The Head Injury Retrieval Trial (HIRT): a single-centre randomised controlled trial of physician prehospital management of severe blunt head injury compared with management by paramedics only

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ABSTRACT

Background Advanced prehospital interventions for severe brain injury remains controversial. No previous randomised trial has been conducted to evaluate additional physician intervention compared with paramedic only care.

Methods Participants in this prospective, randomised controlled trial were adult patients with blunt trauma with either a scene GCS score <9 (original definition), or GCS<13 and an Abbreviated Injury Scale score for the head region ≥3 (modified definition). Patients were randomised to either standard ground paramedic treatment or standard treatment plus a physician arriving by helicopter. Patients were evaluated by 30-day mortality and 6-month Glasgow Outcome Scale (GOS) scores. Due to high non-compliance rates, both intention-to-treat and as-treated analyses were preplanned.

Results 375 patients met the original definition, of which 197 was allocated to physician care. Differences in the 6-month GOS scores were not significant on intention-to-treat analysis (OR 1.14, 95% CI 0.73 to 1.75, p=0.56) nor was the 30-day mortality (OR 0.91, 95% CI 0.60 to 1.38, p=0.66). As-treated analysis showed a 16% reduction in 30-day mortality in those receiving additional physician care; 60/195 (29%) versus 81/180 (45%), p<0.01, Number needed to treat =6. 338 patients met the modified definition, of which 182 were allocated to physician care. The 6-month GOS scores were not significantly different on intention-to-treat analysis (OR 1.14, 95% CI 0.73 to 1.75, p=0.56) nor was the 30-day mortality (OR 1.05, 95% CI 0.66 to 1.66, p=0.84). As-treated analyses were also not significantly different.

Conclusions This trial suggests a potential mortality reduction in patients with blunt trauma with GCS<9 receiving additional physician care (original definition only). Confirmatory studies which also address non-compliance issues are needed.

Trial registration number NCT00112398.

INTRODUCTION

Insults accompanying severe blunt traumatic brain injuries such as hypoxia, hypotension and hypercapnia are associated with poor outcomes. It has therefore been suggested that patients with traumatic brain injury are particularly likely to benefit from advanced prehospital interventions that may treat or prevent these insults.¹ Approaches to providing such interventions have varied across jurisdictions with physicians, or paramedics and nurses with advanced training used as prehospital care providers. However, there have been no randomised trials evaluating effectiveness of an advanced intervention prehospital treatment policy. This clinical trial aimed to determine whether the intervention of a physician led prehospital medical team resulted in better functional outcomes than a standard care regimen of care delivered by paramedics alone for severe blunt traumatic brain injury.

METHODS

Design We designed a randomised controlled, parallel group trial, conducted in the greater Sydney area of New South Wales, Australia, to evaluate the impact of prehospital physician intervention in the setting of severe head injury.² A member of the physician response team monitored the New South Wales Ambulance computerised dispatch system via a web link to identify appropriate cases. Appropriate cases were blunt trauma in adults (>15 years) who were reported to be unconscious, having an altered level of consciousness, or a high-energy mechanism such as fall of more than 5 m or a pedestrian struck by a truck. Incidents with five or more casualties were excluded, as special arrangements applied in these situations. Identified cases were randomised to either standard ground paramedic treatment or standard treatment plus a physician arriving by helicopter.

Inclusion criteria and protocol modification The original severe-head-injury subgroup was defined as patients with a GCS<9 at their first assessment by the treating prehospital team. The
study protocol was modified after the first interim analysis (140
patients enrolled meeting the original criteria), as some patients
who met the criteria for the original severe-head-injury sub-
group did not have severe head injuries, and many patients with
an initial score of 9–12 had anatomically severe injury but
would have been excluded from the primary analysis. The
outcomes of patients identified by both original (GCS<9, ‘original’) and
modified (GCS 3–12 and Abbreviated Injury Scale score
≥3, ‘modified’) definitions are reported. The criteria for random-
ising cases remained unchanged.

Randomisation and masking
Randomisation method was by computer-generated permuted
blocks of 10, stratified by mechanism of injury (fall, transporta-
tion or other) and provided centrally by an automated telephone
system from the National Health and Medical Research Council
Clinical Trials Centre in Sydney. Masking at the time of inter-
vention was not possible due to the nature of the interventions
being studied. The outcome assessor who evaluated patients at
follow-up 6 months post injury was blinded to treatment group
and had no other role in the conduct of the trial.

Interventions
All patients received a ground paramedic response (standard
care). If the patient was allocated to physician care, the physician
team was dispatched by helicopter in addition to the standard
response. The intervention team consisted of physicians with spe-
cialist certification in anaesthesia, emergency or intensive care
medicine with more than 12 months of prehospital experience,
plus a paramedic from the ambulance service. Treatment by the
physician team followed protocols for standard trauma therapy
as determined by the Royal Australasian College of Surgeons
Early Management of Severe Trauma programme. If the phys-
cian team had not arrived when the road crew were ready to
depart, patients were transported without waiting for the phys-
cian team, except in cases of high-grade airway obstruction. The
road ambulance dispatch centre or officers on scene were able to
request a physician team, regardless of treatment allocation,
according to the system that existed before the trial began. Road
paramedics could cancel the physician team if they believed that
the patient did not require a higher level of intervention than
that provided under Ambulance protocols.

Treatment groups
Standard care by paramedics only was according to written proto-
cols of the New South Wales Ambulance including: cannulation
and up to 1-litre intravenous crystalloid infusion; ventilation via
supraglottic airways and bag-valve-mask ventilation; intubation
without neuromuscular blockade; needle chest decompression;
midazolam for seizures or sedation; analgesia with methoxyflurane
and morphine and splinting and spinal immobilisation.
Monitoring consisted of pulse oximetry, ECG and manual BPs.

Interventions additional to standard care by the physician
team included: anaesthesia with neuromuscular blockade and
induction agents such as thiopentone or ketamine; surgical
airways, needle cricothyroidotomy; tube or open thoracostomy;
administration of 7.5% saline or 20% mannitol and administra-
tion of packed red blood cells. Intubated patients were venti-
lated with a portable volume cycled ventilator and monitored
with waveform capnography and automated non-invasive BP
measurement.

All patients were transported to the nearest major (Level 1)
trauma service hospital in accordance with the ambulance service
transport protocols by the fastest available vehicle, given

Follow-up
Surviving, consenting patients who met the trial’s follow-up cri-
tera were interviewed by telephone 6 months after their injury.
Criteria for interview were first GCS≤12; a fall in GCS to ≤12
before arrival in the emergency department unrelated to medi-
cation (including anaesthetics and sedatives) or an Abbreviated
Injury Scale score ≥3 for the head region (2005 definitions).

Outcome assessment interviews were conducted by a single
research nurse, blinded to treatment allocation, or if an inter-
view was not possible, outcome data was obtained from hospital
medical records. Other patients not meeting these definitions for
formal outcome assessment had their survival status at hos-
pital discharge determined.

Outcome measures
The primary endpoint was the score on the Glasgow Outcome
Scale (GOS), 6 months after the injury, categorised as death,
persistent vegetative state or severe disability versus moderate
disability or good recovery. The GOS reflects disability, is the
most widely used outcome measure for assessment after trau-
matic brain injury, and allows different groups of patients to be
compared simply and has been recommended as a measure of
outcome for clinical trials. Secondary outcomes included dur-
ation of stay in hospital and the intensive care unit, 30-day mor-
tality and rates of hypotension (defined as systolic
BP<90 mm Hg) and hypoxia (defined as SpO₂<92%) at first
contact compared with hospital arrival.

Statistical methods
A sample size of 510 patients (255 per group) who met criteria for
primary outcome analysis (GCS<9 on first contact) would provide
80% power to detect a relative increase of at least 23% in the cat-
egories, moderate disability/good recovery of the GOS score,
assuming a 5% level of significance and a two-tailed comparison.
This sample size was based on an ordinal scale, assuming a pro-
portion of 52% in the death, persistent vegetative state or severe
disability category (31%, 2% and 19%, respectively) and 30% and
18% in the moderate disability and good recovery categories,
respectively. A 5% rate of non-compliance was assumed as this was
the historical rate of physician intervention prior to commence-
ment of the trial. Analysis was performed using SAS V9.3.

The effect of intervention within the mechanism of injury
groups was the only planned subgroup analysis. The primary
analysis was conducted according to the intention-to-treat prin-

Patient safety and ethical considerations
Approval for the trial was granted by the Western Sydney Area
Health Service Human Research Ethics Committee (approval
number HS/TG HREC2002/12/4.25(1530)). Due to the nature of

prehospital care


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the intervention, patient consent could not be obtained prior to randomisation. The study, however, complied with the Australian National Health and Medical Research Council criteria for waiver of consent for highly dependent patients and the Human Research Ethics Committee approved the study on this basis. Retrospective consent was obtained from surviving patients for subsequent functional outcome assessment. Study safety was monitored by an independent data and safety monitoring committee, which maintained oversight of trial conduct, patient safety and the two planned interim analyses. Funding was provided by Insurance Australia Group, and the New South Wales Motor Accident Authority.

RESULTS

Patient recruitment

A total of 3124 incidents were randomised between May 2005 and March 2011, from which 3696 patients were identified. The last patient outcome assessment was performed in October 2011. As intervention allocation was required before the level of consciousness was confirmed, many subjects did not meet the criteria for severe head injury when subsequently assessed.

Figure 1 Randomisation, enrolment and outcome data for the original criteria subgroup. Between 14 May 2005 and 14 March 2011, the study randomised 3124 incidents yielding 3696 identifiable patients. Of these, 375 patients met the original criteria for severe head injury.

Figure 2 Randomisation, enrolment and outcome data for the modified criteria subgroup. Between 14 May 2005 and 14 March 2011, the study randomised 3124 incidents yielding 3696 identifiable patients. Of these, 338 patients met the modified criteria for severe head injury.
the scene. This resulted in a much smaller group of patients with severe head injury than the number randomised. Three hundred and seventy-five patients met the original criteria (figure 1) and 338 patients met the modified criteria (figure 2). The Data and Safety Monitoring Committee met twice during this period to consider safety data and on both occasions deemed that there were no safety concerns. No adverse patient events due to enrolment in the trial were reported.

**Compliance and study discontinuation**

The study was discontinued before the target 510 patients had been accrued due to poor recruitment and a greater non-compliance rate than had been planned for in the sample size calculation. The high non-compliance rate was driven by changes in local policy for the standard care group, and the study management committee considered it likely that policy changes by the New South Wales Ambulance would further increase this rate. The numbers of non-compliant patients in each arm by both the original and modified definitions are detailed in figures 1 and 2. In the standard care arm, New South Wales Ambulance physician teams were dispatched to incidents by either road or helicopter depending on proximity to their operations base. In the intervention arm, patients were not treated by physicians when either the road paramedics removed the patient from the scene before the physician team could arrive or they cancelled the physician team in cases where the patient would otherwise have qualified for inclusion.

**Patient characteristics, prehospital intervals and intervention rates**

For both the modified and the original criteria, patients were similar with respect to baseline characteristics (table 1). Prehospital treatment and transport times, rates of physician-only intervention and correction rates of prehospital hypoxia and hypotension are presented in table 2. For both the modified and original criteria, the median time spent at scene was 6 min longer for patients who were allocated to physician intervention (p=0.02 and p<0.01, respectively), although the total prehospital time was significantly longer only in the original definition group. Rates of physician-only interventions were significantly higher in the group allocated to physician care, with occurrence of such interventions in the paramedic only allocation group reflecting the non-compliance rate.

### Table 1 Patient characteristics with patients categorised by modified and original severe head injury criteria, by treatment allocation

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Original head injury definition (GCS&lt;9)</th>
<th>Modified head injury definition (GCS ≥ 3 and AIS ≥ 3)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>178</td>
<td>156</td>
</tr>
<tr>
<td>Age at randomisation (years)</td>
<td>43 (26–55)</td>
<td>45 (26–60)</td>
</tr>
<tr>
<td>Male</td>
<td>124 (70%)</td>
<td>111 (71%)</td>
</tr>
<tr>
<td>At least one comorbidity</td>
<td>74 (50%)</td>
<td>59 (45%)</td>
</tr>
<tr>
<td>Comorbidities unknown</td>
<td>32 (18%)</td>
<td>25 (16%)</td>
</tr>
<tr>
<td>Warfarin</td>
<td>1 (1%)</td>
<td>1 (1%)</td>
</tr>
<tr>
<td>Mechanism of injury</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Transport-related</td>
<td>97 (58%)</td>
<td>89 (60%)</td>
</tr>
<tr>
<td>Fall</td>
<td>38 (23%)</td>
<td>38 (26%)</td>
</tr>
<tr>
<td>Other</td>
<td>33 (20%)</td>
<td>21 (14%)</td>
</tr>
<tr>
<td>Pupil reactivity</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nil reactive</td>
<td>58 (33%)</td>
<td>44 (28%)</td>
</tr>
<tr>
<td>One reactive</td>
<td>16 (9%)</td>
<td>16 (10%)</td>
</tr>
<tr>
<td>Both reactive</td>
<td>104 (58%)</td>
<td>96 (62%)</td>
</tr>
<tr>
<td>Injury measures</td>
<td></td>
<td></td>
</tr>
<tr>
<td>First GCS</td>
<td>3 (3–6)</td>
<td>6 (3–9)</td>
</tr>
<tr>
<td>First systolic BP (mm Hg)</td>
<td>100 (60–125)</td>
<td>111 (80–140)</td>
</tr>
<tr>
<td>Revised trauma score</td>
<td>4.1 (2.9–6.0)</td>
<td>5.0 (4.1–6.2)</td>
</tr>
<tr>
<td>Injury severity score*</td>
<td>24 (6–34)</td>
<td>29 (20–38)</td>
</tr>
<tr>
<td>New injury severity score*</td>
<td>27 (7–48)</td>
<td>41 (27–57)</td>
</tr>
<tr>
<td>Head AIS score*</td>
<td>3 (2–5)</td>
<td>4 (3–5)</td>
</tr>
<tr>
<td>Type of injury—Marshall classification of first CT scan, 10</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diffuse I</td>
<td>52 (40%)</td>
<td>23 (17%)</td>
</tr>
<tr>
<td>Diffuse II</td>
<td>45 (35%)</td>
<td>74 (55%)</td>
</tr>
<tr>
<td>Diffuse III</td>
<td>26 (20%)</td>
<td>30 (22%)</td>
</tr>
<tr>
<td>Diffuse IV</td>
<td>6 (5%)</td>
<td>7 (5%)</td>
</tr>
<tr>
<td>Extradural haematoma</td>
<td>5 (4%)</td>
<td>11 (8%)</td>
</tr>
<tr>
<td>Subdural haematoma</td>
<td>30 (22%)</td>
<td>47 (34%)</td>
</tr>
<tr>
<td>Other haematoma</td>
<td>3 (2%)</td>
<td>4 (3%)</td>
</tr>
<tr>
<td>Traumatic subarachnoid haemorrhage</td>
<td>60 (34%)</td>
<td>84 (54%)</td>
</tr>
<tr>
<td>Base-of-skull fracture</td>
<td>38 (21%)</td>
<td>50 (32%)</td>
</tr>
</tbody>
</table>

Statistics are given as n (%) or median (IQR).

*2005 definition.

AIS, Abbreviated Injury Scale.
Intention-to-treat analyses

The numbers of patients in each subgroup that had their outcomes determined by record review as they were lost to follow-up are detailed in figures 1 and 2. In the intention-to-treat analyses of patients who met the modified criteria, there was no statistically significant difference in the primary endpoint of the GOS score at 6 months post injury (OR 1.14; 95% CI 0.73 to 1.75, p=0.56; figure 3). The results were similar when the analysis was restricted to patients alive at 6 months. There was also no difference in the 30-day mortality rate (OR 1.05, 95% CI 0.66 to 1.66, p=0.84). There was no difference between the groups in the total days in intensive care or hospital stay (table 3). Patients who met the original criteria showed similar results. When the effect of the intervention within injury mechanism subgroups was examined, there was no difference between the groups (figure 3). The tests of interaction were not significant.

Per protocol and as-treated analyses

Separate analyses for those patients who received intervention, according to the protocol (ie, excluding those who did not receive the allocated intervention; ‘compliant group’); who

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### Table 2

Prehospital times, transport mode and physician-only interventions, by treatment allocation

<table>
<thead>
<tr>
<th>Factor</th>
<th>Original head injury definition (GCS&lt;9)</th>
<th>Modified severe head injury definition (GCS 3–12 and AIS≥3)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Standard care (nN (%))</td>
<td>Physician care (nN (%))</td>
</tr>
<tr>
<td>Time intervals (min)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Time from beginning of emergency call to first arrival at patient</td>
<td>9 (7–13) 103 (53)</td>
<td>10 (8–13) 106 (53)</td>
</tr>
<tr>
<td>Time spent at the scene</td>
<td>16 (11–27) 103 (53)</td>
<td>22 (15–33) 106 (53)</td>
</tr>
<tr>
<td>Transport time from scene to hospital</td>
<td>13 (10–21) 60 (33)</td>
<td>13 (9–18) 63 (33)</td>
</tr>
<tr>
<td>Total prehospital time</td>
<td>44 (33–56) 22 (15–33) 90 (49)</td>
<td>48 (38–62) 24 (18–33) 92 (50)</td>
</tr>
<tr>
<td>Patient transport mode</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Helicopter transport</td>
<td>21 (12%) 90 (49)</td>
<td>36 (19%) 92 (50)</td>
</tr>
<tr>
<td>Physician-only interventions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neuromuscular-blockade-assisted intubation</td>
<td>18 (10%) 60 (33)</td>
<td>96 (49%) 63 (33)</td>
</tr>
<tr>
<td>Tube/open thoracostomy</td>
<td>5 (3%) 1 (1%)</td>
<td>5 (3%) 1 (1%)</td>
</tr>
<tr>
<td>Crystalloid &gt;1000 mL</td>
<td>12 (7%) 60 (33)</td>
<td>24 (12%) 63 (33)</td>
</tr>
<tr>
<td>Any Packed Red Blood Cells</td>
<td>6 (3%) 3 (2%)</td>
<td>12 (6%) 6 (2%)</td>
</tr>
<tr>
<td>Any 7.5% saline or 20% mannitol</td>
<td>5 (3%) 29 (15%)</td>
<td>29 (15%) 100 (50)</td>
</tr>
<tr>
<td>Rates of correction or prevention of hypoxia and hypotension</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypoxia at first contact</td>
<td>30 (24%) 60 (33)</td>
<td>37 (22%) 63 (33)</td>
</tr>
<tr>
<td>Hypoxia at hospital arrival</td>
<td>14 (14%) 60 (33)</td>
<td>5 (4%) 63 (33)</td>
</tr>
<tr>
<td>Hypotension at first contact</td>
<td>58 (34%) 60 (33)</td>
<td>63 (32%) 63 (33)</td>
</tr>
<tr>
<td>Hypotension at hospital arrival</td>
<td>13 (7%) 60 (33)</td>
<td>13 (7%) 63 (33)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
| Statistics are given as n (%) or median (IQR). Hypoxia, systolic BP <90 mm Hg; Hypoxia, SpO2 <92%.
| *The Wilcoxon rank-sum test was used for continuous data and χ² tests were used for categorical variables.
| †Conditional binomial test.
| AIS, Abbreviated Injury Scale.

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**Figure 3** Main intention-to-treat results by treatment allocated and mechanisms subgroups. OR>1 indicates higher risk in the physician treated group.
received the intervention to which they were not allocated ('non-compliant group'); and by intervention actually received regardless of allocation ('as-treated group') are presented in figure 4. In the original definition group, there were more deaths at 30 days in non-compliant (p=0.003) and as-treated patients (p=0.005) who received standard care. Those receiving additional physician care had a 16% decrease in mortality representing one fatality prevented for every six patients treated. There were also more patient deaths at 30 days in the modified definition non-compliant group receiving standard care where additional physician care was associated with a 7% mortality decrease needing 14 patients to be treated to prevent one fatality (p=0.04).

**DISCUSSION**

We were unable to demonstrate a significant difference in the primary outcome measure between treatment groups in an intention-to-treat analysis in this study. Prespecified as-treated analyses indicated a potential mortality benefit in patients with GCS<9 receiving physician intervention with the number needed to treat to prevent one fatality being six in this group. Mortality analysis of non-compliant patients in the modified definition group also just reached statistical significance, although this group is the most likely to be biased by selection. The intention-to-treat analysis result could have been due to the high non-compliance rates combined with a smaller than planned sample size. Increasing emphasis by the Ambulance Service on dispatch of physician teams to patients who were considered likely to have severe injuries resulted in non-compliance rates that were much higher than planned and contributed to early termination of recruitment.

Previous cohort studies examining the effectiveness of prehospital physician care compared with paramedic only care for severe head injury patients demonstrated an improvement in outcome associated with physician care.1 11–14 This is the first randomised trial designed to confirm these findings but faced numerous challenges in both conduct and logistical implementation. Although unable to confirm the results of previous studies, the trial was successful in achieving funding to study this high-cost intervention which is of considerable clinical and public health interest. However, Ambulance Service policy changes in introduction of a new proactive tasking system to identify patients for physician team response during trial recruitment resulted in higher than historical rates of dispatch of physicians to patients allocated to paramedic only care.

Recruitment was also slower than planned. Attempts were made to increase the rate by adopting Night Vision Goggle technology 3 years into the trial enabling 12 h of availability to recruit per day in both summer and winter. Extending to more than 12 h per day was not possible due to pilot duty time and funding limitations. Extension of the recruitment area into an adjacent regional area was also explored but excluded due to New South Wales Ambulance tasking policies which would have resulted in dispatch of physician teams to all patients randomised to paramedic only care in this area.

The primary analysis subgroup was modified after the first interim analysis, as some patients who met the original definition were found to not have significant traumatic brain injury. The modified definition including the Abbreviated Injury Scale score was used to retrospectively classify patients into the severe head injury group. The original definition of severe injury which is based only on the GCS is measurable at the incident scene and can be used to allocate patients to treatment groups regimens at the time of prehospital assessment. There was a 16% reduction in mortality in this group on as-treated analysis. Low GCSs can be secondary to factors such as hypotension or hypoxia and brain injury. It is possible that physiological derangement defines the group of patients who are most likely to benefit from advanced prehospital interventions rather than brain injury itself.15 Additionally, the Abbreviated Injury Scale is determined after treatments that may modify the score, so it may not be surprising that patients retrospectively selected from

**Prehospital care**

<table>
<thead>
<tr>
<th>Table 3</th>
<th>Days in ICU and length of hospital stay for the modified and original head injury definition groups</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Original head injury definition</strong> (GCS&lt;9)</td>
<td><strong>Modified severe head injury definition (GCS 3–12 and AIS≥3)</strong></td>
</tr>
<tr>
<td>Standard care</td>
<td>Physician care</td>
</tr>
<tr>
<td>Days in ICU</td>
<td>7 (3–16)</td>
</tr>
<tr>
<td>Days in hospital</td>
<td>5 (0–26)</td>
</tr>
</tbody>
</table>

Statistics given as median (IQR). AIS, Abbreviated Injury Scale; ICU, intensive care unit.

**Figure 4** Sensitivity analyses for the primary outcome and mortality. OR>1 indicates higher risk in the physician treated group. Three missing 6-month outcomes in the Original Head Injury Criteria group (one compliant patient and two non-compliant patients).


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a larger data set on the basis of similar Abbreviated Injury Scale scores were also similar on other post-treatment measures such as the GOS scores. This study specifically addressed the issue of advanced prehospital interventions provided by physician teams. In other jurisdictions, advanced interventions may be provided by non-physician teams such as paramedics and nurses with advanced skill sets. The present study was not designed to compare these treatment models. The results of this trial, however, combined with the previous cohort studies on this subject, indicate that a trial examining the benefit of physician prehospital intervention on mortality in unconscious patients (GCS<9) at the accident scene is now needed.

CONCLUSION
This randomised study of adult blunt injury suggests a reduction in 30-day mortality in patients with blunt trauma with GCS<9 receiving physician prehospital care. Changes in ambulance service policies resulted in high non-compliance rates and difficulty continuing recruitment to the planned sample size in this study making interpretation of the intention-to-treat analyses problematic. Nevertheless, the sensitivity analyses suggest a clinically significant mortality reduction in patients with GCS<9, a finding which should be confirmed by further studies that address non-compliance issues.

Correction notice This article has been corrected since it was published Online First. The provenance and peer review statement has been corrected.

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Contributors AAG: Study guarantor, literature search, study design, data interpretation, and writing of the manuscript. KPM: data analysis, revision of the manuscript. MF: study design, data collection, and revision of the manuscript. EP: data collection, revision of the manuscript. VG: study design, data analysis and interpretation, writing of the manuscript.

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Competing interests All authors have completed the Unified Competing Interests form at http://www.icmje.org/coiDisclosure.pdf (available on request from the corresponding author) and declare that (1) AG has support from CareFlight for the submitted work; (2) AG, KM, MF, EP and VG have no relationships with companies that might have an interest in the submitted work in the previous 3 years; (3) their spouses, partners or children have no financial relationships that may be relevant to the submitted work and (4) AG, KM, MF, EP and VG have no non-financial interests that may be relevant to the submitted work. AG affirms that the manuscript is an honest, accurate and transparent account of the study being reported; that no important aspects of the study have been omitted and that any discrepancies from the study as planned and registered have been explained. All authors, external and internal, had full access to all of the data (including statistical reports and tables) in the study and can take responsibility for the integrity of the data and the accuracy of the data analysis. All researchers are independent of the funding agencies (Insurance Australia Group and the NSW Motor Accidents Authority). AG and EP are employed by CareFlight, a not-for-profit aeromedical service provider who contributed operational infrastructure for the trial. The study funders had no role in study design; in the collection, analysis and interpretation of data; in the writing of the report and in the decision to submit the article for publication. AAG reports grants from Insurance Australia Group, grants from NSW Motor Accident Authority, during the conduct of the study and Medical director of Careflight, a not-for-profit company that provides aeromedical services.

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