Thrombotic Microangiopathy With Granulomatosis Interstitial Nephritis in an Allogenic Bone Marrow Transplant Patient: A Case Report and Review of the Literature

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Abstract

Transplant-associated thrombotic microangiopathy (TA-TMA) is a rare complication of hematopoietic stem cell transplantation (HSCT) with variable presentations. TA-TMA has often been described as a diagnosis of exclusion but a renal biopsy is rarely pursued to confirm the diagnosis, an essential step for our patient with renally limited TMA. We report a case report from the onconephrology clinic and review the literature associated with TA-TMA as it relates to diagnosis and treatment. A 45-year-old woman with acute myeloid leukemia and stage 3 chronic kidney disease underwent a matched unrelated donor allogenic HSCT. Postoperatively, she developed gastrointestinal graft versus host disease (GVHD) and was treated with tacrolimus, sirolimus, budesonide, and beclomethasone. Following discharge, she developed uncontrolled hypertension and required losartan, amlopidine, carvedilol, clonidine patch, and hydralazine as needed. On day 180 post-transplant, she developed lower extremity edema and acute kidney injury (AKI) with creatinine increasing to 2 mg/dL. On day 480 post-transplant, she developed worsening thrombocytopenia, anemia, new hematuria, left flank pain, and worsening renal function with creatinine peaking to 6 mg/dL. Peripheral smear revealed no schistocytes, lactate dehydrogenase of 265 mg/dL, and urinalysis with 100 mg/dL protein. ADAMTS 13 activity was normal (92%) and no inhibitor was detected. She became anuric and was started on hemodialysis. Renal biopsy revealed glomerular changes consistent with TA-TMA. During HSCT, systemic vascular endothelial injury triggers microangiopathic hemolytic anemia, platelet consumption, injury of glomerular endothelial cells and fibrin occluded renal capillaries. Thus, TA-TMA should be considered in HSCT patients with elevated LDH, proteinuria, hypertension, and AKI. However, a diagnosis is difficult to confirm without a renal biopsy. Treatment involves discontinuing potentially toxic agents such as calcineurin inhibitors and sirolimus, prescribing adequate antimicrobial treatment, and using renal replacement therapy if needed. A renal biopsy early in the course of disease not only confirms the diagnosis, but may limit the extent of disease.

Keywords: Thrombotic microangiopathy; Bone marrow transplant; Interstitial nephritis; Renal biopsy

Introduction

Transplant-associated thrombotic microangiopathy (TA-TMA) is a known, but rare complication of hematopoietic stem cell transplantation (HSCT), occurring in about 10-30% of HSCT patients [1-4]. In a bone marrow transplant (BMT) patient, the cause of TMA can be difficult to discern due to coexisting graft versus host disease (GvHD), transplant complications such as infection, disseminated intravascular coagulation, and side effects from immunosuppression [2]. In addition, TMA can present variably, sometimes without findings of hemolytic anemia or acute kidney injury (AKI) [5]. Few published studies document renally limited TA-TMA. We present a novel case in which progressive renal disease was eventually explained by a renal biopsy which revealed TA-TMA with granulomatous interstitial nephritis more than 1 year after HSCT.

Case Report

A 45-year-old woman with an allogeneic stem cell transplant from a matched unrelated donor for chemotherapy-induced high grade myelodysplastic syndrome (MDS) with 5q deletion which subsequently transformed into acute myeloid leukemia (AML) presented to the emergency room with nausea and vomiting. Her dehydration and emesis was treated with intravenous fluids and antiemetics. On admission, her creatinine was 2 mg/dL and her renal function continued to worsen over subsequent days (Fig. 1). About 5 years ago, she was treated for endometrial adenocarcinoma with carboplatin and paclitaxel which was complicated by the development of peripheral neuropathy, and MDS which transformed into AML 3 years later. She underwent induction chemotherapy with cytarabine and high dose daunorubicin (7+3). Due to residual disease on bone marrow biopsy, she was re-induced with salvage chemotherapy...
consisting of cladribine, cytarabine, and filgrastim (CLAG) with subsequent bone marrow biopsy showing hypocellularity with no blasts. She underwent consolidation chemotherapy with CLAG again and underwent allogenic HSCT. She was given a 5-day course of fludarabine and melphalan for transplant conditioning and tacrolimus with sirolimus for GvHD prophylaxis. She tolerated her HSCT without any side effects. Her hospital course was complicated by presumed acute gastrointestinal GvHD, diagnosed clinically due to worsening nausea, vomiting, and diarrhea and was treated with beclomethasone, budesonide, tacrolimus and sirolimus. She later began having neutropenic fevers, bacteremia treated with vancomycin and also started on amphotericin due to concern for a disseminated fungal infection. During this time, her renal function declined with suspicion regarding vancomycin and amphotericin. At the time of discharge, her creatinine was 1.2 mg/dL.

Approximately 180 days after transplant, her gastrointestinal symptoms resolved and her follow-up visits became inconsistent; however, she continued her GvHD prophylaxis regimen. She presented intermittently with hypertension and peripheral edema which was treated with losartan 100 mg
Due to her non-compliance with appointments, she presented to clinic with uncontrolled hypertension and occasional nausea only when she ran out of her immunosuppressants. On subsequent follow-up appointments, her renal function continued to decline and her resistant hypertension required the addition of amlodipine, carvedilol, clonidine patch, and hydralazine.

Approximately 360 days post-BMT, she continued to have uncontrolled hypertension, nausea, vomiting, and weight loss, and developed palpable purpura of her lower extremities. Skin biopsy revealed extravasated erythrocytes consistent with purpura without presence of vasculitis. Tacrolimus was discontinued and she continued on sirolimus and mycophenolate mofetil for immunosuppression. Around 400 days after BMT, she began to experience hypertensive crisis with blood pressure of 198/136 mm Hg and was hospitalized twice due to difficulty tolerating her oral antihypertensives secondary to nausea and emesis. On each admission, she improved with intravenous fluids, antiemetics, and resumed her oral antihypertensives. Losartan was held, but her AKI continued to worsen (Fig. 1).

Around 480 days post-BMT, she developed 2 weeks of hematuria, intermittent lower back pain, and left flank pain. She was seen in nephrology clinic for her chronic kidney disease (CKD). Given her new constellation of anemia, hematuria, flank pain, and persistent proteinuria on urinalysis, there was concern for possible TA-TMA (Fig. 2). An ADAMTS13 revealed activity was normal (92%). Peripheral blood smear was unremarkable without any schistocytes. Her renal function panel showed rapidly progressing renal failure. She became anuric and was started on hemodialysis. Renal biopsy revealed histomorphological features consistent with TMA with granulomatous tubulointerstitial nephritis (Fig. 3). She currently continues to be hemodialysis dependent and follows regularly with nephrology and the BMT clinic.

**Discussion**

**Overview of TA-TMA**

TA-TMA can be caused by a number of factors, making it a difficult etiology to diagnose and treat. It is characterized by systemic vascular endothelial injury that can be triggered by several different elements of the bone marrow transplant process. This endothelial injury causes microangiopathic hemolytic anemia and platelet consumption, which results in thrombosis and fibrin deposition in the microcirculation [6]. TA-TMA is associated with complement system activation; in general, complement dysregulation is associated with poor prognosis. Though common manifestations include hemolytic anemia and thrombocytopenia, organ damage causes the most morbidity and mortality, namely acute renal failure [2]. TA-TMA, in particular, involves injury of glomerular endothelial
cells in which fibrin occludes the renal capillaries [4, 7] (Fig. 3).

This endothelial damage in TA-TMA can be caused by a number of factors. It was once assumed that the myeloablative and intense conditioning regimens prescribed prior to HSCT would increase risk of TMA, but this has not been statistically and consistently proven. Infections from aspergillus, cytomegalovirus, parvovirus B19, human herpes virus-6, and even adenovirus, which bind to vascular endothelial growth factor (VEGF) and BK virus, which causes glomerular and renal tubular cell damage, have all been associated with increased incidence of TMA. In addition, coagulation cascade markers and complement activation mediated by the humoral immune response play integral roles in the development of TMA [6].

TA-TMA is difficult to diagnose due to its variation of criteria (the Bone Marrow Transplant Clinical Trials Network versus the International Working Group criteria) [8]. However, it is understood that elevated lactate dehydrogenase, proteinuria on routine urinalysis, and hypertension are the earliest markers of TMA. Proteinuria and evidence of terminal complement activation (elevated sC5b-9) in the blood at the time of TMA diagnosis were associated with very poor survival (20% at 1 year) [5]. AKI usually occurs 1 month after TMA, whereas hypertension and proteinuria typically occurs 10 - 14 days after TMA [5]. Different criteria signify haptoglobin as normal or decreased in TMA, due to the role of haptoglobin in hemolytic anemia as well as its role as an acute phase reactant [2]. Therefore, one should consider TMA in HSCT recipients if they present with an acute elevation in LDH, proteinuria, and hypertension out of proportion to what would be expected from calcineurin inhibitor and steroid therapy [5].

**GvHD and TA-TMA**

The relationship between GvHD and TA-TMA is known, but not fully understood due to confounding variables. It is thought that during engraftment, donor T lymphocytes cause direct damage to host endothelial cells. Other theories of endothelial injury from transplant include circulating cytokines, low vascular endothelial growth factor (VEGF), and coagulation pathway activation. While the exact pathophysiology of chronic GvHD is unknown, it is often associated with thrombocytopenia [6, 9], which is consistent with our patient (Fig. 4).

Differentiating between intestinal TMA from acute GvHD can be difficult in patients suffering from severe and refractory diarrhea after BMT due to the similar symptomology. When a patient develops diarrhea after transplantation, it is usually a diagnostic component of GvHD. On the contrary, solitary gut TMA should be in the differential because the treatment differs between TMA and GvHD. GvHD is typically treated with increasing immunosuppression and TMA is treated by decreasing immunosuppression [10]. Although it is understood that the first-line therapy for chronic GvHD is steroids, refractory cases have no treatment guidelines resulting in varying combinations of immunosuppressants often based on expert opinion.

**Immunosuppressants: toxicity vs. treatment?**

Many drugs are known to cause nephrotoxicity [11]. While renal failure can be due to chemotherapy, radiation, fluid loss
secondary to diarrhea or vomiting, sepsis, or nephrotoxicity secondary to calcineurin inhibitors and antimicrobials, there are BMT-specific reasons such as marrow infusion toxicity, hepatic veno-occlusive disease, TMA, and GvHD [12, 13]. Research surrounding drug-induced TMA is still growing as the list of insulting agents vary widely. More research shows that calcineurin inhibitors (cyclosporine and tacrolimus) cause renal toxicity. However, antiangiogenic therapy such as sirolimus targeting VEGF or its receptor has been increasingly associated with TMA [14, 15]. This may be due to the fact that sirolimus is a mTOR inhibitor which is a required component for VEGF production and signaling [16]. Similarly, bevacizumab, a more well-known antiangiogenic agent, causes TMA by binding VEGF [17]. Combining both calcineurin inhibitors and sirolimus results in higher rates of TMA [18].

Though some studies show that using both tacrolimus and sirolimus for GvHD prophylaxis increases risk of TA-TMA, other studies are inconclusive [1]. One autopsy study showed that the association between GvHD and renal TMA was independent of the use of calcineurin inhibitors and conditioning with total-body irradiation [19]. Nevertheless, TMA associated with sirolimus had a more favorable prognosis compared to TMA associated with calcineurin inhibitors alone [18]. Thus, sirolimus has remained the first choice for GvHD control for more than a decade given its superiority in renal outcomes over calcineurin inhibitors and other immunosuppressives such as methotrexate despite its increasing association with TA-TMA [20]. Sirolimus has been associated with greater protection against vasculopathy but at the same time induces proteinuria [16]. This downregulation of VEGF by sirolimus prevents repair of injured renal endothelium and is thought to be a potential predisposing factor for TMA, preferentially affecting the kidneys [21]. The balance between risk and benefit is precarious as providers must decide between using sirolimus for GvHD prophylaxis despite the known and increased risk of TMA.

Renal pathology in GvHD and TA-TMA

Patients who receive HSCT experience extensive damage to their kidneys. Chronic kidney disease is estimated to impact 15-20% HSCT recipients [22]. Acute injury is considered to occur when serum creatinine levels are elevated up to 100 days after transplantation, whereas chronic injury represents injury at or after that [13]. This was the case for our patient who presented with a decline in renal function, more than a year after her transplant. Several theories exist as to why TA-TMA tends to preferentially affect the kidneys. One reason may be that fenestrated endothelium of the glomeruli is vulnerable to damage [23]. Yet an autoimmune process may exist given the fact that inflammatory cells such as CD3+ and CD8+ T cells and cytotoxic T cells with natural killer cells were detected in glomeruli, tubules, and the interstitium of patients with TMA. Endothelial damage in TA-TMA has rarely been reported outside the kidney. Another theory states there is more turbulent blood flow through the renal microvasculature than anywhere else [24].

HSCT recipients often experience a number of concurrent renal abnormalities. According to a recent report, kidney-biopsy specimens obtained from patients with TMA showed mesangiolysis and loss of endothelial cells with occlusion of capillary lumens with fibrin, as was the case for our patient [13]. According to one institution’s study of 67 patients, kidney specimens often demonstrated acute tubular injury, interstitial fibrosis, arteriolar hyaline, arteriosclerosis, and rarely membranous nephropathy [25]. Another study showed that the most commonly reported pathology was membranous nephropathy, followed by minimal change disease [22]. However, our patient presented with a rare entity on renal biopsy: the combination of both TMA and interstitial (granulomatous) nephritis (Fig. 3). Granulomatous interstitial nephritis is detected in less than 1% of all renal biopsies and usually associated with analgesics, antibiotics, and granulomatous disorders [26].

Treatment strategies

Autopsy and biopsy studies show that any clinical criteria for TMA lack sensitivity and specificity for renally limited TMA [22]. The question then becomes, should every BMT patient be getting a renal biopsy? Though it presents similarly to other thrombotic diseases, TA-TMA is treated differently. TA-TMA is different than TTP in that these patients often lack suppression of ADAMTS13 (a disintegrin and metalloproteinase with thrombospondin type 1 motif, member 13) and do not respond to plasma exchange [2]. When TA-TMA is suspected, conservative supportive care should be initiated. This includes discontinuing potentially toxic agents such as calcineurin inhibitors and sirolimus, prescribing adequate antimicrobial treatment, and maintaining adequate renal function using renal replacement therapy [4]. Whereas some clinicians discontinue calcineurin inhibitors and opt for switching to mycophenolate mofetil, sirolimus or both, some patients have experienced resolution of their TMA when given increasing doses of calcineurin inhibitors [13]. Furthermore, aggressively managing hypertension, another risk factor for CKD and TA-TMA, is imperative as 70% of children and adults develop hypertension within the first 2 years of their HSCT [13]. Though some studies show blocking the complement system with eculizumab is a possible treatment for TA-TMA, the evidence is still limited as to whether it improves outcomes in patients [27].

Conclusions

This case study emphasizes the importance of TA-TMA as a consideration in the differential diagnosis in patients with HSCT who develop hematuria, proteinuria, hypertension (especially hypertension that is resistant), AKI, and elevated LDH, or a combination of these signs. This case also underscores the importance of renal biopsy when diagnosing the etiology of declining renal function status post HSCT. There is a definite need for biopsy in BMT patients with renal dysfunction. Though biopsy results can reveal type, acuity, and chronicity of the renal disease which facilitate treatment decisions, cancer patients are not often biopsied. TA-TMA is a complex and difficult diagnosis that requires a high degree of suspicion.
from both the hematologist and nephrologist when managing the post-HSCT patient.

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