Case Report

Case report on Plummer Vinson syndrome

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INTRODUCTION

Plummer-Vinson syndrome (PVS) was first described by Patterson and Kelly in 1919.1 It is a rare syndrome defined by the classic triad of dysphagia, iron-deficiency anemia and oesophageal webs.2 Most of the patients are middle-aged women, in the fourth to seventh decade of life but the syndrome has also been described in children and adolescents.2 PVS mainly affects the white females whereas it also observed in black women and rarely reported in males. The exact etiopathogenesis of PVS is unclear, but there is an association of iron deficiency anemia with dysphagia.3,4 The high turnover rate of the epithelium at the upper digestive tract makes the patient vulnerable to iron deficiency because of the deficiency of iron-dependent enzymes. This reduction of oxidative enzymes of epithelial cells, free radicals stress and DNA damage may lead to mucosal atrophy, web formations and malignant changes.5 PVS consists of a triad of dysphagia, iron deficiency anemia, and upper oesophageal webs. Other symptoms include atrophic oral mucosa, cracks or fissuring at the corners of the mouth, glossitis, koilonychia (spoon-shaped nails) or nails that are brittle and break easily.6

CASE REPORT

A 38 year female pale, week, sick looking patient presented with complaints of breathlessness and retrosternal chest pain since one hour history of present illness revealed easy fatigability, palpitations, dysphagia since one year insidious onset gradually progressing to solid foods with odynophagia (painful swallowing) and no complaints of dysphagia to liquids on counseling she
revealed that dysphagia and chest tightness relieved after vomiting. Her past history had no thyroid disease, diabetes, hypertension, tuberculosis and cerebrovascular attack, personal history shows that she was on a mixed diet, not a smoker or alcoholic.

On examination she was found to be gross anaemic with pallor, koilonychia (spoon shaped nails) of fingers (Figure 1) and toes, angular chelitis (cracks or fissuring at the corners of the mouth) and laboratory investigations shows haemoglobin (Hb) 7.8 g/dl, total count 7700 cumm, Blood urea nitrogen 37 mg/dl, serum creatinine 0.9 mg/dl, total bilirubin 0.9 mg/dl, direct bilirubin 0.3 mg/dl, serum albumin 3.2 g/dl, serum protein 6.9 mg/dl, platelets count 4.22 lakh/cumm and serum iron profile includes serum ferritin 7.3 ng/ml and serum iron capacity 298.4 µg/dl, transferrin 203 µg/dl, folate serum 2.93 ng/ml, vitamin B12 400 µg/l, vitamin C 500 µg/l, calcium 9.8 mg/dl, serum albumin 3.2 g/dl, serum protein 6.9 mg/dl, total bilirubin 0.9 mg/dl, direct bilirubin 0.3 mg/dl, serum albumin 3.2 g/dl, serum protein 6.9 mg/dl, platelets count 4.22 lakh/cumm and serum iron profile includes serum ferritin 7.3 ng/ml, total iron binding capacity 298.4 µg/dl, transferrin 203 µg/dl, folate serum 2.93 ng/ml and upper gastrointestinal endoscopy shows mild oesophageal webs in post cricoid region.

After considering the physical examinations, analyzing the laboratory reports and endoscopy results the clinical condition was diagnosed as Plummer Vinson syndrome.

On first and second day treatment included one packed cell transfusion iron supplements tab. Iron-folic acid 200 mg, tab. vitamin B complex od and tablet vitamin C 500 mg supplements, inj. pantoprazole 40 mg and inj. ondansetron 4 mg i.v. taken from day three iron supplements tablets was changed to syrup and folic acid was given as tab. folic acid 5 mg and on the fourth-day, plan for packed cell transfusion and rest of the therapy was same on the fifth-day and fresh complaints were observed, patient was feeling better and she was advised to continue the therapy for about a week and was asked to return for a check-up.

**Figure 1: Koilonychia (spoon-shaped nails).**

**DISCUSSION**

In 1912, Plummer reported that there were some cases of dysphagia associated with severe anemia which had been regarded as a kind of hysteria or neurosis of unknown cause. Subsequently, Vinson reported that this type of dysphagia has three characteristic manifestations: anemia, dysphagia and atrophic glossitis, and Kelly and Paterson pointed out the high incidence of hypochromic anemia in this disease. Since then syndrome associated with such symptoms has been termed as Plummer-Vinson syndrome or Paterson-Kelly syndrome. The syndrome mainly affects white females, in the 40’s to 70s of life, but some cases of pediatric and adolescent have also been reported.

The predominant clinical features of PVS are an upper oesophageal webs, dysphagia, and iron deficiency anaemia. Other oral symptoms like angular chelitis, early loss of teeth, glossitis and anaemic symptoms like weakness, pallor, fatigue, and tachycardia may dominate the clinical picture.

The pathophysiology of PVS is not clear, but various theories like iron deficiency anemia, malnutrition, genetic predisposition and autoimmune etiologies are postulated. Among them, iron deficiency anaemia is most widely accepted etiological factor as dysphagia and oesophageal webs improve with iron supplementation. The theory of iron-deficiency anemia based on the rapid losses of iron-dependent enzymes due to its high cell turnover. As in anemia, there is a reduction of iron-dependent enzymes it causes epithelial atrophy and decreases the repair capacity of the mucosa, it allows the carcinogens and co-carcinogens to act aggressively and predisposes the entire oral cavity and oesophageal area to malignancy. Another theory which has minor importance was vitamin B₆ (pyridoxine) and vitamin B₁₂ (cyanocobalamin) deficiency was also believed to contribute to PVS syndrome. The above case-patient was started with iron supplementation therapy but very few cases have been reported that were responding to iron replenishment. Dysphagia in such settings requires mechanical dilatation in adjunction with iron therapy. In most cases, one session of such dilatation is usually enough for long-term relief but, rarely, multiple sessions may also be warranted.

PVS can be associated with pernicious anemia, thyroiditis or celiac diseases. Celiac disease is one of the recognized causes of chronic iron deficiency and therefore should be considered as an etiological factor in sideropenic dysphagia. PVS has also been identified as a risk factor for developing squamous cell carcinoma of the upper gastrointestinal tract. Three to fifteen percent of the patients with PVS are mostly women between 15-50 years of age, have been reported to develop esophageal or pharyngeal cancer.

Diagnosis of PVS is based on the history of the patient, general clinical examination hematological investigations especially like anemic profile and radiological examinations like barium swallow x-ray test of the chest, upper gastrointestinal tract endoscopy. Supplementary investigations like videofluoroscopy, biopsy for histopathological examination might be helpful in some patients. Differential diagnosis of PVS has to be
performed keeping in view of all the other related causes of dysphagia such as malignant tumors esophagus, benign strictures, spastic motility disorders, scleroderma, diverticula, and gastroesophageal reflux disorders all such conditions have to be evaluated.\textsuperscript{17,18}

The initial step in treating PVS is to detect the cause of iron deficiency. In females, it is mostly due to increased menstrual flow and in males, an underlying malignancy should be ruled out. Treatment consists of correcting underlying iron deficiency anemia, iron supplements, and mechanical dilatation of the webs. Many studies have shown that iron supplementation alone can improve the symptoms and no mechanical dilatation was required.\textsuperscript{19}

This was true in our case as symptoms improved on iron therapy. Mechanical dilatation of webs can be carried out by endoscopic bougies or pneumatic balloons in single or multiple sessions. Surgery is reserved only for recurring webs. Other rare means of disruption of web include neodymium-doped: yttrium aluminum garnet laser therapy or needle-knife electro incision.\textsuperscript{2,6,18} The prognosis of PVS is good as anemia and dysphagia can be effectively treated by iron therapy and webs by dilatation. The condition gets worsen dramatically if this syndrome is associated with complications like squamous cell carcinoma of the hypopharynx and upper esophagus. Other minor supplementary treatments include the patient should be kept on a nutritious diet to maintain the integrity and maturative potential of the oral epithelium due to its malignant potential.\textsuperscript{18}

In this case, the approach has begun with iron supplementation as we observed that patient was feeling better and she was advised to continue the therapy for about a week and was asked to return for the check-up. In the next visit after a month of iron supplementation, the symptoms of dysphagia to solids decreased and the same therapy was continued.

**CONCLUSION**

Plummer-Vinson syndrome is due to chronic anemic condition leading to symptoms like dysphagia and oesophageal webs. PVS always remain a diagnostic challenge because of other chances of dysphagia like malignant tumors esophagus, benign strictures, spastic motility disorders, scleroderma, diverticula, and gastroesophageal reflux disorders. Patients with PVS are to be carefully treated in most of the cases iron supplementation alone can improve the symptoms and no mechanical dilatation was required. If dysphagia persists mechanical dilatation of the esophagus is performed. In this patient treatment started with iron supplementation and vitamin supplements, the patient was feeling better after she was continued on the same therapy and regular visit has been advised to observe the prognosis of the disease.

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