Intracerebral hemorrhage after IV tPA for stroke as early symptom of ANCA-associated vasculitis

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Introduction

Anti-neutrophil cytoplasmic antibody (ANCA)-associated vasculitides (AAV) are rare diseases characterized by a necrotizing small-vessel vasculitis and circulating ANCA that comprise granulomatosis with polyangiitis, microscopic polyangiitis, and eosinophilic granulomatosis with polyangiitis (EGPA). Acute ischemic stroke (AIS) can be a manifestation of central nervous system (CNS) involvement [1]. In addition, intracerebral hemorrhage (ICH) is a potential complication of these necrotizing vasculitides [2]. Intravenous tissue plasminogen activator (IV tPA) is the standard treatment for AIS. Vasculitides are not listed as contraindications for its administration [3]. However, CNS vasculitis could be a contraindication to IV tPA because the risk of ICH. We describe a patient who developed multiple ICHs following IV tPA administration for AIS. The stroke was the initial presentation of AAV. We raise awareness to the dangers of IV tPA use in patients with AAV.

Case report

A 58-year-old man was admitted for myalgia, night sweats, low grade fevers, dark urine and urticarial rash over his trunk and extremities that progressed over two weeks. Twelve hours later, he developed acute right hemiparesis, dysarthria and right facial droop; his National Institutes of Health Stroke Scale (NIHSS) score was 4. Initial evaluation, including computed tomography (CT) scan, did not identify any contraindication and IV tPA was administered at 71 min after symptom onset (Fig. 1A). His symptoms initially resolved with IV tPA but shortly afterward he deteriorated with decreased level of arousal and required intubation. A second CT scan showed multiple bilateral ICHs (Fig. 1B). Magnetic resonance (MR) imaging showed restricted diffusion in the left internal capsule, corresponding to his presenting stroke in addition to multiple other scattered punctate strokes bilaterally (Fig. 1C).

Admission labs showed elevated erythrocyte sedimentation rate (51 mm/h) and C-reactive protein (22.5 mg/dL), leukocytosis (11,500/mm3) and normal renal function. Work up revealed a very high C-ANCA (1:2560) and proteinase 3 antibody titers (> 8.0, reference: 0.0–0.3 AI), whereas myeloperoxidase antibody was negative. Dysmorphic red blood cells were identified on urine microscopy. Head and neck MR angiography, cardiac telemetry, and transesophageal echocardiogram were unremarkable, and blood cultures were negative. A diagnosis of C-ANCA-associated vasculitis was made and treatment was started with corticosteroids and rituximab. His course was complicated by renal failure, attributed to his vasculitis. He was discharged to an inpatient rehabilitation facility with marked neurological and renal improvement at 3 month follow up. His modified Rankin Scale was 3.
3. Discussion

Our case illustrates that AIS may be part of the initial presentation of AAV. If AIS is accompanied by systemic symptoms suggestive of a multisystem vasculitis, one should proceed with caution. In our patient, the diagnosis was not initially suspected. We treated the patient with IV tPA and subsequently multiple hemorrhages not reflecting the area of AIS developed. Because of the multiple hemorrhages, which may not have been secondary to acute ischemia, we evaluated him for a multisystem vasculitis including AAV. Subsequently, the elevated inflammatory markers, the C-ANCA titer, and positive proteinase 3 antibody led to the diagnosis of AAV. Thereafter, the renal and cutaneous manifestations of AAV became apparent. Our patient received immunosuppressive treatment with marked improvement in his non-neurological clinical features strengthening support for that diagnosis.

CNS involvement is rare in AAV and it usually develops as a delayed phase of the disease after onset of systemic symptoms. On occasion, it can be the initial manifestation of the disease, presenting as headache, cranial nerve palsy, seizures, altered mental status, coma, AIS and ICH [2]. Post-mortem exams in these patients have demonstrated AIS with areas of antemortem thrombi and fibrinoid changes with other areas demonstrating damaged vessels with hemorrhage [4]. We believe that in the presence of IV tPA, areas of fibrinoid necrosis are likely to make the patient more prone to hemorrhagic transformation particularly at sites of subclinical ischemia which may not be apparent on the initial non-contrast CT.

Fattahi et al. reported a patient with poorly controlled chronic EGPA who was treated with IV tPA for an AIS and developed a large ICH [5]. They proposed that IV tPA should be considered a relative contraindication in patients with AAV. Our experience with this case further supports their proposal with added caution to include patients who are still within their acute presentation of systemic vasculitis.

4. Conclusion

The risk of hemorrhage following IV tPA administration for treatment of AIS secondary to CNS involvement of AAV is not known and currently, AAV is not listed among the contraindications for IV tPA in guidelines [3]. Our case highlights the high risk of ICH associated with AAV and reinforces that IV tPA may be contraindicated in these patients.

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