Virologic analysis of tissues from a patient with giant cell arteritis (GCA) who was treated with corticosteroids and died of extensive necrotizing granulomatous arteritis revealed widespread varicella-zoster virus (VZV) antigen in multiple large arteries. Long-term treatment with corticosteroids likely potentiated VZV infection.

In 1997, a clinicopathologic case report described a 75-year-old woman with fatal aggressive steroid-refractory GCA that manifested as bilateral vision loss and myelopathy while on treatment with corticosteroids. The woman was not otherwise immunocompromised before becoming ill, and no cutaneous signs of herpesvirus infection developed during her 5-month illness. Bilateral temporal artery (TA) biopsies revealed GCA. Postmortem examination revealed spinal cord infarction secondary to extensive necrotizing granulomatous arteritis of spinal arteries.

Based on detection of VZV in GCA-positive TAs and documented involvement of other large arteries in most patients with GCA, we revisited this case and searched for VZV in the archived TAs and in other large arteries, the spinal cord, and brain. Immunohistochemical examination detected VZV antigen (figure) in both TAs, in the aorta, in the left carotid artery, and in an unidentified artery from the Circle of Willis but not in renal or mesenteric arteries. Viral antigen was not seen in the brain or spinal cord. DNA extracted from every section of each VZV antigen-positive artery was analyzed by PCR with primers specific for VZV and herpes simplex virus (HSV)-1 as described and revealed VZV DNA, but not HSV-1 DNA, in the unidentified cerebral artery from the Circle of Willis despite formalin fixation.

Discussion. The pathologic diagnosis of VZV arteritis comes nearly 20 years after this patient’s death. Reexamination of this case was prompted by the recognition that VZV is commonly found in the TAs of patients with GCA. VZV and extensive granulomatous arteritis in multiple large arteries was most likely due to reactivation of VZV in an elderly woman followed by potentiation of virus infection by several months of high-dose corticosteroids. Widespread vasculopathy occurs in patients with GCA treated with long-term corticosteroids. Recently, virologic studies in
a man with thoracic-distribution zoster and a history of corticosteroid abuse who died suddenly revealed extensive VZV infection in multiple organs and arteries, particularly the coronary arteries and aorta, along with subclinical VZV vasculopathy. Finally, our findings confirm detection of VZV antigen and VZV DNA in GCA-positive TAs and indicate that productive VZV infection in the TAs of patients with GCA parallels productive VZV infection in intracerebral arteries of patients with VZV vasculopathy. In fact, GCA is likely to be a form of VZV vasculopathy that predominately, but not exclusively, affects the TA. Because VZV is triggering the immunopathology of GCA, it is likely that treatment of GCA patients with antiviral agents will not only shorten the course of corticosteroids needed to reduce the immunopathology that produces disease but also prevent spread of VZV and the development of disseminated granulomatous arteritis.

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Widespread arterial infection by varicella-zoster virus explains refractory giant cell 
arteritis

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