Pulmonary extramedullary hematopoiesis involving the pulmonary artery

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Abstract

Extramedullary hematopoiesis (EMH) occurs as a complication of hematologic disorders such as myelofibrosis, sickle cell anemia and thalassemia. The extramedullary tissue usually involves liver, spleen and lymph nodes, less frequently the chest. We present a recent case of a man with myeloproliferative neoplasm who developed pulmonary hemorrhage secondary to EMH in the lung and pulmonary artery. Radiation therapy was considered the best approach, but it didn’t work and the patient died a week after radiation therapy was completed. We also review herein the present literature.

Introduction

Extramedullary hematopoiesis (EMH) occurs as a complication of hematologic disorders such as myelofibrosis, sickle cell anemia, and thalassemia. The extramedullary tissue usually involves the liver, spleen and lymph nodes. Although it has been described as occurring at almost any organ/site in the body, it has been infrequently reported in the chest. When described in chest, it has been generally detected as paravertebral masses seen on chest x-ray. It is important to make this diagnosis as EMH can be associated with intrathoracic cord compression, and rarely fatal hemothorax. It is amenable to therapy with both hydroxyurea and radiation.

Case Report

A 53-year-old Caucasian man presented with a year history of myelodysplastic syndrome/myeloproliferative neoplasm (MDS/MPN) and narrow fibrosis (negative JAK-2 V617F point mutation). He was seen with epistaxis and hemoptysis of two days duration. He had been on ruxolitinib for six months with improvement in appetite and decreased pain secondary to an enlarged spleen. He was being considered for allogeneic stem cell transplantation. Physical examination was significant for a fever of 101.3°F, fine crackles at the right base and an enlarged spleen. Laboratory analyses showed a white blood cell count of 27,500/mm³, hemoglobin level of 7.8 g/dL, hematocrit of 27% and platelet count of 22,000/mm³. Peripheral blood smear review showed markedly increased monocytes many of which appeared atypical. Computed tomographic (CT) scan of the chest with contrast showed a right pulmonary artery filling defect which extended into the right middle lobe (Figure 1A). Enlarged subcarinal, aortopulmonary lymph nodes, splenomegaly and a right liver mass were also seen. Given his long-term smoking history, there was concern that a second malignancy might be present. A recent biopsy of the liver mass had EMH. The differential diagnoses of the right middle lobe and pulmonary artery mass included thromboembolic disease, primary lung malignancy, sarcoidosis, pneumonia or EMH. An endobronchial ultrasound guided aspiration of the subcarinal lymph node and right pulmonary artery mass showed findings consistent with EMH (Figure 1B). A bone marrow biopsy was performed which showed hypercellular bone marrow (>90%) with decreased megakaryocytes, erythrocytosis and left shifted granulopoiesis with 5% blasts, a peripheral monocytosis, and moderate marrow fibrosis consistent with chronic myelomonocytic leukemia-2. He received 14 Gy of the planned 20 Gy external beam radiation therapy in 7 fractions using APPA approach with 10× energy to the right lung mass as outpatient. Ruxolitinib was stopped due to thrombocytopenia (platelet count of 48,000/mm³, PT –15 sec and INR – 1.5). A week later he presented with a hematoma of his right thigh, severe anemia (hemoglobin 5.5 gm/dL), elevated white blood cell count (89,700/mm³) with predominant monocytosis. He was started on hydroxyurea and received a single dose of decitabine for presumed acute leukemia. The patient suddenly became hypoxemic. Chest x-ray showed right sided pleural effusion. A diagnostic thoracentesis revealed bloody pleural fluid. Cytology of the pleural fluid showed hematopoietic cells similar to the ones noted in the pulmonary artery aspirate specimen. At that time his platelets were 76,000/mm³. The patient was deemed to be inoperable given the multiple comorbidities and he died a day later. Autopsy was not performed.

Discussion

Extramedullary hematopoiesis is a compensatory hematopoietic tissue expansion usually involving the reticulo-endothelial system. EMH and myeloid metaplasia have been interchangeably used in the past and refer to the similar pathological process. Whether EMH represents differentiation of embryonic stem cells residing within the vessel wall or implantation of hematopoietic tissue escaping from the bone marrow remains uncertain. Pulmonary EMH (PEMH) has been rarely reported. The majority of the pulmonary EMH masses are asymptomatic. However patients sometimes present with hemoptysis, acute or progressive dyspnea or chest pain. Patients can sometimes present with life threatening complications such as massive pleural effusion, hemothorax, chylothorax, or spinal cord compression (posterior mediastinal EMH). Differentiation between thromboembolic disease and extramedullary hematopoiesis in such cases can be very critical as the treatments strategies are different for each. Several other non-infectious pulmonary complications of MDS/MPN have been described in the literature. Hematopoietic tissue is known to be sensitive to low doses of radiation therapy. The role of radiation therapy in treating patients with EMH causing spinal cord compression has been well established. Radiation therapy to the lung has been successfully used to treat EMH involving the lung parenchyma. The median suggested dose of radiation is 1.25 Gy. Combined modality treatment with radiation and surgery perhaps would be a better approach reserving surgery for only emergent management. Table 1 outlines the published cases of pulmonary EMH, the respective interventions utilized and the corresponding treatment outcomes.
In our patient several factors could have contributed to the pulmonary hemorrhage including the underlying hematological condition leading to thrombocytopenia and coagulopathy. The transbronchial biopsy procedure could have caused the hemothorax as these hematopoietic masses are very likely to bleed upon intervention. However the 14 day time lag between the procedure and bleeding makes it less likely. Radiation therapy to the hematopoietic mass tissue in the lung could have potentially lead to necrosis of the tissue thereby causing hemothorax. Review of the other pulmonary EMH case studies suggests that radiation therapy to the lung has led to the best outcomes.

Table 1. Reported cases of patients with underlying myeloproliferative disease presenting with pulmonary involvement by extramedullary hematopoiesis, respective interventions and their outcomes. Diagnosis of pulmonary extramedullary hematopoiesis made ante mortem.

| Case             | Underlying disease                     | Presentation                             | Treatment for EMH                          | Outcome                          |
|------------------|----------------------------------------|------------------------------------------|--------------------------------------------|-----------------------------------|
| Pinato et al.⁴   | Myelodysplasia                          | Progressive dyspnea                      | None                                       | Unknown                           |
| Chute et al.⁵    | Sickle cell trait/β-thalassemia         | Hypoxia                                  | Thoracostomy                               | Died from hemothorax             |
| Ozbudak et al.⁶  | Myelofibrosis                           | Dyspnea and hemoptysis                   | Prednisone, hydroxyurea, busulfan          | Minimal improvement; died of MI after 2 mths |
| Kuplerschmid et al.³ | Myelofibrosis                        | Dyspnea                                  | Thoracostomy + radiation 1.4 Gy in 10 fractions | Died of pneumonia               |
| Ghosh et al.¹⁰   | None                                    | Progressive dyspnea                      | Thoracostomy with talc pleurodesis + radiation 2 Gy in 4 fractions | Complete resolution               |
| Koch et al.¹⁴    | 2 cases: agnogenic myeloid metaplasia   | Dyspnea, orthopnea, edema weight gain    | 1 Gy in 1 fraction of radiation; 1.5 Gy in 10 fractions | Complete resolution               |
| Weinschenker et al.¹⁵ | Myelofibrosis                     | Dyspnea                                  | Whole lung radiation 200 Gy/4 fractions    | Complete resolution               |
| Runii et al.¹⁶   | Myelofibrosis                           | Progressive dyspnea                      | Hydroxyurea                                | Complete resolution after 3 mths  |
| Ueno et al.¹⁷    | Myelofibrosis                           | Fever and fatigue                        | Steroids started day 90                    | Died 10 days later               |
| Yusen et al.¹⁸   | Myelofibrosis                           | Dyspnea, cough, fevers and night sweats  | 2 days interferon and supportive care      | Died due to respiratory failure  |
| Asakura et al.¹³ | Myelofibrosis; agnogenic myeloid metaplasia | Dyspnea; dyspnea and fatigue               | Steroids and diuretics; unknown            | Died 6 mths later; died 13 mths later |
| Glew et al.²⁰    | Myelofibrosis                           | Fevers                                   | None                                       | Died 5 months later              |

EMH, extramedullary hematopoiesis.

Figure 1. A) Coronal view of post contrast computed tomography imaging of the chest with arrow pointing towards the filling defect within the right pulmonary artery. B) High power view of Hematoxylin and Eosin stain of the endobronchial ultrasound guided aspirate of the right pulmonary artery thrombus showing extramedullary hematopoiesis with immature myeloid cells (box arrow) and focally increased areas of blasts (line arrow).

In conclusion, we present a patient with myeloproliferative neoplasm who developed pulmonary hemorrhage secondary to EMH in the lung and pulmonary artery. Although rare, presentation of hemoptysis in a patient with underlying myeloproliferative neoplasm should prompt clinicians to consider EMH involving the pulmonary artery in the differential diagnosis. Early recognition of the diagnosis remains important. Radiation therapy to the lung mass (or the whole lung if pulmonary hypertension is diagnosed) can be considered as the best approach if patient is hemodynamically stable recognizing that it may not always work.

Conclusions

In summary, we present a patient with myeloproliferative neoplasm who developed pulmonary hemorrhage secondary to EMH in the lung and pulmonary artery. Although rare, presentation of hemoptysis in a patient with underlying myeloproliferative neoplasm should prompt clinicians to consider EMH involving the pulmonary artery in the differential diagnoses. Early recognition of the diagnosis remains important. Radiation therapy to the lung mass (or the whole lung if pulmonary hypertension is diagnosed) can be considered as the best approach if patient is hemodynamically stable recognizing that it may not always work.
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