Late pulmonary tomography assessment in premature infants with bronchopulmonary dysplasia submitted to patent ductus arteriosus management

Avaliação tomográfica pulmonar tardia em prematuros com displasia broncopulmonar e persistência de canal arterial

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Abstract

Objective: To assess through high-resolution computed tomography, the pulmonary parenchyma of children born prematurely with both very low birth weight and patent ductus arteriosus, undergoing medical or surgical treatment who developed bronchopulmonary dysplasia.

Methods: Between December 2006 and January 2007, 14 children born prematurely with a birth weight of less than 1500 g with bronchopulmonary dysplasia (BPD) and patent ductus arteriosus (PDA) were submitted to high-resolution computed tomography (HRCT). Patients were divided into two groups (Group A = medical [n=6] and Group B = surgical [n=8]) and all of them underwent surgical closure of the ductus arteriosus. The pool of patients was comprised of 9 baby boys and 5 baby girls who were 36.5±4.3 month-old. The HRCT were analyzed by two independent observers and quantified in each patient. The statistical analyses were assessed using the Mann-Whitney test and p<0.05 was considered statistically significant.

Results: Three patients presented normal tomography; two of group A and one of B. In Group A, the most frequent finding was multifocal ground-glass opacity. In Group B, multifocal ground-glass opacity, atelectasis, and low attenuation areas with relatively decreased number and caliber of vessels were prevalent (62.5%). There was a statistically significant difference between both groups. Group B presented higher averages in intubation time, use of oxygen, and admission. However, as to the number of injuries found on HRCT there was no statistically significant difference (p=0.0787).

Conclusion: The lately use of HRCT has shown no significant difference between both medical and surgical treatment aiming at to occlude the PDA in pulmonary parenchyma injuries of premature infants with PDA who developed bronchopulmonary dysplasia.

INTRODUCTION

Bronchopulmonary dysplasia is a multifactorial disease, whose etiology is not completely established yet. It results from the synergism of aggression factors to the immature pulmonary tissue, such as mechanical ventilation, oxygen therapy, infection, PDA (patent ductus arteriosus), prematurity, nutrition, and genetics. Bronchopulmonary dysplasia develops from an acute lesion in an immature lung, resulting in an interruption of the normal process of the lung development with impairment of vascular and alveolar growth followed by the abnormal repair process, with the development of chronic pulmonary disease [1].

The basic evaluation method is still the thorax radiography. However, it presents low sensitivity for some lesions when compared to the high-resolution computed tomography (HRCT), which can show subtle abnormalities, such as septal lines, honeycombing, parenchymal bands, and ground-glass opacity, among others [2,3]. Ground-glass opacity is the increased lung attenuation in which it is still possible to identify the contours of the vessels and bronchial structures within the pulmonary area affected by a pathological process that causes a reduction in the aeration of the air spaces [4].

In up to 80% of the times, extremely low birth weight (ELBW) premature infants, i.e., a birth weight less than 1500 g, can present a patent ductus arteriosus (PDA), with ELBW being inversely proportional to gestational age and birth weight at birth [5]. Such children often require treatment for closure of the patent ductus arteriosus, once this leads to an increased pulmonary blood flow and to an interstitial edema with reduced pulmonary compliance besides an increased airway resistance may lead to both higher mechanical ventilation time and risk of bronchopulmonary dysplasia (BPD) [6].

Procedure for closure of the patent ductus arteriosus can either be clinically performed with a prostaglandin inhibitor which proved to be ineffective in up to 40% of patients, or surgically performed by direct PDA ligation. Such approaches are individualized and, currently, there is no consensus regarding the superiority over one another. This fact may have a direct influence on hospitalization time and in the child prognosis as to pulmonary lesions [7].

In this way, we seek to evaluate lately through high-resolution computed tomography the lung parenchyma of the extremely low birth weight (ELBW) prematures infants with PDA undergoing surgical or pharmacologic treatment aiming at PDA closure and who presented with bronchopulmonary dysplasia.

METHODS

Between December 2006 and January 2007, 14 children, mean age of 36.5±4.3 month-old, underwent high-resolution computed tomography to study the lung parenchyma. Nine (64.3%) patients were males and five females (35.7%). Mean birth weight was 939.6±288.4 g.

The study included premature infants with gestational age less than 29 weeks and with a very low birth weight (less than 1500 g), who developed bronchopulmonary dysplasia and required PDA closure treatment for over a year. Patients were divided into two groups: Clinical group or group A (n=6) included patients treated with prostaglandin inhibitors only; Surgical group or group B...
(n=8) included patients with prostaglandin inhibitors failure and required referral to surgical ligation of the ductus or were treated with surgical ligation only.

In group A, median birth weight was 1005 g, ranging from 815 to 1400 g; in group B, median birth weight was 840 g, ranging from 480 to 1380 g.

In the present retrospective study, clinical data were obtained through medical records analysis, gender, age, birth weight, durations of mechanical ventilation (MV), use of oxygen, and admission only.

Table 1 shows intubation time, duration of oxygen dependence, and days of hospitalization. It can be observed that the mean and median values are close to each other, thus disclosing the low influence of clinical cases with atypical and disagreeing values.

The number of lesions found on HRCT was quantified in each patient, i.e., for each patient was added the number of lesions regardless the degree found.

Mann-Whitney test was used to statistically analyze the number of lesions found on HRCT. A p d’ 0.05 was considered statistically significant.

The study was approved by the local Institutional Review Board and Ethics Committee and a written postinformed consent was obtained from the child’s legal representative, under the protocol number 3727/2006.

RESULTS

In the whole group, the median lesion per patient found on HRCT was seven lesions. Three patients presented normal CTs, two in Group A and one in Group B.

In Group A, the most frequent finding was multifocal ground-glass affecting three patients. Atelectases were not found. Figure 1 shows the tomographic findings in this group. In Group B, multifocal ground-glass lesions, atelectasis and areas of low attenuation and reduction in vessel caliber were preponderant in five patients (62.5%); two (25%) presented parenchymal bands, multifocal subsegmental consolidation and focal ground-glass, and one (12.5%) presented bullae.

Figure 2 shows the tomographic findings of Group B in patients who evolved with pharmacological treatment failure and later underwent surgical treatment, and in those who underwent surgical treatment alone.

Hospital dispensary exams were performed at the clínica radiológica Ultra-X Diagnóstico por Imagem de São José do Rio Preto, using a GE LightSpeed (TM) multi-slice CT system (General Electric Medical Systems, Milwaukee, WI), allowing physicians to view 16 images per second with slice intervals of 0.65 cm. All the patients were sedated with propofol (2.0 mg/kg) administered in a peripheral vein and monitored by pulse oxymetry and noninvasive mean arterial pressure, and followed-up during the CT scan by a multidisciplinary staff composed of an anesthesiologist, a pediatrician, and a nurse. There was not observed a reduced respiratory depression and saturation, consequently orotracheal intubation was not required. Sedation was spontaneously reversed after withholding drug therapy. Patients were discharged from hospital after 6 hours.

Pulmonary lesions seen on HRCT were described according to the Brazilian Consensus on Terminology Used to Describe Computed Tomography of the Chest, observing areas of low attenuation and reduction in vessel caliber, atelectasis, parenchymal bands, bullae, multifocal subsegmental consolidation, random small nodules, focal ground-glass, and multifocal ground-glass [8].

Pulmonary lesions analyses were independently performed by two experienced thoracic radiologists and the disagreeing cases were resolved through a posteriori consensus.
Comparing statistical outcomes through the application of Mann-Whitney test, patients who were pharmacologically treated only (Group A) vs those who in some time underwent surgical treatment (Group B), we get subsides to evaluate evidences related to the statistical significance of the difference between the medians obtained for the following variables: intubation time, duration of oxygen dependence, and hospitalization, which seemed to be statistically significant, as shown in Table 2.

These findings indicated that patients undergoing surgical closure of the ductus arteriosus (Group B) had the following higher averages scores: intubation time, duration of oxygen dependence, and hospitalization. In the dispersion plot (Figure 6), correlating the number of lesions and the type of treatment employed between surgical (Group B) and pharmacological (Group B) patients, it is observed a higher number of lesions in those who underwent surgery. There was statistical difference ($p=0.0787$).
DISCUSSION

With the advance of neonatal assistance, survival, PDA and BPD have significantly increased in premature infants. In those with an extremely low birth weight, the outcomes were highly adverse, influencing the cognitive development and the pulmonary function [9].

Several factors can affect the immature lung and develop BPD, such as intubation time, oxygen exposition time, infection, and presence of PDA [10]. As we can observe in Table 1, the mean intubation time was 34±18.4 days and the mean oxygen dependence was 77±24.5 days, which supports the pulmonary insult severity the premature patients suffered in the neonatal period.

The closure of patent ductus arteriosus with hemodynamic repercussion in premature infants should be performed as earlier as possible, whether by pharmacological or surgical intervention in order to try reducing morbidities. Although, a multicenter study does not indicate which of the two therapies should be the first choice in newborn infants [10]. Méier et al. [11] showed that premature newborns with respiratory distress syndrome, who have been administered prostaglandin when underwent surgery as a result of this treatment failure, have the worst outcome in comparison to those who underwent surgery as a first therapy choice.

Pharmacological treatment is undertaken administering oral inhibitors of prostaglandin synthesis, among which are indomethacin, and more recently ibuprofen, besides supportive measures, such as fluid intake restriction and diuretics use [12,13].

Indomethacin is the drug of choice in the majority of neonatology centers and it was administered to our patients. The constrictor therapy effect with indomethacin combined with fluid intake restriction and therapy with diuretics can explain the development of intestinal ischemia and organ dysfunction [7]. Other implications observed are trend to bleeding due to platelet dysfunction, necrotizing enterocolitis (NEC), renal function impairment with diminished urine output, resulting in overhydration with reduced pulmonary compliance and increased ventilation support time and transient heart dysfunction [5,12,14]. Such a fact justified the referral of three children directly to surgical treatment because one already presented intracranial hemorrhage and the other two increased serum creatinine.

In a study carried out with 21 premature infants with PDA, it was observed the closure of patent ductus arteriosus using indomethacin in 19 (90.4%) patients and the re-opening in 6 (31.5%) [14]. The data are not compatible with our sample, once 11 (78.6%) patients who underwent indomethacin treatment, five (45.4%) presented PDA re-opening, what can be explained by PDA association with infection.

In another study comparing premature infants with or without infection, the authors observed a significantly higher failure rate with indomethacin – 68% in infected children and 17% in noninfected ones – regardless birth weight, age, the route used to administer indomethacin (orally or intravenously), and the dose administered [9].

The re-opening rate of ductus arteriosus with pharmacological treatment is high in premature infants with less than 1000 g and lower in those over 1500 g. The success in closure of the patent ductus arteriosus is rather related...
to both birth weight and gestational age than to the indomethacin plasma concentration peak [13,16]. The median birth weight in both groups, surgical and pharmacological, were 840 g and 1055 g, respectively. This confirms that premature infants with less than 1000 g are more likely to have pharmacological treatment failure.

Surgical ligation is usually reserved either for PDA refractory to pharmacological treatment or for the impossibility to perform this treatment. Due to its low morbidity rate, it has been proposed as a primary treatment for PDA in some centers by its higher efficacy and lower morbimortality [14]. However, this remains a controversial procedure, and a more recent study has reported that PDA ligation is largely performed in premature infants, despite the clear evidence of a better outcome regarding pharmacological treatment [13].

In a randomized study comparing surgical vs pharmacological treatment with indomethacin in premature infants with less than 1750 g, there was no significant difference in days of hospitalization, lung lesions, necrotizing enterocolitis, and intracranial hemorrhage [13]. Although, our study was not carried out in a similar fashion, we did not find significant differences as for the number of lung lesions between both treatments; however, in the surgical group there has been found differences in intubation, oxygen dependence, and hospitalization mean times. This can be explained because previous unsuccessful treatment with indomethacin could have delayed surgical intervention and influenced the morbidities. Furthermore, pharmacological treatment was contraindicated to the patient who underwent direct surgical treatment, what could have selected the more critically patients to this group.

HRCT allows a yet more detailed evaluation of the lung parenchyma, by analyzing the distribution of morphofunctional changes throughout the bronchi, lymphatic routes, and inside the secondary lobule [16]. Its use is still limited, despite both sensitivity and specificity in comparison to thoracic radiography, because the radiation dose is higher and its cost is expensive.

A study conducted with 62 extremely low birth weight (ELBW) premature infants, close to their discharge from hospital, showed that most of infants presented more than one alteration on HRCT, described as ground-glass opacity, parenchymal bands, atelectasis, and bullae [17], which are very close to what we have found in our study, once the areas of low attenuation, parenchymal bands, multifocal segmental consolidation, and focal and multifocal ground-glass were present in children treated both surgically and pharmacologically.

Atelectasis, a complication sometimes resulting from surgical treatment drew our attention, even though not statistically significant, because these complications were found in patients who underwent surgical treatment only, suggesting that maybe the surgical aggression could provoke this late consequence in some patients, and that future studies need to be undertaken addressing this significant concern.

On the other hand, of the three patients who presented normal HRCT, two were treated pharmacologically and one surgically only, suggesting that regardless the therapy chosen to prevent late pulmonary outcomes, the bottom line is that early closure of the ductus arteriosus may be achieved pharmacologically using or by surgery.

It is important to emphasize that the goal of our study was to verify lung lesions and correlate them with the treatment for PDA closure. Evaluation of respiratory sign and symptoms was not included in our inclusion criteria, so that there was not any correlation between radiological and clinical image. However, it was naturally observed that the children surgically treated, in consequence of their critically severe health conditions and because most often they have undergone pharmacological treatment previously, presented much more lung infections and, consequently, much more frequent hospitalizations. This fact has drawn our attention and may be eventually addressed in another study.

CONCLUSION

Lately performed high-resolution computed tomography has shown no significant difference between either pharmacological or surgical treatment aiming at the closure of the ductus arteriosus in parenchymal lesions in premature infants with PDA who evolved to bronchopulmonary dysplasia.

REFERENCES


