Fibrillary glomerulonephritis with a favourable prognosis of 26 years

David Micarelli1*, Valentina Pistolesi2, Emanuela Cristi3, Anna Rita Taddei4, Ilaria Serriello1, Santo Morabito3, Konstantinos Giannakakis5

1Nephrology and Dialysis Unit, Belcolle Hospital, Viterbo, Italy
2UO Dialisi, Azienda Ospedaliero-Università Policlinico Umberto I, “Sapienza” Università di Roma, Viale del Policlinico, Rome, Italy
3Division of Surgical Pathology, Belcolle Hospital, Viterbo, Italy
4Department of Radiology, Oncology, Radiology and Pathology, Sapienza University of Rome, Rome, Italy

ARTICLE INFO

Article type:
Case Report

Article history:
Received: 27 July 2019
Accepted: 10 November 2019
Published online: 15 May 2020

Keywords:
Fibrillary glomerulonephritis
Renal outcome
Electron microscopy
End-stage renal disease

ABSTRACT

Fibrillary glomerulonephritis (FGN) is a rare glomerular disease. The prognosis is usually unfavorable with nearly half of patients progressing to end-stage renal disease within 4 years. We report a case of biopsy-proven FGN characterized by an unusual benign clinical course in which a kidney biopsy, repeated after an extended follow-up of 26 years, confirmed the presence of fibrils deposition. In 1993, a 32-year-old Caucasian man was admitted to our nephrology ward because of macroscopic hematuria. Renal function was normal. Kidney biopsy displayed an FGN with mesangial pattern. The patient was treated with lisinopril, titrated for blood pressure; the therapy was maintained during 26 years of follow-up. The yearly slope of estimated glomerular filtration rate was -3.17 mL/min. Starting from March 2018, a rapid worsening of renal function was observed and proteinuria increased up to a nephrotic range. We planned a second renal biopsy to assess the cause of the rapid change of clinical course. The diagnosis of FGN on advanced sclerosis was made, and the severity of glomerular sclerosis. We report a case of FGN with an unusually benign clinical course, characterized by a slow progression to end-stage renal disease over a very extended follow-up time; thus, to better clarify the reason for renal function worsening, a second renal biopsy was performed. The persistence of fibrils deposition confirmed the initial diagnosis of FGN, and a histological pattern characterized by global glomerular sclerosis and interstitial fibrosis has been observed.

Implication for health policy/practice/research/medical education:
Fibrillary glomerulonephritis (FGN) is an uncommon disease, since diagnosis in the past required electron microscopy and the prognosis of the disease was really unfavorable. Here, we described an unusual case of FGN with 26 years of prognosis without progression to end-stage renal disease and not required dialysis.

Please cite this paper as: Micarelli D, Pistolesi V, Cristi E, Taddei AR, Serriello I, Morabito S, Giannakakis K. Fibrillary glomerulonephritis with a favourable prognosis of 26 years. J Nephropathol. 2021;10(4):e44. DOI: 10.34172/jnp.2021.44.

Introduction

Fibrillary glomerulonephritis (FGN) is a rare glomerular disease (less than 1% of glomerulonephritis) (1-4) characterized by the deposition of randomly distributed non-amyloid fibrils in the mesangium and capillary basement membrane (5,6). The prognosis is usually unfavorable with nearly half of patients progressing to end-stage renal disease (ESRD) within 4 years (1-4). The pathogenesis is still unclear, while studies highlighted a possible association with autoimmune disease, malignancy, or hepatitis C infection (4,7,8). Recently, a potential role of DNA homolog subfamily B member 9 (DNAJB9) in causing FGN has been reported (5,9,10). The glomerular deposits usually stain for polyclonal IgG, mostly subtypes IgG1 and IgG4 (3,11). In this disease, immunoelectron studies have shown the co-localization of polyclonal IgG to the FGN fibrils (2,12,13). The clinical feature is usually a therapy-resistant nephrotic syndrome,

*Corresponding author: David Micarelli, Email: davidmicarelli@gmail.com
hematuria, arterial hypertension and a progressive renal impairment, which is present in 50% of all cases of FGN at the time of diagnosis (3,4,8,14).

Here, we report a patient with biopsy-proven FGN characterized by an unusual benign clinical course in which a kidney biopsy, repeated after an extended follow-up of 26 years, confirmed the presence of fibril deposition.

**Case Presentation**

In 1993, a 32-year-old Caucasian man was admitted to our nephrology ward because of macroscopic hematuria. Main laboratory tests are reported in Table 1. Serum creatinine (Scr) was 0.9 mg/dL with an estimated glomerular filtration rate (eGFR) of 112 mL/min/1.73 m². Urinalysis showed 20-40 RBC and 1-2 granular casts in high-power field (HPF) with 24-hour urinary protein excretion of 0.6 g. Serum electrophoresis revealed no abnormalities, and Bence-Jones protein was not detected in the urine sample. Serum C3, C4, and total Ig levels were normal. Additionally, ANA, ANCA, extractable nuclear antigen, and cryoglobulins were absent. Serology markers for hepatitis B and C were negative. Clinical history, as well as the physical examination, was negative. A percutaneous renal biopsy was carried out. Light microscopy showed 10 glomeruli with diffuse mesangial matrix expansion and thickened capillary walls. The arteries displayed mild intimal fibrosis, while arterioles, tubules, and interstitium were normal. Immunofluorescence investigation on frozen sections (5 glomeruli) showed granular mesangial deposition of IgG (2+), C3 (1+), k and l light chains (++). Congo red staining was negative. The staining for IgG4 was positive. Ultrastructural investigation revealed randomly arranged non-branching fibrils of approximately 15 nm in diameter in the mesangium and sub-endothelial space (Figure 1A, 1B) with diffuse foot process effacement (Table 1).

Tangri kidney failure risk equation (16) resulted at 0.2% of risk to progression to ESRD over 5 years.

The patient was treated with lisinopril, titrated for blood pressure. The therapy was maintained during 26 years of follow-up. The yearly slope of eGFR was -3.17 mL/min. Starting from March 2018, a rapid worsening of renal function was observed (Scr 2.1 mg/dL, eGFR 35 mL/min/1.73 m²) (Figure 2) and proteinuria increased up to a nephrotic range (9.8 g/24 h) despite an optimal blood pressure control (<130/80 mm Hg). Immunological tests were still negative, and serum complement profile was average. No paraproteins were detected in serum and urine electrophoresis. Renal ultrasound showed normal-sized kidneys. We planned a second renal biopsy to assess the cause of the rapid change of clinical course (i.e., superimposed glomerulonephritis versus FGN progression) and to evaluate the indication for emergent treatment strategies (e.g., rituximab) (7,15). Light microscopy displayed 15 glomeruli; 12 globally sclerosed, 3 with diffuse mesangial matrix increase associated in 2 of them with global fibrotic crescents. The arteries displayed narrowing of the lumens with wall fibrosis. Some hyaline casts were detected inside the tubules. Polymorphic infiltrates and area of fibrosis were present in the interstitial space. Immunofluorescence examination on paraffin showed segmental, granular deposition of IgG (1+), IgM (1+), fibrinogen (1+) along the capillary walls. Congo red staining was negative. The staining for IgG4 was positive. Ultrastructural investigation revealed randomly arranged non-branching fibrils of approximately 15 nm in diameter in the mesangium (Figure 2A, B, C, D; Table 1).

Tangri kidney failure risk equation (16) resulted in 77.6% in risk to progression to ESRD over two years.

The diagnosis of FGN on advanced sclerosis was made, and the severity of glomerular sclerosis discouraged us to treat the patient with rituximab.

**Discussion**

We report a case of FGN with an unusually benign clinical course, characterized by a slow progression to ESRD over a very extended follow-up time. Indeed, in our patient an advanced chronic kidney disease was evident after 26 years from initial diagnosis; thus, to better clarify the reason for renal function worsening, a second renal biopsy was performed. The persistence of fibril deposition confirmed the initial diagnosis of FGN, and a histological pattern characterized by global glomerular sclerosis and interstitial fibrosis has been observed.

As a rule, FGN has a relentless progression towards ESRD, since 40-50% of patients require dialysis after an average of two years from the diagnosis (2-4,15). In a case series from Columbia University, including 61 patients with FGN, the median time to ESRD was 24.4±15.2
Progressive damage secondary to the deposition of fibrils was generally considered due to the release of profibrotic cytokines as transforming growth factor-beta associated with activation of the NOTCH1 pathway [S8, S9]. Recently, Nasr et al proposed a pathogenic role for DNAJB9, a heat-shock protein that acts as co-chaperone and co-localizes with IgG and components of the complement pathway (5). In particular, DNAJB9 may be pivotal in recognition of misfolded proteins and activate the unfolded protein reaction determining endothelial reticulum stress and inflammation (5). Andeen et al, pointing out the absence of demonstration of other unfolded protein response beside DNAJB9, proposed an alternative hypothesis. The co-localization of DNAJB9 as antigen with IgG and complement fractions suggests autoimmune pathogenesis (10). Thus, questions arise about the therapeutic strategy to adopt in patients with FGN, first of all, the potential role of immunosuppressive treatment.

Our patient has been treated with an angiotensin-converting enzyme inhibitor alone for the entire follow-up, and the presence of positive prognostic factors could explain the long dialysis-free period at the time of diagnosis (i.e., normal renal function, low-grade proteinuria and mesangial pattern). For the absence of definitive evidence on its clinical effectiveness, immunosuppressive therapy has not been taken into consideration after the first renal biopsy.

Different experiences regarding immunosuppressive use in FGN have been later reported. In the above-described cohort, 36% (3) and 48% of patients (4) received immunosuppressive therapy (steroids, rituximab or cyclophosphamide) in various combinations without evidence of a slower progression to ESRD. A subsequent study, including 27 patients with FGN in conservative

| Laboratory data                      | 1993   | 2019   |
|--------------------------------------|--------|--------|
| Serum creatinine (mg/dL)             | 0.9    | 3.32   |
| Urinary protein excretion (g/24 h)   | 0.6    | 10.2   |
| eGFR (mL/min/1.73 m²)                | 112    | 19     |
| BUN (mg/dL)                          | 19.2   | 46.7   |
| Total serum proteins (g/dL)          | 7.3    | 4.8    |
| Serum albumin (g/dL)                 | 3.12   | 2.6    |
| Hemoglobin (g/dL)                    | 14.3   | 11.3   |
| RBC (U/µL)                           | 4.950,000 | 3,740,000 |
| WBC (U/µL)                           | 6.700  | 5,830  |
| PLTs (U/µL)                          | 194,000 | 226,000 |
| Urinary sediment                     | RBC 40 hpf | RBC 10-20 hpf |
| Biopsy findings                      |        |        |
| Glomeruli #                          | 10     | 15     |
| Crescents #                          | None   | 2 (fibrotic) |
| Sclerosis and/or hyalinosis #        | None   | 12     |
| Mesangial matrix increase#           | 10     | 3      |
| Tubular atrophy/interstitial fibrosis| None   | Diffuse |
| Congo red stain                      | neg    | neg    |
| Immunofluorescence                   | IgG (2+), C3 (1+) | IgG, IgM, C1q (0/1+) |
| Deposit type (EM)                    | Fibrils 15 nm | Fibrils 15 nm |
| Deposit location (EM)                | M, BM  | M, BM  |

...
treatment with renin-angiotensin system (RAS) inhibitors, showed that the addition of immunosuppressive therapy with different protocols in 13 out of 27 patients (rituximab-based therapy in 7 cases), resulted in partial remission (>50% decrease in 24-hour proteinuria with <15% decrease in eGFR compared to the baseline value) in about 46% of cases (7). The authors highlighted that those responders were characterized by a higher eGFR at the start of immunosuppressive treatment if compared to non-responders (76 versus 42 mL/min/1.73 m²) (7). More recently, in the more extensive published case series of FGN patients treated with rituximab (n=12), 4 patients, characterized by a median basal eGFR of 71 mL/min/1.73 m², showed a stable or improved sCr with a minimum of 1-year follow-up (non-progressors), while the remaining 8 patients (median basal eGFR 28 mL/min/1.73 m²) showed a progressive renal disease (ESRD in 5 patients after a median follow-up of 17 months) (15). Thus, the therapeutic strategy in FGN remains poorly defined; although it appears reasonable to adopt RAS inhibitors treatment in all patients. The start of an immunosuppressive regimen remains debated and at the same time worthy of further evaluation in well selected and appropriately extended case series.

Conclusion
In conclusion, we could speculate that the benign clinical course observed in our patient, characterized by the maintenance of renal function for more than 25 years, could be explained by a subtype of FGN disease with favourable histologic and clinical prognostic factors along with the protective effects of RAS block (17,18).

Authors’ contribution
DM wrote the paper and revisited the case. KG and EM, as the pathologists, read and reported the first and the second biopsies, respectively. ART was the electron microscopist who read TEM; VP and SM were the physicians of the patient from 1993 to 2018. IS performed the second kidney biopsy All authors read and signed the final paper.

Conflicts of interest
The authors declare no competing interests.

Ethical considerations
Ethical issues including plagiarism, double publication, and redundancy have been completely observed by the authors. The patients gave his consent to publish as a case report.

Funding/Support
The authors declared no funding or support from any institution.

References
1. Iskandar SS, Falk RJ, Jennette JC. Clinical and Pathologic Features of Fibrillary Glomerulonephritis. Kidney Int. 1992;42:1401-7. doi: 10.1038/ki.1992.433.
2. Fogo A, Qureshi N, Horn RG. Morphologic and clinical features of fibrillary glomerulonephritis versus immunotactoid glomerulopathy. Am J Kidney Dis. 1993;22:367-77. doi: 10.1016/0272-6386(93)70138-5.
3. Rosenstock JL, Markowitz GS, Valeri AM, Sacchi G, Appel GB, D’Agati VD. Fibrillary and immunotactoid glomerulonephritis: distinct entities with different clinical and pathologic features. Kidney Int. 2003;63:1450-61. doi: 10.1046/j.1523-1755.2003.00853.x.
4. Nasr SH, Valeri AM, Cornell LD, Fidler ME, Sethi S, Leung N, et al. Fibrillary glomerulonephritis: a report of 66 cases from a single institution. Clin J Am Soc Nephrol 2011;6:775-84. doi: 10.2215/CJN.08300910.
5. Nasr SH, Fogo AB. New developments in the diagnosis of fibrillary glomerulonephritis. Kidney Int. 2019 Apr 9. pii: S0085-2538(19)30407-7. doi: 10.1016/j.kint.2019.03.021.
6. Rosenstock JL, Markowitz GS. Fibrillary glomerulonephritis: an update. Kidney Int Rep. 2019;4(7):917-22. doi: 10.1016/j.ekir.2019.04.013.
7. Javaugue V, Karras A, Glowacki F, McGregor B, Lacombe C, Goujon JM et al. Long-term kidney disease outcomes in fibrillary glomerulonephritis: a case series of 27 patients. Am J Kidney Dis. 2013;62:679-90. doi: 10.1053/j.ajkd.2013.03.031.
8. Payan Schober F, Jobson MA, Poulton CJ, Singh HK, Nickelet V, Falk RJ, et al. Clinical features and outcomes of a racially diverse population with fibrillary glomerulonephritis. Am J Nephrol 2017;45:248-56. doi: 10.1159/000455390.
9. Dasari S, Alexander MP, Vrana JA, Theis JD, Mills JR, Negron V, et al. DnaJ heat shock protein family B member 9 is a novel biomarker for fibrillary GN. J Am Soc Nephrol. 2018;29(1):51-6. doi: 10.1681/ASN.2017030306.
10. Andeen NK, Yang HY, Dai DF, MacCoss MJ, Smith KD. DnaJ Homolog Subfamily B Member 9 Is a Putative Autoantigen in Fibrillary GN. J Am Soc Nephrol. 2018;29(1):231-239. doi: 10.1681/ASN.2017050566.
11. Hemminger J, Nadasy G, Satoaskar A, Brodsky SV, Nadasy T. IgG Subclass staining in routine renal biopsy material. The Am J Surg Pathol. 2016;40(5):617-26. doi: 10.1097/PAS.0000000000000605.
12. Casanova S, Donini U, Zucchielli P, Massuzco G, Monga G, Linke RP. Immunohistochemical distinction between amyloidosis and fibrillar glomerulopathy. Am J Clin Pathol. 1992;97:787-95. doi: 10.1093/ajcp/97.6.787.
13. Yang GC, Nieto R, Stachura I, Gallo GR. Ultrastructural immunohistochemical localization of polyclonal IgG, C3, and amyloid P component on the congo red-negative amyloid-like fibrils of fibrillary glomerulopathy. Am J Pathol 1992;141:409-19.
14. Nasr SH, Vrana JA, Dasari S, Bridoux F, Fidler ME, Kaaki S, et al. DNAJB9 Is a Specific Immunohistochemical Marker for Fibrillary Glomerulonephritis. Kidney Int Rep. 2017;3(1):56-64.
15. Hogan J, Restivo M, Canetta PA, Herlitz LC, Radhakrishnan J, Appel GB, et al. Rituximab treatment for fibrillary glomerulonephritis. Nephrol Dial Transplant.
Fibrillary glomerulonephritis

2014;29(10):1925-31. doi: 10.1016/j.ekir.2017.07.017.

Tangri N, Stevens LA, Griffith J, Tighiouart H, Djurdjev O, Naimark D, et al. A predictive model for progression of chronic kidney disease to kidney failure. JAMA. 2011;305(15):1553-9. doi: 10.1001/jama.2011.451.

Ruggenenti P, Remuzzi G. Angiotensin-converting enzyme inhibitor therapy for non-diabetic progressive renal disease.

Curr Opin Nephrol Hypertens. 1997;6(5):489-95. doi: 10.1097/00041552-199709000-00014.

Sun N, Zhai L, Li H, Shi LH, Yao Z, Zhang B. Angiotensin-converting enzyme inhibitor (ACEI)-mediated amelioration in renal fibrosis involves suppression of mast cell degranulation. Kidney Blood Press Res. 2016;41(1):1. doi: 10.1159/000368549.

Copyright © 2021 The Author(s); Published by Society of Diabetic Nephropathy Prevention. This is an open-access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/4.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.