CURRENT TRENDS IN THE MANAGEMENT OF SICKLE CELL DISEASE: AN OVERVIEW.

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INTRODUCTION
Sickle cell disease is an inherited disorder of hemoglobin that has a worldwide impact on health and longevity. Sickle cell disease, characterized by lifelong hemolytic anemia and a wide variety of painful and debilitating vaso-occlusive events, occurs in 70000 to 80000 Americans of African, Mediterranean, or Middle Eastern extraction¹.

In the United States, the life expectancy for patients with sickle cell disease is shortened by about 30 years², while in Africa, where comprehensive medical care is less available, death in early childhood is usual. In Nigeria the incidence is 1-2% of the population³.

Medical advances in management of sickle cell disease has lead to significant increases in life expectancy of this group of patients. 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 longevity⁴.

ETIOPATHOGENESIS
The term sickle cell disorder refers to states in which the red cell undergoes sickling when it is deoxygenated. The sickle cell diseases are those disorders in which sickling produces prominent clinical manifestations. Included are sickle cell–hemoglobin C disease (hemoglobin SC disease), sickle cell–hemoglobin D disease (hemoglobin SD disease), sickle cell β-thalassemia, and sickle cell anemia. The latter term is reserved for the homozygous state for the sickle cell gene⁶.

There are few diseases of man whose etiology can be traced to as basic a level as sickle cell disease. Sickle cell anemia is due to the substitution of thymine for adenine in the glutamic acid DNA codon (GAG to GTG), which results, in turn, in substitution at β⁶ of valine for glutamic acid⁶. Deoxygenation of HbS leads to a conformational change that exposes a hydrophobic patch on the surface of the β-globin chain. Binding of this site to a complementary hydrophobic site on a β-subunit of another hemoglobin tetramer triggers the formation of large polymers⁷. This leads to the characteristic sickle or elongated shape of erythrocytes. Repeated or prolonged sickling causes progressive
damage to the erythrocyte membrane which is of primary importance in the pathophysiology and presentation of SCD. Various factors are associated with sickling of red cells which include deoxygenation, vascular stasis, low temperature, acidosis, infections.

When a cell sickles and unsickles repeatedly, the membrane is affected and the cell becomes irreversibly sickled; it remains so even when the oxygen pressure is increased. These are the sickled forms seen on air-dried films. An irreversibly sickled cell has a high hemoglobin concentration and a high calcium and low potassium content, and it may be ATP-depleted. These cells appear to be derived directly from reticulocytes but have a short intravascular life span, and the severity of the hemolytic process is directly related to the number of these cells in a patient's circulation.

CLINICAL FEATURES

The newborn infant is protected by the high level of fetal hemoglobin in the red cells during the first 8 to 10 weeks of life. As the level declines the clinical manifestations of sickle cell disease appear, and the hematologic manifestations of sickle disease are apparent by 10 to 12 weeks of age. Many patients with sickle cell anemia are in reasonably good health much of the time, achieving a steady-state level of fitness. This state of relative well-being is periodically interrupted by a crisis that may have a sudden onset and occasionally a fatal outcome. The early recognition and subsequent clinical assessment of sickle crises are greatly facilitated by familiarity with the patient's steady state.

Various types of crises occur, and these may be classified as follows: vasoocclusive (painful) crisis, aplastic crisis, sequestration crisis, and hemolytic crisis. The vasoocclusive crisis is the most common and is the hallmark of the patient with sickle cell disease. The frequency with which such crises occur varies from almost daily to less than once yearly. The vasoocclusive crises result from complex interactions between endothelium, plasma factors, leukocytes, and rigid, sickled red cells leading to the obstruction of blood vessels.

Other features of sickle cell disease are associated with the chronic state of this disease and complications arising thereof. These include: bony abnormalities like avascular necrosis of femoral head, hypostenuria, renal failure, priapism, autosplenectomy, jaundice, acute chest syndrome, stroke, retinopathy, leg ulcers, and infections.

CURRENT TRENDS IN MANAGEMENT

Neonatal Screening

The demonstration in 1986 that prophylactic penicillin markedly reduces the incidence of pneumococcal sepsis provided a powerful incentive for the widespread implementation of neonatal screening for sickle cell disease (SCD). Neonatal screening, when linked to timely diagnostic testing, parental education, and comprehensive care, markedly reduces morbidity and mortality from SCD in infancy and early childhood. It is imperative that all infants, including those born at home, be screened and that the initial screening test always be obtained prior to any blood transfusion, regardless of gestational or postnatal age. Hemoglobins (Hb) identified by neonatal screening are generally reported in order of quantity. Because more fetal hemoglobin (Hb F) than normal adult hemoglobin (Hb A) is present at birth, most normal infants
show Hb FA. Infants with hemoglobinopathies also show a predominance of Hb F at birth. Those with SCD show Hb S in absence of Hb A (FS), Hb S with another hemoglobin variant (e.g., FSC, FSDPunjab), or a quantity of Hb S greater then Hb A (FSA). Hundreds of other Hb variants may also be identified. Most of these variants are associated with few or no clinical consequences, but some are associated with significant anemia or other problems.

**Genetic Counseling**

Sickle cell trait (SCT) is not considered to be a health problem, but individuals who test positive should be informed about the implications for their health and family planning. There are two major circumstances in which adults will learn that they have SCT:

1. Parents of a child with SCT. When a newborn with SCT is identified through screening, at least one of the parents will have SCT.
2. Pregnant women. During prenatal care, women from racial groups with a high prevalence of the sickle cell gene are tested for the gene.

SCT counseling has two components—education and decision-making—but the emphasis differs in the two cases above. For the first group, the focus is on education, that is, to enable individuals to make informed decisions, in their own interest, about future family planning. For the second group, the focus is on education and informed decisions, in their best interest, about the current pregnancy.

**Psychosocial management**

Sickle cell disease (SCD) is a complex condition that affects the patient, the family, and the patient’s and family’s relationship with health care providers and the community. It is imperative that teaching the skills necessary for coping with this illness begin at the time of diagnosis and continue throughout the life of the patient and that providers recognize that including the extended family and the community in the education process will ensure the most positive outcome. Psychosocial issues confronting patients, families, providers, and the community, though multiple and multifactorial, can be addressed and result in positive patient outcomes.

**Pain**

Severe pain should be considered a medical emergency that prompts timely and aggressive management until the pain is tolerable. The following recommendations are for treatment in the emergency room, day treatment center, or hospital if the patient is admitted directly.

- Begin hydration. Total fluids should not exceed 1.5 times maintenance (including volume for drug infusions). Initial fluid should be 5 percent dextrose + half-normal saline + 20 mEq KCl/L, adjusted for serum chemistry results.
- Assess the patient for the cause of pain and complications. Analgesics are the foundation for the management of sickle cell pain, and their use should be tailored to the individual patient. Sedatives and anxiolytics alone should not be used to manage pain, because they can mask the behavioral response to pain without providing analgesia. Management of pain associated with SCD consists of the use of nonsteroidal anti-inflammatory drugs (NSAIDs), opioids, and adjuvant medications. Management of mild-to-moderate pain should include NSAIDs or acetaminophen, unless there is a contraindication; these are nonsedating, so patient activities can
continue. If mild-to moderate pain persists, an opioid can be added. Treatment of persistent or moderate-to-severe pain relies on repeated assessments and appropriate increases in opioid strength or dose. The type of oral preparation used depends on the characteristics and expected duration of the pain. If the patient’s pain typically is of short duration (less than 24 hours), opioids or formulations with a short duration of action are appropriate, with the advantage of quicker onset of action. For patients whose pain requires several days to resolve, a sustained release opioid preparation is more convenient to take and provides a more consistent analgesia. The combination of nonopioid analgesics with opioids can permit lower doses of the latter. If an opioid like codeine is used, pain relief is accompanied by mild sedation that can facilitate rest.

**Infection**

Infection is a major complication of sickle cell disease (SCD). Special preventive measures against infection exist in addition to routine immunizations; treatment regimens are based on local formularies and antibiotic sensitivity tests. The single most common cause of death in children with SCD is *Streptococcus pneumoniae* sepsis. The unusual susceptibility results from two problems: splenic malfunction, and failure to make specific IgG antibodies to polysaccharide antigens. Two prevention strategies are recommended: vaccination and prophylactic penicillin. The single most important clinical study in SCD in the past 20 years was a randomized, placebo-controlled trial that demonstrated that the administration of penicillin twice a day prevents 80 percent of life-threatening episodes of childhood *Streptococcus pneumoniae* sepsis. There are also immunization schedules for Haemophilus influenza, Neisseiria meningitides, Influenza, and Hepatitis B. Virtually all adults with sickle cell anemia are functionally asplenic, but their immune systems have matured to allow type-specific polysaccharide antibody production. Because they are not as susceptible as children to overwhelming sepsis and the incidence of sepsis is relatively low, there is only anecdotal evidence about preventive strategies. *Streptococcus pneumoniae* vaccination is recommended for adults with SCD. Some patients keep penicillin on hand for fever, but most are not prescribed penicillin prophylaxis routinely. Most antibiotic treatments are started empirically, before culture results are available. There is need for the use of broad spectrum antibiotics for SCD patients with fever, before culture results are available.

**Transient Red Cell Aplasia**

Because the life span of red blood cells is greatly shortened in sickle cell disease (SCD), temporary suppression of erythropoiesis can result in severe anemia. Transient red cell aplasia (TRCA) typically is preceded by or associated with a febrile illness. Between 70 and 100 percent of episodes of TRCA are due to infection by human parvovirus B19, also the cause of *erythema infectiosum* (“fifth disease”). Aplasia is the result of direct cytotoxicity of the parvovirus to erythroid precursors, although other progenitors may be affected in some conditions. No experimental trials have been reported regarding the management of TRCA. Although many patients recover spontaneously, red cell transfusions should be considered for those who become symptomatic.
**Stroke and CNS Disease**

Adults presenting with acute ischemic stroke should be evaluated for tissue plasminogen activator (t-PA) treatment. If t-PA is not used, aspirin is appropriate. Adults with transient ischemic attack (TIA) or ischemic stroke should be evaluated for the cause of the ischemia and therapy should be guided by these findings. Alternatives include antiplatelet agents and warfarin. Chronic transfusion is an option, as used in pediatric stroke prevention. Several uncontrolled studies have documented a reduction in recurrent cerebral infarction using chronic blood transfusion with the target of reducing Hb S to less than 30 percent of total hemoglobin\(^{22,23}\). The reduction in recurrent stroke risk is significant, but patients may still have a stroke despite adequate transfusion and low Hb S levels. The use of aspirin 50-325mg/d for prevention of stroke is also recommended\(^{15}\).

**Eye Disease**

Beginning in childhood, all patients with sickle hemoglobinopathies should have yearly dilated examinations by an ophthalmologist with expertise in retinal diseases. Any patient with a sickle hemoglobinopathy who experiences a change in vision should be referred for ophthalmologic consultation immediately. Central retinal artery occlusion, an event which usually results in permanent, devastating loss of vision, is one of the few bona fide ophthalmic emergencies which demands intervention within minutes to hours after the onset of symptoms. Treatment consists of hyper oxygenation combined with rapid reduction of eye pressure utilizing surgical and medical techniques. Vision loss from hemorrhage or retinal detachment also calls for urgent care, but, unlike acute vascular occlusion, can be appropriately addressed within 24 to 48 hours. Any individual with a sickle hemoglobinopathy who sustains ocular or periocular trauma should be examined immediately by an ophthalmologist because of the increased risk of visual loss from elevated eye pressure associated with hemorrhage into the anterior chamber (hyphema)\(^{15}\).

**Acute Chest Syndrome (ACS)**

ACS is an acute illness characterized by fever and respiratory symptoms, accompanied by a new pulmonary infiltrate on a chest x ray. Because the appearance of radiographic changes may be delayed\(^{24}\), the diagnosis may not be recognized immediately. A major risk factor for the development of ACS is the hemoglobin genotype: the highest incidence is seen in Hb SS genotype (12.8 events/100 person-years) and the lowest in Hb-S +β-thalassemia genotype (3.9 events/100 person years)\(^{25}\). ACS is the second most common cause of hospitalization in sickle cell patients and the most common complication of surgery and anesthesia\(^{26}\).

**Oxygen.** Assessment of blood oxygenation requires determination of baseline arterial blood gases (ABG), and estimation of the alveolar arterial (A-a) oxygen gradient and the PaO2/FiO2 ratio. Oxygen should be administered to moderately hypoxemic patients (PaO2 = 70-80 mm Hg, O2 saturation = 92-95 percent) nasally at a rate of 2 liters per min. Chronically hypoxemic patients in whom the admission PaO2 is no lower than in their steady state may still benefit from oxygen because they may not tolerate additional hypoxemia due to ACS. Control of chest pain and incentive spirometry can prevent hypoventilation in most patients\(^{27}\).

**Transfusions.** Simple transfusions (or exchange transfusions) decrease the proportion of sickle red cells and are
indicated for the treatment of ACS\textsuperscript{24,28}. Transfusions will increase the oxygen affinity of blood in sickle cell patients\textsuperscript{29}. The main indication for transfusion therapy is poor respiratory function. **Antibiotics.** Intravenous broad-spectrum antibiotics should be given to febrile or severely ill ACS patients since it is difficult to exclude bacterial pneumonia or super infection of a lung infarct. **Other measures.** Optimal pain control and incentive spirometry are important to prevent chest splinting and atelectasis.\textsuperscript{15}

**Biliary Disease**

Biliary sludge is best managed by serial ultrasound examinations at 12- to 24-month intervals unless cholestasis occurs; at that point, laparoscopic cholecystectomy is indicated. Elective laparoscopic cholecystectomy has become the procedure of choice for symptomatic cholelithiasis\textsuperscript{30} because of the shortened hospital stay, lower cost, and fewer immediate surgical complications. One approach to asymptomatic or minimally symptomatic cholelithiasis is careful observation until symptoms dictate surgery. Bacteremia, ascending cholangitis, empyema, and other hyper acute biliary complications require surgery on a more urgent basis, consistent with good surgical practice.\textsuperscript{15}

**Renal disease**

The kidney in patients with sickle cell disease (SCD) exhibits numerous structural and functional abnormalities, changes that are seen along the entire length of the nephron. These abnormalities include hypostenuria, hematuria, acute renal failure, glomerular abnormalities and chronic renal failure. Patients with SCD should therefore be encouraged to drink liberal amounts of liquids in order to compensate for the fluid loss that is brought on by hypostenuria. The treatment of hematuria in SCD involves bed rest, maintenance of a high urinary flow documented by monitoring of intake and output, and, if blood loss is significant, iron replacement and/or blood transfusion. Vasopressin and epsilon-amino caproic acid (EACA) have both been used with variable success\textsuperscript{31,32}. However, caution must be exercised when using EACA, as this antifibrinolytic agent may predispose to the formation of clots that can obstruct the urinary collecting system. If prolonged and life-threatening bleeding is coming from one kidney, local resection of the bleeding segment is preferred. Unilateral nephrectomy is a last resort, since bleeding may recur from the other kidney. Hematuria that occurs in SCD is not always a consequence of red cell sickling and papillary necrosis. Other, nonsickling causes also should be considered. For example, renal medullary carcinoma in young subjects with SCD and sickle cell trait (Hb AS) has been reported\textsuperscript{33,34}. Therefore, a thorough evaluation is recommended when hematuria is initially found in individuals with SCD and Hb AS. Proteinuria, which can progress to the nephrotic syndrome, is the most common manifestation of glomerular injury in SCD patients. Moreover, as many as 40 percent of SCD-SS patients with nephrotic syndrome may go on to develop end-stage renal disease (ESRD)\textsuperscript{35}. Therefore, patients with persistent proteinuria should have a urine collection obtained for the determination of 24-hour protein excretion, and a nephrology consultation should be requested for consideration of other, nonsickling causes of proteinuria and possible renal biopsy. ACE inhibitors ameliorate pathological changes such as perihilar
focal and segmental glomerulosclerosis. They also decrease urinary protein excretion in patients with early manifestations of sickle cell nephropathy. Renal insufficiency occurs earlier in SCD-SS patients than it does in SCD-SC patients. Factors that appear to predict renal failure in SCD-SS patients include hypertension, proteinuria, increasingly severe anemia, and hematuria. Finally, the risk of renal failure is increased in those SCD-SS patients with the Central African Republic (CAR) s-gene cluster haplotype. As there is no proven treatment for sickle cell nephropathy, every attempt should be made to slow its rate of progression. The amount of proteinuria can be decreased by the administration of ACE inhibitors, and it is conceivable that the progression of sickle cell nephropathy may be slowed by a prolonged course of these drugs. Patients should avoid nonsteroidal anti-inflammatory drugs (NSAIDs), because NSAIDs have been shown to produce significant declines in the rates of glomerular filtration and renal blood flow in patients with SCD. Effective control of blood pressure has been reported to slow the progression of end-stage renal disease (ESRD) in patients with SCD; they should be treated with standard approaches. Because dehydration can precipitate vasoocclusive events, caution should be exercised in the use of diuretic agents in an individual with obligate hyposthenuria. Every effort must be made to avoid additional renal damage due to urinary tract infection. Infection must be recognized and treated vigorously. Follow-up should be maintained longer than for patients without SCD.

Although erythropoietin levels are generally high in steady-state SCD-SS patients, they are not increased to the level that would be expected for the degree of anemia. One explanation for the relatively decreased erythropoietin levels is the right-shifted hemoglobin-oxygen dissociation curve seen in SCD patients. Erythropoietin levels in SCD patients fall still further as renal function worsens and these patients may require substantially higher doses of erythropoietin than are required for patients with other forms of ESRD. If erythropoietin is ineffective, transfusions can be given; they must be done carefully, however, to avoid volume overload. As with all patients who develop ESRD, SCD patients can be treated with both hemodialysis and peritoneal dialysis, and they can undergo renal transplantation.

**Priapism**

Beginning in early boyhood, males need to know that priapism is one aspect of SCD and that this is not an event that should embarrass them. Boys and young men, as well as their families, need to know that they should be prepared to seek medical attention as soon as an episode begins and that if untreated, priapism can result in impotence in the future. The males should know that a full bladder can trigger priapism, and they therefore need to urinate regularly. They also should avoid prolonged sexual activity, which can trigger an episode. If they have had more than one episode, medications can be prescribed that may prevent recurrences. The goal of therapy is to ease pain, make the erection go away, and preserve future erectile function. If treatment is given within 4 to 6 hours, the erection can generally be reduced with medication and conservative therapy. Patients should be advised to drink extra fluids, use oral analgesics, and attempt to urinate as soon as priapism begins.
For episodes lasting more than two hours, patients should go to the emergency room to receive intravenous hydration and parenteral analgesia. According to one protocol, if detumescence does not occur in 1 hour after the patient has arrived in the emergency room, penile aspiration is initiated (procedure should be performed within 4 to 6 hours from onset of priapism). The patient receives conscious sedation and local anesthesia; blood is then aspirated from the corpus cavernosum with a 23-gauge needle followed by irrigation of the corpora with a 1:1,000,000 solution of epinephrine in saline. The concomitant use of automated red cell exchange transfusions to reduce the sickle hemoglobin (Hb S) level to less than 30 percent can also be considered, especially if early intervention with irrigation fails. If there is recurrence despite aspiration and local instillation of vaso-active drugs, shunting may be considered. In this procedure, known as the Winter procedure, a shunt is created between the glans penis and the distal corpora cavernosa with a Tru-cut biopsy needle; this allows blood from the distended corpora cavernosa to drain into the uninvolved corpus spongiosa. Additional medications used for reversal of priapism include -agonists. If impotence persists for 12 months, the patient may wish to consider implantation of a semi rigid penile prosthesis. Some physicians prescribe 30 mg of oral pseudoephedrine at night as an attempt to prevent further episodes in those who have had priapism and have required aspiration and irrigation. Injections of leuprolide, a gonadotropin releasing hormone analogue that suppresses the hypothalamic-testicular axis and the production of testosterone, also has also been used with some degree of success as prophylaxis against further episodes. A small (11 patients) double-blind, placebo-controlled crossover study found that oral stilbestrol in doses of 5 mg daily for 3 to 4 days could abort episodes of priapism and that much smaller doses could prevent recurrence. Although hydroxyurea may potentially be of benefit, clinical studies to determine its efficacy in preventing priapism have not been performed.

Musculoskeletal Abnormalities
Musculoskeletal manifestations of sickle cell disease (SCD) are common and may lead to severe morbidity. Bone and joint involvement result from three main causes: 1) bone marrow hyperplasia, which causes distortion and growth disturbance, particularly in the skull, vertebrae, and long bones; 2) vaso-occlusive events that lead to infarction of metaphyseal and diaphyseal bone and to osteonecrosis of juxta-articular bone; and 3) hematogenous bacterial infection that results in osteomyelitis and septic arthritis. Conservative management by analgesia, rehydration, transfusions and appropriate antibiotics are recommended for uncomplicated cases. Surgical intervention for those with advanced disease and for whom all other conservative measures have failed is recommended. These include, core decompression and osteotomy for osteonecrosis, hip replacement, prosthesis and surgical drainage for septic arthritis and osteomyelitis.

Leg Ulcers
Between 10 and 20 percent of patients with sickle cell disease (SCD) due to a homozygous hemoglobin S (Hb S) genotype (SCD-SS) develop painful, disfiguring, and indolent leg ulcers. The ulcers usually appear between ages 10 and 50 years and are seen
more frequently in males than in females. Studies to prove the efficacy of treatment of leg ulcers are difficult to perform. One reason is that healing depends on blood circulation, and the cumulative time of bed rest and leg elevation is not easily monitored. In addition, the variable extent of wound debridement is difficult to quantify, and a short period of dependency could erase any gains made in the previous period.

Ankle ulcers are painful, and the patient should be given moderately potent analgesics such as oxycodone. Bed rest and elevation of the leg to reduce edema are useful, though not always practical. Wet-to-dry dressings, even if applied only 2 or 3 times a day, can provide gentle debridement; cooperation of patients increases when they are permitted to dampen the dressing slightly before removal, since it is a painful process. Oral zinc sulfate (200 mg 3 times a day) probably does no harm if it does not cause nausea, and may be worth using. The ulcer size should be measured at every clinic visit; seeing the dimensions shrink can provide encouragement to the patient. Some ulcers will not heal. Rigorous studies have not been done to assess the utility of transfusions for treating leg ulcers, but the ulcers seem to correlate with degree of anemia, which suggests transfusions may help. They should be considered for recalcitrant or recurrent skin ulcers if conservative therapy fails. If transfusions are used, they probably should be continued for 3 to 6 months, and a hemoglobin concentration above 10 g/dL with Hb S levels less than 50 percent can be achieved. More complete bed rest, systemic antibiotics, transfusions, and skin grafts sometimes help. If split thickness or pinch grafts are to be used, preoperative preparation of the ulcer bed is probably quite important.

Quantitative bacterial cultures of biopsies of the bed and margin are advocated by some but not all surgeons as a guide to the time for surgery. Intravenous arginine butyrate infusions have been reported to cause rapid healing of ankle ulcers.

**Pregnancy**

Prenatal care for women with SCD should be managed by a multidisciplinary team that includes an obstetrician, nutritionist, primary care physician, and hematologist. The team must decide who will be responsible for each aspect of the patient’s care. Close monitoring, combined with prompt diagnosis and aggressive treatment of complications during the prenatal and neonatal period by a multidisciplinary team, will contribute to better outcomes. The clinical problems of SCD, such as new onset seizures, hepatopathy, acute anemia, and painful episodes should be evaluated and managed for pregnant women in the same fashion as for women who are not pregnant. Women who have had previous pregnancy losses or who have multiple gestations may benefit from the early use of transfusions to maintain a hemoglobin level above 9 g/dL.

**Anaesthesia and Surgery**

- Make sure the operating and anesthesia teams are aware of the diagnosis of a sickle cell syndrome and the need for special attention.
- In patients with SCD-SS, and SCD-SC, simple transfusion to a hemoglobin of 10 g/dL should be performed before all but the lowest-risk procedures.
- Patients with SCD, regardless of genotype, should all receive careful attention, with preoperative monitoring of intake and output, hematocrit,
peripheral perfusion, and oxygenation status.

- Intraoperative monitoring of blood pressure, cardiac rhythm and rate, and oxygenation should be conducted for all surgical procedures.
- Postoperative care should include attention to hydration, oxygen administration with careful monitoring, and respiratory therapy.

**Transfusion**

Used correctly, transfusion can prevent organ damage and save the lives of sickle cell disease (SCD) patients. Used unwisely, transfusion therapy can result in serious complications. Once a decision is made to transfuse, the type of red cells to be given is specified and goals are set for the final post transfusion hematocrit and percent sickle hemoglobin (Hb S) desired. In general, phenotypically matched, sickle negative, leukodepleted packed cells are the blood product of choice, and a post transfusion hematocrit of 36 percent or less is recommended, since a higher value theoretically causes hyper viscosity, which is dangerous to sickle cell patients. Transfusions are used to raise the oxygen carrying capacity of blood and decrease the proportion of sickle red cells. Clinically, they will improve micro vascular perfusion of tissues.

**Induction of Fetal Hemoglobin**

Enhanced concentrations of hemoglobin F (Hb F) can inhibit sickle hemoglobin (Hb S) polymerization and red cell sickling and improve the clinical course of sickle cell disease (SCD). In a cooperative study of the natural history of patients with sickle cell anemia in the United States, the frequency of pain episodes correlated inversely with the Hb F concentration.

Based on these epidemiologic studies and understanding the biophysics of Hb S polymerization, a search was launched for pharmacologic agents that could reverse the switch from γ- to β-globin chain synthesis in erythroid precursors. The first "hemoglobin switching" agent was the nucleoside analog 5-azacytidine, which was followed by butyrate derivatives as compounds which increase Hb F gene expression. Other drugs, such as hydroxyurea and erythropoietin, alter maturation of erythroid precursors and promote Hb F production indirectly. When both hydroxyurea and erythropoietin were given to patients with sickle cell anemia, an increment in Hb F concentration beyond that seen with hydroxyurea alone occurred.

**Hemopoietic cell transplantation**

Hematopoietic cell transplantation (HCT) has curative potential for a broad spectrum of genetic disorders, including sickle cell disease (SCD). The goal is to eliminate the sickle erythrocyte and its cellular progenitors and replace them with donor hematopoietic pluripotent stem cells that give rise to erythrocytes that express no sickle hemoglobin (Hb S), thereby reducing Hb S levels to those associated with the trait condition. The possibility of preventing serious complications from SCD, which can cause extensive morbidity and early death, is balanced by the risk of severe adverse events after transplantation. Children with SCD who experience significant, noninfectious complications caused by vaso-occlusion should be considered for HCT, and if full siblings are available, HLA typing should be performed.

There are other measures which are still in the developmental stage that will be helpful in the management
of patients with sickle cell disease. These include:
Clotrimazole- which inhibits Gardos channel pathway of sickle hemoglobin56.
Nitric oxide- reduces polymerization tendency of sickle hemoglobin57.
Flucor (ReothRx- poloxamer 188) - a new drug under investigation that is said to be useful in reducing the duration of acute pain crises, thereby shortening hospital stay58.
Gene therapy- a process whereby the defective gene of the sickle hemoglobin is replaced with normal gene.

CONCLUSION
Without major breakthroughs in gene therapy or bone marrow transplantation that make these treatments applicable to a large number of patients, drug intervention will remain the major therapeutic option for sickle cell disease. The likelihood is low of finding a "magic bullet" medication that substantially improves sickle cell disease for all or even most patients. Treatment likely will involve the use of different agents alone or in combination to produce optimal results. Combination therapy currently is not an option for sickle cell disease, since only hydroxyurea has been proven to alter the course of the condition. Many, if not most of the agents currently under investigation likely will fall short of investigators' hopes. If a few survive the rigors of testing and join the clinical armamentarium, however, we could mix and match drugs for patients with sickle cell disease. Ideally, the treatment regimens would include drugs with differing modes of action. Hydroxyurea, for instance, combined with clotrimazole would team a drug that enhances fetal hemoglobin production (hydroxyurea) with one that reduces erythrocyte dehydration (clotrimazole). For a particular patient, sickle cell symptoms might be improved substantially by neither drug alone. The combination, however, might significantly ameliorate the condition59. A major goal of investigation should be development of interventions that can be used in very young patients. Many of the problems experienced by adults and adolescents with sickle cell disease reflect incremental organ damage by bouts of hypoxia. The affected areas may initially be microscopic. With time, these foci of injury coalesce to form regions of macroscopic injury, such as avascular necrosis of the femur. Prevention must be the watchword as we seek to improve the management of patients with sickle cell disease.

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