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# Comparative Analysis of Myocardial Protection with HTK Solution and Hypothermic Hyperkalemic Blood Solution in the Correction of Acyanogenic Congenital Cardiopathies – A Randomized Study

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## Abstract

**Objective:** The goal of the present study was to compare the myocardial protection obtained with histidine-tryptophan-ketoglutarate (HTK) cardioplegic solution (Custodiol®) and with intermittent hypothermic blood solution.

**Methods:** Two homogenous groups of 25 children with acyanotic congenital heart disease who underwent total correction with mean aortic clamping time of 60 minutes were evaluated in this randomized study. Troponin and creatine kinase-MB curves, vasoactive-inotropic score, and left ventricular function were obtained by echocardiogram in each group. The values were correlated and presented through graphs and tables after adequate statistical treatment.

**Results:** It was observed that values of all the studied variables varied over time, but there was no difference between the groups.

**Conclusion:** We conclude that in patients with acyanotic congenital cardiopathies submitted to total surgical correction, mean aortic clamping time around one hour, and cardiopulmonary bypass with moderate hypothermia, the HTK crystalloid cardioplegic solution offers the same myocardial protection as the cold-blood hyperkalemic cardioplegic solution analyzed, according to the variables considered in our study model.

**Keywords:** Congenital Heart Disease, CHD. Cardiopulmonary Bypass, CPB (Incl Set-Ups, Equipment, Surface Coatings, Etc.), Pediatric. Myocardial Protection/Cardioplegia. Myocardial Injury (Incl Blunt, Penetrating, Iatrogenic).

## Abbreviations, acronyms & symbols

ANOVA	= Analysis of variance	IPO	= Immediate postoperative
ASD	= Atrial septal defect	PD	= Peritoneal dialysis
AVC	= Atrioventricular canal	PDA	= Patent ductus arteriosus
CAVC	= Complete atrioventricular canal	1PO	= First postoperative
CK-MB	= Creatine kinase-MB	16 PO	= 16 <sup>th</sup> postoperative
CPB	= Cardiopulmonary bypass	ReBEC	= Brazilian Registry of Clinical Trials
d-TGA	= dextro-transposition of the great arteries	RS	= Range sampling
DORV	= Double outlet right ventricle	SD	= Standard deviation
EF	= Ejection fraction	SIRS	= Systemic inflammatory response syndrome
HHB	= Hypothermic hyperkalemic blood	TTE	= Transthoracic echocardiography
HTK	= Histidine-tryptophan-ketoglutarate	VIS	= Vasoactive-inotropic score
ICU	= Intensive care unit	VSD	= Ventricular septal defect

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## INTRODUCTION

The development of the cardiac surgery is directly related to the evolution of myocardial protection techniques, which have made possible complex and long surgeries with adequate preservation of the cardiac function. The already known benefits of hypothermia in reducing the myocardial metabolic demand were associated with chemical cardioplegia, which is characterized by the administration of solutions in coronary arteries that can cause myocardial arrest, preserving the reserves of high-energy phosphate, reducing oxygen consumption, and offering oxygen and other substrates during aortic clamping. Initially proposed by Melrose<sup>[1]</sup>, in 1955, with a solution of 2.5% potassium citrate diluted in blood, this method was then abandoned because of the demonstration of myocardial injury caused by high concentrations of potassium<sup>[2]</sup>. However, it showed great improvement and acceptance in the 1970s, when researchers from Europe<sup>[3,4]</sup> and North America<sup>[5,6]</sup> reintroduced the cardioplegic protection by infusing cold crystalloid solutions containing magnesium chloride and potassium chloride at lower concentrations than those from Melrose solution. In Brazil, Brille was one of the main responsables for the implementation and development of chemical cardioplegia<sup>[7]</sup>.

The myocardial protection techniques remain a reason for great discussions. Questions about the best route of administration (anterograde, retrograde, or both together), ideal temperature (hypothermic, normothermic, or heated), frequency of administration (intermittent or continuous), and its various formulations diluted in blood or crystalloid solutions generate recurrent debates. Solutions diluted in blood are the preference of many surgeons because they have a more physiological composition, with natural buffers and oxygen-carrying capacity. However, crystalloid solutions remain the choice of some groups. One of the most used formulations is that developed by Bretschneider, in 1970, also known as histidine-tryptophan-ketoglutarate (HTK) solution (Custodiol®). This solution is widely used for organ preservation in transplants and, in general, it requires a single administration<sup>[8,9]</sup>.

Initially idealized for cardiac surgeries in adults, cardioplegic solutions were soon adapted for pediatric cardiac surgeries<sup>[10]</sup>. However, the child's heart behaves differently from the adult's heart in relation to ischemia, since there are structural, functional, and metabolic differences, being the child's heart more vulnerable to myocardial injury. This is due not only to the associated malformations, but also to the organ immaturity<sup>[11-13]</sup>. In the Pediatric Heart Surgery Service, at the Hospital do Coração Dr. Carlos Alberto Studart Gomes (Fortaleza, Ceará, Brazil), about 450 pediatric cardiac surgeries are performed per year, most of them using cardiopulmonary bypass (CPB) with aortic clamping and administration of cardioplegic solution. The authors are responsible for approximately half of these surgeries. Initially, a hypothermic blood solution formulation with potassium addition was administered predominantly to the aortic root every 30 minutes. With the advent of the HTK crystalloid hypothermic solution in heart transplants in our hospital, some groups extended its use, due to the practicality of a single administration, to other cardiac surgeries in adults and children.

The objective of this study was to compare the efficacy of

two cardioplegic solutions used for myocardial protection in two homogenous groups of children submitted to surgical correction of congenital heart diseases: HTK solution and hypothermic hyperkalemic blood (HHB) solution.

## METHODS

This is a longitudinal, prospective, clinical, randomized, and double-blind trial. Fifty children consecutively operated on from November 2012 to June 2014 were evaluated by the same surgical team. All of them were carriers of acyanotic congenital cardiopathies, which were corrected with CPB and aortic clamping. Surgeries for correction of atrial septal defect (ASD) exclusively, reoperations, and palliative and emergency surgeries were excluded. The children were randomized into two groups of 25 patients: in Group A, patients received the cold HTK solution (about 5°C), administered as a single dose in the aortic root, for six minutes, with a minimum infusion of 30 mL/kg. In Group B, patients received the HHB solution at a concentration of 25 mEq/L, administered at the root of the aorta with a syringe, at a temperature of 20°C (priming temperature), every 30 minutes, in a volume of 20 mL/kg in induction and 10 mL/kg in maintenance. The 50 numbers were randomized by the Epi Info software version 3.5.1 and placed in separate and sealed envelopes. These envelopes were removed and opened randomly and immediately prior to the application of the solution. In defining the sample size, we considered the number of patients with this profile operated by the specific team during the year.

Researchers who performed the data collection in the medical record and specific protocol did not know which type of solution was used; only the surgeon in charge had access to this information (he knew the randomization seal).

The parameters evaluated in this study were: age (months), weight (kilogram), gender, type of cardiopathy, duration of surgery, CPB, and aortic clamping time (minutes); for each of these variables the mean was calculated and, later, a correlation between the two groups was performed. The evaluation of myocardial damage was made through three variables: dosage of cardiac enzymes, troponin, and creatine kinase-MB (CK-MB). Enzymes were collected preoperatively at the end of surgery and six, 12, 18, 24, and 48 hours after surgery. The type and number of vasoactive drugs used every two hours were analyzed in the first 36 hours after surgery. From these vasoactive drug data, the vasoactive-inotropic score (VIS) was calculated using the formula presented in Table 1<sup>[14]</sup>.

**Table 1.** Vasoactive-inotropic score (VIS) formula.

<b>VIS = (dopamine [mcg/kg/min]) + (dobutamine [mcg/kg/min])</b>
+ (10 x milrinone [mcg/kg/min])
+ (100 x adrenaline [mcg/kg/min])
+ (100 x norepinephrine [mcg/kg/min])

In addition, children underwent transthoracic echocardiography (TTE) in the preoperative period, immediate postoperative period, and between 24-48 hours after surgery, in order to determine the left ventricular function by the percentage of shortening ( $\Delta D$ ). Data were collected from surgical reports and a specific protocol created for the study. In the period of the study, the solutions' efficacy in myocardial protection was compared through the correlation of the estimated marginal means of the previously mentioned parameters, arranged in charts and table.

Categorical quantitative results were presented as percentages and counts, and the numerical results were presented as measures of central tendency. Kolmogorov-Smirnov normality tests were performed for numerical variables. For non-serial measurements, the chi-square test was used for categorical data and the analysis of variance (ANOVA) was used for numerical data. General linear models were used with repeated measurements of the mixed ANOVA type to evaluate serial measurements. Comparisons with *P*-value up to 0.05 were considered significant. Data were tabulated and analyzed by SPSS software (v. 23, SPSS, Inc.), for analysis and evaluation of data obtained in the collection.

The study followed the ethical principles of research involving human beings in the Resolution 466/12 of the National Health

Council, respecting the fundamental principles of autonomy, beneficence, non-maleficence, justice, and equity. It was approved by the Ethics and Research Committee of the Hospital do Coração Dr. Carlos Alberto Studart Gomes, protocol number 892/12. And it was registered in the Brazilian Registry of Clinical Trials (ReBEC), Registration Number RBR-9wqbx9.

## RESULTS

The majority of the patients were female (62%) and had ventricular septal defect (VSD) as the main heart disease (78%), associated or not with other lesions. A total of six (12%) patients required cardiac defibrillation/cardioversion after CPB, one (2%) required peritoneal dialysis (PD), and one (2%) died. There was no statistical difference between the groups for any of these variables (Table 2).

Table 3 shows the range sampling (RS) distribution, means, and standard deviation (SD) of age, weight and time of CPB, and surgery and aortic clamping according to the groups. There was no statistical difference between the groups for any of these variables (Table 3).

Regarding the behavior of the troponin, CK-MB, VIS, and TTE variables, all of them varied significantly in time (all  $P1 < 0.001$ ),

**Table 2.** Distribution of categorical variables according to the groups.

Variables	Solution – N (%)		Total N (%)	P
	A	B		
Sex				
Female	16 (64)	15 (60)	31 (62)	0.771
Male	9 (36)	10 (40)	19 (38)	
Main cardiopathy				
VSD	18 (72)	22 (88)	40 (78)	0.157
CAVC	5 (20)	2 (8)	7 (14)	0.221
Partial AVC	-	1 (4)	1 (2)	0.312
DORV with infundibular and valvular pulmonary stenosis	1 (4)	-	1 (2)	0.312
Subvalvar aortic stenosis	1 (4)	1 (4)	2 (4)	0.952
"Pink tetralogy"	2 (8)	-	1 (2)	0.132
Cardioversion				
No	21 (84)	23 (92)	44 (88)	0.384
Yes	4 (16)	2 (8)	6 (12)	
PD				
No	25 (100)	24 (96)	49 (98)	0.312
Yes	-	1 (4.0)	1 (2)	
Death				
No	25 (100)	24 (96)	49 (98)	0.312
Yes	-	1 (4)	1 (2)	

AVC=atrioventricular canal; CAVC=complete atrioventricular canal; DORV=double outlet right ventricle; PD=peritoneal dialysis; VSD=ventricular septal defect

**Table 3.** Distribution of numerical variables according to the groups.

Variables	Solution A		Solution B		Total		P
	RS	Mean (SD)	RS	Mean (SD)	RS	Mean (SD)	
Age (days)	3-168	28.5 (41.65)	2-108	18.6 (25.61)	2-168	23.5 (34.58)	0.501
Weight (kg)	4-25	11.2 (11.51)	3-25	8.4 (5.47)	3-59	9.8 (9.03)	0.632
CPB time (minutes)	40-125	80.3 (22.25)	42-135	79.8 (22.49)	40-135	80.0 (22.14)	0.816
Surgery time (minutes)	110-270	170.7 (36.76)	75-212	164.0 (31.77)	75-275	167.3 (33.90)	0.662
Clamping time (minutes)	26-90	57.8 (20.46)	20-90	54.2 (17.90)	20-98	56.1 (19.12)	0.640

RS=range sampling; CPB=cardiopulmonary bypass; SD=standard deviation

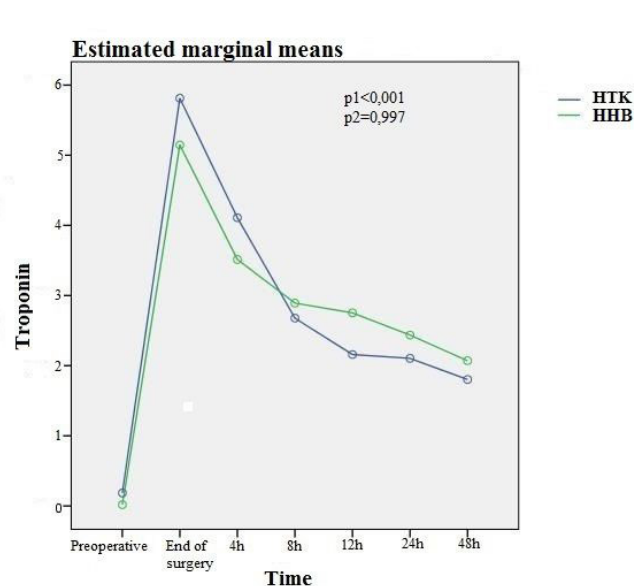
but with no difference between the solutions (P2), as shown in the graphs of the estimated marginal means (Figures 1 to 4).

Table 4 presents the marginal means of troponin, CK-MB, VIS, and TTE between the groups at the different moments analyzed. For all variables, there was significant variation in time, but there was no interaction of time with the type of solution.

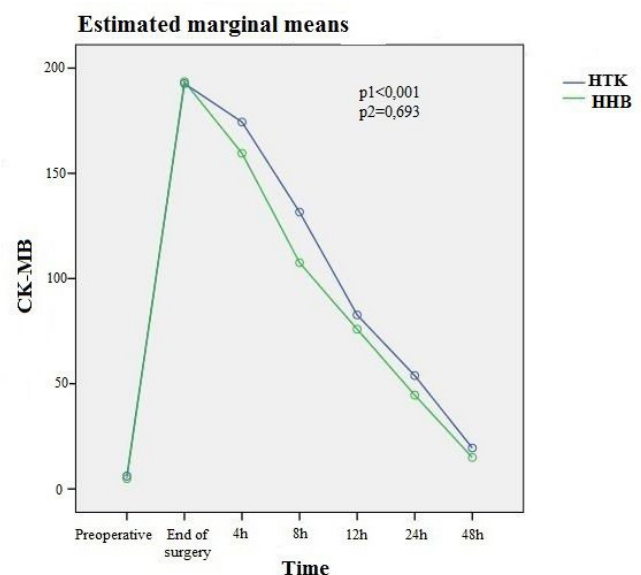
## DISCUSSION

The importance of adequate myocardial protection for a good surgical result, especially in pediatric cardiac surgery, due to the greater susceptibility of the child's heart to ischemic damage, leads to a continuous search for the set of actions that results in a better preservation of the organ during the period of ischemia necessary for the correction of most congenital heart diseases. The type of coronary solution to be administered represents an important factor in this process. Previous studies have demonstrated the benefits of blood cardioplegic solutions

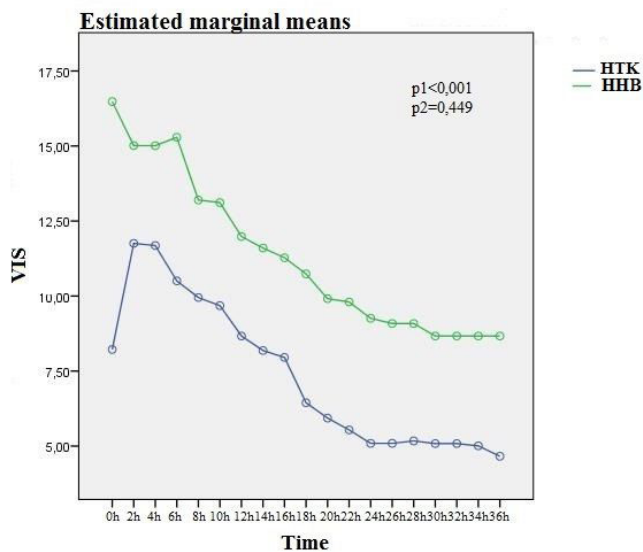
in relation to cold crystalloid solutions, defined by better clinical evolution and lower postoperative troponin values<sup>[15-17]</sup>. All of this makes some sense, considering the expected more physiological properties of blood in relation to crystalloid solutions. However, several surgeons continue to use crystalloid solutions with good results. In our hospital, we have been using the St. Thomas' crystalloid solution for a long time in pediatric cardiac surgeries and heart transplants. Subsequently, we began to use, in pediatric surgeries, the intermittent HHB solution studied here, with good results. Finally, the HTK solution initially used for heart transplants have been standardized as the cardioplegic solution for pediatric congenital heart surgeries by pediatric surgeons of the transplant group. However, despite the apparently good results, the question remained whether this new solution would yield results better than or equal to the ones from the previously used solution, although this new one is more practical. Thus, the purpose of this study arose.



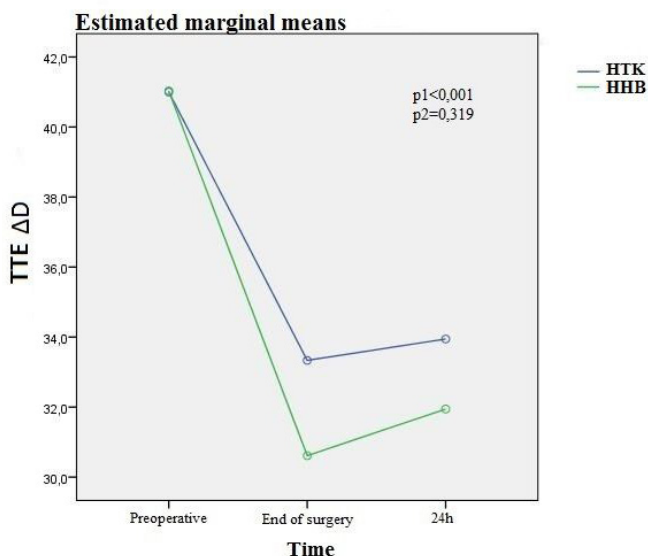
**Fig. 1** – Variation of troponin in time (P1) and between groups (P2). HHB=hypothermic hyperkalemic blood; HTK=histidine-tryptophan-ketoglutarate



**Fig. 2** – Variation of creatine kinase-MB (CK-MB) in time (P1) and between groups (P2). HHB=hypothermic hyperkalemic blood; HTK=histidine-tryptophan-ketoglutarate



**Fig. 3** – Variation of vasoactive-inotropic score (VIS) in time (P1) and between groups (P2).  
HHB=hypothermic hyperkalemic blood; HTK=histidine-tryptophan-ketoglutarate



**Fig. 4** – Variation of transthoracic echocardiography (TTE) in time (P1) and between groups (P2).  
HHB=hypothermic hyperkalemic blood; HTK=histidine-tryptophan-ketoglutarate

Regarding the sample choice, we are aware of the aggressions that the cardiopathy itself can cause to the myocardium. Cyanotic heart diseases, for example, may exert a bias, since different times of chronic hypoxia may characterize hearts more or less susceptible to myocardial damage, trans and postoperatively. Aiming to minimize these biases, we sought a more homogeneous study population and more predictable myocardial behavior in relation to surgical damage. Therefore, we

selected only acyanotic congenital cardiopathies and elective procedures. These surgeries also present a more uniform course and, consequently, more similar surgical times. For the same reason, reoperations were excluded. Other studies with other subgroups of patients may be performed.

The studied group was, in fact, homogeneous, presenting similar distribution of sex, age, and weight. Duration of surgery, echocardiography, and aortic clamping time also did not present a statistically significant difference between the groups. On the latter, the general averages were  $57.8 \pm 20.46$  and  $54.2 \pm 17.90$  minutes for Groups A and B, respectively. Thus, we will assume that the results found in the present study may refer to surgeries in acyanotic cardiopathies with mean aortic clamping time around one hour.

Analyzing the results, the cardiac enzyme curves presented similar behavior among themselves. Both variables varied significantly over time, i.e., they increased significantly in relation to the preoperative values, but without difference between the groups, as shown in Figures 1 and 2 and in Table 4. Regarding troponin, specifically, there was an upper initial elevation in the group that received the HTK solution, although with no statistical difference. We may question whether it would not become significant if the "n" were larger, characterizing a trend. In the CK-MB curve, however, these values were already very close. It is also known that it is the magnitude of the CK-MB values that correlates with the extent of the infarction, not the troponin values.

VIS is a predictor of the short-term evolution of children undergoing cardiac surgery. Davidson et al.<sup>[19]</sup> concluded that a high VIS at 48 hours after pediatric cardiovascular surgery is related to prolonged mechanical ventilation and intensive care unit (ICU) stay. In our study, with respect to VIS curves, these also varied significantly over time, but with no interaction between time and type of solution. However, the VIS values of the group receiving HHB solution were initially higher than the ones of the HTK group and remained higher over time. It occurs that, in this group, we had two patients with special evolutions, needing more drugs and reaching VIS of 111 and 126. The first case was a total correction of complete atrioventricular canal (CAVC) in a 5-month-old infant with Down syndrome, which evolved with an important vasoplegic condition and pneumonia, requiring prolonged use of adrenaline. Its ventricular function in the immediate postoperative (IPO) and 1PO TTEs remained normal. The second case was 2-month-old and 3300-g child undergoing emergency surgery for heart failure. The preoperative TTE showed large VSD with ASD and patent ductus arteriosus (PDA). In the intraoperative period, multiple VSDs were detected, with a larger VSD and ASD and PDA being corrected. The TTE in IPO showed ejection fraction (EF) of 54% and  $\Delta D$  of 25%; in 1PO, those were 61% and 30%, respectively. This child, however, progressed with severe systemic inflammatory response syndrome (SIRS), evolving to death in 16PO, the only death of the study. We chose to maintain these two cases in the study, considering that they were not able to significantly interfere with the behavior of the groups and through the description of the facts. Interestingly, the behavior of VIS was not accompanied by changes in cardiac enzyme values and ventricular function in TTE.



**Table 4.** Values of the marginal means of troponin, CK-MB, VIS, and TTE between the groups at the different moments analyzed.

Variables	Group A/Group B			
	Troponin	CK-MB	VIS	TTE
Preoperative	0.15/0.04	5.44/8.91	-	40.5/41.3
End of surgery (0h)	5.89/5.90	191.28/206.73	8.22/16.48	33.6/31.4
After 2 hours	-	-	11.75/15.01	-
After 4 hours	-	-	11.68/15.01	-
After 6 hours	3.99/3.87	163.45/166.14	10.51/15.29	-
After 8 hours	-	-	9.95/13.20	-
After 10 hours	-	-	9.68/13.11	-
After 12 hours	2.88/3.11	123.18/108.42	8.67/11.99	-
After 14 hours	-	-	8.18/11.60	-
After 16 hours	-	-	7.95/11.28	-
After 18 hours	2.21/2.69	76.51/61.79	6.44/10.74	-
After 20 hours	-	-	5.93/9.91	-
After 22 hours	-	-	5.54/9.80	-
After 24 hours	2.17/2.45	55.33/44.10	5.09/9.26	34.9/34.4
After 26 hours	-	-	5.09/9.09	-
After 28 hours	-	-	5.17/9.09	-
After 30 hours	-	-	5.08/8.67	-
After 32 hours	-	-	5.08/8.67	-
After 34 hours	-	-	5.01/8.67	-
After 36 hours	-	-	4.66/8.64	-
After 48 hours	1.80/1.83	17.91/12.39	-	-
P2	0.997	0.693	0.449	0.319

CK-MB=creatin kinase-MB; TTE=transthoracic echocardiography; VIS=vasoactive-inotropic score

Regarding the ventricular function, although we analyzed the EF and  $\Delta D$  (shortening), we chose the latter for graphic demonstration in our study. We emphasize that, as expected, both variables showed compatible variations. We observed that there was a variation over time with a reduction of  $\Delta D$  in the first 24 hours in relation to the preoperative examination, followed by partial recovery in the next examination, and there was no statistically significant difference between the groups. In summary, the HTK group received fewer vasoactive drugs than the HHB group as well as presented less reduction of ventricular function, although without statistically significant difference. The HHB group (Group B), on the other hand, showed a lower initial peak of troponin and faster recovery of CK-MB, also with  $P>0.05$ , than the HTK group.

Some comparative studies have already been published on other cold crystalloid and intermittent heated blood solutions in pediatric cardiac surgery, showing better results with the latter solution (less release of troponin and clinical evolution)<sup>[15-17]</sup>. Evaluations of the HTK solution in relation to HHB solutions

in adults have also been published with conflicting results and the intermittent HHB solution appeared to be superior to the single dose of HTK in left ventricular preservation after 60 minutes of aortic cross-clamping in a model with juvenile pigs<sup>[20]</sup>. Our study is distinguished by the comparison of HTK solution and intermittent HHB solution among homogeneous groups of children with acyanotic congenital heart disease with mean ischemia time close to 60 minutes. Considering more severe patients and prolonged aortic clamping periods, a non-randomized Chinese study, published in 2016, compared HTK and St. Thomas' solutions in a group of patients undergoing dextro-transposition of the great arteries (d-TGA) correction with VSD and major pulmonary hypertension. In this experiment, the HTK solution promoted better results than the St. Thomas' solution, which was translated into lower mortality and multiple organ failure<sup>[21]</sup>. Bojan et al.<sup>[22]</sup> published in 2013 a French study comparing HTK solution and intermittent tepid blood solution in children undergoing Jatene surgery. In that study, the group that received HTK solution presented higher hemodilution, requiring

higher ultrafiltration, higher release of troponin after surgery and higher mortality rate in 30 days, suggesting that the intermittent tepid blood solution was more effective. However, there was a large difference between the sample size of the groups (30 HTK and 188 blood solution cases) and the HTK group was formed by younger, smaller children than the ones in the intermittent tepid blood solution group, with a higher incidence of coronary anomalies. The time of ischemia varied between mean times of 72 to 124 minutes, depending on whether there were VSD or aortic arch conditions associated with d-TGA. After a risk-adjusted analysis excluding coronary anomalies, the postoperative troponin concentration remained higher in the HTK group than in the intermittent tepid blood solution group. Still in Jatene surgeries, Giordano et al.<sup>[23]</sup> published in 2015 a comparative analysis of HTK and cold blood solutions, showing similar results and being the HTK solution the choice for the group due to its simple use. The group receiving HTK, however, has been operated on in more recent years. In our study, changes in the troponin and CK-MB curves showed no difference. Both groups were composed of 25 patients and the variables characterized by these patients were similar. Our mean duration of ischemia was slightly less than 60 minutes, although some cases had longer times. The difficulty in forming similar groups for comparison was a recurrent aspect in such studies.

In 2001, Careaga et al.<sup>[18]</sup> published a study comparing HTK and HHB solutions in two groups of 15 patients. It has shown benefits of the crystalloid solution in several aspects, including a lower incidence of arrhythmias and need for vasoactive drugs. The study, however, mixes heart diseases acquired in adults and pediatric surgeries, uses more than one blood solution, and evaluates variables that we judge difficult to control and, thus, were excluded from our study, as length of stay in ICU, which may suffer various influences such as infection, incidence of arrhythmias, and use of vasoactive drugs. Regarding this last variable, we find interesting the use of a score, as suggested by Butts et al.<sup>[14]</sup>, for standardization. In 2002, Viana et al.<sup>[24]</sup> published in the European Journal of Cardio-Thoracic Surgery an Australian comparative experience between HTK solution and tepid blood solution. It was a retrospective study of 1900 adult patients who underwent surgery. Only 7% of these formed the group that used HTK. The groups were still very heterogeneous as in the presentation of risk factors and time of aortic clamping. In a first analysis, the benefits favored the blood solution, like lower mortality, incidence of arrhythmias, and time of mechanical ventilation, another variable of difficult consideration due to the biases that involve it. After a "propensity-matched analysis" with two groups of 71 patients, the results were similar for both solutions. Another interesting aspect would be the analysis of variations of the HTK solution. A Brazilian study, published in 2014, found no difference between the original composition and another one with glutamate replacing ketoglutarate in an experimental model with rats.<sup>[25]</sup>

In our study, two homogenous groups of 25 children with acyanotic congenital cardiopathies were prospectively analyzed for two types of cardioplegic solution used. The preoperative variables of the samples were similar. These characteristics classify the study as original and facilitate the correlation of data

and interpretation of the results. New studies will be developed with the objective of comparing subgroups of congenital heart diseases and other methods of myocardial protection following the line of research of our group. The main limitation of the study is the size of the population analyzed. The right ventricular function has not been studied and should be an additional variable in future studies.

## CONCLUSION

We conclude that, in patients with acyanotic congenital cardiopathies who underwent total surgical correction, mean aortic clamping time around one hour, and CPB with moderate hypothermia, the HTK crystalloid cardioplegic solution offers the same myocardial protection as the cold blood hyperkalemic cardioplegic solution, analyzed according to the variables considered in our study model.

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**No conflict of interest.**

## Authors' roles & responsibilities

ASV	Substantial contributions; final approval of the version to be published
GPL	Substantial contributions; final approval of the version to be published
LAMM	Substantial contributions; final approval of the version to be published
AL	Agreement; final approval of the version to be published
FAM	Agreement; final approval of the version to be published
AG	Agreement; final approval of the version to be published
FAR	Substantial contributions; final approval of the version to be published
RGP	Substantial contributions; final approval of the version to be published
KCB	Revising; final approval of the version to be published

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