A Case of Common Variable Immune Deficiency Presenting as Recurrent Pneumococcal Pneumonia

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Common variable immune deficiency (CVID) is the most common primary immune deficiency, which is manifested as chronic recurrent respiratory infections and hypoglobulinemia. CVID usually presents in the second or third decade of life. A 33-year-old woman was admitted with recurrent pneumococcal pneumonia with bacteremia and had very low levels of serum immunoglobulin G, M and A. This case emphasized a high index of suspiciousness for diagnosis of CVID in a mid-adulthood patient presenting with recurrent pneumonia with hypoglobulinemia.

Key Words: Common Variable Immunodeficiency; Pneumonia, Pneumococcal; B-Lymphocytes

Introduction

Common variable immunodeficiency (CVID) is a primary immunodeficiency characterized by B cell dysfunction. CVID generally comprises antibody deficiencies that present in either late childhood or, more typically, early to mid adulthood¹. The diagnosis is based on decreased serum immunoglobulins and a failure to produce antigen-specific antibodies in response to vaccinations or infections². We report a CVID case of a mid-aged woman who had presented with recurrent severe pneumococcal pneumonia.

Case Report

A thirty three-year-old woman admitted our hospital with chilling sense, myalgia and dyspnea. Before admission, she felt chilling and myalgia for 5 days and shortness of breath for 3 days. She suffered recurrent rhinor-

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She appeared chronically ill and showed frequently coughing mixed with purulent sputum. There were no gastrointestinal symptoms. Her height was 160 centimeters and weight was 50 kilograms. Her blood pressure was 120/80 mm Hg, pulse rate 88 beats per minute, respiratory rate 18 per minute, and body temperature 37.6°C. There was no facial anomaly. Cervical and axillary lymph nodes were not palpable. On lung sound, inspiratory coarse crackle was heard on her right upper thorax. There were no clubbing fingers.

Initial white cell count was 13,400/mm³ with 77% segmented neutrophils and 18% lymphocytes. No ab-

rhea, cough, and phlegm during last ten years. She was diagnosed chronic rhinosinutis 5 years ago, and iron deficiency anemia three months ago. Five months before admission, she got myalgia and chilling sense for several days and had taken antibiotics at private clinic. Three months ago, she admitted this hospital because of bilateral pneumococcal pneumonia with bacteremia (Figure 1A). At that time, she received cefotetan (2 g, bid, iv) with klarithromycin (500 mg, bid, oral) for 14 days. With full recovery, she discharged. She had received vaccinations as scheduled except pneumococcal vaccine. She had no regular medication except iron pill. Her family history was not remarkable. She has been married and housewife,

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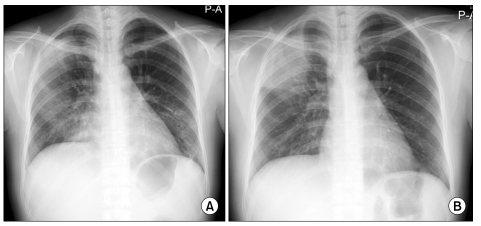


Figure 1. (A) Chest x-ray at previous admission showed pneumonia in right upper, lower lung and left lower lung field. (B) Chest x-ray at this admission demonstrated recurrence of pneumonia in right upper lung field.

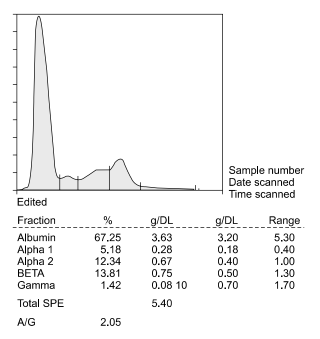


Figure 2. Serum electrophoresis showed a markedly decreased gamma globulin fraction.

normal blood cells were on peripheral blood smear. Hemoglobin was 11.7 g/dL, Hct 36.6%, MCV 82 fL (normal range, 81~101), MCH 26.2 pg (normal ragne, 26.2~34.2) and platelets were 213,000/mm³. In urinalysis, no protein and red blood cells were detected. In blood chemistry, BUN was 11.5 mg/dL, creatinine 0.57 mg/dL, uric acid 3.0 mg/dL, total bilirubin 0.6 mg/dL, AST 14 IU/L, ALT 16 IU/L, alkaline phosphatase 297 IU/L, total protein 6.5 g/dL, albumin 4.3 g/dL and globulin 2.2 g/dL. Serum iron was 6 μ g/dL (normal range, 50~170), total iron binding 305 μ g/dL (normal



Figure 3. Chest CT scan showed right paratracheal lymph nodes enlargements and parenchymal infiltrates on right upper lobe. CT: computed tomography.

range, $262 \sim 474$) and ferritin 65 ng/mL (normal range, $5 \sim 148$). Serum protein electrophoresis showed a markedly decreased gamma globulin 0.08 g/dL (normal range, $0.7 \sim 1.7$) (Figure 2). Precipitated lines about Ig A, G, and M in serum immunoglobulin electrophoresis were vague. Each serum immunoglobulins were decreased as a whole: serum Ig A 43 mg/dL (normal range, $800 \sim 1,800$), Ig G 1 mg/dL (normal range, $90 \sim$ 450), and Ig M 19 mg/dL (normal range, $65 \sim 333$). Her blood type was A and Rh (+). Serum iso-hemagglutinin tests anti-A and anti-B on her red blood cells were negative. Anti-tetanus antibody was not detected on second admission. Anti HIV antibody was negative. In lymphocyte subset flow cytometry, CD4+ T cells were $896/\text{mm}^3$ (normal range, $700 \sim 1,100$), CD8+ T cells 979/mm³ (normal range, $500 \sim 900$) and CD19+ J Jeong et al: A case of CVID with recurrent pneumonia

B cells $351/\text{mm}^3$ (normal range, $200 \sim 400$).

Her chest x-ray showed wedge shaped consolidation in right upper lung field (Figure 1B). Chest computed tomography revealed pneumonic consolidation and ground glass opacities of right upper and lower lung fields with reactive mediastinal and right supraclavicular lymphadenopathies (Figure 3). Transthoracic echocardiogram showed normal cardiac function and no evidence of endocarditis. Esophagogastroduodenoscopy and colonoscopy were normal. Abdominal and pelvic computed tomography showed normal findings.

Her urinary pneumococcal antigen was positive and *Streptococcus pneumoniae* grew in the sputum and blood cultures. After intravenous or oral moxifloxacin (400 mg, qd) for two weeks, she was able to discharge because of clinical and radiological improvement. Her illness was common variable immune deficiency. She has been treated by intravenous immunoglobulin 400 mg/kg monthly. Up to 1 year follow up, there was no serious infection event.

Discussion

CVID has variable clinical manifestations, the most common being recurrent bacterial infections caused by encapsulated bacteria such as Streptococcus pneumoniae, Haemophilus influenzae. Bacterial infections commonly involve the paranasal sinuses and respiratory tract². As CVID is considered as genetic disease, it is the sporadic pattern and relatively late onset. The mean age at diagnosis is between 25 years and 45 years of age. This disorder results from failed B-cell differentiation, with impaired secretion of immunoglobulins³. A lot of defects of T-cell function and deficits in the memory B-cell pool have been identified, but the underlying cause of this defect remains unknown. Therefore, the hallmark of CVID is hypogammaglobulinemia, and the standard treatment is IV immunoglobulin replacement. On the other hand Ig A, Ig G and/or Ig M levels should be less than two standard deviations below the mean for age-adjusted standardized reference, and reduced levels of at least two Ig isotypes².

To confirm the diagnosis of CVID, there were inadequate antibody responses to pneumococcal vaccine and tetanus toxoid or absent isohemagglutinins⁴. She did not receive tetanus vaccine on admission. Unfortunately we did not evaluate the anti-Tetanus titer for response of vaccination. Our patients also showed markedly decreased gamma globulin fraction in serum protein electrophoresis and immunoglobulins despite of negative Anti Tetanus Ab. We observed slightly enlarged mediastinal and right supraclavicular lymph nodes. There were normal sized spleen and abdominal lymph nodes. We considered as reactive lymph node enlargements, so did not perform mediastinoscopic lymph node biopsy. Throughout clinical and radiological examinations, we should evaluate malignancy such as lymphoma. Also there is an increased risk of malignancy in CVID. Chest commuted tomography should be followed as recommended. We excluded other causes of hypogammaglubulinemia as guided⁵. There was no drug exposure and evidence of chromosomal abnormalities. She had no conditions of excessive loss of immunoglobulin such as nephrosis, severe diarrhea. Because of female patient, normal CD19+ B cells and T cells counts, we could exclude X linked aggammaglobulinemia and severe combined immnunodeficiency.

There were three CVID case reports in Korea. One was a seventeen years old girl who had presented continuous infectious events from early childhood⁶. The second case was a 10-year old boy who diagnosed with CVID and a fatal measles⁷. The other was a twenty years old woman who was diagnosed by granulomatous liver disesase but her recurrent infectious episodes had started 14 years ago⁸. But our case was somewhat typically presented in a mid-thirty aged woman whose infections had initiated 5 years ago.

Intravenous immunoglobulin (IVIG) treatment in addition to early diagnosis was thought as important prognostic factor in CVID. IVIG is effective and is currently the mainstay of therapy for CVID⁹. Intravenous immunoglobulin also reduces the incidences of pneumonia and serious recurrent bacterial infections and prevents chronic lung disease and enteroviral meningoencephalitis¹⁰.

Besides recurrent infections, CVID patients have an increased tendency to develop autoimmunity, lymphoproliferative disease and malignancies. Although these disease complications cause severe morbidity, the enormous heterogeneity in the clinical presentation of CVID induced delays in recognizing its illness and complications. As this case presented, CVID should be suspected, if any patient showed recurrent infections, especially of the upper or lower respiratory tract. After diagnosis of CVID, the complications and associated conditions should be followed up.

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References

- 1. Park MA, Li JT, Hagan JB, Maddox DE, Abraham RS. Common variable immunodeficiency: a new look at an old disease. Lancet 2008;372:489-502.
- Cunningham-Rundles C, Bodian C. Common variable immunodeficiency: clinical and immunological features of 248 patients. Clin Immunol 1999;92:34-48.
- 3. Bryant A, Calver NC, Toubi E, Webster AD, Farrant J.

Classification of patients with common variable immunodeficiency by B cell secretion of IgM and IgG in response to anti-IgM and interleukin-2. Clin Immunol Immunopathol 1990;56:239-48.

- Primary immunodeficiency diseases. Report of a WHO scientific group. Clin Exp Immunol 1997;109(Suppl 1): 1-28.
- Conley ME, Notarangelo LD, Etzioni A. Diagnostic criteria for primary immunodeficiencies. Representing PAGID (Pan-American Group for Immunodeficiency) and ESID (European Society for Immunodeficiencies). Clin Immunol 1999;93:190-7.
- Lee MK, Bae SK, Choi SJ, Park MH, Kim YI. A case of common variable immunodeficiency. Korean J Med 1985;28:273-80.
- Cho SJ, Kim SH, Lee SJ, Yoo KH, Kim KH. A case of common variable immunodeficiency with fatal measles. J Korean Pediatr Soc 2001;44:1463-8.
- Oh TH, Lee HC, Lee TY, Song BC, Chung YH, Lee YS, et al. A case of granulomatous hepatitis associated with common variable immunodeficiency. Korean J Hepatol 2001;7:95-9.
- Bonilla FA, Bernstein IL, Khan DA, Ballas ZK, Chinen J, Frank MM, et al. Practice parameter for the diagnosis and management of primary immunodeficiency. Ann Allergy Asthma Immunol 2005;94(5 Suppl 1):S1-63.
- Busse PJ, Razvi S, Cunningham-Rundles C. Efficacy of intravenous immunoglobulin in the prevention of pneumonia in patients with common variable immunodeficiency. J Allergy Clin Immunol 2002;109:1001-4.