REFINE (REducing Falls in In-patieNt Elderly) using bed and bedside chair pressure sensors linked to radio-pagers in acute hospital care: a randomised controlled trial

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Abstract

Background: falls in hospitals are a major problem and contribute to substantial healthcare burden. Advances in sensor technology afford innovative approaches to reducing falls in acute hospital care. However, whether these are clinically effective and cost effective in the UK setting has not been evaluated.

Methods: pragmatic, parallel-arm, individual randomised controlled trial of bed and bedside chair pressure sensors using radio-pagers (intervention group) compared with standard care (control group) in elderly patients admitted to acute, general medical wards, in a large UK teaching hospital. Primary outcome measure number of in-patient bedside falls per 1,000 bed days.

Results: 1,839 participants were randomised (918 to the intervention group and 921 to the control group). There were 85 bedside falls (65 fallers) in the intervention group, falls rate 8.71 per 1,000 bed days compared with 83 bedside falls (64 fallers) in the control group, falls rate 9.84 per 1,000 bed days (adjusted incidence rate ratio, 0.90; 95% confidence interval [CI], 0.66–1.22; \( P = 0.51 \)). There was no significant difference between the two groups with respect to time to first bedside fall (adjusted hazard ratio (HR), 0.95; 95% CI: 0.67–1.34; \( P = 0.12 \)). The mean cost per patient in the intervention group was £7199 compared with £6400 in the control group, mean difference in QALYs per patient, 0.0001 (95% CI: −0.0006 – 0.0004, \( P = 0.67 \)).

Conclusions: bed and bedside chair pressure sensors as a single intervention strategy do not reduce in-patient bedside falls, time to first bedside fall and are not cost-effective in elderly patients in acute, general medical wards in the UK.

Trial registration: isrctn.org identifier: ISRCTN44972300.

Keywords: in-patient falls, falls, bed sensors, older people

Introduction

Falls in hospitals are a major problem and contribute to substantial healthcare burden [1, 2]. Across England and Wales, \(~152,000\) falls are reported in acute hospitals every year [3]. They result in physical and psychological morbidity for patients and large healthcare costs, including the costs of treating injuries, increased hospital stay, complaints and litigation [4–6]. Elderly patients admitted acutely to hospital are particularly vulnerable to falling [2], with more than half of these falls occurring at the bedside [3].

Trials of single interventions to prevent falls in acute hospital care have not demonstrated a convincing reduction in falls [7–10]. Multi-factorial interventions have been
investigated with mixed results [11–14]. A Cochrane review concluded that although multi-factorial interventions prevent falls in hospitals, no specific recommendations could be made regarding the effective components of these interventions [15]. Additionally, multi-factorial interventions can be difficult and costly to implement [16]. Therefore, there is a need to develop and evaluate effective single intervention strategies in acute hospital care.

Advances in assistive technology such as bed and bedside chair pressure sensors afford innovative approaches to reducing in-patient bedside falls [17]. These are increasingly being used in healthcare facilities [18], and have been endorsed by the Joint Commission on Accreditation of Healthcare Organisations [19]. However, a recent cluster randomised trial in the USA found no reduction in bedside falls using these sensors [10]. Whether these findings are valid in other settings and whether using a more advanced radio-pager system is clinically effective and cost effective is unknown. We report the results of a large, pragmatic, parallel arm, randomised controlled trial of bed and bedside chair sensors using radio-pagers to reduce in-patient bedside falls in acute, general medical, elderly care wards in a UK hospital.

Methods

Participants

All patients admitted to the hospital from the medical admissions unit to three acute, general medical elderly care wards (within 24 h) at the Queen’s Medical Centre (1800 beds, serving a population of 680,000), Nottingham, UK were eligible for inclusion into the trial. Participants were individually randomised to receive either a bed and bedside chair pressure sensor linked to a radio-pager (intervention group) or no sensor (control group), for the duration of their admission on the elderly care ward. Exclusion criteria were being permanently bed bound prior to admission, moribund/unconscious, on end of life care or previous inclusion in the trial on an earlier admission. Recruitment occurred between January 2009 and March 2011, with the final patient being discharged 29 March 2011. All patients were asked to give written, informed consent. In those who lacked capacity, subjects were recruited in consultation with family or professional consultees, in keeping with the research provisions of the Mental Capacity Act of England, and as approved by a local Research Ethics Committee.

Randomisation

Subjects were randomised to the intervention group or the control group using a web based randomisation service provided by the Clinical Trials Support Unit, University of Nottingham. The allocation schedule was generated using random permuted blocks of randomly varying size.

Blinding

It was not possible to blind participants or those providing medical or nursing care to the intervention or control group allocation. A feasibility study [20], prior to this research demonstrated that nurses were able to identity which patients had the dummy sensors and which the active sensors by the end of a single nursing shift. Data were extracted from incident forms and analysed by members of the research team, blind to intervention allocation.

Withdrawals

Participants were free to withdraw from the trial at any stage. Data were included in the analyses up to the point of withdrawal. Those who were deemed to be at risk of harm resulting from the sensor equipment (e.g. in cases of confusion where participant mis-handled equipment) had the sensor removed but data collection continued and these participants were included in the analysis in the group to which they had been randomised.

Intervention

The sensor units were leased from an independent manufacturer and consisted of a battery-operated bed and bedside chair pressure sensor linked wirelessly to a handheld battery-operated radio-pager (details of the devices have been previously described [20]). When a patient left the bed or bedside chair, a radio signal alert was transmitted from a transmitter box attached to the foot of the patient’s bed, to the radio-pager carried by a member of the nursing team, which provided the location of the participant. An absence of pressure on the sensor of 5 s or more triggered an alert. A central receiver on each ward recorded all alerts, which were collected by the research team.

The research team fitted the sensor units, reviewed the equipment daily, replaced batteries as required and recorded sensor unit problems, using personal diaries. The pagers were carried, where possible by the nursing aides, who were more flexible in their daily duties (i.e. less likely to be involved in medication rounds, interviews with relatives). The total registered nurse and nurse aide allocation per 28-bedded ward was eight on the morning shift (7 a.m.–2 p.m.), six in the afternoon shift (2 p.m.–7 p.m.) and four overnight (7 p.m.–7 a.m.), with each shift including at least two nursing aides. No ward had more than 10 units linked to two pagers in operation at any one time, therefore the most a single registered nurse or nurse aide would be responsible for was five sensors at any one time. Face-to-face training based on our pilot study [17] was undertaken with the ward staff. New ward staff members were trained as necessary and refresher demonstrations provided monthly.

Standard care for both groups was provided by a geriatric ward-based team comprising geriatricians, nurses,
occupational therapists and physiotherapists delivering routine geriatric medical care.

**Outcome measures**

As the aim of providing bed and bedside chair sensors is to reduce bedside falls, the primary outcome measure was the number of in-patient bedside falls per 1,000 bed days from time of randomisation to date of discharge, death or study withdrawal, whichever occurred sooner. A bedside fall was defined as an unexpected event in which the participant came to rest on the ground, floor or lower level in the area around the bedside, with the bedside being defined as the area encompassed by the curtained area surrounding the bed. For patients in side rooms, the bedside was defined as the area of the room.

Bedside falls were ascertained from incident reporting forms, completed by the ward clinical teams, the use of which was mandatory within the hospital and enforced by systematic quality assurance processes (clinical governance) used in UK hospitals. These forms included details of the fall event, time, injuries sustained and subsequent actions taken. The forms were collected from participating wards each day by the research team, blind to group allocation.

Secondary outcome measures were:

- Number of injurious in-patient bedside falls per 1,000 bed days, defined as falls resulting in abrasion, bruise, swelling, cut, laceration, dislocation, fracture or muscle strain.
- Activities of daily living, measured using the Barthel ADL Index [21].
- Fear of falling, measured using the modified falls efficacy scale (MFES) [22].
- Length of hospital stay (number of days from admission to discharge or death, whichever occurred sooner)
- Residential status on discharge (discharged to same address as on admission or discharged to another address).
- Health related quality of life, measured using the EQ 5D questionnaire [23].

**Statistical analysis**

With 905 patients in each arm, the study had 80% power (with \( \alpha = 0.05 \)) to detect a 35% reduction in the rate of bedside falls among the intervention group, assuming a bedside falls rate of 8.0 per 1,000 bed days in the control group and an over dispersion parameter of 1.5 (to allow for non-independence of falls within individuals). The bedside falls rate was based on an anticipated mean number of falls per hospital admission of 0.25 and an average length of stay of 19 days. The 35% reduction was judged to be clinically meaningful and based on data from our previous 12-month pilot study [17]. This was a two-year (12-month pre-intervention observational study \( n = 209 \) followed by a 12-month intervention study \( n = 153 \)) conducted over a single ward, with bedside falls rates collected by an independent researcher from incident reporting forms, completed by the ward clinical staff.

Analyses were carried out on the basis of intention to treat. Where patients died in hospital or withdrew consent, follow-up was censored at the date of withdrawal/death with any events occurring before this date included in the analysis. On all other occasions, follow-up ended on the date of discharge (Supplementary data are available in *Age and Ageing* online, Appendix A).

Further details on project management, governance, confidentiality and administration may be found in the previously published protocol [20].

**Results**

**Subjects**

The flow of participants through the trial is shown in Figure 1. Of the 3,292 eligible patients, 1,839 were randomised (918 to the intervention group and 921 to the control group). Baseline characteristics are shown in Supplementary data available in *Age and Ageing* online, Appendix B, Table S1. The mean age at randomisation was 84.6 years (range 61–103 years), with a slight predominance of females (55% of total). 65.1% of the subjects had a median MMSE <23/30 and the median Barthel score was 12. The groups appeared well balanced at baseline.

**Outcome measures**

There were 85 bedside falls (65 fallers) in the intervention group, falls rate 8.71 per 1,000 bed days compared with 83 bedside falls (64 fallers) in the control group, falls rate 9.84 per 1,000 bed days, adjusted incident rate ratio 0.90 (95% CI: 0.66–1.22; \( P = 0.50 \)). The rate of minor injuries is shown in Table 1. Sixteen bedside falls in the control group resulted in minor injury (6 bruises, 5 abrasions and 5 lacerations) and 24 in the intervention arm (4 bruises, 5 abrasions and 15 lacerations), (adjusted IRR, 1.60; 95% CI: 0.83–3.08; \( P = 0.15 \)). The number of major injuries was small (three fractures in the control group and two in the intervention group).

There was no significant difference between the two groups with respect to time to first bedside fall, adjusted HR, 0.95; 95% CI: 0.67–1.34; \( P = 0.12 \). Among those who fell, the median time between randomisation and first bedside fall was 7 days in the intervention group and 6 days in the control group. Stratifying by follow-up time showed a non-significant trend towards a greater reduction in early bedside falls risk, with the HR over the first 2 days being 0.60; 95% CI: 0.30–1.20; \( P = 0.11 \) compared with HR, 1.36; 95% CI: 0.82–2.23; \( P = 0.15 \), after 5 days.

None of the secondary outcomes differed significantly between the two groups (Table 2). Among patients who experienced one or more bedside fall \( n = 127 \), the length of stay was significantly longer, median 20 days, IQR 12–31 days than for those who did not fall at the bedside, median 9 days, IQR 5–15 days, \( P < 0.001 \).
The mean cost per patient in the intervention group was £7199 compared with £6400 in the control group (Supplementary data are available in *Age and Ageing* online, Appendix C, Table S4). The mean difference in QALYs per patient, adjusting for baseline, was 0.0001 (95% CI: −0.0006 to 0.0004; \( P = 0.67 \)).

**System problems**

There were a total of 120 problems with sensor system functioning recorded between January 2009 and February 2011. Forty of the records (33%) were classified as de-functioning of the equipment (sensors disconnected, batteries removed, transmitter box power cut), 51 (43%) as pager faults (combination of problems including damage or lost) and 29 (24%) instances where working pagers found unattended (lying on desks or in drawers, taken home).

**Discussion**

The use of bed and bedside chair pressure sensors using radio-pagers did not significantly reduce bedside falls, time to first bedside fall or prove cost effective, in elderly patients admitted to acute, general medical wards in the UK.
There are a number of limitations in our study that need to be recognised. Our study was powered to detect a 35% reduction in the rate of bedside falls, based on the sample size estimates from our pilot study [17]. It is possible that the intervention may be associated with a smaller reduction in bedside falls, which may have been missed. Secondly, we collected our primary outcome data from the mandatory incident reporting system operating within the hospital and it is recognised that under-reporting of falls may occur within such systems [24]. Another limitation was blindness of the staff to the intervention, which may have led to differential reporting of falls between groups, however unless this occurred to a very large extent, it is unlikely to explain our findings of no effect.

Our findings confirm those from a previous small non-randomised study [7] and a recent large cluster randomised RCT [10], extending the generalisability of the findings from previous studies to the UK healthcare setting. This study also adds evidence that the use of radio-paging systems to alert nurses does not enhance the effectiveness of the sensor systems. The findings from RCTs of multi-factorial interventions that have included bed sensors are mixed. One study using sensors as part of a multi-factorial intervention found no effect on falls rate [13], whilst a more recent trial using sensors as part of a multi-factorial intervention significantly reduced in-patient falls, although the actual numbers of patients receiving alarms and the type of alarm were not described [14].

The findings from our study and the recent RCT evaluating sensors as a single intervention would both suggest any reductions in falls rates seen with multi-factorial interventions which include bed/chair sensors, may be attributable to elements of the intervention other than bed/chair sensors.

There are several possible explanations for why bed and bedside chair pressure sensors do not appear to reduce in-patient falls. Complex organisational factors including managing organisational change, leadership, commitment and communication may be important in implementing change within healthcare [25, 26] and these may have been underestimated in translating the findings from our single ward pilot study, to the current RCT. It is also possible that although the sensors alerted, nurses were unable to respond to radio-pagers sufficiently quickly to avert falls and this is supported by findings from a recent U.S. study [27] which suggests faster ‘nurse call light response time’ may contribute to lower fall rates. Nurses may also be able to respond more quickly to radio-pagers if alarms were restricted to a smaller number of people, the majority of whom were at high risk of falls.

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Conclusion

In summary, we found that both bed and bedside chair pressure sensors using radio-pagers did not reduce the rate of in-patient bedside falls, time to first bedside fall and are not cost effective in elderly patients in acute, general medical wards in the UK.

Key points

• In-patient falls are a significant problem in acute hospitals.
• In-patient falls lead to significant injury and increased length of stay.
• Bed and bedside pressure sensors using radio-paging as part of a single intervention do not reduce falls rates.

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Conflicts of interest

None declared.

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Supplementary data

Supplementary data mentioned in the text is available to subscribers in Age and Ageing online.

References

Delay between symptom onset and clinic attendance following TIA and minor stroke: the BEATS study

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Abstract

Background: rapid specialist assessment of patients with transient ischaemic attack (TIA) reduces the risk of recurrent stroke. National guidelines advise that high-risk patients are assessed within 24 h and low-risk patients within 7 days.

Aim: to quantify delay and map pathways taken by patients from symptom onset to specialist assessment.

Design: retrospective cohort study.

Setting: rapid access TIA clinic.

Methods: structured interviews with 278 patients newly diagnosed with TIA (222) or minor stroke (56), and examination of medical records.

Results: of the 133 high-risk TIA patients, 11 (8%) attended the clinic within 24 h of symptom onset; of the 89 low-risk TIA patients, 47 (53%) attended within 7 days. Median delay between symptom onset and seeking help from a healthcare professional (HCP) was 4.0 h (IQR 0.5, 41.3). Delay was less if symptoms were correctly interpreted but not reduced by a publicity campaign (FAST) to encourage an urgent response. Most patients (156, 56%) first contacted a general practitioner (GP) and 46 (17%) called an ambulance or attended the emergency department. Over a third (36%) had a second consultation with an HCP before attending the clinic, and this was more likely in those presenting to paramedics, out of hours GP services or optometry. Time to clinic attendance was less if an emergency pathway was used and greater if patients were seen by a second HCP.

Conclusions: factors contributing to delay include incorrect interpretation of symptoms and failure to invoke emergency services. Delays after presentation could be addressed by direct referral by out of hours services, paramedics and optometrists.

Keywords: acute care, emergency medical services, stroke, transient ischaemic attack, general practice, older people

Introduction

Rapid assessment and treatment of patients with transient ischaemic attack (TIA) or minor stroke reduces the risk of early recurrent stroke [1, 2]. The Royal College of Physicians’ Guidelines suggests that TIA patients should be scored using the ABCD² [3, 4]. Those at high risk (score ≥4) should be assessed by a specialist within 24 h of symptom onset, and