Sickle Cell Disease: Advances in Pathogenesis and Management

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Cardiopulmonary Complications of Sickle Cell Disease: Role of Nitric Oxide and Hemolytic Anemia

Mark T. Gladwin and Gregory J. Kato

Medical advances in the management of patients with sickle cell disease, thalassemia, and other hemolytic anemias have led to significant increases in life expectancy. Improved public health, neonatal screening, parental and patient education, advances in red cell transfusion medicine, iron chelation therapy, penicillin prophylaxis for children, pneumococcal immunization, and hydroxyurea therapy have all likely contributed to this effect on longevity.1,2 Importantly, as a generation of patients with sickle cell disease and thalassemia ages, new chronic complications of these hemoglobinopathies develop. In this context, pulmonary hypertension is emerging as one of the leading causes of morbidity and mortality in adult sickle cell and thalassemia patients, and likely in patients with other hemolytic anemias. A common feature of both sickle cell disease and thalassemia is intravascular hemolysis and chronic anemia. Recent data suggest that chronic intravascular hemolysis is associated with a state of endothelial dysfunction characterized by reduced nitric oxide (NO) bioavailability, pro-oxidant and pro-inflammatory stress and coagulopathy, leading to vasomotor instability and ultimately producing a proliferative vasculopathy, a hallmark of which is the development of pulmonary hypertension in adulthood.3-5 In conclusion, pulmonary hypertension is common in patients with hereditary hemolytic anemias and is associated with a high risk of death in patients with sickle cell disease. New therapies targeting this vasculopathy and aimed at normalizing the vasodilator:vasoconstrictor balance are discussed.

Endothelial Control of Vascular Function

NO is a soluble diatomic gas molecule, much like carbon monoxide. Because of its unpaired electron, NO is a free radical, providing it with unique reactivities and biological properties.6 NO is produced in endothelium by the endothelial NO synthase enzyme, by an oxygen-dependent conversion of L-arginine to citrulline.7 Once produced, NO can diffuse in a paracrine fashion to adjacent smooth muscle, where it binds avidly to the heme moiety of soluble guanylate cyclase. This activates the enzyme, which in turn converts GTP to cGMP, activating cGMP-dependent protein kinases, which ultimately sequester calcium and produce vasodilation.8,9 In addition to this vasodilation, which is tonic in nature and controls approximately 25% of our resting blood flow, NO promotes general vascular homeostasis (Table 1). Importantly, NO also reacts with the oxygenated and deoxygenated heme groups of hemoglobin at nearly diffusion limited rates (10^7 M^-1 sec^-1) to produce methemoglobin and nitrate, and iron-nitrosyl-hemoglobin, respectively (Equations 1 and 2).10

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diffusional barrier around the erythrocyte membrane. Within the erythrocyte membrane, this compartmental bin is limited by compartmentalization of hemoglobin in normal physiology, the reaction of NO with hemoglobin scavenging systems that limit extravasation in the erythrocyte, of laminar flowing blood, and the plasma paradox of biology is solved by the physical properties of bin (10 mM heme concentration in whole blood). This compartmentalization of hemoglobin from endothelium creates two diffusional barriers: a cell-free diffusion barrier along the endothelium in laminar flowing blood and an unstirred bulk diffusional barrier around the erythrocyte membrane. Understanding the role of such barriers and the requirement for a physical separation of hemoglobin from the source of NO production in endothelium helps explain the remarkable morbidity and mortality associated with the use of stroma-free hemoglobin-based blood substitutes and many of the clinical manifestations of hemolytic disease.

Endothelial Dysfunction in Sickle Cell Disease: A Unique State of NO Resistance

In patients who have coronary artery disease, atherosclerosis and its risk factors (obesity, hypertension, diabetes, tobacco smoking, and hypercholesterolemia) a state of endothelial dysfunction is observed, characterized by decreased function of endothelial NO synthase (Figure 1; see color figures, page 546). In patients with sickle cell disease there is a similar dysfunction, characterized by a blunted response to NO synthase inhibition. Unlike patients with atherosclerosis, however, there is also a resistance to exogenously delivered NO donors. This has been demonstrated by multiple investigators in both human and transgenic mouse studies:

- Our group has shown that the blood flow responses to infusions of the NO synthase inhibitor L-NMMA are blunted and that blood flow responses to the NO donor sodium nitroprusside are nearly abolished in patients with high plasma hemoglobin concentrations.
- Eberhardt and colleagues have shown that endothelium-dependent, NO-dependent blood flow is impaired in patients with sickle cell disease, when measured by flow-mediated vasodilation. They also showed that responses to the exogenous NO donor, nitroglycerin, are impaired, compared to control subjects with non-hemolytic anemia.
- Both Nath et al and Kaul et al have described a similar state of resistance to exogenous NO (the NO donor NONOate or sodium nitroprusside) in different transgenic mouse models of sickle cell disease. Kaul and colleagues recently demonstrated that this state of NO resistance correlated with plasma hemoglobin levels and suggested that NO resistance in this model was linked to hemolytic rate and oxidant stress.
- Aslan and Freeman have shown that NO is inhibited in the vasculature of transgenic sickle cell mice with sickle cell disease by a diffusion-limited reaction with superoxide produced from xanthine oxidase on endothelium. Increased xanthine oxidase expression in the lung of the transgenic mouse has also been reported to scavenge NO in this vascular system. Recent studies have suggested a role for vascular NADPH oxidase in aberrant superoxide-mediated NO scavenging in the sickle cell cerebral vasculature.

Because these reactions are so fast and, in the case of Equation 1, irreversible, kinetic calculations would predict that NO produced from endothelium would not survive long enough to diffuse to smooth muscle, becoming inactivated by rapid reaction with intravascular hemoglobin (10 mM heme concentration in whole blood). This paradox of biology is solved by the physical properties of the erythrocyte, of laminar flowing blood, and the plasma hemoglobin scavenging systems that limit extravasation of free plasma hemoglobin into the interstitial space. During normal physiology, the reaction of NO with hemoglobin is limited by compartmentalization of hemoglobin within the erythrocyte membrane. This compartmentalization of hemoglobin from endothelium creates two diffusional barriers: a cell-free diffusion barrier along the endothelium in laminar flowing blood and an unstirred bulk diffusional barrier around the erythrocyte membrane. Understanding the role of such barriers and the requirement for a physical separation of hemoglobin from the source of NO production in endothelium helps explain the remarkable morbidity and mortality associated with the use of stroma-free hemoglobin-based blood substitutes and many of the clinical manifestations of hemolytic disease.

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Equation 1

\[ \text{NO} + \text{oxyhemoglobin (HbFe}^{II}\text{-O}_2) \rightarrow \text{methemoglobin (HbFe}^{III}\text{)} + \text{nitrate (NO}_3^{-}) \]

Equation 2

\[ \text{NO} + \text{deoxyhemoglobin (HbFe}^{II}\text{)} \rightarrow \text{iron-nitrosyl-hemoglobin (HbFe}^{II}\text{-NO}) \]
state and during crisis. In vitro, sickle erythrocytes increase endothelin-1 production by cultured human endothelial cells and endothelin receptor A antagonism decreases the vasoconstrictive effects of conditioned media from pulmonary endothelial cells exposed to sickled erythrocytes on aortic rings. In addition, endothelin-1 activates Gardos channels in human sickle erythrocytes, an effect that may promote sickle cell dehydration and facilitate red blood cell sickling and adhesion.

Intravascular hemolysis has the potential to drive a pro-coagulant state. Platelet activation is profoundly inhibited by NO and such NO-dependent inhibition may in turn be blocked by plasma hemoglobin-mediated NO scavenging. Additionally, hemolytic rate (reticulocytosis) is associated with hemoglobin desaturation (ventilation/perfusion inhomogeneity) and adhesion molecule expression; it is possible that such a hypoxic state can induce hypoxia-inducing factor-1 (HIF-1) dependent factors such as erythropoietin, vascular endothelial growth factor (VEGF), and endothelin-1.

In addition to release of hemoglobin from the red cell into plasma, hemolysis releases erythrocyte arginase, which converts L-arginine, the substrate for NO synthesis, to ornithine. Morris and colleagues found that arginase activities in the plasma of patients correlated significantly with plasma hemoglobin and LDH and was increased in the plasma and red cells of patients with sickle cell disease. Consistent with this observation, in patients with sickle cell disease, the arginine-to-ornithine ratio decreases significantly as pulmonary pressures increase and was independently associated with increasing mortality. Arginine therapy has been shown to decrease pulmonary pressures in patients with sickle cell disease and secondary pulmonary hypertension and has been shown to inhibit endothelin-1 mediated activation of the Gardos channel in the transgenic sickle cell mouse and thus limit erythrocyte dehydration.

These mechanisms likely contribute to the progressive development of sickle cell vasculopathy, characterized by vasoconstriction, intimal and smooth muscle hyperplasia and in situ thrombosis (Figure 2; see color figures, page 546).

Does hemolysis produce a subset of clinical manifestations shared by the hereditary and acquired hemolytic anemias? We have proposed that the clinical manifestations of sickle cell disease may fall into two partially overlapping subphenotypes. The first subphenotype encompasses the more classic clinical manifestations of the disease: vasoocclusive pain crisis and the acute chest syndrome. These clinical morbidities are epidemiologically associated with high white blood cell counts, high steady state hemoglobin levels and low fetal hemoglobin levels (increasing fetal hemoglobin concentration is protective). These “vasoocclusive” complications are widely presumed to be mediated by microvascular obstruction by sickle erythrocytes and the pathogenesis characterized by ischemia-reperfusion injury, infarction and inflammation. The second subphenotype encompasses clinical complications shared by other hemolytic anemias (Table 2) and includes pulmonary arterial hypertension, systemic systolic arterial hypertension, cutaneous leg ulceration, priapism and possibly stroke. Pulmonary hypertension is increasingly observed in hemolytic anemias (Table 2), including sickle cell disease and thalassemia (in particular thalassemia intermedia, such as Hb E-β thalassemia, and inadequately transfused and chelated patients with thalassemia major).

In addition to published case reports of pulmonary hypertension in diseases listed in Table 2 (this extensive list of diseases associated with pulmonary hypertension is not cited due to space limitations, but a pubmed search using the search terms “anemia” and “pulmonary hypertension” will identify reports), our group has received additional reports of patients with pulmonary hypertension associated with hemolytic anemia secondary to unstable hemoglobin variants (personal communication, H. Franklin Bunn and Thomas DeLoughery).

**Pulmonary Arterial Hypertension in Sickle Cell Disease**

### Prevalence

Echocardiographic studies have reported that approximately 30% of screened adult patients with sickle cell anemia have pulmonary hypertension (systolic pulmonary artery pressures (PAP) ≥ 30 mm Hg). Recent autopsy studies suggest that up to 75% of sickle cell patients have histological evidence of pulmonary arterial hypertension at the time of death. Similarly, retrospective studies have demonstrated that 40%-50% of patients with thalassemia

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<th>Table 2. Conditions associated with both intravascular hemolysis and increased risk for pulmonary hypertension.</th>
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<td>Left ventricular assist devices</td>
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<td>Cardiopulmonary bypass devices</td>
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<td>Malaria (?)</td>
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intermedia, and 10%–75% of patients with thalassemia major, have echocardiographic evidence of pulmonary hypertension.

**Risk factors**

In the NIH pulmonary hypertension screening study, all markers of hemolytic anemia, including low hemoglobin and hematocrit, high lactate dehydrogenase (LDH), and high aspartate aminotransferase, but not alanine aminotransferase levels, were associated with elevated pulmonary pressures. Multiple logistic regression analysis identified a history of renal or cardiovascular complications, increased systemic systolic blood pressure, LDH, elevated alkaline phosphatase, and low transferrin levels as independent predictors of pulmonary hypertension. In men, a history of priapism was an additional independent factor associated with pulmonary hypertension. These associated risk factors for pulmonary hypertension suggest that pulmonary hypertension represents one element of the systemic vasculopathy seen in some patients with sickle cell disease (systemic hypertension, renal failure and priapism) and is mechanistically linked to hemolytic rate, iron overload and cholestatic hepatic dysfunction. Interestingly, the development of pulmonary hypertension was not associated with markers of inflammation, fetal hemoglobin levels or platelet counts.

Functional or surgical asplenia may also contribute to the development of pulmonary hypertension in patients with hemolytic disorders. Splenectomy has been reported to be a risk factor for the development of pulmonary hypertension, particularly in patients with hemolytic disorders. It has been speculated that the loss of splenic function increases the circulation of platelet derived mediators and that senescent and abnormal erythrocytes in the circulation trigger platelet activation, promoting pulmonary microthrombosis and red cell adhesion to the endothelium. Intravenous injection of hemolysate promotes the formation of platelet-rich thrombi in the pulmonary vascular bed of rabbits after ligation of the splenic artery, without any thrombus formation in the animals without splenic artery ligation. A role for intensification of intravascular hemolysis following splenectomy (in contrast to predominantly spleen-mediated extravascular hemolysis) has also been suggested by the demonstration of significantly higher plasma hemoglobin and erythrocyte-derived microvesicle levels in patients with thalassemia intermedia who have undergone splenectomy, compared with those who have not.

Interestingly, the number of episodes of acute chest syndrome (a potential cause of chronic lung disease and pulmonary fibrosis) was not associated with pulmonary hypertension in our prospective prevalence study. In addition, a similar prevalence of pulmonary hypertension in patients with thalassemia intermedia, who do not develop the acute chest syndrome, suggests that acute lung injury may worsen pulmonary hypertension but certainly is not etiologic. In our cohort, individuals with pulmonary hypertension have a higher incidence of restrictive lung disease and pulmonary fibrosis on high-resolution chest computed tomography (CT) than age- and hemoglobin-matched patients with sickle cell disease without pulmonary hypertension (Anthi et al, 2005, manuscript under review). Further, in patients with thalassemia, restrictive ventilatory defects and pulmonary fibrosis—associated with pulmonary hypertension—have also been documented. Taken together, these data suggest that similar pathogenic proliferative mechanisms that lead to pulmonary hypertension may underlie the genesis of pulmonary fibrosis in these patients.

**Diagnosis**

Doppler echocardiography provides essential information such as non-invasive estimation of pulmonary artery systolic pressure (via calculation of the tricuspid regurgitant Doppler jet velocity value [TRV]), valvular function and right and left ventricular function. The use of echocardiography to estimate pulmonary artery systolic pressures has been validated in patients with sickle cell disease, and non-invasive assessment correlates well with the measurement of pulmonary arterial pressures by right heart catheterization. The velocity of regurgitant blood across the tricuspid valve during systole is measured, and the pulmonary artery systolic pressure is calculated using the modified Bernoulli’s equation \(4 \times TRV^2 \) plus central venous pressure estimate; method described in detail in 33. To avoid the more subjective estimation of central venous pressures, pulmonary hypertension can be defined by a specific TRV ≥ 2.5 m/sec (based on high risk of death using this value in a prospective cohort study) and moderate-to-severe pulmonary hypertension defined by a TRV ≥ 3.0 m/sec (the more conventional criteria which is consistent with a pulmonary artery systolic pressure of at least 41 mm Hg). The significance of these values has only been defined in adult patients with sickle cell disease; limited information is available for children.

**Prognosis**

Patients with sickle cell disease and pulmonary hypertension have a significantly increased mortality rate compared with patients without pulmonary hypertension. Sutton and colleagues reported a 40% mortality rate at 22 months with an odds ratio for death of 7.86 (2.63–23.4). Powars and colleagues reported a mean 2.5-year survival in sickle cell patients with chronic lung disease with pulmonary hypertension. Castro and colleagues similarly reported a 50% 2-year mortality rate in patients with sickle cell disease with pulmonary hypertension confirmed by right heart catheterization.

Consistent with retrospective studies indicating that pulmonary hypertension is associated with a higher mortality, in the NIH screening study a measured TRV of at least 2.5 m/sec, as compared to a velocity of less than 2.5...
m/sec, was associated with a marked increased risk of death (RR 10.1; 95% CI 2.2–47; P < 0.001) and remained so after adjustment for other possible risk factors in proportional hazards regression analysis. The 18-month mortality was 16% for patients with a TRV of greater than or equal to 2.5 m/sec and was less than 2% in patients without pulmonary hypertension. Further updated follow-up data from this cohort continue to demonstrate that pulmonary hypertension is a strong independent risk factor for mortality (RR 7.4, 95% CI 2.4–22.6, P < 0.001) with 40-month mortality rate of approximately 40% (Figure 3). In addition, De Castro and colleagues reported a remarkably similar 17% mortality rate for patients with pulmonary hypertension over 2 years compared with approximately 2% for subjects without pulmonary hypertension.\(^{59}\) Taken together, the retrospective\(^{40,44,58}\) and prospective\(^{33,41}\) studies strongly support the contention that pulmonary hypertension is the greatest risk factor facing the aging population of patients with sickle cell disease and likely other patients with chronic high-grade intravascular hemolysis.

**Management**

In the absence of clinical guidelines and placebo-controlled therapeutic trials for the evaluation and treatment of pulmonary hypertension in the sickle cell population, we now summarize our empiric and anecdotal diagnostic and therapeutic approach for the adult patient with sickle cell disease diagnosed with pulmonary hypertension. Because we do not yet know if an elevated pulmonary pressure is a direct cause of death or a risk factor for multi-organ disease and generalized sickle cell vasculopathy, for **patients with mild pulmonary hypertension (TRV 2.5–2.9 m/s)** we recommend intensification of sickle cell-specific therapy.

- Consider hydroxyurea treatment at the maximum tolerated dose as defined by the Multicenter Study of Hydroxyurea, with erythropoietin therapy considered if reticulocytopenia limits hydroxyurea therapy.
- Monthly transfusion therapy may be considered for patients with poor responses to hydroxyurea, accompanied by chelation therapy, if indicated. Anecdotally, the TRV has declined in some patients with institution of these treatment measures, although this has not been studied to date.
- Consultation may be considered with a pulmonologist or cardiologist experienced in pulmonary hypertension, the latter especially if the echocardiogram shows evidence of left ventricular dysfunction.
- Identify and treat risk factors associated with pulmonary hypertension such as hypoxemia during rest or exercise and nocturnal hypoxemia, sleep apnea, pulmonary thromboembolic disease, left ventricular systolic and diastolic dysfunction, severe anemia and iron-overload.

In addition to the above measures, we recommend that **patients with TRV ≥ 3 m/s** should undergo:

- Right heart catheterization to assess left ventricular diastolic and systolic function.
- A CT-pulmonary angiogram to exclude chronic thromboembolic pulmonary hypertension.
- Consider systemic anticoagulation. Therapy with warfarin improves outcomes in patients with primary pulmonary hypertension and in-situ thrombosis but no data are available in patients with sickle cell disease.
- Consider specific therapy with selective pulmonary vasodilator and remodeling drugs, particularly if the patient has symptomatic dyspnea on exertion that has progressed in recent months or years. Drugs that are FDA-approved for primary pulmonary hypertension include bosentan (Tracleer\(^8\)) and various forms of prostaglandin therapy, none of which have been comprehensively investigated for sickle cell pulmonary hypertension. We have pilot experience with sildenafil, which has recently gained FDA approval for pulmonary hypertension under the trade name Revatio\(^6,60\). Two multicenter trials using sildenafil and bosentan, for hemolysis-associated pulmonary hypertension are anticipated in the near future. Appropriate consultation and right heart catheterization are recommended at baseline and should be repeated annually. More detailed management recommendations are available in recently published reviews.\(^61-63\)

**Conclusions**

In patients with sickle cell disease, and likely other hemolytic conditions, intravascular hemolysis produces a state of endothelial dysfunction characterized by reduced NO bioavailability and NO resistance. This leads to dysregulation of the endothelium-derived vasodilator:vasoconstrictor system leading to acute vasoconstriction and chronic proliferative vasculopathy. We propose that this vasculopathy is characterized epidemiologically by a clinical subphenotype of pulmonary hypertension, cutaneous leg ulceration, priapism, sudden death, and possibly stroke. Pulmonary hypertension is common in patients with he-

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**Figure 3.** Kaplan-Meier survival curves according to the tricuspid regurgitant jet velocity (TRV). The survival rate is significantly higher among patients with a TRV of less than 2.5 m per second than among those with a TRV of at least 2.5 m per second (P < 0.001). Updated from Gladwin et al. April 2005.\(^33\)
reditary hemolytic anemias and is associated with a high risk of death in patients with sickle cell disease. New therapies targeting this vasculopathy and aimed at normalizing the vasodilator:vasoconstrictor balance are in therapeutic trial.

References

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