Microscopic Polyangitis in Pediatric Systemic Lupus Erythematosus: A Unique Presentation of Pulmonary - Renal Syndrome and Case Report of an Overlap Syndrome

Chen-xing Zhang
    Shanghai Children's Medical Center

Lei Yin
    Shanghai Children's Medical Center

You-ying Mao
    Shanghai Children's Medical Center

Zheng-yu Zhou
    Shanghai Children's Medical Center

He Qin
    Shanghai Children's Medical Center

Wei Zhou (zhouweiscmc@163.com)
    Shanghai Children's Medical Center Affiliated to Shanghai Jiaotong University School of Medicine

Case Report

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Microscopic Polyangitis in Pediatric Systemic Lupus Erythematosus: A Unique Presentation of Pulmonary-Renal Syndrome and Case Report of an Overlap Syndrome

Chen-xing Zhang#, Lei Yin¹, You-ying Mao¹, Zheng-yu Zhou¹, He Qin¹, Wei Zhou¹

Affiliations

1. Department of Nephrology, Shanghai Children's Medical Center, Shanghai Jiao Tong University School of Medicine, Shanghai, China

* Corresponding author: Wei Zhou

Email: zwqq121@sina.com

1678 Dongfang Road, Shanghai, China

Tel/Fax: 86-21-38625600

Abstract

Background: Secondary vasculitis is encountered in about one third of all cases of systemic lupus erythematosus (SLE). Skin is most commonly involved in lupus-related small vasculitis. Although antineutrophil cytoplasmic autoantibodies (ANCA) associated vasculitis (AAV) is relatively uncommon, it can be the most dangerous manifestation associated with high mortality. SLE and AAV are separate diseases with different pathophysiologies and an overlap syndrome has only been reported few times in previous literature.

Case Presentation: We present a unique case of a pediatric patient of pulmonary-renal syndrome, presenting with pulmonary alveolar hemorrhage and rapidly progressive glomerulonephritis. Serological and biopsy findings were suggestive of SLE and AAV occurring simultaneously. Renal biopsy demonstrated necrotizing and crescentic glomerulonephritis, superimposed on diffuse segmental proliferative lupus glomerulonephritis class IV.

Conclusions: Presentations of autoimmune diseases and vasculitis can be multisystemic. Considering overlap syndromes, especially in patients with underlying connective tissue disease or systemic vasculitis, is vital for prompt therapy and prevention of morbidity in this population.
Keywords: Systemic lupus erythematosus (SLE), Antineutrophil cytoplasmic autoantibodies associated vasculitis (AAV), Overlap syndromes

1. Background

Systemic Lupus Erythematosus (SLE) and ANCA associated vasculitis (AAV) are separate diseases with different pathophysiology. While both affect the kidneys and other systemic involvements, SLE can lead to an immune complex glomerulonephritis, while AAV is characterized by pauci-immune necrotizing and crescentic glomerulonephritis [1, 2]. Pulmonary-renal syndrome, for instance, diffuse alveolar hemorrhage (DAH) and rapidly progressive glomerulonephritis can occur in both of the diseases [3, 4]. The occurrence of both SLE and AAV simultaneously has only been depicted few times in previous literature, especially in the pediatric population. Serological tests are extremely helpful when making a diagnosis, but ultimately it must be based on the results of renal biopsy. Treatment with corticosteroids, immunosuppressive agents, and, in severe cases, plasmapheresis has considerably improved the prognosis [5, 6]. Here we have reported the case of a 14-year-old girl with an overlap syndrome of SLE and AAV and reviewed her kidney biopsy finding, the differential diagnosis, diagnosis as well as the management of the disease.

2. Case Presentation

A 14-year-old girl was admitted to our hospital because of arthralgias, rash and fever. The patient had been well until one month before admission, when arthralgias affecting knees and ankles, as well as rash in lower extremities developed. One day before admission, she had a fever with peak temperature of 38.6°C. The patient appeared ill, with pale face and lip. She had known allergy of penicillin, with no history of recent medication, travel, exposure to sick persons, blood transfusions, major illness or previous surgery. She had no previous personal or family history of autoimmune disease, except that her mother suffered from hyperthyroidism.

On physical examination, the temperature was 38.7°C, the blood pressure 108/60
mm Hg, the pulse 90 beats per minute, the respiratory rate 26 breaths per minute, and the oxygen saturation 100% while she was breathing ambient air. The height was 158 cm, the weight 45 kg, and the body-mass index (the weight in kilograms divided by the square of the height in meters) 18.02. There was dependent edema and dispersive purpura like rash over both of the lower extremeties. Auscultation of the chest revealed wet rales in the right lung. Review of other systems was negative. Initial laboratory examinations revealed anemia with hemoglobin of 8.2 g/dL, renal insufficiency with a creatinine of 120 μmol/liter, reduced albumin level of 28.3 g/liter as well as elevated inflammatory markers. Urinalysis revealed proteinuria and hematuria.

A cough developed during the next day. The cough was intermittently productive of thick white sputum that occasionally contained a small amount of blood. The patient also reported fatigue, episodes of shortness of breath and transient chest pain, without headaches, abdomen pain, visual loss, dry eyes or dry mouth. Thus high-resolution computed tomography (HRCT) of the chest was performed. CT of the chest showed multiple patches with increases in density and peripheral opacities (Figure 1A). Axillary lymphadenopathy was also identified. Ceftazidime and Azithromycin was administered intravenously. Supplemental oxygen was also administered through a nasal cannula at a rate of two liters per minute.

Subsequent blood work revealed low C3 (0.87g/liter), ANA (1:100), p-ANCA, anti-myeloperoxidase (MPO), anti-SSA auto-antibody as well as positive Coombs test. Moderate schizocyte were identified in peripheral blood smear test. Anti-glomerular basement membrane (GBM) antibody, and antibody to proteinase 3 (PR-3) were negative. HIV, HBV and HCV were also tested to be negative. Comprehensive laboratory test results are shown in Table 1. Ultrasonography of the abdomen revealed thickening of liver parenchyma. Ultrasonography of the urinary system revealed diffuse alterations in bilateral renal cortex. An electrocardiogram (ECG) showed sinus arrhythmia and nonspecific T-wave abnormalities. Cardiac ultrasonography revealed normal global cardiac function and right-ventricular size, no evidence of a pericardial effusion.
The patient met 4 of the 11 lupus criteria based on the 1997 American College of Rheumatology criteria: hematological disorder (hemolytic anemia), renal disorder (proteinuria of 3.54 grams in 24 hours), arthritis and positive ANA. She also met 5 of the 17 lupus criteria according to the 2012 Systemic Lupus International Collaborating Clinics (SLICC) classification criteria: arthritis, hemolytic anemia, renal disorder (proteinuria of >500 mg per 24 hours), positive ANA, and low complement (low C3), which includes three clinical and two immunological items. Considering involvement of renal and pulmonary system is due to the occurrence of primary disease of lupus, methylprednisolone (500mg, 1 g and 1g daily for 3 days) was administered.

Fever dropped to normal and was stable in the third hospital day. On the fourth hospital day, the patient appeared to be fatigued, the oxygen saturation fell to 93% with accelerated breathing (30 breaths per minute). Because of persistent respiratory symptoms, repeated chest CT scanning was performed. There was massive infiltrates in right lobes, accompanied by pleural effusions (Fig. 1B). Oxygen (5 liters per minute) was administered by face mask and she was transferred to the pediatric intensive care unit (PICU) immediately due to the concurrent deteriorating renal function, implied by the increasing serum creatinine levels. On examination, the patient appeared chronically ill. The temperature was 36.8°C, the pulse 70 beats per minute, the blood pressure 133/88 mm Hg, the respiratory rate 19 breaths per minute, and the oxygen saturation 96% while she was receiving supplemental oxygen through a face mask at a rate of 5 liters per minute. Auscultation of the lungs revealed scattered bilateral crackles. Treatment with ceftazidime and azithromycin was discontinued and treatment with meropenem, pulse cyclophosphamide, and administration of methylprednisolone was initiated. In addition, a femoral vein catheter was placed, plasmapheresis was begun and fentanyl was administered. After two sessions of plasmapheresis, improvement in renal function and resolution of hemoptysis was observed. The patient only underwent 4 units of packed red blood cell transfusion in the previous two days during the PICU stay. She had a total of 7 sessions of plasmapheresis.
High-resolution computed tomography (HRCT) of bilateral alveolar infiltrates is suggestive of diffuse alveolar hemorrhage. HRCT of the chest showed multiple patches with increases in density and peripheral opacities in the 2nd hospital day (Fig.1A). There was massive infiltrates in right lobes, accompanied by pleural effusions in the fourth hospital day, which demonstrated an obvious radiological progression (Fig.1B). There was neither inflammatory exudation nor pleural effusion, as demonstrated by chest CT re-examination in the 4th month of the follow up (Fig. 1C).

Table 1: Laboratory test results upon admission

| Variable                              | Reference Range | On Admission | The 4th Day |
|---------------------------------------|-----------------|--------------|-------------|
| White cell count (per mm³)            | 4000-15,000     | 7250         | 13,150      |
| Differential count (%)                |                 |              |             |
| Neutrophils                           | 50-70           | 75.5         | 85.5        |
| Lymphocytes                           | 20-40           | 17.5         | 11.2        |
| Eosinophils                           | 0.5-5           | 0.4          | 0           |
| Platelet count (per mm³)              | 100,000-550,000 | 193,000      | 253,000     |
| Hematocrit (%)                        | 33-47.3         | 25           | 19.6        |
| Hemoglobin (g/dl)                     | 11-16           | 8.2          | 6.4         |
| Erythrocyte count (per mm³)           | 3700,000-5800,000 | 2740,000     | 2220,000    |
| D-Dimer (mg/liter)                    | < 0.3           | 1.81         | 1.55        |
| Activated partial-thromboplastin time (sec) | 28-45       | 34.5         | 28.9        |
| Prothrombin time (sec)                | 9-14            | 10.9         | 10.7        |
| International normalized ratio for prothrombin time | 0.9-1.4 | 1.01 | 0.99 |
| Fibrinogen (g/liter)                  | 1.5-4.0         | 4.3          | 2.98        |
| Sodium (mmol/liter)                   | 137-145         | 139          | 133         |
| Potassium (mmol/liter)                | 3.5-5.1         | 4.44         | 4.31        |
| Chloride (mmol/liter)                 | 98-107          | 109          | 105         |
| Carbon dioxide combining power (mmol/liter) | 22-30     | 18           | 18          |
| Urea nitrogen (mmol/L)                | 2.5-6.1         | 10.1         | 15.7        |
| Creatinine (µmol/liter)               | 9-88            | 120          | 128         |
| Endogenous creatinine clearance       | 75-115          | 38.32        | /           |
| Parameter                                      | Lower Limit | Upper Limit | Value 1 | Value 2 | Value 3 |
|-----------------------------------------------|-------------|-------------|---------|---------|---------|
| Protein (g/liter)                             |             |             | Total   | Albumin | Globulin |
|                                               | 63-82       |             | 63.9    | 28.3    | 35.6    |
|                                               |             |             |         |         | 27.2    |
| Calcium (mmol/L)                              | 2.23-2.80   |             | 2.01    |         | 1.95    |
| Phosphorus (mmol/L)                           | 1.45-2.10   |             | 1.56    |         | 1.94    |
| Creatine kinase (U/liter)                     | 30-135      |             | 63      |         |         |
| Creatine kinase MB isoenzymes (μg/liter)      | < 3.7       |             | < 0.5   |         |         |
| Troponin I (μg/liter)                         | < 0.02      |             | 0.02    |         | < 0.02  |
| Ferritin (ng/ml)                              | 15-160      |             |         |         | 125.3   |
| C reactive protein (mg/L)                     | < 8         |             | 16      |         | 3       |
| Erythrocyte sedimentation rate (mm/hr)        | 0-20        |             | 92      |         |         |
| Antineutrophil cytoplasmic autoantibody       |             |             |         |         |         |
| Cytoplasmic (Ab to proteinase 3)              | Negative    |             | Negative|         |         |
| Perinuclear (Ab to myeloperoxidase)           | Negative    |             | Positive|         |         |
| Antinuclear antibody                          | Negative    |             | 1:100   |         |         |
| Direct (U/ml)                                 |             |             |         |         |         |
| Anti-RNP antibody (U/ml)                      | Negative    |             | Negative|         |         |
| Anti-Smith antibody (U/ml)                    | Negative    |             | Negative|         |         |
| Anti–SS-A antibody (U/ml)                     | Negative    |             | Positive|         |         |
| Anti–SS-B antibody (U/ml)                     | Negative    |             | Negative|         |         |
| Anti–double-stranded DNA                      | Negative    |             | Negative|         |         |
| Anticardiolipin antibody (U/ml)               | Negative    |             | Negative|         |         |
| C3                                            | 0.9-1.8     |             | 0.87    |         |         |
| C4                                            | 0.1-0.4     |             | 0.15    |         |         |
| Blood gases (venous) while breathing high-flow oxygen by oxygen mask | | | | | |
| pH                                            | 7.35-7.45   |             | 7.455   |         |         |
| Partial pressure of carbon dioxide (mm Hg)    | 32-45       |             |         |         | 26.7    |
| Partial pressure of oxygen (mm Hg)            | 83-110      |             |         |         | 58.9    |
| Base excess (mmol/liter)                      | -3 - +3     |             |         |         | -4.7    |
| Urine                                         |             |             |         |         |         |
| pH                                            | 5-7         |             | 6.0     |         | 5.0     |
| Color                                         | Yellow      |             | Yellow  |         | Yellow  |
| Turbidity                                     | Clear       |             | Clear   |         | Clear   |
| Screening-dipstick test                       |             |             |         |         |         |
| Glucose                                       | Negative    |             | Negative|         | 1+      |
| Bilirubin                                     | Negative    |             | Negative|         | Negative|
Ketones  Negative  Negative  1+
Occult blood  Negative  4+  4+
Protein  Negative  3+  3+
Urobilinogen  Negative  Negative  Negative
Nitrites  Negative  Negative  Negative
White cells  Negative  Negative  Negative

| Sediment | Red cells (per high-power field) | 0-3 | Full field of vision | Full field of vision |
|----------|--------------------------------|-----|---------------------|---------------------|
|          | White cells (per high-power field) | 0-5 | 6-9 | 0 |
|          | Hyaline casts (per low-power field) | 0-2 | 0 | 0 |
|          | Granular casts (per low-power field) | 0 | 0 | 0 |
|          | Muddy casts (per low-power field) | 0-2 | 0 | 0 |
| 24 hour urine protein (mg/24h) | 40-225 | 3540 | / |

| Blood culture | Negative | Negative | / |
| --------------|---------|---------|---|
| Epstein-Barr DNA | < 5*10^3 copies/ml | < 5*10^3 copies/ml | / |
| Cytomegalovirus DNA | < 500 copies/ml | < 500 copies/ml | / |
| Sputum hemosiderin | Negative | / | Negative |

| Coombs test | Direct Coombs test | Negative | Positive | / |
|            | Indirect Coombs test | Negative | Negative | / |

After 7 sessions of plasmapheresis, there was an improvement in renal function. The patient’s condition is relatively stable, without requirements for oxygen. Then renal biopsy was performed. Examination of formalin-fixed tissue by means of light microscopy showed proliferative-appearing glomeruli with focal formation of fibrous and fibrocellular crescents. The GBM was not thickened. Some glomeruli were globally or segmentally sclerosed, indicating disease chronicity. The renal tubules and the interstitium were also abnormal, with mild focal tubular atrophy and mild interstitial focal fibrous hyperplasia and an inflammatory infiltrate composed primarily of monocytes and lymphocytes. Immunofluorescence staining of frozen tissue showed staining with IgG (+~++), IgM (+~++), IgA (+~++), C3 (+), C1q (+~++), Fn (++).

The classification of lupus nephritis is based on the glomeruli alone. In this case, more than half the glomeruli were diseased (indicating diffuse glomerulonephritis) and there was segmental involvement, so the disease was classified as a diffuse segmental proliferative lupus glomerulonephritis, with crescent formation and highly
suspected of focus of necrosis, referred to as class IV-S (A/C) (ISN/RPS2003). The activity score is 9 and the chronicity index is 5.

Since ANCA-associated vasculitis is the most common cause of the pulmonary-renal syndrome, the diagnosis must be considered in this patient. Of patients with granulomatosis with polyangiitis (GPA) and microscopic polyangiitis (MPA), 85 to 90% have ANCA, namely anti-myeloperoxidase (MPO) or PR3 antibodies [7, 8]. Systemic features, such as fever, arthralgias and skin lesions (palpable purpura) are also common and were present in this patient. The pulmonary lesion in ANCA-associated vasculitis is DAH or focal inflammatory vasculitis. In GPA, there may also be lesions in the upper airways (tracheitis, sinusitis and otitis), necrotic pulmonary lesions, and hilar lymphadenopathy[5]. Inflammatory markers such as erythrocyte sedimentation rate and C-reactive protein levels are also elevated. Iron-deficiency anemia may be present if the pulmonary bleeding is severe and chronic. The absence of asthma, sinus disease, and lung nodules rules out GPA or eosinophilic granulomatosis with polyangiitis (EGPA). Histopathological confirmation of the diagnosis of AAV is highly desirable and tissue biopsy can be important for the diagnosis and staging of AAV. In this patient, the renal biopsy demonstrated crescent formation and focus of necrosis, which was accordant with the pathogenic features of MPA. In addition, the case met the 2017 ACR/EULAR classification criteria of MPA.

SLE with both renal and pulmonary involvement should be given serious consideration in this patient, even though DAH occurs in only 0.5 to 5.4% of patients with SLE [9-12]. DAH is usually associated with lupus nephritis and antiphospholipid autoantibodies in SLE patients [13-15]. The exact cause of DAH pathology remains unknown, the lung lesions can be bland alveolar hemorrhage or a localized immune complex-induced pulmonary capillaritis, contributing to damage to basement membranes and leakage of erythrocytes into the alveolar space [16]. This patient had evidence of both hemorrhage and inflammation. Other features that were compatible with SLE were hypocomplementemia, a positive direct Coombs test and ANA, fever, rash and arthralgia in this patient. SLE patients with renal and pulmonary involvement can have anti-GBM disease or disease associated with ANCA (almost always anti-
MPO). These diseases can be diagnosed only through appropriate serologic tests and immunopathological studies of biopsied specimens from a kidney or possibly a lung.

The patient met 4 of the 11 lupus criteria based on the 1997 American College of Rheumatology criteria: hematological disorder (hemolytic anemia), renal disorder (proteinuria of 3.54 grams in 24 hours), arthritis and positive ANA [17]. She also met 5 of the 17 lupus criteria according to the 2012 Systemic Lupus International Collaborating Clinics (SLICC) classification criteria: arthritis, hemolytic anemia, renal disorder (proteinuria of >500 mg per 24 hours), positive ANA, and low complement (low C3), which includes three clinical and two immunological items [18]. We concluded that alveolar hemorrhage and nephritis is due to SLE or AAV. Therefore, the diagnosis of the patient is SLE with a diffuse segmental proliferative lupus nephritis, with crescent formation and highly suspected of focus of necrosis, referred to as class IV-S (A/C) (ISN/RPS2003) as well as MPA.

This patient presented with arthralgias, rash, fever, hematuria, proteinuria, hypocomplementemia and progressive renal insufficiency. Although bronchoscopy was not performed, clinical manifestations and radiological features indicated alveolar hemorrhage and inflammation. Severe anemia suggested acute blood loss into the lung parenchyma, combined with shortened red-cell survival due to hemolytic anemia. The involvement of mucocutaneous, musculoskeletal, respiratory and urinary system suggested a systemic illness affecting multiple systems. The differential diagnosis includes systemic multisystem illness: infectious diseases, malignant tumors, immune-mediated disorders, and other disorders, as depicted in Table 2. Yet the most likely diagnosis was either SLE or microscopic polyarteritis (MPA).

| Table 2: Differential diagnosis |
|-------------------------------|
| **Infectious diseases**       |
| Bacteria Infection           |
| Bacterial endocarditis        |
| Legionnaires disease          |
| Tuberculosis                  |
| Fungus Infection             |
| Histoplasmosis                |

Therefore, the diagnosis of the patient is SLE with a diffuse segmental proliferative lupus nephritis, with crescent formation and highly suspected of focus of necrosis, referred to as class IV-S (A/C) (ISN/RPS2003) as well as MPA.
Aspergillosis

Virus Infection
HIV, EBV, CMV, hepatitis B/C, parvovirus

Malignant tumors
Lymphoma

Immune-mediated disorders
Systemic lupus erythematosus (with or without ANCA, anti-GBM antibody, or anti-phospholipid autoantibody)
ANCA associated vasculitis
  - Microscopic polyangiitis (MPA)
  - Granulomatosis with polyangiitis (GPA)
  - Eosinophilic granulomatosis with polyangiitis (EGPA)
  - Mixed cryoglobulinaemia
Anti-GBM disease
Combined anti-GBM disease and ANCA associated vasculitis
Mixed connective tissue disease
Progressive systemic sclerosis
Idiopathic pulmonary hemosiderosis (with delayed anti-GBM disease)
Hypersensitivity to drugs

Other disorders
Hemolytic–uremic syndrome (with glomerular thrombotic microangiopathy)
Membranoproliferative glomerulonephritis
Focal and segmental glomerulosclerosis
Acute tubular necrosis
Silica dust exposure (with anti-GBM antibody or ANCA)
Multiple pulmonary emboli and a coexistent renal disease (membranous glomerulopathy)
Uremic pneumonitis

The fever, cough, hemoptysis, and new pleural effusion on repeated HRCT examination may reflect an underlying respiratory infection which include those caused by bacteria such as bacterial endocarditis, legionnaires disease, mycobacterium tuberculosis infection, as well as mycoplasma pneumoniae (MP) and fungal infection. Negative results of blood culture and relevant etiology (negative TSPOT and MP mRNA, normal blood levels of 1,3-β-D-glucan) excluded these possibilities. Other viral infections were ruled out by the absence of typical clinical features and negative serologic tests including viruses such as HIV, EBV, CMV, hepatitis B and C.

Malignant tumors that can present as systemic illness, which could be considered
in this case, include non-Hodgkin’s lymphoma and plasma-cell myeloma. The presence of constitutional symptoms, lymphadenopathy, anemia, and an elevation in the erythrocyte sedimentation rate (ESR) all raised concern about the possibility of a lymphoma. However, rash is not a common manifestation of systemic lymphoma. Proteinuria, hematuria, renal insufficiency with an elevated ESR could be the result of a monoclonal gammopathy, as is seen in plasma-cell myeloma [19]. Although lymphoma remained a serious consideration, we believe that other causes are more likely due to presence of autoantibodies in this patient.

The multisystem involvement, progressive nature of the illness and elevated erythrocyte sedimentation rate, without clinical evidence of infectious or neoplastic disease, leads us to consider an immune-inflammatory disorder caused by exogenous noninfectious agents (a systemic reaction to a drug or chemical), a systemic autoimmune disease (systemic lupus erythematosus or its variants), or a systemic vasculitis. Although the patient had known allergy of penicillin, she was not taking any medication and exposed to any chemical toxins during periods before admission. Therefore, the process of elimination leaves us with rheumatic and autoimmune diseases.

Anti-GBM disease primarily affects the pulmonary and renal system and causes symptoms like hemoptysis, hematuria and proteinuria [20, 21]. Although anti-GBM disease is one of the less common causes, it must be considered in this case because of the potential implications for management [22]. Anti-GBM disease is commonly associated with lung hemorrhage. Renal failure is common and usually rapidly progressive, as was suspected in this patient. In anti-GBM disease, the renal lesion is most often a necrotizing and crescentic glomerulonephritis. Systemic symptoms are usually absent and serum complement levels are typically normal, which was not the case in this patient. A useful clinical approach to differential diagnosis involves serologic categorization and renal biopsy. Apart from detection of serum anti-GBM antibody, inflammation of the basement membrane and a liner pattern along the basement membrane is usually observed under a microscope [20, 21]. Approximately 20 to 30% of patients may also be shown to have ANCA [23, 24]. Presence of
systemic symptoms, low serum complement C3 level, and absence of serum anti-
GBM antibody would argue against the diagnosis of anti-GBM disease.

Since the presence of active lupus nephritis (LN) and MPA was confirmed, the patient was then given the second round of methylprednisolone pulse therapy (500mg daily) for 3 days as well as cyclophosphamide intravenously (400mg daily for 2 days) every 4 weeks. The patient did follow up with the nephrology and rheumatology departement and therefore received the additional planned cyclophosphamide infusions while taking oral hydroxychloroquine and prednisone. She has completed the sixth round of cyclophosphamide infusions and was on maintenance therapy with mycophenolatemofetil (MMF, 0.75g+0.5g, bid) up to now. She had a favorable renal response, with drop of proteinuria from initial 6781 mg/24h to present 1089 mg/24h. Her renal function also continued to improve, and 4 months after discharge, the creatinine was 6.67 mg/L (59 μmol/L). Her serum albumin (36 g/L) and hemoglobin (115 g/L) both recovered and has remained stable. In regard to her pulmonary status, there was neither inflammatory exudation nor pleural effusion, as demonstrated by chest CT re-examination (Fig.1C). She adhered well to the medications and the prednisone dose was tapered gradually from 25mg/bid to 15mg/qd.

3. Discussion

The involvement of mucocutaneous, musculoskeletal, respiratory and urinary system suggested a systemic illness affecting multiple systems. The differential diagnosis includes systemic multisystem illness, such as infectious diseases, malignant tumors, immune-mediated disorders, and other disorders. Yet the most likely diagnosis was either SLE or MPA.

SLE is a typical autoimmune disease characterized by chronic inflammation, pathogenic auto-antibodies, and multi-systemic involvement, with a wide range of clinical presentations including rashes, arthralgias, pleuritis, or renal disorders [2]. Our patient tested positive for ANA, p-ANCA, anti-SSA and had low C3 with elevated inflammatory markers and biopsy proven Class IV lupus nephritis. Our patient met both lupus criteria based on the 1997 American College of Rheumatology
criteria and the 2012 Systemic Lupus International Collaborating Clinics (SLICC) classification criteria. Thus SLE was primarily considered when we initially reviewed the case.

While there are no specific classification criteria for pediatric MPA, the Chapel Hill nomenclature provides a useful definition for clinical studies in children[25]. The additional clinical features of pulmonary-renal syndrome and laboratory findings in this case also pointed strongly toward the underlying MPA. The diffuse changes in bilateral renal cortex implied that the disease may be chronic, as evidenced by ultrasound of urinary system. Since abnormal kidney echotexture and progressive renal insufficiency are ominous signs, a renal-biopsy specimen would be crucial in the determination of the diagnosis, the prognosis, and the best course of therapy.

The renal biopsy revealed immune-complex deposits consistent with WHO class IV diffuse segmental proliferative lupus glomerulonephritis [26]. Seventy percent of glomeruli (7/10) was circumferential fibrous crescents and fibrocellular crescents in our patient, the proportion of which is much higher than that observed in the general SLE population. Nevertheless, necrotizing and crescentic glomerulonephritis (including cellular and fibrocellular and fibrous crescents), one of the features and usually present in renal biopsy of AAV patients, was also observed in our patient. Based on the Chapel Hill consensus conference of 2012, this patient’s AAV is most consistent with the MPA subtype due to the absence of granulomatous inflammation or eosinophils [25]. In addition, The draft classification criteria of AAV was put forward by ACR/EULAR in 2017, which is applicable to patients with small or medium vasculitis. Although it applies “subtraction method” to rule out other small vasculitis and remains to be validated, it is useful and instructive in clinical practice because it entails multiple indicators such as clinical manifestations as well as imaging, pathological and serological features. Our case met the 2017 ACR/EULAR classification criteria of MPA. The immune deposits in the biopsy of our patient implied lupus nephritis diagnosis but did not exclude the possibility of MPA, which is classically considered to be pauci-immune, with focus of necrosis. SLE and AAV are separate diseases with different pathophysiology and an overlap syndrome has only
been reported few times in the literature, especially in the pediatric population. We present a unique clinical case of a 14 year old girl, without a previous personal or family history of systemic autoimmune disease, with serological and biopsy findings of both diseases occurring simultaneously.

ANCA, typically p-ANCA, can be detected in patients with lupus nephritis, ranging from 3.69% to 16.9%. Although this finding is rarely of clinical significance since most patients do not get concomitant vasculitis. This confounding effect can be in part due to a cross reaction with ANA. Previous studies have demonstrated that classes IV and III were predominant in ANCA-positive LN. Furthermore, MPO-ANCA-positive LN patients were more likely to have lower titers of ANA, higher serum concentrations of complement, and worse baseline renal function, all of which were consistent in our case [27-31]. Distinguishing AAV from SLE vasculitis may be difficult, as SLE vasculitis can manifest in small vessels and occurs in up to one third of patients with underlying SLE [32]. Therefore, renal histopathological findings are paramount in making the final diagnosis.

With only few cases of overlap syndrome of SLE and AAV in previous literature, this case report provides further insight into this unique presentation with pulmonary-renal syndrome. Although AAV is rare in childhood, renal involvement is common and occurs in 60%-90% of reported pediatric cases. The most severe form of renal involvement in AAV is the rapidly progressive ANCA-associated necrotizing crescentic GN. Among children, 30%-40% will progress to CKD and about 10% to ESRD by adulthood [33-35]. Prognostic factors of SLE and AAV are not invariably uniform. In 2010, an international working group of renal pathologists developed a new histopathologic classification system for ANCA-associated glomerulonephritis (focal, crescentic, mixed, and sclerotic) [36]. Noone et al has analyzed its association with renal outcomes in the largest cohort of pediatric patients (40 patients) so far. There is a significantly poor renal survival in the crescentic/mixed group and sclerotic category, with an adjusted hazard ratio of 3.14 and 23.6 respectively, compared with the focal category [37]. Negative prognostic factors for lupus nephritis was associated with an initial Class IV histology (relative risk, 1.78), hypertension at presentation
A French nationwide survey was conducted to identify cases of SLE/AAV overlap syndrome and only 8 cases was identified, among which only 2 cases occur concomitantly. A severe clinical presentation (rapidly progressive glomerulonephritis and frequent pulmonary involvement) is an almost universal feature in SLE/AAV overlap syndrome, as exhibited in our patient. In addition, a majority have both ANA and anti-MPO antibodies [39].

Considering the lack of evidence in children and the high level of evidence in adult studies for AAV, the EULAR recommendations on adult-onset AAV can be applied in pediatric AAV patients. An approach for the treatment of crescentic glomerulonephritis/rapidly progressive glomerulonephritis was described in 2018 European consensus-based recommendations for the diagnosis and treatment of rare paediatric vasculitides. Treatment is divided into two phases: a period of intense treatment (induction therapy), followed by a period of less intense maintenance therapy. Induction regimens typically include high dose corticosteroids, intravenous cyclophosphamide, and therapeutic plasmapheresis. Recommended first-line maintenance options are azathioprine, methotrexate and mycophenolate mofelil[5]. Early diagnosis and prompt commencement of therapy is of vital important to maximize chances of renal recovery.

Our patient was initially treated with pulse doses of intravenous steroids, cyclophosphamide and plasmapheresis. The patient is following up with the nephrology and rheumatology department and has received the additional planned cyclophosphamide infusions while taking oral hydroxychloroquine and prednisone. She has completed the sixth round of cyclophosphamide infusions and was on maintenance therapy with MMF up to now. The serum creatinine level and 24-hour proteinuria declined continuously, indicating a dramatic improved renal function as well as a favourable renal response. Besides, dramatic improvement was observed in pulmonary imaging and the hemoglobin level has restored and remained stable in the normal range. Thus the prednisone dose was tapered gradually, with stable blood
pressure. We will continue to have a follow up with the patient. Further investigations are needed to elucidate the possible shared immune pathogenesis involved in co-existence of SLE and AAV in these patients to seek for new therapeutic options.

4. Conclusion

For patients who present with acute pulmonary-renal syndrome with atypical lupus serology (relatively lower titers of ANA, serum complement of threshold value), prominently elevated proportion of crescents in renal biopsy, along with ANCA positivity, clinicians should be aware of concurrent AAV. Proper vigilance and timely diagnosis of AAV in a lupus patient will lead to early initiation of proper disease-oriented treatment.

Abbreviations

SLE: Systemic lupus erythematosus
ANCA: Antineutrophil cytoplasmic autoantibodies
AAV: Antineutrophil cytoplasmic autoantibodies associated vasculitis
DAH: Diffuse alveolar hemorrhage
HRCT: High-resolution computed tomography
MPO: Anti-myeloperoxidase
GBM: Glomerular basement membrane
PR-3: Proteinase 3
ECG: Electrocardiogram
SLICC: Systemic Lupus International Collaborating Clinics
PICU: Pediatric intensive care unit
GPA: Granulomatosis with polyangiitis
MPA: Microscopic polyangiitis
EGPA: Eosinophilic granulomatosis with polyangiitis
MP: Mycoplasma pneumoniae
ESR: Erythrocyte sedimentation rate
LN: lupus nephritis
MMF: Mycophenolate mofetil

Availability of Data and Material
The data and material used to support the findings of this study are available from the corresponding author upon request.

Ethical Approval and Consent to Participate
This study was approved by the Ethics Committee of Shanghai Children's Medical Center, Shanghai Jiao Tong University School of Medicine.

Consent for Publication
All authors have finally approved the submitted version to be published.

Competing Interests
The authors declared no potential conflicts of interest with respect to the authorship and publication of this article.

Authors’ Contributions
The authors as listed on the title page of the manuscript have all made substantial contributions which qualifies them as authors.

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High-resolution computed tomography (HRCT) of bilateral alveolar infiltrates is suggestive of diffuse alveolar hemorrhage. HRCT of the chest showed multiple patches with increases in density and peripheral opacities in the 2nd hospital day (Fig. 1A). There was massive infiltrates in right lobes, accompanied by pleural effusions in the fourth hospital day, which demonstrated an obvious radiological progression (Fig. 1B). There was neither inflammatory exudation nor pleural effusion, as demonstrated by chest CT re-examination in the 4th month of the follow up (Fig. 1C).

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