Wegener's Disease in a Black Malian

Dr. Sanogo A1,4, Diaby Lm2,3, Diollo L1, Maiga3,4, Kane AST1,3, Camara Y6, Sow S1, Diakite Y3, Kaya Assetou Soukho5

1Infirmary of the Internal Medicine Service of BAMAKO hospital
2Infirmary of the Internal Medicine services of Kati hospital
3Hospital of Sikasso Mali
4D.C.S.S.A Mali
5Internal Medicine Service CHU Point G Bamako
6Central Military Hospital Niger

DOI: 10.36347/sjmer.2021.v09i01.004 | Received: 13.12.2020 | Accepted: 24.12.2020 | Published: 09.01.2021

*Corresponding author: Dr. Abasse Sanogo

Abstract

Wegener’s disease is a very rare condition in Africa. In 1931, a German student, Heinz Klinger, reported an unusual observation of peri-arteritis, this was characterized by pneumatic-renal syndrome, severe airway involvement including rhinitis, nasal saddle, pansinusitis, middle ear, laryngo-tracheal and bronchial ulcerations and one Exophthalmos. Autopsy examination revealed the presence of necrotizing and granulomatous vasculitis in the upper and lower airways, associated with increasing glomerulonephritis[1]. The objective of study is to show the existence of Wegener’s cases in our country. Observation: this is a 28-year-old patient followed by ENT for nasal-sinus polyposis, hospitalized for chronic cough with sputummuco-purulent sometimes streaked with blood with deterioration of the general condition; pathological examination of the biopsy nasal specimen concluded with the diagnosis of Wegener’s granulomatosis. Discussion: GW can be seen at all ages the average age at diagnosis is between 40 and 50 years, with a slight male predominance in some series (table 23-11), which does not appear to be significant in the studies epidemiological[6]. Less than 15% of cases are pediatric. The presence of high titer ANCA with antiproteinase 3 specificity is a strong argument for the diagnosis of GW; on the other hand, the negativity of ANCA should not make the diagnosis rejected, in particular in the face of a form of limited appearance. Histology remains a powerful and often essential diagnostic element, because, as often in systemic granulomatosis, the diagnosis can be discussed at the borders of infections caused by intracellular bacteria and lymphoproliferative syndromes (in particular lymphoid granulomatosis of liebow)[80].

Keywords: Wegener, granulomatosis; black; Malian.

Copyright © 2021 The Author(s): This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0 International License (CC BY-NC 4.0) which permits unrestricted use, distribution, and reproduction in any medium for non-commercial use provided the original author and source are credited.

Review Article

INTRODUCTION

Wegener’s disease is an extremely rare condition in Africa. In 1931, a German student, Heinz Klinger, reported an uncommon case of Polyarteritisnodosa. It was characterized by a pulmonary renal syndrome, a severe impairment of the airwayswith rhinitis, saddle nose, pan sinusitis, otitis media, laryngotracheobronchitis and exophthalmos. Autopsy examination revealed the presence of necrotizing and granulomatous vasculitis of upper and lower airways, associated with ascending glomerulonephritis[1]. It was in 1936 and 1939 that Friedrich Wegener, affected by the precocity and intensity of the damage to upper airways, described three cases of the disease which bears his name[2]. The main anatomical features of Wegener granulomatosis (GW) were documented in a study of 7 cases by Godman and Churg [3] in 1954, which emphasized the rapidly fatal nature of the disease. Carrington and Liebow [4] proposed in 1966 the concept of limited form of GW, characterized by predominant pulmonary involvement, the absence of renal damage and a better prognosis compared to the classical diffuse form: with a decline of 4 to 150 months, half of their patients were alive. By 1973, Fauci et al. [5] suggested the efficiency of the combination of corticosteroids and oral cyclophosphamide, which allowed remission in 93% of the cases. A renewed interest for this disease came from the demonstration of specific autoantibodies against a constituent of the cytoplasm of neutrophils (ANCA Anglo-Saxon authors). ANCAcs have emerged as a new diagnostic tool and an etiopathogenic component of systemic vasculitis in the broad sense and GW in particular.

Epidemiological data come only from Anglo-Saxon sources. In the United States, the prevalence of GW, estimated from hospital registers, is 3 per 100000 population [6]. Geographically, it decreases according to...
GW histologically associates 3 main lesions: giant cell granuloma, necrosis and vacuities. These 3 elements, analyzed in isolation; are encountered in 20 to 30 % of the upper airways biopsy however, the complete triad is observed in only 3 to 16 percent of the cases [12]. The diagnostic usefulness of the biopsy depends mainly on the size of the swab, which is the maximum for open-chest biopsy. Heterogeneity in the distribution of lesions, noted over serial sections within a single biopsy, appears in itself a diagnostic guide. The cellular inflammatory infiltrate is a polymorphic granuloma that typically has a palissadichistiocytic infiltrate and many multinu key giant cells. The presence of neutrophils or eosinophils and plasma cells is less specific. The granuloma may be intra-, peri- or extravasularly situation. It is sometimes centered on non-caseous necrosis. Necrotizing vasculitis is acute, circumferential; it concerns arterioles and small veinlets. It may appear isolated, without granuloma or tissue necrosis. Capillarity is possible, mainly involved in necrotizing glomerulonephritis and intra-alveolar hemorrhage. Thrombosis of large arteries, responsible of ischemic necrosis seems underestimated as it has a significant role in the case of CO fail fatal multiorganance [13,14]. Necrosis can adopt extravasular tissue topography, without coexisting with a phenomenon of vasculitis. Moreover, the presence of micro Polynuclear abscess is quite common and especially precocious, constituting a good presumptive argument in cases of incomplete histological evidence. Lesions predominate in the upper airways, lung and kidney. The respective distribution of vasculitis and granuloma varies according to organs: at autopsy, a granuloma is present in the lung in 81% of the cases, in the kidney in 66% of the cases, in the spleen in 55% of the cases, in the upper airways in 51% of the cases and in the heart in 11% of the cases. Vasculitis lesions predominate in the lung (87%), kidney and spleen (77%), and are present in the upper airways only 25% of the cases [14].

The accountability of an infectious agent colonizing the upper airways has been raised since the first descriptions [15] but no evidence of causal infection has been found so far, including the study of the washing liquidbronchoalveolar [15]. However, a starting role is likely:

- a viral or bacterial infection would precede a push in 45% of renal relapse cases [15];
- a seasonal predominance (spring, winter) of the inaugural thrust has been noted in some series [11];
- chronic nasal portage of staphylococcus coagulase positive is associated with a significantly higher frequency of relapses [17];
- an antibiotic, cotrimoxazole, appears to be effective in the treatment of certain limited forms of GW [18] and in the prevention of relapse [19]. Chronic parvovirus B19 infection, suspected to be associated with GW [20, 21], was "cleared" by the constant negativity of viral nucleic material research in 42 adults with Wegener disease of recent appearance [22]. The existence of a genetic predisposition remains controversial: a significant association, in small rule, was found, in a variable way according to the authors, the studied ethnic groups and the techniques used, as well with the HLA antigens of class 1: HLA-B7, B8 [23], B50 [24], than Class 2 [25]: DR1 [26], DR2 [23] and DR9 [28]. Family forms are also exceptional [29-31]. Autoantibodies against cytoplasm of neutrophils and monocytes (antineutrophilcytoplasmicantibodies, ANCA) have acquired their status as a marker of GW [32, 33]. ANCs are not just a diagnostic test or scalability of vasculitis.They are also a conceptual and nosological tool: GW is now an autoimmune disease [34]. The presence of ANCA in the serum of patients with necrotizing glomerulonephritis was initially observed in 1982, but the link with Wegener's disease was only formally established in 1985 [35, 36]. The antigenic targets of ANCs are mainly enzymes contained in the cytoplasmic granules of neutrophils.

The aim of this study is to show the existence of Wegener cases in our county.
**Observation**

I mean a 28 year old patient followed ENT for nasal polyposis - sinussen hospitilazed for chronic cough with muco-purulent sputum sometimes streaked with blood and a poor general condition. Her temperature was 40°C with oscillation. On the ENT examination there was ulceration of the cornet with pus at the left sinus punctiform; parotid bilateral inflammatory hypertrophy.

The pleurapulmonary examination had found diffuse bilateral crackles. Dermatologically, papular macules was noted on the thorax and on the face. Elsewhere there was diffuse bilateral and peripheral joint pain.

We had found the paraclinical examination: Biology showed leukocytosis 262,100 cells / mm3 and anemia to 6.7g / l; an absence of proteinuria to the test strips. The dosage of c-ANCA by IF1- ino- IL was positive at 28 IU / L.

The radiograph of the frontal thorax showed multiple bilateral macromolecular images, diffuse, excavated peripheral. Sinus CT showed osteolysis and chondrolysis of the nose.

Pathological examination of nasal biopsy at a Wegener granulomatosis. We used concomitant prednisolone at a dose of 1.5 mg / kg / day and methotrexate at a weekly dose of 25 mg / week for injection plus 15 mg folic acid per week.

There was a clear improvement of the disease during the first two (weeks) of the hospitalization. After release of the patient, she had a pulmonary complication that led to the emergency and the death was found on arrival.

**Discussion**

GW can be observed at all stages of life. Average age during diagnosis is between 40 to 50 years, with a slight male predominance in some series (Table 23-11), which does not appear to be significant in epidemiological studies [6]. Less than 15% of the cases are pediatric [37]. The delay of average diagnostic is one year. This delay is in fact extremely variable, significantly longer in forms without renal involvement [4, 38] where the evolution is often insidious, interspersed with spontaneous remissions of several months or years. In the study of NIH (National Institutes of Health, Bethesda, USA), the diagnosis was delayed by 5 to 16 years in 8% of the cases [39]. The onset is generally marked by nasal or respiratory symptoms of commonplace, including recurrent evolution, resistance to antibiotic treatment but sensitive to short corticosteroid therapy, association with fever, progressive deterioration of general condition or visceral involvement should receive attention. Elsewhere, the evolution can be done in an acute mode with the installation in a few weeks of a diffuse attack from the outset, both ENT, pulmonary and renal.

The radiological aspect is of a great diversity. The CT scan of the chest is essential, to highlight non visible lesions on X-rays and to clarify their appearance. Typically, it is about nodular opacities, of variable size (from a few mm to several centimeters), of sometimes pseudo-tumoral aspect, most often multiple, bilateral, well limited, without particular topographic distribution and evolving towards the excavation in half of the cases. Infiltrates are also common and can be distinguished into 3 types [40]:

- diffuse alveolar opacities, bilateral, sparse and asymmetrical, which lead to intra-alveolar haemorrhage
- Opacity low density but localized, variable from one cliche to another and finally
- Dense opacities, localized and able to develop towards the excavation.

The spontaneous disappearance of parenchymal lesions is possible, but rarely is it durable [4, 41]. In some cases, radiological abnormalities remain stable for months or even years without evidence of systemic spread of the disease [41]. The simple excision of nodules can exceptionally remove the pulmonary symptomatology and general signs in the absence of anti-inflammatory treatment in a limited form [41] but such an evolution cannot be expected. A diffuse interstitial infiltrate is exceptional and must seek a diagnostic alternative. Treatment progress is usually towards the rapid disappearance of lesions, sometimes at the cost of scarring emphysema where aspergilloma may be grafted [42], or rarely calcification [40, 5]. Intravenous haemorrhagealveolar, responsible for acute respiratory failure sometimes fatal, affects 5 to 13%of the patients with a lethality of the order of 50%.100 [40, 43, 44, 45, 46], The possibility of pneumorenal syndrome with concomitant presence of glomerular basal antinuclear antibody and ANCA has been reported [47].

Bronchial stenosis is the same mechanism as subglottic stenosis.

**Mucocutaneous damage**

Mucocutaneous involvement affects about half of the patients [48, 49, 50, 51, 52, 53]. It is present initially in 4 to 73 % of the cases according to the series. Very polymorphic, more common in diffuse forms [49, 50], it is dominated by infiltrated purpura, papules, subcutaneous nodules, pustules and hyperplastic gingivitis. Macular or maculopapular rash [5, 4], genital ulceration, gangrene of the extremities are more rare. Cutaneous ulcers may cause a pyoderma gangrenosum [55]. Other manifestations are anecdotally reported: painful induration of the
postoperative scar, palpebral xanthoma accompanying a pseudotumor of the ipsilateral orbit [50, 56], erythema elevatum diutinum [57].

Histological examination, easy to perform, reveals lesions of non-granulomatous vasculitits in more than three quarters of the cases. Granulomatous extravascular lesions occur only in 5% of the cases. A palisadic granuloma, centered by extravascular necrosis occurs in 10% of the cases [48-51]. The presence of granulomatous vasculitis seems rare. The immune fluorescence study may highlight deposits of IgG, IgA or IgM, C3, of fibrin on dermal vessels [49-51] or may be negative [49]. Purpura lesions are never granulomatous and correspond to a non-specific leukocytoclastic vasculitits; their biopsy is thus of poor diagnostic efficiency and we will rather seek to take a nodular lesion [48, 50].

**Joint and muscular damage**

Arthralgia, or less commonly arthritis, occurs in 60 to 76 per cent of the cases, [3, 59], it occurs initially in half of these cases. The damage is polyarticular and symmetrical in two thirds of the cases, more rarely monoarticular. The wrists, knees and ankles are the joints most commonly affected. If an arthritis symmetrically affects the extremities and if is accompanied by rheumatoid factor, it may lead to an early case of rheumatoid arthritis. However, the development is neither deforming nor destructive, although discrete erosive arthritis has been described [58, 59]. Synovial biopsy may reveal granulomatous synovitis. Myalgia is frequent, rarely reported to granulomatous acute myositis [60]. It may suggest Horton's disease or rhizemelic pseudo-rheumatoid arthritis, especially as temporal arteritis is possible [61, 62]. The diagnosis is then rectified by a secondary appearance of the damage of the airways.

**Biology**

Neutrophil leukocytosis, inflammatory anemia, thrombocytosis and elevation of inflammation proteins are the rule in diffuse forms. A leukopenia is exceptional [63] and must call into question the diagnosis. Eosinophilia occurs in 10% of the cases [13, 54, 67]. It is generally moderate, between 500 and 800 / mm3, higher numbers being nevertheless possible [54]. A form with predominant tissue eosinophilia, but without blood eosinophilia or asthma, has been described [65]. It is thought to be a borderline form with Churg and Strauss syndrome, especially since familial or personal allergic terrain (85 p. 100) would be more common in patients with GW [66]. Serum rheumatoid activity is detected in approximately half of the cases, rarely associated with the presence of cryoglobulin. There is usually no Hypergammaglobulinemia polyclonal. The serum complement and its actions are not lowered [5]. Search for antinuclear factors [5, 54, 64] and antigenemia HBs [5, 64, 67] is negative. Hypercalcemia by secretion of a vitamin D metabolite or by association with hyperparathyroidism has been described. The cytokine dosing has been studied in short series. An increase in the level of interleukin2 and interferon alpha [68] occurred. The serum level of the soluble receptor for interleukin-2 would be correlated with the progression of the disease, the increase of which could lead to a clinical relapse [6-9]. A rise in plasma level of thrombomodulin in the active phase of the disease is evidence of the extension of vascular lesions [71]. The increase in serum neopterin, secreted by macrophages under the influence of gamma interferon, seems correlated with the activity of the disease and the presence of infectious complications [72], the search for ANCA (see Pathogenesis and Diagnosis) is first performed by immune fluorescence screening. The ELISA technique subsequently makes it possible to type the antigenic reactivity of the positive sera and to titrate antiproteinase 3 antibodies.

Histological evidence of GW may be difficult to obtain because biopsies rarely associate the characteristic triad, which may warrant multiple and sometimes surgical specimens. As a result of these difficulties, the ACR proposed classification criteria to support the diagnosis of GW with 88.2% sensitivity to demonstrated vasculitis and a specificity of 92 % [73]. The Chapel Hill consensus conference emphasized the histological aspect, defining GW by the presence of granulomatous inflammation of the airways and necrotizing vasculitis of vessels of small and medium caliber [74]. In fact, these classifications offer little diagnostic value at the individual level, but they make it possible to analyze relatively homogeneous groups of patients in clinical studies [75]. The detection of ANCA uses indirect immunofluorescence, the value of which depends very much on the experience of the operator. In practice, solid phase methods that use purified antigens are the indispensable complement of the immunofluorescence screening technique; they make it possible to determine the specificity and to quantify the ANCA [76]. antiproteinase c-ANCA 3 have 90 % sensitivity in generalized and active Wegener's disease, which decreases to 60% when the disease is localized; overall it is 66% [75 ], c-ANCA meet with great specificity, of 98%, in a spectrum mainly covering GW but also micro polyangeite and some glomerulonephritic croissants. They are virtually absent in "classic" polyarteritis nodosa. The positive predictive value of c-ANCA is only 63% [77 ]. A significant, as a rule, weak title of c-ANCA has been found in some patients with various conditions: uveitis, AIDS, hepatitis C, mycobacteriosis, aspergillosis, cystic fibrosis, infectious endocarditis, non-Hodgkin's lymphoma, rheumatoid arthritis. In these "entire false positive" cases, the search for antiproteinase 3
antibodies was negative, or unrealized. In contrast, the majority of sera tested in patients with hepatic amoebiasis appear to be associated with specific ANCA activity of proteinase 3 [78]. The p-ANCA are less specific and can be observed in some systemic vasculitis, such as allergic vasculitis, Churg-Strauss, in glomerulonephritis croissant, rheumatoid arthritis, induced lupus, polychondritisatrophic or inflammatory colitis, or even in healthy subjects [32, 79]. The differential diagnosis between GW and PCA, sometimes difficult, is discussed in the Polychondritechapteratrophic (637). In practice, in front of a suspect patient of GW, one must be able to identify subclinical ENT, pulmonary, neurological or renal impairment by proposing:

- A sinus scan and an ENT examination;
- A thoracic scanner of high resolution in thin sections;
- A possibly electromyogram;
- Repeated analysis of urinary sediment and proteinuria

The presence of high ANCA with a specificity of antiproteinase 3 is a strong argument for GW diagnosis; on the other hand, the negativity of the ANCA does not have to reject the diagnosis, especially in a "limited" appearance. Histology remains a powerful and often essential diagnostic element because, as often in granulomatosis systemic, the diagnosis may be discussed in the borders of intracellular bacterial infections and lymphoproliferative disorders (especially granulomatosis lymphomatoid of Liebow) [80]. The negativity of ANCAs and the lack of improvement during treatment should suggest the condition recently described in subjects with a genetic deficiency of TAP complex (transporter associated with antigen presentation) [81].

**Treatment**

Before the use of glucocorticoid therapy, GW was considered fatal within an average of 5 months, mainly due to end-stage renal failure before the advent of hemodialysis. These old statistical data do not account for the clinical polymorphism of the disease, with fluctuations in activity and sometimes prolonged spontaneous survival, especially in localized forms. The current treatment of GW has an overall satisfactory efficacy at the cost of a significant iatrogenesis, not only in terms of morbidity but also of mortality related to its administration duration. In addition, some of these complications (infections, allergies, renal failure, hematuria ...) are difficult to distinguish from a specific disease. Therapeutic innovation therefore aims above all to develop, in addition to new drugs with a more eradicator aim, administration techniques or even less toxic sequential schemes. Thus the distinction between treatment of attack and maintenance, if it seems a little artificial, has the merit of designating 2 essentially different therapeutic situations of purpose: the obtaining of a complete remission and the prevention of relapses.

**Initial treatment**

**Corticosteroids**

Conventional therapy involves the combination of prednisone at the initial dose of 1 mg / kg / day and a cytotoxic agent. Indeed, the use of corticoids alone in the treatment of stroke, especially in cases of diffuse disease with renal damage, does not allow to obtain a complete remission, the median survival being only 5 months (without corticosteroids) to 12 months [82]. The initial dose of prednisone is maintained for a duration of an average of 1 month [5, 44], which most often makes it possible to objectify a resolution of signs of activity. Fauci then calls the passage to an alternating corticosteroids [5], the total duration of the corticosteroid therapy is 12 months [5, 83], shorter than in other teams.

**Methotrexate**

The prednisone and methotrexate combination was proposed first line in 42 patients with histologically proven GW patients but not involving life-threatening speedily (creatinine less than 220 umol/l, absence of acute hypoxemic intra-alveolar haemorrhage area...) [83]. In 15 cases, it was an inaugural push. Methotrexate was administered orally, initially at a dose of 0.3 mg/ kg/ week without exceeding 15 mg/ week for 1 to 2 weeks, and then gradually increased by weekly increments of 2.5 mg to 20-25. mg/ week. Decreases in prednisone doses were based on the standard NIH protocol, with complete discontinuation occurring after an average of 7 months. Complete remission was obtained in 71% of the cases within an average of 4.2 months. Pneumocytosis of early onset was observed in 3 cases including 2 deaths in patients treated with methotrexate and high doses of corticosteroids. The authors suggest the systematic prophylactic use of cotrimoxazole (at a low dose of 3 tablets per week), which is theoretically contraindicated because of the expected bone marrow toxicity associated with these antifolicdrugs. Methotrexate may therefore be an alternative to the cyclophosphamide, either immediately in some limited forms of GW, or secondarily after conventional short-term induction therapy in diffuse forms of the disease [84].

**Cotrimoxazole**

Cotrimoxazole was proposed in 1985 in monotherapy, mostly on empirical grounds, [85] as granulomatous initial treatment of subacute and localized Wegener. The use of this treatment, as a first-line treatment, has been the subject of controversy which remains valid, in the absence of a methodologically satisfactory assay [86]. A recent, prospective, open, non-randomized study was conducted in 19 patients with ENT-localized disease, without systemic vasculitis or significant impairment of general health state. The use of Cotrimoxazole alone, at
a daily oral dose of 2 tablets of Bactrim forte®, resulted in 11 remissions (6 complete and 5 partial) with an average duration of 17 months (6-88 months). In 5 patients, locoregional progression was clear and in 3 other non-responders the disease was secondarily generalized within 24 to 60 months. In this study, the overall effectiveness was 58% [87]. The use of cotrimoxazole as a first-line treatment appears “reasonable” only in the rare forms localized to the upper airways, with no general signs [88].

Conversely, because of the high risk of potentially fatal pneumocysts [89-92], co-trimoxazole prophylaxis should be added to corticosteroid and immunosuppressive therapy during the attack phase. It is to say during the first year of treatment, and especially as there is a global lymphopenia (largely iatrogenic) sometimes deep. Pentamidine aerosols have also been proposed, particularly in patients treated with methotrexate [83].

**MAINTENANCE THERAPY**

The evolution of the GW is marked by a high rate of relapse, of the order of 30 to 50 p. [5], these relapses occurring at a variable distance from the first thrust, until 20 years after stopping any treatment. The increase in c-ANCA titers was initially considered a predictor of GW clinical outcome [93]. Some authors have even proposed to strengthen treatment on this single biological argument [94]. The NIH study, which covers the largest published series of Wegener’s disease, has actually shown that ANCA titers are paralleling 64% clinical activity. In 100% of the cases, the ANCA elevation precedes the push in only 24% of the cases [9 5]. Conversely, in 36% of the cases, the clinical and biological evolutions were uncorrelated or even completely discordant. The increase in ANCA titers was re-evaluated: this is a relapse index with a sensitivity of 43% with a positive predictive value of 23%, therefore quite weak [96].

This significant risk of relapse explains why immunosuppressive therapy is usually prescribed for at least 1 year after obtaining complete remission [83, 97]. Dose reduction is always gradual before complete cessation. To avoid the cumulative toxicity of cyclophosphamide, sequential alternative treatment regimens, first using oral cyclophosphamide for a limited period of time to achieve complete remission, followed by relay therapy with another drug (cotrimoxazole, methotrexate, azathioprine), are under evaluation.

In a double-blind, placebo-controlled study, cotrimoxazole (2 tablets of Bactrim forte daily for 2 years) significantly reduced relapse rate in patients in complete remission, whether they were treated in combination (approximately half of all cases) or not with prednisolone and / or cyclophosphamide. At 2 years old, 82% of patients treated with cotrimoxazole were still in remission compared to 60% of those treated with placebo. This significant reduction in the rate of relapse was a parallel to the decrease in the frequency of infections and unrelated to the evolution of the ANCA titer [98].

The use of methotrexate has also been proposed after obtaining complete or partial remission by conventional treatment. An open study compared four groups of patients receiving methotrexate intravenously or cotrimoxazol, with or without corticosteroids [99]. The previous duration of treatment with oral cyclophosphamide was unfortunately very heterogeneous and variable depending on the groups. Methotrexate appeared to be more effective than cotrimoxazole in patients treated with or without corticosteroids (91% vs. 0% and 86% vs. 58%, respectively). However, the limitations of this open-label study (small groups, short duration) reduce its scope [99].

**CONCLUSION**

Wegener is a rare granulomatosis in Africa. It is characterized by a clinical polymorphism. Its diagnosis is very difficult, leading to complications that can affect the functional and vital prognosis.

Management is multidisciplinary with early corticosteroid therapy and the use of other immunosuppressive therapy.

**REFERENCES**


92. Tervaert JW, van der Woude FJ, Fauci AS, Ambrus JL, Velosa J, Keane WF, Meijer S, van der Giessen...


