Assessment of Genetics Mutations in Genes CDAN1, SEC23B and Dell-15q22 in Inducate Congenital Dyserythropoietic Anemia Syndrome

Keywords: Generalized congenital dysrhythmia anemia syndrome; CDAN1; SEC23B genes; Blood disorders

Abstract

CDA syndrome is a genetic disorder that affects the growth of red blood cells. This syndrome is one of the anemia disorders characterized by a lack of red blood cells. Researchers have identified three types of CDA syndromes, each of which has different genetic causes and patterns of signs and symptoms. Type 1 CDA syndrome is characterized by moderate to severe anemia. CDA1 syndrome is caused by the mutation of the CDAN1 gene, which is based on the long arm of chromosome 15 as 15q15.2. The CDA2 syndrome is caused by the mutation of the SEC23B gene, which is based on the short arm of chromosome number 20, 20p11.23.

Generalized Congenital Dysrhythmia Anemia Syndrome (CDA)

CDA syndrome is a genetic disorder that affects the growth of red blood cells. This syndrome is one of the anemia disorders characterized by a lack of red blood cells. This deficiency prevents the transfer of oxygen from the blood to the tissues of the body. Symptoms include fatigue, weakness, dullness, and other complications [1] (Figure 1).

Clinical Signs of Congenital Dysrhythmia Anemia Syndrome (CDA)

Researchers have identified three types of CDA syndromes, each of which has different genetic causes and patterns of signs and symptoms. Type 1 CDA syndrome is characterized by moderate to severe anemia. This disorder is usually diagnosed in childhood or adolescence, although in some cases the syndrome may also be diagnosed before birth. Many people with CDA1 syndrome experience jaundice in the skin and the white bowel of the eye, and the size of the liver and spleen (hepatosplenomegaly). This disorder also causes excessive iron intake to the body, causing damage to the tissues and organs. In particular, excess iron can lead to abnormal heart rhythm (arrhythmia), congestive heart failure, diabetes, and chronic liver disease (cirrhosis). Rarely, people with CDA1 syndrome are born with skeletal disorders, which often include fingers or toes [2] (Figure 2).

Anemia associated with CDA2 syndrome varies from mild to severe, and most people with this disorder, jaundice, enlarged liver and spleen (hepatocellular membranes) and gallstones. CDA2 syndrome is usually diagnosed in adolescence or childhood. Abnormal iron deficiency usually occurs after the age of 20, leading to complications such as heart disease, diabetes and cirrhosis of the liver [3].

Symptoms and Symptoms of CDA3 Syndrome are different from the two previous ones. Adults with CDA3 syndrome have retinal disturbances that can cause visual impairment. Some people with CDA3 syndrome have a blood disorder called monoclonal gammopathy that can lead to white blood cell (multiple myeloma) cancer [4] (Figure 3).

Etiometry of Congenital Dysrhythmia Anemia syndrome (CDA)

CDA1 syndrome is caused by the mutation of the CDAN1 gene, which is based on the long arm of chromosome 15 as 15q15.2. The function of this gene is still not well understood, and it is unclear how the mutation in this gene leads to the signs and symptoms of CDA1 syndrome.

The CDA2 syndrome is caused by the mutation of the SEC23B gene, which is based on the short arm of chromosome number 20, 20p11.23. The gene provides instructions for protein synthesis that interfere with the transport of other proteins within the cells. During the growth of red blood cells, the protein produced from the SEC23B gene may help to ensure that proteins are transferred to the required sites. Researchers are trying to determine which mutations in the SEC23B gene lead to symptoms and symptoms of CDA2 syndrome [5] (Figure 4 and 5).

The genetic cause of CDA3 syndrome has not yet been determined. This syndrome is probably caused by a mutation in a gene in the long arm of chromosome 15 in the 15q22 region. The genetic changes responsible for the development of CDA syndrome cause disturbances in the normal growth of red blood cells, known as the erythropoiesis process. The term disperitopoietic is a condition in which red blood cells form abnormally. In people with CDA, abnormal red blood cells are called erythroblasts in an unusual form, and other disorders are like additional nuclei. These abnormal
CDA1 and CDA2 syndrome follow an autosomal recessive hereditary pattern. Therefore, in order to produce this syndrome, two copies of the mutant gene CDAN1 and SEC23B (one of the father and mother) are needed. The deficiency of healthy red blood cells leads to signs and symptoms of anemia, as well as hepatoplasmosis and abnormal iron elevation [6] (Figure 6).

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Frequency of congenital diarticular anemia syndrome (CDA)

CDA2 syndrome is the most common form of this disorder, and so far, more than 300 cases of CDA2 syndrome have been reported from medical literature throughout the world. A rare CDA3 syndrome is reported in only a few families in Sweden, Argentina and the United States. Frequency of CDA1 syndrome is unknown. Because CDA syndrome is very rare and its signs and symptoms indicate overlap with other disorders, so many cases are unlikely to be diagnosed or indirectly diagnosed as other disorders [9] (Figure 7).

Congenital Dysrhythropoietic Anemia Syndrome (CDA)

The CDA syndrome treatment and management strategy is symptomatic and supportive. Treatment may be done by a team of experts, including pediatricians, eye specialists, cardiologists, liver specialists, hematologists, dietitians and other healthcare professionals. There is no valid treatment for this syndrome, and all clinical measures are needed to reduce the suffering of the sufferers. Genetic counseling is also a special place for all parents who want a healthy baby [11] (Figure 9).

Discussion and Conclusion

CDA syndrome is a genetic disorder that affects the growth of red blood cells. This syndrome is one of the anemia disorders characterized by a lack of red blood cells. Researchers have identified three types of CDA syndromes, each of which has different genetic causes and patterns of signs and symptoms. Type I CDA syndrome is characterized by moderate to severe anemia. CDA1 syndrome is caused by the mutation of the CDAN1 gene, which is based on the long arm of chromosome 15 as 15q15.2. The CDA2 syndrome is caused by the mutation of the SEC23B gene, which is based on the short arm of chromosome number 20, 20p11.23. There is no valid treatment for this syndrome, and all clinical measures are needed to reduce the suffering of the sufferers.

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