A Diagnosis Missed for Several Years-
Wegener’s Granulomatosis

Paudyal BP, Pantha S, Ranjitkar N, Manandhar A, Arjyal A

Department of Medicine
Patan Academy of Health Sciences
Patan Hospital, Lalitpur, Nepal

Department of Community Health Sciences
National Academy of Medical Sciences
Kathmandu, Nepal

ABSTRACT
Wegener’s granulomatosis is a form of systemic vasculitis of small to medium sized vessels and affects upper respiratory tract, lungs and kidneys along with various organs. It causes necrotizing granulomatous inflammation of the affected parts and presents with positive antineutrophil cytoplasmic antibodies in more severe forms. Being a systemic disease with the potential to affect any organ-systems with a wide range of clinical presentations, it is associated with a risk of delay in diagnosis with resultant setback in institution of appropriate treatment. Confusion may arise due to an extent of histological similarity between Wegener’s granulomatosis and the more prevalent tuberculosis, both causing granulomatous inflammation of the affected parts. Here, we present two cases of this rare disorder where the diagnosis was missed for several years in the beginning causing a delay in institution of specific therapy which led to the development of complications.

KEY WORDS
delayed diagnosis, multisystem disease, systemic vasculitis, Wegener’s granulomatosis

INTRODUCTION
Wegener’s granulomatosis (WG), first described in 1936, is a chronic systemic vasculitis that classically affects the upper respiratory tract, lower respiratory tract and kidneys. It affects small to medium sized blood vessels and causes necrotizing and granulomatous inflammation in the nose, paranasal sinuses, eyes, and lungs, necrotizing glomerulonephritits, and systemic involvement of many organ systems. It can manifest in two forms: limited disease without and generalized disease with renal involvement; most of the cases of generalized disease are associated with the presence of cytoplasmic anti-neutrophil cytoplasmic antibodies or c-ANCA.

Owing to the systemic nature of the disease with a wide range of clinical presentations and the potential to involve any organ system in the body, patients often present to different specialists and the absence of a unifying hypothesis that explains all the features leads to piecemeal diagnosis delaying the already difficult diagnosis and appropriate treatment is often delayed. On the other hand, because of a low index of suspicion of WG due to its rarity, and confusion with other granulomatous diseases, particularly widely prevalent tuberculosis (TB), many patients are treated with antitubercular drugs causing a further delay in appropriate treatment. So it is not uncommon to see complications like irreversible organ damage like end stage renal disease by the time the disease is diagnosed. Here, we present two cases of WG, which were both histologically and serologically confirmed. Both of them offered many clues for diagnosis in earlier hospital visits, but it took two years for the first patient and eleven years for the second patient before a final diagnosis was made and appropriate therapy was started.
CASE ONE

A 61 year-old female presented to the Department of Ear, Nose, and Throat of a tertiary care hospital with a history of progressive throbbing headache and nasal bleeding of three months’ duration. Examination revealed a left sided nasal mass and a CT scan of Paranasal Sinuses (PNS) showed a soft tissue lesion in the left nasal cavity and in all of the left sided PNS suggesting polyposis. Histopathological examination of the endoscopically debulked mass revealed dense inflammatory cells with multinucleate giant cells and the presence of neutrophilic inflammatory cells infiltrating the vascular wall with areas of necrosis. Anti-tubercular therapy (ATT) was started on the basis of histological finding of granulomatous inflammation.

However, her headache and nasal bleeding did not improve even after several weeks of ATT. Six weeks after the onset of nasal symptoms, the patient experienced painful red eyes. Ophthalmic evaluation revealed a bilateral scleritis and oral prednisolone was started. After a few weeks, she developed low grade fever, cough with blood streaked sputum, and joint pains. The patient was taken to a tertiary care hospital in a neighboring country where as an inpatient she developed mononeuritis multiplex in both feet, and a large necrotic ulcer at the site of intravenous canula in left forearm. As the patient’s condition progressively deteriorated, she was asked to return back to Nepal. A repeat ophthalmic consultation in Kathmandu suggested bilateral ulcerative keratitis with anterior scleritis (Fig 1). USG of orbit suggested a pseudotumour within extraocular muscles. She was immediately referred to our hospital for further management.

At the time of admission in our hospital her BP was 150/90mmHg; with normal pulse and temperature. She had bilateral ocular inflammation. A polypoid growth was visible in the left nasal cavity. There were multiple ulcers in the oral cavity, with another 3x3cm necrotic ulcer in the ventral aspect of the left forearm. The breath sounds were diminished in both lung bases. There was bilateral foot drop. Investigations revealed WBC 13,200/μL (neutrophils 81%, lymphocytes 13%), Hb 11.1g/dL, platelets 539000/μL, and ESR 118 mm in 1st hour. The creatinine was 1.4mg/dL while the electrolytes and liver enzymes were normal. Chest x-ray showed multiple nodular opacities in the right lung with slightly enlarged cardiac silhouette. An echocardiography revealed mild pericardial effusion. The Mantoux test revealed 15 mm induration after 72 hrs of inoculation. Urine samples showed trace albumin, and persistence of microscopic haematuria. CT scan of the chest showed peripherally located irregular, non-enhancing nodules in both lungs with pleural and pericardial effusion (Fig 2).

MRI of lumbosacral spine showed mild degenerative changes in the lumbar spine without any compromise of the central canal and neural foramina.

Further investigations revealed negative serologies for hepatitis B, C, and HIV. The rheumatoid factor (RF) was positive and the anti-nuclear antibody (ANA) was negative. The c-ANCA was positive by indirect immunofluorescence method.

On the basis of multisystem nature of the disease, raised inflammatory markers, positive c-ANCA and biopsy findings, a diagnosis of generalized Wegener’s granulomatosis was made. ATT was discontinued and Methylprednisolone pulse therapy was administered followed by tapering dose of steroids, and oral cyclophosphamide was instituted. The patient steadily improved and all of the features of active disease gradually disappeared. Now after 18 months of initiation of treatment, she enjoys a disease-free state maintained with low dose Azathioprine.

CASE TWO

A 38 year-old woman was referred to our hospital for the management of recently developed acute renal failure. The patient reported that she had a sudden onset of decreased urine output followed by progressive increasing swelling of body and shortness of breath. This followed a febrile illness 2 months ago. The renal function at the time of presentation was: urea 158mg/dL, creatinine 10.6mg/dL,
sessions of maintenance haemodialysis every week. One month after the initiation of immunosuppressive treatment, she was much better with improvement in the systemic symptoms and tendency towards lesser requirements for haemodialysis.

DISCUSSION

WG is a rare disease, with the involvement of multiple organ systems; hence it is not uncommon for the affected patients to seek help from many physicians in many specialities. Majority of the patients with WG present with upper airway disease, occurring in more than 90% of cases. Common upper airway presentations include recurrent rhinosinusitis, nasal polyposis, nasal chondritis leading to saddle nose deformity, nasal septal perforation, serous otitis media, hearing impairment, and stridor due to subglottic stenosis. Lung involvement is one of the cardinal manifestations of Wegener’s granulomatosis and is seen in up to 87% of patients. Common pulmonary manifestations include cough, haemoptysis, and shortness of breath with the underlying lesions as cavities, nodules, and fixed infiltrates. Renal involvement on kidney biopsy is present in up to 85% patients during the course of disease but in the majority the extra-renal disease often precedes the renal disease. The common renal manifestations are asymptomatic haematuria and proteinuria; however, it may progress to rapidly progressive glomerulonephritis (RPGN) with varying degrees of renal impairment as the disease becomes more severe. Immunohistologically, the glomeruli have no deposits of immunoglobulins; hence ‘pauci-immune necrotizing and crescentic glomerulonephritis’ is sometimes used to describe this pattern. Eyes can be affected with episcleritis, retinal vasculitis and
Case Report

pseudotumour formation being common manifestations. Inflammatory arthritis, muscle pain, and variety of skin lesions ranging from palpable purpura to pyoderma gangrenosum are musculoskeletal manifestations of WG. Apart from these, any other organ systems could be affected by the vasculitic process, for example, nervous system, heart, gastrointestinal tract, etc. Constitutional symptoms like fever, weight loss, fatigue can be present at any time during the course of the active disease.

Both of our patients had presented with upper airway symptoms at the onset of the disease. As the disease progressed, these cases developed features of other organ involvement particularly eyes, skin, lungs, peripheral nerves, heart, and kidneys. Both the cases had prominent musculoskeletal symptoms during the course of disease. The first case developed prominent pulmonary involvement with lung nodules whereas the second case had a very severe renal disease (RPGN) requiring maintenance haemodialysis.

The serological marker of WG is the presence of c-ANCA. The c-ANCA pattern is seen by indirect immunofluorescence in a majority of active cases of WG (55% with limited and 88% with systemic disease). These antibodies are directed specifically against proteinase - 3 (PR-3) in the azurophilic granules of neutrophils and the specificity of anti PR3 antibodies in the detection of WG is 99%. Both of our patients were positive for c-ANCA, meaning the disease process was active at the time of presentation at our centre.

Biopsy remains the gold standard for diagnosis of several types of vasculitis, particularly the small- to medium-vessel vasculitis like WG. Inflammatory lesions in WG typically include necrosis, granulomatous changes, and features of vasculitis. Though, vasculitis and granuloma formation can occur in the same lesion, it is unusual to see both in the same biopsy specimen. Moreover, renal biopsy in WG typically shows necrotizing inflammation without granuloma formation. Granulomatous inflammation, with features of vasculitis was present in both of our cases with features of necrosis as well in the first case. However, incorrect interpretation of the biopsy findings led to the delay in the initiation of specific treatment in both the cases.

WG and TB may present with similar clinical and histological findings; and one may confuse while taking care of such cases. Incorrect diagnosis of TB in a patient with WG may lead to the delay in institution of appropriate therapy with the development of resultant complications; on the other hand incorrect diagnosis of WG in a patient with TB leads to inappropriate treatment with immunosuppressive drugs, and this may even herald the development of military tuberculosis. However, with careful assessment of the progression of the disease, radiological characteristics, presence of c-ANCA and careful review of the results of the histological examination, it was established that these patients had WG.

These cases highlight the importance of considering WG in differential diagnosis of patients with multisystem disease. Close communication between physicians of different specialties is essential; at the same time an effective interaction between a clinician and a pathologist is equally important to avoid an erroneous diagnosis in the presence of equivocal biopsy test results.

REFERENCES
11. Harrison NK and Knight RK. Tuberculosis of the nasopharynx misdiagnosed as Wegener’s granulomatosis. Thorax 1986; 41: 219-220