Out-of-Hospital Cardiac Arrest

Studies on aetiology, treatment and outcome

ACADEMIC DISSERTATION
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To my wife Tarja and our children Salla, Tuomas and Aapo.
Abstract

The survival rate after out-of-hospital cardiac arrest (OHCA) has not improved much over the last decade. The aim of this thesis was to study treatment and outcome of OHCA in the emergency medical service (EMS) systems in Tampere District EMS and the physician-staffed helicopter emergency medical service (HEMS) in the Helsinki and Turku areas in Southern Finland.

Gastric regurgitation and pulmonary aspiration are serious adverse events in OHCA. The objective was to determine whether there is an association between bystander mouth-to-mouth ventilation and regurgitation in prehospital cardiac arrest patients. In this prospective material, the study population consisted of 529 consecutive prehospital cardiac arrest patients with attempted resuscitation. Exclusion criteria were cardiac arrest due to trauma or drug overdose. The EMS personnel documented if regurgitation was present in OHCA patients on the scene. Regurgitation occurred in a fourth of patients. Bystander cardiopulmonary resuscitation (CPR) with mouth-to-mouth ventilation was associated with a significantly increased risk of regurgitation compared with no CPR and CPR without ventilation. The mode and role of bystander CPR in cardiac arrest needs to be further evaluated.

The association between clinical signs of regurgitation and radiological findings consistent with aspiration in resuscitated OHCA patients admitted to hospital were studied. The incidence of regurgitation was studied in 182 successfully resuscitated OHCA patients. The inclusion criterion was the restoration of spontaneous circulation (ROSC) after OHCA not caused by trauma or drug overdose. The incidence of regurgitation was 20%. Regurgitation was associated with radiological findings consistent with aspiration with high specificity (81%) and low sensitivity (46%). Although there was a strong association between clinical regurgitation and radiological findings consistent with aspiration, our data suggest that regurgitation is not invariably followed by radiological findings compatible with aspiration. Radiological findings consistent with aspiration appeared to be relatively infrequent without preceding signs of regurgitation in resuscitated patients.

The objective in Study III was the cause of death (COD) in patients who died after an unsuccessful attempt at out-of-hospital resuscitation when the initial cardiac rhythm had been pulseless electrical activity (PEA). The aim was to determine whether there is a difference in the distribution of CODs between those who underwent autopsy and those whose COD was estimated based on clinical and previous medical history. Data were collected from 91 patients treated by the emergency medical service systems. An autopsy was performed on
59 patients, while the COD was determined without autopsy in 32 patients. There were significantly more diagnoses of acute myocardial infarction (AMI) and less pulmonary embolism (PE), aortic dissection and rupture among those without autopsy compared with those who underwent autopsy. The conclusion was that in unsuccessful resuscitation from OHCA with PEA as initial rhythm, an autopsy should be performed to determine the exact cause of death.

Mild therapeutic hypothermia improves neurological outcome after cardiac arrest. The cooling and haemodynamic effects of prehospital infusion of ice-cold Ringer’s solution were studied in 13 adult patients after successful resuscitation from non-traumatic cardiac arrest. After haemodynamic stabilisation, 30 ml/kg of Ringer’s solution was infused at a rate of 100 ml/min into the antecubital vein. Arterial blood pressure and blood gases, pulse rate, end-tidal CO₂ and oesophageal temperature (T_{esof}) were monitored closely. The mean core temperature decreased significantly from 35.8 ± 0.9 °C at the start of infusion to 34.0 ± 1.2 °C on arrival at hospital. No serious adverse haemodynamic effects occurred. It was concluded that the induction of therapeutic hypothermia using this technique in the prehospital setting is feasible.
Sairaalan ulkopuolella tapahtuneen sydänpysähdyksen ennuste ei ole parantunut paljoa viime vuosikymmenen aikana. Tämän väitöskirjan tarkoituksena oli tutkia hoitoa ja selviytymistä sairaalan ulkopuolella tapahtuneesta sydänpysähdyksestä Tampereen aluepelastuslaitoksen, Medi-Heli 01:n ja Medi-Heli 02:n alueilla Etelä-Suomessa.


Toisessa osatyössä tutkittiin kemiallisesti havaitun elvytyksen aikaisen regurgitaation vaikutusta sairaalassa tehtyihin radiologisiin löydöksiin. Tutkimukseen otettiin mukaan 182 potilasta, jotka olivat onnistuneesti elvytetty sairaalan ulkopuolella ja joiden elottomuuden syy ei ollut trauma tai myrkytys. Näillä potilailla regurgitaatio ilmeni 20 %. Kentällä todetun regurgitaation yhteydessä radiologisiin löydöksiin todettiin korkealla spesifiteetillä (81 %) ja matalalla sensitiviteetillä (46 %). Vaikka todetun regurgitaation ja aspiraation sopivien radiologisten löydösten täällä oli vahva riippuvuus, näin tulostemme mukaan aspiraatista ei aina seuraa radiologisesti todennettavia muutoksia. Aspiraation vaarantaa radiologiset löydökset ovat suhteellisen harvinaisia elvytetyillä potilailla, ellei elvytyksen aikana kemiallisesti havaintoa regurgitaatiosta.

Sairaalan ulkopuolella epäonnistuneesti PEA-alkurytmistä elvytetyjen sydänpysähdyspotilaan kuolinsyyt tutkittiin. Tarkoituksena oli selvittää, poikkeavatko ruumiinnavaulta perusteella määritettyjen kuolinsyyjen jakauma potilaan lääketieteellisen historian ja elvytyksen kulun perusteella kliinisesti määritettyjen kuolinsyyjen jakaumasta. Tutkimukseen otettiin mukaan 91 epäonnistuneeseen elvytyseen päättynyt sydänpysähdyspotila PEA-alkurytmillä, joista ruumiinavaus suoritettiin 59 vainajalle. 32 vainajan kuolinsyyt määritettiin edellä mainituin kliinisin perusteella. Sydäninfarktien osuus kuolinsyistä oli merkitsevästi yleidestettuna ja keuhkoveritulpat ja aortan repeämät tai dissekaatiot aliedustettuina niillä vainajilla, joiden kuolinsyy oli
määritetty kliinisin perustein. Voidaankin todeta, että PEA-alkurytmillä alkaneseen sydänpyhsähdykseen menehtyneen vainajan kuolinsyy tulee määrittää ruumiinavauksella todellisen kuolinsyyn selvittämiseksi.

Hypotermiahoito sydänpyhsähdyksen jälkeen parantaa ennustetta. Ensimmäisessä osatyössä selvitetiin jääkylmällä Ringerin nesteellä toteutetun hypotermiahoidon toteutumista ja verenkiegamma vaikutuksia 13 potilaalla sairaalan ulkopuolisen sydänperäisen elottomuuden jälkeen. Verenkiegymen jälkeen, 30ml/kg Ringerin nestettä 100ml/min infusoihin kyynärtaipeen laskimoon. Verikasuja, verenpainetta, pulssia, hengitysilman ulostulevaa hiilidioksidia ja ruokatorven lämpötilaa mitattiin tarkasti. Potilaan ydinlämpö laski merkitsevästi 35.8 ± 0.9 °C:sta 34.0 ± 1.2 °C:een. Vakavia verenkiegymen häiriöitä ei havaittu. Todettiin, että sairaalan ulkopuolella indusoitu hypotermiahoito on tällä menetelmällä toteuttamiskelpoinen.
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## Abbreviations

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<th>Definition</th>
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<tr>
<td>ALS</td>
<td>Advanced life support</td>
</tr>
<tr>
<td>AMI</td>
<td>Acute myocardial infarction</td>
</tr>
<tr>
<td>ARDS</td>
<td>Acute respiratory distress syndrome</td>
</tr>
<tr>
<td>ASY</td>
<td>Asystole</td>
</tr>
<tr>
<td>ATP</td>
<td>Adenosine triphosphate</td>
</tr>
<tr>
<td>BLS</td>
<td>Basic life support</td>
</tr>
<tr>
<td>°C</td>
<td>Symbol for degree Celsius</td>
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<tr>
<td>CA</td>
<td>Cardiac arrest</td>
</tr>
<tr>
<td>COD</td>
<td>Cause of death</td>
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<tr>
<td>CNS</td>
<td>Central nervous system</td>
</tr>
<tr>
<td>CO$_2$</td>
<td>Carbon dioxide</td>
</tr>
<tr>
<td>COPD</td>
<td>Chronic obstructive pulmonary disease</td>
</tr>
<tr>
<td>CPC</td>
<td>Cerebral performance category</td>
</tr>
<tr>
<td>CPP</td>
<td>Coronary perfusion pressure</td>
</tr>
<tr>
<td>CPR</td>
<td>Cardiopulmonary resuscitation</td>
</tr>
<tr>
<td>ECG</td>
<td>Electrocardiogram</td>
</tr>
<tr>
<td>EMD</td>
<td>Electromechanical dissociation</td>
</tr>
<tr>
<td>EMS</td>
<td>Emergency medical service</td>
</tr>
<tr>
<td>ERC</td>
<td>European Resuscitation Council</td>
</tr>
<tr>
<td>EVO</td>
<td>Competitive research funding of the Pirkanmaa Hospital District</td>
</tr>
<tr>
<td>GCS</td>
<td>Glasgow coma score</td>
</tr>
<tr>
<td>H$^+$</td>
<td>Hydrogen ion</td>
</tr>
<tr>
<td>HACA</td>
<td>Hypothermia after Cardiac Arrest Study Group</td>
</tr>
<tr>
<td>HEMS</td>
<td>Helicopter emergency medical service</td>
</tr>
<tr>
<td>H$_2$O</td>
<td>Water</td>
</tr>
<tr>
<td>ICH</td>
<td>Intracranial haemorrhage</td>
</tr>
<tr>
<td>ICU</td>
<td>Intensive Care Unit</td>
</tr>
<tr>
<td>ILCOR</td>
<td>International Liaison Committee on Resuscitation</td>
</tr>
<tr>
<td>LES</td>
<td>Lower oesophageal sphincter</td>
</tr>
<tr>
<td>min</td>
<td>minute</td>
</tr>
<tr>
<td>ml</td>
<td>millilitre</td>
</tr>
<tr>
<td>NNT</td>
<td>Number needed to treat</td>
</tr>
<tr>
<td>NPV</td>
<td>Negative predictive value</td>
</tr>
<tr>
<td>OHCA</td>
<td>Out-of-Hospital Cardiac arrest</td>
</tr>
<tr>
<td>OPC</td>
<td>Overall performance category</td>
</tr>
<tr>
<td>OR</td>
<td>Odds Ratio</td>
</tr>
<tr>
<td>PCO$_2$</td>
<td>Partial pressure of carbon dioxide</td>
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</table>
PE Pulmonary embolism
PEA Pulseless electrical activity
PPV Positive predictive value
ROSC Restoration of spontaneous circulation
SBP Systolic blood pressure
SCA Sudden cardiac arrest
SCD Sudden cardiac death
SD Standard deviation
SpO₂ Saturation of peripheral blood oxygen
ΔT Change in temperature
T eso Oesophageal temperature
VF Ventricular fibrillation
VT Ventricular tachycardia
X-ray Roentgen ray; here radiological imaging study
This thesis is based on the following original publications referred to in the text by Roman numerals I-IV:


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Introduction

Ischaemic heart disease is a leading cause of death in the industrial world and sudden cardiac arrest (SCA) is the cause of death in 60 % of adult deaths from coronary disease (Zheng et al. 2001). Based on data from Finland, the annual incidence of resuscitation for out-of-hospital cardiac arrest (OHCA) of cardiac aetiology is 53 per 100,000 population (Kuisma et al. 1996). Mortality among these patients admitted to hospital after OHCA remains high.

Gastric regurgitation and pulmonary aspiration are serious adverse events in OHCA. Assisted ventilation without a secured airway is often associated with regurgitation, leading to increased morbidity and mortality (Pepe 1996). It is not known how often documented gastric regurgitation during treatment of cardiac arrest leads to radiographic findings compatible with aspiration.

A recent study has suggested that ventilation may not be needed for several minutes after onset of cardiac arrest since outcome after CPR with chest compressions only has been shown to be similar to that with conventional CPR including mouth-to-mouth ventilation (Hallstrom et al. 2000). Also, compression only CPR has been shown to be better than no CPR at all (Bossaert et al. 1989, Van Hoeyweghen et al. 1993).

The distribution of primary cardiac rhythm in cardiac arrest is changing. Although ventricular fibrillation (VF) has been considered the most common initial rhythm (50-83 %) (Weaver et al. 1986, Bayes de Luna et al. 1989) in OHCA, a major decline (50 %) in its incidence has occurred in recent decades. Concomitantly, the number of patients with pulseless electrical activity (PEA) as initial cardiac rhythm has increased (Herlitz et al. 2000, Kuisma et al. 2001, Cobb et al. 2002). Recent studies have shown the incidence of primary PEA to be 22-27 % in OHCA (Engdahl et al. 2001), and as high as 32 % in in-hospital cardiac arrest. The aetiology behind PEA is not very well known and needs further investigation.

Therapeutic induced hypothermia is reported to improve survival and neurological outcome in patients with VF (Bernard et al. 2002; Hypothermia after Cardiac Arrest Study Group 2002). Hypothermia should be induced as soon as possible after return of spontaneous circulation (ROSC) (Safar et al. 2002). Ideally, the technique should already be available in the prehospital setting and should also be easily managed by non-physician prehospital care providers. Medical experience of induced hypothermia after cardiac arrest is based on in-hospital studies and needs to be studied in the field for wider utilisation of this promising technique.
The purpose of the study was to determine whether there is an association between bystander mouth-to-mouth ventilation and regurgitation in prehospital cardiac arrest and to assess the association between prehospital regurgitation and subsequent radiological findings of resuscitated patients in the hospital. In addition, the aetiology behind PEA in unsuccessful resuscitation was studied. A feasibility trial of induced hypothermia soon after ROSC was undertaken.
Cardiac arrest is the sudden, abrupt loss of heart function. It has been estimated that incidence of sudden cardiac death (SCD) is 1 per 1,000 inhabitants annually in USA and Europe (Myerburg et al. 1992, Priori et al. 2001). The incidence of OHCA in Helsinki is 80/100,000 inhabitants/year (Kuisma et al. 1996). Resuscitation is attempted in 50-66/100,000 inhabitants/year (Herlitz et al. 1999). SCA is responsible for more than 60 % of adult deaths from coronary disease (Zheng et al. 2001). The purpose of CPR is to reverse sudden unexpected cardiac arrest from a potentially reversible cause and to restore prearrest life.

1. Historical perspective of resuscitation

Possibly the earliest record of mouth-to-mouth resuscitation can be found in Old Testament, in the Septuagint (LXX) version of Kings, where reads in 21 verse: “Then he blew air into the boy three times” (Paraskos 1992). On October 4, 1858, János Balassa reported an 18 year old woman suffering cardiac arrest due to airway obstruction from ulcerated laryngitis. Tracheotomy was made immediately and Balassa “exerted bellows-like rhythmic pressure to the front of her chest imitating breathing. Air entered the lungs with a sharp whistling sound.” After 6 minutes of resuscitation she begun to breathe and after 15 minutes she regained consciousness (Robicsek et al. 2004). Although all effective therapies were described before the year 1900 (Table 1), it took many decades to integrate these techniques into modern CPR. Zoll reported the first successful defibrillation of human VF with external paddles in 1956 (Zoll et al. 1956). Safar and Elam described mouth-to-mouth ventilation and effective airway techniques in 1958 (Elam et al. 1958, Safar et al. 1958). Kouwenhoven rediscovered closed chest cardiac massage in 1960 (Kouwenhoven et al. 1960). It took six years of synthesis to introduce the first recommendation on CPR (Anonymous 1966). Since 1973 the American Heart Association (AHA) has published “Standards for Cardiopulmonary resuscitation and Emergency Cardiac Care”. Guidelines were updated with publications in 1980, 1986 and 1992. In 1993 on the basis of worldwide co-operation the International Liaison Committee on Resuscitation (ILCOR) was formed to identify and review international science and knowledge relevant to CPR, and to offer consensus on treatment recommendations. In 2005 the European
Figure 1. Current Adult Advanced Life Support (ALS) algorithm. Modified from (Nolan et al. 2005).

The actions performed to rescue the cardiac arrest victim is called the Chain of Survival. It includes early recognition of the emergency and activation of the emergency service, early CPR, early defibrillation and ALS (Nolan 2005).

Table 1
Effective cardiopulmonary resuscitation manoeuvres described before 1900.

<table>
<thead>
<tr>
<th>Technique</th>
<th>Investigator or provider</th>
<th>Year</th>
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<tbody>
<tr>
<td>Intubation</td>
<td>Vesalius</td>
<td>1500s</td>
</tr>
<tr>
<td>Jaw thrust</td>
<td>Esmarch and Heiberg</td>
<td>1800s</td>
</tr>
<tr>
<td>Mouth-to-mouth ventilation</td>
<td>Midwives, Tossach</td>
<td>1700s</td>
</tr>
<tr>
<td>Open-chest cardiac massage</td>
<td>Shiff</td>
<td>1870s</td>
</tr>
<tr>
<td>Closed-chest cardiac massage</td>
<td>Boehm</td>
<td>1874</td>
</tr>
<tr>
<td>Adrenaline</td>
<td>Crile</td>
<td>1898</td>
</tr>
<tr>
<td>Defibrillation</td>
<td>Prevost</td>
<td>1899</td>
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</table>

2. Out-of-Hospital Cardiac arrest

SCA is one of the most frequent CODs in industrialised countries (Herlitz et al. 1999). Despite CPR after OHCA and post-resuscitation care mortality remains high at 44-66% (Langhelle et al. 2003). Survival from cardiac arrest depends on a sequence of interventions, all of which have to be optimised to maximize survival (Cummins et al. 1991). The chain of survival is early recognition and call for help, early CPR, early defibrillation and post resuscitation care (Nolan et al. 2006).

Dispatching centres handle all emergency calls in Finland and dispatch units according to medical risk analysis protocols. Basic Life Support (BLS) units are capable of performing CPR and using an automated external defibrillator. ALS units are capable of securing the airway with intubation tube or supraglottic airway and they can administer intravenous drugs. An emergency physician staffed ambulance unit or helicopter emergency medical service (HEMS) unit is capable of initiating intensive care level procedures on the patient. The dispatching centres have instructions to dispatch the nearest possible unit as a first responding unit in suspected cardiac arrest. Simultaneously a BLS unit and ALS unit are dispatched. In areas where an emergency physician manned ambulance unit or HEMS unit is available, it will also be dispatched (Väisänen et al. 2006).

The risk factors affecting survival from cardiac arrest are several. The most important factors are short time intervals in the treatment of cardiac arrest. Bystander CPR has a beneficial effect on surviving from OHCA (Van Hoeyweghen et al. 1993) and the unfavourable effect of delay in initiation of resuscitation is well documented, especially in initiation of ALS. The chance of survival decreases with each passing minute without defibrillation, CPR or ALS. In cardiac arrest patients with witnessed VF, survival decreased 3% with each minute until CPR was started and 4% with each minute to first shock after initiation of CPR (Weaver et al. 1986).

Initial cardiac rhythm is an important predictor of outcome. VF with early defibrillation as a primary initial rhythm is associated with more favourable outcome than other rhythms (Silfvast 1990, Cummins et al. 1991). Patients with ventricular tachycardia (VT) or VF have several times better outcome than with PEA or asystole (ASY) (Herlitz et al. 1999). Aetiology of cardiac arrest does play a role in outcome. Cardiac arrest due to presumed cardiac origin is associated with over three times better outcome compared to non-cardiac origin cardiac arrest (Pell et al. 2003). The rescuer performance has an impact on survival after OHCA. In a retrospective observational study conducted in the United Kingdom, the experience of the ambulance crew and the level of their training influenced outcome after OHCA (Soo et al. 1999). High socioeconomic status is associated with a 1.6 fold increase in survival rate after VF, after
adjustment for other factors (e.g. age, time from call to paramedic arrival, activity, location, witnessed collapse, bystander CPR, and chronic morbidity) (Hallstrom et al. 1993). In the last part of the chain of survival is in-hospital care of patients resuscitated from OHCA. In Sweden and Norway the outcome after OHCA varies between different hospitals. Optimised in-hospital factors are associated with improved outcome after OHCA (Engdahl et al. 2000, Langhelle et al. 2003).

VF has the best prognosis, but the incidence of VF has been declining 50 % from 50-83 % incidence (Weaver et al. 1986, Bayes de Luna et al. 1989) in OHCA during the last two decades. At the same time, there has been an increase in the number of patients with PEA as initial cardiac rhythm. The proportion of these patients increased by more than 60 % during the 1980s and 1990s (Herlitz et al. 2000, Kuisma et al. 2001, Cobb et al. 2002) and the factors behind this phenomenon are unclear.
3. Regurgitation and aspiration in cardiac arrest

After the genesis of modern CPR in the early 1960s, the first reports of complications were reported in the mid-1960s. Attention was directed to multiple rib fractures, haemothorax, pneumothorax, contusion and laceration of lungs, and fat and bone-marrow embolism. In the early years of resuscitation outside the operating theatre, vomitus and subsequent aspiration was often seen because of lack of reflexes protecting the airway during resuscitation (Greenberg 1967).

Regurgitation is defined as a passive flow of gastric contents to the pharynx. Aspiration is defined as aspiration of gastric contents into the lungs, as a subsequent phenomenon to the regurgitation. This complication has been reported several times during the last three decades during resuscitation in OHCA patients. A study where rescue units detected and treated a VF during resuscitation was reported to have an 11% incidence of aspiration (Liberthson et al. 1974). A prospective autopsy study of 705 cases concerning complications after unsuccessful resuscitation reported an incidence of oropharyngeal vomitus in 10.1% and tracheal vomitus in 9.4% (Krischer et al. 1987). A group of patients who were resuscitated after OHCA and who died within 24h after admission were studied retrospectively to determine the incidence of pulmonary aspiration. The incidence of aspiration of gastric contents or blood in autopsy material after CPR was found to be 29%. No details are available on the mode or duration of ventilation prior to the intubation in this study. The aetiology of cardiac arrest within these patients included cardiac and non-cardiac causes. The authors stated that the incidence of pulmonary aspiration (29%) may underestimate the problem, because 46% of the patients studied had full stomachs (Lawes et al. 1987). In Vienna, a prospective observational study was conducted to discover whether bystander CPR increases mouth-to-mouth ventilation related complications and adverse effects caused by chest compressions. The chest radiographs on admission were studied from patients surviving cardiac arrest, and no difference was found between bystander CPR group and no bystander CPR group. A 17–18% incidence of severe gastric insufflation after CPR was revealed (Oschatz et al. 2001). In a Swedish study the experiences of bystanders were studied shortly after performing CPR. The rescuers most frequently had problems regarding the patient with mouth-to-mouth ventilation (20%) and vomiting (18%) (Axelsson et al. 1996). An in-hospital study compared the incidence of gastric regurgitation between the bag valve mask and laryngeal mask airway (Stone et al. 1998). The details of gastric regurgitation were prospectively recorded from 797 patients. Regurgitation occurred at some stage of resuscitation in 180 (23%) patients.
3.1 The role of ventilation in cardiopulmonary resuscitation

Since the beginning of the modern CPR in the 1960’s, mouth-to-mouth ventilation and subsequent assisted ventilation and intubation of the trachea have been the cornerstones of CPR.

The benefits of ventilation during respiratory arrest were already demonstrated in the 1950’s (Elam et al. 1958, Safar et al. 1958). Elam and Safar showed the feasibility of direct mouth-to-mouth ventilation by a layperson on curarised patients and that exhaled air is a resuscitative gas. The “victims” of respiratory arrest were healthy volunteers with normal haemodynamics.

The international guidelines on the role of ventilation remained virtually unchanged from 1966 to 1986. However, as cumulating data showed an increased likelihood of gastric inflation and subsequent pulmonary aspiration, new recommendations for ventilation in CPR were introduced (Melker 1985).

The assisted ventilation manoeuvres without a secured airway are also often linked with regurgitation and increased morbidity and mortality (Pepe 1996). In the delivery of artificial ventilation with “bag and mask” the pressure in the hypopharynx may exceed 25 H2O cm, a pressure causing the opening of the gastro-oesophageal sphincter in most patients (Ruben et al. 1961). Gastric insufflation and subsequent regurgitation of gastric contents with aspiration usually follows (McIntyre et al. 1978). The role of the lower oesophageal sphincter (LES) in regurgitation is crucial but there is little evidence on its pressure and function during resuscitation. In a laboratory trial using domestic swine, a rapid and severe decrease in LES tone was demonstrated during prolonged cardiac arrest. The LES tone decreased from mean baseline 21 cm H2O to mean 3.3 cm H2O during seven minutes of cardiac arrest. When ROSC occurred after defibrillation the LES tone was then measured for a further seven minutes. It increased rapidly but only to half of the prearrest baseline. Unfortunately the impact of CPR on LES tone was not studied during CPR (Bowman et al. 1995).

The need for initial mouth-to-mouth ventilation and subsequent assisted ventilation has been challenged in CPR (Berg et al. 1993, Van Hoeyweghen et al. 1993, Hallstrom et al. 2000, Berg et al. 2001). CPR performed with 15 chest compressions (at a rate of 100/min) and 2 rescue breathings compared to continuous chest compressions at the same rate showed a compromised effect on haemodynamics in ventilated swine (Berg et al. 2001). Since ventilation has been considered an essential part of CPR, the impact on the survival of the cardiac arrest patient should be positive. In a clinical study, the survival was better when bystanders performed chest compression only i.e. cardiac-only resuscitation, instead of conventional CPR (Hallstrom et al. 2000). The existing guidelines indicate that chest compression only CPR should be performed only if bystander is unwilling or unable to give mouth-to-mouth ventilation (Handley et al. 2005).

There are several reasons why bystander mouth-to-mouth ventilation may not be conductive to survival. The first obstacle in survival from OHCA is lack of
bystander CPR. The need for mouth-to-mouth ventilation greatly reduces the willingness to initiate bystander CPR (Ewy 2005) and in a swine model of bystander resuscitation a prompt initiation of chest compressions alone was as effective as chest compressions plus ventilation (Berg et al. 1993). When mouth-to-mouth ventilation cannot be applied, chest compression only CPR is better than no CPR at all with respect to outcome (Van Hoeyweghen et al. 1993). The coronary perfusion pressure is the difference between the aortic diastolic pressure and the right atrial diastolic pressure. In a clinical observation study professional rescuers were shown to ventilate OHCA patients excessively during resuscitation. In a subsequent animal study aortic, right atrial and thoracic pressure were measured during CPR. Three different ventilation rates were studied (12, 20 and 30 breaths per minute) and the ventilation was initiated during the decompression phase. These results showed that excessive ventilation rates significantly decreased coronary perfusion and survival rates. Furthermore, the venous return was shown to be reduced to the right heart in this setting. (Aufderheide et al. 2004a, Aufderheide et al. 2004b). This situation is exacerbated, if powerful ventilation is given during chest compressions, because of a further increase in intrathoracic pressure (Aufderheide et al. 2004a). There is also evidence that air on the alveolar level is equivalent to the room air when the airway is open; therefore blood in the arterial system is already oxygenated without artificial ventilation (Mithoefer et al. 1967, Meursing 1983) and enables chest compression to circulate oxygenated blood (Meursing et al. 2005).

3.2. Radiological findings of gastric aspiration

Predisposing conditions to pulmonary aspiration of gastric contents are reduced levels of consciousness, which is evident during OHCA and CPR (Adnet et al. 1996, Bartlett et al. 1975). However, it is not known how often documented gastric regurgitation during the treatment of cardiac arrest leads to radiographic findings compatible with aspiration. The consequence of the aspiration of gastric contents is known as Mendelson’s syndrome: Initially there is abrupt onset of acute respiratory distress. Bronchospasm is a characteristic feature in all patients. Chest X-ray film changes consisting of soft, irregular, mottled densities in the right lower lobe or both lower lobes, are associated with frothy nonpurulent sputum (Mendelson 1946). Hypoxia, together with normal to lowered PCO₂, indicates ventilation-perfusion disturbances (Bartlett et al. 1975).

Specific studies concerning the role of documented aspiration during resuscitation and survival after OHCA have not been conducted. In a study on the acute aspiration of gastric contents, altered state of consciousness played a role in aspiration, but not cardiac arrest. On the day of aspiration 54 out of 60 patients had abnormalities in the first chest X-ray. In the appropriate clinical setting, any radiographic infiltrates should raise the suspicion of aspiration (Landay et al. 1978). Even in well documented gastric aspiration into the lungs with tachypnea, cough, cyanosis, and wheezing, the only clinical sign may be
fever in 10 % of cases i.e. the consequences of aspiration may be minor (Bartlett et al. 1975, Landay et al. 1978).

4. Pulseless electrical activity

PEA is defined as cardiac electrical activity in the absence of any palpable pulse. There may be myocardial contractions, but they are too weak to produce pulse or blood pressure (Nolan et al. 2005). PEA is usually caused by potentially reversible causes and may be treated if these conditions are identified and corrected. A practical approach to the aetiology of PEA has been proposed by Kloeck. The mnemonic includes the 10 most likely and treatable conditions (five H’s and five T’s) that may be associated with PEA. These conditions are listed as follows: Hypoxia, Heart attack, Hypovolaemia, H+ and electrolyte abnormality, Hypothermia, Test other pulses (if carotic pulse is absent, other pulses should be palpated), Tension pneumothorax, Tamponade (cardiac tamponade), Toxins and therapeutic agents (overdosaging of therapeutic or non-therapeutic agents), Thrombo-emboli (Kloeck 1995).

Whereas it is well established that VF is associated with coronary heart disease (Baum et al. 1974), the conditions that cause PEA as initial cardiac rhythm in OHCA have been much less investigated. Pulmonary embolism (PE) has been shown to be common (36 %) in patients with unexpected cardiac arrest presenting with PEA (Comess et al. 2000), and a study on patients who died after primary PEA (previously called electromechanical dissociation, EMD) reported that cardiovascular rupture, pulmonary embolisation or tension pneumothorax was the cause of death in 19 out of 50 patients (Pirolo et al. 1985). Another study involving 28 patients with PEA as initial rhythm who died after unsuccessful resuscitation documented 11 cardiovascular ruptures and 6 PEs (Silfvast 1991). Courtney et al. suggested that PE was the COD in 52 % of the patients with primary PEA in a study on witnessed prehospital cardiac arrest (Courtney et al. 2001).

Obviously, the CODs presented above are based on autopsy findings. However, only a minority of patients who die after attempted resuscitation undergo autopsy (Silfvast 1991, de la Grandmaison et al. 2002, Vanbrabant et al. 2006). In most cases, the patient’s physician determines the COD based on clinical grounds, and only if the cause is unknown, if there are reasons to suspect a specific condition, or if it is required by law or regulations, will an autopsy be requested.
5. Post-resuscitation disease

Although in OHCA the primary resuscitation is often successful, the major obstacle to good neurological survival is post-resuscitation disease. Negovsky described post-resuscitation disease as a specific multiorgan pathophysiological state of the resuscitated cardiac arrest patient. These post-resuscitation processes do not involve only the CNS system, but also the rest of the body, and may lead to severe disability or even death after otherwise successful resuscitation. This clinical syndrome affects the cardiovascular, neurological, pulmonary, renal and metabolic systems. Such disorders in these systems are caused by marked endotoxemia washed out from ischaemia affected organs and tissues, altered haemodynamics after resuscitation, and changes in neuroendocrine profile and rheological characteristics of the blood (Negovsky 1988). There is a notable inter-relationship between the pathological processes developing in the brain and the extracerebral system. The pathological changes in the brain after successful resuscitation are discussed further in the next chapter. Survival from cardiac arrest is dependent on how rapidly CPR, defibrillation, and ALS have been initiated (Cummins et al. 1991). The quality of CPR has also been on focus in the literature. Chest compressions appear to be the most important factor in resuscitation of a human being (Van Hoeyweghen et al. 1993). The coronary perfusion pressure (CPP) is the difference between the aortic diastolic pressure and the right atrial diastolic pressure. The importance of chest compressions has been studied and an investigation established that interrupting chest compression for rescue breathing causes a 7 mmHg drop in mean CCP during CPR (Berg et al. 2001). Recently, quality of out-of-hospital CPR has been studied and the main finding was that during CPR chest compression were not delivered for half of the time. Furthermore, most compressions were too shallow (Wik et al. 2005). Skrifvars et al. demonstrated that multiple factors affect 6-month outcome following resuscitation from cardiac arrest. Strict glucose control in the Intensive Care Unit (ICU), serum potassium level and beta-blocking agents were independently associated with survival (Skrifvars et al. 2003). There are cumulating data showing that in-hospital factors are associated with outcome after OHCA (Engdahl et al. 2000, Langhelle et al. 2003).

6. Hypothermia after cardiac arrest

Interventions to mitigate neuronal injury after cardiac arrest have been studied with different approaches, but only therapeutic hypothermia has been shown to reduce mortality and morbidity. Induced hypothermia has been in use since the 1950s to protect the brain against global ischaemia during open-heart surgery. A case report of the successful use of hypothermia after cardiac arrest outside the operating theatre was published at the end of the 1950’s. Two children and two
adults were treated, and three of them recovered completely and one with moderate neurological impairment (Williams et al. 1958). One year later a study with a control group was published, where nineteen patients (including those previously reported two children and two adults) were resuscitated after cardiac arrest with resultant neurological damage. These patients were divided into a normothermia group and a hypothermia group. Survival was 14% and 50% respectively (Benson et al. 1959). The method was subsequently abandoned due to uncertain benefit and difficulties with its use. Interest in induced hypothermia after return of spontaneous circulation rose again in the 1990s and has been associated with improved functional recovery and reduced cerebral histological defects in this setting (Sterz et al. 1991). The timing of induction of therapeutic hypothermia has been shown to be critical. In a canine study with induced mild hypothermia after normothermic cardiac arrest, hypothermia improved cerebral functional and morphologic outcome. However, if the induction of cooling was delayed for 15 min after ROSC, it did not improve functional outcome, although it may have mitigated histological tissue damage (Kuboyama et al. 1993).

The exact mechanism of induced therapeutic hypothermia is not clear. A reduction of cerebral oxygen consumption has been proposed (Hegnauer et al. 1954) and other multifactorial physical and chemical mechanisms during and after low-flow induced ischaemia have also been postulated (Hypothermia after Cardiac Arrest Study Group 2002). These include reduction of intracellular acidosis (Chopp et al. 1989), reducing cerebral oedema and protection of lipoprotein membranes (Dempsey et al. 1987), inhibition of biosynthesis and release of excitatory neurotransmitters (Busto et al. 1989). ATP concentration in brain tissue has been shown to most consistently reflect biochemical activity among available biochemical indicators. Profound hypothermia has been shown to result in a three to fourfold increase in survival of cerebral ATP during circulatory arrest (Kramer et al. 1968).

Promising preliminary human data accumulated at the turn of the century. Surface cooling after OHCA maintained for 12 hours in the ICU significantly improved outcome compared to retrospective controls (Bernard et al. 1997). The use of mild hypothermia after OHCA yielded better outcome, but also more pneumonias after 48-hour hypothermia and very slow re-warming at a rate no greater than 1 °C per day (Yanagawa et al. 1998). In a pilot study of the HACA-Study group, external cooling of the head and trunk after ROSC in the emergency department was feasible and safe (Zeiner et al. 2000). Felberg et al. reported a feasibility trial where external cooling was feasible and safe. However, external cooling was slow and imprecise and efforts to speed up the start of cooling and to improve the cooling process are needed (Felberg et al. 2001).

A first out-of-hospital prospective randomized trial was conducted by Hachimi-Idrissi et al., where patients with cardiac arrest due to ASY or PEA were enrolled and randomized to a normothermic and a hypothermic group. Hypothermia was induced using a helmet device containing an aqueous glycerol solution and was found feasible, easy to use, inexpensive and effective with no
additional complications (Hachimi-Idrissi et al. 2001). Callaway et al., however, reported that application of ice to the head and neck during ongoing CPR failed to produce a significant cooling effect on cerebral or core temperatures. Furthermore, it was found to be moderately cumbersome and necessitated additional personnel in the field. The authors proposed the use of cold intravenous fluids in further studies (Callaway et al. 2002).

In February 2002 two randomised clinical trials, one in Europe (Hypothermia after Cardiac Arrest Study Group 2002) and another in Australia (Bernard et al. 2002), were reported. These studies showed a fundamental improvement in both neurological outcome and reduction of mortality in OHCA with VF as a primary rhythm. These two studies yielded similar results, thus making the important conclusions more convincing. In the HACA study the number needed to treat (NNT) for favourable neurological outcome (good recovery or moderate disability) was 6, while NNT to prevent a death was 7. In the study by Bernard et al., NNT for normal or minimal disability at the time of discharge from the hospital was 4, and NNT for avoidance of death at the same time was 6. In both studies the complication rates did not differ significantly between the hypothermia and normothermia groups. Based on these two studies, a strong recommendation was made: “Although we await further studies with great interest, we recommend the use of mild hypothermia in survivors of cardiac arrest – as early as possible and for at least 12 hours (Safar et al. 2002).” ILCOR made the following recommendations in October 2002 (Nolan et al. 2003): “Unconscious adult patients with spontaneous circulation after OHCA should be cooled to 32-34 °C for 12-24 h when the initial rhythm is VF. Such cooling may also be beneficial for other rhythms or in-hospital cardiac arrest.” A recent meta-analysis yielded a statement that one patient would leave the hospital with favourable neurological recovery by treating 4 to 13 OHCA patients with mild hypothermia (Holzer et al. 2005).

Bernard et al. conducted a pilot study using a rapid infusion of large volume (30 ml/kg), ice-cold (4 °C) intravenous fluid. The method was found to be a safe, rapid and inexpensive technique to induce mild hypothermia in OHCA patients. Regardless of quite a large and fast volume load, no patient developed pulmonary oedema (Bernard et al. 2003). The effect of large volume, ice-cold fluid intravenous infusion for the induction of moderate hypothermia on younger and older healthy volunteers has also been studied. A volume load of 40ml/kg was infused at a rate of 70-100ml/min and no pulmonary oedema was reported among patients (Frank et al. 2000, Rajek et al. 2000).

The most optimal timing of the induction of hypothermia remains uncertain. It is surprising that the clinical benefits associated with hypothermia occurred despite long delays in attaining target body temperature in the above mentioned studies (Bernard et al. 2002, Hypothermia after Cardiac Arrest Study Group 2002).
7. Techniques for the induction of hypothermia after cardiac arrest

A variety of techniques can be applied to induce hypothermia after OHCA (Table 2). In the field there are not so many feasible methods to induce therapeutic hypothermia after OHCA. Surface cooling can be done with ice packs applied to the torso and head of the patient. An approximately 0.9 °C/h cooling effect has been achieved (Bernard et al. 2003). Mild hypothermia has been induced in the field by a helmet device (Hachimi-Idrissi et al. 2001). A thympanic temperature of 34 °C was reached in 60 min. and bladder temperature in 180 min. The technique using a large volume ice-cold intravenous fluids was introduced in the previous chapter in detail. The remaining methods mentioned in Table 2 are used in the in-hospital setting and they are not currently feasible in the field.

Table 2. Different techniques for the induction of hypothermia after cardiac arrest.

<table>
<thead>
<tr>
<th>Method</th>
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<tr>
<td>Surface cooling</td>
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<tr>
<td>Large volume ice cold intravenous fluid</td>
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<td>Intravascular catheter cooling</td>
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<tr>
<td>Extracorporeal cooling</td>
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<tr>
<td>Partial liquid ventilation with cold fluorocarbons</td>
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<tr>
<td>Pharmacological approaches</td>
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<td>Isolated brain cooling</td>
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<td>Body cavity lavage</td>
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Aims of the study

The aim of this thesis was to study aetiology, treatment and outcome in OHCA by the emergency medical service (EMS) systems in Tampere District EMS and the physician-staffed Helicopter EMSs in the Helsinki and Turku areas in Southern Finland. The specific aims were the following:

1. To determine whether there is an association between bystander mouth-to-mouth ventilation and regurgitation in prehospital cardiac arrest patients (I) and to assess the association between clinical signs of prehospital regurgitation and radiological findings in resuscitated patients. (II)

2. To study the causes of death after witnessed cardiac arrest followed by pulseless electrical activity and unsuccessful out-of-hospital resuscitation; and to detect any differences between causes of death determined at autopsy and those inferred from clinical history. (III)

3. To evaluate the haemodynamic and cooling effects of infusing ice-cold Ringer’s solution in the field immediately after return of spontaneous circulation. (IV)
Material and methods

Patients and methods

Altogether 542 patients who suffered an OHCA were studied. 77 patients were excluded because of missing data on regurgitation or other parameters or because cardiac arrest was due to trauma or intoxication. The remaining patients were distributed in the area of Tampere District EMS (n=114) and the physician-staffed Helicopter EMSs in the Helsinki (n=287) and Turku areas (n=64) in Southern Finland between 13 July, 2001 and 12 July, 2003. The patients were treated according to current European Resuscitation Council resuscitation guidelines. The inclusion criteria are shown in Table 3.

<table>
<thead>
<tr>
<th>Inclusion criteria</th>
<th>Study I</th>
<th>Study II</th>
<th>Study III</th>
<th>Study IV</th>
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<tr>
<td>- All adult OHCA patients with</td>
<td>- All adult OHCA</td>
<td>- Witnessed OHCA</td>
<td>- All adult OHCA</td>
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<tr>
<td>attempted CPR</td>
<td>patients with ROSC</td>
<td>with primary PEA rhythm</td>
<td>patients with ROSC</td>
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<td></td>
<td></td>
<td>- Unsuccessful CPR</td>
<td>- ROSC ≥10 min</td>
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<td>- GCS ≤ 5/15</td>
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</table>

<table>
<thead>
<tr>
<th>Exclusion criteria*</th>
<th>Study I</th>
<th>Study II</th>
<th>Study III</th>
<th>Study IV</th>
</tr>
</thead>
<tbody>
<tr>
<td>- Trauma</td>
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<td>- Trauma</td>
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<td>- Drug overdose</td>
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<td>- Drug overdose</td>
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<tr>
<td>- Missing chest x-ray</td>
<td>- Airway obstruction</td>
<td>- Bleeding</td>
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<td></td>
<td>- Drowning</td>
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<td>- Pregnancy</td>
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<td>- SBP &lt;90 mmHg</td>
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<td></td>
<td>not responding to therapy</td>
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<td></td>
<td></td>
<td></td>
<td>- T_ao &lt; 34 °C</td>
<td></td>
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Table 3. Inclusion and exclusion criteria of Studies I-IV. *Missing data was an exclusion criterion in all studies.

1. Regurgitation during CPR (I) and radiological findings (II)

Data from all adult OHCA patients with attempted resuscitation not due to trauma or drug overdose were collected prospectively in the Tampere, Helsinki and Turku areas. Moreover, in Study II patients with ROSC and chest X-rays obtained during the first 2 days were included. The EMS crew secured the
airways of all patients with endotracheal intubation and registered data on a separate study sheet. The EMS personnel documented the presence and nature of bystander CPR on arrival by observation and by questioning bystanders. Data were recorded as no CPR, compressions only, ventilations only, or conventional CPR. The status of the pharynx at the moment of intubation was classified as no signs of regurgitation on laryngoscopy, or clinical findings compatible with regurgitation or aspiration, i.e., gastric contents present in the pharynx or visible in or suctioned from the intubation tube. The time of regurgitation was determined based on clinical observations and interviews made on the scene by EMS personnel (before EMS arrival, after EMS arrival but before intubation, after intubation). Data on 30-day survival were later obtained from Statistics Finland.

In order to assess the radiological findings produced by regurgitation of gastric contents, and not hospital-acquired or ventilator-associated pneumonia, only the chest X-rays obtained during the first 2 days after cardiac arrest were reviewed in a retrospective manner. According to general practice, a chest x-ray is routinely obtained on admission and on the first and second days after admission, and thereafter as indicated. Radiological examinations of the patients were ordered at the discretion of the treating physicians in hospital and not dictated by the study protocol. Radiological examinations were therefore not obtained in all patients, and those patients were excluded. Together with a senior consultant in radiology at the receiving hospital (Helsinki, Turku and Tampere University Hospitals), the principal investigator (I.V.) evaluated all chest x-rays obtained during the first two days after admission.

![Flow diagram of patients in Studies I and II.](image)

One radiologist at each hospital evaluated all x-rays. The findings were classified as normal, suspicion of aspiration, or radiological signs consistent with
aspiration. The radiologist and the principal investigator were blinded to the clinical findings of regurgitation at the time of the radiological evaluation.

2. PEA in unsuccessful out-of-hospital resuscitation – the cause of death (III)

In this prospective study EMS personnel determined the initial cardiac rhythm at the beginning of the resuscitation attempt and documented the data on a separate study sheet. All patients more than 17 years of age who suffered a witnessed prehospital cardiac arrest of presumed cardiac origin with PEA as initial cardiac rhythm and in whom resuscitation was unsuccessful in the field were included.

![Flow chart of patients in Study III.](image)

According to the Finnish legislation (Act 459/73), a police inquiry must be conducted and a medicolegal autopsy performed to determine the COD: 1) when death is not known to be due to illness, or if the deceased has not been treated by a physician during the period of his/her last illness; 2) when death is caused or suspected to have been caused by a crime, accident, suicide, poisoning, occupational disease or medical treatment procedure; or, 3) in cases of otherwise unexpected death. In the event of prehospital death, the police authorities are responsible for necessary further actions and determine whether any of the above mentioned conditions apply. If a medicolegal autopsy is required, it will be performed at the local department of forensic medicine. In the remaining cases, the treating physician of the deceased will be contacted, and he decides whether sufficient data exist in the patient records to allow a death certificate to be issued without an autopsy. If considered necessary, the treating physician may also
request a routine autopsy to determine the COD before issuing the death certificate.

In this study, two strategies were designed. In the Helsinki area, the EMS physician on the scene made a request for an autopsy on the EMS run sheet in all cases where resuscitation was terminated as unsuccessful when the first recorded rhythm had been PEA, irrespective of the suspected COD. In the Tampere and Turku areas the EMS crew made no requests for autopsy on the EMS run sheets, and the COD was determined according to general practice. In these areas, unless a medicolegal autopsy was required by law, the police contacted the deceased’s treating physician. The treating physician decided whether the death certificate could be issued on the basis of previous history and clinical notes of the cardiac arrest. If not, he ordered a medical autopsy. The EMS run sheets of all patients in the study areas were collected and evaluated. Special attention was focused on any notes regarding the treating EMS crews’ observations on or suspicion of the cause of arrest. The autopsy referrals made by the treating physicians to the pathologists were also retrieved and reviewed for purposes of requesting an autopsy. Data on the COD stated on the death certificates of the patients who did not undergo an autopsy were obtained from Statistics Finland. The corresponding data of those who were autopsied were retrieved from the autopsy protocols.

3 Induction of therapeutic hypothermia (IV)

This prospective study was conducted in the Helsinki Area HEMS between 23rd April 2002 and 12th July 2003. The inclusion criteria were OHCA not due to trauma or drug overdose, age over 18 years and ROSC later than 10 minutes from the onset of cardiac arrest, and Glasgow Coma Score (GCS) ≤ 5. Exclusion criteria were pregnancy, systolic blood pressure < 90 mmHg not responding to volume or inotropes, or oesophageal temperature (TeSO2) < 34.0 °C.

After ROSC patients’ lungs were manually ventilated and end-tidal CO2 (Life-Cap; Medtronic PhysioControl, Redmond, Washington, USA) was monitored continuously to achieve normocapnoea. An arterial blood gas measurement was undertaken using the i-STAT (i-STAT Corporation, Windsor, New Jersey, USA) portable blood gas analyser with the EC6+ cartridge to obtain electrolyte values, pH, and blood gases, and to find out the possible difference between ETCO2 and arterial CO2. A TeSO2 probe was inserted and connected to the monitor. When the patient was stabilised and found eligible, informed consent was obtained from relatives.

Mild hypothermia was induced with ice-cold Ringer’s acetate. The fluids were stored in an insulated box with ice cubes to maintain + 4 °C temperature. Pressure bags were used to infuse the target volume of 30 ml/kg at a rate of 100 ml/min. TeSO2 was monitored continuously and the infusion stopped if the core temperature of 33 °C was reached or adverse haemodynamic events (i.e.
arrhythmias or hypotension) occurred before the calculated volume had been infused. Blood pressure, heart rate, SpO₂, ECG, and end tidal CO₂ were closely monitored and data was collected every five min. The haemodynamic effects were defined to be rhythm observation (especially in the case of new arrhythmia e.g. VT, VF or other rearrest) and changes in arterial blood pressure.

At the end of infusion, arterial blood gas analysis was repeated. After that, the patient was carried to the ambulance and transported to hospital with all monitoring in place. On arrival at hospital, arterial blood gases were analysed, the last temperature was recorded and the study ended. Further care in hospital was at the discretion of the treating physicians.

4. Ethical aspects

The study protocols were approved by the institutional review board of Helsinki University Hospital.
The need for informed consent from relatives was waived due to the observational nature of the studies I-III.
The HEMS physician explained the study protocol to the relatives of the patients and written informed consent was obtained before induction of hypothermia in study IV.

5. Statistical methods

Statistical calculations were made using the SPSS versions 9, 11, 12 or 15 (SPSS Inc, Chicago, IL, USA). In Study I the association between CPR and clinical findings of regurgitation was analysed with the Pearson Chi square test, where a p-value <0.05 was considered significant. Also, Odds Ratios (OR) for achieving ROSC and 30-day survival if the patient regurgitated were calculated with a confidence interval of 95%. In Study II the association between clinical signs of regurgitation of gastric contents or pulmonary aspiration documented at the time of intubation and radiological signs consistent with pulmonary aspiration was analysed with the Chi-square test. The null hypothesis was that no such association exists. Inter-group differences in demographics, rhythm and ROSC data were analysed with the Kruskal-Wallis test, followed by the Mann-Whitney independent sample test. In addition, the sensitivity, specificity and positive (PPV) and negative (NPV) predictive values of clinical regurgitation to predict radiological signs consistent with pulmonary aspiration were calculated. In Study III the association between determination the COD in the clinical history group and the autopsied group was analysed with the Pearson Chi-square test and Fisher's exact test, where appropriate. The null hypothesis was that no such association exists. Analysis of variance and t-tests were used in Study IV where appropriate.
Due to the nature of the Studies I-IV, no preliminary sample size calculations were performed.
A p-value <0.05 was considered significant.
Results

Regurgitation and bystander CPR

To determine the relation between regurgitation and bystander CPR 452 patients with complete data were included in this study. In 62 of these patients, cardiac arrest was witnessed by EMS personnel. The data of those 62 patients were not analysed further because of absence of bystander CPR. Thus, the data of altogether 390 patients who suffered a cardiac arrest before EMS arrival were subjected to further analysis. Their mean age was 63.7 ± 16.6 years, and 71.5 % were males. The EMS crew reached 64 % of the patients in less than 10 minutes from the beginning of the emergency phonecall. The incidence of regurgitation was 28.9 %. In the majority of these patients (83 %), regurgitation had occurred before the arrival of the EMS personnel, whereas the remaining of the patients who regurgitated did so in the presence of the professional care providers. 13 patients regurgitated before and 5 patients after intubation. Patients in the no bystander CPR- group were significantly older and were reached earlier than the two other groups. Bystander CPR including mouth-to-mouth ventilation was associated with a significantly increased risk of regurgitation compared with no CPR (p<0.013) or compressions only CPR (p<0.01) (Table 4). ROSC was achieved in 162 patients who were all subsequently admitted to hospital.

The OR for achieving ROSC was 0.82 (95% CI 0.52-1.28, p=0.37) and that for death within 30 days was 1.79 (95% CI 0.91-3.51, p=0.08) if the patient regurgitated during cardiac arrest.

Table 4. Patients who suffered cardiac arrest before arrival of EMS and incidence of regurgitation in relation to mode of CPR.

<table>
<thead>
<tr>
<th></th>
<th>N</th>
<th>regurgitated</th>
<th>% regurgitated</th>
</tr>
</thead>
<tbody>
<tr>
<td>No bystander CPR</td>
<td>220</td>
<td>54</td>
<td>25</td>
</tr>
<tr>
<td>Conventional CPR or ventilation only</td>
<td>127</td>
<td>50</td>
<td>39</td>
</tr>
<tr>
<td>Compressions only CPR</td>
<td>43</td>
<td>9</td>
<td>21</td>
</tr>
<tr>
<td>Total</td>
<td>390</td>
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</table>
Aspiration and radiological findings in pre-hospital cardiac arrest

Radiological findings after out-of-hospital regurgitation during CPR were studied. Resuscitation was successful in 202 patients, and they were admitted to hospital. No radiological examinations were performed on 20 patients, leaving 182 patients for further analysis. Their mean age was 62.4 ± 15.5 (mean ± Standard Deviation (SD)) years and 74 % were males. ROSC had been achieved within 21.6 ± 12.8 (mean ± SD) min. Altogether 256 chest x-rays were available for further analysis, one for 108 patients and two for 74 patients. The timing of the first chest x-ray was the day of admission in 36%, the day after admission in 60 % and the second day after admission in 4 % of the patients.

In 20 % of the patients (n=37), EMS personnel documented signs of regurgitation on the scene. In hospital, the chest x-ray showed suspicion of or findings consistent with aspiration in 24 % of patients (n=44). The chest x-ray revealed findings compatible with pulmonary aspiration in 46 % of patients with clinical signs of regurgitation on the scene compared with 19 % of patients without such findings. Thus, clinical signs of regurgitation in the prehospital phase resulted in radiological signs consistent with pulmonary aspiration with 81 % specificity and 46 % sensitivity. The PPV was thus 0.39 and NPV 0.86.

Cause of death in resuscitation with PEA

To determine the COD in patients who died after an unsuccessful attempt at out-of-hospital resuscitation with PEA as a primary rhythm, 91 patients were included during the study period, 31 in the Helsinki area and 60 in the Tampere and Turku areas. The mean age (± SD) of the patients was 73.5 ± 11.9 years, and 62 % were males. Cardiac arrest occurred in the presence of the EMS crew in 24 % of the patients; in the remaining patients the arrest was witnessed by a bystander. An autopsy was performed on 59 patients (65 %), in 81 % of those in the Helsinki area and in 57% of those in the Tampere and Turku areas.

Non-cardiac CODs were diagnosed almost entirely in autopsy and there were significant difference between diagnoses in the cardiac COD and non-cardiac COD groups (Table 5). The distribution of diagnoses was significantly different between the patients whose cause of death was determined by autopsy compared with those whose cause of death was determined on clinical grounds (Table 6).

When the COD was determined based on the clinical course of the resuscitation and previous clinical history of the deceased, there were significantly more AMIs and significantly fewer PEs and aortic dissections or ruptures compared with those who underwent autopsy. There were no differences between these two groups in intracranial haemorrhage (ICH), ischaemic coronary disease or in the other COD group.
There was a suspicion of a specific cause for the arrest mentioned on the EMS run sheet or on the referral from the treating physician in only 6 of the deceased patients. In five of these six, the clinical suspicions appeared to be correct. Due to protocol violation, the treating physician determined the CODs of six patients in Helsinki and their CODs were determined based on medical history and the course of the resuscitation attempt. These CODs (in the treating physicians’ opinion) were AMI in 4 patients and chronic obstructive pulmonary disease (COPD) in 2 patients.

Table 5. Cardiac and non-cardiac causes of death in relation to mode of determination of death.

<table>
<thead>
<tr>
<th>Cause of death</th>
<th>Autopsy Yes</th>
<th>Autopsy No</th>
<th>Total</th>
<th>p-value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cardiac</td>
<td>23 (48)</td>
<td>25 (52)</td>
<td>48</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Non-cardiac</td>
<td>36 (84)</td>
<td>7 (16)</td>
<td>43</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Total</td>
<td>59 (65)</td>
<td>32 (35)</td>
<td>91</td>
<td></td>
</tr>
</tbody>
</table>

* A p-value less than 0.05 was considered significant with Chi-square test and Fisher's exact test

Table 6. Cause of death in relation to mode of determination of death.

<table>
<thead>
<tr>
<th>Cause of death</th>
<th>Autopsy Yes</th>
<th>Autopsy No</th>
<th>Total</th>
<th>p-value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute myocardial infarction</td>
<td>11</td>
<td>19</td>
<td>30</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Aortic dissection or rupture</td>
<td>17</td>
<td>1</td>
<td>18</td>
<td>0.002</td>
</tr>
<tr>
<td>Pulmonary embolism</td>
<td>15</td>
<td>0</td>
<td>15</td>
<td>0.001</td>
</tr>
<tr>
<td>Coronary heart disease</td>
<td>8</td>
<td>4</td>
<td>12</td>
<td>1.000</td>
</tr>
<tr>
<td>Other cardiac cause</td>
<td>4</td>
<td>2</td>
<td>6</td>
<td>1.000</td>
</tr>
<tr>
<td>Intracranial haemorrhage</td>
<td>2</td>
<td>1</td>
<td>3</td>
<td>1.000</td>
</tr>
<tr>
<td>Other cause</td>
<td>2</td>
<td>5</td>
<td>7</td>
<td>0.092</td>
</tr>
<tr>
<td>Total</td>
<td>59</td>
<td>32</td>
<td>91</td>
<td></td>
</tr>
</tbody>
</table>

Other cardiac causes include two congestive heart failure, undefined pericardial disease, rheumatic valvular disease (mitral and aortic valve), undefined pulmonary hypertension, aortic valve stenosis.

Other causes include two chronic obstructive pulmonary disease, motoneuron disease, cancer, pneumonia and unexplained sudden death.

* A p-value less than 0.05 was considered significant with Chi-square test and Fisher's exact test.
Induction of therapeutic hypothermia after OHCA

The course and efficiency of hypothermia induction, and the demographics of the subjects are shown in Table 7. The mean age of patients was 60.8±12.5 years. The gender distribution was 62% males vs. 38% females. ROSC was achieved at 26±10 min. Initial cardiac rhythm was VF (53.8%), ASY (30.8%) or PEA (15.4%). Mean core temperature decreased from 35.8 ± 0.9 °C at the start of infusion to 34.0 ± 1.2 °C on arrival at hospital. The mean infused volume was 2188 ± 754 ml. Infusion started at 27 ± 12 min after ROSC and mean duration of infusion was 25 ± 11 min. One patient (Patient 2) experienced a transient drop in blood pressure. It responded well to inotrope therapy and there was no obvious cause for it. No other adverse changes in haemodynamics were observed.

Therapeutic hypothermia was continued in 5 patients for 24 hours in the ICU according to the same protocol as in the HACA study.

Table 7. Effect of hypothermia induction and demographics.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Sex</th>
<th>Initial rhythm</th>
<th>Age (years)</th>
<th>Permanent ROSC (min)</th>
<th>T before induction</th>
<th>T on admission</th>
<th>ΔT °C</th>
<th>Infused amount (ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>male</td>
<td>Asystole</td>
<td>63</td>
<td>22</td>
<td>35.1</td>
<td>33</td>
<td>-2.1</td>
<td>2500</td>
</tr>
<tr>
<td>2</td>
<td>female</td>
<td>VF</td>
<td>72</td>
<td>21</td>
<td>35.9</td>
<td>33,4</td>
<td>-2.5</td>
<td>2450</td>
</tr>
<tr>
<td>3</td>
<td>female</td>
<td>Asystole</td>
<td>53</td>
<td>23</td>
<td>35.8</td>
<td>33,1</td>
<td>-2.7</td>
<td>1800</td>
</tr>
<tr>
<td>4</td>
<td>male</td>
<td>VF</td>
<td>55</td>
<td>27</td>
<td>35.0</td>
<td>34,7</td>
<td>-0.3</td>
<td>500</td>
</tr>
<tr>
<td>5</td>
<td>male</td>
<td>VF</td>
<td>79</td>
<td>32</td>
<td>35.1</td>
<td>33,2</td>
<td>-1.9</td>
<td>2000</td>
</tr>
<tr>
<td>6</td>
<td>female</td>
<td>PEA</td>
<td>79</td>
<td>42</td>
<td>35.8</td>
<td>32,7</td>
<td>-3.1</td>
<td>2200</td>
</tr>
<tr>
<td>7</td>
<td>male</td>
<td>Asystole</td>
<td>64</td>
<td>22</td>
<td>37.2</td>
<td>35,3</td>
<td>-1.9</td>
<td>3000</td>
</tr>
<tr>
<td>8</td>
<td>male</td>
<td>VF</td>
<td>41</td>
<td>46</td>
<td>36.1</td>
<td>33</td>
<td>-3.1</td>
<td>3000</td>
</tr>
<tr>
<td>9</td>
<td>female</td>
<td>Asystole</td>
<td>52</td>
<td>19</td>
<td>34.2</td>
<td>32,7</td>
<td>-1.5</td>
<td>1000</td>
</tr>
<tr>
<td>10</td>
<td>male</td>
<td>VF</td>
<td>65</td>
<td>13</td>
<td>36.0</td>
<td>36,4</td>
<td>0.4</td>
<td>2000</td>
</tr>
<tr>
<td>11</td>
<td>male</td>
<td>VF</td>
<td>53</td>
<td>15</td>
<td>35.9</td>
<td>34,3</td>
<td>-1.6</td>
<td>2500</td>
</tr>
<tr>
<td>12</td>
<td>male</td>
<td>VF</td>
<td>43</td>
<td>26</td>
<td>36.9</td>
<td>35,3</td>
<td>-1.6</td>
<td>2500</td>
</tr>
<tr>
<td>13</td>
<td>female</td>
<td>PEA</td>
<td>71</td>
<td>29</td>
<td>36.7</td>
<td>34,3</td>
<td>-2.4</td>
<td>3000</td>
</tr>
</tbody>
</table>

VF = ventricular fibrillation, PEA = pulseless electrical activity, ROSC = return of spontaneous circulation
T = oesophageal temperature, ΔT °C = change of oesophageal temperature
Discussion

It was found in Study I that more than a fourth of patients who suffered prehospital cardiac arrest expressed clinical signs of regurgitation at some point during the prehospital phase. This concurs with previous findings (Stone et al. 1998). Although the bystanders were not asked at which point after cardiac arrest regurgitation occurred, it seemed that bystander CPR including mouth-to-mouth ventilation increased the risk of aspiration of gastric contents. In most patients, regurgitation had occurred already before the arrival of the EMS crew, soon after cardiac arrest. This finding was confirmed later in a recent study, where paramedics determined the presence and timing of emesis in the field. The incidence of regurgitation was 32 %, and in a majority of these patients (66 %), regurgitation had occurred before the arrival of EMS personnel (Simons et al. 2007). In the same study the patients who received bystander CPR expressed emesis more frequently than those who did not receive bystander CPR. Of special interest in this context is whether the victim received bystander ventilation during CPR, because it seems to be associated with increased risk of regurgitating gastric contents during resuscitation. Unfortunately Simons et al. did not report detailed data on the mode of bystander CPR (ventilation only, chest compression only or conventional CPR) which would have shed more light on the impact of bystander ventilation on regurgitation of gastric contents during resuscitation.

Thirteen regurgitated before intubation when EMS personnel were present and five patients regurgitated after intubation. In the former situation there is a possibility to prevent the regurgitation and subsequent aspiration into the lungs with the use of the Sellick manoeuvre in the intubation of the patient. In the latter situation the regurgitation should be quite well tolerated if the cuff of the intubation tube is correctly inflated and the intubation tube is correctly placed into the trachea.

A possible explanation for increased incidence of regurgitation is that regurgitation may have been provoked by air being ventilated into the stomach in the patients who received conventional bystander CPR or ventilation only, causing gastric distension. The risk of regurgitation may be further increased due to loss of tone of the lower oesophageal sphincter. This has been found to occur in experimental cardiac arrest, where a rapid and marked decrease in lower oesophageal sphincter (LES) tone was shown to occur within a few minutes after onset of cardiac arrest (Bowman et al. 1995). This could also explain why the patients who suffered an EMS crew witnessed cardiac arrest regurgitated less
frequently because of rapid response to cardiac arrest and securing the airway before the loss of tone of LES.

The current resuscitation guidelines discuss the risk of gastric distension and possible regurgitation during mouth-to-mouth ventilation in cardiac arrest, but little information has been provided on the incidence of regurgitation or advice on how to act in case of regurgitation. So far, conventional CPR including compressions and ventilations has been the general recommendation in resuscitation guidelines (Anonymous 2000). However, compressions-only CPR has been shown to be associated with similar outcome to that of conventional CPR in patients with a short delay from the onset of cardiac arrest to the arrival of the EMS crew (Hallstrom et al. 2000). Although compressions-only CPR currently is recommended only in dispatcher assisted CPR or if the lay rescuer is unwilling to perform mouth-to-mouth ventilation (Anonymous 2000), it seems that more consideration should be given to this option if the dispatcher estimates that qualified help will arrive within minutes.

New data on the mode of bystander CPR have been published in Japan, where a study reported that cardiac-only resuscitation performed by a bystander is the preferable approach to the resuscitation of adult patients (SOS-Kanto study group 2007). The technical reasons support cardiac-only resuscitation because of interruptions to chest compressions during ventilation and subsequent interruption of cerebral blood flow (Assar et al. 2000, Ewy 2007). Unfortunately this study was also lacking the information about the regurgitation of gastric contents and bystander performed ventilation. An editorial was published in the same journal, where an opinion that the current guidelines regarding bystander mouth-to-mouth ventilation should be changed without delay in light of new evidence published (Ewy 2007), but the European Resuscitation Council immediately responded by retaining current guidelines. The study by the SOS-Kanto study group was undertaken in 2002-2003, that is, under the previous version of the international resuscitation guidelines (Zideman et al. 2007). The guidelines will be revised after an international review of resuscitation science in 2010.

Study II showed an association between clinical regurgitation documented during OHCA and early radiological findings consistent with aspiration. The incidence of clinically recognised regurgitation of gastric contents among the patients admitted to hospital after ROSC in this study (20 %) was close to the incidence of aspiration reported in an autopsy material (Lawes et al. 1987) showing findings of blood or gastric contents in the airways in 29 % of subjects. Lawes et al. studied the incidence of pulmonary aspiration in patients admitted to hospital after resuscitation from OHCA. The aetiology of OHCA included both cardiac and non-cardiac causes. The authors stated that the incidence of pulmonary aspiration (29 %) may underestimate the problem, because 46 % of patients studied had full stomachs. It seems that not all patients with full stomach always regurgitate during CPR.
Oschatz et al. found in a prospective observational study that bystander CPR did not increase the mouth-to-mouth ventilation related complications and adverse effects caused by chest compressions. A 17–18% incidence of severe gastric insufflation after CPR was revealed (Oschatz et al. 2001). In a Swedish study the experiences of bystanders were studied shortly after performing CPR. The rescuers most frequently had problems with patient’s vomiting (18%) and mouth-to-mouth ventilation (20 %) (Axelsson et al. 1996)

Despite the strong association between clinical signs of regurgitation and the subsequent development of radiological findings of aspiration, the sensitivity was relatively poor, 46 %. When regurgitation was detected on the scene there was a 39 % (PPV) chance of finding radiological signs consistent with pulmonary aspiration. When there were no signs of regurgitation there was an 86 % chance that no radiological signs consistent with pulmonary aspiration would develop (NPV). To what extent the early development of radiological signs of aspiration should be interpreted as infection has not been studied. Previous work reporting infectious complications after cardiac arrest has documented that radiological signs of pneumonia develop after 7 days (Bartlett et al. 1975), and that progressive deterioration of radiographic infiltrates after 3 days suggests secondary bacterial pneumonia, acute respiratory distress syndrome (ARDS) or PE (Landay et al. 1978), but the role of aspiration was not specifically addressed in these studies. Because the subsequent development of infectious complications was not the aim of this study, we only evaluated x-rays obtained on the first and second days after admission.

No radiological signs of aspiration appeared in 54 % of those who regurgitated. One explanation evinced has been the pH of the aspirate, since it has been shown that only when the pH of the aspirated matter is less than 2.4 does it cause acidic pneumonitis, manifesting as radiological signs of aspiration (Rello et al. 1995). The nature of the regurgitated matter was not characterised. Therefore, we do not know whether it was liquid or food, an issue that may affect the clinical consequences.

Also, in other documented cases of aspiration of gastric contents, 8-10 % of the patients developed no other symptoms than fever, and the radiographic findings were extremely variable (Landay et al. 1978). Besides the pH, another factor influencing the development of radiological signs of aspiration is probably the volume of the aspirate. Thus, the overall incidence of radiological findings consistent with aspiration in this study is close to the 18 % reported by Oschatz et al. in their survey (Oschatz et al. 2001).

In the majority of patients, only one chest X-ray was obtained during the first 2 days after cardiac arrest. This was because the study protocol did not dictate the timing of radiological examinations, and there were no strict protocols to guide such examinations after cardiac arrest in the participating hospitals. Considering that all hospitals were university teaching hospitals, the small number of radiological examinations was surprising. One explanation for the lack of X-ray examinations may be that the physicians in charge have a particularly pessimistic view with regard to the prognosis of resuscitated
patients. Because two chest X-rays were not obtained in all patients, the true incidence of radiological findings may be greater than documented here. As we evaluated only the chest X-rays obtained during the first 2 days, it is possible that regurgitation-related alterations may have manifested later.

It is well known from earlier studies that VF as an initial rhythm is associated with coronary heart disease (Baum et al. 1974). This has also been documented with an autopsy study (Soo et al. 2001). It has also been shown that PEA is associated with primary non-coronary causes of cardiac arrest (Hess et al. 2007), but systematic efforts to document the causes behind primary PEA in SCA have not been undertaken. In a former Finnish study, the COD was verified by autopsy in 39 % of patients (Silfvast 1991). At autopsy, coronary heart disease was considered to be the COD in 78 % of patients with VF. The COD in patients presenting with PEA was massive PE in 21 % of patients, and cardiovascular rupture in 39 % of patients who underwent autopsy. Recent studies have shown the incidence of primary PEA to be rising in OHCA (Engdahl et al. 2001, Wik et al. 2005, Hallstrom et al. 2006). Incidence of primary PEA has been as high as 32 – 52 % in in-hospital cardiac arrests (Gajic et al. 2004, Nadkarni et al. 2006).

Study III revealed a striking difference between the causes of death determined by autopsy compared with those based on the opinion of the treating physician in patients who died after unsuccessful prehospital resuscitation after witnessed prehospital cardiac arrest with PEA as an initial rhythm. This difference does not seem to be due to selection bias for autopsy. Determination of the correct cause of death is essential for reliable mortality statistics, to guide research and, in the future, treatment efforts, and potentially also guidelines in preventing deep venous thrombosis. Patients who suffer a SCA in the prehospital setting are a very challenging group to diagnose. That is because frequently the final event is rapid and very limited information is available in the scene. The tool for reliable determination of the cause of death is autopsy.

However, the number of autopsies performed in hospitalized patients has been decreasing in recent decades (Marwick 1995, Hull et al. 2007), and the number of autopsies performed on patients who die outside hospital has been reported to be 26-39 % (Silfvast 1991, Vanbrabant et al. 2006). In a large number of patients, the COD is a clinical estimation and true CODs may be missed.

In light of the present findings, it seems that an autopsy should be performed on patients who die after unsuccessful resuscitation from primary PEA. If not, large number of patients with aortic rupture (dissection or rupture of an aortic aneurysm) and PE will probably be misdiagnosed to have died from coronary heart disease (Table 3). Greater awareness of the association between primary PEA and PE may also have clinical implications, because thrombolysis during CPR has resulted in survival and good neurological outcome (Bottiger et al. 2001). There are data on thrombolysis during CPR due to massive PE and good neurological survival when prolonged resuscitation is combined with therapeutic hypothermia as post-resuscitation treatment (Bartels et al. 2007).
The environment is demanding in the prehospital HEMS system for keeping a fluid at +4°C 24 hours per day, when there is no continuous electricity available to run a refrigerator. In summer, the temperature may easily rise to +40°C in a grounded helicopter or an emergency vehicle. Both vehicles need their own cool storage. Ringer’s acetate has to be cold (+4°C) to induce hypothermia when cold fluids are infused intravenously. Unfortunately there is no simple and inexpensive method to achieve this goal. We ended up using common modified insulated transport boxes with ice cubes made by an ice cube machine. The fluids were placed in the box with ice cubes, so when there were any ice and water left the box the temperature of the Ringer’s acetate would be 0°C. This method is simple, inexpensive and reliable, but needs daily maintaining i.e. removing melted water and adding new ice cubes to the box.

A strong recommendation to initiate therapeutic hypothermia as soon as possible after successful resuscitation after OHCA was made at the same time when two large randomised controlled studies was published (Hypothermia after Cardiac Arrest Study Group 2002, Safar et al. 2002, Bernard et al. 2003). The results of Study IV suggest that induction of therapeutic hypothermia with rapid infusion of ice-cold Ringer’s solution soon after ROSC is well tolerated and already feasible in the prehospital setting. An unconscious adult patient after ROSC with OHCA should be cooled to 32-34°C. Induction of hypothermia should be initiated as soon as possible after ROSC according to the international guidelines on resuscitation. The international resuscitation guidelines recommend the use of 30mg kg⁻¹ of 4°C-saline to initiate cooling after cardiac arrest (Nolan et al. 2005).

Later 2 litres of 4°C normal saline was used to induce hypothermia in hospitalised patients and it was also found safe and effective in lowering body temperature with hospitalised patients after OHCA (Kim et al. 2005). A larger prospective interventional study showed that induction of hypothermia with ice-cold intravenous fluids combined with ice-water cooling blanket was safe, efficacious and quick. In this study the patients had various neurological injuries, postanoxic encephalopathy, subarachnoid haemorrhage, or traumatic brain injury (Polderman et al. 2005). After these studies therapeutic hypothermia after cardiac arrest was recommended as a standard for care (Bernard 2006). In 2007 Kim et al. published a prehospital study of 125 patients who were randomised to receive standard care with or without intravenous cooling (Kim et al. 2007). Sixty three patients were randomised to cooling, and their temperature decreased 1.24 ± 1°C while 62 patients in the normothermia group had no alteration in their temperature. The study was unfortunately only prehospital and the protocol was not continued in the hospital. In the outcome there was only a trend to better outcome in the patients resuscitated from VF.

In a recent article induced hypothermia was shown to be underused after resuscitation from OHCA (Abella et al. 2005). In that study, 87% of physicians practising with patients resuscitated from OHCA with VF in United States had never used hypothermia as a postresuscitation treatment. Merchant et al. found
that hypothermia was used as a postresuscitation treatment after OHCA with VF by only 26-36% of physicians practising in units treating resuscitated cardiac arrest patients in the UK, Finland and the United States (Merchant et al. 2006).

The rather low survival rate in Study IV, although not the study endpoint, may be explained by the large number of patients with known factors for poor outcome (Kuisma et al. 1997). There were four patients with non-coronary causes of arrest, and ROSC times were long. Therefore, the infusion of cold saline was unlikely to contribute to the number of non-survivors (Hallstrom et al. 2000). On the other hand in study IV, 57% of patients survived from OHCA with VF, which is well in line with a recent study where standardised treatment protocols for post resuscitation care after OHCA were implemented (Sunde et al. 2007). The current resuscitation guidelines also suggest that OHCA patients with non-shockable rhythm may also benefit from induced hypothermia after cardiac arrest (Nolan et al. 2005). Post-resuscitation care has been proposed to be the missing link in the chain of the survival of cardiac arrest patient (Peberdy et al. 2005). A fundamental effect of the implementation of standardised post-resuscitation care protocols to guide post-resuscitation care and strong commitment to the protocols has shown a significantly improved outcome after OHCA (Sunde et al. 2007).

We used an oesophageal thermal probe as a fast and simple method for continuous measurement of core temperature. This method has been shown to reliably reflect core temperature with intact circulation (Lenhardt 2003, Insler 2006). There are, however, weaknesses in its use, which may explain the findings in two of our patients. The probe may come into close contact with the superior vena cava and thus register falsely low “core” temperatures as blood mixed with cold saline passes by. This may explain why the temperature of patient # 4 decreased rapidly after only 500 ml of fluid, and why the temperature rose again shortly after the infusion was stopped. The probe may also be inadvertently advanced too deep, eventually to enter the stomach, thus not reflecting correct values (e.g. patient # 10, whose temperature remained unchanged).

There are also other methods to induce hypothermia. Cold infusions are a feasible and safe way to initiate therapeutic hypothermia, but for maintenance additional cooling systems are needed (Kliegel et al. 2007). External cooling methods, ice packs and cold-air blankets, ice-water blankets and closed loop intravenous cooling catheters have been used to maintain therapeutic hypothermia (Merchant et al. 2006). Cool packs are a simple and inexpensive method to induce hypothermia, but were very unpopular with medical and nursing staff (Bernard 2006). In a Norwegian study, prehospital cooling was initiated in the field with sport ice-packs and continued with ice-water soaked towels over the torso in ICU. This method is also inexpensive and easily implemented in any system (Busch et al. 2006).

Now ice-cold intravenous fluids should be applied to induce hypothermia in survivors of OHCA (Mayer 2005). Clinical trials that compare out-of-hospital
paramedic cooling with hospital cooling are under way in Melbourne, Australia and in Seattle, WA (Bernard 2006).
Limitations of the study

The main limitation of Study I is that it is an observational one and therefore cannot produce the same level of evidence as a randomised trial. We did not include patients in whom an EMS crew-initiated attempt of CPR was considered inappropriate. Therefore we do not know the exact number of patients who actually regurgitated. Also, the percentage of patients with incomplete or missing data was 14.6 %, which was rather high. However, data collection in prehospital studies is a challenging task, and it is difficult to achieve the designed level of completeness.

The retrospective collection and variable number of chest X-rays were major weaknesses in Study II. The study protocol did not dictate the timing of radiological examinations, and there were no strict protocols to guide such examinations after cardiac arrest in the participating hospitals. It is also possible that patients regurgitated and subsequently aspirated gastric contents on the hospital ward.

Study III was a prehospital study, and the causes of death may be different in hospitalised patients. In the Helsinki area, a protocol violation resulted in the loss of 6 cases. Their death certificates were issued on the basis of clinical judgment, and their true CODs remain open. In the Turku and Tampere areas, the police authorities and physicians were unaware of the study, and their actions in dealing with these cases were supposed to reflect common practice. A selection bias is nevertheless possible. On the other hand, the striking difference in the distribution of diagnoses between those who underwent autopsy and those who did not strongly suggests that the correct diagnosis remained undetermined in a significant number of patients whose cause of death was estimated without autopsy.

The small number of patients in Study IV increases the risk of a beta error, but the lack of observed adverse effects of this treatment renders further studies warranted. Second, we did not have a control group. In a study with a control group using the same cooling method the temperature in the control group did not change (Kim et al. 2007). Third, the measurement of $T_{esof}$ is prone to errors, but the decrease in temperature was continuous and relevant in the majority of patients.
Conclusions

In conclusion, it was found that bystander CPR including mouth-to-mouth ventilation was associated with a significantly increased risk of regurgitation. In view of these findings the role of bystander ventilation during CPR in cardiac arrest needs to be further evaluated.

Regurgitation was associated with subsequent development of radiological findings consistent with aspiration; with moderate sensitivity and high specificity. PPV was 0.39 and NPV was 0.86. It seems that clinical signs of regurgitation of gastric contents and subsequent aspiration into the lungs do not always lead to the development of radiological signs consistent with pulmonary aspiration.

Aortic dissection or rupture, pulmonary embolism and acute myocardial infarction account for 73% of causes of deaths after witnessed cardiac arrest followed by pulseless electrical activity in an out-of-hospital setting. In the death certificates issued, a higher portion of diagnoses of AMI, and fewer PEs, aortic dissections, or ruptures were present among those without autopsy than in those who underwent autopsy. Therefore, in unsuccessful resuscitation from OHCA with PEA as initial rhythm an autopsy should be performed to determine the exact cause of death.

Induction of therapeutic hypothermia with peripheral infusion of ice-cold Ringer’s solution seems feasible soon after ROSC in the prehospital setting. No serious adverse haemodynamic effects occurred, and the mean core temperature decreased significantly.
Acknowledgements

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I wish to thank my mother for her encouraging attitude and belief in me, who, to my sorrow, never saw this dissertation completed and to my father for the support which made this dissertation possible.

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Pirkkala, May 2008
References


Bernard S, Buist M, Monteiro O and Smith K (2003): Induced hypothermia using large volume, ice-cold intravenous fluid in comatose survivors of


hospital cardiac arrest resuscitated by the same Emergency Medical Service and admitted to one of two hospitals over a 16-year period in the municipality of Goteborg. Resuscitation 43: 201-11.


Original publications
Bystander mouth-to-mouth ventilation and regurgitation during cardiopulmonary resuscitation

I. VIRKKUNEN 1,2, S. KUJALA 3, S. RYYNÄNEN 4, A. VUORI 5, V. PETTILÄ 6, A. YLI-HANKALA 1,4 & T. SILFVAST 2,6

Abstract. Virkkunen I, Kujala S, Ryynänen S, Vuori A, Pettilä V, Yli-Hankala A, Silfvast T (Tampere University Hospital, Tampere; Helsinki Area Emergency Medical Air Service, Vantaa; Faculty of Medicine, University of Turku, Turku; Medical School, University of Tampere, Tampere; Administration, Hospital District of Southwest Finland and Department of Anaesthesiology, Turku University Hospital, Turku; and Department of Anaesthesiology and Intensive Care Medicine, Helsinki University Hospital, Helsinki; Finland). Bystander mouth-to-mouth ventilation and regurgitation during cardiopulmonary resuscitation. J Intern Med 2006; 260: 39–42.

Objectives. To determine whether there is an association between bystander mouth-to-mouth ventilation and regurgitation in prehospital cardiac arrest patients.

Design. Prospectively conducted observational study.

Setting. Data were collected from patients treated by the emergency medical service (EMS) systems in three middle-sized or large Finnish urban communities, the Tampere District EMS and the physician-staffed Helicopter EMSs in the Helsinki and Turku areas in southern Finland.

Subjects. The study population consisted of 529 consecutive prehospital cardiac arrest patients with attempted resuscitation. Exclusion criteria were cardiac arrest due to trauma or drug overdose.

Main outcome measures. Regurgitation in prehospital cardiac arrest patients documented by EMS personnel on the scene.

Results. Regurgitation occurred in a fourth of patients. Bystander cardiopulmonary resuscitation (CPR) with mouth-to-mouth ventilation was associated with a significantly increased risk of regurgitation compared with no CPR (P < 0.013) and CPR without ventilations (P < 0.01).

Conclusions. The mode and role of bystander CPR in cardiac arrest needs to be further evaluated.

Keywords: cardiac arrest, mouth-to-mouth ventilation, prehospital, regurgitation, resuscitation.

Introduction

Gastric regurgitation and pulmonary aspiration are serious adverse events in out-of-hospital cardiac arrest. Assisted ventilation manoeuvres without a secured airway are often associated with regurgitation, leading to increased morbidity and mortality [1]. A recent study has suggested that ventilation may not be needed for several minutes after onset of cardiac arrest as outcome after cardiopulmonary resuscitation (CPR) with chest compressions only has been shown to be similar to that with conventional CPR including mouth-to-mouth ventilation [2]. Also, compression-only CPR has been shown to be better than no CPR at all [3, 4]. We reasoned that bystander CPR with mouth-to-mouth ventilation could be associated with an increased risk of regurgitation.

Methods

Data were prospectively collected in three middle-sized or large Finnish urban communities. The study population consisted of 529 consecutive prehospital cardiac arrest patients treated by the emergency medical service (EMS) system in the Tampere District...
and the physician-staffed Helicopter EMSs in the Helsinki and Turku areas in southern Finland. The institutional review board of the Helsinki University Hospital approved the study protocol and waived the need for informed consent from relatives due to the observational nature of the study. All adult patients who suffered prehospital cardiac arrest not due to trauma or drug overdose and in whom an attempt of CPR was considered appropriate were included.

The EMS crew secured the airways of all patients with endotracheal intubation and registered data on a separate study sheet. EMS personnel documented the presence and nature of bystander CPR on arrival by observation and by questioning bystanders. Data were recorded as no CPR, compressions only, ventilations only, or conventional CPR. The status of the pharynx at the moment of intubation was classified as no signs of regurgitation on laryngoscopy, or clinical findings compatible with regurgitation or aspiration, i.e. gastric contents present in the pharynx or visible in or suctioned from the intubation tube. The time of regurgitation was determined based on clinical observations and interviews made on the scene by EMS personnel (before EMS arrival, after EMS arrival but before intubation, after intubation). Data on 30-day survival was later obtained from Statistics Finland. The association between CPR and clinical findings of regurgitation was analysed with the Pearson chi-square test, where a \( P \)-value <0.05 was considered significant. Also, Odds Ratios (OR) for achieving return of spontaneous circulation (ROSC) and 30-day survival if the patient regurgitated were calculated with a confidence interval of 95%. Statistical analyses were performed using the SPSS \(^{®} \) 11.5.1 for Windows (SPSS Inc., Chicago, IL, USA).

**Results**

Of 529 patients with attempted resuscitation, 77 were excluded because of missing data on regurgitation or other parameters. Of the remaining 452 patients with complete data, 62 suffered an EMS-witnessed cardiac arrest. Their data were not analysed further because CPR and advanced life support was immediately available. Regurgitation occurred in eight of these patients. Thus, the data of altogether 390 patients who suffered a cardiac arrest before EMS arrival were left for further analysis. Their mean age was 63.7 ± 16.6 years, and 71.5% were males. The EMS crew reached 64% of the patients in <10 min from the beginning of the emergency phone call. The incidence of regurgitation was 28.2% \((n = 110)\). In the majority of these patients (83%), regurgitation had occurred before the arrival of the EMS personnel, whereas the remaining of the patients who regurgitated did so in the presence of the care providers (13 patients before and five after intubation). Demographics of the groups are shown in Table 1. Bystander CPR including mouth-to-mouth ventilation was associated with a significantly increased risk of regurgitation compared with no CPR \((P < 0.013)\) and compressions-only CPR \((P < 0.01)\) (Fig. 1). ROSC was achieved in 162 patients who all were subsequently admitted to hospital.

The OR for achieving ROSC was 0.82 (95% CI: 0.52–1.28, \( P = 0.37 \)) and that for death within 30 days was 1.79 (95% CI: 0.91–3.51, \( P = 0.08 \)) if the patient regurgitated during cardiac arrest. The outcomes of those who regurgitated and those who did not are provided in Table 2.

**Discussion**

We found that more than a fourth of patients who suffered prehospital cardiac arrest expressed clinical signs of regurgitation at some point during the prehospital phase. This agrees with previous findings [5]. Of concern is that bystander CPR with mouth-to-mouth ventilation was associated with a substantially increased risk of regurgitation. In most patients, regurgitation had occurred already before the arrival of the EMS crew, soon after cardiac arrest. Although the bystanders were not asked at which point after cardiac arrest regurgitation occurred, it seemed that bystander CPR including mouth-to-mouth ventilation increased the risk of aspiration of gastric contents. A possible explanation is that regurgitation may have been provoked by air being ventilated into the stomach in the patients who received conventional bystander CPR or ventilation only, causing gastric distension. The risk of regurgitation may be further increased due to loss of tone of the lower oesophageal sphincter. This has been found to occur in experimental cardiac arrest, where a rapid and marked decrease in lower oesophageal sphincter tone was shown to occur within a few minutes after onset of cardiac arrest [6]. This could also explain why the patients who
suffered an EMS crew witnessed cardiac arrest regurgitated less frequently.

Current resuscitation guidelines discuss the risk of gastric distension and possible regurgitation during mouth-to-mouth ventilation in cardiac arrest, but little information is provided on the incidence of regurgitation or advice on how to act should this occur. So far, conventional CPR including

---

**Table 1** Demographics of the groups

<table>
<thead>
<tr>
<th>1. No bystander CPR</th>
<th>2. Ventilations only or conventional CPR</th>
<th>3. Compressions only CPR</th>
<th>P-value&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients</td>
<td>220</td>
<td>127</td>
<td>43</td>
</tr>
<tr>
<td>Mean age (SD)</td>
<td>65.7 (16.1)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>61.6 (17.1)</td>
<td>59.4 (16.2)</td>
</tr>
<tr>
<td>Males, n (%)</td>
<td>155 (71)</td>
<td>95 (75)</td>
<td>28 (65)</td>
</tr>
<tr>
<td>Initial cardiac rhythm, n (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VF/VT</td>
<td>86 (39)</td>
<td>57 (45)</td>
<td>16 (37)</td>
</tr>
<tr>
<td>PEA</td>
<td>56 (26)</td>
<td>29 (23)</td>
<td>4 (9)</td>
</tr>
<tr>
<td>ASY</td>
<td>76 (35)</td>
<td>40 (32)</td>
<td>22 (51)</td>
</tr>
<tr>
<td>Data missing</td>
<td>2 (1)</td>
<td>1 (1)</td>
<td>1 (2)</td>
</tr>
<tr>
<td>Patient reached within 10 min, n (%)</td>
<td>159 (72)&lt;sup&gt;c&lt;/sup&gt;</td>
<td>63 (50)</td>
<td>28 (65)</td>
</tr>
<tr>
<td>Duration of EMS CPR, min (SD)</td>
<td>17.7 (21.1)</td>
<td>16.6 (12.7)</td>
<td>13.8 (8.3)</td>
</tr>
<tr>
<td>ROSC, min (SD)</td>
<td>23.8 (12.9)</td>
<td>23.5 (13.3)</td>
<td>20.6 (10.3)</td>
</tr>
<tr>
<td>ROSC, n</td>
<td>85</td>
<td>55</td>
<td>22</td>
</tr>
<tr>
<td>Regurgitated, n (%)</td>
<td>54 (25)&lt;sup&gt;d&lt;/sup&gt;</td>
<td>50 (39)&lt;sup&gt;e&lt;/sup&gt;</td>
<td>9 (21)</td>
</tr>
<tr>
<td>Time of regurgitation, n (%)</td>
<td>41 (76)</td>
<td>42 (84)</td>
<td>8 (89)</td>
</tr>
<tr>
<td>Before EMS arrival</td>
<td>41 (76)</td>
<td>42 (84)</td>
<td>8 (89)</td>
</tr>
<tr>
<td>EMS present, but before intubation</td>
<td>9 (17)</td>
<td>4 (8)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>After intubation</td>
<td>4 (7)</td>
<td>0 (0)</td>
<td>1 (11)</td>
</tr>
<tr>
<td>Data missing</td>
<td>0 (0)</td>
<td>4 (8)</td>
<td>0 (0)</td>
</tr>
</tbody>
</table>

Data are mean ± SD or absolute numbers (%). ROSC, return of spontaneous circulation; CPR, cardiopulmonary resuscitation; EMS, emergency medical service; VF, ventricular fibrillation; VT, ventricular tachycardia; PEA, pulseless electrical activity; ASY, asystole. <sup>a</sup>Kruskall–Wallis test for continuous variables, chi-square test for numeral. <sup>b</sup>P = 0.01 between groups 1 and 2 and 1 and 3 (Mann–Whitney). <sup>c</sup>P = 0.001 between groups 1 and 2 (chi-square). <sup>d</sup>P = 0.004 between groups 1 and 2 (chi-square). <sup>e</sup>P = 0.03 between groups 2 and 3 (chi-square).

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**Fig. 1** Flow diagram of 452 patients and incidence of regurgitation in relation to mode of CPR. CPR, cardiopulmonary resuscitation; EMS, emergency medical service.

compressions and ventilations has been the general recommendation in resuscitation guidelines [7]. However, compressions-only CPR has been shown to be associated with similar outcome as conventional CPR in patients with a short delay from the onset of cardiac arrest to the arrival of the EMS crew [2]. Although compressions-only CPR currently is recommended only in dispatcher-assisted CPR or if the lay rescuer is unwilling to perform mouth-to-mouth ventilation [7], it seems that stronger consideration should be given to this option if the dispatcher estimates that qualified help will arrive within minutes.

The main limitation of this study is that it is an observational one and therefore cannot produce the same level of evidence as a randomized trial. We did not include patients in whom an EMS crew-initiated attempt of CPR was considered inappropriate. Therefore we do not know the exact number of patients who actually regurgitated. Also, the percentage of patients with incomplete or missing data was 14.6%, which was rather high. However, data collection in prehospital studies is a challenging task, and it is difficult to achieve the level of completeness that is aimed at.

In conclusion, we found that bystander CPR including mouth-to-mouth ventilation was associated with a significantly increased risk of regurgitation. In the view of these findings and previous data [2], the mode and role of bystander CPR in cardiac arrest needs to be further evaluated.

Table 2 Relationship between regurgitation and achieving ROSC and regurgitation and death 30-days (data missing in one patient)

<table>
<thead>
<tr>
<th>Regurgitation</th>
<th>Yes</th>
<th>No</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ROSC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>43</td>
<td>70</td>
<td>113</td>
</tr>
<tr>
<td>No</td>
<td>119</td>
<td>158</td>
<td>277</td>
</tr>
<tr>
<td>Total</td>
<td>162</td>
<td>228</td>
<td>390</td>
</tr>
<tr>
<td></td>
<td>Death</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>100</td>
<td>12</td>
<td>112</td>
</tr>
<tr>
<td>No</td>
<td>228</td>
<td>49</td>
<td>277</td>
</tr>
<tr>
<td>Total</td>
<td>328</td>
<td>61</td>
<td>389</td>
</tr>
</tbody>
</table>

Conflict of interest statement

No conflict of interest was declared.

Acknowledgements

We would like to thank the crews of the Helsinki and Turku Area HEMS and Tampere District Emergency Medical Services for their valuable help in enrolling the study patients. This study was supported by institutional EVO grant no. 9B102 from Special State Allocation via Tampere University Hospital. Funding sources had no role in the design and conduct of the study; collection, management, analysis, and interpretation of the data; and preparation, review, or approval of the manuscript. We also thank Statistics Finland for providing data on 30-day survival.

References


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(fax: +358 3 31164363; e-mail: ilkka.virkkunen@uta.fi).
Erratum

Publication I, page 40, right column, fourth line should be:

“tion was 29.0% (n=113). In the majority of these”
Incidence of regurgitation and pulmonary aspiration of gastric contents in survivors from out-of-hospital cardiac arrest

I. Virkkunen1,2, S. Ryyinen3, S. Kujala4, A. Vuori5, A. Piironen6, J.-P. Kääriä7, V. Kahara8, V. Pettila9, A. Yli-Hankala1,3 and T. Silfvast2,9

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Background: The regurgitation of gastric contents and subsequent pulmonary aspiration remain serious adverse events in cardiac arrest and cardiopulmonary resuscitation. The aim of this study was to determine the association between clinical signs of regurgitation and radiological findings consistent with aspiration in resuscitated out-of-hospital cardiac arrest (OHCA) patients admitted to hospital.

Methods: The incidence of regurgitation was studied in 182 successfully resuscitated OHCA patients. The inclusion criterion was the restoration of spontaneous circulation after OHCA not caused by trauma or drug overdose.

Results: The incidence of regurgitation was 20%. Regurgitation was associated with radiological findings consistent with aspiration with a high specificity (81%) and a low sensitivity (46%).

Conclusions: Although there was a strong association between clinical regurgitation and radiological findings consistent with aspiration, our data suggest that regurgitation is not invariably followed by radiological findings compatible with aspiration. Radiological findings consistent with aspiration are relatively infrequent without preceding signs of regurgitation in resuscitated patients.

Accepted for publication 10 October 2006

Key words: aspiration; cardiac arrest; emergency medical services; out-of-hospital cardiopulmonary resuscitation; regurgitation; resuscitation/adverse effects.

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CARDIAC arrest and cardiopulmonary resuscitation (CPR) are associated with a risk of regurgitation (passive flow) and pulmonary aspiration of gastric contents (1–3). Previous studies have documented a 29% incidence of aspiration of gastric contents or blood in autopsy material after CPR (1), a 17–18% incidence of severe gastric insufflation after CPR (2) and an 18% incidence of regurgitation of gastric contents as a reported problem by bystanders in association with bystander CPR (3). In a previous study conducted by this group (4), we showed that regurgitation occurred in 28% of patients at some stage during pre-hospital cardiac arrest. Although this complication is associated with increased morbidity and mortality (5), it is not known how often documented gastric regurgitation during the treatment of cardiac arrest leads to radiographic findings compatible with aspiration. Therefore, we decided to determine the association between clinical signs of regurgitation and early radiological findings consistent with aspiration in patients who were successfully resuscitated from pre-hospital cardiac arrest and admitted to hospital.

Methods

The study population consisted of adult patients who were successfully resuscitated after suffering pre-hospital cardiac arrest not caused by trauma or drug overdose between August 2001 and March 2003 (4). The patients were treated by the emergency medical service (EMS) system in Tampere District and the physician-staffed helicopter EMSs in the Helsinki and Turku areas in southern Finland.
The institutional review board of the Helsinki University Hospital approved the study protocol. EMS personnel intubated all patients in the field and documented prospectively the status of the pharynx at the moment of intubation on a separate study sheet. Findings were classified as follows: no signs of regurgitation on laryngoscopy; gastric contents present in the pharynx; or gastric contents visible in or suctioned from the intubation tube. After the restoration of spontaneous circulation (ROSC) and stabilization in the field, the patients were admitted to hospital.

In order to assess the radiological findings produced by the regurgitation of gastric contents, and not hospital-acquired or ventilator-associated pneumonia, only the chest X-rays obtained during the first 2 days after cardiac arrest were reviewed. Radiological examinations of the patients were ordered at the discretion of the treating physicians in hospital and not dictated by the study protocol. For that reason, radiological examinations were not performed in all patients, and patients who did not undergo radiological examination were excluded. According to general practice, a chest X-ray is routinely obtained on admission and thereafter as indicated. Together with a senior consultant in radiology at the receiving hospital (Helsinki, Turku and Tampere University Hospitals), the principal investigator (I.V.) evaluated all chest X-rays obtained during the first 2 days after admission. One radiologist evaluated all X-rays at each hospital. The findings were classified as follows: normal; suspicion of aspiration; or radiological signs consistent with aspiration. The radiologist and principal investigator were blind to the clinical findings of regurgitation at the time of the radiological evaluation.

The association between clinical signs of regurgitation of gastric contents or pulmonary aspiration documented at the time of intubation and radiological signs consistent with pulmonary aspiration was analysed using the chi-squared test. The null hypothesis was that no such association exists. Inter-group differences in demographics, rhythm and ROSC data were analysed using the Kruskal-Wallis test, followed by the Mann-Whitney independent sample test. \( P < 0.05 \) was considered to be significant. In addition, the sensitivity, specificity and positive (PPV) and negative (NPV) predictive values of clinical regurgitation to predict radiological signs consistent with pulmonary aspiration were calculated. Statistical analyses were performed using SPSS® 11.5.1 for Windows (SPSS Inc., Chicago, IL).

## Results

During the study period, resuscitation was successful in 202 patients, and they were admitted to hospital. No radiological examinations were performed in 20 patients, leaving 182 patients for further analysis. Their mean age was \( 62.4 \pm 15.5 \) years [mean \( \pm \) standard deviation (SD)] and 74% were males. ROSC had been achieved within \( 21.6 \pm 12.8 \) min (mean \( \pm \) SD). Altogether, 256 chest X-rays were available for further analysis, one in 108 patients and two in 74 patients. The timing of the first chest X-ray was the day of admission in 36%, the day after admission in 60% and the second day after admission in 4% of patients.

In 37 patients (20%), EMS personnel documented signs of regurgitation on the scene. In hospital, the chest X-ray showed a suspicion of or findings compatible with aspiration in 44 patients (24%). Chest X-ray revealed findings compatible with pulmonary aspiration in 17 of the 37 patients (46%) with clinical signs of regurgitation on the scene, compared with 27 of the 145 patients (19%) without such findings (\( P < 0.001 \) by chi-squared test) (Table 1). Thus, clinical signs of regurgitation in the pre-hospital phase resulted in radiological signs consistent with pulmonary aspiration with an 81% specificity (118 of 145) and a 46% sensitivity (17 of 37). PPV was 0.39 and NPV was 0.86.

<table>
<thead>
<tr>
<th>Table 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Relationship between clinical observations on regurgitation of gastric contents and radiological findings compatible with the aspiration of gastric contents (( P &lt; 0.001 )). Sensitivity, 0.46; specificity, 0.81; positive predictive value (PPV), 0.39; negative predictive value (NPV), 0.86.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Regurgitation on scene</th>
<th>Yes</th>
<th>No</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Radiological findings compatible with aspiration</td>
<td>17</td>
<td>27</td>
<td>44</td>
</tr>
<tr>
<td>No</td>
<td>20</td>
<td>118</td>
<td>138</td>
</tr>
<tr>
<td>Total</td>
<td>37</td>
<td>145</td>
<td>182</td>
</tr>
</tbody>
</table>

## Discussion

This study revealed a strong association between clinical regurgitation documented during out-of-hospital cardiac arrest (OHCA) and early radiological findings consistent with aspiration. The incidence of clinically recognized regurgitation of gastric contents in the patients admitted to hospital after ROSC in this...
study (20%) was close to the incidence of aspiration reported in autopsy material (1), which showed findings of blood or gastric contents in the airways in 29% of subjects. Other studies have shown severe gastric insufflation and regurgitation of gastric contents in 17–18% of patients who received bystander CPR after suffering OHCA (2, 3).

Despite the strong association between clinical signs of regurgitation and the subsequent development of radiological findings of aspiration, the sensitivity was relatively poor: 46%. When regurgitation was detected on the scene, there was a 39% (PPV) chance of finding radiological signs consistent with pulmonary aspiration. When there were no signs of regurgitation, there was an 86% chance that no radiological signs consistent with pulmonary aspiration would develop (NPV). To what extent the early development of radiological signs of aspiration should be interpreted as infection has not been studied. A previous investigation reporting infectious complications after cardiac arrest has documented that radiological signs of pneumonia develop after 7 ± 6.2 days (6), and that progressive worsening of radiographic infiltrates after 3 days suggests secondary bacterial pneumonia, acute respiratory distress syndrome (ARDS) or pulmonary embolism (7), but the role of aspiration was not specifically addressed in these studies. Because the later development of infectious complications was not the aim of this study, we only evaluated X-rays obtained on the first and second days after admission.

No radiological signs of aspiration appeared in 26% of those who regurgitated. One explanation offered is the pH of the aspirate, as it has been shown that only when the aspirated matter has pH < 2.4 does it cause acidic pneumonitis, manifesting as radiological signs of aspiration (8). The nature of the regurgitated matter was not characterized. Therefore, we do not know whether it was liquid or food, an issue that may have an influence on the clinical consequences.

Moreover, in other well-documented cases of the aspiration of gastric contents, 8–10% of patients developed no symptoms other than fever, and the radiographic findings were extremely variable (7). In addition to the pH, another probable factor influencing the development of radiological signs of aspiration is the volume of the aspirate. Thus, the overall incidence of radiological findings consistent with aspiration in this study is close to the 18% reported by Oschatz et al. (2) in their survey.

There are some limitations in this study. The retrospective collection and variable number of chest X-rays were major weaknesses. In the majority of patients, only one chest X-ray was obtained during the first 2 days after cardiac arrest. This was because the study protocol did not dictate the timing of radiological examinations, and there were no strict protocols to guide such examinations after cardiac arrest in the participating hospitals. Considering that all hospitals were university teaching hospitals, this finding was surprising. One explanation for the lack of X-ray examinations may be that the physicians in charge have a particularly pessimistic view with regard to the prognosis of resuscitated patients. Because two chest X-rays were not obtained in all patients, the true incidence of radiological findings may be greater than documented here. As we evaluated only the chest X-rays obtained during the first 2 days, it is possible that regurgitation-related alterations may have manifested later.

In conclusion, the incidence of regurgitation during pre-hospital cardiac arrest was 20% in patients admitted to hospital after ROSC in this study. Regurgitation was associated with the subsequent development of radiological findings consistent with aspiration with moderate sensitivity and high specificity. It seems that clinical signs of regurgitation of gastric contents and subsequent aspiration into the lungs do not always lead to the development of radiological signs consistent with pulmonary aspiration.

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References


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Pulseless electrical activity and unsuccessful out-of-hospital resuscitation: What is the cause of death?

Ilkka Virkkunen, Laura Paasio, Sanna Ryynänen, Arno Vuori, Antti Sajantila, Arvi Yli-Hankala, Tom Silfvast

**Aims:** To study the cause of deaths after witnessed cardiac arrest followed by pulseless electrical activity and unsuccessful out-of-hospital resuscitation; and to detect any differences between causes of death determined at autopsy and those inferred from clinical history.

**Methods:** In this prospective observational study, data were collected from 91 individuals treated by the emergency medical services in three urban communities in southern Finland.

**Results:** Cause of death was determined at autopsy in 59 cases and without autopsy in 32 cases. There were significantly more diagnoses of acute myocardial infarction and fewer of pulmonary embolism and aortic dissection and rupture among cases without autopsy compared with those followed by autopsy.

**Conclusion:** In unsuccessful resuscitation from out-of-hospital cardiac arrest with pulseless electrical activity as initial rhythm, an autopsy should be performed to determine the correct cause of death.

**Introduction**

Although ventricular fibrillation (VF) has been considered the most common initial rhythm (37–83%) in out-of-hospital cardiac arrest (OHCA), a major decline (50%) in its incidence has occurred during recent decades. Concomitantly, the number of cases of pulseless electrical activity (PEA)...

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**Keywords:** Out-of-hospital; Cardiac arrest; Cause of death; Autopsy; Cardiopulmonary resuscitation; Pulseless electrical activity

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**Summary**

Aims: To study the cause of deaths after witnessed cardiac arrest followed by pulseless electrical activity and unsuccessful out-of-hospital resuscitation; and to detect any differences between causes of death determined at autopsy and those inferred from clinical history.

Methods: In this prospective observational study, data were collected from 91 individuals treated by the emergency medical services in three urban communities in southern Finland.

Results: Cause of death was determined at autopsy in 59 cases and without autopsy in 32 cases. There were significantly more diagnoses of acute myocardial infarction and fewer of pulmonary embolism and aortic dissection and rupture among cases without autopsy compared with those followed by autopsy.

Conclusion: In unsuccessful resuscitation from out-of-hospital cardiac arrest with pulseless electrical activity as initial rhythm, an autopsy should be performed to determine the correct cause of death.

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as initial cardiac rhythm has increased.\textsuperscript{4–6} Recent studies have shown the incidence of primary PEA to be 22–30\% in OHCA,\textsuperscript{7–11} and even 32–52\% in in-hospital cardiac arrest.\textsuperscript{12,13}

Whereas VF is strongly associated with coronary heart disease,\textsuperscript{14} the conditions that cause PEA as initial cardiac rhythm in OHCA often reflect a non-coronary aetiology. Pulmonary embolism (PE) has been shown to be common (36\%) in cases of unexpected cardiac arrest presenting with PEA,\textsuperscript{15} and a study of people who died after primary PEA (previously called electromechanical dissociation) reported that cardiovascular rupture, PE or tension pneumothorax was the cause of death (COD) in 19 of 50 cases.\textsuperscript{16} Another study involving 28 people with PEA as initial rhythm, who died after unsuccessful resuscitation, documented 11 cardiovascular ruptures and 6 PEs.\textsuperscript{17} Courtney et al.\textsuperscript{18} found that PE was the COD in 52\% of cases of primary PEA in a study of witnessed prehospital cardiac arrests.

Obviously, the CODs presented above were based on autopsy findings. However, only a minority (26–39\%) of persons who die after attempted resuscitation undergo autopsy.\textsuperscript{17,19,20} In most cases, the patient’s physician determines the COD according to clinical history. Only if the cause is unknown, if there are reasons to suspect a specific condition or there are legal requirements is an autopsy requested. We hypothesised that if the COD was determined on clinical grounds, based on previous medical history and the course of OHCA, this estimated COD would not be correct in considerable number of cases where the initial cardiac rhythm was PEA. Therefore, we conducted a prospective study of all cases of OHCA where the first recorded rhythm had been PEA and resuscitation was terminated as unsuccessful, and compared CODs determined after autopsy with CODs estimated without autopsy.

Materials and methods

The study included patients in the emergency medical service (EMS) systems in the city of Tampere and Helsinki urban areas in southern Finland. The organisation of these EMS systems has been described in recent papers.\textsuperscript{21,22} In the Tampere EMS a person with cardiac arrest is reached by a basic life support unit within a mean 7.5 min, and the distribution of initial rhythms is ventricular fibrillation/tachycardia (VF/VT) 30\%, PEA 34\% and asystole (ASY) 36\%.\textsuperscript{11} The corresponding data are not available for the Turku and Helsinki areas because they consist of several individual EMSs covered by a helicopter unit. All patients more than 17 years of age, who suffered a witnessed prehospital cardiac arrest of presumed cardiac origin with PEA as initial cardiac rhythm and for whom resuscitation was unsuccessful in the field, were prospectively included. The cause of arrest was presumed to be cardiac if no obvious other cause was evident, e.g. trauma, intoxication, airway obstruction, bleeding or drowning. The patients were treated according to current European Resuscitation Council resuscitation guidelines.\textsuperscript{23}

Finnish legislation states that a police inquiry must be conducted and a medicolegal autopsy performed to determine the cause of death:

- when death is not known to be due to illness, or if the deceased has not been treated by a physician during the period of his/her last illness;
- when death is caused or suspected to have been caused by a crime, accident, suicide, poisoning, occupational disease or medical treatment procedure;
- in cases of otherwise unexpected death. In the event of prehospital death, the police are responsible for necessary further actions and they determine whether any of the above-mentioned conditions apply.

In this study, in the Helsinki area an autopsy was requested for all persons with primary PEA when resuscitation had been unsuccessful. The EMS physician on the scene recorded a recommendation for medicolegal autopsy because of a possible non-coronary cause for the arrest. In the two other study areas, the COD was determined according to current general practice, and no efforts to influence this practice were made. If considered by the police to be required by law, a medicolegal autopsy was performed, and in the remaining cases the treating physician of the deceased decided whether sufficient data existed to allow for a death certificate to be issued without an autopsy. The treating physician could request an autopsy to determine the COD before issuing the death certificate if considered necessary. Data on the COD as stated on the death certificates of individuals who did not undergo an autopsy were obtained from Statistics Finland. The corresponding data of those who were autopsied were retrieved from the autopsy records. The institutional review board of the Helsinki University Hospital approved the study protocol.

The association between CODs and mode of determining COD (autopsy or no autopsy) was analysed with the Pearson Chi-squared test and Fisher’s exact test, where appropriate. The null hypothesis was that no such association exists. Statistical analyses were performed using SPSS\textsuperscript{29} for Windows; \(p\)-values <0.05 were considered significant.

Results

A total of 91 patients were included, 31 in the Helsinki area and 60 in the control areas. Their mean age (±S.D.) was 73.5 ± 11.9 years, and 62\% were men. In the Helsinki area the mean age (±S.D.) was 73.3 ± 7.7 years, and 68\% were men, whereas the corresponding figures in the combined control areas were 73.6 ± 13.7 years and 58\%, respectively. An autopsy was performed in 59 cases (65\%), i.e. 81\% of those in the Helsinki area and in 57\% of those in the Tampere and Turku areas. The distribution of diagnoses between those who were autopsied and those who were not differed (Table 1). There were more diagnoses of acute myocardial infarction (AMI) and less PEs, aortic dissections and ruptures among cases without autopsy compared with those including autopsy. A suspicion of a specific COD was stated on the referral sheets for autopsy in six cases in the control areas. Five of these suspected diagnoses proved to be correct: two AMIs, two PEs and one ruptured acute abdominal aneurysm which was incorrectly suspected to be AMI. The clinical diagnoses in the six cases in the Helsinki area without autopsy (because of protocol violation) were AMI in four and chronic obstructive pulmonary disease in two.
Table 1 Cause of death in relation to mode of determination of death

<table>
<thead>
<tr>
<th>Cause of death</th>
<th>Autopsy</th>
<th>Total</th>
<th>p-Value*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Yes</td>
<td>No</td>
<td></td>
</tr>
<tr>
<td>Cardiac</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acute myocardial infarction</td>
<td>11 (37)</td>
<td>19 (63)</td>
<td>30</td>
</tr>
<tr>
<td>Coronary heart disease</td>
<td>8 (67)</td>
<td>4 (33)</td>
<td>12</td>
</tr>
<tr>
<td>Other cardiac</td>
<td>4 (67)</td>
<td>2 (33)</td>
<td>6</td>
</tr>
<tr>
<td>Non-cardiac</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aortic dissection or rupture</td>
<td>17 (94)</td>
<td>1 (6)</td>
<td>18</td>
</tr>
<tr>
<td>Pulmonary embolism</td>
<td>15 (100)</td>
<td>0 (0)</td>
<td>15</td>
</tr>
<tr>
<td>Intracranial haemorrhage</td>
<td>2 (67)</td>
<td>1 (33)</td>
<td>3</td>
</tr>
<tr>
<td>Other</td>
<td>2 (29)</td>
<td>5 (71)</td>
<td>7</td>
</tr>
<tr>
<td>Total</td>
<td>59 (65)</td>
<td>32 (35)</td>
<td>91</td>
</tr>
</tbody>
</table>

Other cardiac includes two congestive heart failure, undefined pericardial disease, rheumatic valvular disease (mitral and aortic valve), undefined pulmonary hypertension, aortic valve stenosis.

Other includes two chronic obstructive pulmonary disease, anoxic brain damage due to prolonged cardiac arrest, motoneuron disease, cancer, pneumonia and unexplained sudden death. Data are absolute numbers (%).

*A p-value less than 0.05 was considered significant with Chi-Square test and Fisher’s exact test.

Discussion

This study revealed a striking difference in COD after unsuccessful prehospital resuscitation from primary PEA between deaths followed by autopsy and those without an autopsy to confirm clinical opinion. This suggests that when the COD is estimated without autopsy, a significant number of people with aortic rupture (dissection or rupture of aortic aneurysm) or PE may be misdiagnosed to have died from coronary heart disease.

The determination of the correct cause of death is a prerequisite for reliable mortality statistics, and an essential tool in this process is autopsy. However, the number of autopsies performed for hospital in-patients has been decreasing in recent decades,24,25 and the number of autopsies performed for patients who die out-of-hospital has been reported to be 26–39%.17,19 In a large number of cases, the COD is an estimation and true CODs may be missed.

It has been shown that PEA is associated with primary non-coronary causes of cardiac arrest,26 but there have been no systematic efforts to document the causes behind primary PEA in sudden cardiac arrest. As the proportion of cases with VF as a primary rhythm is declining and that of cases with primary PEA is increasing, one way to explore this phenomenon further is to request more frequently an autopsy to determine the COD. Although several of the conditions that cause primary PEA are fatal, the correct diagnosis helps to understand the mechanism behind sudden cardiac arrest.

There are several limitations to this study. First, this was a prehospital study, and the causes of death may be different for hospital in-patients presenting with PEA. In the Helsinki area, a protocol violation resulted in the certification of six cases without autopsy; their death certificates were based on clinical judgment, and their true CODs remain a guess. In the Turku and Tampere areas, the police authorities and physicians were unaware of this study, and their actions in dealing with these cases were supposed to reflect common practice. Despite this, a selection bias is possible. On the other hand, the striking difference in the distribution of diagnoses between those based on autopsy and those based on clinical judgement strongly suggests that the correct diagnosis remained undetermined in a significant number of the latter.

Conclusion

We conclude that in unsuccessful resuscitation from OHCA with PEA as initial rhythm an autopsy should be performed to determine the correct cause of death.

Conflict of interest

None.

Acknowledgments

We thank the crews of the Helsinki and Turku Area HEMS and Tampere District Emergency Medical Services for their valuable help in enrolling the study participants.

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References

Induction of therapeutic hypothermia after cardiac arrest in prehospital patients using ice-cold Ringer’s solution: a pilot study

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Abstract
The cooling and haemodynamic effects of prehospital infusion of ice-cold Ringer’s solution were studied in 13 adult patients after successful resuscitation from non-traumatic cardiac arrest. After haemodynamics stabilisation, 30 ml/kg of Ringer’s solution was infused at a rate of 100 ml/min into the antecubital vein. Arterial blood pressure and blood gases, pulse rate, end-tidal CO2 and oesophageal temperature were monitored closely. The mean core temperature decreased from 35.8 ± 0.9 °C at the start of infusion to 34.0 ± 1.2 °C on arrival at hospital (P < 0.0001). No serious adverse haemodynamic effects occurred. It is concluded that the induction of therapeutic hypothermia using this technique in the prehospital setting is feasible.

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Keywords: Out-of-hospital CPR; Cardiac arrest; Hypothermia; Haemodynamics; Emergency medical services; Fluid therapy

1. Introduction
Mild therapeutic hypothermia improves neurological outcome after cardiac arrest [1,2]. It has been suggested that hypothermia should be induced as soon as possible after return

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of spontaneous circulation (ROSC) [3]. The earliest possible induction of hypothermia implies that the technique should be available in the prehospital setting and that it should be capable of being undertaken easily by non-physician prehospital care providers. The purpose of this study was to investigate the haemodynamic and cooling effects of infusing ice-cold Ringer’s solution immediately after ROSC in closely monitored prehospital patients using a technique described recently [4], but which has not been used in the prehospital setting before.

2. Material and methods

We enrolled 13 prehospital cardiac arrest patients treated by the physician staffed Helsinki Area Helicopter Emergency Medical Air Service (Helsinki Area, HEMS). The study protocol was approved by the institutional review board of Helsinki University Hospital, Finland. Written informed consent was obtained from relatives of the patients before induction of hypothermia. Inclusion criteria were age more than 18 years, cardiac arrest not due to trauma or drug overdose, ROSC later than 10 min from the onset of cardiac arrest, and Glasgow Coma Score \( \leq 5 \) before induction of hypothermia. Exclusion criteria were pregnancy, systolic blood pressure less than 90 mmHg not responding to volume or inotropes, or oesophageal temperature (Tesof) \( < 34.0 \, ^\circ \text{C} \). Patients were treated according to current European Resuscitation Council guidelines [5], and all patients were intubated before ROSC. After ROSC, the patients’ lungs were ventilated manually and end-tidal CO\(_2\) (Life-Cap; Medtronic PhysioControl, Redmond, WA, USA), ECG and pulse oximetry (Zoll M-Series CCT; ZOLL Medical Corp., MA, USA) were monitored continuously. The arterial blood pressure was measured non-invasively every 5 min. An arterial blood sample was drawn and blood gas analysis was performed using the i-STAT portable blood gas analyser (i-STAT Corporation, Windsor, NJ, USA) with the EG6 + cartridge. When the patient had been found eligible and informed consent had been obtained, hypothermia was induced by infusing ice-cold Ringer’s acetate, which had been stored in an insulated box with ice cubes. Pressure bags were used to infuse the target volume of 30 ml/kg at the rate of 100 ml/min via the antecubital vein. Oesophageal temperature was monitored continuously and the infusion was stopped if a core temperature of 33 \( ^\circ \text{C} \) was reached or adverse haemodynamic events occurred before the calculated volume had been infused.

At the end of infusion, arterial blood gas analysis was repeated. The patient was then transported to hospital with all monitoring in place. On arrival at hospital, arterial blood gases were measured and further care in hospital was at the discretion of the treating physicians. Outcome was determined according to the Utstein guidelines [6].

Table 1

<table>
<thead>
<tr>
<th>Patient characteristics and study parameters</th>
<th>Sex</th>
<th>Age (years)</th>
<th>Initial rhythm</th>
<th>BLS (min)</th>
<th>ACLS (min)</th>
<th>Permanent ROSC (min)</th>
<th>T before induction (( ^\circ \text{C} ))</th>
<th>T on admission (( ^\circ \text{C} ))</th>
<th>Delta ( T ) (( ^\circ \text{C} ))</th>
<th>Infused amount (ml)</th>
<th>Outcome</th>
<th>Cause of death</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Male</td>
<td>63</td>
<td>Asystole</td>
<td>2</td>
<td>2</td>
<td>22</td>
<td>35.1</td>
<td>33.1</td>
<td>−2.1</td>
<td>2500</td>
<td>Subarachnoidal haemorrhage</td>
<td>3</td>
<td>Cerebral infarction</td>
</tr>
<tr>
<td>2 Female</td>
<td>72</td>
<td>VF</td>
<td>6</td>
<td>6</td>
<td>21</td>
<td>35.9</td>
<td>33.4</td>
<td>−2.5</td>
<td>2450</td>
<td>Acute myocardial infarction</td>
<td>7</td>
<td>Acute myocardial infarction</td>
</tr>
<tr>
<td>3 Female</td>
<td>53</td>
<td>Asystole</td>
<td>13</td>
<td>14</td>
<td>23</td>
<td>35.8</td>
<td>33.1</td>
<td>−2.7</td>
<td>1800</td>
<td>Airway obstruction</td>
<td>49</td>
<td>Pneumonia</td>
</tr>
<tr>
<td>4 Male</td>
<td>55</td>
<td>VF</td>
<td>5</td>
<td>11</td>
<td>27</td>
<td>35</td>
<td>34.7</td>
<td>−0.3</td>
<td>500</td>
<td>Acute myocardial infarction</td>
<td>1</td>
<td>Acute myocardial infarction</td>
</tr>
<tr>
<td>5 Male</td>
<td>79</td>
<td>VF</td>
<td>7</td>
<td>7</td>
<td>32</td>
<td>35.1</td>
<td>33.2</td>
<td>−1.9</td>
<td>2000</td>
<td>Acute myocardial infarction</td>
<td>5</td>
<td>Arrhythmia</td>
</tr>
<tr>
<td>6 Female</td>
<td>79</td>
<td>PEA</td>
<td>5</td>
<td>5</td>
<td>42</td>
<td>35.8</td>
<td>32.7</td>
<td>−3.1</td>
<td>2200</td>
<td>Acute myocardial infarction</td>
<td>5</td>
<td>Acute myocardial infarction</td>
</tr>
<tr>
<td>7 Male</td>
<td>64</td>
<td>Asystole</td>
<td>6</td>
<td>7</td>
<td>22</td>
<td>37.2</td>
<td>35.3</td>
<td>−1.9</td>
<td>3000</td>
<td>Coronary heart disease</td>
<td>1</td>
<td>Coronary heart disease</td>
</tr>
<tr>
<td>8 Male</td>
<td>41</td>
<td>VF</td>
<td>7</td>
<td>17</td>
<td>46</td>
<td>36.1</td>
<td>33.1</td>
<td>−3.0</td>
<td>3000</td>
<td>Acute myocardial infarction</td>
<td>3</td>
<td>Acute myocardial infarction</td>
</tr>
<tr>
<td>9 Female</td>
<td>52</td>
<td>Asystole</td>
<td>5</td>
<td>8</td>
<td>19</td>
<td>34.2</td>
<td>32.7</td>
<td>−1.5</td>
<td>1000</td>
<td>Acute myocardial infarction</td>
<td>0</td>
<td>Acute myocardial infarction</td>
</tr>
<tr>
<td>10 Male</td>
<td>65</td>
<td>VF</td>
<td>4</td>
<td>9</td>
<td>15</td>
<td>35.9</td>
<td>34.3</td>
<td>−1.6</td>
<td>2500</td>
<td>Acute myocardial infarction</td>
<td>2</td>
<td>Acute myocardial infarction</td>
</tr>
<tr>
<td>11 Male</td>
<td>53</td>
<td>VF</td>
<td>0</td>
<td>7</td>
<td>26</td>
<td>36.9</td>
<td>35.3</td>
<td>−1.6</td>
<td>2500</td>
<td>Myocarditis</td>
<td></td>
<td>Survived</td>
</tr>
<tr>
<td>12 Male</td>
<td>43</td>
<td>VF</td>
<td>7</td>
<td>17</td>
<td>46</td>
<td>36.9</td>
<td>35.3</td>
<td>−1.6</td>
<td>2500</td>
<td>Myocarditis</td>
<td></td>
<td>Survived</td>
</tr>
<tr>
<td>13 Female</td>
<td>71</td>
<td>PEA</td>
<td>3</td>
<td>3</td>
<td>29</td>
<td>36.7</td>
<td>34.3</td>
<td>−2.4</td>
<td>2500</td>
<td>Pulmonary embolism</td>
<td>3</td>
<td>Pulmonary embolism</td>
</tr>
</tbody>
</table>

3. Results
Research data and demographics are shown in Table 1. The mean age of eight male and five female patients was 60.8 ± 12.5 years. ROSC was achieved at 26 ± 10 min. Initial cardiac rhythm was ventricular fibrillation (53.8%), asystole (30.8%) or pulseless electrical activity (15.4%). The oesophageal temperature decreased significantly during treatment (Fig. 1). The mean infused volume was 2188 ± 754 ml. Infusion started at 27 ± 12 min after ROSC and mean duration of infusion was 25 ± 11 min. One patient (patient 2) experienced a transient decrease in blood pressure, but no other haemodynamic side effects were observed (Fig. 1). Therapeutic hypothermia was continued in patients 3, 4, 8, 10 and 11 for 24 h after admission to hospital according to the same protocol as in the HACA study [1].

4. Discussion
The results of this pilot study suggest that induction of therapeutic hypothermia with rapid infusion of ice-cold Ringer’s solution soon after ROSC is well tolerated and feasible in the prehospital setting. This treatment has been documented in hospitalised patients starting, on average, 73 min after ROSC [4]. However, the cardiovascular situation might be different immediately after ROSC, and the haemodynamic effects of induction of cooling by infusion at this stage have not been reported, although recommended [3]. This feasibility study suggests that further evaluations of this technique seem justified. We used an oesophageal thermal probe as a fast and simple method for continuous measurement of core temperature. There are, however, weaknesses of its use, which may explain the findings in two of our patients. The probe may come in close contact with the superior vena cava and thus register falsely low “core” temperatures when blood mixed with cold saline passes by. This may explain why the temperature of patient 4 decreased rapidly after only 500 ml of fluid, and why the temperature rose again shortly after the infusion was stopped. The probe can also inadvertently be advanced too far, eventually entering the stomach, thus not reflecting correct values (e.g. patient 10 whose temperature remained unchanged).

The rather low survival rate, although not the study endpoint, may be explained by the large number of patients with known factors for poor outcome [7]. There were four patients with non-coronary causes for arrest, and times to ROSC were long. Therefore, the infusion of cold saline is unlikely to have contributed to the number of non-survivors. There are some obvious limitations in our study. First, the small number of patients increases the risk of a beta error, but the lack of observed adverse effects of this treatment renders further studies warranted. Second, we did not have a control group. Third, the measurement of oesophageal temperature is prone to errors, but the decrease in temperature was continuous and relevant in the majority of patients.

5. Conclusion
We conclude that induction of therapeutic hypothermia with peripheral infusion of ice-cold Ringer’s solution seems feasible soon after ROSC in the prehospital setting. Because this technique is also readily available to non-physician care providers, the induction of therapeutic hypothermia can be started earlier if found safe and well tolerated in larger trials.

Acknowledgements
We would like to thank the crew of the Helsinki Area HEMS for the valuable help in enrolling the study patients. This study was supported by institutional EVO grant #9B102 from Special State Allocation via Tampere University Hospital.

References


