ACUTE LYMPHOID LEUKEMIA (LLA): A PEDIATRIC CASE REPORT IN DIZYGOTIC TWINS

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ABSTRACT

Acute leukemias are considered neoplasias that reach the hematopoietic system, resulting in rapid clonal expansion and accelerated proliferation of blood cells. In this way, the cells lose their capacity for maturation and differentiation, accumulating young cells (blasts) and without function in the bone marrow. Acute lymphoblastic leukemia (ALL) is a neoplasm that affects cells of the lymphoid lineage, but its prevalence occurs in B-cell precursors compared to T cells. It is common to occur in children between 1 and 5 years of age, comprising about 80% of the cases of all leukemias in this age group, but in adults, the probability of being present is lower. The present study aimed at the accomplishment of a case report on acute lymphoid leukemia in children, based on the comparison of risk factors in the family, and the evaluation of socioeconomic interference in the treatment, where, through an explanatory booklet, health professionals and the community about the major signs and symptoms of ALL in children. The research was submitted to the Ethics Committee of the Catholic University Center of Quixadá, through the Brazil Platform where it was analyzed and approved (01673218.2.0000.5046). In this study, we reported the case of two twin sisters who entered the Hospital São Sebastião de Pedra Branca, Ceará at different times, where one was admitted with fever and boulders, and another presenting only fever, after laboratory tests they were referred to a referral hospital in Fortaleza, Ceará, and were then diagnosed with acute lymphoid leukemia. Due to the poor prognosis of this pathology and a possible worsening of the patients’ clinical status, they were referred for bone marrow transplantation. Fortunately, after the treatments performed, the twin had a positive evolution, no longer needing to undergo bone marrow transplantation.

INTRODUCTION

Leukemia is a neoplasm that affects thousands of people worldwide, has characteristics and changes that specifically affect hematopoietic cells, to cause their rapid and unregulated development. In the initial phase of the disease, these cells will be present in the bone marrow, later on going to extra spinal areas (MOURA, 2014). Acute leukemias result from rapid clonal expansion and accelerated blood cell proliferation.

To lose the maturation and differentiation capacity, accumulating young cells (blasts) and without functions in the bone marrow. Blasts are first produced in the bone marrow, where they subsequently reach the peripheral blood and can enter hematopoietic organs such as the liver, spleen, and lymph nodes, and also reach non-hematopoietic organs such as the central nervous system, the skin, and the reproductive organs. The affected immature cells may come from the lymphoid or myeloid lines (KAVEH et al., 2017; CASADO et
al., 2016). Among the various forms of this pathology, there is acute lymphoid leukemia (ALL), which is characterized by being a neoplasm that affects cells of the lymphoid lineage, with prevalence in the precursors of B lymphocytes, in comparison to T lymphocytes. Occurrence in children, from 1 to 5 years old, comprising about 80% of all cases of leukemia in this age group. Children who develop ALL when diagnosed early have a 90% chance of total cure, demonstrating a good prognosis (NASCIMENTO et al., 2013; CRUZ, 2017). However, the big challenge is to make the diagnosis quickly and efficiently, because ALL symptoms are quite confused with other types of pathologies. Several factors contribute to the delay in diagnosis, among them are the lack of information and knowledge of parents, who often postpone the search for a diagnosis, as well as the inefficiency in the anamnesis process, the lack of trained professionals and the delay in service.

Public health system. These factors are extremely important, as the prognosis of these patients can be considered good, in the face of an early diagnosis, starting the treatment quickly and thus increasing the chances of cure (MATTOS, 2017). At the same time, the lifespan of children with ALL depends entirely on the efficiency and speed of diagnosis, as well as on the appropriate and specialized treatment for each type of patient, according to the ALL subtype and the stage of the disease. For this, information for a better understanding is necessary for a good understanding of the pathology, and thus help the most in the diagnosis, so that these children have more chances of cure and better quality of life. Although LLA is widely reported, information that helps health professionals is still lacking, especially in cities that do not offer specialist doctors and specialized exams at LLA (CRISTOVAM, 2014). In the meantime, the present study aimed to carry out a case report about ALL in children, based on the comparison of risk factors existing in the family, and the assessment of socioeconomic interference in the treatment.

METHODOLOGY

A research carried out was of the exploratory descriptive type with retrospective character and qualitative, transversal approach and source of the documentary base. A preceded institution went to the Municipal Hospital São Sebastião - HMSS located in the city of Pedra Branca, Ceará, Brazil, between September to December 2018. For the research, the following criteria were taken into account: having the ALL, patients are dizygotic twins (2 years old), the socioecological life style of the patient, the treatment, starting the treatment quickly and thus increasing the chances of cure (MINAYO, 2010), exercising two types of modalities: the instrumental one, because “it is developed to help in the knowledge or redefinition of the problem “and the collective, for having” the purpose of studying characteristics of a population”(GIL, 2008). The research was submitted to the Ethics Committee of the Centro Universitário Católica de Quixadá (UNICATÓLICA), with due approval by opinion 01673218.2.0000.5046. Respecting the precepts established by resolution 466/12 of the National Health Council, which regulates the guidelines and standards of research involving human beings.

CASE REPORT

Patient 1: M.A.A.M. patient, female, aged 1 year and 4 months, born in Pedra Branca - CE, coming from the city of Fortaleza - CE, was admitted to the Municipal Hospital São Sebastião, showing yellow spots on the nails of her hands, being immediately afterward diagnosed as nail fungus, the doctor then requested a complete blood count, the result of this examination showed all the vestments within the reference value, then the patient was medicated and discharged. Two months later, the patient was admitted to the São Sebastião Municipal Hospital again, showing purple spots on her body and constant fever, a blood count was requested, the biochemistry responsible for the examination, verified a large number of young lymphocytes, and the number of leucocytes elevated, after dialogue with the doctor and analysis of exams by the same, the patient was transferred to Hospital Albert Sabin, Fortaleza-CE. On December 7, 2016, the patient had bruises on the upper and lower limbs and fever, then a new blood count was requested, which presented bicicopenia (decrease of two blood series) in the case of the patient in question, her bicicopenia was anemia and thrombocytopenia: platelets (8,901/mm³), erythrocytes (2,078/mm³), hemoglobin (5.6g / dl) and hematocrit (19%) and leukocytosis (52,000/mm³), being then hospitalized on December 8, 2016. On December 9, 2016, a myelogram was performed that showed lymphoblastic B-cell leukemia, with 98% of bone marrow blasts, with a final diagnosis of high-risk Acute Lymphoblastic Leukemia. Chemotherapy treatment was started on December 9, 2016, with myelogram with 3% blasts, markedly hypocellular bone marrow, and DRM 2.36% (lumbar puncture: the absence of neoplastic cells). A new myelogram was then performed, which showed bone marrow apparently in cytological remission and less than 1% of the blast (lumbar puncture: the absence of neoplastic cells).

On January 3, 2017, he presented a positive culture for S. aureus, resistant to oxacillin and sensitive to vancomycin: blood cultures were performed on 02/01, 01/05, and 01/11 performed at the tip of the catheter and showed contamination by Candida albicans. He underwent antibiotic treatment, improving his condition, a new blood culture was performed on January 18, 2017, with a negative result. During hospitalization, she presented with febrile neutropenia, with abdominal distension and abdominal wall edema. On January 11, 2017, ultrasound was performed, obtaining a normal abdominal result, normal amylase and lipase. The last febrile peak was on January 13, 2017, with improvement in the condition the patient was discharged, this being on January 23, 2017, with a return review for January 26, 2017, to perform a
new blood count and biochemical tests. A summary of the therapy used can be seen in Figure 1.

Motor rehabilitation through physical therapy and Total Parental Nutrition (TPN) were also part of the treatment. Some surgical procedures were part of the treatment, such as the puncture of the central venous access in the right subclavian vein on December 21, 2016, which was removed on January 14, 2017.

Patient 2: M.C.A.M. patient, aged 1 year and 8 months, born in Pedra Branca - CE, from the city of Fortaleza - CE, was admitted to the Municipal Hospital São Sebastião with a fever, the doctor on duty aware of the conditions of his twin sister, the forwarded to Albert Sabin Hospital, where a blood count was performed on February 6, 2017, with the patient showing bruises on the upper and lower limbs, and fever, the blood count showed bicitopenia, with anemia and thrombocytopenia: platelets (341,900/mm³), red blood cells (3,394/mm³), hemoglobin (9.8g / dl), hematocrit (29.1%) and leukocytosis (28,000/mm³). A myelogram was performed with a 57% blast count, closing the diagnosis of low-risk L1 Acute Lymphoid Leukemia. She was then referred to the ICU where she stayed for 6 days, evolved with Tiflite (inflammation of the cecum) and pneumonia, being treated with antibiotic prophylaxis (ATB), meropenem and vancomycin evolving with stability. On March 3, 2017, an abdominal ultrasound was performed showing a large amount of intra-abdominal fluid, his TGO value being 120u / L and TGP 242u / L. These results show liver changes, accumulation of fluid in the intra-abdominal region, one of the side effects that chemotherapy can cause. Chemotherapy aims to cause the death of leukemic cells, but chemotherapy is not only specific for altered cells, but it also affects cells that renew quickly, as is the case of cells of the digestive system and skin (RUSCHEL, 2016). Patient 02 had a TGO result of 42.0u / L and TGP 242u / L. These results show liver changes, possibly originating from chemotherapeutic drugs that were metabolized in the liver, showing the overload that the liver suffered during treatment, plus a side effect, as hematotherapists have hepatotoxicity (MATTOS, 2017). The chemotherapy of patient 01, since it was more prolonged, there was a need to implant a venous catheter, constituted of silicone or polyurethane of single lumen, or multiple, classified as semi or implantable, used for the administration of the chemotherapy approached by Júnior et al., (2010). However, there are some complications in its implantation, handling, and maintenance of this equipment with the occurrence of frequent infections.

DISCUSSION

The reports described corresponding to the diagnostic criteria of the Brazilian Childhood Leukemia Treatment Group (GBTLI-99), which classifies the patient according to their bone marrow blast count, peripheral blood leukocyte count and the age at which the diagnosis was made. However, patient 02 presented a finding in the OM of 57% of blasts, her leukocyte count was 28,000 mm³, and her diagnosis was established at 1 year and 8 months of age, two months after her sister's diagnosis, but it fits in the low-risk group established by Rocha (2012). The treatment of the twins is different because even though they also have ALL, the sisters belong to different groups according to the Brazilian Childhood Leukemia Treatment Group (GBTLI-99), which separates patients into two groups: high risk and low risk if differentiating according to the criteria already mentioned, with this information the treatments of the patients are elaborated and which drugs will be used, the dose and days that should be administered as explana Casado (2016).

Patients 01 and 02 presented abdominal distension, due to the accumulation of fluid in the intra-abdominal region, one of the side effects that chemotherapy can cause. Chemotherapy aims to cause the death of leukemic cells, but chemotherapy is not only specific for altered cells, but it also affects cells that renew quickly, as is the case of cells of the digestive system and skin (RUSCHEL, 2016). Patient 02 had a TGO result of 42.0u / L and TGP 242u / L. These results show liver changes, possibly originating from chemotherapeutic drugs that were metabolized in the liver, showing the overload that the liver suffered during treatment, plus a side effect, as hematotherapists have hepatotoxicity (MATTOS, 2017). For Avila (2013) these results occur from the 3rd cycle of chemotherapy sessions, it is important to note that even with this change, the values of TGO and TGP can remain within the reference value, in most treatment. The chemotherapy of patient 01, since it was more prolonged, there was a need to implant a venous catheter, constituted of silicone or polyurethane of single lumen, or multiple, classified as semi or implantable, used for the administration of the chemotherapy approached by Júnior et al., (2010). However, there are some complications in its implantation, handling, and maintenance of this equipment with the occurrence of frequent infections.

Figure 1. flowchart summarizing the treatment protocol used on the patient 1

Figure 2. flowchart summarizing the treatment protocol used on the patient 2
caused by pathogenic microorganisms that can aggravate the general condition of patients. In the case of patient 01, there was no apparent sign of infection by the catheter, but she presented fever and chills, then a blood culture was performed at the tip of the catheter, which was positive for *Cândida* formota. According to Dansi (2017) fungi are responsible for 10-30% of all infections that occur in catheters. The fact that the patient is immunosuppressed due to chemotherapy makes her more susceptible to infections. During the treatment, both patients had the Hand-Foot-to-Mouth Syndrome, which is caused by the highly contagious Coxsackie virus, common in children from 1 to 5 years of age, causing fever, blisters on the hands, feet, and mouth, and maybe acquired from food contaminated with the virus, possibly due to the depletion of patients' immunity (CRISTOVAM, 2014). In terms of risk factors, such as alcoholism, smoking, contact with ionizing radiation, age of the mother, and family history of hematopoietic diseases, there is no relation of such factors on the appearance or aggravation of the pathology, in the patients of the study. However, socioeconomic factors can influence the application of effective treatment for patients with neoplasms who require intense pharmacological treatment, however, the sisters received assistance for the treatment of the city hall of the city in which they reside (MAIA, 2014; LAKN, 2017). Patient 01, had a low platelet count, this fact is manifested when blood cell production is disrupted by changes in OM affecting the morphology, quantity, and function of all cells in the hematopoietic system. Platelet concentrate administration was required. Both patients had the number of red blood cells well below the reference value, also resulting in a drop in hemoglobin, thus presenting normocytic and normochromic anemia, requiring the administration of red blood cell concentrates as stated by Campregher (2017).

ALL is easily confused with a bacterial or fungal infection, anemia, and other types of neoplasms. This shows the great importance of a differential diagnosis, which guarantees the most accurate conclusion. The first symptoms that arise, the doctor may be suspicious of any common infection in children, so it is necessary to request tests. The blood count will indicate the first necessary parameters, the high leukocyte count can indicate an infection and the low amount of hemoglobin indicate anemia as explained by Moreira (2018). Other parameters are analyzed, such as the presence of young cells, which have ALL. This can cause the doctor to rule out infections and anemias. Evidencing the importance of requesting more specific tests, such as myelogram, where an OM puncture is performed and the percentage of blasts is verified, this test is confirmatory for leukemia. But it is not the last test requested, it is essential to know what type of leukemia that patient has so that the treatment is as specific and effective as possible, thus performing a cytochemical examination, showing the type of blood cell that cancer has affected. Immunophenotyping identifies cell subtype (DANTAS, 2015).

**Conclusion**

Based on the results described above, it was possible to diagnose ALL. Due to the poor prognosis of this pathology and a possible worsening of the patients' clinical condition, they were referred for bone marrow transplantation. The assistance granted by the city hall in the city where the patients live was of great value for the treatment, since, even though the treatment was provided by the Brazilian Health System, travel costs were incurred since the city of the patients did not undergo pharmacotherapy proper. After the treatments performed, the twins’ condition evolved positively, and it is no longer necessary to undergo bone marrow transplantation. The presentation and explanation of the case allowed highlighting the clinical and laboratory criteria, and the therapeutic alternatives recommended for patients with Acute Lymphoid Leukemia, demonstrating the importance of research in this category for the enrichment of science. In this way, the folder will serve as an information tool for health professionals and the community, assisting in the knowledge and perception of the main signs and symptoms of ALL, as well as contributing to the stages of diagnosis and prognosis.

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