Is mitral stenosis associated with gastrointestinal bleeding: A twist on Heyde’s Syndrome

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Abstract

Introduction: Heyde’s syndrome is the link between Aortic stenosis (AS) and angiodysplasia leading to Gastrointestinal (GI) bleed. There have been multiple cases and theories leading to this phenomenon, including the loss of von Willebrand factor (vWF) through shear factor of a stenotic valve. This association is further validated by the cessation of GI bleed after aortic valve replacement. The question posed is, if AS can lead to GI bleeding through loss of vWF or other mechanisms, would mitral stenosis (MS) cause a similar presentation?

Methods: A single center, retrospective chart analysis was performed on patients, age 18 and over, with evidence of MS on echocardiogram for any signs of GI bleeding. The outpatient clinic notes and admission notes, along with colonoscopies to detect the presence of angiodysplasia were reviewed for GI bleeding. Patient’s with concomitant AS were excluded.

Results: MS group were 35% males and 65% female. Average age was 61 years old. Approximately 4% had MS secondary to rheumatic heart disease and 45% due to calcified annulus. Of 162 patients with MS, 7 (4.3%) patients had evidence of gastrointestinal bleed versus 16 (10%) of non-MS group (p=0.06). Patients with MS and GI bleed were found to have arteriovenous malformation (AVM) (35%), gastric or duodenal ulcer (35%), colon cancer (3%) and diverticulitis (37%).

Conclusion: MS does not have an increase incidence of GI bleeding when compared to the control group, though the P value was not statistically significant. vWF is thought to be decreased because of increased shear force through a stenotic valve. Flow through the stenotic mitral valve is orders of magnitude lower than the flow through a stenotic aortic valve given the force of contractility in the atria compared to the ventricle, therefore is unlikely to cause decrease in vWF to lead to GI bleeds.

Introduction

Heyde’s syndrome is the link between Aortic stenosis (AS) and angiodysplasia leading to Gastrointestinal (GI) bleed. It was first noted by Dr. Edward C. Heyde back in 1958 as GI bleeding of idiopathic sources was significantly more prevalent among patients with AS [1]. There have been multiple theories behind Heyde’s syndrome; however, it is now believed to be caused by a depletion of Von Willebrand factor (vWF), similar to Von Willebrand disease type IIA (vWD-2A). Bleeding is thought to occur due to angiodysplasia colonic vascular ectasias that is unable to clot due to destruction and depletion of vWF; this occurs as vWF flows through the narrowed valve and cleaved by metalloproteinase [2]. This association is further validated by the cessation of GI bleed after aortic valve replacement. The question posed is, if AS can lead to GI bleeding through loss of vWF or other mechanisms, would MS cause a similar presentation?

Methods

Study design

A single-centered, retrospective chart analysis was performed between 2004 and 2014 of patients who had evidence of MS, and matched with non-MS group. This study was conducted at a major hospital center in one of the most diverse communities in the United States, providing a culturally and epidemiologically significant advantage. An approved retrospective chart analysis using QuadraMed Computerized Patient Record (QCPR) and accessed with data-input and calculations formulated in a computerized software. This study has been approved by Institutional Review Board of the Mount Sinai School of Medicine (HS#: IRB-16-00393).

Selection criteria

One hundred and sixty-two patients had evidence of MS based on 2D echocardiogram. Patient’s with concomitant AS were excluded. Among these patients, the chart was reviewed for admissions suspected of upper or lower GI bleed with signs of anemia, and had either Esophagogastroduodenoscopy (EGD), Colonoscopy or both. The underlying cause of MS was identified, and a random matched control group without MS was selected (Table 1). For all statistical analysis, the results were considered significant when two-tailed p-values were <0.05. The distributions of age, sex and baseline co-morbidities were compared between the two cohorts.

Results

MS group were 35% males and 65% female with average age of 61 years old. Approximately 4% had MS secondary to rheumatic heart disease and 45% due to calcified annulus. Of 162 patients with MS, 7 (4.3%) patients had evidence of gastrointestinal bleed versus 16 (10%)
of non-MS group (p=0.06). The causes of GI bleed in patients with MS were as follows: arteriovenous malformation (AVM) (35%), gastric or duodenal ulcer (35%), colon cancer (3%) and diverticulitis (37%).

**Discussion**

**MS is caused by five main conditions**

Rheumatic disease, congenital abnormalities, active infective endocarditis, metabolic or enzymatic abnormalities such as gout or whipples massive annular calcification [3] The severity of MS can be broken down into: mild with gradient 8 mm Hg, moderate with gradient 8 to 12 mm Hg and severe with gradient 12mm Hg or higher [4]. According to the data we gathered, there is no association between MS and angiodysplasia or GI bleeding. The accepted pathogenesis of Heyde’s syndrome is acquired vWF deficiency due to shearing. Shear rate is directly proportional to the blood flow velocity and inversely proportional to vessel diameter. Under normal physiologic situations, the highest wall shear rates occur in the arterioles 4,000 seconds⁻¹, which enhance platelet/vWF interactions [5,6].

The vWF abnormality is related to the severity of stenosis [7]. Severe AS can produce shear rates as high as 10,000 seconds⁻¹, which induces a conformational change to the high molecular weight (HMW) vWF multimers, exposing the bond between amino acids Tyr842 and Met843, leading to proteolysis by ADAMTS13, an important disintegrin and metalloprotease that cleaves vWF at the A2 domain [5-9]. This effectively decreases the number of HMW multimers leading to an acquired VWD-2A and impairing hemostasis at angiodysplastic vessels, causing to a tendency to bleed, as HMW multimers are the most platelet reactive, and aid clotting [6,9,10]. In MS, the velocity increases from a normal value of less than 1 m/sec to a value greater than 1.5 m/sec [11] and even with significant decrease in mitral valve area in patients, peak and mean trans mitral gradient do not significantly change [12]. In addition, the atrial contraction or atrial kick accounts for roughly 10% of ventricular filling, and up to 24% across a stenotic mitral valve, depending on degree of stenosis and mitral valve area, which fluctuates with flow [13,14]. When considering force of contraction of the atrium to the force of contraction of the ventricle, where normal ejection fraction is 65%, there is large discrepancy in force and velocity generated between the two chambers, showing a clear difference in shear force. Finally, atrial fibrillation occurs in 40-75% of patients with MS, [15] which effectively removes the force produced by the atrium, further effectively decreasing shear rate. Therefore, shear in MS is negligible when compared to AS, and would not cause the conformational change and destruction that would lead to an acquired vWF deficiency. In a study of 500 operatively excised mitral valves, 45% of the valves had concomitant stenotic and regurgitation features [3], which would prompt a patient to become further symptomatic and receive mitral valve replacement. In patients with GI bleed and AS, shear stress was significantly reduced after valve replacement [16] and further studies showed that aortic valve replacement stopped bleeding in as high as 93% of patients [10,17,18] and thought to be the solution for this syndrome. Therefore, even if MS had enough shear force to cause an acquired vWF deficiency, a fair number of patients would have had the required treatment early on and not experience GI bleed, unless from anticoagulation for valve replacement.

**Conclusion**

Finally, with MS, there is a risk of left atrial thrombus due to increased systemic levels of byproducts from the coagulation cascade. Studies have shown that there are increased prothrombin fragments in the left atrium [19,20] and this hypercoagulable state is likely due to sluggish blood flow and low shear rate conditions. Therefore, MS may actually have less incidence of bleeding which is what our data showed, though the data set is small and statistically not significant. High shear rates, which also occur pathologically when blood flows rapidly through larger but stenotic vessels, MS does not have an increase incidence of GI bleeding when compared to the control group, though the P value was not statistically significant and may actually have less incidence given a possible hypercoagulable state from having MS. vWF is thought to be decreased because of increased shear force through a stenotic valve, leading to degradation; however, flow through the stenotic mitral valve is orders of magnitude lower than the flow through a stenotic aortic valve given the force of contractility in the atria, or lack of, in atrial fibrillation, as compared to the force of contraction in the ventricle. Therefore, is unlikely to cause decrease in vWF that leads to GI bleeding from angiodyplastic vessels.

**References**


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