Thalassemia, a human blood disorder

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Abstract

A group of inherited blood defects is known as Thalassemia is among the world’s most prevalent hemoglobinopathies. Thalassemias are of two types such as Alpha and Beta Thalassemia. The cause of these defects is gene mutations leading to low levels and/or malfunctioning α and β globin proteins, respectively. In some cases, one of these proteins may be completely absent. α and β globin chains form a globin fold or pocket for heme (Fe++) attachment to carry oxygen. Genes for α and β-globin proteins are present in the form of a cluster on chromosome 16 and 11, respectively. Different globin genes are used at different stages in the life course. During embryonic and fetal developmental stages, γ globin proteins partner with ε globin and are later replaced by β globin protein. Globin chain imbalances result in hemolysis and impede erythropoiesis. Individuals showing mild symptoms include carriers of alpha thalassemia or the people bearing alpha or beta-thalassemia trait. Alpha thalassemia causes conditions like hemolytic anemia or fatal hydrops fetalis depending upon the severity of the disease. Beta thalassemia major results in hemolytic anemia, growth retardation, and skeletal aberrations in early childhood. Children affected by this disorder need regular blood transfusions throughout their lives. Patients that depend on blood transfusion usually develop iron overload that causes other complications in the body systems like renal or hepatic impairment therefore, thalassemias are now categorized as a syndrome. The only cure for Thalassemias would be a bone marrow transplant, or gene therapy with currently no significant success rate. A thorough understanding of the molecular basis of this syndrome may provide novel insights and ideas for its treatment, as scientists have still been unable to find a permanent cure for this deadly disease after more than 87 years since it is first described in 1925.

Keywords: thalassemias, beta globin genes, hemoglobinopathies.

Resumo

Um grupo de defeitos sanguíneos hereditários é conhecido como talassemia e está entre as hemoglobinopatias mais prevalentes do mundo. As talassemias são de dois tipos, como talassemia alfa e beta. As causas desses defeitos são as mutações genéticas que levam a níveis baixos e/ou proteínas de globina com mau funcionamento, respectivamente. Em alguns casos, uma dessas proteínas pode estar completamente ausente. As cadeias de globina α e β formam uma dobra ou bolsa de globina para a fixação de heme (Fe++) para transportar oxigênio. Os genes das proteínas α e β-globina estão presentes na forma de um cluster nos cromossomos 16 e 11, respectivamente. Diferentes globin genes são usados em diferentes estágios do curso de vida. Durante os estágios de desenvolvimento embrionário e fetal, as proteínas γ globina se associam à ε globina e, posteriormente, são substituídas pela proteína β globina. Os desequilíbrios da cadeia de globina resultam em hemólise e impedem a eritropoiese. Indivíduos que apresentam sintomas leves incluem portadores de talassemia alfa ou as pessoas com traços de talassemia alfa ou beta. A talassemia alfa causa condição como anemia hemolítica ou hidropsia fetal fatal, dependendo da gravidade da doença. A beta talassemia principal resulta em anemia hemolítica, retardo de crescimento e aberrações esqueléticas na primeira infância. As crianças afetadas por esse distúrbio precisam de transfusões de sangue regulares ao longo da vida. Os pacientes que dependem de transfusão de sangue geralmente desenvolvem sobrecarga de ferro que causa outras complicações nos sistemas do corpo, como insuficiência renal ou hepática, portanto as talassemias...
A dentist named Cooley first reported a progression of iron deficiency in infants leading to splenomegaly and bone deformation soon after their birth in 1925 (Cunningham, 2010).

In 1932, Whipple and Bradford explained the pathology of the disease for the first time, and as most of the patients were found to be native to the Mediterranean range, they termed the condition as “thalassemia” (Rachmilewitz and Giardina, 2011).

Blood disorder types characterized by low levels or missing normal globin chains in the normal red blood cell protein hemoglobin are now characterized as thalassemia. There are four types of globin chains present named alpha (α) beta (β) gamma (γ) and delta (δ). Depending on which chain production is disturbed, the thalassemias are termed as α-, β-, γ-, δ-, δβ-, or γδβ-thalassaemias. Mostly inherited as a recessive trait, the most common types are α- and β-thalassaemias, that result from the deficiency of α- or β-globin proteins that are necessary for the production of normal hemoglobin molecule (HbA, α2β2) in an adult human (Taher and Cappellini, 2014).

As defined, thalassemia is a complex of various hereditary disorders of hemoglobin combination featuring insufficient production of at least one of the globin chains driving imbalanced globin-chain production, damaged hemoglobin eventually causes anemia (Cappellini et al., 2014). Hemoglobin is a metalloprotein (Hb or Hgb) found inside the red blood cells (RBCs) of all the vertebrates (except a fish family Channichthyidae) and some invertebrates, functions as an oxygen-transporting vehicle (Burmester and Hankeln, 2014).

Inside the blood, hemoglobin acts as a carrier that conveys oxygen from lungs or gills to entire body tissues and upon reaching there, it unloads the oxygen that in turn is used in aerobic respiration to produce ATPs to run the metabolic processes of an organism (Zhao et al., 2019).

A hemoglobin molecule inside the mammalian RBCs could bind up to four oxygen molecules at a time, which in turn increases the blood oxygen binding capacity (OBC) seventy times, as the OBC (C₉₉₉) of hemoglobin alone is 1.34 O₂ mL/gm Hb (Ali, 2018). Some amount of oxygen may also dissolve directly into the blood during respiration but that is only 1.5% of total oxygen carried through the blood (Molnar and Gair, 2013).

Hemoglobin is involved in transporting other gases too, it binds a portion of carbon dioxide (CO₂) gas which is produced as a result of metabolic activities, in the form of carboxyhemoglobin, i.e., hemoglobin loaded with CO₂, and contributing a fraction of 20–25% of the total CO₂ exhaled (Hughes and Pride, 2012). Hemoglobin additionally carries nitric oxide (NO), which is an essential regulatory molecule, NO is bound to a thiol group of globin protein, discharging along with oxygen (Biagioli et al., 2009).

Hemoglobin is not only confined to the red blood cells (RBCs) but it can also be found in progenitor cells of RBCs, dopaminergic neurons of group A9 present in the ventrolateral midbrain of some rodents and primates, macrophages, cells of alveoli, and mesangial cells supporting the glomerular tuft inside the kidney are the other cells which contain hemoglobin. In these tissues, hemoglobin functions as an antioxidant and metabolic regulator to some extent rather than an oxygen supplier (Hughes and Pride, 2012).

The functional, practical exercises performed by hemoglobin other than gas transport include fertilization, signaling, adjustment of inflammatory reactions to defend and protect the cell. These exercises are proficiently executed while Hb is sequestered securely inside the boundaries of the RBCs. Outside the RBC limits, Hb disaggregates and jeopardizes the cell life as in severe cases, antioxidants become overwhelmed to clear free radicals that are produced during Hb oxidation (Mairbäurl and Weber, 2012).

Hemoglobin is a protein with four polypeptide subunits with 64,500 Da molecular weight. It consists of two α- and two β-chains of 141 and 146 amino acids each, respectively (Figure 1). In adults, the hemoglobin’s secondary structure of all polypeptide chains is mainly α-helical. HbA does not contain any β-strands or disulfide bonds. Unusually, it also lacks isoleucine. The β-chain of HbA comprises eight helical fragments, designated by letters A–H. The α-chain is quite similar but lacks helix D. Each polypeptide chain of HbA maintains a three-dimensional conformation, known as globin fold, just like a closely related monomeric protein, myoglobin (Mb). The globin fold is, in fact, an arrangement of helices due to which a cavity is formed that holds and attaches to a heme (Fe⁡⁺⁺) prosthetic group. A covalent bond is present between the Nᵦ atom of the F8His residue and the heme iron inside each polypeptide of the globin subunits. This heme/iron atom that is in the ferrous state has the potential to reversibly bind with specific gas ligands.

**Figure 1.** Ultrastructure of the hemoglobin molecule.
like oxygen, carbon monoxide, and nitric oxide (Hassan and Alhajouj, 2011). As already mentioned, hemoglobin comprises two α and two β-chains. Therefore, it can be said that it is a dimer of alpha-beta dimers (αβ dimers). The αβ, and αβ dimers are connected by a 2-fold pivot of symmetry. The interfaces of subunits αβ, and αβ, dimers are identical, and αβ, and αβ, subunit images are mirror images of each other (Hamamy and Al-Allawi, 2013).

The human embryo and fetus normally develop with different hemoglobin variants i.e., Hb Portland-1 (Gφ2), Hb Gower-1 (Gφ2), Hb Gower-2 (Gφ2) and fetal Hb or HbF (Gφ2) (Kohne, 2011). Stage-specific combinations of globin genes are transcribed within the fetus during gestation, which leads to producing various forms of hemoglobin molecules such as several embryonic hemoglobins, hemoglobin A (HbA) and, hemoglobin F (HbF) (Shown in Table 1). Immature erythrocytes produce the embryonic hemoglobins (Gower 1, Gower 2, and Portland) in the yolk sac. They persist up to the 12th week of pregnancy. In the fifth week, Hemoglobin F (Hb F) appears which predominates during fetal life. Hb F is synthesized initially in the liver of the fetus and remains there for several weeks after birth. HbA prevails after birth, which originates in the bone marrow. Most common among other variants is Hemoglobin A (Gφ2) usually with an amount of around 95% (Kaufman et al., 2021). The normal range of hemoglobin A2 (Gφ2) is 1.5–3.5% of the total hemoglobin in an adult. Though Hb A2 has no physiological importance in an adult its level may rise in case of alpha or beta chain reduction (Figueiredo, 2015) the mutations in delta globin genes along with beta-globin gene variations may also interfere with the beta-thalassemia diagnosis especially in the cases of beta-thalassemia trait (Harirahan et al., 2016). The gamma chain is synthesized late in the last trimester and at this stage, the Hb F (Gφ2) is only restricted in a small number of RBCs known as ‘F-cells’. Interestingly, in individuals having sickle cell anemia and beta-thalassemia, the Hb F level is quite elevated (Kaufman et al., 2021).

All genetic hemoglobin disorders fall under the broad term “hemoglobinopathy”. These are split further into two main categories, which are:

- Thalassemia syndromes (resulting in severe anemia and other related complications);
- Structural variants of hemoglobin (malfunctioning hemoglobin proteins).

Both occur due to mutated and deleted segments of α- or β-globin genes. Thalassemia is caused due to defects in the globin genes, which results in Hb synthesis disorders i.e., under or overproduction of globin chains. The structure of Hemoglobin in such cases remains normal. Abnormal hemoglobin only occurs when there is a change in the structure of Hb. There are many combinations of both forms like ββ′-thalassemias, Hb/Sickle disease, and HbE/α-thalassemias. The conventional capabilities of the pathophysiology and various ailments patterns are restrained, resulting in limited possibilities to summarize them (Giardine et al., 2014).

The pathogenic hemoglobin protein also indicates the cells and organ types before and after birth in which it is originated (Table 2) (Mehdi and Al Dahmash, 2011; De Sanctis et al., 2017). Based on the affected globin chain, the thalassemias are arranged into alpha (α) or beta (β) types. The β thalassemia is found throughout the world while α thalassemia is more prevalent in the Mediterranean region, (Lippi and Mattiuzzi, 2020, De Sanctis et al., 2017) and China (Qin and Wu, 2009; Muncie Junior and Campbell, 2009).

1. Alpha Thalassemia

There are 2 copies of the alpha-globin gene in the human genome both located on chromosome 16, therefore in a normal diploid cell, 4 copies of the gene are available, to produce the protein. Alpha-thalassemia is caused by an underproduction of α-globin proteins due to mutation or deletion of one of the four α globin genes (Leung and Lao, 2012).

There are two phenotypes of α thalassemia recorded so far. α thalassemia I or minor and α thalassemia II with no clinical symptoms of thalassemia. It is now known that α thalassemia I is associated with the complete absence of α globin proteins and the other is merely a reduction in a globin expression. These two α thalassemia variants are now designated as thalassemia α° and thalassemia α+, respectively as shown in Table 3 (Haley, 2017).

**Alpha (0) thalassemia** – there are more than 20 different mutations described, resulting in the deletion of all the sets of α-globin genes. People having this deficiency are unable to synthesize normal α-globin and therefore cannot make any normal functioning A, F, or A2 hemoglobin. This prompts the onset of hydrops fetalis or “hemoglobin Bart”; children born with this disorder do not survive outside the uterus.

**Alpha (+) thalassemia** – more or less fifteen genetic mutations are reported, which result in limited α-globin protein synthesis generally because of the functional deletion of at least one alpha-globin gene.

Alpha (+) thalassemia is further sub-classified into four categories:

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**Table 1.** Various globin genes and their subunits.

| Subunit | Gene | Chromosome | Expression |
|---------|------|------------|------------|
| ζ-Globin | HBφ1 | 16 | Embryonic |
|         | HBφ2 |   |           |
| α-Globin | Hβα1 | 16 | Fetal and adult |
|         | Hβα2 |   |           |
| ε-Globin | Hβε | 11 | Embryonic |
| γ-Globin | HBG1 (Ay) | 11 | Fetal |
|         | HBG2 (GY) |   |           |
| β-Globin | HBB | 11 | Adult |
| δ-Globin | HB D | 11 | Adult |
A- Thalassemia (-a/aa) occurs when three out of four functional α-genes are inherited. The individuals are referred to as asymptomatic carriers for α-thalassemia. Various terms used for this disorder are “alpha thalassemia minima”, “alpha thalassemia-2 trait”, and “heterozygosity for alpha (+) thalassemia minor”. These carriers are clinically normal or may have mild anemia;

B- Thalassemia (-a/-a) condition is known as a trans-deletion because two healthy alpha genes are inherited, one from each of the two chromosomes (-a/-a) whereas homozygosity for alpha (+) thalassemia (aa/--) (two on the same chromosome) is known as a “cis deletion” resulting in “alpha thalassemia minor” or “alpha thalassemia-1 trait”. Parents who are carriers of the cis deletion can have one out of four (25%) babies affected with alpha thalassemia major in every pregnancy;

C- Hemoglobin H: when only one healthy alpha gene (-a/-a) is inherited, it results in the massive production of hemoglobin H (Hb H) comprising tetramers of surplus β-chains. The disorder is named “Hb H disease”;

D- Hemoglobin Bart’s disease: When all four alpha genes are lost, a situation occurs, that is life-threatening. Four gamma-globin chains are formed during the life of the fetus inside the womb which unites to frame irregular hemoglobin known as ‘hemoglobin Bart’s’ (Camacho et al., 1999).

Alpha thalassemia is different from beta-thalassemia in its pathophysiology. The shortage of α chain results in excess production of gamma or beta chains, which form Hb Bart’s
Table 1

| Subtype                          | Description                                                                 |
|----------------------------------|-----------------------------------------------------------------------------|
| **1. Alpha thalassemia minor**   | It is an asymptomatic carrier condition that occurs due to the deletion of the one α-globin gene. This condition usually causes no symptoms or signs of anemia and does not need treatment due to negligible alpha protein deficiency; therefore, the hemoglobin appears to be normal (Leung and Lao, 2012). The term “silent carrier” is usually used to describe this condition because it is not easily diagnosed through standard hematological investigations. Only DNA analysis could detect this condition (Camacho et al., 1999). |
| **1.1. Alpha thalassemia minor** | The trait is also known as mild alpha-thalassemia. The patients are deficient in two alpha-globin genes. The affected individuals have RBCs smaller than usual and are mildly anemic but do not show any symptoms and may only be diagnosed by routine tests (Leung and Lao, 2012). |
| **1.2. Alpha thalassemia trait** | The trait is also known as mild alpha-thalassemia. The patients are deficient in two alpha-globin genes. The affected individuals have RBCs smaller than usual and are mildly anemic but do not show any symptoms and may only be diagnosed by routine tests (Leung and Lao, 2012). |
| **1.3. Alpha thalassemia intermedia** | Also, it is known as hemoglobin H disease. Individuals lacking three alpha globin genes become severely anemic and mostly cannot survive without blood transfusion. Newborns who inherited alpha thalassemia intermedia seem healthy at birth but mainly develop anemia and splenomegaly as they approach the second year of his life. Hepatomegaly is not commonly reported, and there might be some relationship to the mental retardation in affected individuals. As hemolysis occurs in this type of anemia the tendency to develop respiratory infections, gallstones, and leg ulcers increases. Bone deformities are not usually found in hemoglobin H disease (Lee et al., 2010). The imbalanced alpha and beta chain synthesis (that is usual) induces aggregation of beta chains inside the RBCs. Usually, beta chains are coupled with alpha chains only. Alpha thalassemia with three-gene deletions causes the beta chains to accumulate in gatherings of four, creating unusual hemoglobin, called “hemoglobin H”. This condition leads to “hemoglobin H disease”. This hemoglobin variant has two issues. First, it does not convey oxygen efficiently, making it practically useless to the cell. Secondly, hemoglobin H protein harms the cell membranes of RBCs, accelerating cell death. A combination of reduced alpha chain synthesis and red cell lysis in hemoglobin H disease creates severe and fatal anemia. Without treatment, most individuals do not survive and expire in their early teens or before (Camacho et al., 1999). |
| **1.4. Alpha thalassemia major** | “Hydrops fetalis” or alpha thalassemia major is a condition in which no alpha genes are found in the patients’ genome, resulting in four gamma-globin chains production by the fetus that produces malfunctioning hemoglobin known as hemoglobin Bart’s. Most affected individuals having Hemoglobin Bart’s cannot survive or otherwise die in just a few hours after birth (Lee et al., 2010). Alpha thalassemia with four deletions in the gene has rarely been diagnosed in the uterus, especially in a family with a history of the disorder occurring in early childhood. Reportedly, some of these children have been saved through blood transfusions during pregnancy (Camacho et al., 1999). |
| **2. Beta Thalassemia**          | Around 200 mutations of the beta-globin gene have been identified worldwide which produce beta-thalassemia. Unlike alpha thalassemia syndromes where deletion is usually the root cause, beta-thalassemia occurs due to mutations that influence all stages of beta-globin protein synthesis including transcription, translation, and beta-globin production durability. Two types of β-thalassemia, β° and β+ thalassemia are identified so far that lack beta chain production altogether (Table 1). Beta thalassemia major usually results when β° or β+ thalassemia occurs in homozygous condition. Occasionally, however, the compound heterozygous state for both β° and β+ thalassemia results in beta-thalassemia. In the case of homozygous β° thalassemia, there is no Hb A, an abundance of HbF, and variable amounts of Hb A2. In individuals with homozygous β+ thalassemia, the amount of Hb A is variable, Hb F is increased and distributed heterogeneously among RBCs whereas Hb A2 is normal, decreased, or elevated (Taylor et al., 2012). The molecular variations in β thalassemia result in missing or diminished β chain generation. Alpha chain synthesis remains unaffected, and therefore there is an unequal amount of globin chain generation that prompts an abundance of α chains. They are not stable in the absence of their normal partners and precipitate in the RBC precursors, which interferes with RBC processing. As a result, there is a variable level of intramedullary destruction of RBCs precursors (i.e. ineffective erythropoiesis). The RBCs having α chain incorporated when enter the bloodstream interfere with their segment via microcirculation, exclusively in the spleen. Such cells demonstrate a high variability in the structure of membrane and penetrability and are short-lived. Hence, anemia occurs due to both abnormal erythropoiesis and shortened cell survival. The anemia stimulates erythropoietin production resulting in bone marrow expansion, which consequently causes deformed skull and large bones ref needed. Since the spleen is overburdened and required to remove a continuous stream of abnormal red cells, enlarges before it exhausts ultimately (Borgna-Pignatti and Gambirini, 2011). Generally, three categories of beta-thalassemia have been recognized, ranging from mild to severe by affecting the patient’s body. |
2.1. Beta thalassemia minor

Also known as thalassemia trait due to one of malfunctioning beta-globin genes, but this generally causes no significant problem in the proper functioning of hemoglobin protein (Hay and Weatherall, 2017). When there is an excess of alpha chains, the mechanism that switches off gamma chain expression does not work effectively, thus the levels of Hb F remain somewhat elevated in these patients. The alpha chains consolidate with the accessible beta chains bringing about diminished levels of hemoglobin rest of the surplus alpha chains empower the over-production of delta chains (Jha and Jha, 2014) Affected individuals have a 1:1 chance to pass the thalassemia minor trait to their child (Memon, et al., 2017).

2.2. Beta thalassemia intermedia

A condition where the absence of beta polypeptide in the hemoglobin is sufficient to bring about more extreme anemia and serious medical issues, including shortness of breath, bone disfigurements, mild jaundice, and an enlarged spleen. The condition is characterized by having two abnormal genes in affected individuals while still producing some beta-globin. Depending on the level and functional competence of beta-Globin is a broad range in the clinical severity of this disease (Hay and Weatherall, 2017).

2.3. Beta thalassemia major

It is also known as “Cooley's anemia” and is the most severe form of beta-thalassemia with absent beta-globin synthesis thus preventing the production of significant amounts of Hb A. The severe irregularity of globin chain synthesis (alpha >> beta) brings about extreme microcytic hypochromic anemia. Within the RBCs surplus unpaired alpha-globin chains precipitate, this harms the plasma membranes of RBCs and brings about intravascular hemolysis. Besides, premature death/apoptosis/lysis/necrosis of erythroid precursors reduces the number of RBCs even further. The severe anemia results in hypoxia and the resulting EPO causes hyperplasia in the bone marrow and will lead to extramedullary hematopoiesis (Cunningham, 2010).

During childbirth, the infant with thalassemia major appears to be healthy. This is because there is a predominance of fetal hemoglobin (Hb F) during gestation, which lacks any beta chains. Anemia starts to appear a few months after birth, as the infant switches over from gamma to beta globulin. The infant's growth retards and the resulting EPO causes hyperplasia in the bone marrow and will lead to extramedullary hematopoiesis (Cunningham, 2010).

Regular blood transfusions and extensive continuous therapeutic care are required throughout life in this type of anemia. After some time, these successive transfusions prompt iron overload in the body. Without treatment, this overabundant iron will be stored in the liver, heart, spleen, and other organs and could prompt a sudden death due to major systemic failure (Cunningham, 2010).

Silent carriers of Alpha thalassemia lack signs or side effects of the disorder. Individuals affected by alpha or beta thalassemia disorders might have a slight iron deficiency. It might be an indication. Side effects might be more adverse in the expecting women, or the people with anxiety, or malnourished. Evidence may include fatigue. This might be the main side effect that a person with beta-thalassemia minor shows. Exhaustion is created by the diminished oxygen-conveying limit of the RBCs, bringing about reduced oxygenation for cells and tissues and causing pale skin tone due to insufficient oxygen in the blood (Bhatia et al., 2014).

Over the past three decades, regular blood transfusions have dramatically eliminated the complicity of thalassemia and bone marrow transplantation, enhanced the quality of life-permitting normal development throughout childhood and extended life span. But, transfusion results in a complication due to iron overload (Haidar et al., 2011). Regular blood transfusion leads to iron overload-related complications including hormonal complications such as growth retardation, sexual immaturity, diabetes mellitus, and insufficiency of the parathyroid, thyroid, pituitary, adrenal glands, dilated myocardopathy, liver fibrosis, and cirrhosis (Behera et al., 2014; Mokhtar et al., 2013; Goulas et al., 2012; Borgna-Pignatti and Gambineri, 2011).

In non-transfused thalassemic patients, the spleen, liver, heart, and bone marrow become significantly enlarged was stated before. Expansion of marrow cavities and thinning of cortices produce a variety of bone abnormalities in patients who are not optimally transfused (Bhatia et al., 2014). The result of bone biopsies from non-transfused thalassemic patients shows osteoporosis with increased bone resorption, decreased mineralization, and fewer bone-forming sites (Behera et al., 2014).

Over the past few decades, there has been a tremendous advancement in the field of clinical and genetics research. Many countries have nearly wiped out the disease by making better decisions like mass testing for hemoglobinopathies before marriage or childbirth. Thalassemia is now being considered to be treated beyond bone marrow transplantation which has always been the last hope to survive for a thalassemia patient. Today, allogeneic hematopoietic stem cell transplant (HSCT) from human leukocyte antigen (HLA)-matched sibling or other donors is the only treatment for thalassemia patients with >90% transfusion independent survival rate in the patients transplanted with sibling matched donors. However, the treatment is possible in the individuals at a very young age. The disease-free survival rate depends upon the factors like HLA- matching, age and iron overloading etc. (Soni, 2020). Scientists have developed better tools to treat genetic disorders like stem cell technology and gene therapy to avoid tissue rejection in recipients. The therapy adds a corrective gene whose product combines with α-globin to produce functional hemoglobin, thereby reversing the ineffective red blood cell production seen in β-thalassemia. However, the technique is much costly.
(~ $1.6 billion) which could not be afford to be by the patient’s family or the healthcare providers (Harrison, 2019).

A novel CRISPR technology to edit faulty genes is a new game changer that showed promising results in disease models that made it a new hope to the diseased. The CRISPR (Clustered Regularly Interspaced Short Palindromic Repeats) along with the CRISPR-associated system, Cas (known as Crispr-Cas) is a powerful gene editing tool which has revolutionized the field of molecular biology in terms of gene therapies to treat hereditary genetic disorders (Lino et al., 2018). CRISPR Cas system enables programmable targeting of single base insertion or deletion (Indels) at a particular site of the genomic DNA. Frangoul et al. (2021) has successfully used the Crispr tool to edit the hematopoietic stem cells in order to downregulate the β globin gene to reduce the faulty beta globin chains and enhance the reactivation of fetal hemoglobin (G globin protein) to overcome the anemia eventually reducing the blood dependency in the thalassemia patients. The patients under trial were transplanted with these edited stem cells. The trial had shown promising results and no discrepancy has been reported so far.

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