Adrenalin can restart the heart, but is no good for the brain

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Abstract

The study
A randomised trial of epinephrine in out-of-hospital cardiac arrest


This project was funded by the National Institute for Health Research HTA Programme (project number 12/127/126).

To read the full NIHR Signal, go to: https://discover.dc.nihr.ac.uk/content/signal-000639/adrenaline-can-restart-the-heart-but-is-no-good-for-the-brain

Why was this study needed?

Adrenalin (epinephrine) injections have been commonly used during cardiopulmonary resuscitation (CPR) for cardiac arrest for more than 60 years, without clear evidence if they are helpful or harmful. Adrenalin can increase the likelihood that the heart will regain a normal rhythm as it directs blood flow to the heart. However, it also causes constriction of small blood vessels, which can reduce blood flow to other organs—including the brain—and may lead to neurological damage.

Use of adrenalin has been linked to better short term survival in observational studies, so it has remained in the cardiac arrest guidelines. Randomised controlled trials have been needed to determine if the benefits outweigh the potential harms. This study assessed routine adrenalin use in cardiac arrests occurring outside hospital.

What did this study do?

The PARAMEDIC2 randomised controlled trial allocated 8014 people with cardiac arrest (average age 70) to receive either 1 mg adrenaline or placebo saline injections as part of advanced CPR treatment by paramedics at the scene. On average, people in the adrenaline group received a total dose of 4.9 mg.

In 59% of each group, bystanders performed CPR. Ambulances took just over six minutes to arrive and paramedics gave the injections on average 21 minutes after emergency services were first called. Major outcomes were adjusted for variations in these factors.

More than one third of patients had unwitnessed cardiac arrest, more than half had no heart electrical activity (asystole), and only 19% had a shockable rhythm. This may have contributed to the low survival rates.

This large, well designed trial provides much needed UK based, high quality evidence to inform practice.

What did it find?

• People who received adrenaline had a slightly higher rate of survival at 30 days, 130/4105 (3.2%) compared with 94/3999 (2.4%) who received the saline placebo (adjusted odds ratio 1.47, 95% confidence interval 1.09 to 1.97).

• Return of spontaneous circulation was much more likely with adrenaline, occurring in 1457 (36.3%) people in the adrenaline group versus 468 (11.7%) people in the placebo group. A substantially higher number of people given adrenaline survived until hospital admission, 947 (23.8%) compared with 319 (8%) of the placebo group.

• Although the overall rate of survival at 30 days was slightly better with adrenaline, 39/126 (31%) people had severe neurological disability in the adrenaline group, compared with 16/90 (17.8%) in the placebo group. Severe neurological disability was defined as a score of 4 or 5 on the modified Rankin scale.

• Only 27 people in total had no neurological symptoms at discharge. There were similar numbers of people in each group when combining those with no symptoms, mild or moderate neurological disability, or severe disability (adjusted odds ratio 0.87, 95% confidence interval 0.59 to 1.29).

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CPR, and defibrillation. Strategies to increase public training in CPR and make more defibrillators available may increase survival rates at 30 days. More of those survivors had increased survival from prompt recognition, bystander CPR, and defibrillation. Strategies to increase public training in CPR and make more defibrillators available may increase the number of people surviving out-of-hospital cardiac arrest.

The overall survival rate is low, so the jury is still out on whether adrenaline should be reserved for use in certain heart rhythms or within a particular time period and not for others.

What are the implications?

Adrenaline improved the return of spontaneous circulation and likelihood of survival to reach hospital, but only slightly increased survival rates at 30 days. More of those survivors had severe neurological problems.

It remains unclear if out-of-hospital protocols should change as a result of this trial. The findings are also not able to inform hospital cardiac arrest protocols, as use of adrenaline typically occurs within three minutes of cardiac arrest.

Other studies of cardiac arrest taking place out of hospital have shown better outcomes from prompt recognition, bystander CPR, and defibrillation. Strategies to increase public training in CPR and make more defibrillators available may increase the number of people surviving out-of-hospital cardiac arrest.

The modified Rankin scale scores for disability

- 0 No symptoms
- 1 No major disability despite symptoms
- 2 Slight disability; unable to perform all previous activities but able to look after own affairs without assistance
- 3 Moderate disability; requires some help but able to walk unassisted
- 4 Moderate severe disability; unable to walk and attend to bodily needs without assistance
- 5 Severe disability; bedridden, incontinent, and requiring constant nursing care
- 6 Death.

The number needed to treat (NNT) with adrenaline injections to prevent one death in this trial was 112 people. Other trials of out-of-hospital cardiac arrest found much lower numbers for

- early recognition, NNT 11 people
- bystander CPR, NNT 15 people
- early defibrillation, NNT 5 people.

What does current guidance say on this issue?

The 2015 Resuscitation Council UK guideline recommends giving adrenaline every three to five minutes during CPR if a normal heart rhythm is absent. Guidance on the early management of people who have had a cardiac arrest by the Faculty of Pre-Hospital Care at the Royal College of Surgeons of Edinburgh (2017) emphasises the importance of recognising that the person is having a cardiac arrest and beginning CPR as soon as possible.

Education into practice

Are you comfortable with managing cardiac arrest out of hospital?
Do you know where the nearest defibrillators are to your places of work?
Have you checked the emergency drugs, including adrenaline, where you work and do you know the protocols for administering them?

Competing interests The BMJ has judged that there are no disqualifying financial ties to commercial companies. The authors declare the following other interests: none.

Further details of the BMJ policy on financial interests is here: https://www.bmj.com/about-bmj/resources-authors/forms-policies-and-checklists/declaration-competing-interests

Contributors All authors contributed to development and review of this summary, as part of the wider NIHR Signals editorial team (https://www.bmj.com/NIHR-signals). RC is guarantor.

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