cT4NxM1b). The patient received local palliative radiation therapy and chemotherapy with platinum/pemetrexed (Fig. 1B1 and B2). The third case was a 73-year-old man receiving active treatment guided by sensitivity testing results for documented Mycobacterium xenopi infection. Positron emission tomography showed a lesion measuring 40 × 25 mm with central cavitation in the left upper lobe (SUVmax 28.38) (Fig. 1C) and a hypermetabolic focus located in the left ilioptic muscle with SUVmax 13.43, suggestive of MSM.

In all 3 cases, histological specimens were obtained for characterization, and the results were consistent with high grade undifferentiated tumor, striated muscle infiltrated with adenocarcinoma, and squamous carcinoma, respectively, all originating in the lung. The clinical progress of the patients differed: death 2 weeks after diagnosis, pain control, and reduced tumor size (Fig. 1B2) after targeted oncological treatment; clinical stabilization was achieved in the last 2 cases described.

Given the low prevalence of MSM, a detailed differential diagnosis that includes the more common malignant and benign entities (sarcomas, primary muscle lymphomas, and myxomas/hemangiomas) must be made. Although no clinical guidelines are available for the specific management of MSM, treatment is based on general oncological principles guided by clinical picture, site, and life expectancy, and approaches include observation, surgical excision (persistent solitary lesions after a period of remission), chemotherapy and radiation therapy (useful for pain control and for the reduction of tumor size).2–5

The correct identification of MSM in LC patients is essential for clinical management and prognosis. For this reason, the possible neoplastic etiology of any muscle lesion, whether symptomatic or not, detected in LC patients must be evaluated with combined radiological procedures and histological confirmation of the lesion.

Hyponatremia in COPD: A Little Known Complication

La hipoventruria en la EPOC, una complicación poco conocida

To the Editor,

Chronic obstructive pulmonary disease (COPD) is a chronic inflammatory disease, and in some patients, extrapulmonary manifestations can worsen prognosis.1

COPD patients often have other concomitant diseases, particularly cardiovascular conditions, and it has been postulated that COPD plays a significant role in the pathogenesis of these processes.2

Hyponatremia developing during hospitalization for a COPD exacerbation is relatively common, and is associated with a poorer clinical course.3 Low sodium in blood may be a sign of water retention due to other comorbidities, such as heart or kidney failure, drug treatments, adrenal insufficiency after withdrawal of corticosteroids, or syndrome of inappropriate ADH secretion (SIADH). Diseases that may present with SIADH include lung infections (pneumonia, pulmonary abscess, tuberculosis, aspergillosis), asthma, COPD, lung tumors, cystic fibrosis, and acute respiratory failure.4 Hypoxia is associated with ADH secretion,5 but hypercapnia is more commonly associated with this phenomenon.

Both in stable COPD and during exacerbations, hyponatremia, due to its prevalence, its impact on prognosis, and varying etiologies (which can coexist), is a challenge for the clinician and requires appropriate follow-up and treatment. Although the relationship between COPD and SIADH is often mentioned in the literature, we did not find any references to hyponatremia caused by SIADH in COPD (Medline and Pubmed searches, keywords: SIADH and COPD).

For this reason, we believe that our report of a patient with SIADH due to COPD is of interest and provides a good illustration of certain aspects of the differential diagnosis and treatment.

Our patient was an 84-year-old man, active smoker, with a history of benign prostate hypertrophy and exacerbator phenotype COPD with emphysema, and very severe obstruction (FEV1 27%), receiving treatment with salasosin, omeprazole, and glycopyrroinum/indacaterol. He presented with intense dyspnea and cough with greenish expectoration. The only finding of note on physical examination was the presence of disperse rhonchi in both hemithoraxes and the absence of edema or signs of fluid overload. O2 saturation was 89% with home oxygen therapy at 21 per minute (lpm). Clinical laboratory tests showed microcyt anemia (hemoglobin 10 g/dl), plasma sodium 111 mEq/l (normal value [NV]: 135–155 mEq/l), plasma osmolality 229 mOsm/l (NV: 280–300), PCR 64 mg/l (NV: 0–5), normal creatinine levels, urinary sodium 76 mEq/l (NV: 54–150), and urine osmolality 273 mOsm/l

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Footnotes

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It (particularly urine) showed saline. Treatment with intravenous saline, restricted to 123 mmol/l, was necessary, and normal saline was used instead of saline. The patient's sodium levels in blood did not normalize after completion of COPD treatment and no further changes were observed on physical examination. Repeat continuous laboratory tests showed plasma sodium 124 mmol/l, plasma osmolarity 247 mOsm/l, urine sodium 62 mmol/l, and urine osmolarity 372 mOsm/l, with normal thyroid and adrenal function. After ruling out other possible causes, a diagnosis of SIADH due to an exacerbation of emphysematous phenotype COPD was given, and treatment began with tolvaptan 15 mg/24 h, leading to normalization of blood sodium in 4 days. The dose was reduced to 7.5 mg/24 h, and sodium levels in blood were normal in subsequent visits.

SIADH is characterized by the sustained release of arginine-vasopressin (ADH) in the absence of the usual stimuli, particularly hyperosmolarity and hypovolemia. Diagnosis is based on hyponatremia, plasma hypoosmolarity, urinary sodium >40 mmol/l, and normal osmolarity >100 mmol/kg, after ruling out processes that involve loss of effective blood volume (heart failure, cirrhosis with ascites, etc.), and normal renal, adrenal, and thyroid function has been reported.

Symptoms are non-specific, and can range from nausea, dizziness, general malaise, agitation and confusion to seizure or coma in cases of sudden onset or very low blood sodium levels. SIADH is the most common cause of hyponatremia, and its heterogeneous etiology includes mainly important infections, drugs, tumors (particularly small cell lung carcinoma), COPD, and asthma. Hyponatremia due to SIADH occurs quite frequently in small cell carcinoma, sometimes as a first manifestation and is associated with decreased survival.

Both COPD and small cell lung carcinoma are a cause of SIADH and strongly associated with smoking, so blood sodium must also be monitored in patients with COPD and a heavy accumulated consumption of tobacco.

Treatment of hyponatremia due to SIADH differs, depending on two different clinical scenarios. In acute situations with moderate/severe symptoms (sleepiness, confusion, stupor, respiratory distress), and plasma Na+ <120 mmol/l, treatment should begin with 3% hypertonic saline. The initial infusion rate will be 0.5 mg/kg/h or 1–2 ml/kg/h, depending on neurological signs. Treatment with tolvaptan (a selective vasopressin V2-receptor antagonist) may be considered, depending on progress. When SIADH develops with mild hyponatremia, water restriction and furosemide should be considered, and in patients who are not candidates for these measures or whose clinical symptoms persists, the use of tolvaptan is recommended.

COPD exacerbation is a cause of SIADH, as demonstrated by Chanela et al. However, in our patient, hyponatremia persisted for almost 10 days after resolution of the infectious COPD exacerbation, and in the follow-up visit the patient continued to require tolvaptan to maintain normal sodium levels. A more typical course in SIADH due to COPD exacerbation would have been transient hyponatremia that normalized after resolution of the exacerbation.

We believe that this case of SIADH associated with a COPD exacerbation illustrates the need to include blood sodium monitoring in the management of these patients, in order to detect and reduce the morbidity and mortality of this fluid-electrolyte imbalance. It is also important to remember that this syndrome can, albeit rarely, be the first manifestation of lung cancer.

References


Response to Omalizumab in a Patient With Chronic Eosinophilic Pneumonia and Poor Response to Corticosteroids

Respuesta a omalizumab en paciente con neumonía eosinófila crónica y mala respuesta al tratamiento con corticoides

To the Editor,

We report the case of a 55-year-old woman, former smoker since September 2012 (18 pack-years), with a history of allergic rhinoconjunctivitis and bronchial asthma. No infiltrates were observed in a chest X-ray performed in 2012. Skin prick tests showed sensitization to cat dander and grass and olive pollen.

She was admitted in 2014 for an acute episode of dyspnea, breath sounds, and non-productive cough without fever. Chest X-ray revealed an alveolar-interstitial pattern predominantly in the lung bases. Chest computed tomography showed bilateral patchy ground glass infiltrates. Bronchoscopy was performed with transbronchial biopsy and bronchoalveolar lavage; neutrophils: 11%; eosinophils: 85.9%; lymphocytes: 0.6%. Biopsy showed a histological pattern of pulmonary eosinophilia suggestive of chronic eosinophilic pneumonia (CEP), with a focal pattern typical of organizing pneumonia. Lung function tests highlighted a restrictive pattern with carbon monoxide diffusion changes: FVC 1830 – 59%; FEV1 1410 – 54% and FEV1/FVC 77%; DLCO 32%; KCO 63%; TLC 61%; and RV 86%. All immunological tests were negative; IgE 493 UI/ml. In the 6-min walk test, initial saturation was 97%, distance walked 440 m, and final saturation 85%.

CEP was diagnosed, with no known occupational or environmental risks, although a relationship with the administration of anti-inflammatories (celecoxib) could not be ruled out. This drug was discontinued, and treatment began with corticosteroids, resulting in both clinical and radiological worsening.

Given the poor response to treatment with corticosteroids, a surgical biopsy was performed, and the pathology study reported squamous-like interstitial pneumonia with eosinophils, consistent with CEP. The patient was placed on the lung transplantation waiting list. Lung function tests prior to the introduction of omalizumab were: FVC 1430 – 47%; FEV1 1120 – 43%; FEV1/FVC 78%; DLCO 24%; KCO 55%; TLC 44%; RV 81%.

In a review of the literature, 2 cases1,2 of CEP with a favorable response to omalizumab were identified, so treatment began with this drug, producing clinical improvement and functional stabilization, so we decided to gradually withdraw the corticosteroids. Twenty-four months after starting omalizumab, the patient has shown a remarkable improvement in her symptoms without corticosteroids, confirming the reversal of her previous functional decline noted from the introduction of this drug (Fig. 1). The patient was taken off the lung transplantation waiting list when the adverse effects of steroid treatment had resolved. When mepolizumab came on the market, this therapeutic option was proposed to the patient, but she refused it in view of her good progress.

CEP is an idiopathic disease characterized by an abnormal accumulation of eosinophils in the interstitial and alveolar space. Standard treatment is systemic corticosteroids. Response is usually so rapid and favorable, that if no response is observed, alternative diagnoses must be considered.

Our patient presented a poor response to high-dose corticosteroids, so a surgical biopsy was performed, which confirmed the initial diagnosis. We identified 2 cases of CEP with response to omalizumab in the literature1,2 involving patients with elevated IgE and poor response to corticosteroids (or a requirement for high doses). In both cases, treatment with omalizumab reversed deterioration, and produced improvements in lung function tests.

Omalizumab is a humanized anti-IgE antibody, usually indicated for the treatment of persistent allergic asthma. Numerous effects that go far beyond IgE immunomodulation have also been described.3-11 Omalizumab reduces basophil Fc epsilon RI alpha receptor expression.2,6 It also increases a decrease in triphosphate, Th2 cytokines (IL-4 and IL-13), and chemokines (IL-8 and RANTES), irrespective of IgE levels. Similarly, it reduces cytokine expression (IL-5, IL-10, and IL-13) in dendritic cells,10 and interferes with their ability to activate CD4. It also reduces eosinophil numbers3,9 in both sputum and lung tissue, monocytes,3 fraction of expired nitric oxide (FENO),3,10 and endothelin-1 concentrations in exhaled breath condensate.10 Moreover, stimulation of IgE increases the deposit of collagen I, III, and IV, and fibronectin by the Erk1/2 MAPK

Fig. 1. Functional progress of our patient. The arrow indicates the time of starting omalizumab.

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