cells, as well as to renal mesangial cells and epithelial cells (podocytes and tubular cells). Cell apoptosis results from Gb3 binding, endocytosis, retrograde transport, cytosolic translocation of Shiga toxin, and subsequent ribosomal inactivation. Shiga toxin is also cell-activating, proinflammatory, and prothrombotic and facilitates thrombosis by inducing endothelial secretion of von Willebrand factor. |
| **PRESENTATION AND DIAGNOSIS** |
| Severe abdominal pain and diarrhea, often bloody, begin several days after contaminated food is consumed. Thrombocytopenia and renal...
failure begin as gastrointestinal symptoms resolve.\textsuperscript{41} Shiga toxin is identified by means of stool analyses during the acute colitis phase but may not be identifiable when ST-HUS begins.\textsuperscript{54}

**TREATMENT**

Treatment remains supportive. Early, aggressive hydration has a renal protective role.\textsuperscript{41} Patients commonly require dialysis.\textsuperscript{47} The benefits of plasma exchange and anticomplement treatment are uncertain.

**LONG-TERM OUTCOMES**

Hypertension and neurologic abnormalities may persist after the acute phase has resolved. End-stage renal disease rarely occurs.\textsuperscript{55}

**FUTURE NEEDS**

The prevention of enterohemorrhagic infections and ST-HUS by public health measures (e.g., food safety and hygiene education) is the greatest need.\textsuperscript{46} New agents that neutralize Shiga toxin may provide effective treatment for ST-HUS.

**PRESENTATION AND DIAGNOSIS**

Drug-mediated TMA may be suspected by the sudden onset of severe systemic symptoms, often with anuric acute kidney injury, within hours after drug exposure. There may be a history of illness after previous exposures to the suspected drug. The association with quinine may be overlooked because exposure may occur over many years and may not be reported by the patient without explicit questions. Exposure may include tablets or quinine-containing beverages. Documentation of drug-dependent antibodies supports the clinical diagnosis; however, a negative test does not exclude a drug association.

**TREATMENT**

Supportive care and drug avoidance may be the only beneficial management. Plasma exchange is often begun because TTP is suspected and a drug-mediated cause is uncertain.

**LONG-TERM OUTCOMES**

Chronic kidney disease with hypertension is common.\textsuperscript{59} End-stage renal disease may occur.

**FUTURE NEEDS**

Rapidly available testing for drug-dependent antibodies is needed to assist in the clinical diagnosis. Understanding the mechanism of acute kidney injury may provide insight for targeted treatment.

**Drug-Mediated TMA (Toxic Dose-Related Reaction)**

**BACKGROUND**

Many drugs, including immunosuppressive and chemotherapeutic agents and vascular endothelial growth factor (VEGF) inhibitors, have been reported to cause TMA through dose- and time-dependent toxicity. Evidence supporting a causal role is limited.

**CAUSE**

There may be multiple mechanisms for toxic drug-mediated kidney injury. Among the likely roles of calcineurin inhibitors (such as cyclo-
sporine and tacrolimus) is their ability to cause endothelial dysfunction and increased platelet aggregation, possibly through the inhibition of prostacyclin. The inhibition of VEGF function in renal endothelial cells and podocytes causes gradual development of glomerular TMA.\textsuperscript{64,65}

**PRESENTATION AND DIAGNOSIS**

The typical presentation is gradual loss of kidney function with hypertension.\textsuperscript{65} Abrupt, severe TMA may occur, as with intravenous abuse of the opiate oxymorphone.\textsuperscript{66}

**TREATMENT**

Supportive care and drug avoidance may be the only beneficial management. For some drugs, such as calcineurin inhibitors, dose reduction, rather than drug avoidance, may be sufficient.

**LONG-TERM OUTCOMES**

Microangiopathic hemolytic anemia and thrombocytopenia often resolve. Renal failure may persist.

**FUTURE NEEDS**

Determining how drug toxicity causes TMA, or identifying alternative explanations for TMA in these patients, is essential.

### Metabolism-Mediated TMA

**BACKGROUND**

Cobalamin C disease is a hereditary disorder of cobalamin (vitamin B\textsubscript{12}) metabolism that may cause TMA and multiple organ dysfunction in infants.\textsuperscript{67,68} In addition, TMA has been reported in one adult.\textsuperscript{69}

**CAUSE**

Disorders of cobalamin metabolism result from homozygous or compound heterozygous mutations in a gene encoding the methylmalonic aciduria and homocystinuria type C protein (\textit{MMACHC}). The resulting deficiency in methylcobalamin causes hyperhomocysteinemia, decreased plasma methionine levels, and methylmalonic aciduria. Abnormal cobalamin C metabolism is associated with platelet activation, generation of reactive oxygen species, endothelial dysfunction, increased tissue factor expression, and coagulation activation.\textsuperscript{70}

**PRESENTATION AND DIAGNOSIS**

Infants with cobalamin C disease have diverse developmental abnormalities. The one reported adult presented with microangiopathic hemolytic anemia, thrombocytopenia, acute kidney injury, and hypertension; he had no relevant personal or family history.

**TREATMENT**

Parenteral hydroxycobalamin is the principal treatment for infants. When the above-mentioned adult patient did not have a response to anticomplement treatment, an intracellular defect of vitamin B\textsubscript{12} metabolism was suspected and confirmed by routinely available tests, which showed hyperhomocysteinemia, decreased plasma methionine, methylmalonic aciduria, and normal plasma vitamin B\textsubscript{12} levels. He had a response to treatment with hydroxycobalamin, betaine, and folinic acid.\textsuperscript{69}

**LONG-TERM OUTCOMES**

Neurologic sequelae are common in affected infants. Chronic kidney disease with hypertension and proteinuria has been reported in up to 40% of patients. End-stage renal disease and the recurrence of TMA appear to be prevented by replacement treatment.

**FUTURE NEEDS**

Greater awareness of cobalamin C–induced TMA and greater use of routine screening tests for cobalamin deficiency are required.

### Coagulation-Mediated TMA

**BACKGROUND**

Genetic abnormalities of thrombomodulin,\textsuperscript{71} plasminogen,\textsuperscript{32} and a protein kinase C–associated protein, diacylglycerol kinase \(\varepsilon\) (DGKE),\textsuperscript{72,73} have been identified in patients with TMA. These findings suggest that there may be a primary role for coagulation proteins in the pathogenesis of TMA syndromes.

**CAUSE**

The question of whether the role of thrombomodulin in TMA is primarily related to coagulation or is mediated solely by complement requires further study. The role of DGKE has been documented in two reports describing 22 patients in 12 families.\textsuperscript{72,73} All the patients had homozygous
or compound heterozygous DGKE mutations. Patients with heterozygous mutations had no clinical abnormalities. In one report, no mutations of complement regulatory genes were identified.\textsuperscript{73} It has been proposed that a homozygous mutation in the gene encoding DGKE leads to a loss of function, resulting in protein kinase C (PKC) activation. PKC activation facilitates the up-regulation of prothrombotic factors (von Willebrand factor, tissue factor, and plasminogen activator inhibitor 1) and the down-regulation of the VEGF receptor. Decreased VEGF function with resulting renal podocyte injury may also be a consequence of DGKE mutations. The sum of these events is a shift to a prothrombotic state.\textsuperscript{74} Communication between the coagulation system and the complement system may have a role in the pathogenesis of TMA.

**PRESENTATION AND DIAGNOSIS**

Patients with DGKE mutations present with acute kidney injury. Most of these patients who have been studied were under the age of 1 year.\textsuperscript{72,73}

**TREATMENT**

The benefits of plasma infusion or exchange and immunosuppression have been inconsistent. Relapses have occurred while patients were receiving anticomplement therapy.

**LONG-TERM OUTCOMES**

End-stage renal disease is common. Four children with this disorder have undergone kidney transplantation; one has had recurrent kidney injury.\textsuperscript{72}

**FUTURE NEEDS**

Determining the prevalence of these syndromes and understanding the mechanisms leading to TMA are essential.

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**MECHANISMS OF ORGAN INJURY**

The above-mentioned causes of the primary TMA syndromes do not explain why kidney injury is predominant in all except for TTP. TTP causes extensive kidney microvascular thrombi,\textsuperscript{75} yet severe acute kidney injury and chronic kidney disease are rare.\textsuperscript{16,25} The difference may be that the pathogenesis of TTP is primarily related to vascular thrombosis, whereas the pathogenesis of the other primary TMA syndromes also includes injury to resident renal cells and endothelial cells. For example, in patients with ST-HUS, Shiga toxin causes injury to epithelial podocytes,\textsuperscript{50} mesangial cells,\textsuperscript{49} and renal tubular cells.\textsuperscript{51} VEGF inhibitors affect glomerular epithelial podocytes,\textsuperscript{64,65} and enhanced protein kinase C activation in patients with DGKE mutations may cause podocyte injury.\textsuperscript{74} Understanding the mechanism of injury to resident renal cells in complement-mediated TMA, by quinine-dependent antibodies, or in cobalamin C disease requires further study.

There are also distinct differences with respect to the risk of chronic kidney disease among the seven primary TMA syndromes in which acute kidney injury is typically present. Among the acquired syndromes, chronic kidney injury rarely develops in patients with ST-HUS,\textsuperscript{59} perhaps because ST-HUS does not recur. However, patients with only a single recognized episode of quinine-induced TMA commonly have chronic kidney injury.\textsuperscript{59} Among the hereditary syndromes, there is also substantial variety in the severity of kidney injury. Patients with complement mutations may present at any age, whereas those with DGKE mutations typically present with acute kidney injury in infancy.\textsuperscript{73} These variations in the clinical manifestations of TMA syndromes emphasize the need for greater understanding of the mechanisms of disease.

**CONCLUSIONS**

In recent years, there has been a dramatic acceleration in our understanding of the primary TMA syndromes. New discoveries with respect to causal mechanisms have created opportunities for specific treatments. The availability of specific treatments has created the need for rapid, specific diagnoses. The use of effective treatments has decreased mortality but has also revealed previously unrecognized long-term morbidities. These advances predict continuing acceleration of our understanding of the primary TMA syndromes.

Dr. George reports receiving fees for serving on advisory boards from Alexion; and Dr. Nester receiving grant support from Celldex and fees for serving on advisory boards from Alexion. No other potential conflict of interest relevant to this article was reported.

Disclosure forms provided by the authors are available with the full text of this article at NEJM.org.
SYNDROMES OF THROMBOTIC MICROANGIOPATHY

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