Idiopathic Membranous Nephropathy: Diagnostic and Therapeutic Challenges

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Introduction

The association between human immunodeficiency virus (HIV) and renal glomerular disease has been well recognized [1–9]. Significant focus in literature has been placed on HIV-associated nephropathy (HIVAN). HIVAN has been reported as the most common cause of kidney disease in patients with HIV, and it must be considered in all HIV positive patients presenting with proteinuria [1–4]. However, biopsy studies of patients with HIV and kidney disease have found substantial prevalence of other glomerular diseases, including membranous nephropathy (MN) [1, 5–9]. MN is a glomerular pathology characterized microscopically by subepithelial immune complex deposits and glomerular basement membrane (GBM) thickening without hypercellularity [10]. The idiopathic subtype of MN has been closely associated with positive tests for anti-phospholipase A2 receptor (PLA2R) antibodies [11–14]. To our knowledge, there are no reports in the literature of patients with well-controlled HIV who developed MN without an underlying etiology for secondary MN and tested positive for anti-PLA2R antibody to indicate idiopathic disease. In this study, we present a case of an HIV-positive patient with anti-PLA2R antibody-associated MN effectively treated with ACTH immunosuppression.
Case Report

A 36-year-old, asymptomatic, Caucasian male with a history of hemophilia A, HIV secondary to blood transfusion in childhood and cleared hepatitis C infection was incidentally found on urinalysis to have proteinuria. The patient had normal serum creatinine of 0.74 mg/dl, serum albumin of 2.4 g/dl and nephrotic range proteinuria of approximately 7 g/24 h without peripheral edema on exam. Urinalysis showed 6 RBC/hpf without cellular casts. Repeated random urine protein/creatinine (UP/C) ratio was ranging between 6 and 8 g/g and he had normal complement, and negative anti-nuclear, anti-double stranded DNA, anti-neutrophil cytoplasmic, anti-myeloperoxidase, anti-proteinase 3 antibodies. Tests for hepatitis C revealed the presence of positive antibodies to hepatitis C with negative hepatitis C RNA by PCR. HIV viral load was also undetectable with CD4 count of 1,234/μl. He had been treated with abacavir/lamivudine/zidovudine (Trizivir) for more than 10 years with excellent long-term control. During the workup of his proteinuria, highly active antiretroviral therapy (HAART) regimen was switched to efavirenz/emtricitabine/tenofovir (Atripla) for simplification and then to abacavir/lamivudine/efavirenz to avoid insult to kidneys. Quantitative free kappa and lambda light chains were within normal limits in the normal ratio. Serum protein electrophoresis with immunofixation showed no monoclonal proteins. Kidney ultrasound revealed no abnormalities.

The patient was started on lisinopril and maximized up to 20 mg twice daily as tolerated by the blood pressure, and proteinuria was consistently above 6 g daily for 6 months. The patient was admitted to the hospital for trans-jugular kidney biopsy with continuous infusion of factor VIII and monitoring of factor VIII activity levels as he had a baseline activity level of 4%. The biopsy was performed without complications.

Biopsy results revealed global glomerulosclerosis involving 2/42 glomeruli (4.7%). There were normal-sized glomeruli with normal to slightly increased cellularity and no glomerular proliferative lesions, cellular crescents, GBM double contours, or breaks on light microscopy (fig. 1a). Silver stain highlighted GBM spikes (fig. 1b). There was mild tubular atrophy and minimal focal lymphocytic inflammatory infiltrate. Trichrome stain highlighted mild interstitial fibrosis with moderate arterial fibro-intimal thickening. Immunofluorescence microscopy demonstrated 2+ granular staining of the GBM for IgG, kappa and lambda light chain, and trace granular staining for C3. Linear nonspecific staining in the GBM for albumin was also seen (fig. 1c). Electron microscopy revealed GBM thickening, diffuse effacement of the foot processes, and numerous subepithelial electron dense deposits of complex immune type, partially surrounded by new GBM matrix formation (fig. 1d). These findings are consistent with stage II MN.

Workup was initiated for causes of secondary MN. CT of chest, abdomen, and pelvis was unremarkable. Colonoscopy revealed no masses or polyps. Prostate-specific antigen and creatinine kinase levels were within normal limits. Serologic tests for hepatitis B, chlamydia, gonorrhea, and syphilis were negative. The patient tested weakly positive for anti-PLA2R antibodies. Because of negative hepatitis C virus (HCV)-RNA, undetectable HIV viral load and absence of any evidence of secondary MN in renal biopsy, we considered his MN as likely to be idiopathic.

Therapy was initiated with twice weekly 80 U injections of adrenocorticotropic hormone (Acthar) gel. UP/C was 6.0 g/g when therapy was initiated. After the therapy was completed, UP/C was down to 4.0 g/g after 2 months, 0.9 g/g at 8 months, 0.5 g/g at 12 months and 0.4 g/g at 18 months of follow-up. Acthar gel was decreased to once weekly injections after 8 months and then stopped after 16 months of starting therapy (fig. 2). The patient was maintained on lisinopril throughout this time. The patient has remained asymptomatic with regard to kidney disease and additionally, his CD4 count was around 2,000/μl; also, he had undetectable HIV and HCV viral load for 18 months of follow-up. Repeated western blot after completion of the Acthar gel therapy showed significantly decreased PLA2R antibody (fig. 3) and negative anti-thrombomodulin type-1 domain-containing 7A (THSD7A).

Detection of Anti-PLA2R and THSD7A Antibodies by Western Blotting

Human glomerular extract (HGE) and extracts of HKE293 cells transfected with either recombinant human PLA2R or human THSD7A were electrophoresed on SDS-PAGE 4–15% gradient gels, transferred to nitrocellulose membrane, and western blotted overnight with human serum (at a 1:10 dilution for this case, or 1:25 dilution for a positive control serum with a high titer of anti-PLA2R antibodies). The blot was then incubated with sheep anti-human IgG antibodies (1:3,000; The Binding Site) followed by peroxidase-conjugated donkey anti-sheep IgG (1:10,000; Jackson ImmunoResearch). After exposure to a chemiluminescent substrate, the blots were exposed to film for 10, 30 and 120 s and developed. Blots were scanned on a CanonScan Lide 25 scanner and the image assembled using Photoshop CS4 for Windows.

Discussion

MN is a common form of glomerular disease presenting with proteinuria. A majority of patients develop nephrotic syndrome and a significant number go on to develop the end-stage renal disease [15]. MN can be subdivided into idiopathic, without an identifiable etiology, and secondary types with idiopathic being more common [10]. Secondary MN has been associated with chronic infections, drugs, autoimmune disease, and malignancies, and therapy focuses on treating or removing the underlying cause [16–18]. In contrast, treatment of idiopathic MN is more poorly defined. Polanco et al. [19] reported that 31.7% of patients with idiopathic MN will see spontaneous remission of proteinuria when treated conservatively with a majority of spontaneous remission occurring within the first 2 years. In this study, spontaneous remission was more common among patients with lower levels of baseline proteinuria, and a gradual decrease to <50% of baseline proteinuria within the first 12 months predictive of spontaneous remission. Patients with severe proteinuria, persistent proteinuria, or reduced kidney function are typically treated with immunosuppressive therapy and this therapy typically involves immunosuppression...
Therefore, it is important in the workup of a patient with MN to elucidate the underlying diagnosis as idiopathic or secondary to guide treatment [16].

The case presented here provides many diagnostic nuances. Although it has been well documented that numerous glomerular diseases other than HIVAN, including MN, exist in patients with HIV, these associations have not been very well studied. Most cases of HIV-associated MN in literature have been reported generally without being specified as idiopathic or secondary [1, 5–9]. Due to common risk factors, patients with HIV are commonly coinfected with hepatitis B, hepatitis C, and syphilis, which are associated with secondary MN [17, 18]. Furthermore, patients with glomerular disease other than HIVAN are more likely to be coinfected with hepatitis B or hepatitis C [1].

Multiple reports of HIV-positive patients with MN demonstrated that they responded well to treatment of an underlying etiology consistent with secondary disease. Aydin et al. [21] reported a case of MN in a patient with recently diagnosed HIV without coinfections whose proteinuria decreased following treatment with ACE and HAART, indicating that MN may have been secondary to HIV. Alarcón-Zurita et al. [22] reported a similar case of MN in a patient positive for HIV and hepatitis C whose kidney disease responded to HAART. Chen et al. [23] reported a case of MN in a patient with HIV who was diagnosed with syphilis and who responded positively to

Fig. 1. a Rigid-appearing capillary walls (H&E ×400). b Membrane spikes, black arrow (Jones silver satin, ×400). c Granular capillary loop staining with IgG, white arrow (anti-IgG immunofluorescence, ×400). d Subepithelial electron dense deposits (transmission electron microscopy, ×6,800).
renal disease when treated with penicillin. Lopez-Lopez et al. [24] identified a patient diagnosed with HIV and systemic lupus erythematosus with MN that improved after the administration of hydroxychloroquine and HAART. Unlike these cases, our patient’s workup did not reveal any underlying etiology for secondary MN. Our patient’s HIV infection was extremely well controlled on HAART with normal CD4 counts and an undetectable viral load, which points away from a connection between his infection and his renal disease. Likewise, our patient’s MN is likely unrelated to his history of hepatitis C infection due to negative tests for hepatitis C by PCR dating back more than 1 year prior to MN diagnosis. Moreover, our patient tested positive for PLA2R antibodies, which has been identified as a sensitive and specific marker associated with idiopathic MN [11]. While anti-PLA2R antibodies have been identified in patients with secondary MN related to lupus, hepatitis B and cancer, the prevalence of anti-PLA2R antibodies in these patient groups is low [12]. Testing for anti-PLA2R antibodies has been demonstrated as effective in differentiating idiopathic from secondary MN with a specificity of 89% [12–14]. Thus, it is evident that this patient represents a case of idiopathic MN in a patient with HIV. A blotting technique was used because western blot is the most sensitive assay for the detection of low levels of anti-PLA2R antibody [11]. Both commercially available and easy-to-perform immunofluorescence and ELISA methods would have likely resulted in a negative or ambiguous staining.

In an epidemiologic biopsy study of patients with HIV and renal disease, Szczech et al. [1] described HIV-positive patients with MN benefiting primarily from treatments other than HAART. Although these findings are consistent with other case reports and with our present case, they are in contrast to the cases mentioned earlier in which renal disease with MN was effectively treated with HAART [21, 23, 25, 26]. It is likely that these conflicting reports represent patients with different subtypes of MN. Renal disease responding to HAART therapy may signify instances of MN secondary to HIV. On the other hand, renal disease not responding to HAART but effectively treated with immunosuppression likely represents idiopathic MN that has developed in a patient infected with HIV. This thought is consistent with our current case of idiopathic MN as identified by positive anti-PLA2R antibodies that went into remission following immunosuppression as opposed to HAART.

Immune suppression via corticosteroids, cyclosporine, and alkylating agents is generally required for MN patients with high-grade proteinuria [27]. Unfortunately,
such therapies can be associated with significant toxicity, and many patients experience recurrence or develop resistance to treatment [28]. Acthar gel is obtained from the processing of the porcine pituitary gland and is currently the only Food and Drug Administration (FDA)-approved therapy for the treatment of nephrotic syndrome [29]. Despite the severity of proteinuria in our case, we tried to avoid immunosuppressive medications giving the history of HIV and hepatitis C. We initially used the angiotensin-converting enzyme inhibitor with no significant reduction of proteinuria for more than 6 months. We elected to use Acthar gel, because its safety profile appears to be much better than that of the other immunosuppressive and cytotoxic agents [29–31]. ACTH has been used to treat glomerulonephritis based on its steroidogenic effects via the stimulation of adrenal cortisol production. In addition to this steroidogenic mechanism, there is growing evidence that some renoprotective and antiproteinuric consequences of ACTH are related to steroid-independent melanocortin effects on extra-adrenal tissue, including the kidneys. These nonsteroidal results include anti-inflammatory, anti-apoptotic, and immunomodulatory effects that are important in treating glomerular disease [32, 33]. These mechanisms could help explain the therapeutic benefit of ACTH for our patient without producing significant immunosuppression and activating his underlying infections. One limitation in the present case is that the patient’s cortisol levels were not tested before or during ACTH therapy, which may have supported whether his observed remission was more related to steroidogenic or melanocortin effects of therapy.

It cannot be ruled out that the patient’s improvement in this case may have represented spontaneous remission of disease given that his improvement occurred within 2 years of diagnosis when most spontaneous remission occurs [19]. However, there was no change in the patient’s proteinuria for over 6 months prior to the initiation of ACTH followed by a significant decline in proteinuria in the months following the initiation of ACTH (fig. 2) indicating that remission was more likely related to therapy. The weekly positive serum anti-PLA2R with strongly positive anti-glomerular extracts could suggest the possible coexistence of autoantibodies other than anti-PLA2R. But the relative strength of the PLA2R band between the HGE lane and the recombinant PLA2R lane is more likely due to the relative amounts of the protein in each preparation. Note that the strong positive control also has a stronger band in glomerular extract vs. the recombinant PLA2R lane. The 2 molecules (native vs. recombinant PLA2R) are slightly different, as evidenced by the faster mobility of the recombinant band, which reflects under-glycosylation of the recombinant, cell-expressed band (a common problem with cell-expressed proteins). We know that a given anti-PLA2R-positive serum reacts less well with PLA2R that has been de-glycosylated [11], and thus it is possible that the anti-PLA2R antibodies react less well with the under-glycosylated, cell-expressed PLA2R as well. Although there may be other, unrecognized antigens in the glomerular extract, they would need to run at approximately the same molecular weight to yield such a pattern.

In conclusion, with advancements in HAART and improved control of HIV, cases of glomerular disease other than HIVAN, are likely to be seen more frequently in the HIV-positive population, and MN should be considered in such patients [8]. Patients with MN who have an apparent underlying cause of secondary MN may benefit from

Fig. 3. Western blot of HGE and extracts of cells transfected with human PLA2R or THSD7A were blotted with serum from the patient at a dilution of 1:10. The initial sample shows weak reactivity for native HGE and recombinant PLA2R. Insignificant amount of reactivity was detected in his follow-up sample (lower). In contrast, western blotting with a serum from a patient with primary MN with high anti-PLA2R titer shows strong signals (top). None of these sera reacted with the THSD7A autoantigen.
further diagnostic efforts to differentiate secondary from idiopathic etiologies. Testing for anti-PLA2R antibodies might help to identify cases of idiopathic MN that will require immunosuppression. Acthar gel might be useful in the management of complex cases of idiopathic MN.

Disclosure Statement
None.

References


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