Paroxysmal Nocturnal Hemoglobinuria (PNH) — Symptoms and Diagnosis

Paroxysmal nocturnal hemoglobinuria (PNH) is an acquired defect in the myeloid stem cell lineage and can be seen as a rare, chronic, morbid disorder. Formally known as Marchiafava-Micheli syndrome, it received its current name as a descriptive term for the disease. Individuals afflicted with the disease wake up to dark, “cola” colored urine due to RBC breakdown and release of hemoglobin in the urine overnight. The discoloration of urine is seen in several other disorders, which is why it is important to discern the disease from other hemolytic disorders as the article depicts.

Epidemiology of PNH

Paroxysmal nocturnal hemoglobinuria as a rare disorder

Paroxysmal nocturnal hemoglobinuria typically presents in males and females in early adulthood and manifests throughout. It has an incidence of **10 cases per million with a 50 % mortality rate.**
Etiology of PNH

PNH is an **acquired defect in the myeloid stem cells** with a mutation in the PIG-A gene located in the stem cells of the bone marrow. With a **PIG mutation, affected red blood cells** are known as “**PNH RBCs.**“ The PNH RBCs lack the shield of proteins that protect normal red blood cells from the complement system. This is all due to deficiency in **glycosylphosphatidylinositol (GPI).** GPI are linked proteins on RBCs, neutrophils and platelets. If absent it renders cells susceptible to destruction.

Pathophysiology and Pathogenesis of PNH

The predisposing factor in anyone with PNH is the **inability to synthesize GPI.** GPI is located on the X chromosome, rendering only one mutation necessary to eliminate the expression of GPI-linked proteins. One essential GPI-membrane-linked protein is decay accelerating factor (DAF). This protein interacts with complement proteins to neutralize the complement attached to RBCs, neutrophils and platelets. In the absence of this protein, RBCs are susceptible to complement-mediated intravascular hemolysis. The destruction of RBC membranes by the complement releases hemoglobin into the bloodstream and circulation.

The body has a certain threshold in which it can degrade plasma hemoglobin. However, once levels are reached, any extra hemoglobin leads to increased heme in the urine and plasma. This sequelae leads to further pathologies as increased hemoglobin levels deplete nitric oxide levels in circulation. Nitric oxide is needed for vasodilation, smooth muscle relaxation and vascular homeostasis. **Decreased levels of nitric oxide** lead to vasoconstriction, smooth muscle contractions and spasms.

When it comes to mortality and morbidity, PNH, unfortunately, has a high incidence of almost 50 %. This is due to the pathophysiology of **thrombosis,** secondary to PNH. **Hemolysis** increases the number of thrombotic events, which, when logged in multiple organs, increases organ failure and insufficiency. Thrombotic events are also due to hypercoagulable states induced by free hemoglobin in the bloodstream.
Paroxysmal nocturnal hemoglobinuria is named due to episodic (paroxysmal) hemolysis occurring usually at night due to acidotic activation. It is believed that mild respiratory acidosis develops with shallow breathing during sleep. This activates the complement mechanism leading to RBC, WBC and platelet lysis. This lysis leads to hemoglobinemia and hemoglobinuria in the morning.

Symptoms of PNH

Signs of paroxysmal nocturnal hemoglobinuria

Associated with a plethora of clinical findings, PNH is usually symptomatic, with a small percentage being asymptomatic. The first classic sign of PNH is hemolysis due to red blood cell lysis. This leads to symptoms like fatigue, jaundice and hematuria.

The side effects of hemolysis can be broken down by different manifestations in the body’s organs and pathways. Renal insufficiency arises due to intravascular hemolysis in PNH. Toxicity increases due to direct free heme in the kidney. Chronic hemolysis increases renal iron deposition leading to cortical scarring and infarcts.

Additionally, lysis leads to depletion of nitric oxide levels. Symptoms of decreased nitric oxide leads to smooth muscle dystonia, abdominal pain, cramping and finally, erectile dysfunction. Abdominal pain is due to smooth muscle dystonia, leading to contraction of the visceral organs and vasculature, which causes spasms and pain. Lack of nitric oxide also leads to decreased vascular dilation in genital tissue, specifically the corpora cavernosa, leading to erectile dysfunction.

The most detrimental complication with PNH is thrombosis. It is the primary cause of death in afflicted patients, occurring in over 40% of the individuals. Thrombi may lodge in the venous or arterial system, such as hepatic, portal or cerebral veins, causing cirrhosis, splenic congestion and cerebral strokes, respectively. Symptoms may be insidious as clots aren’t found immediately. Skin necrosis, vein thrombosis and pulmonary embolisms are some of the most frequent symptoms. Some clots are found incidentally, and unfortunately, many go undetected leading to increased mortality.

Iron deficiency from blood loss leads to anemic like signs and symptoms, i.e., fatigue, pallor (Image 2), dyspnea, headaches, tachycardia. Patients with aplastic anemia (a disorder where the body halts production of new red blood cells due to bone marrow damage), studies show, are more prone to develop PNH.
Diagnosis of PNH

Paroxysmal nocturnal hemoglobinuria detected by flow cytometry

Several laboratory measures are used to detect PNH. Typically, an evaluation consists of testing the patient for hemolytic anemia and to rule out other causes of hemolysis. This includes autoimmune pathologies or injuries. Most commonly performed tests are:

- CBC
- RBC smear
- Reticulocyte count
- Direct Coombs testing
- Urine hemoglobin or hemosiderin.

Flow cytometry (Image 3) is the most specific test used to confirm PNH. Patients at risk for PNH, aplastic anemia, or myelodysplastic disorder patients, are screened yearly to monitor development of subclinical PNH. The patient’s blood is incubated with tagged fluorescent antibodies that bind to GPI-linked proteins. The most commonly used antibodies are Fluorescent AERolysin (FLAER). This state-of-the-art laboratory test sends the patient’s blood for flow cytometry to detect CD59 (MIRL), a glycoprotein, and CD55 (DAF), in regulation of complement action. Absence or reduced expression of both CD59 and CD55 on PNH RBCs is diagnostic.

A second diagnostic test would be the sugar water or sucrose lysis test. This test uses the ionic strength of serum that is reduced by adding an iso-osmotic solution of sucrose, which then activates the classic complement pathway, and complement-sensitive cells are lysed.
Lab values typically show anemia, increased reticulocyte count, increased LDH, free serum hemoglobin with red serum, a negative Coomb’s test and iron deficiency. Paroxysmal nocturnal hemoglobinuria (PNH) leukocytes have a low leukocyte alkaline phosphatase (ALP) score.

**Therapy of PNH**

Therapy aimed at treating PNH is focused on targeting the underlying hemolytic defect and monitoring for progression. Patients with PNH have yearly screens for increased or decreased PNH clone sizes. As of now, the only therapies for classic PNH include hematopoietic cell transplantation (HCT) or complement inhibition with medications such as eculizumab. Those with multiple thrombi are treated with anticoagulation therapeutics. Iron supplementation is used for iron deficient patients, in conjunction with folate. Those with bone marrow failures, or malignancies, are treated with immunosuppressive therapy and weekly WBC monitoring.

**Opioids and analgesics** are used for those suffering from smooth muscle dystonia and spasms. Women suffering from PNH who are interested in pregnancy pose a risk of increased maternal and fetal morbidity and mortality. Therefore, strict iron and folate supplementation, along with transfusions, may be needed during pregnancy.

**Summary**

Overall, paroxysmal nocturnal hemoglobinuria (PNH) is a rare, but life-threatening, disorder. Unexplained hemolytic anemia, thrombi and hematuria all lead to a painful increase in mortality. All patients should have a baseline testing which should continue annually thereafter. Management may be aggressive or mild depending on symptoms and severity. Overall, there is no cure for PNH at the moment, but management and research is being done to help treat this hemolytic disease.

**Review Questions**

The answers are below the references.

1. A 26-year-old male comes into the clinic for lab work due to suspected PNH. Past medical history shows recent onset of labored breathing, stomach pains and neuropathies. After diagnostic, his lab results would read all of the following, except?

   - A. Hemolytic anemia
   - B. Leukopenia
   - C. Hemoglobinuria
   - D. Neutropenia

2. A 56-year-old female comes into the hospital for hemolytic anemia, red urine and difficulty breathing. She was diagnosed with paroxysmal nocturnal hemoglobinuria. Red blood cells in PNH patients are sensitive to which of the following?

   - A. Complement proteins
   - B. Warm antibodies
   - C. Cold antibodies
   - D. Nonoxidative stress
3. A 37-year-old male visits your office for dyspnea and fatigue for 1 month. Past medical history is significant for hospitalization for hepatic vein thrombosis. On physical exam, patient was seen to have slight skin necrosis. Lab studies showed the following:

- WBC: 2900/µl
- HGB: 5.9 gm%
- MCV: 80 fl
- Platelets: 82 k/µl
- Reticulocyte count: 4.3%
- Haptoglobin: Undetectable
- Lactic Dehydrogenase (LDH): 900 U/L
- Direct Coomb’s Test: Negative

Which is the next best step in diagnosing the suspected condition?

A. urine analysis  
B. flow cytometry  
C. blood smear  
D. PT/PTT/INR analysis

References


Correct answers: 1D, 2A, 3B

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