Dental health in sickle cell disease

S. M. AlDallal*, M. M. AlKathemi1, W. H. Haj2 and N. M. AlDallal4

1Haematology Laboratory Specialist, Amiri Hospital, Kuwait
2Dentist, General Practitioner, Dental One Clinic, Kuwait
3Paediatric & Handicapped Dental Specialist, Basma Clinic, Kuwait
4General Surgeon, Farwaniya Hospital, Kuwait

Abstract

Sickle cell disease (SCD) is one of the most common blood disorders typically inherited from one’s parents. It is presented with a wide variety of clinical symptoms, and varied degrees of severity that can be determined based on the phase during which the disease is diagnosed, the age of the patient, number of hospitalisations in the past, requirement for continuous drug use and for blood transfusions, in addition to several other factors. It is highly critical that the physicians should be aware of the oral manifestations and physiopathology of the disease. Additionally, the dental surgeons should cautiously obtain the patient’s clinical history and collect information about specific features so that they can build up a plan for any dental treatment that is in accordance to the patient’s limitations and requirements. Maintaining a complete chart recording the general patient information along with periodically updating the medical history of the patient should be practise by all the physicians. The treatment strategy should focus on the achievement and maintenance of oral health and to decrease the risks of dental complications. The literature summarizes the treatment of dental complications in patients with SCD.

Introduction

Sickle cell disease (SCD) is one of the most prevalent genetic disorders worldwide. Studies have reported a count of approximately 100,000 Americans being affected by this disease. It is also estimated that one in 50 US African American births is affected with sickle cell anaemia. The prevalence of the disease is highest in sub-Saharan Africa and is also widely spread through Middle East, Southeast Asia, and Mediterranean regions [1]. Around 5-7% of the population worldwide carries an abnormal haemoglobin gene with the frequency of SCD noted as one in four in every 50 West Africans and making itself the commonest genetic disease in UK and France [2,3]. SCD is the most prevailing form of hemoglobinopathy. The disease is characterized by morphologic changes in red blood cells (RBCs), triggered by abnormal haemoglobin (Hb) polymerization. The sickle mutation is the consequence of single base change, GAG to GTG, in the sixth codon of exon 1 of the β-globin gene in charge for the synthesis of the β-globin polypeptide of the Hb molecule (α2β2). This change cause substitute of the normal glutamic acid with valine at position 6 of the β-globin chain and the development of sickle Hb (βsβs) [4-6]. Acute infections can activate sickle cell crises. Therefore, it is imperative that dental infections should be prevented but, if there is an occurrence of infection, then effective ways of dealing with it should be devised immediately. A clear understanding of the dental implications of SCA must be gained in order to successfully treat SCA patients. The treatment should always begin only after a thorough investigation on the patient’s background has been performed.

Clinical presentation of SCD

The signs and symptoms of SCD vary from one patient to the other and may change over time. A key aspect of the disease is vaso-occlusive crises of the microcirculation, which leads to inadequate blood supply to tissues and consequently results in tissue necrosis [7]. Pain is the main and the most chronic feature of this disease and dominates its clinical representation throughout the life of the patient. Patients frequently report pain in form of acute pain crises. It has a substantial impact on the quality of life of the patient and their families. Therefore, several medical and psychological treatment techniques are widely devised and used for pain management in patients affected with SCD [8,9].

In SCD, all body organs are affected or remain at the constant risk of being affected. As such, the most prevalent complications include diseases of nervous, cardiopulmonary, musculoskeletal, hepatobiliary, endocrine and genitourinary systems [10]. The most frequent oral manifestations of SCD greatly affect the oral mucosa, gingival tissue, mandible, osteonecrosis, facial swelling, increased risks for caries nerve supply, and tooth enamel and pulp [11-19] (Table 1). In addition, paleness of the oral mucosa, delayed tooth eruption, depapillation leading to atrophic alteration of the tongue, high degree of abnormality in the hypophosphatemic teeth, odontogenic infection, orofacial pain, craniofacial disorders such as protrusion of the midface area, maxillary expansion, mandibular retrusion, and maxillary protrusion. Dentists play a significant role in avoiding these complications and improving the quality of life in SCD patients as the SCD patients are more vulnerable to infections and periodontal disease [20,21]. Furthermore, these patients are at a higher risk of developing dental caries leading to elevated occurrence of dental opacities arising due to the unremitting...
Table 1. Dental complications of SCD [9].

<table>
<thead>
<tr>
<th>Dental Complications of SCD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dental Caries</td>
</tr>
<tr>
<td>Infractions</td>
</tr>
<tr>
<td>Hypodontia</td>
</tr>
<tr>
<td>Dental Erosions</td>
</tr>
<tr>
<td>Malocclusions</td>
</tr>
<tr>
<td>Pulp Necrosis</td>
</tr>
<tr>
<td>Abnormal Trabecular Spacing</td>
</tr>
<tr>
<td>Infection</td>
</tr>
</tbody>
</table>

use of medication containing sucrose owing to the high incidence of complications and hospitalization required by the lack of proper oral cleanliness [22]. Managing dental complications is frequently ignored as SCD patients are more dedicated towards maintaining a standard general health because of the serious blood disorder. Ignoring minor dental health matters under these conditions not only worsen the problem but can also cause a painful sickle-cell crisis, leading to emergency hospital admission. Therefore, the management of oral complications in SCD patients require to be modified in accordance to their blood disorder, in order not to cause any additional deterioration to their overall health.

Dental complications in SCD

The dental complications in sickle cell anaemia must be understood at first to efficiently treat SCD patients. The management and treatment of the patients should begin with a thorough assessment of the patient’s background. Studies have revealed that common oral findings associated with the disorder include mucosal pallor, dental hypoplasia, delayed eruption, and radiographic changes [23]. Furthermore, hypercementosis, which leads to excessive generation of normal calcified tissue on the roots of one or more teeth, has also been reported in SCD patients [24]. Malocclusion of the teeth, intrinsic opacity of enamel, dental caries, and diastemata are other dental remarks found in the disease in sicklers [25]. Details on some of these are described below.

Dental caries

Dental caries are the most frequent dental complication worldwide. Patients with SCD are more susceptible to dental caries having more chances of tooth decay [26]. Caries is identified as an infectious disease of teeth causing progressive demineralization and destruction of the enamel, dentin, and cementum of the teeth [9]. The key source of caries is the acidification of the oral environment, which is caused by the fermentation of remaining food particles mainly sugars or carbohydrates on tooth surfaces [27]. Untreated caries can cause slow destruction and tooth fractures. It may also further lead to infection of the surrounding oral soft tissues and may also transfer to ear, neck and jaw [28]. Sometimes, it may be as fatal as causing cavernous sinus thrombosis (CST) leading to the blood clot in the cavernous sinus, the reason being complication of an infection in teeth [29].

SCD patients appear to be more vulnerable to dental caries than healthy people. Also, the occurrence of caries in SCD children and adults are known to associated with the socioeconomic status of the affected families [22-30]. Low income most frequently affects the amount of value given on lifestyle, health and access to medical care and health information. Past results have shown that low-income individuals are more susceptible to decayed teeth problems as compared with individuals without the disease. On the other hand, Passos et al. [31] reported that the disease itself does not particularly predispose to caries or periodontal disease. Risk the contributing factors including daily smoking, older age, and lack of daily dental flossing can attributed towards dental caries and periodontal diseases.

Hypodontia

Hypodontia, also known as tooth agenesis, refers to missing teeth as a result of failure for them to develop. It has been described in one patient with HbS SC disease [32]. These patients must be subjected to preventive care to lessen the chances of any functional complications.

Dental erosion

Dental Erosions, also known as acid erosions, result from acidity that is not caused by bacteria but by food particularly acidic fruit juices. Intrinsic causes of acidity comprise certain disorders such as gastroesophageal reflux where gastric acid is in contact with teeth. Erosions have not been described in SCD patients [33,34].

Malocclusion

Malocclusion, in dentistry, refers to the manner opposing teeth meet. The father of modern orthodontics, Edward Angle, introduced the term malocclusion which is defined as a misalignment of the teeth between the two corresponding dental arches (Figure 1A) [35]. He divided malocclusion into three classes in accordance to the severity and the relation between the teeth involved. In Class I, the occlusion in a SS patient is normal for the first maxillary molar but abnormal for the other teeth with crowding, rotation, and spacing and over or under eruption (Figure 1B). In Class II, inconsistent jaw growth is observed in a patient with HbSS whereas the lower jaw is deficient in forward growth causing an overjet<3 mm (Figure 1C). This increases the chances of dental trauma as compared to those with an overjet<3 mm In Class III, malocclusion in a patient with HbSS, it has been observed that the lower jaw extends too far forward ahead of the upper jaw causing an under-bite, which is also known as mandibular prognathism (Figure 1D) [36]. Reports have shown that Class II malocclusion was most frequent in SCD patients [9]. Furthermore, facial growth alterations which are frequent in SCD might also result in malocclusion because of protrusive maxilla and forward growth tendency of the mandible [37,38].

Malocclusion can be corrected by the application of braces to the protruding teeth. The braces are metallic or ceramic sets of wire that are cemented to the teeth for 2-3 years or a particular time period as stated by the orthodontists [9]. It is advised that the braces procedure and treatments must be done at childhood stage in order to avoid speech problem and bite issues. It should be noted that when a SCD affected patient is undergoing orthodontic treatment, the practitioner be completely aware of the disease and the respective treatment procedure as a complete blood supply is highly important during and after the application of intraoral and extroral forces. The orthodontist should be very careful about the chances of pulpal necrosis involving healthy teeth, the alterations in the bone turnover during orthodontic movements, the mandibular painful episodes, and the increased susceptibility to infections during the treatment.

Infection

Poor dental cleanliness can lead to tooth decay or cavities and also results in to the formation of plaque on teeth when food particles interact with bacteria normally found in the mouth. Plaque that remains on the teeth for a long time hardens and forms calculus. The
formation of plaque and the associated bacteria can not only lead to infection in the gums and teeth, but also affect the gum tissue and bone that support the teeth. Persistent plaque and calculus cause also gingivitis, thereby leading to inflammation of the gums and bleeding. If proper treatment is not adopted, gingivitis can proceed to have serious complications such as chronic periodontitis, dental abscess, and bone destruction culminating in tooth loss [40]. Periodontal infection may precipitate painful vaso-occlusive crises and increase the frequency of hospital admissions among adult SCD patients [41,42].

**Pulp necrosis**

Pulp necrosis, or dead pulp, is defined as necrotic dental pulp due to infection, trauma or chemical reaction characterized by no response to thermal stimulation. A necrotic dental pulp causes toothache, acute apical periodicities, discoloration of the tooth or dental abscess [43,44]. Previous reports suggest that HbSS is a potential risk factor for pulp necrosis in intact permanent teeth [44,45]. A reduced blood supply to teeth results in necrosis of the dental pulp in patients with SCD. Such abnormal blood flow to the dental pulp might result in increased toothaches in SCD patients. Javed et al. have also stated that there are also chances that a certain proportion of SCD patients may remain asymptomatic to pulp changes, which may make them unsuspecting of the ongoing dental pulp tissue damage.

**Conclusion**

This mini review was designed to present brief background information on SCD, and describing the course of oral complications in SCD patients. The dentist’s aim should be to treat the SCD patient with a thorough understanding and knowledge of the disorder and the consequences of the disease must be considered carefully before dental treatment is started. One of the dentist’s goals should be to instil a positive attitude in the SCD patient and their parents toward maintaining good dental health. In addition, it is always advisable that sickle-cell anaemia carriers should be encouraged to have their oral health under control by practicing preventive procedures as directed by the physicians.

**Conflicts of interest**

None to be declared.

**Acknowledgments**

The authors are thankful to www.manuscriptedit.com for providing English language editing and proofreading services for this manuscript.

**References**

4. Ingram VM (1956) A specific chemical difference between the globins of normal human and sickle-cell anaemia haemoglobin. Nature 178:729-794. [Crossref]
5. Ingram VM (1959) Abnormal human haemoglobins III. The chemical difference between normal and sickle cell haemoglobins. Biochim Biophys Acta 36:402-411. [Crossref]
6. Ingram VM (1957) Gene mutations in human haemoglobin: the chemical difference between normal and sickle cell haemoglobin. Nature 180: 326-328. [Crossref]
24. Okafor LA, Nomoo DC, Ojehanon PI, Aikhionbare O (1986) Oral and dental complications of sickle cell disease in Nigerians. Angle Orthodontics 56: 43-49. [Crossref]
32. Passos CP, Santos PR, Aguiar MC, Canguussu MC, Toralles MB, et al. (2012) Sickle cell disease does not predispose to caries or periodontal disease. Spec Care Dentist 32: 55-60. [Crossref]
41. Laurence B, Haywood C Jr, Lanzkron S (2013) Dental infections increase the likelihood of hospital admissions among adults with sickle cell disease. Community Dent Health 30:168-172. [Crossref]