

Coexistence of Giant cell arteritis with aortitis and Sweet's syndrome: is it a coincidence?

Abstract:

Sweet's syndrome (SS) is a rare disorder characterized by dermal infiltration by neutrophils. It ~~was~~ has been reported in association with drugs, malignancies, infections, rheumatoid diseases, inflammatory bowel diseases. Its association with ~~Giant-giant~~ cell arteritis (GCA) hasn't been reported ~~in~~ to our knowledge. The diagnosis of ~~Giant-giant~~ cell arteritis was based on inflammatory biological syndrome with aortitis with negative infectious investigations. Herein, we present an unusual case of SS associated with GCA treated with steroids with good outcomes.

Key words:

Aortitis, Sweet's syndrome, Giant cell arteritis

Introduction:

Sweet's syndrome (SS) is a rare disorder characterized by fever, neutrophilic leukocytosis, painful plaques ~~or~~ nodules on skin and dermal infiltration by neutrophils. Many associations have been reported in the literature, including malignancies, drugs, HIV infections, inflammatory bowel diseases, Behçet's disease, rheumatoid arthritis, thyroid disease. Its association with ~~g~~ Giant cell arteritis (GCA) has ~~not~~ been reported ~~to~~ in our knowledge. This case represents ~~an~~ unusual case of SS associated with GCA treated with steroids with good outcomes.

Case report:

A 67 year-old woman who had no medical history was hospitalized in April, 2018 in our department of internal medicine, for ~~exploration~~ survey of aortitis.

The patient suffered ~~since a month from~~ general weakness, weight loss, ~~pain tenderness over~~ of large joints and insomniac paroxysmic headaches ~~for one month~~. She had no ~~visual-ocular~~ manifestations. In our department, her blood pressure was 100/60 mmHg, ~~and~~ her pulse was at 80 ~~beats per battement~~ /minute (bpm). ~~Vascular-Physical~~ examinations including measurement ~~of~~ palpitation and auscultation of pulses in all major vascular regions as well as the temporal artery ~~found~~ ~~showed~~ wide and symmetric pulses. Skin examination found bilateral infiltrated erythematous plaques on the palms of the hands. Skin biopsy confirmed the diagnosis of Sweet's syndrome with evidence of dense dermal neutrophil infiltration: neutrophilic infiltrate with dermal edema, perivascular and interstitial infiltrate composed predominately of neutrophils and histiocytes (**Figure 1**).

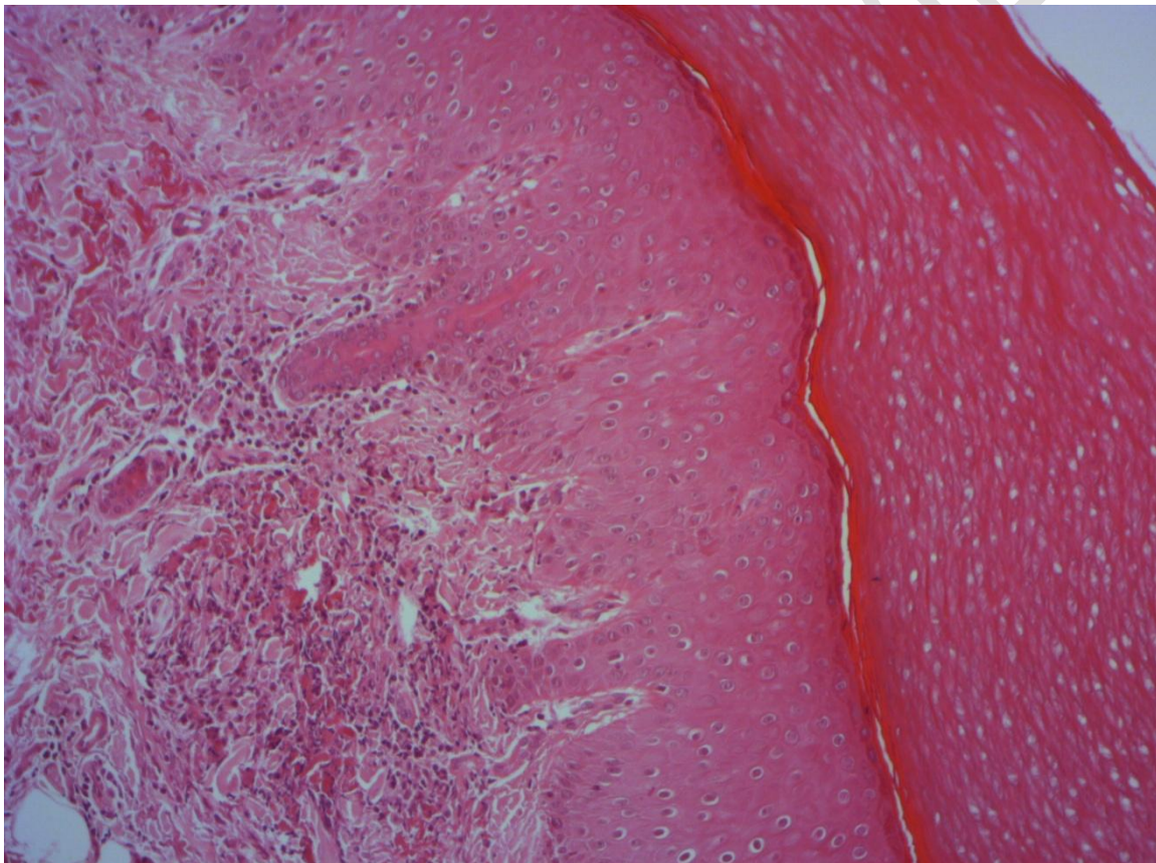


Figure 1: Diffuse neutrophilic and histiocytic dermal infiltrate in reticular dermis. Epidermis is normal. (HE x 200)

Biochemical findings revealed increased Erythrocyte sedimentation rate (125mm/hour) and elevated C-reactive protein (CRP) (146 mg/dL); Blood cell count showed white cell count at 11000/ μ L, haemoglobin ~~was~~ at 10.5 g/dL,

platelets of 320000/ μ L. Liver function tests were normal. Abdominal ~~m~~Magnetic ~~r~~Resonance ~~i~~maging (MRI) revealed thickening of the aortic wall with late peripheral contrast uptake and ~~peri-peri~~peri-aortic fat infiltration, with hyposignal T1-~~weighted sequence~~ and hyper-signal T2-~~weighted sequence~~, from T12 to L3. These findings were concordant with aortitis (**Figure 2**). Infectious investigations including viral hepatitis, ~~VHB, VHC, VIH, HBV, HCV, HIV,~~ tuberculosis, syphilis serology, ~~wright~~and widal-~~wright~~ tests were negatives. Temporal artery biopsy was normal. The diagnosis of GCA with ~~S~~Sweet's syndrome was retained. Corticosteroids were started at ~~a~~ high ~~dosagees~~ (1mg/Kg/day) for 1 month then progressively digressed. The outcome was favorable and she ~~had~~ no headaches. Inflammatory markers were within normal ranges (CRP = 5). During her check-up, she was free of complaints (prednisone 10 mg/day) however she developed diabetes and osteoporosis. After total duration of 32-~~months~~ of steroids intake, ~~steroid is stopped for~~ no flare up ~~of her disease was~~ noticed, ~~decision to stop steroids was proposed~~. Radiological control was programmed before their stop.



Figure2: Abdominal MRI, Axial section, aortitis

Discussion

GCA is a granulomatous vasculitis involving large and medium vessels, especially the extracranial branches of the carotid arteries. It ~~is's~~ the most common vasculitis in adults affecting people more than 50 years [1].

The frequency of aortitis in GCA is under-estimated. Aortitis represents a serious complication because of the risk of ~~anevrism~~aneurism, rupture and/ or dissection. It can be observed in initial presentation or occurs as a delayed complication. When it²s presents at the time of diagnosis of GCA, it seems to be associated with high risk of relapses and higher long-term vascular mortality rate [2]. Therefore, the screening of aortitis lesions at GCA diagnosis by an aortic CT-scan ~~was is~~ recommended [2]. Imaging studies showed signs of infra-clinical aortitis in 20 to 65% of cases at diagnosis [3]. Aortitis secondary to GCA is characterized in histological analysis by inflammation in media and adventitia with the presence of necrosis in the media ~~associated~~ and multifocal or transmural infiltrates of lymphocytes, histiocytes, neutrophils and polynuclear giant cells [4]. ~~Only aortic biopsy and histologic findings could relate aortitis to GCA.~~ In our case, clinical findings including headaches, deteriorated general status, and inflammatory biological syndrome were suggestive of GCA. Temporal artery biopsy wasn't contributive. Good outcome with steroid therapy was an ~~agreer~~argument in favor of diagnosis. Our patient ~~also had~~also Sweet's syndrome (SS) which was diagnosed simultaneously with GCA. ~~It's defined as an acute febrile neutrophilic dermatosis.~~ SS is characterised by fever and erythematous painful nodules, plaques, and/or papules localized on the face, trunk and limbs [5]. It may be associated with systemic diseases such as dermatomyositis, Sjögren's syndrome, Behçet's disease, and ~~T~~takayasu arteritis. Its pathogenesis is still unknown. Autoimmune and infectious factors seem to be involved [6, 7]. ~~Theis~~ ~~exceptionnal~~—association of SS with GCA and involvement of the aorta has not been reported. The validity of the coexistence of these two diseases remained to be established. Few cases reported the coexistence of aortitis associated with SS had been published [8, 9]. In fact, vasculitis may be observed in SS, [10] leading us to question if the aortitis is a part of SS or GCA or both. Arthralgia also could be multifactorial. They are observed in both SS and GCA. Articular manifestation could be seen in 20 to 30% of adult patients with SS [11, 12]. Although the etiology of SS is not completely understood, an inflammatory component was involved. There is no consensus about treatment of aortitis in GCA or SS, but ~~they are sensible~~it may ~~be effective~~ for systemic steroids [2, 13]. In our case, we noticed persistent clinical and biological improvement after total duration of 32 months of steroids. Radiological control is scheduled before their stop.

Conclusion:

The association of inflammatory diseases such as Sweet's syndrome and giant cell arteritis is exceptional-unclear and even with involvement of the aorta. Biological-inflammatory syndrome Neurological manifestations, abnormal blood pressure with specific skin findings in elderly must lead us to suspect giant cell arteritis Horton arteritis and often normal temporal artery biopsy couldn't eliminate-totally exclude the diagnosis. Both diseases had good outcome with steroids if diagnosed and treated early.

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