Pulmonary Hypertension as Presenting Manifestation of Systemic Lupus Erythematosus

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ABSTRACT

Pulmonary hypertension as presenting manifestation of SLE is extremely rare. Here we report a case of 29 year old woman presenting with progressive shortness of breath associated with pedal edema which on detailed examination found to be secondary to pulmonary hypertension. Further examination of patient fulfilled diagnostic criteria of SLE.

KEY WORDS: Systemic Lupus Erythematosus (SLE), Pulmonary Hypertension (PH).

INTRODUCTION

Pleurapulmonary manifestations of SLE present 4-5% of patients at presentation. These include pleural effusion, pulmonary infections, alveolar hemorrhages, diffuse interstitial lung disease, pulmonary embolism and pulmonary hypertension. PH is third leading cause of death in SLE⁴. PAH is defined as mean arterial pressure more than 25 mmHg on right heart catheterization. It is associated with many autoimmune diseases like SLE, systemic sclerosis, scleroderma. SLE associated PAH is a rare condition and uncommon cause of death after cardiovascular disease, infection, renal failure and non hodgkins lymphoma. It usually occurs 3-5 yrs after disease onset, it is a life threatening condition due to insidious nature of the disease.

CASE REPORT

We report a case of 29 year old female who presented to emergency department with complaints of progressive shortness of breath on exertion since 3 weeks associated with bilateral pitting type of ankle edema. History revealed that
she used to get waxes and wanes of dyspnea that interfere with daily activity and polyarthritis involving hand joints of 6 months duration. Physical examination revealed raised jvp, left parasternal heave of grade 2/3 which is sustained. Auscultation revealed pansystolic murmur of grade 3/6 and loud p2. Electrocardiogram showed poor R-wave progression and persistence of the S wave from V3 through V6. Transthoracic echo revealed dilated right atrium and right ventricle with severe tricuspid regurgitation, an estimated systolic arterial pressure of 90 mm of Hg. Right heart catheterisation revealed systolic pulmonary arterial pressure of 76 mm of Hg and mean pulmonary arterial pressure of 52 mm of Hg. Connective tissue screening revealed rheumatoid factor- positive, Anti smith -positive, Anti cardiolipin antibodies-positive, Anti dsDna - positive, ANA - moderately positive with speckled pattern. Diagnosis of SLE was made as it fulfilled American college of Rheumatology criteria.

DISCUSSION
Systemic lupus erythematosus is an autoimmune disease, events that considered in pathogenesis of SLE is autoimmune vasculitides, thromboembolic events, and increased ratio of vasoconstrictor to dilator ratio. prominent role of immunological events in endothelial damage is supported by response to immunosuppressive therapy in earliest phase of SLE(6). Thromboembolic events in SLE contribute to small vessel damage as indicated by presence of antcardiolipin antibodies 2,4. This cardiolipin antibodies can trigger inflammation by binding to endothelial cells and monocytes by specifically targeting a multiprotiensignaling complex toll like receptors and annexin which ultimately activate TNF receptors, IL-1 receptors and nuclear factor kappa light chain enhancers. These ultimately lead to irreversible changes in and around small vessels refered as vascular remodeling.

PH may occur at any stage during SLE with prevalence being 1-4% and its presentation in SLE is varied. Early therapy is essential because advanced disease may be less responsive to therapy. Based on data obtained with cardiac catheterisation, the gold standard for the diagnosis of PAH, the SLE-PAH prevalence has been estimated to range between 0.005 to 14%. This wide range of reported percentages may be due to different cut-offs (mmHg) adopted for the diagnosis in the different studies – i.e. 25 mmHg vs. 30 mmHg and/or different patients ethnicity. The prognosis of patients with PH has been reported very poor and duration of survival from onset of PH was within 2 years. Early diagnosis and treatment to lower PH may improve the prognosis of patient.(1) Heart lung transplantation and lung transplantation for PAH resulted in long term survival(5).

The case reported here illustrates several points. Severe pulmonary hypertension in association with SLE may present at a time when other manifestations of SLE are quiescent. Our patient also illustrates the point that patients with SLE may present with pulmonary hypertension without...
any evidence of parenchymal lung disease by chest x ray.

CONCLUSION
SLE as a cause of pulmonary hypertension should be suspected and evaluated accordingly in all patients particularly young females which is most common age group for both these conditions.

REFERENCES