Hereditary hemorrhagic telangiectasia: A rare cause of severe anemia

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Abstract

Hereditary hemorrhagic telangiectasia (HHT) is an uncommon autosomal dominant inherited multisystem disorder in which abnormal arteriovenous communications (telangiectasias) develops which characteristically affects small blood vessels in the skin, mucous membranes, gastrointestinal tract & organs such as liver, lungs and brain causing vascular dysplasia and tendency for bleeding. This is a rare condition which often goes unrecognized & misdiagnosed because of its non specific symptoms which vary among affected population. HHT, though usually presents as recurrent epistaxis, gastrointestinal bleeding & arteriovenous malformations, iron deficiency anemia is found in few patients of HHT and is usually mild. Concomitant severe anemia in HHT is rarely seen and is found in elderly. This article presents a case of a 30 year old man presented with recurrent epistaxis since childhood. Investigating results showed multiple telangiectasias in gastrointestinal tract, hemangiomas in liver & arteriovenous malformations (AVMs) in lungs. Based on Curacao criteria, it was diagnosed as severe iron deficiency anemia with hereditary hemorrhagic telangiectasia.

Keywords: Hereditary hemorrhagic telangiectasia, iron deficiency anemia, arteriovenous malformation, epistaxis

Introduction

Hereditary hemorrhagic telangiectasia also known as Osler Weber Rendu syndrome is a rare (1 in 5000-8000) autosomal dominant genetic disorder, despite the fact that about 20 % cases are unaware of a positive family history. It leads to abnormal blood vessels formation in skin, mucous membranes & visceral organs due to defective angiogenesis. Defective angiogenesis causes alteration in elastic & muscle layer of vessel walls, making them more prone to spontaneous rupture & injuries. HHT is attributed to aberrant signaling TGFβ1 receptor involved in angiogenesis. HHT 1 is caused by mutation gene ENG encoding endothelin on chromosome 9q and HHT 2 is caused by mutation in gene ALK-1 (Activin Receptor like Kinase-1) on chromosome 12-13. HHT1 & 2 accounts for approximately 85% of cases.

Clinical manifestations

Clinical manifestations of HHT are variable, recurrent epistaxis is most common presenting symptom seen in about 90 % of patients. Mucocutaneous telangiectasia are seen in 80 % of patients with involvement of skin lip, nail
bed or oral mucosa. Half of patients manifest cutaneous lesions by age of 30 years. Pulmonary AVMs seen in 40-50% of cases, large multiple AVMs may result in dyspnea, fatigue, cyanosis, clubbing & polycythemia. Hepatic AVMs occur in 30-40% patients of HHT, hepatic AVMs can lead to right upper quadrant pain, jaundice, hepatomegaly, high output heart failure, portal hypertension, encephalopathy & liver failure. Cerebral AVMs seen in 10-20% HHT patients may lead to stroke, brain abscesses or intracellular hematoma with focal neurological signs. Spinal AVMs are seen in 1% cases. GI telangiectatic lesions most commonly involved mucous surface of stomach & small bowel. GI bleeding develops in 15-20% cases of HHT.

**Diagnosis of HHT**

The diagnostic Curacao criteria were developed based on four criteria of spontaneous recurrent nosebleeds, mucocutaneous telangiectasia (multiple & at lips, oral cavity, fingers, nose) visceral involvement (Gastrointestinal telangiectasia, pulmonary, hepatic, cerebral or spinal AVM) and an affected first degree relative, to allow a high degree of suspicion without overdiagnosis. The diagnosis of HHT is definite when three criteria are present, suspected is two criteria are present, most commonly family history and nose bleeds, or unlikely when only one criterion is present.

**Management of HHT**

Treatment of HHT is largely conservative. Anemia may require multiple blood transfusions & iron supplementation. Topical as well as systemic estrogen therapy reduces mucosal bleeding. Nose bleeds often require Nd:Yag or Argon laser treatment or local injection of sclerosing agents. Antiangiogenesis agents like bevacizumab & thalidomide have also proven to be effective in reducing mucosal bleeding. Treatment options of GI bleeding includes endoscopic Argon Plasma Coagulation (APC) & Nd-YAG laser ablation of mucosal telangiectactic lesions. Management options for skin lesions include long pulsed Nd-YAG laser ablation & skin grafting. Pulmonary hepatic & cerebral AVMs needs embolisation of the feeding vessels by clipping or coiling. Liver transplantation is indicated in cardiac failure, biliary necrosis & severe portal hypertension. Asprin & anticoagulant agents are contraindicated in patients with active bleeding.

Anemia in HHT can be due to recurrent epistaxis or chronic gastrointestinal bleeding and is usually mild. However concomitant severe anemia is a rare presentation of Hereditary Hemorrhagic telangiectasia, which needs high index of suspicion for diagnosis. This rare condition often goes unrecognized & misdiagnosed because of its non specific & variable presentation among affected population. Here we present a case report where a thirty years old male with Hereditary Hemorrhagic telangiectasia presented with severe anemia.

**Case Report**

A 30 year, chronic smoker, male, presented with easy fatigability for one year, generalized weakness for 6-8 months & dizziness for last 1 month and skin lesions which first appeared on nail bed of both hands, gradually progressing to finger, tongue, oral cavity, face, trunk & lower abdomen. Significant past history was recurrent episodes of epistaxis since age of 9 years, spontaneous in nature, lasting for 3-5 minutes only, it had no seasonal or diurnal variation. There was no history of hemarthrosis and similar problem in any other member of his family. On examination patient had sign & symptom of anemia, his nasal mucosa was inflamed. Skin lesions were reddish purple in colour, slightly elevated, sharply demarcated maculopapular & punctate, size ranging from 0.25 to 3.5 mm, blanchable & non tender present on nail bed of both
hands, finger, tongue, oral cavity, face, trunk & lower abdomen (Fig 1&2). Patients haemogram suggested severe iron deficiency anemia (Table-1). Stool was negative for occult blood. Chest X Ray showed a Nodular prominence over right hilar region(Fig.3). Differentials for this patient were hereditary hemorrhagic telangiectasia, hemophilia & Von willebrand disease. Hemophilia& von willebrand disease were excluded due to no muscle joint hematoma or swelling, no sign of easy bruising & no family history of bleeding disorder. Patient was further investigated to find other possible causes of anemia Upper GI endoscopy revealed multiple mucosal telangiectasias from oral cavity up to second part of duodenum (Fig-4), which were not bleeding at the time of study. Lower GI mucosa was found normal on colonoscopy. Triple phase CT abdomen revealed two hepatic hemangiomas (Fig-5), CT angiography of chest showed small AVM in lower lobe of right lung (Fig-6). No similar lesions detected in brain. So in the view of recurrent epistaxis, mucocutaneous telangiectasia & visceral lesions a diagnosis of HHT was made based on Curacao criteria. Patient was treated with 5 units of PRC transfusion, injection iron sucrose, oral haematinics & Hormone Replacement Therapy (Tab Tamoxifene 20 mg OD).With this patient’s general condition improved significantly. Patient has been planned for ablation of GI lesions by endoscopic Argon plasma coagulation & intervention embolization for pulmonary AVM.

Discussion:
Association of severe anemia , like the one present in our patient of HHT is highly unusual. After extensive internet search only 26 cases reports of severe anemia due to HHT were found till date 2,4,11,12,13. Previous studies have shown that most common cause of severe anemia in patients of HHT were chronic gastrointestinal bleeding in 70% followed by epistaxis in 25%, both of which are evident in this reported case12. Iron deficiency anemia found in HHT is usually mild and if severe is found in elderly14. Our reported case is also unique in the sense that this patient develops severe anemia at the age of 30 years probably due to recurrent epistaxis & GI bleeding. Patients with HHT presents with normal hemostasis and normal platelet function; hence recurrent bleeding is related to telangiectasia. Number of telangiectasia decides the severity of anemia.

Conclusion: HHT is rare disease. Further it rarely causes severe anemia & therefore is not included in differential diagnosis of anemia. Severe anemia and negative family history make the disease more rarer as in our case. HHT should be suspected in patients with early onset recurrent nasal or GI bleeding with normal coagulation profile. Though HHT is a rare disease, we need to have very high index of suspicion to establish its relation to iron deficiency anemia, this can be recognized in most patients due to its classic presentation and diligent general physical examination.

References


**Figures**

Figure 1: Telangiectatic lesions over (a) back (b & c) fingers

Figure 2: Telangiectatic lesions over oral mucosa
Figure 3: Chest X-ray PA view

Figure 4: Telangiectasias on esophageal mucosa

Figure 5: Triple phase CT abdomen showing hepatic hemangioma.

Figure 6: CT chest angiography showing right pulmonary arteriovenous malformations

Table 1: Investigations

<table>
<thead>
<tr>
<th>Investigations</th>
<th>Value</th>
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<tbody>
<tr>
<td>Hb</td>
<td>4.7 g/dl (14-18 g/dl)</td>
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<tr>
<td>TLC</td>
<td>8.8<em>1000/mm$^3$ (4.3-10.0</em>1000/mm$^3$)</td>
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<tr>
<td>Platelets</td>
<td>2.96 (1.4-4.4 lac/mm$^3$)</td>
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<tr>
<td>MCV</td>
<td>67.2 (70-99 fl)</td>
</tr>
<tr>
<td>MCH</td>
<td>18.1 (26-32 pg)</td>
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<tr>
<td>MCHC</td>
<td>27 (32-36 g/dl)</td>
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<tr>
<td>Ferritin</td>
<td>14.4 ng/ml (22-322 pg/ml)</td>
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<tr>
<td>S. Iron</td>
<td>12 ug/dl (40-150 ug/dl)</td>
</tr>
<tr>
<td>TIBC</td>
<td>640 (255-450 ug/dl)</td>
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<tr>
<td>Vitamin B$_{12}$</td>
<td>457 (211-911 pg/ml)</td>
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<tr>
<td>Folate</td>
<td>8.5 (3-17 ng/ml)</td>
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<tr>
<td>BT</td>
<td>2.27 (2-5 min)</td>
</tr>
<tr>
<td>CT</td>
<td>4.29 (3-5 min)</td>
</tr>
<tr>
<td>Stool for occult blood</td>
<td>negative</td>
</tr>
</tbody>
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