RESOLUTION OF RETINAL BLEEDING AND EXUDATIVE RETINAL DETACHMENT AFTER CHEMOTHERAPY WITH MELPHALAN IN MULTIPLE MYELOMA

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ABSTRACT

Introduction: Multiple myeloma is the second most common hematologic cancer. It is characterized by the clonal proliferation of malignant plasma cells and associated organ dysfunction. Ophthalmic manifestations include hyperviscosity retinopathy and retinal detachment. The aim of this study is to report retinal manifestation in Multiple Myeloma.

Methods: A man 45 years old, complained loss of vision on both eyes especially left eye over 1 month. He also feels weakness and 3 times spontaneous nose bleeding. His visual acuity is 6/12 on the right and 2/60 left eye with Retinal Bleeding and Retinal Detachment. His Laboratory examination result increase in WBC 65.91 x 10^3/µL (Neutrophil, Monocyte, Basophil and Lymphocyte) decrease RBC 3 x 10^6/µL, HGB 6.85 g/dL, no normal PLT 193.60 10^3/µL, Normal Bleeding and Clotting Time and increase ESR 140 mm/hour. Multiple myeloma was diagnosed by Internal Department with bone marrow biopsy and increase of Gamma Globulin.

Results: The patient was treated with melphalan 3 tablet 2 times a day and prednison 4 mg 3 times a day for 5 days every 28 days. After 8 month the VA 6/6 on the right and 6/10 on the left with complete resolution of on the right eye and there are resolution on the left eye with fibrosis, the laboratory examination normal WBC 3.8 x 10^3/µL with Neutrophil 59.3%, Monocyte 6.1%, Eosinophil 6.3%, Basophil 0.3% and Lymphocyte 28.0%, normal RBC 4.6 x 10^6/µL, slightly decrease HGB 12.8 g/dL, normal PLT 177.0 x 10^3/µL and decrease Gamma Globulin.

Discussion: This case of bilateral Retinal Bleeding and Exudative Retinal Detachment are remarkable for both its presentation and response to melphalan therapy. Treatment with melphalan and prednison alone was sufficient to clear the Retinal Bleeding and restoration of Exudative Retinal Detachment.

Conclusion: Complication of Multiple Myeloma can caused organ disfunction such as retinal bleeding and exudative retinal detachment. Early Diagnosis is very crucial. In our case combination chemotherapy with Melphalan and Prednison showed clinically significant resolution. Workship Internal Department and Ophthalmology Department are very important for management and monitoring ophthalmology manifestation in multiple myeloma.

Keywords: Multiple Myeloma, Retinal Bleeding, Exudative Retinal Detachment, Melphalan Chemoteroaphy.


INTRODUCTION

Multiple myeloma is the second most common hematologic cancer. It is characterized by the clonal proliferation of malignant plasma cells in the bone marrow microenvironment, monoclonal protein in the blood or urine, and associated organ dysfunction.1 Multiple Myeloma usually occurs between 40 and 70 years of age.2 Multiple myeloma occurs slightly more frequently in men than in women (1.6:1).3,4
Nearly 74,800 Americans currently have MM, and an estimated 10,700 will die from the disease in 2012.\(^5\) Multiple Myeloma accompanied by nonspecific symptoms such as generalized fatigue, weight loss, and bone pain on movement. Osteolytic lesions and pathological fractures may be seen on skeletal survey.\(^2\) Multiple Myeloma can cause many complications, including Calcium elevation (hypercalcemia), Renal (kidney) dysfunction, Anemia and Bone disease. This constellation of signs and symptoms is commonly referred to as “CRAB”.\(^5\) Multiple Myeloma can present as pathology in practically all the orbital and ocular tissues. Early identification and early treatment is necessary to prevent irreversible visual loss. Ophthalmic manifestation in multiple myeloma is rare. Retinal vascular disorders have been reported in up to 66% of patients with multiple myeloma. Retinal features of hyper viscosity include venous dilatation, segmentation and tortuosity, superficial and deep hemorhages and detachment of both the sensory retina and retinal pigment epithelium.\(^6\) The two pathophysiological processes that give rise to a majority of the ophthalmic signs are tissue infiltration by plasma cells and hematological abnormalities.\(^2\) Laboratory tests characteristically reveal a normocytic normochromic anemia, raised erythrocyte sedimentation rate, hypercalcemia, renal dysfunction, Bence Jones proteinuria and monoclonal globulin spike on serum electrophoresis. Bone marrow biopsy usually confirms the diagnosis by showing an uncontrolled proliferation of plasma cells.\(^7\)\(^8\)

There is no one standard treatment for MM, and choice of therapy depends on many factors, including physical exam and laboratory test results, the specific stage or classification of the disease, age and general health, symptoms, presence of complications and prior treatment. Commonly used regimens for initial therapy include traditional agents such as corticosteroids (e.g., dexamethasone) and alkylating agents such as melphalan.\(^5\)

METHOD

This is about a case report. A man 45-year-old, complained loss of vision on both eyes especially left eye over 1 month. He also feels weakness and 2 times spontaneous nose bleeding. His visual acuity is 6/12 on the right and 2/60 left eye. Anterior Segment examination in normal limit. Dilated Fundus examination showed dilatation and tortuosity vein (sausage shape) with dot, blot, flame shape, roth spot and decrease macular reflex on the right eye. We found vitreous hemorrhage with dilatation and tortuosity vein (sausage shape), subhyaloid bleeding and decrease macular reflex on the left eye. Ocular Computed Tomography was done and showed increase in Central Retinal Thickness (CRT) 317µm on the right eye and 482 µm on the left.

We diagnosed patient with Retinal Bleeding and Eksudatif Retinal Detachment and Vitreous Hemorrhage on the left eye. His Laboratory examination result RBC 3.25 x 10^6/µL (4.4-5.9), Hemoglobin 7.8 g/dL (13.2-17.3), Hematocrit 26% (40-52), MCV 80 fl (80-100), MCH 24 pg (26-34), MCHC 30 g/dL (32-36), PLT 171 x10^9/µL (150-440), WBC 8.7x 10^3/µL (3.8-10.6) , Basophil 0.3% (0-1) Absolut 0.03x 10^3/µL, Eosinophil 4.4% (2-4) Absolut 0.38x 10^3/µL, Neutrophil 43.2% (50-70) Absolut 3.74 x 10^9/µL, Lymphocyte 45.2% (25-40) Absolut 3.92 x 10^3/µL, Monocyte 6.9% (2-8) Absolut 0.60 x 10^3/µL, ESR 140 mm/hour (0-15), BT 1.5 minute (1-6), CT 9.0 minute (5-15), PT 24.2 second (10.8-14.4), APTT 43.2 second (26.4-37.6), INR 2.4, SGOT 18 U/L (<33), SGPT 12 U/L (<50), Ureum 14 mg/dL (6-20), Creatinin (Cr) 1.18 mg/dL (0.7-1.2), Chlorida (Cl) 110 mmol/L (98-107), Natrium( Na) 137 mmol/L (136-145), Kalium (K) 4.1 mmol/L (3.5-5.1), IgM, IgG anti Toxoplasma and anti CMV were negative, IgM anti HSV1 was negatif, VDRL and TPHA were nonreactif, Fasting Blood Sugar 94 mg/dL (<100), 2hPP Blood Sugar 106 mg/dL (<140), Uric Acid 12.0 mg/dL (<7.0).

We consulted this patient to Internal department but unfortunately the patient came to emergency room with spontaneous nose bleeding on the next day. His Laboratory result increase in WBC 65.91x10^9/µL (Neutrophil, Monocyte, Basophil and Lymphocyte) decrease RBC 3x10^6/µL, HGB 6.85g/dL, normal PLT 193.60x10^3/µm. The patient was consulted to Hemato Oncology and was planned to bone marrow aspiration with diagnosed suspect Acute Myeloid Leukemia.

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Patient was consulted to our division. On our examination we found visual acuity is 6/12 on the right and 5/60 left eye. Anterior Segment examination in normal limit. Dilated Fundus examination showed dilatation and tortuosity vein (sausage shape) with dot, blot, flame shape, Roth spot and decrease macular reflex on the right eye. We found decrease on vitreous hemorrhage with dilatation and tortuosity vein (sausage shape), subhyaloid bleeding and decrease macular reflex on the left eye. We planned to observation and waited for the bone marrow aspiration resulted.

Bone marrow aspiration was done 3 days after the patient was consulted to our division. The resulted are Multiple Myeloma with Hypercelluler, Erytroid System Activity, Myeloid System Activity, Megakaryocites System Activity decrease and 40% Plasma Cell Infiltration. Electrophoresis and bone survey was planned.

We found increased in Gamma Globulin 8.62 g/dL (0.9-1.5), 71.8% (12.1-21.1) with total protein 12 g/dl. Bone Survey examination showed didapatkan Lytic soliter lesion on left frontal bone. Multiple Myeloma Stage IIIA was diagnosed by hemato-oncology division and the patient was given Melphalan 6 mg two times per day and prednisone 20 mg three time per day for 5 days and repeated for 28 days.

RESULT

After 10 cycles of melphalan therapy, patient came to eye polyclinic. On our examination we found visual acuity is 6/6 on the right eye and 6/10 with pinhole not improve on the left. Dilated funduscopy examination we found complete resolution on the right eye with no bleeding and tortuosity. Macular reflex are positive. On the left eye we found weiss ring and fibrosis in the vitreous, with no subhyaloid bleeding, but unfortunately the macular reflex still decreased.

We took fundus photograph and OCT to the patient and showed normal fundus image on the right eye and Weiss ring with fibrotic lesion on the left. Ocular Computed Tomography showed Central Retinal Thickness (CRT) on the right eye was normal (224 µm) and 324 µm on the left.
Multiple myeloma is characterized by bone pain, skeletal punched out lesions, anemia, renal insufficiency, hypercalcemia, and increased susceptibility to infections. Ocular involvement with multiple myeloma can be found throughout the eye and visual pathways as well as the orbit, although rarely are these the presenting signs. Specifically, the retina may have dilated and tortuous veins, retinal hemorrhages, Roth spots, vitreous hemorrhages, cotton wool spots and exudates, and central and branch retinal vein occlusions. Uncommonly, retinal and choroidal detachments can be seen in multiple myeloma.9,11,12

This case is unusual in that the initial presentation of the patient was significant decrease in visual acuity 6/12 on the right and 2/60 left eye. Posterior examination we found with roth spot and intraretinal bleeding with venous dilatation and turtuosity. Macula reflex decrease on the right and hazy (bleeding) on vitreous with subhyaloid bleeding, intraretinal bleeding, venous dilatation and turtuosity. Macula reflex decrease. OCT macula show CRT on RE 317µm and LE 482 µm.

Retinal bleeding is caused by hyperviscosity syndrome of Multiple myeloma. The shear wall stress due to increased blood viscosity is greatest on vessels with the smallest cross-sectional area and highest flow rates, providing an explanation for localization of haemorrhage to the arteriolar beds. Retinal vein swelling and haemorrhage is probably secondary to compression of the central optic vein due to retinal artery engorgement as a result of increased viscosity.13 Exudative retinal detachment that happen in this case is unusual. Dogan et al speculate that Ig accumulation in the subretinal space increases the oncotic pressure, while fluid accumulation in the subretinal area separates the RPE from the photoreceptors.11

Patient feel weakness and history of spontaneous nose bleeding was positive. Laboratory Examination show increase ESR 140 mm/hour, decrease HGB 6.85 g/dL. Renal Function in normal limit. Electrophoresis show increase in Gamma 8.62 g/dL (0.9-1.5), 71.8% (12.1-21.1). Skull AP Lateral on the bone survey showed Soliter Lytic Lession in Left Frontal. Bone marrow aspiration result Plasma Cell Infiltration 40% leading to Multiple Myeloma

Multiple Myeloma is classified as asymptomatic or symptomatic, depending on the absence or presence related organ damage.1 Laboratory tests characteristically reveal a normocytic normochromic anemia, raised ESR, hypercalcemia, renal dysfunction, Bence Jones proteinuria and monoclonal globulin spike on serum electrophoresis. Bone marrow biopsy confirms the diagnosis (uncontrolled proliferation of plasma cells).2
Our patient was diagnosed Multiple Myeloma Stage IIIA by hemato-oncology division and the patient was given Melphalan 6 mg two times per day and prednisone 20 mg three time per day for 5 days and repeated for 28 days.

Since the 1960s, melphalan and steroid have been the standard of therapy. Chemotherapy consisted of oral melphalan (8 mg/m) on days 1–4 and oral dexamethasone (40 mg/day) on days 1–4 and days 9–12. The cycles were repeated every 28 days for a total of 12 cycles or until intolerance or disease progression was observed.¹³⁻¹⁴ This case of bilateral retinal bleeding and exudative retinal detachment are remarkable for both its presentation and response to therapy. Our case is also remarkable in that there was a favorable response to therapy by way of visual acuity improvement, resolution on retinal bleeding and exudative retinal detachment after 10 cycles therapy with melphalan and prednisone although there is PVD complete and fibrosis after vitreous hemorrhage on the left eye.

CONCLUSION
Complication of Multiple Myeloma can caused organ disfunction such as retinal bleeding and exudative retinal detachment. Early Diagnosis is very crucial. In our case combination chemotherapy with Melphalan and Prednison showed clinically significant resolution. Workshop Internal Department and Ophthalmology Department are very important for management and monitoring ophthalmology manifestation in multiple myeloma.

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REFERENCES