Case report: Focal segmental glomerulosclerosis in a pediatric atypical progeroid syndrome

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Atypical progeroid syndrome (APS) is a rare type of progeroid syndrome mainly caused by heterozygous missense mutations in the LMNA (MIM 150330) gene. APS has heterogeneous clinical manifestations, and its kidney manifestations, particularly in children, are rarely documented. Here, we report the first pediatric case of APS with focal segmental glomerulosclerosis (FSGS). A 10-year-old boy with progeroid features was referred to the nephrology clinic because of hyperuricemia. He had dark skin, protruding eyes, and beaked nose and was very thin, suggesting lipodystrophy. He had been treated for recurrent urinary tract infection during infancy, and liver biopsy for persisting hepatitis showed steatohepatitis. He also had hypertrophic cardiomyopathy (HCMP) with mitral and tricuspid valve regurgitation. Genetic studies were performed considering his multisystem symptoms, and he was diagnosed as having APS according to exome sequencing findings (c.898G > C, p.Asparagine300History of LMNA).

During the first visit to the nephrology clinic, he had minimal proteinuria (urine protein/creatinine ratio of 0.23 mg/mg), which worsened during follow-up. In three years, his urine protein/creatinine ratio and N-acetyl-b-D-glucosaminidase/creatinine ratio increased to 1.52 and 18.7, respectively. The kidney biopsy result was consistent with findings of FSGS, showing segmental sclerosis of 1 (5%) glomerulus out of 21 glomeruli. An angiotensin receptor blocker was added to manage his proteinuria. This is the first pediatric report of FSGS in an APS patient with confirmed LMNA defect, who manifested progeroid features, lipodystrophy, HCMP with heart valve dysfunction, and steatohepatitis. Our case suggests that screening for proteinuric nephropathy is essential for managing APS patients since childhood.

KEYWORDS
focal segmental glomerular sclerosis (FSGS), atypical progeroid syndrome, lipodystrophy, LMNA, TGF - β1

Introduction

Progeroid syndromes are a group of rare genetic disorders characterized by clinical features that mimic physiologic aging. Progeroid syndromes share similar clinical features such as hair loss, short stature, skin tightness, cardiovascular diseases, and osteoporosis. However, the underlying mechanism can vary according to the causative
gene (1). The progeroid syndrome can be classified into two groups according to its molecular pathophysiology: alterations in components of the nuclear envelope and mutations in genes involved in DNA-repair pathways (1, 2). The nuclear envelope component involved in progeroid syndrome is the nuclear lamina, a thin protein meshwork between chromatin and the inner nuclear membrane (3, 4), composed of lamins. Lamins contribute to the maintenance of nuclear shape and structure, chromatin organization, and other aspects of nuclear metabolism by interacting with regulatory molecules (5–8). Lamin A and lamin C (encoded by LMNA, MIM 150330) are widely expressed in somatic cells, and diverse LMNA mutations cause various disorders, laminopathies, including diseases affecting striated and cardiac muscle, lipodystrophy syndromes such as familial partial lipodystrophy (FPLD), peripheral neuropathy, and premature aging (progeroid syndromes) (9, 10).

Among the laminopathies, FPLD encompasses abnormal fat distribution and insulin resistance disorders. FPLD type 2 (Dunnigan-type, MIM 151660) is known to be caused by LMNA mutation and is characterized by the progressive lipoatrophy of the limbs, buttocks, and trunk sparing the neck and face. Metabolic alterations are common and cardiovascular comorbidities, and hepatic steatosis is often reported (11). Hutchinson-Gilford progeria syndrome (HGPS, MIM 176670) is another form of laminopathy, an early-onset premature aging disorder. It typically presents at 1–2 years of age with severe growth retardation, lipodystrophy, and skeletal and cardiovascular features (12, 13). The average life span is 13 years (12, 14, 15). Typical HGPS is caused by de novo heterozygous silent mutation in the LMNA gene c.1924C > T (p.Gly608Gly) (16, 17). This activates a cryptic splicing site and results in abnormal splicing of the prelamin A, producing a truncated protein called progerin, which is known to cause toxic effects when accumulated (18–21). Atypical progeroid syndrome (APS) is another type of progeria from LMNA mutation. APS is very rare, and until now, 69 patients with LMNA mutation have been reported worldwide (22–33). It is characterized by not having an accumulation of the Lamin A precursors (14, 15) and is caused by heterozygous LMNA mutations other than c.1924C > T. The onset of APS symptoms is relatively late, and the life span of APS patients is generally longer than those of HGPS (12, 14, 15). Though its clinical phenotypes have not been well established, typically, affected patients have growth retardation, joint contractures, and progeroid features, including a prominent nose with beaking, partial alopecia, dental crowding, and skin anomalies (14, 15, 34). Also, most of the patients suffered from marked metabolic abnormalities such as insulin resistance, diabetes mellitus, hypertriglyceridemia, steatohepatitis, various degrees of lipodystrophy, and cardiomyopathy (15, 35, 36).

Kidney involvement as proteinuria has been seldomly described in APS (26), and the pathophysiology of kidney manifestations remains unclear. Here, we report a case of a clinical presentation of an APS patient who presented with proteinuria and pathologically confirmed focal segmental glomerulosclerosis (FSGS).

Case presentation

A 10-year-old boy visited the nephrology clinic for further evaluation of incidentally found hyperuricemia. Previously, his perinatal medical history was unremarkable, with full-term vaginal delivery and birth weight of 3.64 kg, but he had four episodes of urinary tract infection during infancy. Bilateral vesicoureteral reflux (left grade 4 and right grade 1) was identified at three months, and he was given nitrofurantoin prophylaxis until 22 months. It was discontinued as there was no further urinary tract infection for a year. After the discontinuation, his vesicourethrogram at the age of 28 months showed no reflux. He was the only child, devoid of any remarkable family history of diseases. At 2 years of age, he complained of discoloration of extremities after exposure to cold, but evaluations for autoimmune disease and arteriography of upper and lower extremities did not reveal any abnormal findings. At 7 years of age, he was noticed to have waddling gait and knee and hip flexion limitations. Two years later, marked coxa valga with relative coxa magna with a small pelvis were found on orthopedic evaluation. He visited a genetic specialist at 9 years of age. While he had normal growth with a height of 143 cm (88.6 percentile) and weight of 33 kg (47.5 percentile), with normal development, decreased subcutaneous fat tissue in his trunk and both extremities with pale and dark skin were noted. He also had acanthosis nigricans at his neck and both axillary areas, a short neck, protruding eyes, and a beaked nose. Raynaud’s phenomenon was still present. Liver function abnormality was found with elevated liver transaminases (AST 102 IU/L, normal range 15–50 IU/L; ALT 248 IU/L, normal range 5–45 IU/L) and cholesterol (LDL-cholesterol 186 mg/dl, normal range 60–140 mg/dl, triglyceride 305 mg/dl, normal range 31–108 mg/dl; HDL-cholesterol 42 mg/dl, normal range ≥40 mg/dl). His HbA1c was 6.5% (normal range: 4.0–6.4%), with a fasting blood glucose level of 83 mg/dl (normal range: 70–99 mg/dl), and increased insulin (94 μIU/ml, normal range 1.9–15.97 μIU/ml). He was referred to the endocrinologist for elevated HbA1c and dyslipidemia, however, his fasting glucose was lower than 100 mg/dl, but HbA1c and insulin were still elevated (6.1%, 55.7 μIU/ml) showing insulin resistance. Liver biopsy at 10 years showed a fatty change of hepatocytes and portal and periportal fibrosis. Mild mitral and tricuspid valve regurgitation was noted, which worsened during follow-up along with the development of hypertrophic cardiomyopathy (HCMP), requiring mechanical mitral valve replacement at 13 years of age. His diagnosis remained elusive despite
multisystem symptoms. Various studies targeting progressive storage disorders, including glycogen storage disease, were performed, with no meaningful results. Eventually, he was found to have a de novo heterozygous mutation c.898G > C (p.Asp300His) of LMNA, genetically diagnosing the patient as APS. During regular follow-up for the steatohepatitis, hyperuricemia (serum uric acid 7.6 mg/dl, normal range: 3.0–7.0 mg/dl) was noticed, and he was referred to the nephrology clinic.

His hyperuricemia was managed well with benzbromarone. Initially, minimal proteinuria [urine protein/creatinine (Cr) ratio of 0.23 mg/mg] was present, which gradually worsened during follow-up. At 13, his urine protein/Cr ratio (normal range: 0–0.2 mg/mg) and N-acetyl-b-D-glucosaminidase/Cr ratio (normal range: 0–5.6 IU/gCr) increased to 1.52 and 18.7, respectively. Urine ß2 microglobulin was 0.33 ug/ml, and serum uric acid level was 12.4 mg/dl. However, he was normotensive with a height of 160.4 cm (50–75 percentile), a weight of 37.7 kg (5–10 percentile), and a BMI of 14.65 kg/m², with normal serum albumin (4.4 g/dl, normal range: 3.3–5.2 g/dl) and serum Cr (0.71 mg/dl, normal range: 0.31–0.88 mg/dl). Doppler kidney sonography was unremarkable with normal size, parenchymal echogenicity, and intact internal perfusion. His kidney biopsy revealed FSGS, peri-hilar type, showing segmental sclerosis of 1 (5%) glomerulus out of 21 glomeruli. In microscopic findings, glomerular size was mildly increased, and there was focal mild hypercellularity involving mesangial and endothelial cells (Figure 1). Tubules were slightly atrophic, and there was no interstitial fibrosis, with mild focal infiltration of mononuclear cells. Immunofluorescence staining for immunoglobulins (IgG, IgM, and IgA), Kappa light chains, Lambda light chains, complement C3 and C1q, were all negative. Electron microscopy revealed a normal glomerular basement membrane, and effacement of the foot process was mild. There were no electron-dense deposits.

FIGURE 1
Kidney biopsy findings. (A–B) H&E and PAS staining show segmental sclerosis at the vascular pole of the glomerulus. The patient was diagnosed as having focal segmental glomerulosclerosis, perihilar variant. (A) H&E stain, × 400, (B) PAS stain, × 400. (C) Mild endocapillary hypercellularity and glomerular enlargement was observed in other glomeruli, with a maximum diameter of 302 μm. (PAS stain, × 250). (D) Electron microscopy showing mild effacement of foot processes. (EM, × 12,000).
An Angiotensin receptor blocker (Losartan) was added to manage his proteinuria (0.7 mg/kg), but it was difficult to continue due to dizziness. Since his proteinuria worsened during follow-up (urine protein/Cr ratio: 3.7 mg/mg, serum Cr 0.69 mg/dl, eGFR 98.22 ml/min/1.73 m²), losartan was restarted. His proteinuria waxed and waned with his cardiac condition, and the dosage was adjusted according to the symptoms (up to 1.2 mg/kg) (Figure 2). While taking losartan, his proteinuria did not aggravate, and his kidney function stayed stationary (urine protein/Cr ratio: 1.34 mg/mg, serum Cr 0.83 mg/dl, eGFR 84.43 ml/min/1.73 m²). During his last follow-up at 15 years of age, he did well without complaints and worked out daily. He has been prescribed benzbromarone, losartan, and sodium bicarbonate at the nephrology clinic (serum uric acid 6.8 mg/dl, serum Cr 0.79 mg/dl, cystatin C 1.74 mg/l, eGFR 84.43 ml/min/1.73 m², and urine protein/Cr ratio 0.83 mg/mg). He has taken warfarin for mitral valve replacement and a pacemaker for postoperative sinus node dysfunction. He was also taking a beta-blocker, diuretic, and amlodipine for HCMP and omega-3 for hyperlipidemia.

Discussion

This is the first report of FSGS in a pediatric APS patient with confirmed LMNA defect, who manifested progeroid features, lipodystrophy, HCMP with heart valve dysfunction, and steatohepatitis. APS is an extremely rare disease, and kidney involvement in APS is not a typical finding. Previous reports of kidney manifestation showed proteinuria, including the nephrotic range (26). While the mutation of our case, c.898G>C (p.Asp300His) in LMNA (Figure 3), was previously reported in two cases (25, 29), kidney involvement was not described. It is not a novel mutation, as the mutation was reported previously (25, 29). However, it is an extremely rare mutation, so its frequency has not been reported in genomic databases, including GNOMAD or Clinvar. One case with this variant was a 24-year-old Chinese man with multiple vascular lesions, progeroid features, hypertension, numerous intracranial calcifications, peripheral artery disease, and dyslipidemia (29). The other case is a 23-year-old woman from Myanmar with progeroid features, including short stature, thin scalp hair, absent eyebrow and eyelashes, and a beaked nose. She suffered from hypertension, secondary amenorrhea, generalized lipodystrophy, bilateral carotid artery stenosis, and left ventricular hypertrophy (25). Another missense variant of the same nucleotide locus (c.898G>A, p.Asp300Asn) was found in a 31-year-old French man with progeroid features, osteoporosis, premature atheromatosis, lipoatrophy, and cerebral ischemic disease (28). APS with cardiovascular diseases and dyslipidemia were common in all these patients, including our patient, but proteinuria was not described. Therefore, it is unclear if the FSGS of our case is a true manifestation of his APS.

However, proteinuria has been reported in several cases with LMNA mutations, and to our knowledge, nine patients were confirmed with FSGS by kidney biopsy (23, 24, 26, 37–41) (Table 1). The primary diagnosis of these cases was lipodystrophy with or without APS. Their presentations of
proteinuria were all during adulthood, and two of them with APS features required kidney replacement therapy (23, 26). Since the others with the same mutation (p.Arg349Trp) had normal to impaired kidney function, the kidney outcome seems variable (23, 26, 37, 41), and proteinuria might have been present since their childhood as in our case, but not detected earlier. Therefore, if possible, our patient also needs careful follow-up and intervention since some required dialysis in their 30s, although their variant site was different from ours.

However, the pathophysiology of FSGS in LMNA mutations remains unclear. Transforming growth factor-beta1 (TGFβ1), which is activated in lipodystrophy in laminopathies (42, 43), might play a role in the disease mechanism. It is well known that activation of TGFβ1, the central regulator of fibrotic responses (44, 45), leads to mesangial cell matrix overproduction and glomerulosclerosis (46, 47) in diseased glomeruli. Interestingly, lamin A or C was essential for inhibiting fibroblast proliferation by TGFβ1 (48). Therefore, LMNA mutation might be linked to FSGS. Also, metabolic alteration of laminopathies, including our patient, might contribute to podocyte injury leading to kidney damage (49, 50, 51). Our patient also had a history of urinary tract infection and vesicoureteral reflux; FSGS might come from reflux nephropathy. However, his kidney pathology was incompatible with typical pathologic findings of reflux nephropathy, such as interstitial scarring, tubular atrophy, or loss of nephron mass.

Regarding the treatment, as TGFβ1 plays a crucial role in pathogenesis, targeting this cytokine appears promising. However, its therapeutic application is held back because of its multifunctional and pleiotropic actions. Fresolimumab, a human monoclonal antibody neutralizing human isoforms of TGFβ, was proven ineffective in clinical trials in FSGS (52, 53). Other approaches, including decreasing the production of prelamin A or clearing progerin (54) showed limited effect (55), as APS is not associated with the accumulation of lamin A precursors (15, 56). Therefore, so far, early recognition and treatment of the manifestations is the mainstay of treatment, which makes identifying the phenotypes of the disease more crucial. Therefore, in this case, early recognition and intervention might improve kidney outcomes.

In conclusion, this is the first pediatric APS patient with FSGS. Though kidney manifestation of the disease has not been emphasized before, accompanying proteinuria and
| N | Literature, year of publication | LMNA mutation | Sex | Age at diagnosis of kidney disease (years) | Progeroid features | Lipodystrophy | Others | Kidney parameters at the time of biopsy | Remarks |
|---|--------------------------------|---------------|-----|------------------------------------------|------------------|-------------|--------|--------------------------------------|---------|
| 1 | Rankin et al. (40), 2008      | c.1930C>T (p.Arg644Cys) | F   | 32                                       | N/D              | FPLD, Dunnigan variety | DM, HTN, dyslipidemia | UPR 595 mg/mmol, sALB 27 g/L, eGFR 33 ml/min/1.73 m², sCr 1.7 mg/dl | N/D |
| 2 | Thong et al. (41), 2013       | c.1045C>T (p.Arg349Trp) | F   | 35                                       | N/D              | FPLD, Non-Dunnigan | HTN, AR, paroxysmal atrial fibrillation | UPR 3.67 g/24 h, sALB 35 g/L, eGFR 141 ml/min/1.73 m², sCr 0.52 mg/dl |         |
| 3 | Thong et al. (41), 2013       | c.1045C>T (p.Arg349Trp) | F   | 27                                       | N/D              | FPLD, Non-Dunnigan | HTN, mild MR, trivial AR, bilateral hearing loss | UPR 121 mg/mmol, sALB 38 g/L, eGFR 115 ml/min/1.73 m², sCr 0.77 mg/dl |         |
| 4 | Thong et al. (41), 2013       | c.1045C>T (p.Arg349Trp) | M   | 35                                       | N/D              | FPLD, Non-Dunnigan | Cardiomyopathy, dyslipidemia, erectile dysfunction, Rt. Hearing loss | UPR 121 mg/mmol, sALB 41 g/L, eGFR 73 ml/min/1.73 m², sCr 0.7 mg/dl |         |
| 5 | Hussain et al. (24), 2018     | c.29C>T (p.Thr10Ile)   | F   | 33                                       | N/D              | Generalized lipodystrophy | DM, dyslipidemia, AR, MR, TR, heart failure, left ventricular hypertrophy, atherosclerosis, primary amenorrhea, steatohepatitis | UPR >2 g/dl |         |
| 6 | Fountas et al. (37), 2017     | c.1045C>T (p.Arg349Trp) | F   | 27                                       | N/D              | FPLD, Non-Dunnigan | DM, dyslipidemia, myopathy, hepatic steatosis | UPR 2.2 g/24 h, sALB 4.2 g/dl, eGFR 118 ml/min/1.73 m², sCr 0.67 mg/dl | USG normal, ACEi improved kidney function, proteinuria |
| 7 | Hussain et al. (23), 2020     | c.1045C>T (p.Arg349Trp) | M   | 34                                       | beaked nose, prominent eyes, partial alopecia, skin atrophy, thin lips, small mandible | FPLD, Non-Dunnigan | HTN, dyslipidemia, moderate MR, TR, neuroendocrine tumor, sensorineural hearing loss, hepatic steatosis | UPR 9 g/24 h, eGFR 30 ml/min/1.73 m², sCr 7 mg/dl | HD, KT |
| 8 | Hussain et al. (23), 2020     | c.1045C>T (p.Arg349Trp) | F   | 38                                       | Pointed nose, thin lips | FPLD, Non-Dunnigan | DM, HTN, dyslipidemia, cardiomypathy, mild AR, MR, TR | UPR 4.5 g/24 h | dialysis |

N/D, not described; DM, diabetes mellitus; HTN, hypertension; AR, aortic regurgitation; MR, mitral regurgitation; TR, tricuspid regurgitation; UPR, urine protein; sALB, serum albumin; eGFR, estimated GFR; sCr, serum creatinine; HD, hemodialysis; KT, kidney transplantation; USG, ultrasonography.
FSGS might further deteriorate the prognosis, especially when detected belatedly after the advancement of sclerosis. Therefore, screening for proteinuria and kidney function should be considered when managing patients with APS. Further studies are needed for novel treatment strategies.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author/s.

Ethics statement

The studies involving human participants were reviewed and approved by Institutional Review Board of Seoul National University Hospital. Written informed consent to participate in this study was provided by the participants' legal guardian/next of kin.

Written informed consent was obtained from the minor(s)’ legal guardian/next of kin for the publication of any potentially identifiable images or data included in this article.

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Author contributions

SJ and HGK drafted the initial manuscript and reviewed and revised the manuscript. JSK, JMK, and YHA reviewed the manuscript for important content. SL reviewed the pathologic findings of the patient. All authors contributed to the article and approved the submitted version.

Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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