The neuroprotective role of therapeutic hypothermia after cardiac arrest

Papel neuroprotector da hipotermia terapêutica pós paragem cardio-respiratória

ABSTRACT

Objectives: Therapeutic hypothermia following cardiorespiratory arrest has been demonstrated to have cardio- and neuroprotective effects, resulting in improved survival and better neurological outcomes. The objective of this study was to assess the outcomes of patients undergoing therapeutic hypothermia following cardiorespiratory arrest.

Methods: A prospective, 10-month observational study of patients admitted to an intensive care unit and undergoing therapeutic hypothermia after cardiorespiratory arrest was undertaken. Therapeutic hypothermia was induced by cold fluid administration and body surface cooling in patients admitted no more than 12 hours after resuscitation from cardiorespiratory arrest. A target temperature of 33ºC was maintained for 24 hours.

Results: Overall, 12 patients were included (median age 64 years, 58% male). Half of the cardiorespiratory arrests were in-hospital. The median first-day Charlson Index, Sequential Organ Failure Assessment (SOFA) and Acute Physiology and Chronic Health Evaluation II scores were of 2.9, 11 and 24.5, respectively. The intensive care unit mortality rate was 42% (N=5). Five of the 7 surviving patients recovered their pre-cardiorespiratory arrest neurological status. Hypothermia was initiated 120 min (median) after recovery of spontaneous circulation. Most patients (75%) required vasopressor support. During the first 3 days after cardiorespiratory arrest and therapeutic hypothermia, a progressive SOFA score decrease (median 11 on day 0, 10 on day 1 and 7 on day 2) was observed.

Discussion: In this study, therapeutic hypothermia was applied to all post-cardiorespiratory arrest patients and demonstrated good neurological outcome in surviving patients.

Keywords: Hypothermia, induced; Heart arrest/therapy; Neurological recovery

INTRODUCTION

Cardiorespiratory arrest (CRA) is associated with increased morbidity and mortality. The survival rate of in-hospital CRA cases is below 20%, and this rate is reduced by half in out-of-hospital cases.1 Notwithstanding the past 50 years of progress in both basic and advanced life support, the mortality rate and severity of neurological sequelae following the recovery of spontaneous circulation (RSC) remain high,2 jeopardizing long-term quality of life.

In CRA patients with RSC, the mortality rate and neurological sequelae following CRA are primarily ascribed to two pathophysiological mechanisms:
anoxic encephalopathy resulting from brain circulation impairment and reperfusion syndrome. This syndrome is characterized by a systemic inflammatory response triggered during reperfusion that is a consequence of the activation of several biochemical cascades, leading to release of free oxygen radicals and other molecules that are harmful to brain cell mediators. This inflammatory response may be prolonged for 48-72 hours.\textsuperscript{1,3,4}

Moderate therapeutic hypothermia (TH) consists of cooling the body to reach a central temperature of 33 ± 0.5ºC, which is aimed to prevent and/or reverse the mechanisms responsible for the reperfusion syndrome and neurological injury. TH has been used since the mid-1940s, when the first cases were described\textsuperscript{5} and was shown to be effective in animal studies in the mid-1980s.\textsuperscript{6,7} However, TH has only recently been validated as an effective CRA therapy. The cardio-protective effects of TH and the reduced mortality rates and neurological sequelae following CRA have been clearly demonstrated in two randomized and controlled trials.\textsuperscript{8,9} These trials have led to the inclusion of TH in the CRA care recommendations by the International Liaison Committee on Resuscitation (ILCOR) since 2003 with an I-B evidence level.\textsuperscript{9} Despite the inclusion of TH in this recommendation, the practical use of TH continues to be less frequent than anticipated\textsuperscript{10} in many European countries, including Portugal.

The most recent ILCOR recommendations state that TH should be used in adult coma patients after CRA due to outpatient ventricular fibrillation and can be beneficial after CRA due to in-hospital, non-shockable rhythms.\textsuperscript{11} This study assessed the feasibility of a TH protocol and outcomes in CRA patients undergoing this therapeutic intervention.

METHODS

A prospective observational study was conducted that included patients admitted to the general intensive care unit (ICU) of the Hospital de São Francisco Xavier, Centro Hospitalar de Lisboa Ocidental (CHLO) undergoing TH after CRA during a 10-month period from August 2009 to May 2010. The study was approved by the hospital's ethics committee. Informed consent was waived because of the observational nature of this trial, with no additional interventions being made in addition to the usual therapeutic approach for these patients.

Inclusion criteria were spontaneous circulation maintained for ≥ 5 minutes, systolic blood pressure (SBP) > 80 mmHg, time from CRA ≤ 12 hours, time until advanced life support (ALS) ≤ 10 minutes, ALS time ≤ 45 minutes, time of hypotension after RSC ≤ 30 minutes (SBP < 80 mmHg or mean blood pressure (MBP) < 45 mmHg) and a Glasgow Coma Scale (GCS) score ≤ 9.\textsuperscript{12}

Exclusion criteria were primary coagulopathy, terminal disease or indications for non-resuscitation, aorta dissection, intracranial bleeding, significant bleeding or myoclonic status epilepticus.

TH was performed according to the local protocol. TH was induced by fast fluid infusion, preferably with polyelectrolyte solutions at 4ºC, 30 mL/kg at 100 mL/min, external cooling by ice packs applied to the axilla and groin areas and eventually wet sheets covering the hands and feed. Additionally, the ventilator’s circuit heating and humidification was suspended, and a gastric and vesical lavage with 4ºC saline was performed. This cooling can be performed using peripheral venous access, the process was not delayed by central venous catheter (CVC) placement. The temperature was maintained using wet towels or thermal blankets. Ventilation was maintained on control mode, aiming to maintain peripheral oxygen saturation between 93 and 95%, thereby preventing hypoxemia and hypocapnia. Continuous sedation and analgesia were maintained using midazolam and alfentanil; propofol was avoided, due to its negative inotropic effects. In addition, a neuromuscular blockade was maintained using vecuronium for shivering prevention. The protocol included invasive blood pressure monitoring (arterial line), which also provided access for blood gas control and blood samples for additional laboratory testing. The temperature was continuously monitored using a pharyngeal thermometer.

For maintaining a MBP above 80 mmHg, a venous line was maintained for fluid replacement and/or vasopressor drug administration. In cases of renal failure, metabolic acidosis or insufficient cooling with the above described measures (temperature reduction > 1.5ºC or temperature < 35ºC after 90 minutes), continuous renal replacement was started (hemofiltration or continuous veno-venous hemodiafiltration) with active extracorporeal circuit cooling with ice. Laboratory panels were performed daily, including specific neural enolase (SNE) dosage.

All patients underwent stress ulcer prophylaxis, deep venous thrombosis and decubitus ulcers. Additionally, magnesium sulfate perfusion and strict potassium, and blood glucose monitoring were performed. A temperature of 33 ± 0.5ºC was targeted and maintained for 24 hours.

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Reheating was slow and passively conducted for at least 8 hours at no more than a 0.5°C/hour pace with hourly blood glucose monitoring, due to the increased risk for hypoglycemia. After the body temperature was ≥ 36ºC, the neuromuscular blockade and then sedation were suspended.

The following clinical and demographic data were collected from the patients undergoing TH: age, gender and Charlson index. Acute Physiology and Chronic Health Evaluation II (APACHE II) and Sequential Organ Failure Assessment (SOFA) scores. CRA was characterized according to the occurrence site, ALS maneuver duration, time until RSC (arrest time) and RSC heart rhythm. The TH parameters assessed were as follows: time from RSC to TH initiation, time from TH initiation until reaching the target temperature (33ºC), TH duration, minimum temperature achieved, volume of cold fluid infused, requiring vasopressor drug support, requiring extracorporeal circulation, SNE values, C-reactive protein and white blood cell count. Changes in SOFA scores and neurological outcomes according to the Glasgow Coma Scale (GCS) (1. death; 2. vegetative state; 3. severe disability; 4. moderate disability; 5. good recovery) were also assessed within the 3 days after CRA and TH (D0. TH day; D1. reheating phase; D2. day after reheating), as well as in-hospital mortality.

Descriptive statistical analyses were performed with Microsoft Office Excel (version 2007) software. Although survival and non-survival group variables were analyzed, we chose to present only descriptive analyses, due to the small sample size. The results are expressed as the median and interquartile interval (IQI) or mean ± standard deviation, depending on the variable distribution.

**RESULTS**

Twelve patients with a median age of 64 years (IQI 29) were included during the trial period. Most patients were male (58.3%). The mean APACHE II scores were 24.5 (IQI 15.25), and the Charlson score was low in five (41.6%), moderate in three (25%) and high in four (33%) patients. The admission GCS was 5 (IQI 3.75).

CRA occurred in six patients (50%) in-hospital and two in the ICU (Table 1). The CRA time was 12.5 minutes (IQI 13.3 minutes), and the time to RSC (duration of ALS maneuvers) was 7 minutes (IQI 7.5 minutes). Sinus rhythm was predominant upon RSC in 75% of cases, with atrial fibrillation in 16.6% and other supra-ventricular rhythms being observed in 8.3% of cases.

<table>
<thead>
<tr>
<th>Table 1 – Cardiorespiratory arrest site</th>
</tr>
</thead>
<tbody>
<tr>
<td>CRA site</td>
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<tr>
<td>--------------</td>
</tr>
<tr>
<td>ICU</td>
</tr>
<tr>
<td>Regular ward</td>
</tr>
<tr>
<td>Home</td>
</tr>
<tr>
<td>Total</td>
</tr>
</tbody>
</table>

CRA – cardiorespiratory arrest; ICU – intensive care unit.

**Hypothermia use**

The median time from RSC until initiating TH was 120 minutes (IQI 75 minutes). The time required to reach the target body temperature varied but was 240 minutes (IQI 327.5 minutes) on average, and the duration of hypothermia was 27 hours on average. All patients reached the target temperature, and the minimal temperature was 32.2ºC.

To reach the target temperature, each patient was infused with 5.3 ± 2.9 liters of cold fluids. In most patients, the target temperature was reached using external cooling and cold fluids. In one extreme case, 12.5 liters were required, and in two other patients, continuous renal replacement therapy was necessary.

In most cases (9 patients, 75%), vasopressor support was required during TH. Vasoactive amines selection was conducted under the discretion of the treating physician. The maximal doses given during TH were 5.82 µg/kg/min noradrenalin and 7.5 µg/kg/min dopamine.

SNE was 21.4 (IQI 14 ng/mL) on D1 and 27.15 (IQI 28.9 ng/mL) on D2. However, in some patients (N = 5), SNE was not assessed or was incompletely assessed during TH.

**Clinical outcome**

Of the included patients, 5 died in the ICU (41.6% mortality rate). The neurological outcome among the survivors was full recovery in 5 and severe disability and vegetative state in the remaining patients. One patient eventually died on the regular ward, resulting in a 50% in-hospital mortality rate.

The SOFA scores improved during TH within the first 3 days in the ICU: 11 (IQI 2.75) on the TH initiation day, 10 (IQI 3) on the reheating day and 7 (IQI 4.5) after reheating (Figure 1).

Survival with full neurological recovery was seen in all patients experiencing CRA in the ICU (N = 2; 16%). Half of the patients experiencing in-hospital (N = 4; 33.3%) and out-of-hospital CRA (N = 6; 50%) died; the survivors recovered with moderate to severe disability in the first group and with no sequela in the second group (Table 2).
Age, Charlson index scores, arrest times (CRA + ALS times) and time before starting the TH protocol (median [IQI]) were higher among patients experiencing mortality compared with those with good neurological outcomes (score 5) (66 [33] vs. 56 [43] years; 4.5 [6.8] vs. 1.6 [7]; 30 [25] vs. 24 [8] minutes and 135 [70] vs. 120 [90] minutes, respectively). The time until achieving the target temperature was widely variable both in the good neurological outcome and deceased patient groups.

The SNE value progression could not be assessed during the TH because dosages were partially missing in some patients. The C-reactive protein values increased during TH in all patients (2.3, 2.7 and 4.1 mg/dL on D0, D1 and D2, respectively).

Complications

No evidence of skin injury associated with external cooling was found. Three infective complications were diagnosed after being discharged from the ICU: nosocomial pneumonia in one patient with a good neurological outcome and two patients who died with nosocomial pneumonia and invasive candidiasis.

Figure 1 - Sequential Organ Failure Assessment score progression during therapeutic hypothermia.

D0 – day of therapeutic hypothermia; D1 – time of reheating; D2 – day after reheating; CRA – cardiorespiratory arrest; SOFA - Sequential Organ Failure Assessment.

Table 2 - Clinical and demographic characteristics of patients undergoing therapeutic hypothermia, characterization of cardiorespiratory arrest, hypothermia variables and neurological outcomes

<table>
<thead>
<tr>
<th>N</th>
<th>Age and gender</th>
<th>Site of CRA</th>
<th>Cause of CRA</th>
<th>ALS time (minutes)</th>
<th>Initial rhythm</th>
<th>Charlson</th>
<th>APACHE II</th>
<th>Vasopressors</th>
<th>Volume of Fluids (mL)</th>
<th>Time of CRA (minutes)</th>
<th>Time to start TH (minutes)</th>
<th>Time to target temperature (min)</th>
<th>Time of TH duration (h)</th>
<th>Neurological outcome</th>
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<tbody>
<tr>
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<td>SVT</td>
<td>7</td>
<td>17</td>
<td>Y</td>
<td>6890</td>
<td>10</td>
<td>20</td>
<td>5</td>
<td>28</td>
<td>11</td>
</tr>
<tr>
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<td>ICU</td>
<td>Asystole</td>
<td>5</td>
<td>AF</td>
<td>7.4</td>
<td>42</td>
<td>Y</td>
<td>3000</td>
<td>5</td>
<td>30</td>
<td>120</td>
<td>24</td>
<td>12</td>
</tr>
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<td>SR</td>
<td>4.5</td>
<td>32</td>
<td>Y</td>
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<td>45</td>
<td>135</td>
<td>0</td>
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<td>21</td>
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<td>SR</td>
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<td>28</td>
<td>N</td>
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<td>10</td>
<td>180</td>
<td>420</td>
<td>44</td>
<td>9</td>
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<td>6</td>
<td>80. ♂</td>
<td>Hospital</td>
<td>Ventricular fibrillation/AMI</td>
<td>?</td>
<td>SR</td>
<td>7</td>
<td>42</td>
<td>Y</td>
<td>3550</td>
<td>2</td>
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<td>17</td>
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<td>10</td>
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<td>6</td>
<td>N</td>
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<td>120</td>
<td>240</td>
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<tr>
<td>11</td>
<td>27. ♂</td>
<td>Home</td>
<td>Asystole</td>
<td>15</td>
<td>SR</td>
<td>0</td>
<td>16</td>
<td>Y</td>
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<td>20</td>
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<td>70. ♂</td>
<td>Hospital</td>
<td>Aspiration pneumonia</td>
<td>2</td>
<td>SR</td>
<td>1.6</td>
<td>33</td>
<td>Y</td>
<td>5800</td>
<td>2</td>
<td>120</td>
<td>240</td>
<td>22</td>
<td>9</td>
</tr>
</tbody>
</table>

APACHE II - Acute Physiology and Chronic Health Evaluation; D0 – day of therapeutic hypothermia; D1 – reheating period; D2 – day after reheating; AMI -acute myocardial infarction; VF- ventricular fibrillation; GCS - Glasgow Coma Scale; TH- therapeutic hypothermia; N – no; NA – not applicable; CRA – cardiorespiratory arrest; ALS- advanced life support; Y – yes; SOFA - Sequential Organ Failure Assessment; and Neurological outcome (Glasgow Outcome Scale: 1. death; 2. vegetative state; 3. severe disability; 4. moderate disability; 5. good recover); SVT- supraventricular tachycardia; ICU- intensive care unit.
DISCUSSION

In the current study, TH was successfully used in CRA patients. The in-hospital mortality rate was 50%, i.e., less than the 71.4% reported by Nolan et al.\(^2,9\) and 66% reported by Ravetti et al.\(^18\) However, this finding was influenced by the small sample size, inclusion of both in-hospital and out-of-hospital CRA cases and inclusion of all arrest rhythms. Another aspect that may have influenced this result was the lower number of cardiovascular origin CRA.

In our series, the deceased patients were older, had more comorbidities and had longer CRA times. A full neurological recovery was observed in 41.6% of patients; however, if only the surviving patients are considered, a full recovery was seen in 71.4% of patients. Patients with the best neurological recovery after TH were younger and had fewer comorbidities (higher GCS and lower Charlson index scores at admission).

Similar to other small series, a direct relationship was identified between good neurological outcomes and shorter CRA times and times to initiating TH.\(^{19}\) Similarly, in our study, the CRA time and time to initiating TH were shorter among survivors.

Although evidence for the benefit of TH is clear in out-of-hospital cases of ventricular fibrillation or tachycardia,\(^{20,21}\) in our series, this therapeutic approach was used independently of the pre-CRA rhythm or arrest site.

The neurological outcome was better for in-ICU CRA patients, which agrees with other studies.\(^{18,22}\) It is reasonable to assume that continuous hemodynamical monitoring is beneficial for prompt arrest recognition and therapy, the recognition of potentially reversible causes and significantly shortening the time to initiating basic and advanced life support measures and is additionally advantageous for assessing the effectiveness of treatment. After HACA and Bernard\(^4,8\) demonstrated the effectiveness of TH on neurological outcome in CRA patients, TH was included in the 2003 and 2010 ILCOR recommendations.\(^9,21\) However, these recommendations do not state an exact method or protocol for reaching the target temperature. Several external and internal methods are available for achieving the target temperature.\(^21\) One of the most frequently used methods is the perfusion of cold fluids (usually 4ºC saline at 30 mL/kg), which is able to reduce central temperature by 1.5ºC. Other possible methods for inducing and maintaining TH are ice bags, cool towels, cooling pillows or blankets, gel pillows with circulating water, intravascular heat exchangers and extracorporeal circulation.\(^{21}\) To the best of our knowledge, no study has compared the effectiveness of these different TH induction methods.

The use of TH in our study was previously defined by a local protocol based on external cooling and cold fluids with no special technology requirements and therefore no incremental costs. It is of note that the target temperature was achieved for all patients. Internal (4ºC fluid infusion) and external (ice bags to the axilla and groin areas and eventually wet sheets, gastric and vesical lavage with cold fluids) methods were used, frequently requiring vasopressor drug support. Only two patients required dialysis to reach the target temperature. In most patients, the hypothermia protocol was both simple and affordable. On average, the target temperature was reached after 5 hours (293 minutes); this finding agrees with those of published literature reports.\(^{4,8}\)

One case of nosocomial pneumonia was identified among the surviving patients and two cases among the deceased patients. No arrhythmia, coagulopathy, skin ice-burning injury or severe hydroelectrolytic changes were detected.

Few studies on TH are available in Portugal, suggesting that this therapy is not common. Our study has clearly shown that this proven, beneficial and recommended technique is feasible and safe and can be used in any ICU.

Although our results agree with those of larger series, the limited number of patients, the short follow-up period, this study’s retrospective nature and the inclusion of different CRA rhythms and sites render our findings merely descriptive. Additionally, it was not possible to assess the SNE level progression; therefore, no prognostic assessment based on this parameter could be made.

CONCLUSIONS

ILCOR recommends TH for CRA patients with an I-B evidence level. The method used in this study required no special technology need or expensive resources and was proven simple and effective, providing good neurological outcomes for most surviving patients. The implementation of TH was simplified by the availability of a previous protocol.
RESUMO

Objectivos: A hipotermia terapêutica demonstrou ter efeitos neuro e cardioprotectores, com melhoria da sobrevida e redução das sequelas neurológicas em doentes vítimas de paragem cardio-respiratória. O objectivo deste estudo foi avaliar a evolução dos doentes submetidos a hipotermia terapêutica após paragem cardio-respiratória.

Métodos: Estudo prospectivo observacional dos doentes submetidos a hipotermia terapêutica após paragem cardio-respiratória numa unidade de cuidados intensivos polivalente durante 10 meses. Aos doentes admitidos até 12 horas após paragem cardio-respiratória foi induzida a hipotermia terapêutica através da administração de fluidos arrefecidos e arrefecimento corporal externo e mantida a temperatura alvo, 33°C, durante 24 horas.

Resultados: Foram incluídos 12 doentes, idade (mediana) de 64 anos, 58% do sexo masculino. A paragem cardio-respiratória ocorreu em meio hospitalar em 6 doentes. O índice de Charlson, o Sequential Organ Failure Assessment (SOFA) e o Acute Physiology and Chronic Health Evaluation II, no primeiro dia, foram 2.9 [IIQ 6.8], 11 [IIQ 2.75], e 24.5 [IIQ 15.25], respectivamente. A taxa de mortalidade na unidade de cuidados intensivos polivalente foi de 42% (N=5). Dos 7 sobreviventes, 5 recuperaram o estado neurológico prévio e uma diminuição do valor mediano de SOFA constatado, nos 3 dias subsequentes à paragem cardio-respiratória. A hipotermia terapêutica foi iniciada cerca de 120 minutos [IIQ 78.75], após recuperação de circulação espontânea. A maioria dos doentes (75%) necessitou de suporte vasopressor. Foram incluídos 12 doentes,idade (mediana) de 64 anos, 58% do sexo masculino. A paragem cardio-respiratória ocorreu em meio hospitalar em 6 doentes. O índice de Charlson, o Sequential Organ Failure Assessment (SOFA) e o Acute Physiology and Chronic Health Evaluation II, no primeiro dia, foram 2.9 [IIQ 6.8], 11 [IIQ 2.75], e 24.5 [IIQ 15.25], respectivamente. A taxa de mortalidade na unidade de cuidados intensivos polivalente foi de 42% (N=5). Dos 7 sobreviventes, 5 recuperaram o estado neurológico prévio e uma diminuição do valor mediano de SOFA constatado, nos 3 dias subsequentes à paragem cardio-respiratória. A hipotermia terapêutica foi iniciada cerca de 120 minutos [IIQ 78.75], após recuperação de circulação espontânea. A maioria dos doentes (75%) necessitou de suporte vasopressor.

Conclusão: A aplicação de um protocolo de hipotermia terapêutica revelou ser simples e eficaz e permitiu obter em doentes com indicação, boa recuperação neurológica.

Descritores: Hipotermia induzida; Parada cardíaca/terapia; Recuperação neurológica

REFERENCES


