Case Study

**Japanese Variant of Multicentric Castleman’s Disease Associated With Serositis and Thrombocytopenia — A Report of Two Cases: Is TAFRO Syndrome (Castleman-Kojima Disease) a Distinct Clinicopathological Entity?**

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Multicentric Castleman’s disease (MCD) is a polyclonal lymphoproliferative disorder that manifests as marked hyper-γ-globulinemia, severe inflammation, anemia, and thrombocytosis. Recently, Takai et al. reported a new disease concept, TAFRO syndrome, named from thrombocytopenia, anasarca, fever, reticulin fibrosis, and organomegaly. Furthermore, Kojima et al. reported Japanese MCD cases with effusion and thrombocytopenia (Castleman-Kojima disease). Here, we report two cases of MCD associated with marked pleural effusion, ascites, and thrombocytopenia, and discuss the independence of the TAFRO syndrome (Castleman-Kojima disease). Case 1: A 57-year-old woman had fever, anemia, anasarca, and some small cervical lymphadenopathy. Although she had been administered steroid therapy, and full-coverage antibiotics, her general condition, including fever, systemic inflammation, and anasarca, deteriorated steadily. We administered chemotherapy [CHOEP (cyclophosphamide, doxorubicin, vincristine, etoposide, and prednisolone) regimen], but despite a transient improvement, she died due to septic shock. Case 2: A 73-year-old man with a history of aplastic anemia and remission presented with fever, severe inflammation, and anasarca. Prednisolone was administered (15 mg daily), and his hyperinflammation once improved. Nevertheless, his general condition, including pleural effusion and ascites, worsened, and C-reactive protein and interleukin-6 levels showed marked increases. The patient died due to multiorgan failure. Cases of TAFRO syndrome (Castleman-Kojima disease) are still rare. Therefore, it is necessary to conduct multicenter clinical surveys including similar cases, such as ours, to reach a consensus regarding diagnostic criteria, therapeutic strategy, and pathophysiological etiology for this syndrome. [J Clin Exp Hematop 53(1): 79-85, 2013]

**Keywords:** Castleman’s disease, POEMS syndrome, anasarca, interleukin-6, polyclonal hypergammopathy

**INTRODUCTION**

Multicentric Castleman’s disease (MCD) is a polyclonal lymphoproliferative disorder (LPD) that manifests as marked hyper-γ-globulinemia, severe inflammation, anemia, and thrombocytosis.1,4 In western countries, this syndrome is usually associated with human herpes virus-8 (HHV-8) infection as well as human immunodeficiency virus (HIV) infection.4,5 On the other hand, most Japanese patients are negative for both HHV-8 and HIV. Therefore, MCD in western countries and Japan seem to be different disorders.7,8 Severe inflammation due to overproduction of interleukin-6 (IL-6) is a finding common to patients in western countries and Japan.

Takai et al. reported a new disease concept, TAFRO syndrome, named from thrombocytopenia, anasarca, fever, reticulin fibrosis, and organomegaly.9 Furthermore, Kojima et al. reported Japanese MCD cases with effusion and thrombocytopenia (Castleman-Kojima disease).10 These conditions seem to be different from hyaline-vascular type of MCD or...
POEMS syndrome (polyneuropathy, organomegaly, endocrinopathy, M protein, and skin changes).

Here, we report two cases of MCD associated with marked pleural effusion, ascites, and thrombocytopenia, and discuss the independence of the TAFRO syndrome (Castleman-Kojima disease).

Case 1

A 57-year-old woman with high fever and dry cough was diagnosed with pneumonia and anemia (hemoglobin 9.6 g/dL) and thrombocytopenia (52 × 10^3/μL) one mon before admission to Kanazawa Medical University Hospital (KMU) (Fig. 1). Although antibiotics, antifungal drugs, and γ-globulin were administered, there were no improvements in her high inflammation, anemia, and thrombocytopenia. Due to suspicion of hemophagocytic syndrome, methylprednisolone pulse therapy (500 mg × 3 days) followed by prednisolone at 60 mg daily was administered. However, with the exception of dry cough, her condition worsened, and hypoalbuminemia and anasarca (pleural effusion and ascites) emerged and she was admitted to KMU.

The patient gained 14 kg body weight in one mon. She had fever, anemia, anasarca, and some small cervical lymphadenopathy (< 1 cm in diameter) on admission to KMU. On laboratory examination, mild proteinuria and microscopic hematuria were detected by urinalysis. On hematological examination, red blood cell (RBC) count was 2.32 × 10^6/μL, hemoglobin 7.1 g/dL, hematocrit 20.7%, reticulocytes 11.4 × 10^6/μL, white blood cell count 6.87 × 10^3/μL (neutrophil granulocytes 84.3%, lymphocytes 10.4%, eosinophils 0.8%, and monocytes 1.6%), and platelets (PLT) 13 × 10^3/μL. Erythrocyte sedimentation ratio (1 hr) was elevated to 109 mm, and C-reactive protein (CRP) was elevated (17.73 mg/dL). The patient showed prothrombin time-international normalized ratio 1.44, activated partial prothromboplastin time 40.6 sec, hepaplastin time 34.9%, fibrinogen 461 mg/dL, fibrinogen degradation product 32.1 µg/mL, D-dimer 22.85 mg/mL, and antithrombin III 46.5%. Blood urea nitrogen was 30 mg/dL, creatinine 1.34 mg/dL, total protein 5.0 g/dL, albumin 1.5 g/dL, IgG 1,860 mg/dL, IgA 288 mg/dL, IgM 40 mg/dL, IgE 97.5 IU/mL, IgG 97.5 IU/mL, C3 75 mg/dL, C4 24 mg/dL, CH50 31.1 U/mL, total bilirubin 2.3 mg/dL, direct bilirubin 1.8 mg/dL, lactate dehydrogenase (LDH) 130 U/L, aspartate aminotransferase 20 U/L, alanine aminotransferase 10 U/L, alkaline phosphatase (ALP) 710 U/L, ferritin 640 ng/mL, β2-microglobulin 9.6 mg/mL, soluble IL-2 receptor 4,310 U/mL, IL-6 65.5 pg/mL, and vascular endothelial growth factor (VEGF) 91 pg/mL. Direct and indirect Coombs’ tests were both positive (+), antinuclear antibody ×160 speckled pattern, and platelet-associated IgG 86.7 ng/10^7 cells, but other autoantibodies were negative. Serum antibodies against HIV, human T-lymphotropic virus-1, and HHV-8 were all negative. Bone marrow aspiration and biopsy revealed slightly hypercellular bone marrow without any tumor cells, dysplastic changes, hemophagocytosis, or fibrosis. Furthermore, no monoclonality was detected by flow cytometric analysis and no karyotypic abnormalities were detected by G-banding analysis. Massive pleural effusion and ascites were detected by whole-body computed tomography (CT) scan (Fig. 2a), and weak accumulation (standardized uptake value; SUVmax < 3) of 18F-fluorodeoxy glucose (18FDG) to cervical and abdominal lymph nodes by 18FDG-positron emission tomography (PET)-CT (Fig. 2b). On histological examination of cervical lymph node (LND) biopsy specimen, the basic structure of LND was unclear, and massive plasma cell growth around the atrophic germinal center was seen but no neoplastic features were detected (Fig. 3). Therefore, MCD was the most likely diagnosis but lymphadenopathy associated with autoimmune diseases, such as systemic lupus erythematosus, or angioimmunoblastic T-cell lymphoma (AITL) were also considered.

Although she had been administered and continued steroid therapy, including pulse-therapy, and full-coverage antibiotics and antifungal drugs since before and after admission to KMU, the patient’s general condition, including fever, systemic inflammation, pancytopenia, and anasarca, deteriorated steadily. Albumin administration and RBC/PLT transfusion did not improve her condition or clinical data. Acute renal failure of pre-renal type occurred, and steroid pulse therapy (methylprednisolone 1 g × 3 days) was commenced. Al-
though the result of LND biopsy indicated MCD as the most likely diagnosis, we administered chemotherapy by the CHOEP regimen (cyclophosphamide, doxorubicin, vincristine, etoposide, and prednisolone) because we suspected that she may have lymphoma, especially AITL, due to the progressively deteriorating clinical course. Despite transient improvement of general condition, including renal function, the patient died due to septic shock associated with febrile neutropenia. Bilateral pleural effusion (right 800 mL and left 700 mL) and 3,000 mL of ascites, splenomegaly (200 g), systemic candida infection were documented, but no particular neoplastic cells were detected at autopsy.

**Case 2**

A 73-year-old man with a history of aplastic anemia since the age of 69 years old had been treated with anti-thymocyte globulin, cyclosporin A, granulocyte colony stimulating factor, RBC and PLT transfusion, and anabolic steroid administration (Fig. 4). Due to the efficacy of this treatment, his course of aplastic anemia itself had been become stable with no transfusion requirement since the age of 71 years old, but he had been followed up due to iron overload-induced liver cirrhosis, diabetes mellitus, hypothyroidism, and chronic heart failure. Since September 2008, when he was 72 years old, the patient showed remittent fever, general malaise, CRP elevation, polyclonal hypergammopathy, and hyponatremia. His performance status was grade 3 according to the Eastern Cooperative Oncology Group (ECOG) criteria and he showed marked emaciation. On physical examination, anemia and...
horizontal nystagmus toward the left side and clumsiness of bilateral finger-nose test were seen. However, the patient showed neither lymphadenopathy nor hepatosplenomegaly.

Urine test indicated mild proteinuria and microscopic hematuria without casts. On hematological examination, RBC was $1.84 \times 10^{12}/\mu L$, hemoglobin 7.6 g/dL, hematocrit 21.5%, reticulocyte 8.6 $\times 10^{9}/\mu L$, white blood cell 4.73 $\times 10^{9}/\mu L$, neutrophil granulocyte 70.0%, lymphocyte 10.5%, eosinophil 15.5%, and monocyte 3.5%), and PLT 24 $\times 10^{9}/\mu L$. Erythrocyte sedimentation ratio (1 hr) was elevated to 153 mm, and CRP was elevated to 4.17 mg/dL. Prothrombin time-international normalized ratio was 1.25, activated partial prothromboplastin time 36.7 sec, hepaplastin time 59.4%, fibrinogen 243 mg/dL, and D-dimer 1.48 mg/mL. Blood urea nitrogen was 54 mg/dL, creatinine 1.45 mg/dL, sodium 127 mEq/L, potassium 4.7 mEq/L, chlorine 94 mEq/L, total protein 11.1 g/dL, albumin 1.8 g/dL, IgG 7.080 mg/dL, IgA 61 mg/dL, IgM 74 mg/dL, IgE 686 IU/mL, C3 18 mg/dL, C4 < 2 mg/dL, CH50 13.0 U/mL, total bilirubin 1.1 mg/dL, LDH 127 U/L, aspartate aminotransferase 57 U/L, ALP 630 U/L, ferritin 2,217 ng/mL, β2-microglobulin 11.4 μg/mL, IL-6 49.0 pg/mL, and VEGF 25 pg/mL. Serum antibodies against HIV, human T-lymphotropic virus-1, and HHV-8 were all negative. Rheumatoid factor was weakly positive (13 IU/mL), but the patient was negative for all other autoantibodies. Bone marrow aspiration revealed normocellular bone marrow without any tumor cells, dysplastic changes, and hemophagocytosis. Furthermore, no monoclonality was detected by flow cytometric analysis and no karyotypic abnormalities were determined by G-banding analysis. Mild fibrosis was seen in bone marrow biopsy (Fig. 5a & 5b). CT scan showed bilateral pleural effusion (Fig. 6b) and ascites, but no particular lymphadenopathy or hepatosplenomegaly. Magnetic resonance scan of the brain showed an elliptical mass measuring 2 × 1 cm with high intensity on T2 imaging at cerebellopontine angle (neurinoma) (Fig. 6a). Weak accumulation of $^{18}$FDG in PET-CT was seen on the cerebellopontine angle mass, but no other accumulation, such as lymphadenopathy, was seen (Fig. 6c).

Although no particular mass formation was documented by imaging study, he was suspected to have an MCD-like hyper-IL-6 condition due to severe inflammation and marked polyclonal hyper-γ-globulinemia.

After admission, prednisolone was administered (15 mg daily), and his hyperinflammation, including CRP, IgG, IgG4, and IL-6, improved once. Open brain surgery was performed for brain tumor (neurinoma) of the cerebellopontine angle followed by stereotactic radiosurgery. Nevertheless, his general condition, including pleural effusion and ascites, worsened and CRP and IL-6 increased markedly. The patient died due to multiple organ failure.

At autopsy, secondary hemochromatosis due to transfu-
sion, pyogenic spondylitis, small multiple lymphadenopathy showing mixed-type MCD histopathology (HHV-8 negative by immunohistochemical staining) (Fig. 5c & 5d), and massive pleural effusion (right 1,600 mL and left 1,200 mL), 180 mL of pericardial effusion, 900 mL of ascites, 160 g of splenomegaly, and 1,810 g of hepatomegaly were documented.

**DISCUSSION**

Castleman’s disease (CD) was first described by Castleman et al. CD is a LPD, and three disorders bearing the eponym CD have been identified and were reviewed by Frizzera et al.: localized CD of the hyaline-vascular (HV) type, localized CD of the plasma cell (PC) type, and MCD.1-4 However, several reports have indicated that MCD is composed of several disease entities, including idiopathic MCD, and secondary MCD due to HIV infection, autoimmune disease-associated lymphadenopathy, POEMS syndrome, and lymphomas.12-19 MCD is a clinicopathological entity of generalized LPD that manifests as polyclonal hypergammopathy, anemia, thrombocytosis, and severe inflammation.

The etiology of MCD remains uncertain. MCD in western countries develops in HIV-positive patients associated with HHV-8 and exhibits an aggressive and usually fatal course associated with infectious complications and the development of malignant tumors, such as Kaposi’s sarcoma or B-cell lymphoma.4-6 Viral IL-6 expression due to HHV-8 infection causes hyper-IL-6 syndrome.4-6 Hyper-IL-6 induces VEGF production resulting in angiogenesis, induces B-cell differentiation resulting in plasma cell expansion and hyperγ-globulinemia, induces megakaryocyte differentiation in bone marrow resulting in thrombocytosis, and induces acute inflammatory protein production in the liver resulting in increases in CRP, fibrinogen, serum amyloid A, etc. Furthermore, hepcidin is induced in the liver resulting in microcytic anemia due to both inhibition of iron absorption and a decline of iron recycling from the reticuloendothelial system (Table 1).

On the other hand, MCD in Japan is a hyper-IL-6 syndrome without HIV and HHV-8 infection in most cases, and usually exhibits a chronic disease course. Moreover, MCD in Japan does not appear to progress to Kaposi’s sarcoma or B-cell lymphoma.

Kojima et al. reported that idiopathic MCD in Japan consisted of two variants with distinct clinicopathological features, i.e., idiopathic plasmacytic lymphadenopathy with polyclonal hyperimmunoglobulinemia (IPL) type and non-IPL type.21-24 IPL type MCD resembles PC type MCD in western countries characterized by prominent polyclonal hyperimmunoglobulinemia, systemic manifestations, such as malaise, fever, weight loss, and high serum IL-6 levels.1-4 Non-IPL type MCD is characterized by mixed type or HV type CD histology, high incidence rate of pleural effusion and ascites, and is frequently associated with autoimmune diseases during the clinical course. Therefore, they suggested that a portion of non-IPL type cases may be secondary MCD, i.e., autoimmune disease-associated LPD.21,24

Recently, Takai et al. reported three cases of TAFRO syndrome, a new disease concept that is similar to but is distinct from AITL, PC type MCD, or POEMS syndrome.9 Furthermore, Kojima et al. reported seven cases of MCD with effusion at initial clinical presentation, five of which also showed thrombocytopenia (Castleman-Kojima disease).10 At a recent meeting, Japanese hematologists and pathologists collaborated to discuss the features of TAFRO syndrome (Castleman-Kojima disease). A number of particular findings were raised at this meeting, such as anasarca (pleural effusion and ascites), thrombocytopenia and microcytic anemia on hemograms, renal dysfunction, ALP elevation, low level of LDH, relatively mild polyclonal hypergammopathy (IgG level is usually < 4,000 mg/dL), some immunological abnormalities (positivity of rheumatoid factor, platelet-associated IgG, anti-

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<th>Table 1. Possible mechanism of multicentric Castleman’s disease (MCD) and Castleman-Kojima disease (TAFRO syndrome)</th>
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<tr>
<td>HIV infection → immune deficiency → HHV infection → HHV-8 vIL-6 expression</td>
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<tr>
<td>→ IL-6 overproduction</td>
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<tr>
<td>→ induction of VEGF production → angiogenesis</td>
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<td>→ B-cell differentiation → plasma cell expansion, hyper-γ-globulinemia</td>
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<td>→ megakaryocyte differentiation → thrombocytosis</td>
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<td>→ liver; production of acute inflammatory proteins → CRP, fibrinogen, SAA, etc</td>
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<tr>
<td>→ induction of hepcidin</td>
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<td>→ inhibition of iron absorption in gut</td>
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<tr>
<td>→ inhibition of iron recycling in reticuloendothelial system → anemia</td>
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<tr>
<td>→ other hit! → cytokine storm other than IL-6 and/or VEGF</td>
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<tr>
<td>→ anasarca, thrombocytopenia; Castleman-Kojima disease (TAFRO syndrome)</td>
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<tr>
<td>HIV, human immunodeficiency virus; HHV, human herpes virus; vIL-6, viral interleukin-6; VEGF, vascular endothelial growth factor; CRP, C-reactive protein; SAA, serum amyloid A</td>
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thyroid antibodies, or Coombs’ test), elevation of IL-6 and/or VEGF, myelofibrosis, and increased levels of megakaryocytes in bone marrow, small or unclear lymphadenopathy (> 1.5 cm in diameter) with mild accumulation of 18FDG by 18FDG-PET-CT, and most cases show mixed type, but fewer with HV type of MCD histology. Furthermore, glucocorticoid, immunosuppressive therapy, or tocilizumab may be effective as treatment for patients with this condition.

Our cases did not meet all the criteria for TAFRO syndrome, but they had the most specific findings, i.e., anasarca (pleural effusion and ascites) and thrombocytopenia, which cannot be explained due only to hyper-IL-6 syndrome. Therefore, they were similar to cases of TAFRO syndrome (Castleman-Kojima disease).

Several questions remain regarding the mechanism and etiology of TAFRO syndrome (Castleman-Kojima disease); is it different from MCD, or some part of MCD, or changes seen in a certain process in MCD (Table 1)? Furthermore, are these conditions neoplastic, autoimmune, infectious, or some other type of disease entities?

Cases of this syndrome are still rare, and it is therefore necessary to conduct multicenter clinical surveys including similar cases such as ours, to reach a consensus regarding diagnostic criteria, therapeutic strategy, and pathophysiological etiology of TAFRO syndrome (Castleman-Kojima disease).

ACKNOWLEDGEMENT

We thank all participants of the TAFRO syndrome (Castleman-Kojima disease) meeting including for critical discussion.

The sources of support in the form of grants:

This work was partially supported by the Research Program of Intractable Disease (Research on IgG4-related disease) provided by the Ministry of Health, Labor, and Welfare (MHLW) of Japan, and by the Japanese Ministry of Education, Culture, Sports, Science and Technology (Grant Nos. 13557160, 15024236, 15390313, and 13877075 to Kanazawa Medical University (Grant No. K 2011-7) and Grant for Project Research from High-Tech Research Center of Kanazawa Medical University (Grant No. H 2011-11).

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