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COPA Syndrome as a Cause of Lupus Nephritis



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INTRODUCTION

Genetic kidney diseases are rare situations resulting from mutations in genes expressed in major structures of the kidney, including podocytes, tubular cells, and basement membrane.¹ More recently, a new field of monogenic inflammatory diseases has been identified²; these immune-mediated diseases can affect all organs, including the kidney. Here we report on a young girl diagnosed with lupus nephritis and carrying a mutation in the *COPA* gene encoding for coatamer protein complex subunit alpha.

CASE PRESENTATION

A 10-year-old girl was referred because of fever, facial edema, and fatigue. Upon admission, her blood pressure was 117/70 mm Hg and her temperature was 39 °C. Clinical examination revealed facial edema. Her medical history was unremarkable. Two other second-degree relatives were suspected of having systemic vasculitis because of lung hemorrhage and antineutrophil cytoplasmic antibody positivity. Initial laboratory testing performed at the emergency department revealed acute renal failure with increased serum creatinine at 106 µmol/L, a urinary

protein-to-creatinine ratio of 1.76 g/g, and microscopic hematuria with $>1 \times 10^6$ red blood cells/ml. C-reactive protein was 3 mg/l, and the erythrocyte sedimentation rate was 52 mm. The hemoglobin level was 9 g/dl. Tests for antinuclear antibodies, antineutrophil cytoplasmic antibody, and antiglomerular basement membrane were negative, and C3 and C4 complement levels were 1.23 g/l and 0.39 g/l, respectively. Infectious screening was normal, and a kidney biopsy was performed. Light microscopy and immunofluorescence studies were consistent with lupus nephritis (Figure 1). Treatment of putative lupus nephritis was started with a steroid pulse, i.v. cyclophosphamide, and hydroxychloroquine. Proteinuria persisted in the following months and angiotensin-converting enzyme inhibitors were introduced, but the disease was progressive.

Transient arthralgia of the knee was later reported without joint lesions or synovitis. Two additional biopsies were performed and showed the same immunopathologic pattern with progression of chronic lesions. Mycophenolate mofetil was introduced but without significant impact on the chronic course of the disease and she developed end-stage renal failure. While living donor transplantation was

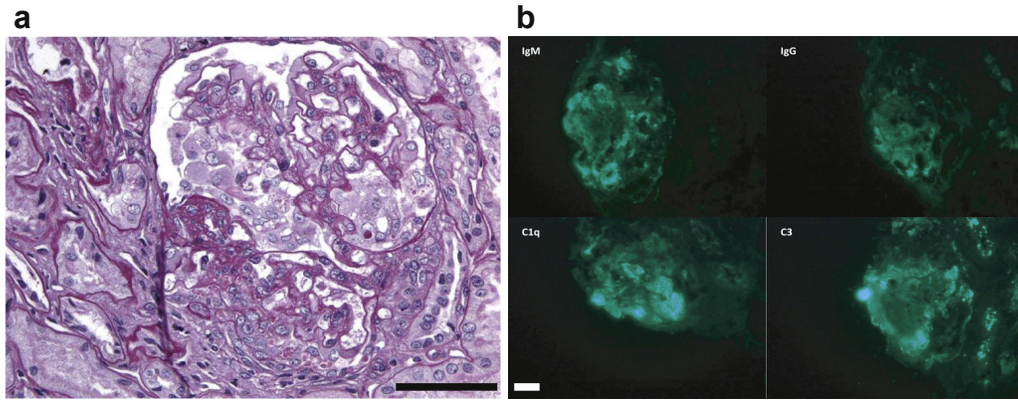


Figure 1. Kidney biopsy specimen of patient 1 (case index). (a) Proliferative lesions associated with focal and segmental glomerulosclerosis. (b) Immunofluorescence study revealed endomembranous deposits of C1q, C3, IgM, and IgG (bar = 50 μm).

being considered, exploration in the father, age 50 years, revealed chronic proteinuria. A kidney biopsy revealed membranous glomerulonephritis. Meanwhile, the 2 second-degree relatives were diagnosed with COPA syndrome, a monogenic disease presenting

as antineutrophil cytoplasmic antibody-associated vasculitis. The patient and her father then underwent screening for COPA mutation, and they both carried the familial mutation. Diagnosis of COPA syndrome was made.

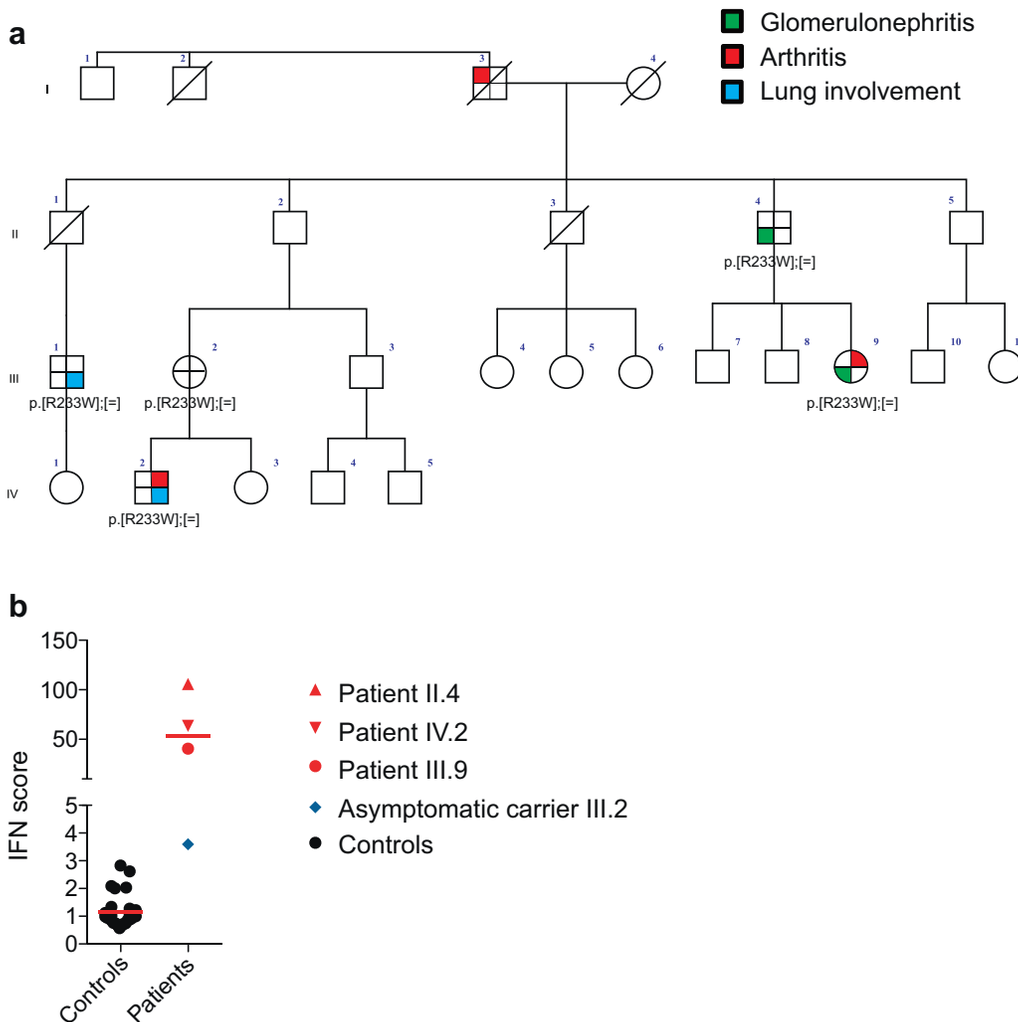


Figure 2. Genetic and immunologic description. (a) Family tree of the index case (III.9) with mutation segregation and clinical features. (b) Type I interferon (IFN) assessment in whole blood of members of the family. The IFN signature was increased in all symptomatic patients carrying the familial mutation.

Table 1. Teaching points

COPA syndrome is an inherited disease presenting with systemic autoimmunity from antineutrophil cytoplasmic antibody vasculitis to lupus phenotype
Expressivity and penetrance are variable in a single family
Type I interferon is increased in symptomatic patients and may play a role in the pathogenesis of the disease

COPA, coatamer protein.

DISCUSSION

COPA syndrome represents a newly recognized autosomal-dominant cause of autoinflammatory/auto-immune disease characterized by variable expression and a high frequency of clinical nonpenetrance. Arthritis, interstitial lung disease, and glomerulonephritis represent the main characteristic features.³ The COPA syndrome occurs as a result of a heterozygous mutation in the *COPA* gene encoding for the alpha subunit of the COPI coatamer protein complex. The COPI protein complex is involved in the trafficking of membranes containing proteins and lipids from the Golgi to the endoplasmic reticulum.⁴ To date, the link between intracellular trafficking and immune dysregulation has not been characterized, but endoplasmic reticulum stress and the activation of the unfolded protein response have been evoked.⁴ Here, the family history was informative (Figure 2a). Despite the absence of symptoms in the siblings of the father, a phenotypic link between the cousins, the father, and index case was suspected, and genetic screening for the known familial *COPA* mutation confirmed that both the father and the index case were heterozygous for the same amino acid substitution (p.Arg233His), a mutation already described as causal in 2 families.³ The family tree in this pedigree illustrates the variable expression and incomplete penetrance of COPA syndrome, suggesting that additional environmental or genetic modifiers might influence the phenotype. To date, 31 cases of COPA syndrome have been reported, 14 of which presented with renal lesions; the main histologic features are crescentic glomerulonephritis and focal mesangial hypercellularity with immune complex deposits, ranging from isolated IgA deposits to “full-house” immunofluorescence (IgM, IgG, and C1q deposits).

The patient benefited from a live-donor transplantation from her mother, and renal function was satisfactory 3 months after transplantation. Remarkably, patient IV.2 displayed a positive response to

rituximab, suggesting a possible contribution of B cells to some aspect of the disease.

Underlying pathogenesis of COPA syndrome is still unknown, but a positive type I interferon signature has been identified in some patients with COPA syndrome.⁵ We also explored type I interferon signaling in the peripheral blood of different members of the family and identified a positive transcriptomic signature in all symptomatic individuals tested (Figure 2b). Of note, type I interferon is a key cytokine in systemic lupus erythematosus. Taken together, a history of arthritis, interstitial lung disease, and immune-mediated glomerulonephritis with dominant inheritance is evocative of COPA syndrome (Table 1). Type I interferon assessment might represent an additional biologic marker of this syndrome and an explanatory factor for the overlapping features of COPA syndrome with both lupus renal immunopathology and the phenotype associated with gain-of-function of the gene-encoding Stimulator of Interferon Genes (*STING*).⁶ Thus Janus kinase inhibitors or type I interferon inhibition may represent promising therapies in this context.

DISCLOSURE

All the authors declared no competing interests.

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