Polycythaemia Vera with Abdominal Pain - An Unusual Association

BINOY KRISHNA TARAFDER,1 MOHAMMAD ASHIK IMRAN KHAN,2 MRINMAY KUMAR PODDAR,2 ESTAKUR RAHMAN2, MD. AMIRUZZAMAN3, MD. TITU MIAH4

Abstract:
Polycythaemia vera is difficult disease to diagnose when it has an atypical presentation. Here we present a case where a middle aged man who presented to us with abdominal pain and recurrent peptic ulceration and was later diagnosed as a case of polycythaemia vera. To best of our knowledge this is the first report of such association from Bangladesh.

Keyword: Polycythaemia, vera, Abdominal pain, Peptic ulcer, Bangladesh

Introduction:
Abdominal pain is a common complaint presenting to primary care physicians, and accounts for 4–8% of all adult visits to a hospital emergency department.1,2 While approaching a patient with abdominal pain, one needs to be cautious as many causes can be missed if not kept in mind. The above statement is more true when one is working in a setting where reliance is mostly on clinical findings. Polycythaemia vera is a condition characterized as a panhyperplastic, malignant marrow disorder. The most prominent feature is the rise in red cell mass, often accompanied by rises in white blood cell and platelet counts. Leukaemic conversion occurs in 1–3% and the median survival with treatment is 13 years.3 We encountered a patient with upper abdominal pain, who had undergone repeated endoscopy for abdominal pain but was ultimately diagnosed with polycythaemia vera.

Case Report:
A 50 year old married gentleman presented to our OPD with upper abdominal pain for 2 years. Pain was burning in nature, episodic, related with food but no specific pattern, subsided by taking antiulcerant drugs. But over the last 3 months pain was almost on a daily basis, not adequately responding to antiulcerant drugs. Pain was not associated with fever or weight loss. He gave no history of hematemesis, melena or vomiting, chronic cough or respiratory distress. He was non alcoholic but smoker and smoked about 5 pack years. On query, gave history of generalized itching for last 3 yrs. His examination revealed plethoric appearance, conjunctiva: Congested. Pulse: 72b/m BP: 130/80 mm of Hg, palmer erythema present. Systemic Examination revealed mild tenderness over epigastric region, mild splenomegaly. other systemic examinations were unremarkable.

His complete blood count revealed Hb -21.7 gm/dl, WBC -16.42, RBC-10.63, MCV - 61.9, MCH -20.4, MCHC- 33. Platelet - 369000, ESR - 18 mm in 1st hour. Neutrophil - 70%, Lymphocyte- 19.2%, Monocyte-3.2%,Eosinophil-7.4%,Basophil-0.2%. Unfortunately patient discarded all his previous endoscopy images keeping only his papers. Review of his medical records showed duodenal ulcers in all his endoscopies but H. Pylori was absent in two of the three occasions. Even then he received eradication therapy.

His CXR was normal and echocardiography revealed no underlying heart disease. Persistently maintained oxygen saturation (SpO2) above 90% range ruled out secondary polycythemia indirectly as well. Due to inadequate facility in the ward, his oxygen saturation was measured in the OR where pulse oxymetry machine is available.

1. Junior Consultant, Medicine, 250 Bed Hospital, Sadar, Gopalganj.
2. Medical Officer, Medicine, 250 Bed Hospital, Sadar, Gopalganj.
3. Resident Physician, Sir Salimullah Medical College & Mitford Hospital, Dhaka.
4. Associate Professor, Dhaka Medical College Hospital, Dhaka

Correspondence: Dr. Binoy Krishna Tarafder, Junior Consultant, Medicine, 250 Bed Hospital, Sadar, Gopalganj.
Discussion:
While approaching a patient with abdominal pain in a medicine ward, one needs to be very cautious, especially while working in a setting where adequate investigation facilities are not present. There is no substitution for careful history taking and meticulous and thorough clinical examination. Although medial causes do not account for more than 5% of all abdominal pains, correct diagnosis is only reached by only those who are looking for a clue in all body systems, especially outside the abdomen.4

Diagnosis of polycythemia vera is largely based on parameters set by polycythemia study group (PVSG). It can be established if the following criteria are met:5

But there are many controversies regarding these criteria. But the cornerstone of diagnosis is excluding the diagnosis of familial and secondary forms of polycythemia and thrombocytosis. Serum Epo assays are useful in the initial evaluation of erythrocytosis, particularly when related to simultaneous hemoglobin/ hematocrit determinations. Although PV remains a diagnostic possibility in the presence of low or normal serum Epo levels, erythrocytosis associated with increased Epo levels is usually due to secondary causes. Low Epo levels can also be seen in ET. Initial experience indicates that mutation screening for JAK2-V617F can accurately distinguish between PV and secondary polycythemias.6-9

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<thead>
<tr>
<th>CATEGORY A Lab Tests</th>
<th>CATEGORY B Lab Tests</th>
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<tr>
<td>A1: Total RBC Volume (REDMAS) Male &gt;= 36 ml/kg BW Female &gt;= 32 ml/kg BW</td>
<td>B1: Thrombocytosis (WBC) &gt;400,000/mm³</td>
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<tr>
<td>A2: Arterial saturation (OSAT) O2 &gt; 92%</td>
<td>B2: Leukocytosis &gt;12,000/mm³ (PLAT) (no fever or infection)</td>
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<tr>
<td>A3: Splenomegaly (SPLEEN)</td>
<td>B3: Leukocyte alkaline phosphatase (LAP) Score &gt; 100</td>
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<td></td>
<td>B4a: Serum vitamin B12 &gt; 900 pg/ml or (B12)</td>
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<tr>
<td></td>
<td>B4b: Serum UB12BC &gt; 2,200 pg/ml (UBBC)</td>
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DIAGNOSIS OF POLYCYTHEMIA VERA:
A1 + A2 + A3 or A1 + A2 + any two from category B
In our patient, many of the criteria of polycythemia were fulfilled. Due to lack of diagnostic facility, we could not go for the complex laboratory testing. The interesting fact was patient fled the hospital when he was told about the treatment modalities, especially venesection probably out of fear.

But his mode of presentation makes this case unique and very interesting. Many medical conditions have been discussed as causative factor for abdominal pain in adults. They have been described under broad headings such as referred abdominal pain from heart such as myocarditis, endocarditis, congestive heart failure, myocardial ischaemia and infarction. Respiratory causes such as pneumonia, pneumonitis, pulmonary, embolism and infarction can also cause abdominal pain.

Metabolic and endocrine causes such as ketoacidosis (diabetic and alcoholic), acute adrenal insufficiency, uraemia, hyperthyroidism, porphyria hyperparathyroidism/hypercalcaemia, narcotic withdrawal. Even heat stroke, psychiatric disorders abdominal wall spasm/haematoma can cause abdominal pain. Many haematological cause such as sickle cell anemia, haemolytic anaemia, Henoch-Schönlein purpura, acute leukaemia and polycythemia rubra vera can also cause abdominal pain. 10,11 Various neurological causes and even toxins can cause abdominal pain.

Our patients recurrent abdominal pain led him to various physicians where medical cause was overlooked. The underlying mechanism for abdominal pain in polycythemia has been attributed mostly to hyperviscosity syndrome but it has also been argued due to coagulopathies as well. 4 It has been proved that qualitative platelet defect is present in polycythemia contributed to various factors such as thrombocytosis, loss of VWF. 12 Circulating, bone marrow hematopoietic stem cell–derived endothelial progenitors can home to areas of vascular injury or ischemia and repopulate the intimal surfaces of the vessel wall as differentiated endothelial cells. TPO and JAK2 mutations may also be responsible for increased platelet aggregation. The above mentioned factors might have been the underlying cause of the recurrent ulcers in our patient. 13-15 Though the polycythemia itself is not usually seen as an underlying cause of peptic ulceration or a common cause of abdominal pain, it must be kept in mind. This association is a rare finding and has not been reported earlier to the best of our knowledge.

It has been seen that the diagnostic accuracy of physicians assessing abdominal pain using clinical signs and symptoms and preliminary investigations is not very good. When preliminary and final diagnoses are compared, clinical accuracy has been reported to be 50–65% overall, and as low as 30% in the elderly. 16,17 So, as physicians we must keep an open mind, and remain suspicious even when there is no reason to be so.

Conflict of Interest: None

References:


