# World Journal of Clinical Cases

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CASE REPORT

## Integrated Chinese and Western medicine in the treatment of a patient with podocyte infolding glomerulopathy: A case report

Mei-Ying Chang, Yu Zhang, Ming-Xu Li, Fang Xuan

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#### **Abstract**

Podocyte infolding glomerulopathy (PIG) is a newly described and rare glomerular disease. To date, only approximately 40 cases have been reported globally.

#### CASE SUMMARY

A 26-year-old female patient presented to our hospital with a complaint of intermittent edema of both lower limbs over the past 2 years. The patient was diagnosed with PIG. She was prescribed corticosteroid therapy in other hospitals during the initial stage, to which she had responded poorly and had developed femoral head necrosis. Therefore, we administered immunosuppressants, reninangiotensin system inhibitors, combined with traditional Chinese medicine. The patient was followed for 1 year, during which her clinical condition improved.

Integrated Chinese and Western medicine may be effective for PIG treatment, which requires active intervention to improve prognosis.

Key Words: Corticosteroid therapy; Immunosuppressive agents; Podocyte-infolding glomerulopathy; Renin-angiotensin system; Traditional Chinese medicine; Case report

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**Core Tip:** Due to the limited number of reported cases, insufficient information on the characteristics, diagnosis, and treatment of podocyte infolding glomerulopathy is available. Based on our case and those reported in PubMed, we believe that treatment with corticosteroids, immunosuppressants, and reninangiotensin system inhibitors is effective. Some patients cannot tolerate corticosteroids and immunosuppressants. When adverse effects occur, clinicians should avoid making negative treatment. Doctors should actively intervene and offer patients treatment suggestions, among which traditional Chinese medicine may be an effective treatment method.

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#### INTRODUCTION

The concept of podocyte infolding glomerulopathy (PIG) was put forward by the Japanese Society of Nephrology in 2008[1]. PIG can be diagnosed based on histopathological findings. Its pathological feature is the presence of microspheres, microtubules, or both in the glomerular basement membrane (GBM) under electron microscopy, and podocyte infolding into the GBM[1]. Knowledge of the pathogenesis and progression of PIG is rare, due to the limited number of reported cases[1]. Thus, integrated and definitive immunological therapies are not yet available. Some case reports have suggested that corticosteroid therapy can be used to treat this disease. However, it is difficult to assess its efficacy because of the limited number of cases and the lack of long-term follow-up of cases [2-15]. Overall, 15 consecutive cases of PIG, most of which were from Asia, have been reported since the initial study conducted by Joh et al[1] in 2008. Due to the unclear etiology and varied clinical presentations, PIG is more likely a pattern of injury than an etiological diagnosis[7,11].

Therefore, further clinical data need to be accumulated to gain a better understanding of this type of glomerulopathy. Herein, we report a case of PIG in a 26-year-old woman with nephrotic syndrome without renal functional impairment. Along with a literature review of previous cases, we summarize the clinical features and treatment methods for PIG.

#### CASE PRESENTATION

#### Chief complaints

A 26-year-old Chinese woman presented to our hospital with intermittent edema of both lower limbs of 2 years' duration.

#### History of present illness

The patient developed pitting edema of the lower extremities in January 2020. She went to the Sixth Medical Center of Chinese PLA General Hospital. Laboratory data are presented in Table 1. A renal biopsy was performed. She was prescribed 80 mg valsartan and 60 mg prednisone daily. After 8 wk of treatment, the dose of prednisone was reduced by 10 mg every 2 wk. However, she had poor compliance and stopped the medication when the dose had been reduced to 20 mg daily in May 2020. Laboratory tests were not performed thereafter. The patient later developed intermittent edema but ignored it.

In December 2021, she was re-hospitalized due to bilateral lower extremity edema, and her 24-h proteinuria level was 4840 mg and serum albumin level was 16.8 g/L. This time, she was treated with 40 mg of intravenous methylprednisolone and 240 mg of oral allisartan isoproxil daily. Seven days later, she developed hip pain and weakness. Hip magnetic resonance imaging showed bilateral ischemic necrosis of the femoral head with joint cavity effusion; therefore, the intravenous methylprednisolone was discontinued, and oral methylprednisolone 36 mg/day was initiated and rapidly tapered (1 tablet per week, or 4 mg). Simultaneously, she was administered 100 mg cyclosporine twice a day while continuing allisartan isoproxil. Her 24-h proteinuria was 5803-7650 mg. In February 2022, her 24-h proteinuria was 6370.65 mg and she came to our hospital for treatment. At admission, the patient had mild edema of both lower limbs, hip pain, and foamy urine. The tongue was pale, with a thin white coating, and her pulse was weak.

#### History of past illness

One year earlier, the patient had developed an increase in blood pressure, reaching 170/100 mmHg. She

Table 1 Laboratory data before the renal biopsy							
Test	Results						
24-h proteinuria, mg	8730						
Hemoglobin, g/L	101 (normal: 115-150)						
White blood cells, $\times 10^9/L$	3.06 (normal: 3.5-9.5)						
Platelets, $\times 10^9/L$	386 (normal: 125-350)						
Cholesterol, mmol/L	6.96 (normal: 3-5.7)						
Triglyceride, mmol/L	2.36 (normal: < 1.7)						
Creatinine, µmol/L	67.4 (normal: 53-97)						
Albumin, g/L	13 (normal: 40-55)						
ANA	Negative						
ANCA	Negative						
Anti-double stranded DNA antibody	Negative						
Rheumatoid factor	Negative						
C3, mg/dL	98.4 (normal: 90-180)						
C4, mg/dL	33.6 (normal: 10-40)						
IgA, mg/dL	292 (normal: 70-400)						
IgM, mg/dL	119 (normal: 40-230)						
Anti-PLA2R antibody	Negative						

ANA: Antinuclear antibodies; ANCA: Antineutrophil cytoplasmic antibodies; IgA: Immunoglobulin A; IgM: Immunoglobulin M; PLA2R: Phospholipase A2 receptor.

was treated with 240 mg of oral allisartan isoproxil daily.

#### Personal and family history

The patient denied any family history of kidney disease.

#### Physical examination

On physical examination, her vital signs were as follows: Body temperature, 36.8 °C; blood pressure, 120/86 mmHg; heart rate, 90 beats per min; respiratory rate, 18 breaths per min. Furthermore, mild edema was observed in both lower limbs.

#### Laboratory examinations

The results of laboratory examination were as follows: Urinary protein = 3+, red blood cells =  $50/\mu$ L, 24h proteinuria = 6165 mg, serum creatinine = 70.7 μmol/L, serum albumin = 24.2 g/L, cholesterol = 8.03 mmol/L, and triglyceride = 3.24 mmol/L.

#### Imaging examinations

Under a light microscope, 28 glomeruli with no spherical sclerosis and three small cell crescents were visible. The mesangial area showed mild focal segmental proliferation and matrix thickening. Capillary loops were well open, and focal epithelial cell vacuolar degeneration was present. The basement membrane was mildly thickened in focal segments. No significant atrophy or inflammatory cell infiltration was seen in the renal interstitium. No significant thickening of the renal vasculature was observed (Figure 1A). Under an electron microscope, extensive podocyte foot-process effacement with infolding into the GBM was seen. The GBM was diffusely thickened with numerous microspherical and microtubular structures (Figure 1B-D). Immunofluorescent staining of the biopsy specimen showed granular deposition of immunoglobulin (Ig)G along the mesangial areas, but with no staining for C3, IgA, IgM, C1q, or fibrinogen (Supplementary Figure 1).

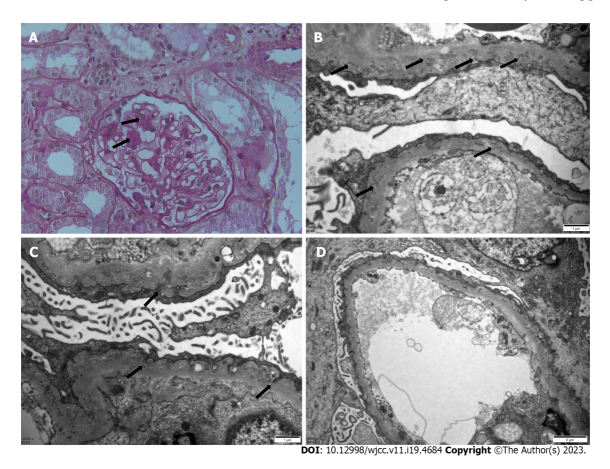


Figure 1 Kidney biopsy findings. A: Light microscopic findings on periodic acid-Schiff staining. The mesangial area showed mild focal segmental proliferation and matrix thickening (arrows). The basement membrane is mildly thickened (original magnification, × 400); B: Electron microscopy of kidney biopsy specimen. Thickened glomerular basement membrane (GBM) with numerous diffusely scattered microspherical or microtubular structures (arrows), characteristic of podocyte infolding glomerulopathy (original magnification, × 12000); C: Extensive podocyte foot-process effacement with infolding into the GBM (arrows) (original magnification, × 12000); D: There are no electron-dense deposits (original magnification, × 8000).

#### FINAL DIAGNOSIS

Combined with the patient's medical history, a final diagnosis of PIG was made.

#### **TREATMENT**

We continued allisartan isoproxil treatment and increased the cyclosporine dose to 150 mg in the morning and 100 mg at night. Oral mycophenolate mofetil (500 mg twice a day) was also administered.

In addition, we treated the patient with traditional Chinese medicine. First, we diagnosed the patient as having blood stasis and water stagnation, and treated her with Danggui-Shaoyao-San, a compound formula in traditional Chinese medicine, comprised of six herbs: Paeonia lactiflora Pall. 20 g, Alisma orientalis (Sam.) Juzep. 10 g, Angelica sinensis (Oliv.) Diels. 10 g, Poria cocos (Schw.) Wolf. 20 g, Atractylodes macrocephala Koidz. 20 g and Ligusticum chuanxiong Hort 10 g. Over the course of treatment, the edema of the patient was gradually reduced, and the patient's condition was subsequently mainly characterized by deficiency of the spleen and kidney, manifested as fatigue. We differentiated the syndrome as deficiency of the spleen and kidney, combined with wind evil, and treated her with Modified Huangqi Chifeng decoction to strengthen qi further. The whole prescription was composed of seven herbs: Astragalus membranaceus Bge 30 g, Euryale ferox Salish 20 g, Rosae Laevigatae Fructus 10 g, Radix Paeoniae Rubra 10 g, Saposhnikoviae Radix 10 g, Rhizoma Dioscoreae Nipponicae 20 g, and Hedyotis Diffusae Herba 20 g.

### OUTCOME AND FOLLOW-UP

The patient was followed-up for 1 year, during which we adjusted the drugs slightly according to her symptoms. As shown in Figure 2, her serum albumin levels gradually increased and her urinary protein

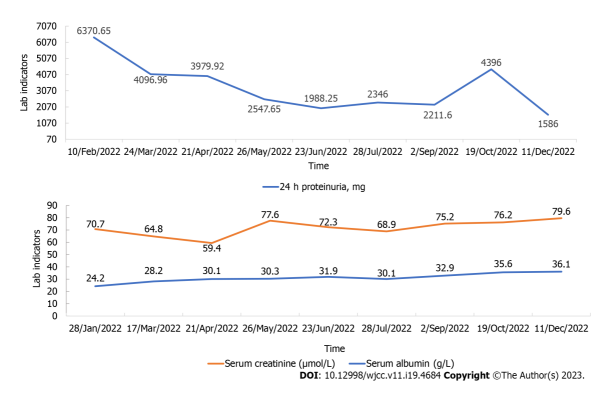


Figure 2 Changes in 24-h proteinuria (mg), serum creatinine (µmol/L), and serum albumin (g/L) over time.

excretion gradually decreased, with an increase in October 2022, which was considered to be related to an upper respiratory infection. Overall, her clinical condition improved.

#### DISCUSSION

The concept of PIG, as a newly recognized form of glomerular disease, was first proposed by Joh et al[1] in 2008. Although PIG cases are increasing, it remains unclear whether PIG is a new disease entity or simply a transient morphological phenomenon [2,15]. Hence, it is important to discuss the salient features of this case to gain a better understanding of the nature of the disease.

Microstructures typical of PIG were first described in the 1970s[13], referred to as rounded extracellular particles (REPs)[16]. According to Dales and Wallace, REPs in a patient with membranous nephropathy (MN) are nuclear pore organelles based on their morphologic similarity and the presence of autoantibodies in cytoplasmic and nuclear organelles of the patient's serum[17]. Zhang et al[8] considered that PIG was actually first identified in 1992 when Sato et al[18] described patients with collagen diseases whose renal biopsies demonstrated fine electron-dense deposits in the GBM. It is difficult to determine when PIG first appeared; however, in 2008, a new terminology for PIG was proposed by Joh et al[1], who defined it as microstructures (microspheres and/or microtubules) associated with podocytic invagination (large cytoplasmic projections from podocytes) into the GBM. In contrast to granular debris commonly seen in diseases such as MN, these microstructures are thought to originate from the primary podocyte cell membranes.

To date, 40 cases of PIG have been reported globally: 36 from Asia (28 from Japan, 6 from China, and 1 each from Korea and India), 2 from North America (1 from the United States, 1 from Canada), 1 from Argentina, and 1 from Europe [1-15]. We summarized the clinical profiles of the 40 cases, 25 of which were included in the study by Joh et al[1], while the remaining 15 cases are shown in Table 2.

As shown in Table 2, the age range of the remaining 15 cases was 4-79 years, which was greater than that reported by Joh et al[1], indicating a higher prevalence of PIG and an incidence in a broader population. Significantly more female than male patients were noted (female:male 13:2). Proteinuria reached 16798.8 mg/day, which was markedly higher than the maximum value of 7500 mg/day reported by Joh et al[1]. Three patients, with a mean age of 42.7 years, had increased serum creatinine levels exceeding 1.5 mg/dL (133 µmol/L). On the one hand, this suggests the need for early follow-up of patients and early intervention to slow down the progression of the disease. On the other hand, as PIG is not yet widespread, health education and regular medical check-ups of healthy individuals are important.

In Joh et al's report, 17 of the 25 cases were treated with corticosteroids alone or in combination with immunosuppressants[1]. In the 15 newly reported cases of PIG summarized in Table 2, prednisolone (or

#### Table 2 Clinical profile of podocyte infolding glomerulopathy (15 cases, Case No. 26-40)

Case No.	Country of the study	Year of the study	Age (years)	Sex	Initial proteinuria (mg/day)	Hematuria	Renal function (Creatinine, µmol/L)	Blood pressure (mmHg)	Treatment	Concomitant diseases
Case 26[2]	Japan	2013	14	F	3060	1-4	48.6		PSL (40 mg/day)	First biopsy: np; Second biopsy: FSGS
Case 27[3]	Japan	2014	79	M	1426		113.2	140/67	PSL (20 mg/day)	HTN, MM
Case 28[4]	South Korea	2016	44	F			39.8	100/70	PSL (10 mg/day); ARB (15 mg/day)	SLE
Case 29[5]	India	2018	45	F	5800	20	145.9	130/80	High-dose PSL, MMF, 1 g of rituximab	UCTD
Case 30[6]	China	2018	52	F						pSS, HT
Case 31[7]	Germany	2019	56	F		35 U/L	386.3		High dose Pred, rituximab (1 g × 2)	RA
Case 32[8]	China	2019	27	F	629		168.0		PSL (48 mg/day), HCQ (0.2 g twice a day)	pSS, HT
Case 33[8]	China	2019	23	F	16796.8	+	46.9		PSL (40 mg/day), HCQ (0.2 g twice a day) tacrolimus (1 mg twice a day)	SLE
Case 34[9]	Argentina	2020	38	F						SLE
Case 35[10]	China	2020	4	M	3670				Given the resistance to steroids and tacrolimus, only treated him with diuretics and (ACEIs)	SIOD, hypothyroidism
Case 36[11]	China	2020	33	F	2124	-	36.2		ARB, Radix Astragali and Huangkui	UCTD
Case 37[12]	Japan	2020	35	F						pSS, scleroderma
Case 38[13]	United States	2021	60	F	2286		61.9		Rituximab (1 g × 3)	PLA2R antibody(+)
Case 39[14]	Canada	2021	52	F		11-20	111.4	158/97	ARB (300 mg/day); spironolactone (25 mg/day); Pred (1 mg/kg)	HBcAb(+); HBsAb(+)
Case 40[15]	China	2021	61	F	2060	19	75.1		ARB (100 mg/day), HCQ (100 mg twice a day), PSL (40 mg/day), CTX (600 mg/month)	SLE, HTN

ACEIs: Angiotensin-converting enzyme inhibitors; ARB: Angiotensin II receptor blocker; CTX: Cyclophosphamide; F: Female; FSGS: Focal segmental glomerulosclerosis; HCQ: Hydroxychloroquine; HT: Hashimoto's thyroiditis; HTN: Hypertension; M: Male; MM: Multiple myeloma; MMF: Mycophenolate mofetil; pSS: Primary Sjögren's syndrome; PLA2R: Phospholipase A2 receptor; Pred: Prednisone; PSL: Prednisolone; RA: Rheumatoid  $arthritis; SIOD: Schimke\ immuno-osseous\ dysplasia; SLE: Systemic\ lupus\ erythematosus;\ UCTD:\ Undifferentiated\ connective\ tissue\ disease.$ 

> prednisone) was administered to nine patients (two were treated with prednisolone or prednisone alone [2,3], five were treated with prednisolone/prednisone combined with immunosuppressants, such as mycophenolate mofetil, rituximab, hydroxychloroquine, tacrolimus, and cyclophosphamide[5,7,8,15], one was treated with prednisolone combined with the angiotensin II receptor blocker (ARB) fimasartan [4], one was treated with prednisolone combined with the ARB irbesartan and spironolactone[14]).

Immunosuppressants were administered to six patients (one received rituximab alone)[13]. One patient was administered an angiotensin-converting enzyme inhibitor (ACEI) alone[10], and another was treated with the ARB olmesartan medoxomil combined with the traditional Chinese medicines Radix Astragali and Huangkui[11]. The treatments of the remaining three cases were not mentioned in the original articles[6,9,12]. Although treatment guidelines for PIG have not been established to date, a review of the literature above showed that corticosteroids are most widely used for PIG, followed by immunosuppressants and ACEI/ARB.

Analysis of concomitant diseases showed that PIG was mostly associated with collagen diseases, such as systemic lupus erythematosus (SLE) (4/15), rheumatoid arthritis (1/15), and primary Sjögren's syndrome (2/15). In Joh et al's report[1], 13 of the 25 cases had combined SLE. It was also seen in patients with non-collagen diseases, such as undifferentiated connective tissue diseases (2/15), Schimke immuno-osseous dysplasia (1/15), hypertension (1/15), multiple myeloma (1/15), and Hashimoto's thyroiditis (1/15). The exact relationship between collagen diseases and PIG requires further research.

In addition, in one case[2], two renal biopsies were performed; the first renal biopsy showed only PIG, but the second also suggested focal segmental glomerulosclerosis. Thus, in PIG cases, we should pay attention to follow-up, particularly when the patient's condition does not improve for an extended period or worsens, which may indicate the need for a repeat renal biopsy.

Our patient was a 26-year-old woman without systemic autoimmune conditions, based on serological findings. The mesangial area showed mild proliferation and matrix thickening on light microscopy; however, electron microscopy confirmed the diagnosis of PIG. PIG has been classified into three groups based on electron microscopy findings: Invagination of only primary podocytes in type A, invagination and microstructures in type B, and only microstructures in type C[13]. Our patient was classified as type B (Figure 1B-D). After a period of treatment with corticosteroids, she developed femoral head necrosis. Thereafter, she was treated with immunosuppressants and renin-angiotensin system inhibitors, but the results were somewhat disappointing. She came to our hospital for treatment in February 2022. Along with Western medicine treatment, we began to treat her with Chinese herbal medicine treatment based on syndrome differentiation. Her 24-h proteinuria decreased from 6370 mg to 1586 mg over the course of a year.

Nonetheless, the treatment outcomes attained cannot be generalized as this is a case report. Future research in prospective studies and case series are required. accumulation of further case reports will yield the experience required to provide patients with better counseling.

#### CONCLUSION

Although PIG is a very rare glomerular disease, the combination of our case and those reported in PubMed leads us to believe that treatment with corticosteroids, immunosuppressants, and reninangiotensin system inhibitors is effective. However, some patients cannot tolerate corticosteroids and immunosuppressants. When adverse effects occur with these treatments, clinicians should avoid making negative treatment. Doctors should actively intervene and suggest treatments to patients, among which traditional Chinese medicine is an effective treatment.

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#### **FOOTNOTES**

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