Extramedullary Multiple Myeloma With Pleural Involvement: A Rare Clinical Entity

Mieloma múltiple extramedular con afectación pleural: una rara entidad clínica

To the Editor,

Thoracic involvement has been described in the course of multiple myeloma (MM), in the form of bone lesions, extraosseous plasmacytomas, pulmonary infiltration and, exceptionally, pleural effusion (PE). Myelomatous pleural effusion (MPE) occurs in only 1% of cases of PE in patients with MM, and is associated with a poor prognosis (median survival of 1.5–3 months after appearance). We report a patient with MM with extramedullary thoracic involvement who developed secondary MPE.1,2

This was a 67-year-old man with a history of quiescent MM IgA kappa diagnosed in 1998 who attended the emergency room of our hospital in November 2017 due to fever, general malaise, asthenia, and cough. The patient had presented several plasmacytomas in the right chest wall (Fig. 1A) and the spinal canal in 2012 and 2014, but without PE, and was treated with different therapeutic options (chemotherapy, autologous hematopoietic stem cell transplantation, local radiation therapy), resulting in complete resolution of the lesions. In 2015, he presented a new extramedullary relapse in the form of a paravertebral mass in the right hemithorax, which was treated with chemotherapy. Given the patient’s clinical picture and radiographic findings (not present in previous studies) of right PE observed in the emergency department (Fig. 1B), we decided to admit him and perform a positron emission tomography/computed tomography (PET/CT).

Fig. 1. (A) Axial CT scan of the chest (mediastinum window) performed in 2012 that identifies a solid mass in the upper right hemithorax (asterisk) infiltrating the chest wall and penetrating the spinal canal through the right T2–T3 intervertebral foramen (arrow). (B) Posteroanterior chest X-ray performed in November 2017 showing right pleural effusion for the first time in this patient. (C) Chest axial image of a PET/CT scan performed in November 2017 which identifies bilateral opacities of pneumonic aspect (arrows) and right pleural effusion (asterisk). (D) Thoracic axial image of another PET/CT scan performed in March 2018 which shows the disappearance of pneumonic opacities and the presence of minimal pleural effusion.

PET/CT showed bilateral hypermetabolic pneumonic pulmonary opacities and confirmed a moderate amount of right PE (Fig. 1C). Serosanguineous pleural fluid with the following characteristics was obtained: pH 7.43, glucose 104 mg/dl (serum glucose 91 mg/dl), lactate 3.2 mmol/l, pleural fluid protein/serum protein ratio: 0.70, pleural fluid LDH/serum LDH ratio: 2.37, LDH pleural fluid: 524, hematocrit <15%, lymphocytes 28.7%, neutrophils 0.0% (criteria for lymphocytic exudate) and negative microbiological tests, thus ruling out PE due to infection. Flow cytometry of pleural fluid detected 60% large neoplastic plasma cells (plasmablasts), CD 138+ and CD 56−, which confirmed the malignant nature of the PE (myelomatous). Jugal mucosa biopsy ruled out the presence of amyloid deposits. The patient was initially treated with antibiotics (pipericillin–tazobactam) and subsequently underwent right pleural drainage followed by chemical pleurodesis with talc, local radiotherapy and chemotherapy (pomalidomide–dexamethasone–cyclophosphamide), presenting excellent clinical and radiological progress (Fig. 1D).

During the course of MM, 15%–30% of patients may develop extramedullary involvement.1 The pleural cavity is an unusual site for MM recurrence; in fact, MPE occurs in only in 1% of cases of PE in patients with MM. In a recently published series, MPE represented only 0.6% of malignant PEs.2,3 Diagnostic criteria for MPE are: (1) presence of atypical plasma cells in pleural fluid (neoplastic plasma cells or a monoclonal component); (2) pleural biopsy consistent with neoplastic plasma cells, or (3) demonstration of monoclonal proteins in the pleural fluid by electrophoresis. In doubtful cases, flow cytometry helps establish the immunophenotype of neoplastic plasma cells compared to that of reactive cells. When a patient with MM develops PE, it is important to rule out common etiologies, such as paraneoplastic PE, heart failure, kidney failure, and amyloidosis. The latter may cause PE due to cardiac (heart failure), renal (nephrotic syndrome), liver (ascites), or pleuropulmonary involvement.4 MPE can be caused by an abnormal proliferation of plasma cells from an extramedullary plasmacytoma of the chest wall, invasion from an adjacent bone lesion, or direct invasion of the pleura by myeloma.5 While various treatments are available for MPE (chemotherapy, therapeutic thoracentesis, chest drain, or pleurodesis), there is no consensus about how we should manage these patients. Extramedullary involvement is associated with an adverse prognosis, especially when it is recurrent MM. Pleural infiltration is usually fatal, with a median survival of 1.5–3 months. Therefore, more aggressive chemotherapy regimens can be indicated in MM with involvement of the pleural cavities. Our patient responded well to multimodal treatment consisting of a combination of radiation therapy, chemotherapy, and chemical pleurodesis, achieving clinical remission that lasted 6 months after diagnosis of the MPE.

This case reminds us that we must investigate all the causes of PE in patients with a history of MM and that, although the incidence of MPE is low, it must be taken into account as a diagnostic possibility. Its grim prognosis and aggressive natural course require us to make a quick and appropriate diagnosis in order to start treatment as soon as possible.

References

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Overnight Change in Urinary Prostaglandin and Thromboxane in Obstructive Sleep Apnea

Cambio en la prostaciclina y el tromboxano urinario durante la noche en la apnea obstructiva del sueño

Dear Editor,

Obstructive sleep apnea (OSA) is a common disorder1 eliciting sympathetic alterations and intermittent hypoxia (IH) resulting in oxidative stress and inflammation. As a result, OSA has been linked to enhanced cardiovascular (CV) disorders and hypercoagulability,2 endothelial function, intima-media thickness, and high blood pressure.3 Prostanoids (PG) are products of arachidonic acid catabolism by cyclooxygenase (COX) isoenzymes COX-1 and COX-2. Among PG, Thromboxane (TXA2) and Prostacyclin (PGI2) are known for their role as regulators of vascular tone, remodeling and angiogenesis. TXA2 is mainly generated by platelets through COX-1 and quickly metabolized into Thromboxane B2 (TXB2). TXA2 induces platelet activation, vasoconstriction, and vascular smooth muscle cell proliferation. On the other hand, PGI2 mostly depends on endothelial COX-2 and prostacyclin synthase enzymes. PGI2 is metabolized into 6-keto Prostaglandin F1α (6-ketoPGF1α). PGI2 inhibits platelet aggregation and vasoconstriction. Therefore, TXA2 and PGI2 have antagonist properties and are both excreted in urine and plasma.4 Aspirin (acetyl salicylic acid, ASA) is a non-selective COX inhibitor with beneficial anti-thrombotic effects by inhibiting the release of TXA2. Although ASA can also inhibit the synthesis of PGI2 which has anti-thrombotic effect, more pronounced inhibition of TXA2 versus PGI2 has been detected in humans after low-dose ASA.4

Recently, our group reported that pre-atherosclerotic aorta remodeling induced by chronic IH mimicking OSA in mice can be prevented by ASA treatment.5 We here hypothesize, that ASA preventive effects are related to its capacity to inhibit COX-1 and COX-2 pathways. Thus, the aim of the present study is to characterize TXA2 and PGI2 overnight change according to OSA severity, and to investigate the effect of ASA treatment in this overnight change.