Abstract. Mycoplasma is one of the most common pathogens causing community-acquired pneumonia in pediatric patients. In recent years, the number of refractory or severe cases with drug resistance has been gradually increasing and cases that developed embolism after Mycoplasma pneumoniae (M. pneumoniae) infection have been reported. The present study retrospectively analyzed the clinical features, diagnosis and treatment of M. pneumoniae pneumonia (MPP) combined with pulmonary embolism (PE) in a series of 7 cases encountered between January 1st, 2016 to August 1st, 2019 at the Department of Pediatric Intensive Care Unit of The First Hospital of Jilin University (Changchun, China). Combined with relevant Chinese and international studies published during the last two decades, a comprehensive analysis was performed. All of the pediatric patients of the present study had fever, cough and dyspnea respiratory symptoms at onset and the disease progressed rapidly. Thereafter, PE was confirmed by a series of examinations. Pulmonary CT indicated patchy inflammations and significantly elevated D-dimer levels, accompanied by positive anticardiolipin antibodies. Furthermore, a filling defect in the pulmonary artery branch was observed on CT pulmonary angiography (CTPA) examination. In 2 cases, the condition was improved with anti-infection and anticoagulation treatment with low-molecular-weight heparin and warfarin, respectively, and the pulmonary embolism disappeared after 3-4 months. A total of 5 cases, who were not responsive to the drug treatment, underwent surgical resection. During the operation, the local tissues were determined to be infarcted and the pathological diagnosis was consistent with pulmonary infarction. Among the 5 cases, 2 died of Acute Respiratory Distress Syndrome at 3-8 days after the operation. The remaining patients underwent 6-12 months of follow-up and respiratory rehabilitation and their quality of life is now good. In conclusion, compared with healthy individuals, pediatric patients with critical MPP have an elevated risk of embolism. It is necessary to be vigilant regarding whether MMP is combined with PE and perform timely CTPA examination. Early detection, early treatment and surgical intervention (if necessary) may significantly reduce the risk of mortality and disability.

Introduction

Mycoplasma pneumoniae (M. pneumoniae) is the most common pathogen in pediatric patients with community-acquired pneumonia (1,2) which breaks out every 3-5 years (3). In recent years, the number of refractory or severe cases with drug resistance has been gradually increasing (4-6). In addition to pulmonary inflammation as the most common manifestation, M. pneumoniae infection may also cause damage to multiple systems and organs (7-9). In the last 20 years, ~60 cases of M. pneumoniae infection with thrombotic disease in pediatric patients have been reported worldwide (10-15). The present study reported on a series of 7 cases of M. pneumoniae pneumonia (MPP) accompanied by pulmonary embolism (PE) encountered January 1st, 2016 to August 1st, 2019 at the Department of Pediatric Intensive Care Unit of The First Hospital of Jilin University (Changchun, China), and the clinical data of these cases were reviewed. The present study aimed to improve the understanding of clinicians regarding the laboratory examinations, diagnosis and treatments for pediatric patients with MPP-associated PE.

Case report

Cases. MPP-associated PE was confirmed in 7 cases by radiological examination combined with serological tests and the corresponding clinical data were collected and summarized. The present study had been approved by the Ethics Committee of the First Hospital of Jilin University (Changchun, China; approval no. 2019-253).
Baseline data. The seven cases were aged between 6 and 11 years (median, 8.0 years) with a male/female ratio of 4:3, as presented in Table I. All patients were otherwise physically healthy. Patients with a family history of thrombophilia and a history of allergy were excluded.

Clinical symptoms and physical signs. All of the patients had a cough and fever as the initial symptoms, typically irritable cough with viscous sputum, and remittent fever. Among them, 5 cases developed dyspnea within 2 to 6 days and were hospitalized on day 3-12 from onset. These cases developed PE on day 10-14 and their condition soon deteriorated. Older pediatric patients complained of chest pain, chest tightness or sudden dyspnea; younger patients were unable to describe their symptoms, but physical examinations revealed spiritlessness, aggravating dyspnea, flapping of nasal wings, reduced respiratory movement amplitude on the affected side and weak breath sounds. One case was combined with swelling in the right lower limb.

Results of auxiliary examinations

Laboratory tests. The serum biochemistry results of the cases are presented in Table II. The serum *M. pneumoniae* antibody titers (Particle Agglutination assay; SERODIA-MYCOII; Fujirebio) were increased by varying degrees [from negative titer at presentation to positive titer at the 2nd examination (n=1) or >4-fold increased antibody titers during admission (n=6)]. All cases had a significant increase in the platelet count and D-dimer level. The levels of protein C and protein S first transiently decreased and were then restored to normal. Furthermore, two cases were weakly positive for antilipin antibody (ACA). In addition, four cases were positive for ACA as detected by ELISA (16) (QUANTA Lite ACA IgG III; Inova). A total of four cases received bronchoscopy, through which endobronchitis and necrotizing pneumonia were revealed. The bronchoalveolar lavage fluid was tested positive for *M. pneumoniae* (DNA sequence copy number, 7,880-16,343) and positivity for the macrolide resistance gene was detected in 2 cases.

Radiological examination. All 7 cases received dynamic monitoring by chest X-ray, pulmonary CT scan and computed tomographic pulmonary angiography (CTPA) during hospitalization. Furthermore, 6 cases were indicated to have extensive diffuse inflammatory changes in the two lungs upon chest X-ray or pulmonary CT scan (Figs. 1 and 2) and 1 case had subcutaneous emphysema (Fig. 3). Furthermore, 4 cases were combined with a moderate amount of pleural effusion and 1 case was combined with mild pericardial effusion (Fig. 4). CTPA indicated that 2 cases had a pulmonary arterial embolism in multiple branches bilaterally (Fig. 5, arrows); 2 cases had a pulmonary arterial embolism in the upper lobe of the right lung and one of the two lesions was located at the distal end of the right upper lung; 1 case had filling defects in the pulmonary artery branches in the upper lobe of the left lung; 2 cases had distal pulmonary artery embolism in the lower lobe of the right lung. Furthermore, 1 case with swelling in the lower limbs received local vascular ultrasound examination, through which thrombosis in the common femoral vein was detected.

Table I. Baseline demographic and clinical characteristics of the patients (n=7).

| Characteristic | Value |
|---------------|-------|
| Demographics  |       |
| Age (years)   | 8 (6, 11) |
| Male sex      | 4 (57.14) |
| Anthropometry |       |
| Body weight (kg) | 24.2 (21.3, 30) |
| Body height (cm) | 123.4 (119.5, 130) |
| BMI z-score   | 0.5 (-0.5, 1) |
| Clinical symptoms |   |
| Cough         | 7 (100) |
| Fever         | 7 (100) |
| Dyspnea       | 5 (71.43) |
| Swelling in limb | 1 (14.29) |
| Radiological examination | |
| Pulmonary CT   |       |
| Extensive diffuse inflammatory | 6 (85.71) |
| Subcutaneous emphysema | 1 (14.29) |
| Pleural effusion | 4 (57.14) |
| Pericardial effusion | 1 (14.29) |
| Pulmonary arterial embolism | |
| Bilateral multiple branches | 2 (28.57) |
| Upper lobe of the right lung | 2 (28.57) |
| Lower lobe of the right lung | 2 (28.57) |
| Upper lobe of the left lung | 1 (14.29) |

Values are expressed as the median (interquartile range) or n (%). BMI, body mass index.

Treatment. After admission, all of the cases were given the standard anti-infection therapy of macrolides and certain patients received concurrent antibiotic therapy with third-generation cephalosporins or carbapenems. Over the same period, moxifloxacin was selected for the 2 cases with positivity for the drug resistance gene. Those patients with dyspnea were treated by tracheal intubation and mechanical ventilation. In the meantime, other systemic treatments, such as organ protection and nutritional support, were given. Risk stratification was performed based on the guidelines of the American College of Chest Physicians (17), along with the thrombolysis and anti-coagulation therapies. Low-molecular-weight heparin calcium was injected subcutaneously at the dose of 50 IU/kg per time, twice daily. During the treatment, the pediatric patients were properly immobilized to avoid violent cough and movement. Since these pediatric patients did not have a basic history of congenital heart disease and presented with no pulmonary thrombosis, no shock due to PE and no deep vein thrombosis, thrombophilia was excluded and thrombolytic therapy was therefore not selected. However, anticoagulant therapy had no curative effect in 5 cases and the disease progressed to pulmonary infarction; thus, surgical resections were conducted.
### Table II. Results of auxiliary examinations of the patients.

| Patient | WBC (x10^9/l) | PLT (x10^9/l) | CRP (mg/l) | Serum M. pneumoniae resistance | BALF-DNA (M. pneumoniae) | Serum S. aureus antibody (µg/l) | CT | CTPA Pathology | CTPA Treatment | Outcomes |
|---------|---------------|---------------|------------|--------------------------------|-------------------------|--------------------------------|----|----------------|----------------|----------|
| 1       | 34.62         | 755           | 265        | Negative                        | 1:320                   | 16343                         | OP | -              | -              | Died     |
| 2       | 27.55         | 810           | 210        | Positive                        | 1:160                   | Positive                      | OP | -              | -              | Alive    |
| 3       | 5.93          | 480           | 80         | NA                              | NA                      | NA                            | NA | -              | -              | Alive    |
| 4       | 12.33         | 550           | 110        | NA                              | NA                      | Cpn                            | NA | -              | -              | Alive    |
| 5       | 20.12         | 610           | 164        | NA                              | NA                      | Cpn                            | NA | -              | -              | Alive    |
| 6       | 9.84          | 490           | 714        | NA                              | NA                      | Cpn                            | NA | -              | -              | Alive    |
| 7       | 18.13         | 680           | 123        | NA                              | NA                      | Cpn                            | NA | -              | -              | Alive    |

* Determined during the 2nd examination, which was performed 6.7±1.4 days after the 1st examination. ACA, anti-cardiolipin antibody; BALF-DNA, Mycoplasma pneumoniae DNA sequence copy number in BALF; BALF, bronchoalveolar lavage fluid; M. pneumoniae, Mycoplasma pneumoniae; Cpn, Chlamydia pneumoniae; CRP, C-reactive protein; CTPA, computed tomographic pulmonary angiography; OP, operation; PLT, platelets; S. aureus, Staphylococcus aureus; WBC, white blood cells; NA, not available.

**Figure 1.** Bilateral pulmonary extensive diffuse inflammation, with bilateral atelectasis of the dorsal lobes of a 7 year old male.

**Figure 2.** Bilateral pulmonary extensive diffuse inflammation with consolidation in the right lung of an 8 year old male.

**Figure 3.** Severe inflammatory changes in the bilateral lungs with extensive subcutaneous emphysema of an 11 year old male.

**Figure 4.** Inflammation in the middle lobe of the right lung and all lobes of the left lung of a 6 year old female, with partial atelectasis in the lower lobe of the left lung (upper panel) and mild pericardial effusion (lower panel).
Outcomes and follow-up. Anticoagulant therapy is the first choice for all pediatric patients with PE if there is no contraindication (18). A total of 2 patients achieved significant improvement after 15-21 days of treatment as detected by chest radiological examination and D-dimer test. They were discharged after the symptoms improved, and the PE disappeared 3-5 months later as detected during the follow-up. The 5 remaining cases exhibited no improvement in the local PE, which was confirmed by pulmonary CT scan on day 14-21 during the anticoagulation therapy. Pulmonary infarction was considered in certain patients who had local nodular solid lesions in the lungs with cavitation and the condition deteriorated to high dependence on oxygen. Among them, 2 patients had considerable pleural effusion (pus) and required surgical resection. Intraoperative findings included dark-red resected pulmonary tissues or yellowish-white infarct-like changes. These tissues had no contraction and dilation functions, with high tension and pus coating on their surface. The pathological diagnosis of the resected pulmonary tissues was pulmonary infarction (Figs. 6 and 7). Among them, 2 cases were combined with Acinetobacter baumannii infection after surgery and Acute Respiratory Distress Syndrome occurred 3-8 days later. Although the patients were treated with extracorporeal membrane oxygenation, their condition did not improve and they eventually died. The remaining surviving pediatric patients underwent 6-12 months of follow-up and respiratory rehabilitation and they recovered to a normal state.

Literature search and review. Using ‘Pulmonary Embolism’ and ‘Mycoplasma pneumoniae pneumonia’ with ‘pediatric’ as the keywords, relevant articles were searched in the PubMed, MEDLINE, Update, Web of Science and Embase databases. The Chinese subject heading terms used in the Wanfang, Chinese National Knowledge Infrastructure and Chongqing VIP databases were the same as those above. The inclusion criteria were as follows: i) Pediatric patients with a confirmed diagnosis of MPP and PE during the treatment process; and ii) Studies published within the last 20 years. Exclusion criteria was incomplete clinical information. Ultimately, the clinical data of 10 pediatric cases with MPP and PE were reported and their details are listed in Table III (19‑26).

These cases were aged between 6 and 13 years (median age, 9.0 years) with a male/female ratio of 5:1. The levels of M. pneumoniae antibody were significantly increased, along with a transient decrease in protein S and protein C. The lesions were located at the lower lobe close to the hilus of the lung. After receiving anti-infective and anti-coagulant treatments, 8 cases improved but 1 patient died.

Discussion

Pediatric patients with critical MPP complicated with PE was rarely reported. Cases with mild PE may be asymptomatic, while severe cases may suffer from pulmonary arterial hypertension, unstable hemodynamics or even sudden death (27,28). The common symptoms include shortness of breath, chest pain and even dyspnea (19‑26). Missed diagnosis may occur if young pediatric patients are not able to properly describe
Table III. Details of previous studies.

| Author / Year | Case no. | Sex | Age (years) | Interval (days) | Antibody to *M. pneumoniae* | Agglutination test | D reg | Treatment | Outcome | (Refs.) |
|---------------|---------|-----|-------------|----------------|----------------------------|---------------------|-------|-----------|---------|--------|
| Graw-Panzer (2009) | 1 | M | 13 | 5 | ELISA IgM (1:128) | Increased D-dimer, protein S deficiency and positive ACA. | Left popliteal vein embolism and PE. | Heparin + warfarin | Radiographic chest findings returned to normal after 3 months and the anemia resolved gradually over 5 months. | (19) |
| Chen (2013) | 2 | F | 12 | 12 | PA (1:160) | Increased D-dimer, positive ACA | Thrombosis in right lower limb and PE (left lower lobe). | Low-molecular-weight heparin + warfarin | The chest X-ray was almost normal at follow-up after 6 months. | (20) |
| Brown (2008) | 3 | M | 6 | 16 | Complement binding (1:640) | Positive ACA and acquired activated protein C resistance. | Femoral vein embolism and PE (left lower lobe) | Not mentioned | Alive. | (21) |
| Su (2012) | 4 | M | 6 | 17 | ELISA (1:128) CA (1:1,024) | Increased D-dimer, positive ACA and decreased activity of plasma protein C. | PE (left lower lobe) | Heparin + warfarin | At the 3-month follow-up, Aca was negative, plasma protein C activity recovered and lung lesions were absorbed. | (22) |
| Wei (2015) | 5 | M | 9 | 23 | Not mentioned (1:1,280) | Increased D-dimer and positive ACA. | PE (right lower lobe). | Heparin + warfarin | Chest radiographic findings returned to normal after 3 months. | (23) |
| Zhuo (2015) | 6 | M | 9 | 10 | PA (1:160) | Increased D-dimer. | PE (mainly on the right side). | Low-molecular-weight heparin + warfarin | Died on the eighth day after admission. | (24) |
| Qin (2019) | 7 | F | 10 | 14 | Not mentioned (1:320) | Increased D-dimer, anticardiolipin IgM antibody was positive, plasma protein C/S activity was not mentioned. | PE (bilateral lung). | Low-molecular-weight heparin calcium + warfarin | At the 8-month follow-up, chest CT indicated old lung lesions in both lungs, segmental atelectasis of the right upper lung accompanied by bilateral lower lung filaments and a small amount of pleural lesions in the left lung. | (25) |
their symptoms. Therefore, if PE is not discovered in a timely manner, the anti-coagulation treatment is delayed and the disease may progress into acute pulmonary infarction or even death. When encountering pediatric cases with MPP, the patient or the parents should be asked whether there is a family history of protein C/protein S deficiency, recent history of surgeries or presence of congenital vascular malformation, so as to preliminarily assess the risk of PE. In the present study, evaluation at the early stage of admission indicated a low risk of thrombosis in all cases. However, their symptoms kept on deteriorating during the treatment and chest radiological examination indicated poor recovery. In combination with laboratory tests and chest radiological examination, the diagnosis of PE was confirmed and the standard treatment was provided.

For such pediatric cases, physicians should begin early dynamic monitoring and examination, which may be able to effectively control disease progression, reduce surgical rates and mortality. However, with the current technological standards available, it is still limited to perform the interventional treatment and implement thrombolysis therapy for young pediatric patients with PE. More work, such as a more appropriate design using more sophisticated instruments, still needs to be done to overcome this deficiency in the future.

The pathogenesis of M. pneumoniae infection with thrombosis remains to be fully elucidated, but it may be associated with immune damage mediated by infection (7,29-31). Since the membrane proteins and glycolipids of M. pneumoniae have certain common antigens in the heart, liver, lung, brain, kidney and smooth muscle tissues of the human body, upon infection of the host with M. pneumoniae, the corresponding antibodies are produced and the immune complex is formed to activate complement, which produces neutrophils. Previous studies reported that embolism may affect multiple sites of the body after M. pneumoniae infection, including the brain, lower extremity veins, spleen and pulmonary arteries (10-15). Chemokines, which attract a large number of white blood cells to invade the lesion, release a large number of inflammatory mediators and lysosomal enzymes, causing inflammatory damage to target organs. It was reported that patients with MPP and thrombus were positive for ACA (32,33). ACA is an autoantibody that targets antigens in platelets and cardiolipin on endothelial membranes and is associated with thrombogenesis. The present study concluded that M. pneumoniae infection may cause vascular endothelial cell injury and ACA positivity, leading to a temporary hypercoagulability state that induces thrombosis. Under severe conditions, M. pneumoniae infection further affects the synthesis of coagulation factors and thrombin (e.g., protein C, protein S and antithrombin III), resulting in embolism. Certain patients may acquire protein C or protein S deficiency/resistance (21,34). The D-dimer test is an important screening method for PE and a negative result may exclude PE with 100% certainty (35,36). While CTPA is considered as the gold standard for PE diagnosis (37). Based on the above points, in a child with severe MPP, the D-dimer test, ACA test, Protein C test, Protein S test and CTPA should be considered to prevent the occurrence of PE.

At present, the major treatment for pediatric patients with acute PE is anticoagulant therapy, the purpose of which

| Author       | Case no. | Age (years) | Interval (days) | Antibody to M. pneumoniae | Agglutination test | D-reg, damaged region | Treatment | Outcome |
|--------------|----------|-------------|----------------|--------------------------|--------------------|----------------------|-----------|---------|
| Zhang (2019) | 8        | F           | 8              | ELISA (1:1,280)          | Increased D-dimer, II, protein C, S were normal. | Increased D-dimer, II, protein C, S were normal. | Methylprednisolone + Nadroparin calcium | Thrombosis of the posterior tibial vein in both lower limbs and PE (bilateral lung). |
| Zhang (2019) | 9        | F           | 5              | ELISA (1:1,280)          | Increased D-dimer, II, protein C, S were normal. | Protein S was decreased. | Methylprednisolone + Nadroparin calcium | Thrombophlebitis of the great saphenous vein in the right lower limb and PE (lower lobe in the bilateral lung). |
is to prevent acute thrombosis and expansion. If available, local thrombus therapy or interventional thrombus therapy may be performed. During the anticoagulation therapy with low-molecular-weight heparin, dynamic monitoring of the coagulation status and treatment outcome is required to avoid bleeding, whose risk is considerable. Furthermore, cooperation between the pediatric thoracic surgery department and the vascular surgery department is preferred and surgical intervention may be provided if necessary.

Critical *M. pneumoniae* infection in pediatric patients is associated with a high risk of PE. During clinical treatment, such cases should be screened for high-risk factors and patients could be closely monitored for any manifestations of PE. Necessary examinations, particularly CTPA, should be performed to confirm the diagnosis and to initiate standard treatment as early as possible. Surgical intervention is another important salvage to reduce poor prognosis, if the disease progresses to pulmonary infarction, and therefore results in serious and life threatening complications.

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Availability of data

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

Authors' contributions

CQS conceived the current study and drafted and revised the manuscript. CFY collected the literature and reviewed and revised the manuscript. YA and ZYZ collected the data and performed initial analyses. YML coordinated and supervised data collection, and critically reviewed the manuscript. CQS and YML confirm the authenticity of all the raw data. All authors read and approved the final manuscript.

Ethics approval and consent to participate

The research followed international and national regulations in accordance with the Declaration of Helsinki. The study was approved by the Ethics Committee of the First Hospital of Jilin University. (Changchun, China; approval no. 2019-253).

Patient consent for publication

Written informed consents were obtained from the patients' legal guardians for the publication of any accompanying images prior to submission.

Competing interests

The authors declare that they have no competing interests.

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