A case of rheumatic fever with acute post-streptococcal glomerulonephritis and nephrotic syndrome caused by a cutaneous infection with beta-haemolytic streptococci

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Case Report

A 44-year-old woman of Greenlandic origin was referred to Queen Ingrid’s Hospital. She had been healthy apart from excessive alcohol consumption and pulmonary tuberculosis diagnosed two years earlier for which she had received full antitubercular treatment. The patient initially complained of a painful traumatic skin lesion of the right lower extremity with a diameter of approximately 2 cm. A few days later, her temperature rose to 39.5°C and bullous erysipelas of the area of initial trauma was noted. Laboratory tests revealed a markedly elevated sedimentation rate of 111 mm/hour [2-21], an elevated C reactive protein of 144 mg/L [150-450]. Neutrophilia and leucocytosis were noted. Culture from the lesion of the right lower extremity was positive for group A beta-haemolytic streptococci (GABHS). In spite of relevant antibiotic treatment, the clinical condition rapidly deteriorated with involvement of the deeper layers of the skin and subcutaneous tissue, spreading across the fascial planes of the subcutis (Figure 1). Necrotising fasciitis (NF) was diagnosed. Based on positive skin cultures, this condition was considered to be due to GABHS. The patient had no other predisposing factors to NF, specifically no diabetes mellitus, no intravenous abuse and no daily intake of non-steroidal anti-inflammatory drugs (NSAID). Radiological examination of the right lower extremity and foot showed no signs of osseous involvement. Extensive surgical revision of the NF-lesion was performed and meropenem, ciprofloxacin and clindamycin were administered. Approximately three weeks after the initial presentation, impairment of renal function with azotemia and severe nephrotic syndrome with periorbital and peripheral oedema were noted.

Laboratory tests showed an elevated creatinine of 386 μmol/L [44-115], BUN: 23.6 mmol/L [2.5-6.7], normochromic, normocytic anaemia with hemoglobin: 4.9 mmol/L [7.5-9.0], as well as macroscopic hematuria, renal-creatinine clearance: 30 ml/min and U-protein: 5.8 g/24 h [<0.15]. Anti nuclear antibodies (ANA), anti-neutrophil cytoplasmic antibodies (ANCA), IgA, IgG, IgM and protein electrophoresis were all normal. Renal biopsy was performed. The biopsy demonstrated endocapillary glomerulonephritis with proliferation of endothelial cells and infiltration of polymorphonuclear cells. Using immunofluorescence techniques, granular deposits of IgG and C3 were identified as typically seen in acute post-streptococcal glomerulonephritis (APSGN) (Figure 2). During the same period, the patient complained of chest pain. Electrocardiogram (ECG) showed sinus rhythm with no signs of ischemia and no elevation of cardiac troponin I was found. Echocardiography showed moderate reduced left ventricular ejection fraction (LVEF): 35% [50-65%] and moderate mitral regurgitation (grade II). Pulmonary pressure was significantly increased >60 mm Hg [18-25]. Brain natriuretic peptide (BNP) was significantly increased >8000 pg/mL [<100]. There were complaints of arthralgia of the left elbow. There were no other signs of acute rheumatic fever (ARF) and no signs of pharyngitis in the whole period. Chest X ray and HR-CT of the thorax showed changes in both lungs consistent with old tuberculosis. Two weeks after the first analysis, renal function had improved and U-protein had decreased to 2.6 g/24 h, creatinine 139 μmol/L, BUN 9.5 mmol/L. Ultrasound of the kidneys was normal. Echocardiography still...
showed moderate reduced left ventricular ejection fraction (LVEF): 35% with mild mitral regurgitation (grade 1). Continued follow-up of renal and cardiac function is planned.

**Discussion**

This case illustrates the simultaneous development of carditis, arthralgia and endocapillary glomerulonephritis with severe nephrotic syndrome preceded by a cutaneous GABHS in the lower right leg. The patient fulfilled the revised and updated Jones criteria for classification of ARF based on fever, arthralgia, carditis, increased acute phase reactants (CRP, ESR) and a positive culture of GABHS. ARF develops after group A streptococcal pharyngitis and is only seldom described after GABHS skin infections. Our patient had no signs of pharyngitis but developed NF presumably from a traumatic skin lesion. Culture from the initial lesion of the right lower extremity was positive for GABHS. We propose that our patient developed ARF from her skin infection. We are aware of no history of sore throat described in the literature in cases of ARF. Also throat swab will be positive in only a third of patients infected with group A streptococcus before antibiotic treatment and in only a tenth of patients afterwards. GAS pyoderma rather than pharyngitis as a driving force behind ARF has been described in aboriginal communities in Northern Australia. Repeated exposure to GABHS can play a central role in the development of ARF as a sort of immune priming. In our case there was no previous history of pharyngitis or other focus of GABHS infections. ARF and APSGN, two important sequelae of streptococcal throat or skin infections, according to current concepts may be elicited by autoimmune mechanisms due to molecular mimicry between GABHS and human tissue. APSGN follows infection with a limited number of GABHS serotypes. Type 12 is the most frequent M serotype causing APSGN after pharyngitis or tonsillitis, whereas M-49 is the type most frequently related to pyoderma-associated nephritis. The patient had no earlier symptoms of heart disease and so we strongly suspect the high BNP and and moderate mitral regurgitation to be carditis related to ARF. In affluent societies where GABHS disease is uncommon apart from pharyngitis in childhood, increasing numbers of NF and streptococcal toxic shock syndrome (STSS) have been seen, as well as an acute upsurge of rheumatic fever apparently restricted to parts of the United States. In some of these locations, a virulent M1 serotype GABHS clone has been found. We suggest a possible relation between a virulent GABHS clone causing NF and ARF.

**References**