Conjoined twins with a single heart with complex disease

Gêmeos unidos com um único coração com doença complexa

Gemelos unidos con un solo corazón con enfermedad compleja

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Abstract

Conjoined twins are a rare presentation with an estimated incidence of 1: 50000 pregnancies. They occur due to a random event, not related to heredity, maternal age or delivery conditions. This article is a case-report of a pair of female thoraco-omphalopagus conjoined twins, who were diagnosed during prenatal period through a morphological ultrasound. After a 24-week gestation period, the female newborns had a total weight of 1100g, and were born of a cesarean section. Both fetuses scored 3 in the Apgar scale in the first minute, getting down to a score of 1 in the fifth minute. Soon after birth, the newborns were referred to the Intensive Care Unit, where supportive measures were taken and transthoracic Doppler echocardiography was performed for diagnostic confirmation. The echocardiogram showed that there was only one heart for both newborns with a complex heart disease. In 75% of the cases of thoraco-omphalopagus conjoined twins, structural heart disfigurements are found in at least one of the fetuses. Besides, in approximately 80-90% of the cases, the cardiac abnormalities in the heart junction are far too complex, making it impossible to surgically separate the fetuses. This way, a structural and functional analysis of the heart is considerably more efficient post-birth, and it’s shown to be more accurately done through an echocardiography. This reveals that cardiac changes are a crucial factor in the fetal and postnatal evaluations of conjoined twins. Such procedures are a way of providing adequate medical recommendations, which may always be honest and based on scientific evidence.

Descriptors: Diseases in Twins; Twins, Conjoined; Heart Diseases.

Resumo

Os gêmeos unidos são uma apresentação rara, com incidência estimada em 1: 50000 gestações. Eles ocorrem devido a um evento aleatório, não relacionado à hereditariedade, idade materna ou condições de parto. Este artigo é um relato de caso de um par de gêmeos unidos toraco-omphalopágos do sexo feminino, diagnosticados durante o período pré-natal por meio de ultrassonografia morfológica. Após um período de 24 semanas de gestação, as recém-nascidas do sexo feminino tinham um peso total de 1100g e nasciram de uma cesariana. Ambos os fetos marcaram 3 na escala de Apgar no primeiro minuto, chegando a 1 no quinto minuto. Logo após o nascimento, os recém-nascidos foram encaminhados para a Unidade de Terapia Intensiva, onde foram tomadas medidas de apoio e a ecocardiografia transtorácica com Doppler foi realizada para confirmação diagnóstica. O ecocardiograma mostrou que havia apenas um coração para os dois recém-nascidos, com uma doença cardíaca complexa. Em 75% dos casos de gêmeos unidos toraco-omphalopágos, alterações cardíacas estruturais são encontradas em pelo menos um dos fetos. Além disso, em aproximadamente 80-90% dos casos, as anormalidades cardíacas na junção cardíaca são complexas, impossibilitando a separação cirúrgica dos fetos. Dessa forma, uma análise estrutural e funcional do coração é consideravelmente mais eficiente após o nascimento e é demonstrada com mais precisão por meio de uma ecocardiografia. Isso revela que as alterações cardíacas são um fator crucial nas avaliações fetais e pós-natais de gêmeos unidos. Tais procedimentos são uma maneira de fornecer recomendações médicas adequadas, por serem autênticas e baseadas em evidências científicas.

Descritores: Doenças em Gêmeos; Gêmeos Unidos; Cardiopatias.

INTRODUCTION

Conjoined twins are a rare presentation with an estimated incidence of 1: 50000 pregnancies, although the incidence of live births is 1: 25000 with female predominance (3: 1)¹².

They occur due to a random event, not related to heredity, maternal age or delivery conditions. The cases may be classified into eight different types, each one corresponding to the junction’s specific anatomical location³ and according to the terminology proposed by Spencer et al.⁴. This report refers to a case of single-heart thoraco-omphalopagus conjoined twins, who carried a complex heart disease which was incompatible with the fetuses’ lives.

CLINICAL CASE

Through a morphological ultrasound, a pair of female conjoined twins was diagnosed as the thoraco-omphalopagus type at their prenatal period. Besides the junction, this had already been identified.
as shown in Figure 1. The diagnosis indicated a severe congenital heart disease which was incompatible with the fetuses’ lives and, although they did not have other alterations, it was decided to interrupt the pregnancy.

The serologies collected during prenatal showed that the woman carrying the twins was free of toxoplasmosis, HIV, Hepatitis B and syphilis. At the time, the pregnant woman was 34 years old, had blood type O+ and no previous abortions or miscarriages. After a 24-week gestation period, the female newborns had a total weight of 1100g, and were born of a cesarean section, having the amniotic sac broken in the act. Both fetuses scored 3 in the Apgar scale in the first minute, getting down to a score of 1 in the fifth minute. The first examination showed that the fetuses were hypotonic, hypoactive, with a single umbilical stump and no bleeding. The doctors also identified agonal breathing which evolved into cyanosis and bradycardia. Through a pulmonary auscultation, vesicular murmurs were identified along with fine crepitations and rales. The doctors were also able to identify cardiac fusion, as well as other major flaws in the heart. Moreover, in the beginning of the second trimester of the pregnancy, it is possible to make a more detailed analysis of the heart’s anatomy and the extent of the cardiac fusion. Nonetheless, external cardiac details involving circulatory anastomosis are less likely to be identified before birth.

During the first three months of the pregnancy, the exact details of the heart’s interior anatomy are difficult to pinpoint. However, in this gestational phase, it is already possible to identify a cardiac fusion, as well as other major flaws in the heart. Moreover, in the beginning of the second trimester of the pregnancy, it is possible to make a more detailed analysis of the heart’s anatomy and the extent of the cardiac fusion. Nonetheless, external cardiac details involving circulatory anastomosis are less likely to be identified before birth.

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Figure 1: RX performed after the birth in intensive care unit.

Figure 2: Caption: Four-chamber echocardiogram demonstrating the presence of complex congenital heart disease.

DISCUSSION

According to the previously developed hypothesis on the division of joint germination, identical twins are formed when a single fertilized egg divides itself into two embryos - a phenomenon which occurs between the 13th and 15th day after fertilization. Thus, a flaw in this process may lead to the formation of conjoined twins. Other theories affirm that this type of formation may come as a result of a secondary fusion of single-ovary embryos that has been previously separated.

In 75% of the cases of thoraco-omphalopagus twins, structural heart disfigurements are found in at least one of the fetuses. Besides, in approximately 80-90% of the cases, the cardiac abnormalities in the heart junction are far too complex, making it impossible to surgically separate the fetuses. The twins’ cardiac anatomy may be categorized into four fundamental types: 1) Both heart and pericardium are separated, 2) Separated heart and single pericardium, 3) Separate ventricles and single atrium, 4) Single atrium and ventricle.

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also run a computed tomography which provides contrast for the analysis of the three dimensional planes - a method that is highly recommended for the examination of venous and arterial anatomy.

From the points taken into consideration so far, it is safe to assume that, in the case of any sort of ventricular fusion, surgical procedures become considerably more hazardous and should not be performed1. It is also important to note that some complications (e.g. a defect in one of the twins’ heart or a significant arterial perfusion that involves both twins) may be caught up in an extensive ethical debate regarding the surgical separation of the twins, considering that such flaws may lead to the death of one of the fetuses, as it has been the case brought up to the United Kingdom’s Supreme Court6.

This reveals that cardiac changes are a crucial factor in the fetal and postnatal evaluations of conjoined twins. Such procedures are a way of providing adequate medical recommendations, which may always be honest and based on scientific evidence6.

During pregnancy, it is quite usual that the physical and psychological alterations in the woman’s body lead to the building of great expectations about the child that she’s carrying, which often results in a sort of idealization of the “perfect child”. When this idealization is disrupted by the knowledge that said “perfect child” is, instead, a child that needs special care or a child that may not survive due to their special conditions, the baby’s parents may have to deal with the strong feelings of frustration, sadness and anguish that come along with the dismantle of the dreams and expectations they had been nurturing. If the child, for some reason, remains alive, the psychological impact of their birth may reverberate in the relationship that the parents are to build with the child - affecting, for example, the quality of the interaction between mother and baby, and in the relationship that other members of the family and other social groups may have with this infant9.

The presence of a child diagnosed with a disability or a congenital anomaly in a family implies a great concern over the several special care measurements that are given by the doctors along with the diagnosis because, for the child’s parents, who, in most cases, carry the greatest deal of responsibility in this process, there may be consequences regarding their quality of life and bio psychosocial well-being. As far as psychological aspects go, they may be overcome with feelings of depression, sadness, self-devaluation, and guilt as well as show symptoms of disturbance and post-traumatic stress10,11.

It is relevant to note, however, that such diagnosis might result in distinct reactions from each of the child’s parents. Because the connection between mother and child is inherently stronger, the impact of the birth and presence of a disabled child is often stronger in the mother’s perspective, which may also make her adaptation to this situation considerably more difficult. In the father’s perspective, the weight of the stigma and the social and financial impacts that the disabled child may bring to the family are usually the most critical factors that end up affecting his life10,11.

Taking these factors into consideration, it is evident that one must stay aware of the possible repercussions that such situation may bring to the physical, psychological and social aspects of the parents’ lives. Such repercussions must be dealt with in an interdisciplin ary and multidisciplinary way, in an attempt to provide the family with tools that may minimize the suffering that comes along with this process.

CONCLUSION

From the arguments that have been exposed in this article, it is possible to conclude that cardiac alterations are a crucial factor in the fetal and postnatal evaluation of conjoined twins. Such procedure enables the doctor to give adequate advice to the children’s family, and such advice should always be honest and based on scientific evidence in an interdisciplinary and multidisciplinary way, in order to provide solid, comprehensive care to all those involved in the process.

REFERENCES

7. Lopes LM, Brizot ML, Schultz R, Liao AW, Krebs VLJ, Francisco RPV et al. Twenty-five years of fetal echocardiography in conjoined...

CONFLICTS OF INTERESTS
The authors declare no conflicts of interests.

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