Training approaches for the deployment of a mechanical chest compression device: a randomised controlled manikin study

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ABSTRACT

Objectives To evaluate the effect of training strategy on team deployment of a mechanical chest compression device.

Design Randomised controlled manikin trial.

Setting Large teaching hospital in the UK.

Participants Twenty teams, each comprising three clinicians. Participating individuals were health professionals with intermediate or advanced resuscitation training.

Interventions Teams were randomised in a 1:1 ratio to receive either standard mechanical chest compression device training or pit-crew device training. Training interventions lasted up to 1 h. Performance was measured immediately after training in a standardised simulated cardiac arrest scenario in which teams were required to deploy a mechanical chest compression device.

Primary and secondary outcome measures Primary outcome was chest compression flow fraction in the minute preceding the first mechanical chest compression. Secondary outcomes included cardiopulmonary resuscitation quality and mechanical device deployment metrics, and non-technical skill performance. Outcomes were assessed using video recordings of the test scenario.

Results In relation to the primary outcome of chest compression flow fraction in the minute preceding the first mechanical chest compression, we found that pit-crew training was not superior to standard training (0.76 (95% CI 0.73 to 0.79) vs 0.77 (95% CI 0.73 to 0.82), mean difference −0.01 (95% CI −0.06 to 0.03), P=0.572). There was also no difference between groups in performance in relation to any secondary outcome.

Conclusions Pit-crew training, compared with standard training, did not improve team deployment of a mechanical chest device in a simulated cardiac arrest scenario.

Trial registration number ISRCTN43049287; Pre-results.

BACKGROUND

Mechanical chest compression devices provide a method to deliver high-quality chest compressions to patients in cardiac arrest.1 Despite evidence that manual chest compression delivery is often suboptimal and that high-quality chest compression delivery is associated with improved survival, randomised controlled trials in out-of-hospital cardiac arrest (OHCA) have found that use of a mechanical chest compression device is not superior to manual chest compressions in relation to patient outcome.2–5 In contrast, very low quality evidence from in-hospital cardiac arrest (IHCA) suggests that use of devices may be associated with improved patient outcome.6

The key risk associated with use of mechanical chest compression devices is the pause in chest compressions associated with their deployment.1 In the LINC randomised controlled trial of the use of mechanical chest compression devices in OHCA, a substudy found that the median chest compression pause associated with device deployment was 36.0 s (IQR 19.5–45.5).7 Such pauses are associated with a reduction in coronary pressure during the early part of a cardiac arrest and may therefore offset the potential benefit of improved chest compression delivery associated with devices.8

Training as a team may be an effective strategy to optimise the delivery of care that is usually provided by a clinical team.9 Pit-crew resuscitation describes a concept where clinicians in a team act in a predetermined...
way to undertake set tasks in a specific order, akin to a formula one motor racing team. In a clinical before–after quality improvement study, the use of this concept alongside other interventions was associated with a marked reduction in the pause associated with device deployment in OHCA. However, the impact of the use of a pit-crew training approach has not, to date, been tested against other team training approaches in a randomised controlled trial.

**METHODS**

We conducted a randomised controlled parallel group trial to evaluate the impact of pit-crew training, compared with standard training, on mechanical chest compression deployment in a simulated cardiac arrest. The mechanical chest compression device used was the LUCAS-2 mechanical chest compression device (Jolife AB/Physio-Control, Lund, Sweden). Outcomes were evaluated through a video-recorded simulation test that was undertaken immediately after the training intervention.

All participants provided written informed consent prior to receiving any study intervention.

**Participants**

Teams of three clinicians were recruited to participate in the trial. To be eligible to participate, individual clinicians were required to be registered health professionals with current Immediate Life Support or Advanced Life Support (ALS) certification, and to have completed the online manufacturer’s device training package. Participants were not eligible if they had an injury or disability that prevented use or handling of the device, or if they had received practical training in the use of the device in the preceding 6 months.

Each team was required to be composed of three eligible clinicians, of which one was required to be an ALS-certificated provider or instructor. As such, teams were broadly reflective of the standard required of IHCA teams in the UK.

**Study process**

We advertised the study at the hospital site through posters in staff areas, emails to staff and face-to-face discussions. Staff that consented to participate in the study were asked for their availability and allocated a training slot. As such, teams were created based on convenience, in that we established teams based on individual clinician’s availability to attend training at a specified time. This broadly reflects how hospital cardiac arrest teams are created in clinical practice, in that the designated team will be drawn from clinicians who have been scheduled to work a particular shift.

On attendance at the training session, the team was randomised using a simple randomisation system provided by an internet-based randomisation service, which ensured allocation concealment (Sealed Envelope, London, UK). Teams were randomly allocated in a 1:1 ratio to receive either pit-crew training or standard training. After randomisation, a researcher (KC), with experience in teaching ALS courses, immediately delivered the allocated training intervention. The team was not blinded to their allocated training intervention.

Following the training, the teams undertook a standardised manikin-based simulation test. The team acted as a hospital cardiac arrest team. The scenario, given in the SBAR format (situation, background, assessment, recommendation), described a 62-year-old male who had undergone tracheal intubation due to hypoxia secondary to a probable pulmonary embolus and who had subsequently become haemodynamically unstable leading to cardiac arrest.

During the scenario, a mechanical chest compression device arrived during the second cardio-pulmonary resuscitation (CPR) cycle, and the team was required to use the device. The length of the scenario was approximately 8 min (four cycles of CPR).

The simulation test was video-recorded. Two digital video recorders were used to mitigate against the risk of data loss and possible obstruction of a camera by participants.

**Study intervention**

An overview of the two training approaches, based on the TIDieR (template for intervention description and replication) framework, is included as an electronic supplement (online supplementary tables S1 and S2). Both training approaches incorporated a presentation on device deployment and use, followed by an opportunity for the team to practise these skills. Skill practice was scenario-based, with feedback following each scenario. It was anticipated that training would last approximately 45 min, although teams were allowed to practice for as long as required.

The key difference between the training approaches was that teams randomised to pit-crew training received an overview of the concept and potential value of the pit-crew system in the training presentation, together with guidance on how to operationalise the pit-crew system in deploying the mechanical chest compression device. This pit-crew training system requires team members to adopt a nominated role that is associated with specific tasks, with team members trained to perform tasks in a coordinated manner. These concepts were also highlighted in feedback during practice scenarios.

In both training approaches, teams were trained to deploy the device in two stages, such that there was a pause for the deployment of the device back plate followed by the resumption of CPR and then a subsequent pause to enable deployment of the upper part of the mechanical device.

**Outcome measures**

The primary outcome was the chest compression flow fraction in the minute preceding the first mechanical chest compression. Flow fraction describes the proportion of time in which chest compressions are being delivered.
over a designated period. This specific outcome was selected as it enabled us to capture all pauses that may be attributable to device deployment.

There were a number of secondary outcome measures including chest compression flow fraction (prior to the first mechanical chest compression, following the first mechanical chest compression, whole event), the duration of chest compression pauses associated with device deployment and non-technical skills measured using the Team Emergency Assessment Measure (TEAM) tool.14 Participating teams were not routinely informed as to the precise outcome measures being collected. However, a key focus of training interventions was the minimisation of pauses during deployment, and outcomes were recorded in the study entry on the trial registration website.

**Data management**

Videos were reviewed by two researchers (KC, RMV) independently. The first reviewer (KC) delivered the training intervention and so was not blinded to training allocation. The second reviewer (RMV) was blinded to training allocation. Videos were viewed using software that enabled timings to be derived to the nearest one-tenth of a second.

We assessed agreement between video reviewers by computing the average difference and 95% CI or median difference and IQR for each outcome and plotting data using a Bland-Altman plot.15 Bland-Altman plots visually depict the level of agreement between two reviewers, with good agreement represented by a small difference between the upper and lower limit of agreement and the majority of measurements falling between the limits of agreement. We used the mean value of the two reviewers in the analysis.

**Statistical analysis**

Our planned sample size of 20 teams (three clinicians per team) was based on demonstrating an absolute increase in flow fraction in the minute preceding the

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**Table 1** Team characteristics

|                  | Pit-crew training (10 teams) | Standard training (10 teams) |
|------------------|-----------------------------|-----------------------------|
| Teams with n doctors, n (%) |                             |                             |
| 0                | 3 (30)                      | 3 (30)                      |
| 1                | 5 (50)                      | 2 (20)                      |
| 2                | 2 (20)                      | 3 (30)                      |
| 3                | 0 (0)                       | 2 (20)                      |
| Teams with n Advanced Life Support instructors, n (%) |                             |                             |
| 0                | 4 (40)                      | 5 (50)                      |
| 1                | 2 (20)                      | 4 (40)                      |
| 2                | 3 (30)                      | 1 (10)                      |
| 3                | 1 (10)                      | 0 (0)                       |
| Teams with at least one member that has previously received device training, n (%) | 5 (50)                      | 4 (40)                      |
| Teams with at least one member with experience of using a device in clinical practice, n (%) | 9 (90)                      | 10 (100)                     |
Table 2  Individual participant characteristics

|                                | Pit-crew training (N=30) | Standard training (N=30) |
|--------------------------------|--------------------------|--------------------------|
| **Gender**                     |                          |                          |
| Male, n (%)                    | 12 (40.0)                | 14 (46.7)                |
| **Health profession, n (%)**   |                          |                          |
| Doctor                         | 9 (30)                   | 14 (46.7)                |
| Nurse                          | 21 (70.0)                | 12 (40.0)                |
| Operating department practitioner | –                      | 1 (3.3)                 |
| Paramedic                      | –                        | 2 (6.7)                  |
| Other                          | –                        | 1 (3.3)                  |
| **Professional experience, years, median (IQR)** | 13.5 (6.3–21.3) | 8.0 (3.8–13.5) |
| **Grade, n (%)**               |                          |                          |
| Band 5/FY1                     | 3 (10.0)                 | 5 (16.7)                 |
| Band 6/SHO                     | 8 (26.7)                 | 11 (36.7)                |
| Band 7/registrar               | 14 (46.7)                | 9 (30.0)                 |
| Band 8+/consultant             | 5 (16.7)                 | 5 (16.7)                 |
| **Current speciality, n (%)**  |                          |                          |
| Acute medicine                 | 6 (20.0)                 | 5 (16.7)                 |
| Critical care                  | 11 (36.7)                | 20 (66.7)                |
| Emergency medicine             | 6 (20.0)                 | 2 (6.7)                  |
| Surgery                        | 3 (10.0)                 | 2 (6.7)                  |
| Other                          | 4 (13.3)                 | 1 (3.3)                  |
| **Resuscitation qualification, n (%)** |                      |                          |
| Immediate Life Support provider | 7 (23.3)                | 7 (23.3)                 |
| Advanced Life Support provider  | 12 (40.0)                | 17 (56.7)                |
| Advanced Life Support instructor | 11 (36.7)              | 6 (20.0)                 |
| **Approximate number of cardiac arrests attended in last 6 months, median (IQR)** | 4.5 (2.0–10.0) | 8.0 (1.8–15.8) |
| Previously received device training, yes, n (%) | 7 (23.3) | 4 (13.3) |
| **Previous use of a device in clinical practice, yes, n (%)** |                      |                          |
| If yes, which device, n (%)    | 18 (60.0)                | 17 (56.7)                |
| LUCAS                          | 10 (55.6)                | 8 (47.1)                 |
| AutoPulse                      | 11 (61.1)                | 10 (58.8)                |
| Thumper                        | 2 (11.1)                 | 1 (5.9)                  |
| If yes, how many times used, median (IQR) | 2.5 (1.0–5.8) | 3.0 (1.5–4.5) |

first mechanical chest compression of 0.15 (baseline 0.58, SD 0.10) at 90% power with a P value of 0.05. Due to the nature of the study, we did not increase the sample size to account for drop-outs. If a team member did not attend their allocated training session, then the team would not be eligible and so would not be randomised.

Data analysis was based on intention-to-treat principles. For baseline team and individual data, we report categorical variables as number and percentage, while continuous variables are described as mean (95% CI) or median (IQR), depending on the normality of the data. All outcomes are assessed at the team level. For normally distributed continuous outcomes, we summarise team performance as mean (95% CI), and compare groups using an independent t-test and report the mean difference, 95% CI and P value. For non-normally distributed continuous outcomes, we summarise group performance as median (IQR) and compare groups using a Mann-Whitney U test and report the P value.

**RESULTS**
Between June 2016 and September 2016, 78 clinicians consented to participate in the study (figure 1). Of these, 60 participated in randomised teams. Four participants were excluded after giving consent to participate but prior to randomisation (one identified that their
resuscitation certification had expired, one developed an injury that prevented use of the device and two received previous practical mechanical device training). For the remaining 13 participants, we were unable to identify a convenient time for training prior to randomising the 20 teams required.

Demographic data at the team and individual level are shown in table 1 and table 2, respectively. Team characteristics were comparable between groups. There were some differences between groups at the individual participant level in relation to, for example, clinical experience and specialty.

For the primary outcome, the average difference between reviewers for the 20 cases was 0.01 (95% CI: −0.01 to 0.02). Bland-Altman and average differences for other outcomes are included in the electronic supplement (online supplementary table S3 and figure S1). Based on these data, outcome analyses are based on the average data from the two reviewers.

In relation to the primary outcome, we found no difference in the flow fraction preceding the first mechanical chest compression between study groups (0.76 (95% CI: 0.73 to 0.79) vs 0.77 (95% CI: 0.73 to 0.82), mean difference −0.01 (95% CI: −0.06 to 0.03), P=0.572) (table 3).

The chest compression pause associated with the deployment time of both the back plate and upper part of the device being similar between groups. We also observed no difference between groups in relation to any other secondary outcome.

There were no study adverse events, and all training interventions were delivered as planned, with time taken to deliver training interventions being similar between groups (table 3). During one simulation test (pit-crew training arm), the device battery failed as due to

| Table 3 | Outcome measures (based on average from two videotape reviewers) |
|---------|---------------------------------------------------------------|
|         | Pit-crew training (n=10) | Standard training (n=10) | Mean difference (95% CI) | P value* |
| Device deployment, mean (95% CI) |                     |                     |                      |
| Flow fraction in minute preceding first mechanical CC | 0.76 (0.73 to 0.79) | 0.77 (0.73 to 0.82) | −0.01 (−0.06 to 0.03) | 0.572 |
| Time to deploy back plate (s) | 3.80 (2.83 to 4.76) | 3.82 (2.62 to 5.02) | −0.03 (−1.46 to 1.41) | 0.971 |
| Time to deploy upper part of device (s) | 9.99 (8.84 to 11.14) | 9.67 (8.02 to 11.32) | 0.32 (−1.55 to 2.19) | 0.724 |
| Total pause for mechanical device deployment (s) | 14.33 (12.62 to 16.03) | 13.56 (11.05 to 16.06) | 0.77 (−2.04 to 3.58) | 0.572 |
| Time from device arrival to first mechanical CC (s), median (IQR) | 55.25 (51.63 to 75.24) | 60.43 (52.70 to 73.99) |                     | 0.912† |
| Compliance with manufacturer’s guidelines (out of 8), median (IQR) | 8 (8–8) | 8 (8–8) |                     | 0.739† |
| CPR quality |                     |                     |                      |
| Flow fraction, mean (95% CI) |                     |                     |                      |
| Whole episode | 0.94 (0.93 to 0.95) | 0.94 (0.93 to 0.95) | 0.00 (−0.01 to 0.01) | 0.790 |
| Predevice deployment | 0.90 (0.89 to 0.92) | 0.90 (0.88 to 0.91) | 0.01 (−0.01 to 0.03) | 0.538 |
| Postdevice deployment | 0.97 (0.96 to 0.98) | 0.97 (0.96 to 0.98) | 0.00 (−0.01 to 0.01) | 0.681 |
| Number of pauses>5 s, median (IQR) | 2.00 (1.75 to 3.00) | 2.50 (1.00 to 4.13) |                     | 0.853† |
| Number of shocks delivered, median (IQR) | 2 (2–2) | 2 (2–2) |                     | 1.000† |
| Shocks delivered appropriately, median (IQR) | 2 (2–2) | 2 (2–2) |                     | 1.000† |
| Perishock pause (s), median (IQR) |                     |                     |                      |
| Preshock pause | 0.00 (0.00–0.00) | 0.00 (0.00–0.00) |                     | 1.000† |
| Postshock pause | 0.00 (0.00–0.00) | 0.00 (0.00–0.00) |                     | 1.000† |
| Non-technical skills, TEAM tool |                     |                     |                      |
| TEAM tool, component score (out of 44) | 38.0 (35.7 to 40.3) | 37.6 (34.8 to 40.4) | 0.40 (−2.94 to 3.74) | 0.804 |
| TEAM tool, global (overall) score (out of 10) | 8.1 (7.2 to 8.9) | 7.9 (7.3 to 8.6) | 0.15 (−0.87 to 1.17) | 0.760 |
| Training delivery |                     |                     |                      |
| Duration of training, mean (95% CI)‡ | 49.0 (44.0 to 54.0) | 45.3 (40.5 to 50.1) | 3.1 (−2.7 to 10.1) | 0.244 |

*By independent t-test unless stated. †By Mann-Whitney U test. ‡Data point measured directly during training intervention, so not based on assessment from two reviewers.

CC, chest compression; CPR, cardiopulmonary resuscitation; TEAM, Team Emergency Assessment Measure.
researcher error the device battery was not charged prior to training. This battery failure did not affect the primary outcome, but did marginally reduce flow fraction across the whole scenario and following device deployment.

**DISCUSSION**

In this randomised controlled manikin trial, we found that the use of a pit-crew training approach, compared with a standard training approach, did not affect the deployment of a mechanical chest compression device in simulated IHCA. Similarly, the use of pit-crew training did not affect any other device deployment, CPR quality or non-technical skills metric.

Our study, in contrast to previous pit-crew cardiac arrest studies, directly compared pit-crew training with a standard training approach. While not directly comparable, it is noteworthy that previous studies have typically reported an association between implementation of pit-crew principles and improvements in process and patient-focused outcomes. For example, in a before–after OHCA clinical study, the implementation of pit-crew training alongside real-time feedback and post-event debriefing was associated with an improvement in neurologically intact survival following OHCA (OR 2.3, 95% CI 1.3 to 4.0).

There are four possible explanations for this apparent contrast in findings. First, previous studies have tended to implement pit-crew training alongside other interventions, such as real-time feedback, additional training and cardiac arrest debriefing. As such, the findings of previous studies may reflect the impact of the overall package of interventions, such that pit-crew training either did not have an independent effect or complementary interventions enhanced its effect. Second, previous studies have adopted an observational design, such that the findings may be attributable to unmeasured confounders rather than the intervention that was implemented.

A third explanation is that where team performance is high, as observed in this study, pit-crew training provides no additional benefit. In Levy et al's before–after study, where the pit-crew training intervention was associated with a marked improvement in care delivery, the median preintervention pause between the final manual chest compression and first mechanical chest compression prior to implementation of pit-crew training was 21 s (IQR 15 to 31). In contrast, the mean total pause associated with device deployment in the standard training group in this study was about 14 s, such that the opportunity for pit-crew training to have a measurable effect on team performance may have been limited.

Finally, it may be that pit-crew concepts provide more value in the clinical setting, compared with simulated cardiac arrest conditions. Replicating the real-life complexity of cardiac arrest management in the simulation laboratory is challenging. It is possible that pit-crew training provides most advantage in the unpredictable clinical setting, where the system provides a clear framework to direct effective team functioning.

Our study has several weakness. First, we tested performance in a simulated cardiac arrest scenario that occurred immediately after the training intervention, such that we do not know how training interventions affected performance either in the longer term or in the clinical setting. Second, participants were often experienced clinicians with frequent exposure to cardiac arrest, so it is unclear whether findings are generalisable to less-experienced clinicians. Third, we found that performance in both groups was markedly better than that estimated in our power calculation, so our study was underpowered to reliably detect a clinically important difference between groups. Fourth, we did not capture participant views on the impact of training approaches on team functioning. Such information may have captured the personal learning style of individuals within the team and how receptive they were to the different training approaches. Finally, while agreement between video reviewers was generally good, there was some variation in relation to the TEAM tool outcomes. However, analysis at an individual reviewer level produced similar results to our overall planned analysis (data not presented).

**CONCLUSION**

In this randomised controlled manikin trial, we found that pit-crew training, compared with standard device training, did not improve the deployment of mechanical chest compression devices in a simulated cardiac arrest. The time taken to deliver training was similar. Future controlled trials should examine the effect of pit-crew training in the clinical setting and on long-term performance.

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**Contributors** KC conceived and designed the study, acquired the data, analysed the data, contributed to interpretation of the data, critically revised the manuscript for important intellectual content, drafted the manuscript and approved the final manuscript. RMV analysed the data, contributed to interpretation of the data, critically revised the manuscript for important intellectual content and approved the final manuscript. AD conceived and designed the study, contributed to interpretation of the data, critically revised the manuscript for important intellectual content and approved the final manuscript. RL conceived and designed the study, contributed to interpretation of the data, critically revised the manuscript for important intellectual content and approved the final manuscript. BO conceived and designed the study, contributed to interpretation of the data, critically revised the manuscript for important intellectual content and approved the final manuscript. GDP conceived and designed the study, contributed to interpretation of the data, critically revised the manuscript for important intellectual content and approved the final manuscript.

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