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Chapter 1

Introductory Chapter: Introduction to the History, Pathology and Clinical Management of Sickle Cell Disease

Baba Inusa, Maddalena Casale and Nicholas Ward

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1. Introduction

Sickle cell disease (SCD) is due to a single point mutation (Glu6Val) that causes polymerisation of the mutant hemoglobin (Hb) S, resulting in sickling of erythrocytes [1]. Inflammation, haemolysis, microvascular obstruction and organ damage characterise the clinical expression of SCD, which is highly variable in individual patients. Environmental and multiple genetic factors modify many aspects of SCD and therefore contribute to the clinical variability [2, 3].

This chronic, complex and monogenetic haematological condition that leads to haemolytic anaemia, severe acute complications and chronic organ damage was formally reported in 1910 [4]. One hundred years after Herrick’s initial report, there have been significant advances in the diagnosis and management [5].

Forty years ago only half of children with sickle cell anaemia were expected to reach adulthood [6]. However, advances in the diagnosis and management of SCD have improved the survival and quality of life significantly. Recent figures suggest survival of 53 years for men and 58.5 years for women [7, 8] and over 94% surviving to adulthood [8, 9].

SCD is now one of the most common genetic disorders in the world with high prevalence in sub-Saharan Africa (SSA), the Middle East, the Mediterranean and the Indian subcontinent [1, 10, 11]. In the United States, almost 1 in 10 African Americans has the sickle cell trait, with 1 in 600 being homozygous for the sickle hemoglobin allele [12, 13]. In the UK, it is diagnosed at birth, through routine Newborn screening with incidence of about 1 in every 2000 births [3, 14]. However despite the high prevalence in sub-Saharan Africa (SSA) essential public health programmes have not been implemented due to limited medical resources and infrastructures [10, 11, 15, 16]. As a consequence, infant and childhood mortality due to SCD remains high. It is estimated that without intervention, up to 50-90% of affected infants may die by 5 years of age from SCD [11, 15, 17, 18]. While high regional disease prevalence would
be expected to facilitate epidemiologic, translational and clinical research, the majority of SSA countries lack the capacity to provide the comprehensive care for SCD [11].

2. Early description

James Herrick is widely acknowledged as the first physician to describe a case of sickle cell. However, the term ‘sickle cell anaemia’ was not coined until 1922 [19], when a review of the first four cases of the disease was conducted, including that described by Herrick [20, 21].

However, it is argued that case reports, up to 60 years prior to that by Herrick, may have described sickle cell disease [6, 17]. A post-mortem, performed in 1898, of a patient who died in hospital after being admitted for pains, jaundice and previous leg ulcers shows that these symptoms are suggestive of SCD [22]. It has also been proposed that the clinical illness had been previously recognised in Africa as “cold-season rheumatism” and that the possibility of genetic inheritance of the illness had even been noted [10].

3. Discovery of molecular and genetic basis

It was not until 40 years after the first description of sickle cell anaemia that a homozygous pattern of inheritance was confirmed simultaneously by two separate studies, one a pedigree study in Africa [11, 12]. These studies demonstrated that sickle cell anaemia was inherited in an autosomal recessive pattern. In the late 1940s, Pauling discovered that blood from sickle cell patients had differing mobility during electrophoresis [23, 24]. This indicated that there were chemical differences between the hemoglobin of the different cells. Thus, Pauling and colleagues proposed that sickle cell anaemia was a molecular disease, the first disease of this kind [23].

Vernon Ingram was to be the discoverer of the exact difference between sickle cell and normal adult hemoglobin. Ingram identified that sickle hemoglobin (HbS) was more positively charge hemoglobin A (HbA) [21, 25]. The amino acid sequence of hemoglobin was shown to differ by only one substitution of glutamic acid with valine in the β chain of HbS [26]. As knowledge of the pathophysiology of SCD is a complex phenomenon between vaso-occlusion and hemolysis [27–29].

4. Prophylactic penicillin and pneumococcal vaccination

Studies have shown that infant mortality is particularly high in sickle cell disease. Scott’s observations revealed that the number of children reaching adulthood was only approximately 50% [4] in the 1960s. The Cooperative Study of Sickle Cell Disease (CSSD) has shown that the peak incidence of death was between the ages of 1 and 3 [30]. Furthermore, the CSSD showed that the largest killer in SCD was infection, with this being the cause of death in 50% of subjects being followed below the age of 20 [20]. Infection was also shown to be a predominant factor in
other studies of mortality \cite{21, 22} with pneumococcal disease being identified as a major killer \cite{19, 20}. It has even been calculated that the risk of developing meningitis from \textit{Streptococcus pneumoniae} is 30 times more likely in the sickle cell population than in the general population \cite{23}.

The increased risk of infection may be attributed to many factors, including decreased immunoglobulin function, poor cell-mediated immunity and reduced opsonisations \cite{24–27}. Importantly, there is a decrease in splenic function as a result of microvascular occlusions and infarctions \cite{24}. The decrease in immune function leads to a vulnerability to encapsulated organisms particularly \textit{Streptococcus pneumoniae}, \textit{Haemophilus influenzae} type B and \textit{Salmonella} \cite{28}.

Following the recognition of infection playing a major role in the disease, a study commencing in 1983 showed the benefit of prophylactic penicillin. Gaston and colleagues recruited sickle cell patients under the age of 3 to a randomised control study to determine if daily prophylactic penicillin would be effective at reducing childhood infections. Astonishingly it was found to reduce septicaemia by 84\% and the trial was terminated early. The Penicillin Prophylaxis in SCD study (PROPS) therefore recommended that all children should be screened and started on penicillin by the time they reached 4 months of age \cite{31}.

In addition to prophylactic antibiotics, pneumococcal vaccination has also become an important part of sickle cell management. Those immunised had lower incidence of pneumococcal septicaemia than the control group \cite{32, 33}. The introduction of new conjugated vaccines against other encapsulated bacteria (13 strains of \textit{Streptococcus pneumoniae}; \textit{Meningococcus} A, C, W, Y and B and \textit{Haemophilus influenzae}) has reduced the rate of pneumococcal bacteraemia by 93.4\% in children aged <5 years to 134 cases per 100,000 person-years, $P<0.001$ \cite{33–35}. Incidence of invasive pneumococcal disease among individuals with sickle cell disease before and after the introduction of the pneumococcal conjugate vaccine \cite{32, 35}.

5. Prenatal and Neonatal SCD Screening

SCD is associated with high infant mortality rate; therefore, early preventive intervention is essential in reducing these deaths \cite{14, 36}. A maternal blood screen during or preceding pregnancy seems the most cost-effective method of initiating a screening programme, as if the mother does not carry the sickle allele, then there is no risk of the child having sickle cell disease and additionally this initial screen does not need repetition for future pregnancies. If maternal screening reveals a sickle cell trait or SCD, the father can be invited for investigation \cite{14, 36}.

Since 1978 it has been possible to obtain foetal DNA to diagnose sickle cell disease prenatally \cite{18}, via amniocentesis or chorionic villus sampling (CVS). CVS offers the advantage that the family is given the choice early in pregnancy. Prior to this it was only possible to diagnose sickle cell disease via direct viewing of foetal blood. In some countries it has been in use for national screening programmes and rates of termination of pregnancy are as high as 74\% in Cuba \cite{37–40}.
The importance of neonatal identification of SCD is emphasised [36, 41, 42].

By 1994 the majority of United States cities had implemented universal neonatal hemoglobinopathy screening [12, 43]; the UK has only fully implemented its universal sickle cell screening programme in the last few years [14, 44]. In Africa, only Ghana and the Republic of Benin have established selective and pilot Newborn screening programme, respectively [37, 41, 45, 46].

5.1. Pain management

The clinical severity of SCD is variable and usually defies genetic or phenotypic explanation. The cooperative study of the natural history of sickle cell disease showed that about 5% of patients accounted for one-third of hospital days devoted to pain control [19]. Acute VOC and chronic pain syndrome could be disabling to the patient, and studies have demonstrated an association of acute pain syndromes with other complications of SCD, including death [19, 47].

Pain management guidelines and the need to rule out life-threatening comorbidities suggest the need for rapid physician evaluation of patients who present to the emergency department (ED) with complaints related to SCD [47]. The American Pain Society guidelines recommend initial assessment and analgesia within 30 min of arrival [5, 48].

5.2. Hydroxyurea (HU)

From the difficulty in identifying sickle-shaped erythrocytes in neonates, Janet Watson postulated, in 1948, that high fetal hemoglobin leads to a decrease in cells that could sickle and therefore a decrease in sickle hemoglobin in those red blood cells [49]. Several years later it was suggested that by increasing or prolonging the concentration of foetal hemoglobin in erythrocytes, it would be possible to decrease the frequency and severity of the clinical manifestations of SCD [50–52].

HU, an anti-leukaemia agent with the capacity to increase foetal Hb, was introduced to treat sickle cell disease with significant clinical and symptomatic improvement. In 1992 a randomised double-blind study, the Multicentre Study of HU in Sickle Cell Anaemia (MSH), showed that this new drug could decrease episodes of painful crises, acute chest syndrome and the need for transfusion [53, 54]. The study recruited 299 adults with more than three painful crises in the year prior to the start of HU, and the clinical effect was great enough for the trial to terminate 3 months early [54, 55].

The paediatric studies on the use of hydroxyurea reflected the results of the adult trials, with reduced acute complications and with no toxicity or decreased growth, and suggested reduced end organ damage if administered early [56–58].

6. Stroke and stroke prevention

Stroke is an important cause of morbidity and mortality of SCD with an estimated 11% having had a stroke by the age of 20 [59]. Although the treatment of strokes in SCD had been
improving, it was a ground breaking trial in the final decade of the twentieth century that changed the way SCD was monitored [60, 61] to prevent stroke in this group of patients.

Landmark's work into the use of transcranial Doppler (TCD) ultrasounds showed that severe sickle cell disease could lead to the narrowing of cerebral blood vessels. These would appear on ultrasonography as increased flow in that region and patients with abnormal flow are at increased risk of stroke [60, 62]. Silent cerebral infarcts are the most common neurologic injury in children with sickle cell anemia and are associated with the recurrence of an infarct (stroke or silent cerebral infarct), and regular blood transfusion therapy significantly reduced the incidence of the recurrence of cerebral infarct in children with sickle cell anemia by 58% [63, 64].

7. Pulmonary hypertension

Pulmonary hypertension (PAH) is recognized to be a common complication of SCD and other hemolytic disorders and is associated with increased mortality and morbidity. Retrospective studies suggest a prevalence of PAH ranging from 20 to 40% [65, 66] among adults, and a prospective study showed that prevalence of PAH in pediatric population is 31%, and the two-year mortality rates approach 50% [65, 66].

Chronic hemolysis represents a prominent mechanistic pathway in the pathogenesis of SCD-associated PAH via nitric oxide (NO) scavenging and abrogation of NO salutary effects on vascular function. These processes lead to acute and chronic pulmonary vasoconstriction [65]. Many known infectious risk factors for PAH, i.e. human immunodeficiency virus (HIV) infection, chronic hepatitis B and C viral infections and possibly malaria, are hyper endemic in African countries where the prevalence of SCD is very high. Interactions between these infectious complications and SCD-related hemolysis could yield an even higher incidence of PAH among the African SCD patients.

Self-reported history of cardiovascular and renal complications, systolic hypertension, high lactate dehydrogenase levels (index of hemolysis), high level of alkaline phosphatase, low transferrin concentration (indicating iron overload) and priapism in men were found to be independent correlates of PAH [20, 67].

Therapy for SCD-related PAH remains challenging. Treatment with HU at maximum tolerated dose and judicious use of blood transfusion support and iron-chelating agents where indicated is recommended [66]. While HU is a potential NO donor effect, it has not been shown to impact mortality of SCD-related PAH [51]. There is increasing interest in the use of sildenafil in SCD-related PAH. Sildenafil was well tolerated by both male and female patients and it reduced the estimated pulmonary artery systolic pressure and increased the 6-min walk distance [68, 69].

Gene therapy has the potential to correct the underlying defect leading to the clinical manifestations of sickle cell disease. It would reduce the need for many of the preventative treatments and invasive therapies. A few years ago, this may not have seemed possible; however, gene
therapy has been proven to be effective in mouse models [1, 2], and there are currently several teams working on phase II trials using viral vectors [70–72].

8. Global perspectives

The World Health Organization has estimated that there may be approximately 216,000 babies born with HbSS disease in Africa each year and that the disease may account for 10–20% of neonatal mortality in West Africa, very little is known about the overall global burden of SCD [68]. Although there have been improvements in the management of SCD in developed countries, much less progress has been made in the developing world in which the disease is common, where it is still an important cause of childhood mortality. In low-income countries, basic facilities for management are lacking, systemic screening is not a common practice, and diagnosis is made late.

Research collaborations between sickle cell research groups in rich and poor countries would be of considerable benefit to the health and well-being of patients with SCD in the developing countries. In particular, interactions of this kind offer an opportunity to improve clinical and diagnostic facilities and hence the management of patients in the developing world. We cannot yet cure SCD, but we have learnt that simple interventions significantly improve morbidity and mortality.

9. Conclusion

Although there has been dramatic improvement in the diagnosis, management and monitoring of sickle cell disease over the last century, bone marrow transplantation (BMT) is currently the only cure for SCD with a curative success rate of approximately 90–93% [68, 69]. Conversely, there are severe risks and complications including rejection, marrow aplasia, neurological disorders, graft-versus-host disease and death, and standard BMT is limited by a requirement for human leukocyte antigen (HLA)-matched sibling donors [70]. Currently transplantation is only considered for those children with HLA-identical siblings and in whom the clinical manifestation of the disease does not respond to standard care.

The past 100 years have shown great accomplishments for SCD including newborn screening, penicillin prophylaxis, primary stroke prevention using blood transfusion, stroke prevention trial in sickle cell anaemia (STOP) and controlled trial of transfusions for silent cerebral infarct in sickle cell anaemia. The role of hydroxyurea therapy in adults and children to reduce pain, hospitalisations and most recently as an alternative for controlling cerebral blood velocity— transcranial Doppler (TCD) with Transfusion Changing to Hydroxyurea (TWiTCH) therapy. Although a risk-free cure has not been found in this last and current century, the following 100 years look to be extremely promising for sickle cell anaemia; stem cell transplantation is now more commonly used in SCD.
Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

MC and NW reviewed the literature and wrote the draft of the manuscript, BI designed, reviewed and made the substantial changes the manuscript. All authors discussed, read and approved the manuscript.

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Author details

Baba Inusa¹²*, Maddalena Casale³ and Nicholas Ward²

*Address all correspondence to: baba.inusa@gstt.nhs.uk

1 Paediatric Haematology, Guy’s and St Thomas’ Hospital, London, United Kingdom
2 King’s College London, London, United Kingdom
3 The Second University of Naples, Caserta, Naples, Italy

References


[34] Preventing infections in sickle cell disease: The unfinished. 2016;63(5):25911.


