Blockade of Osteopontin Inhibits Glomerular Fibrosis in a Model of Anti-Glomerular Basement Membrane Glomerulonephritis

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Introduction

End-stage renal disease is characterized by the development of fibrotic lesions in the glomerular and other renal compartments. Many diseases such as diabetes, hypertension and various inflammatory glomerulonephritis (GN) eventually lead to glomerular fibrosis, which permanently destroys glomerular architecture. Intensive studies in the past decades have shown glomerular fibrosis to be a very complicated process, involving many molecules and cells [1–4]. Similar to fibrosis occurring in other organs, TGF-β/Smad pathway plays a central role in mesangial cell activation and differentiation of fibroblasts, which in turn produce extracellular matrix (ECM) proteins [5–7]. It is less clear how the ECM proteins, once produced, are incorporated into the architecture of fibrotic tissue and whether those proteins can regulate gene expression in fibroblasts.

Osteopontin (OPN), or SPP1, is a highly modified integrin-binding ECM glycoprophosphoprotein. OPN participates in many physiological and pathological events, including maintenance or re-establishment of tissue integrity, inflammatory processes and tumor metastasis [7–9]. OPN also functions as both an anti- and pro-inflammatory molecule in autoimmune pathogenesis, as well as in other immune responses [10]. More recently, a critical
role of OPN in fibrosis has been reported [11–13]. One study showed that OPN is required for differentiation of myofibroblasts, a major player in fibrosis [14]. In addition to its regulatory function, OPN is an important structural molecule.

The role of OPN in both renal development and disease has been explored [15, 16]. Constitutive expression of OPN in renal tubular epithelium is required for prevention of kidney stone formation. Roles of OPN in renal fibrosis, especially in renal interstitial tissue, have been well investigated in OPN null mice. OPN may modulate angiotensin II-induced inflammation, oxidative stress, and fibrosis of the kidney [17]. Several studies reported an association of upregulation of glomerular OPN expression with macrophage accumulation and progressive renal injury [18, 19]. Anti-OPN antibody treatment reduced proteinuria and prevented loss of renal function, suggesting OPN to be a pro-inflammatory molecule [20]. However, experiments using OPN null mice showed a lack of any in vivo functions of OPN in an anti-GBM GN model [21]. Although it still lacks direct evidence, several studies have suggested a potential role of OPN in glomerular fibrosis [22, 23].

We have established a rat model for autoimmune anti-GBM GN, which is induced by immunization with a T cell epitope pCol(28–40) derived from α3 chain of type IV collagen (Col4α3) [24, 25]. The animals undergo two distinct overlapping stages after disease induction: (1) CD4+ T cell-mediated glomerular inflammation followed by (2) severe glomerular fibrosis. We have previously reported upregulation of OPN in a novel multinucleated myofibroblasts in the glomeruli and massive deposition of OPN on the fibrotic tissue during fibrotic stage [26]. In the present study, we investigated whether blockage of OPN activity at early inflammatory/early fibrotic stage would alter fibrosis progress in our model.

**Evaluation of GN Severity and Glomerular Fibrosis**

Urine samples were collected daily for evaluation of albuminuria. In some cases, 24-hour urine was collected using metabolism cages and urine albumin concentrations were determined with a Rat Albumin ELISA Quantitation Set (Bethyl Lab, Montgomery, Tex., USA). GN severity in experimental rats was graded by histology. A portion of kidney tissues were fixed in Bouin’s solution, and glomerular injury score was calculated as described previously [24]. The tissue sections were processed for PAS, trichrome staining. Glomeruli with fibrotic crescent lesions were identified and expressed as a percentage of all glomeruli. Tissue sections stained with anti-collagen1 antibody were used for calculation of collagen positive area in glomeruli with a program (Nikon). A portion of renal tissue was processed for transmission electron microscopy [24]. Glomerular fibrosis was also evaluated by RT-PCR on expression of collagen 1α1 following a previously published method [5]. In addition to acidic ribosomal phosphoprotein P0, housekeeper gene GAPDH was also used as another control. Real-time PCR was performed for the collagen 1α1 gene using P0 as a comparison.

**In vivo Blockage of OPN Activity with OPN Antibody**

A rabbit anti-OPN polyclonal antibody, which has been demonstrated to react with OPN and block its activity, was provided by Dr. William Butler, a pioneer in OPN research, University of Texas Health Science Center at Houston. A normal rabbit serum (NRS) was used as a negative control. Rats were immunized with pCol(28–40). A total of 800 μl of antisera or normal sera were injected i.v. into each immunized rat at days 26, 28, 33 and 38 postimmunization. The rats were sacrificed at day 40 or day 50, or at their death. Kidneys of 3 rats from each group were used for isolation of glomerular cells at days 40 and 50. The cells were analyzed by flow cytometry or for overnight culture to detect fibroblast-like cells in wells of a 96-well plate. Approximately 6–7 pictures were taken to cover an entire well under ×100 magnification. Fibroblast spread area was determined by computer-assisted area calculation (NIS-Element 2.3, Nikon). A portion of the kidney was fixed for histological evaluation. Sera OPN concentrations were determined by Rat OPN Assay Kit from Immuno-Biological Laboratory (IBL, Guma, Japan). Rabbit IgG concentration in recipients’ sera were measured by ELISA using a pair of polyclonal anti-rabbit IgG antibodies (Bethyl Lab, Montgomery, Tex., USA). OPN-mediated peritoneal macrophage invasion assay was performed following a published method with minor modifications. Rat recombinant MCP-1 (R&D, Minneapolis, Minn., USA) was used as a chemoattractant.

**Methods**

**Antigen Preparation and GN Induction**

Nephritogenic T cell peptide pCol(28–40) was commercially synthesized and dissolved in milli-Q water at a 1-mM concentration. Female Wistar Kyoto (WKY) rats (4–6 weeks of age) were purchased from Harlan (Indianapolis, Ind., USA). The rats were maintained in the animal facility at the University of Texas Health Science Center and allowed to acclimate for a minimum of 3 days. Rats were immunized with a peptide (0.125 μmol) emulsified in CFA, in one hind footpad and at the base of the tail. Rats immunized with CFA alone served as controls. All animal procedures were approved by the IACUC.

**Purification and Analysis of Glomerular Cells**

Glomerular cells were isolated at days 40 or 50 postimmunization for detection of OPN expression and determination of leukocyte populations. After kidneys were perfused in situ with PBS, glomeruli were isolated to a high purity as previously described using repeated low-speed centrifugations. The isolated glomeruli with purity over 95% were digested with a mixture of collagenase IV and AccutaseTM (eBioscience, San Diego, Calif., USA) at room temperature for 30 min with gentle stirring. The cells were released by repeated pipetting and collected by centrifugation. Dead cells were further removed by Ficoll density centrifugation. The cells were used for immunofluorescence, flow cytometry or further culture with antibody or normal sera. For immunofluorescence, the cells were placed in each well of a 4-well chamber slide.
overnight (Fisher, Rochester, N.Y., USA). For flow cytometry, the isolated cells were stained with paired antibodies (PE-anti-CD4 (OX4) vs. FITC-anti-rat IgG, or PE-anti-CD8 (OX8) vs. FITC-anti-CD11b/c). The stained cells were analyzed with a flow cytometer (FACSCalibur, Becton Dickinson, San Jose, Calif., USA). Glomerular fibroblasts were isolated from early stage of fibrosis. In some cases, attached fibroblasts were collected by trypsin digestion and used for other purposes. Fibroblast proliferation was determined by the Quick cell proliferation Assay Kit (BioVision).

Statistical Analysis
One-way ANOVA test was used for comparisons between experimental groups. All calculations were performed using Prism version 3.0.

Results

OPN Antiserum Neutralizes OPN in vivo
The glomerular inflammation in our model peaks at approximately 28–35 days, and gradually vanishes. On the other hand, the glomerular fibrosis is histologically detectable as early as days 25–28, and rapidly progresses after day 35. We tested whether blockage of OPN activities at a late inflammatory/early fibrotic stage could still affect inflammation and/or progress of fibrosis. A rabbit OPN antiserum, which effectively inhibited OPN-mediated migration of peritoneal macrophage in in vitro invasion assays, was used to block OPN activity at late inflammatory early fibrotic stage (fig. 1a). Since glomerular fibrosis in our model starts as early as 25 days postimmunization, OPN antiserum was injected intravenously into 8 immunized rats at days 26, 28, 33 and 38 postimmunization to ensure a constant level of the antibody (fig. 1c). Another group of immunized rats (total 8) received normal rabbit serum (NRS) as controls. Ten immunized rats, without having received any sera, were used as positive controls.

Two assays were performed to evaluate effectiveness of the injected OPN antiserum. First, sera OPN concentrations from three groups of rats at day 40 were determined by ELISA. As compared to normal rabbit serum-treated rats, a reduction (>85%) in the sera OPN concentration was observed in the OPN antiserum-treated rats, suggesting neutralization of OPN in vivo by the antiserum (fig. 1b). Second, final sera from the OPN antibody-treated rats at day 50 postimmunization showed the presence of rabbit IgG at 26.3 ± 6.75 μg/ml.

OPN Antiserum Treatment Does Not Reduce Glomerular Inflammation
Flow cytometry analyses were performed on glomerular cells, which were isolated from the OPN- or normal serum-treated groups at days 40 and 50. At day 40, 3 rats/group were analyzed. Both the OPN and normal antiserum groups showed the presence of CD4+ T cells and CD11+ cells (mostly macrophages) (fig. 2a–c). Due to high mortality, only 2 rats were available for analysis from the normal sera group at day 50. In contrast, all rats from the anti-OPN group survived and 3 were used for analysis. At day 50, there were a few, if any, glomerular CD4+ T cells (0.1%) and CD11+CD8+ cells (0.2%) in 2 normal rabbit serum controls. On the other hand, the rats from the OPN antiserum group showed the presence of a significant number of CD4+ T cells (5.1%) and CD11+CD8+ cells (1.1%) (fig. 2a–c). Persistent glomerular inflammation in OPN antiserum-treated rats was further confirmed by immunofluorescence. ED1+ macrophages were observed within the glomeruli of OPN antiserum-treated rats (fig. 2d). However, no macrophages were present within the glomeruli of normal rabbit sera-treated rats except for a few surrounding fibrotic crescents (fig. 2e). Albuminuria was closely monitored daily. There were no significant differences in albu-
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Fig. 2. Treatment of immunized rats with OPN antiserum at late inflammatory/early fibrotic stage does not fully inhibit glomerular inflammation. a, b Flow cytometry on infiltrating leukocytes in the glomeruli of NRS- or α-OPN-treated rats at days 40 and 50 postimmunization with pCol(28–40). c Summary of glomerular various cell populations detected by flow cytometry in different groups as indicated. d, e A pair of immunofluorescent images shows the presence of many ED1+ macrophages within a glomerulus of an α-OPN-treated rat (e), but not in NRS-treated rats (d). Fibrotic tissue in d is outlined by arrowheads.

Fig. 3. OPN antiserum treatment does not improve albuminuria in the experimental rats. a Albuminuria, expressed as mg/24 h, NRS- and OPN antiserum-treated rats at day 45 after immunization at their death. b Urine albumin, as detected by SDS-PAGE, in representative rats from different groups.

minuria/24-hour or urine albumin concentrations among the two groups during the experiments (fig. 3). However, the majority of normal serum-treated rats developed oliguria after day 40 as indicated by the low urine volumes (<5 ml) per 24 h. On the other hand, oliguria did not occur in OPN antiserum-treated rats. Serum creatinine concentrations at day 50 were determined. Serum creatinine in the OPN antiserum group was elevated (2.8 ± 1.7 mg/dl, n = 5). However, it was much lower than for the NRS group (4.6 ± 0.9 mg/dl, n = 2, others died before day 50) and the no-treatment group (4.9 ± 1.1 mg/dl, n = 5, others died before day 50). The results suggested that OPN antiserum had a limited influence on glomerular inflammation when given at the late inflammatory/early fibrotic stage.

Treatment with OPN Antiserum Reduces Glomerular Fibrosis

Histopathological analyses, based on specially stained fibrotic tissues, showed that OPN antiserum treatment led to a significant reduction in glomerular fibrosis, both quantitatively and qualitatively (fig. 4a, b). The rats from the OPN antiserum-treated group had a much lower frequency of glomeruli with fibrotic crescents (39%) than those from the normal rabbit sera group (79%) (fig. 4c).

Although rats from the normal rabbit sera group showed a slightly lower frequency of fibrotic glomeruli than in the group without any sera treatment (86.6%), there was no significant difference between the two (p > 0.05). Furthermore, fibrotic crescents in the OPN antiserum-treat-
ed group were structurally different from the two controls. First, in the two control groups (normal serum or no serum), an entire glomerulus was often occupied by fibrotic tissue without any normal tissue left (Fig. 4, 5). Severe glomerular fibrosis greatly affected surrounding tissues, especially the tubules, which had greatly dilated lumens with thin, flattened walls. In contrast, fibrotic tissue was limited to a small area of Bowman’s capsule of affected glomeruli in the OPN antiserum-treated animals. Circular crescentic lesions were not observed in any OPN antiserum-treated rats, and no surrounding tissue was affected. Second, in the OPN antiserum-treated rats, glomeruli remained largely intact with most of capillary lumens patent and GBM was not significantly thickened (Fig. 5b, d). On the other hand, there was profound destruction of glomeruli in the control groups (Fig. 5a, c).

Glomerular fibrosis was next evaluated by collagen 1α1 deposition and expression. Three rats were randomly selected from each group. The renal sections from these rats were stained by anti-collagen 1α1 antibody (Fig. 6a, b). Collagen 1α1-positive areas in five randomly selected glomeruli from each rat were calculated (Fig. 6c). OPN antiserum treatment led to a significantly reduced collagen 1α1-positive area. We next calculated the net increase in the collagen-positive areas using normal glomeruli as a baseline; the positive area in the antiserum-treated rats reduced to 30% of those from normal serum controls. RT-PCR on glomerular RNA showed that all 5 rats treated with OPN antiserum showed a reduced collagen 1α1 expression as compared to those treated with normal rabbit serum, although the reduction level in each OPN antibody-treated individual was variable (Fig. 6d). Quantitative RT-PCR showed a 6.6-fold reduction in collagen 1α1 expression in the OPN antiserum-treated group as compared to the normal rabbit sera group.

**Discussion**

OPN has been widely recognized as both pro- and anti-inflammatory molecules in various diseases including autoimmune diseases [7–9]. More recently, a critical role of OPN in scar formation or fibrosis in several organs, including kidney, has been reported [10–13]. Previous studies have speculated that OPN played a role in glomerular fibrosis [22]. In the present study, we provided additional evidence that OPN plays an important role in glomerular fibrosis in rat anti-GBM GN. Treatment with an OPN antiserum at the late inflammatory/early fibrosis stage led to a significant reduction in glomerular fibrosis, in both quantity and quality. OPN antibody has been reported to reduce severity of GN in a rat nephrotoxin model when given at an early inflammatory stage [19]. In our study, administration of anti-OPN at the late inflammatory stages did not significantly reduce glomerular inflammation, especially CD4+ cells. Thus, reduction in glomerular fibrosis induced by OPN antiserum in this study may not be mediated by the inhibition of inflammation.

Although several studies, including the present study, suggested a critical role of OPN in glomerular inflammation or renal fibrosis, a previous paper reported that OPN may have no effect on murine anti-GBM GN based on an OPN KO strain [21]. Although it is difficult to address this question, there are a few possibilities which may ex-
plain the conflicting results between the OPN KO model and the antibody blockage model. First, the pathological mechanisms may be different in the two models. In the murine model, glomerulonephritis is induced by an immune response to the 'embedded' foreign antibody bound to GBM [21]. In our model, glomerular inflammation is mediated by the T cell response to a well-characterized self T cell epitope from autoantigen collagen 4a3 [24, 25]. Second, the pathological mechanism may differ in different species. Our unpublished data suggest that T cell response alone may not be sufficient to mediate severe anti-GBM GN in mice. Third, it is a well-known phenomenon that in some cases, genetic manipulation may lead to an unexpected result due to complicated regulation/interaction among genes. For example, bone development in OPN KO mice is largely normal.

Different approaches have been tested in experimental models of renal fibrosis. It is clear that TGF-β/Smad, as well as angiotensin II and ET1, play a central role in mesangial cell activation and differentiation of fibroblasts to myofibroblasts [4–7]. However, TGF-β may no longer be critical once fibrosis is initiated. The fibrotic mechanism downstream of the TGF-β pathway has been studied. Several studies suggest a role of ECM, such as SPARC, in renal fibrosis [27]. Several studies have determined OPN's role in renal fibrosis due to ureteral obstruction, ischemia, and cyclosporine toxicity using OPN knockout mice. These studies suggested that OPN may have mul-

Fig. 5. Histopathology showing fibrotic tissue in glomeruli in OPN antiserum-treated rats (anti-OPN) (b, d, f, h), as compared with NRS-treated controls (a, c, e, g). All rats were previously immunized with pCol(28–40). a, b HE staining; c, d PAS staining; e, f trichrome staining; g, h Jones silver staining. × 300.

Fig. 6. Reduced collagen 1α1 expression and its deposition in the glomeruli in OPN antiserum-treated rats. a, b Immunofluorescent micrographs show massive deposition of collagen 1α1 (red) in an NRS-treated rat (a), but much less in an α-OPN-treated one (b). GBM was counter-stained by SR-13 antibody (green). × 500. c Comparison of collagen 1α1-positive areas in glomeruli in different groups. The kidney sections were stained for collagen 1α1 (see a, b), and positive areas in a glomerulus were captured using a computer-aided program and calculated as percentage of entire area of a glomerulus. d RT-PCR detection of collagen 1α1 (Col1A) in NRS- or α-OPN-treated rats. Two housekeeper genes, GAPDH and acidic ribosomal phosphoprotein (PO), were used for comparison. Each lane represents one individual.
tiple functions including recruitment of macrophages, induction of apoptosis, and modulation of angiotensin II [17]. Our study again demonstrated that OPN could be critical for fibrosis.

However, the mechanism by which OPN antiserum inhibits fibrosis when given at a late inflammatory stage of GN remains unclear in the present study. It is unlikely that inhibition of fibrosis by OPN antiserum is directly related to the TGF-β pathway, because OPN antiserum was given after fibrosis had been initiated. One possible mechanism is that reduction in fibrosis is due to inhibition of glomerular inflammation by OPN antibody. However, it is also less likely, as glomerular inflammation was not significantly affected by the antibody. Alternatively, OPN antibody may act at two levels on OPN after fibrosis has been initiated. First, fibrosis, including glomerular fibrosis, is often defined as a wound-healing response that has gone out of control, resulting in excess deposition of ECM. The key cellular mediators are the myofibroblasts, which differentiate from fibroblasts (or glomerular cells such as mesangial cells) and are the main source of excessive ECM proteins [3, 28]. Interaction with ECM may be required for fibroblasts to differentiate [29]. Thus, blockage of OPN activities in the present study may inhibit differentiation of myofibroblasts. This was evidenced by a much smaller number of fibroblast-like cells in glomeruli in OPN antiserum-treated rats [unpubl. data]. Furthermore, our in vitro experiments demonstrated a significantly reduced proliferation rate and altered adhesion in CD90+ glomerular fibroblast-like cells freshly isolated at the early fibrotic stage. Through the analysis of global gene expression, our recent results showed altered expression patterns in fibrosis-related genes [unpubl. data]. In fact, the requirement of OPN for the differentiation of fibroblasts has been demonstrated in the cardiac fibrosis model [14]. Second, OPN may facilitate assembly of ECM protein to form fibrotic tissue. OPN has been recognized as a structural protein with multiple binding sites for collagen and other molecules such as adhesion molecules [30, 31]. A previous study using OPN mutant mice showed a disorganization of matrix in skin incisions, suggesting involvement of OPN in ECM protein assembly [32]. Thus, antibody inhibition of OPN may affect organization of collagen fibers and/or adherence of fibroblasts. However, our antibody blockage experiments were not able to provide direct evidence to support this possibility. We have shown previously a massive deposition of OPN on fibrotic tissue at the mid to late stage of fibrosis [26]. Binding of antibody to OPN may disturb the architecture of fibrotic tissue. This hypothesis can be tested if the OPN antibody is administered after fibrosis has been initiated.

In summary, our study showed that OPN antibody given at the late inflammatory/early fibrotic stage inhibited the progression of fibrosis. The effectiveness of the antibody was not due to inhibition of glomerular inflammation. Several results demonstrated that reduction in fibrosis was through inhibition of glomerular fibroblast-like cell growth/differentiation. OPN has been a therapeutic target in several other fibrotic diseases. Our data showed that the OPN antibody given at late stage of GN was able to halt or inhibit progression of fibrosis after it had been initiated. It remains to be determined, however, whether the antibody treatment may reverse the diseases. Nevertheless, our model will surely be an excellent tool for investigation of the fibrosis mechanism downstream of the TGF-β pathway.

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References

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