Paroxysmal nocturnal hemoglobinuria (PNH) is an acquired stem cell disorder characterized by three cardinal clinical manifestations: 1) hemolysis and hemoglobinuria, 2) thrombosis, and 3) bone marrow failure indistinguishable from aplastic anemia. Paroxysmal nocturnal hemoglobinuria is a rare disease with an incidence of 2-6 per million persons. Only 220 reported cases of PNH were found in one of the largest retrospective epidemiological studies in France from 1946 to 1995.

Case Report

A 54-year-old woman presented to the Emergency Department complaining of sudden onset of right-sided tinnitus followed by dizziness, dysarthria, nausea, and vomiting. Her only additional complaint was that of “cola” colored urine and dysuria. The patient denied a history of fever, seizures, focal weakness or sensory abnormalities. There was no history of cough, night sweats, weight loss or diarrhea. The patient’s neurologic symptoms resolved within minutes, followed only by a self-limiting bi-frontal headache.

Her past medical history was significant for a diagnosis of paroxysmal nocturnal hemoglobinuria made at age 27 when she presented to hospital with symptoms of anemia. Investigations at that time, including a HAMs test, confirmed the diagnosis of PNH. After her initial management, she had not been followed by a haematologist.

Four years prior to the current admission she presented to hospital with a seizure and severe left hemiparesis. Imaging revealed a right frontal intracerebral haemorrhage (ICH) secondary to superior sagittal sinus thrombosis. She had full neurologic recovery from the ICH, which was treated conservatively. She also had two non-ST-elevation myocardial infarctions (NSTEMI), the first in 2000 and the second was 20 days prior to her current hospitalization. The NSTEMI in 2000 had been treated conservatively with acetylsalicylic acid (ASA). The second NSTEMI in 2004 prompted an angiogram, which revealed an occlusive thrombus in the circumflex artery (see Figure 1). She was transferred to a tertiary care center for angioplasty after initial medical management but the second angiogram showed no further clot in the circumflex artery. Clopidogrel was added to ASA for secondary cardiac prevention. The patient had no other vascular risk factors including no history of hypertension, diabetes, or hyperlipidemia and she was a non-smoker.

With her current presentation, neurologic examination was completely normal. The general physical exam was also unremarkable. Her initial lab work was significant for hemoglobin 84 g/L (mean corpuscle volume 96 fl), elevated bilirubin (total 43 uM and indirect 34 uM), reticulocyte count 191 g/L, elevated AST 58 U/L and LDH 714 U/L, and an undetectable haptoglobin (<0.07 g/L). Electrolytes and fasting serum glucose were normal; creatinine was 46. Peripheral blood film showed a normocytic anemia with no fragments consistent...
with the laboratory studies for an intravascular haemolytic anemia. Brain CT showed no acute changes; an old right frontal parasagittal hemorrhage was seen.

Brain MRI, MR-angiogram (MRA) and MR-venogram (MRV) were done. The MRI revealed an acute right cerebellar infarct in the superior cerebellar artery (SCA) territory (see Figure 2a). Also noted were chronic changes of an old right frontal ICH as well as old infarcts in the left cerebellar hemisphere and left caudate head (see Figures 2b and 2c). The MRV demonstrated no acute venous thrombosis but showed features of a prior sagittal sinus thrombosis (see Figure 3). The MRA revealed no evidence of atherosclerotic disease within the carotid or vertebral vessels (not shown).

The patient’s diagnosis of PNH was re-confirmed by flow cytometric analysis showing CD59 deficient erythrocytes of 74% and granulocytes of 79%. Laboratory tests for other causes of hypercoagulability were negative (protein C 1.22 U/ml, protein S 0.53 U/ml, anti-thrombin III 1.0 U/ml, homocysteine 8.8 uM, antiphospholipid antibody screen, prothrombin variant, and factor V Leiden). A transthoracic echocardiogram with saline bubble study was normal. Her fasting lipid profile was normal.

The patient had a transfusion of three units of packed red blood cells. For secondary stroke prevention she was continued on combined antiplatelet therapy with ASA and Clopidogrel but had a recurrence of neurological symptoms in hospital causing moderate left arm ataxia. She was then anticoagulated with unfractionated heparin, then coumadin. Low dose ASA was continued. Upon hospital discharge she had only mild residual incoordination in the left arm. Follow-up was arranged with hematology for consideration of eculizumab therapy or bone marrow transplant, the internal medicine ambulatory clinic, and the Stroke Prevention Clinic.

**DISCUSSION**

Paroxysmal nocturnal hemoglobinuria is the only haemolytic anemia caused by an intrinsic defect that is acquired and not inherited. Defective stem cells arise from clonal expansion of a totipotent stem cell containing a somatic mutation in the...
phosphatidylinositol glycan complementation group A gene (PIG-A gene) on the X chromosome. More than 100 PIG-A mutations have now been described. The PIG-A gene produces the glycosylphosphatidylinositol (GPI) anchor that serves to attach a number of proteins to the cell membrane. Two such proteins, CD55 (Decay Accelerating Factor – DAF) and CD59 (Membrane Inhibitor of Reactive Lysis – MIIRL), missing from the red blood cell surface in PNH, constitute the normal red cell’s ability to resist complement-mediated lysis. Approximately 50% of PNH patients present with nocturnal hemoglobinuria. Darkening of the urine is most noticeable in the morning, either because the urine is more concentrated or there is increased hemolysis at night. Increased venous thrombotic episodes have been noted in patients with PNH and thrombosis related death occurs in 40% of patients. These thrombotic episodes can occur in any site throughout the body, however venous thrombosis has been reported more frequently than arterial. Classically, hepatic venous thrombosis (Budd-Chiari syndrome) has been associated with PNH. Arterial thrombosis has been reported rarely with involvement of cerebral, renal, splenic, iliac and coronary arteries, based on results of autopsy findings. The increased thrombosis is due to platelet activation via complement, an increase in circulating procoagulant activity resulting from erythrocyte hemolysis and release of adenosine diphosphate (ADP).

Cerebral venous sinus thrombosis is seen in PNH and often carries a poor prognosis. Less frequently, arterial ischemic strokes occur within medium to large artery territories either in isolation or in patients with prior systemic venous thrombosis. At presentation, these patients are often younger than typical stroke patients and warrant complete work-up of rare stroke etiologies. These include hypercoagulable states (antiphospholipid antibodies, protein C or S deficiency, antithrombin III deficiency, prothrombin gene mutation, and MTHFR mutation), oral contraceptive use, paradoxical emboli related to an intra-cardiac shunt (patent foramen ovale or atrial septal defect), arterial dissection and primary central nervous system vasculitis. Atherosclerotic risk factors and cardiac arrhythmias must also be addressed if present. In some patients, a co-existent procoagulant factor is found to guide therapy, though in most cases PNH is the only risk factor. No randomized trials have evaluated the efficacy of antiplatelet agents or anticoagulation for primary or secondary prevention of stroke and sinovenous thrombosis in these patients.

A diagnosis of PNH should be entertained in all patients presenting with pancytopenia of unknown origin with reticulocytosis, non-immune hemolysis and venous thrombosis at an unusual anatomic site. Paroxysmal nocturnal hemoglobinuria can be differentiated from other haemolytic anemias by the presence of hemosiderinuria. Definitive diagnosis is established by the demonstration of GPI linked protein deficiencies on erythrocyte and neutrophil surfaces by flow cytometry.

Testing for PNH includes the sucrose-hemolysis test (HAMS test) based on complement-mediated lysis of erythrocytes in a low ionic-strength solution. Greater sensitivity and specificity may be achieved through the use of fluorescence activated cell sorter (FACS) to quantitate the percentage of cells deficient in CD59 or CD55. The percentage of cells completely deficient in CD59 (Type III PNH cells) can be correlated with the degree and severity of clinical symptoms in PNH. Also, at least two GPI linked proteins (CD55 and CD59) should be analysed to exclude the rare inherited deficiencies of single antigens (selective CD55 deficiency).

The only known cure for PNH is haematopoietic stem cell transplantation, which replaces the abnormal clone(s) present in the bone marrow. Supportive therapy includes red blood cell transfusion, which is believed to suppress marrow production of new PNH clones, thereby decreasing thrombotic complications. Some evidence supports the use of anticoagulation for patients who have experienced a thrombotic event. A non-randomized study by Hall et al. showed anticoagulation of patients with large PNH clones reduced the frequency of thrombotic events by 30%. Iron and folic acid supplementation are recommended as in any haemolytic anaemia. Recombinant tissue plasminogen activator (rtPA) may be used in the management of acute ischemic stroke following current guidelines for its use. Eculizumab, a recently developed complement inhibitor has entered clinical trials and has been shown to be safe and effective at reducing hemolysis in PNH patients.

The clinical course of PNH is quite variable. Patients with PNH may have a long-term chronic illness but the disease does shorten life. The median survival from diagnosis was 10 to 15 years in two large historical studies. Patients most commonly die of thrombosis or progressive cytopenias. Malignancy, such as myelodyplasia or acute myeloid leukemia supervenes in 2-3% of cases. Due to a relatively hypercoaguable state, oral contraception and hormone replacement therapy should be avoided. Pregnancy can be considered with prophylactic anticoagulation in a high-risk obstetrical clinic.

This case report provides a description of a rare hematologic disorder, PNH, its clinical presentation, pathophysiology, diagnostic tests and recommended treatment. The case also documents a rare neurologic presentation of PNH with both arterial and venous thrombosis.

References


