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CLINICAL INVESTIGATION

Evaluation of an intervention to reduce tidal volumes in ventilated ICU patients[†]

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Abstract

Background: There is considerable evidence that the use of tidal volumes <6 ml kg $^{-1}$ predicted body weight (PBW) reduces mortality in mechanically ventilated patients. We evaluated the effectiveness of using a large screen displaying delivered tidal volume in ml kg $^{-1}$ (PBW) for reducing tidal volumes.

Methods: We assessed the intervention in two 6-month periods. A qualitative study was undertaken after the intervention period to examine staff interaction with the intervention. The study was conducted in a mixed medical and surgical intensive care unit at University Hospitals Bristol, UK. Consecutive patients requiring controlled mechanical ventilation for more than 1 h were included. Alerts were triggered when tidal volume breached predetermined targets and these alerts were visible to ICU clinicians in real time.

Results: A total of 199 patients with 7640 h of data were observed during the control time period and 249 patients with 10 656 h of data were observed in the intervention period. Time spent with tidal volumes $<6 \,\mathrm{ml\,kg^{-1}\,PBW}$ increased from 17.5 to 28.6% of the period of controlled mechanical ventilation. Time spent with a tidal volume $<8 \,\mathrm{ml\,kg^{-1}\,PBW}$ increased from 60.6 to 73.9%. The screens were acceptable to staff and stimulated an increase in attendance of clinicians at the bedside to adjust ventilators. Conclusions: Changing the format of data and displaying it with real-time alerts reduced delivered tidal volumes. Configuring information in a format more likely to result in desired outcomes has the potential to improve the translation of evidence into practice.

Key words: acute respiratory distress syndrome; behavioural economics; computerised decision support systems; lung protective ventilation; quality improvement

The translation of evidence-based interventions into clinical practice in patients receiving mechanical ventilation remains a challenge. Randomized controlled trials and meta-analyses have shown that the use of lung protective ventilation (LPV)

reduces mortality in patients with acute respiratory distress syndrome (ARDS). $^{1-4}$ A large randomized controlled trial demonstrated reduced mortality (31% vs 40%) in patients ventilated with low tidal volumes [<6 ml $\rm kg^{-1}$ predicted body weight

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Editor's key points

- The implementation of proven interventions into clinical practice is often delayed.
- This study evaluated the effect on clinicians' behaviour of using a large visual screen to display real-time delivered tidal volumes in ICU patients.
- Use of the real-time displays led to an increase in clinicians' adjustments to ventilator settings at the bedside. Consequently, compliance with recommended standards
- These data suggest that simple changes in the way in which information is displayed to clinicians can change their

(PBW)] compared with patients ventilated with higher tidal volumes (>12 ml kg⁻¹ PBW).² More recently, several trials have demonstrated that the use of lower tidal volumes benefits patients both with and without ARDS at the onset of mechanical ventilation.5-11

Although reducing tidal volumes remains one of the few proven interventions to reduce mortality in mechanically ventilated patients, this intervention is inconsistently applied to patients who may benefit. 12-15 A recent observational study on patients with ARDS undertaken in four academic teaching hospitals found that only 41% of eligible ventilator settings were adherent to LPV criteria and 37% of patients never received LPV. 16 In the control group of the High Frequency OSCillation in ARDS (OSCAR) trial undertaken in patients with ARDS in UK intensive care units (ICUs), patients were ventilated with an average tidal volume of 8.3 ml kg⁻¹ PBW for the first 3 days after enrolment despite clinicians being encouraged to use LPV. 17

Difficulty diagnosing ARDS; lack of education; lack of a protocol; concerns over hypercarbia, acidosis, and hypoxemia; and physician perceptions of the contraindications to LPV have all been suggested as reasons for the underuse of LPV. 14 18-21 Interestingly, although physicians may document an intention to use LPV and state that they deliver it frequently or always, they still often fail to deliver it to their patients. 19 22 Failure to implement LPV is clearly not just a failure of intent.

To test whether the way clinical information is presented to staff can improve the compliance with low tidal volume ventilation, we deployed two large screens that display delivered tidal volume in the format of ml kg-1 PBW at either end of our ICU. Information was displayed for all patients in the ICU with values derived from the clinical information system (CIS), using realtime alerts when volumes breached 6 and 8 ml kg⁻¹ PBW. A qualitative study of the acceptability and impact of the displays among staff was undertaken after the intervention period in this study was complete.

Methods

We performed a prospective before and after evaluation of the effect of the displays on delivered tidal volume to all patients receiving controlled mechanical ventilation in two distinct 6-month periods in the ICU at University Hospitals Bristol. The first 6-month period (November 1, 2010-April 30, 2011) acted as the control period and during this time there was no access to the alerting displays. In the second 6-month period (November

1, 2011-April 30, 2012), the displays were in use. The intervention period commenced after the displays had been installed and checked over a period of 6 months. No other interventions were introduced between the two time periods. No protocols or quality improvement processes were changed and ventilation equipment was the same in both the control and intervention period.

The ICU at University Hospitals Bristol is a closed-format tertiary medical and surgical ICU with 13 consultants, 10 senior fellows, and 5 junior fellows. Consultants are permanent and fellows rotate through the unit every 3 months. There were two rotations of fellows during each study period. Ventilator settings are predominantly adjusted by the consultants and senior fellows. A small number of experienced nurses also make adjustments to ventilator settings, but they tend to check these with a doctor shortly afterwards. The unit does not employ respiratory therapists. Pressure-controlled ventilation is the preferred method for delivery of controlled mechanical ventilation. The institutional research board classified the study as a service evaluation and waived the requirement for individual patient consent and formal ethical review.

The unit has used the Innovian Solution Suite clinical information system (Dräger, Lübeck, Germany) since 2008. This is an electronic charting system that automatically collects all information relating to patient care, including physiological data, laboratory results, and data from ventilators. This information is displayed on a computerised chart and is also stored on a database (SQL Server 2008; Microsoft, Redmond, WA, USA). The data in the database are available for immediate analysis. An algorithm was constructed to take real-time data from the database and automatically calculate delivered tidal volumes in the format ml kg⁻¹ PBW using the formulas PBW=[height in cm-154]×0.9+50 for males and PBW=[height in cm-154]×0.9+45.5 for females.²³ Staff can access the CIS at the bedside, on central desktop computers, and remotely on computers housed away from the ICU. The display screens described in this study are not visible from within the CIS; they operate completely separately and are mounted on the wall.

Intervention

Two large 48-inch display screens were configured to display a number of metrics derived from the CIS database using freely available reporting software (Visual Studio 2008, Microsoft) (Fig. 1). They were mounted on the wall at either end of the ICU and were visible to most staff working in the unit.

Tidal volume alerts

The screens alerted to increased tidal volume by turning yellow when the tidal volume was ≥ 6 and < 8 ml kg⁻¹ PBW. When the tidal volume was ≥ 8 ml kg⁻¹ PBW the display turned red. The alerts did not cancel until the measured tidal volume returned to <6 ml kg⁻¹ PBW. The alerts were only applied to patients receiving controlled mechanical ventilation. Tidal volumes for patients on spontaneous breathing modes were displayed in ml kg⁻¹ PBW, but without coloured alerts. The information on the screens was refreshed every 5 min. The latest Po, from an arterial blood gas was also displayed on screens among other metrics. If the Po. was >100 mmHg the box turned red. This was the only other metric displayed that had a coloured alert. For the purposes of this study we only analysed the effect of the screens on tidal volume.

All ICU physicians were notified by email that the screens would be introduced at the start of the second study period. Rotating fellows received one standard email when they joined the

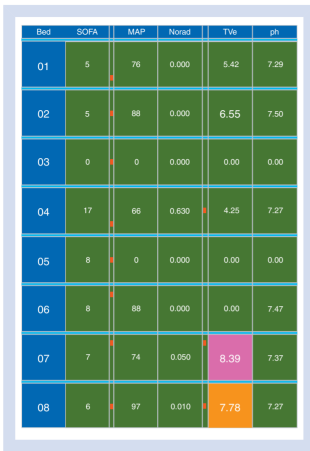


Fig 1 Display screen appearance.

unit informing them of the function of the screens. Nurses received a similar notification via the staff communication book. No other education or information campaign relating to low tidal volume ventilation occurred during the two study periods.

All patients receiving conventional controlled mechanical ventilation for more than 1 h during the two study periods were eligible for inclusion in the study. Patients receiving highfrequency oscillatory ventilation or airway pressure release ventilation were excluded since tidal volume is not directly manipulated in these modes.

Data collection

Data were collected on the CIS database. The configuration of the displays did not alter the format of the database and data were collected in the same way for both the control and intervention period. The tidal volumes for every hour of controlled mechanical ventilation were available for analysis; tidal volumes when patients were on spontaneous modes were excluded. Data were extracted for tidal volume (ml kg-1 PBW), patients' age (categorized as <50, 50-<60, 60-<70 and ≥70 years), sex, Acute Physiology and Chronic Health Evaluation (APACHE) II score, Sequential Organ Failure Assessment (SOFA) score, peak inspired pressure (PIP) set (in mbar), positive end expiratory pressure (PEEP) (in mbar), pH (multiplied by 100), day of the week (classified into hours falling on weekdays and hours falling on weekends), time of day (8 AM-6:59 PM and 7 PM-7:59 AM), and patient status (improved or died at ICU discharge).

Statistical methods

Mixed effects logistic regression models that take repeated measures within individual patients into account were used to quantify associations of exposures with the binary outcome of a tidal volume <6 ml kg⁻¹ PBW (i.e. consistent with LPV and coded 1 for the analysis). The first model included all variables that either stayed constant for each patient or were measured for every hour of data collection. Unadjusted and fully adjusted odds ratios (ORs) and 95% CIs are presented. Models were run on the sample with complete data for all variables of interest. A second model, on a reduced sample size, included the same variables as the first model, as well as variables that were only measured sporadically (pH and Pa_{O2}/FI_{O2}). The 3-month moving averages were calculated for mean tidal volume for a time period of 1 year prior to implementation and 1 year after full implementation to account for linear trends in tidal volume with time. The association between the time period (control or dashboard intervention) and patient status was examined using a chi-squared test. All analyses were carried out using Stata version 12 (StataCorp, College Station, TX, USA).

Qualitative analysis of staff interaction with the screens

An interview-based qualitative study was undertaken after the screens had been deployed for 9 months (i.e. 3 months after the intervention period in this study) to assess staff reactions to the intervention. An independent qualitative researcher interviewed a representative sample of staff (consultants, fellows, senior and junior nurses). Interviews were audio recorded and transcribed. A thematic analysis of the interview data was performed by an experienced qualitative researcher using an iterative approach.²⁴

Results

Effect of screens on tidal volume

A total of 199 patients with 7640 h of data were observed during the control time period and 249 patients with 10 656 h of data were observed in the dashboard intervention period. Baseline characteristics were similar between the control and intervention groups (Table 1). Overall, the majority (69%) of patients were male and 24.1 and 31.7% of patients were ≥70 years of age in the control and intervention periods, respectively. The mean APACHE II score was 16.7 (SD 5.5) in the control period and 17.6 (SD 6.0) in the intervention period. The proportion of time spent with a tidal volume <6 ml kg⁻¹ PBW was lower in the control period (17.5%) compared with the dashboard intervention period (28.6%). The proportion of time spent with a tidal volume <8 ml kg⁻¹ PBW was 60.6% in the control period and 73.9% during the intervention period. The mean tidal volume decreased from 7.7 (SD 2.1) to 7.0 (SD 2.0) ml kg^{-1} PBW from the control to the intervention period.

In the fully adjusted model, the dashboard intervention period was associated with an approximate two-fold increased odds of achieving a tidal volume <6 ml kg⁻¹ PBW (OR 1.91; 95% CI 1.29, 2.82; P=0.001; Table 2). The estimate for the dashboard intervention was not substantially changed in the model that also controlled for the sporadically measured variables (OR 2.00; 95% CI 1.18, 3.39; P=0.010; Table 3). Other variables that were associated with achieving a tidal volume <6 ml kg⁻¹ PBW in fully adjusted

Table 1 Study characteristics. APACHE: Acute Physiology and Chronic Health Evaluation; IQR: interquartile range; PEEP: positive end expiratory pressure; PIP: peak inspired pressure; SOFA: Sequential Organ Failure Assessment.

Characteristics by patient	Control period	Dashboard period	
Patients, n	199	249	
Age group (years), n (%)			
<50	47 (23.6)	58 (23.3)	
50-<60	55 (27.6%)	50 (20.1)	
60-<70	49 (24.6)	62 (24.9)	
≥70	48 (24.1)	79 (31.7)	
Sex, n (%)			
Female	62 (31.2)	78 (31.3)	
Male	137 (68.8)	171 (68.7)	
APACHE II score, mean (sp)	16.7 (5.5)	17.6 (6.0)	
Median length of stay, h (IQR)	120 (61–242)	116 (69–233)	
Median length of ventilation, h (IQR)	34 (10–93)	34 (10–84)	
Characteristics by hour of ventilation			
Total hours of ventilation	7640	10 656	
Days of week, h (%)			
Falling on weekdays	5382 (70.4)	7721 (72.5)	
Falling on weekends	2258 (29.6)	2935 (27.5)	
Time of day, h (%)			
8 am-6:59 pm	3333 (43.6)	4553 (42.7)	
7 pm-7:59 am	4307 (56.4)	6103 (57.3)	
SOFA score, n [mean (sd)]	6201 [8.9 (3.3)]	9043 [8.9 (3.9)]	
PIP set, n [mean (sp)]	7640 [23.6 (5.8)]	10 655 [21.0 (5.0)]	
PEEP, n [mean (sD)]	7638 [8.0 (3.0)]	10 655 [8.3 (3.0)]	
pH, n [mean (sD)]	3071 [7.35 (10.8)]	4099 [7.35 (10.8)	
Pa _{O2} /Fi _{O2} , n [mean (sD)]	3050 [212 (108)]	4090 [228 (116)]	
Tidal volume, ml kg ⁻¹ , n [mean (sp)]	7628 [7.7 (2.1)]	10 652 [7.0 (2.0)]	
Tidal volume group, n (%)			
<6	1333 (17.5)	3045 (28.6)	
6 to <7	1510 (19.8)	2677 (25.1)	
7 to <8	1778 (23.3)	2147 (20.2)	
≥8	3007 (39.4)	2783 (26.1)	

models were the PIP set, PEEP and pH. An increase in the proportion of hours with a tidal volume <6 ml kg⁻¹ PBW was seen for the dashboard intervention compared with the control period and was observed across all levels of Pa_{O_2}/FI_{O_2} at the start (Table 4). For example, for patients with a Pa_{O2}/FI_{O2} at the start of <100, 230/1490 (15.4%) of their hours of data achieved a tidal volume of <6 ml kg⁻¹ PBW in the control period compared with 326/ 1252 (26.0%) hours in the intervention period. For patients who had the highest Pa_{O_2}/FI_{O_2} (\geq 300), the percentage of hours with a tidal volume <6 ml kg⁻¹ PBW increased from 23.5% in the control period to 31.0% in the intervention period.

The mean tidal volume dropped after the introduction of the display screens, but appeared to have risen slightly 6 months after full implementation (Fig. 2). There was no evidence that the control/intervention period was associated with patient outcome (chi-square P=0.538).

Factors affecting staff interaction with the screens—qualitative analysis

Four consultants, three fellows, eight nurses, and one physiotherapist were interviewed and their data extracted and analysed. LPV was accepted as a fundamental concept by all staff. The tidal volume alerts were considered helpful. Two clear staff groups with contrasting views emerged; doctors and senior nurses who have responsibility for the entire ICU, and bedside nurses who are primarily responsible for one patient at a time. The screens were used differently by each group. The doctors and senior nurses with responsibility for the whole unit found the screens more useful than did the bedside nurses. They indicated that the most useful aspect of the screens was that they stimulated attendance of either a doctor or senior nurse to the bedside to assess the patient, often leading to a subsequent recalculation of the target tidal volume. Bedside nurses found the display less useful and commented that they would prefer a bedside display of target tidal volume in order to incorporate this into their patient checks. A greater explanation of the rationale for the screens would have been beneficial for the bedside nurses.

Discussion

This study demonstrated an improvement in the delivery of low tidal volumes to patients receiving controlled mechanical ventilation. Time spent at <6 ml kg $^{-1}$ PBW and 8 ml kg $^{-1}$ PBW as a percentage of time on controlled ventilation was increased. The average tidal volume was reduced during the intervention period and the findings were sustained throughout the intervention period. The reduction in tidal volumes was still apparent more than 1 yr after the screens were deployed. Increased deployment of low tidal volumes was achieved independently of any attempt to change the intentions of clinicians to deploy lower tidal volumes. No educational interventions or quality improvement Table 2 Odds ratios for achieving tidal volume <6 ml kg $^{-1}$, from mixed effects logistic regression models. Binary outcome of Tve <6 coded as 1. APACHE: Acute Physiology and Chronic Health Evaluation; OR: odds ratio; PEEP: positive end expiratory pressure; PIP: peak inspired pressure; SOFA: Sequential Organ Failure Assessment.

	Unadjusted (n=305 people, 12 991 h)			Fully adjusted (n=305 people, 12 991 h)		
	OR	95% CI	P	OR	95% CI	P
Time period						
Control (reference group)	1.00		< 0.001	1.00		0.00
Dashboards	2.19	(1.50, 3.20)		1.91	(1.29, 2.82)	
Age, years						
<50 (reference group)	1.00		0.408	1.00		0.30
50-<60	0.93	(0.54, 1.59)		1.03	(0.60, 1.78)	
60-<70	0.81	(0.47, 1.40)		0.82	(0.47, 1.41)	
≥70	0.65	(0.38, 1.11)		0.65	(0.37, 1.12)	
Sex						
Female (reference group)	1.00			1.00		0.08
Male	0.81	(0.54, 1.22)	0.319	0.70	(0.46, 1.05)	
APACHE II (per score unit)	1.01	(0.98, 1.05)	0.412	1.01	(0.97, 1.05)	0.61
SOFA (per score unit)	1.03	(1.01, 1.06)	0.018	1.05	(1.02, 1.08)	0.00
Days of week						
Weekday (reference group)	1.00		0.876	1.00		0.61
Weekend	0.99	(0.88, 1.12)		0.99	(0.86, 1.09)	
Time of day						
8 ам–6:59 рм (reference group)	1.00		0.561	1.00		0.50
7 рм–7:59 ам	0.97	(0.88, 1.07)		0.97	(0.88, 1.07)	
PIP set (per mbar)	0.95	(0.94, 0.96)	< 0.001	0.95	(0.92, 0.95)	< 0.00
PEEP (per mbar)	1.02	(1.00, 1.05)	0.092	1.02	(1.05, 1.11)	< 0.00

Table 3 Odds ratios for achieving tidal volume <6 ml kg^{-1} , from mixed effects logistic regression models including the variables pH and Pa_{O_2}/Fi_{O_2} which were only measured sporadically. *pH×100. APACHE: Acute Physiology and Chronic Health Evaluation; OR: odds ratio; PEEP: positive end expiratory pressure; PIP: peak inspired pressure; SOFA: Sequential Organ Failure Assessment.

	Unadjusted (N=297 people, 5093 h)			Fully adjusted (N=297 people, 5093 h)		
	OR	95% CI	P	OR	95% CI	P
Time period						
Control (reference group)	1.00		< 0.001	1.00		0.010
Dashboards	2.17	(1.44, 3.27)		2.00	(1.18, 3.39)	
Age, years						
<50 (reference group)	1.00		0.251	1.00		0.30
50-<60	0.95	(0.54, 1.68)		0.85	(0.41, 1.74)	
60-<70	0.74	(0.42, 1.33)		0.64	(0.31, 1.33)	
≥70	0.59	(0.33, 1.05)		0.52	(0.25, 1.09)	
Sex						
Female (reference group)	1.00		0.651	1.00		0.22
Male	0.90	(0.58, 1.41)		0.71	(0.41, 1.24)	
APACHE II (per score unit)	1.03	(1.00, 1.07)	0.079	1.03	(0.98, 1.08)	0.22
SOFA (per score unit)	1.03	(1.00, 1.07)	0.087	0.99	(0.95, 1.03)	0.64
Days of week						
Weekday (reference group)	1.00		0.495	1.00		0.22
Weekend	1.07	(0.88, 1.29)		0.88	(0.71, 1.08)	
Time of day						
8 ам–6:59 рм (reference group)	1.00		0.385	1.00		0.98
7 рм–7:59 ам	0.94	(0.8, 1.09)		1.00	(0.85, 1.18)	
PIP set (per mbar)	0.95	(0.93, 0.97)	< 0.001	0.91	(0.89, 0.94)	< 0.00
PEEP (per mbar)	1.04	(1.00, 1.08)	0.040	1.08	(1.03, 1.13)	0.002
pH*	0.913	(0.905, 0.921)	< 0.001	0.899	(0.888, 0.909)	< 0.00
Pa _{O2} /FI _{O2} (per mmHg)	0.999	(0.998, 1.00)	0.098	1.000	(0.999, 1.000)	0.41

initiatives relating to the use of lower tidal volumes were conducted during the study period. The screens were accepted by a cross section of staff and they appear to have stimulated a greater discussion of target tidal volume between doctors and senior nurses and nurses at the bedside.

This study analysed tidal volumes for every consecutive hour of controlled mechanical ventilation applied to all our patients over a total of 12 months and, as such, gives excellent insight into the ventilation practice within our ICU. Most studies examining compliance with low tidal volumes only quote discrete ventilator settings measured at a few predetermined time points during the day. Compliance with low tidal volume ventilation measured in this way is subject to error if clinicians pay more attention to ventilator settings at these time points. An important limitation is that the intervention in this study was not randomized and the before and after effects reported originate from one ICU. As with any before-and-after design, our results may be influenced by other unmeasured factors. Every attempt was

Table 4 Whether Tve <6 ml kg⁻¹ PBW was achieved by Pa_{O₂}/FI_{O₂} at the start of ventilation. Tve, tidal volume (of expired breath)

Pa _{O2} /FI _{O2}	Control pe	eriod	Dashboards period		
at start	Total	n (%) with	Total	n (%) with	
	hours of	Tve <6	hours of	Tve <6	
	data		data		
<100	1490	230 (15.4)	1252	326 (26.0)	
100 to <200	2476	334 (13.5)	3595	950 (26.4)	
200 to <300	1634	311 (19.0)	1909	577 (30.2)	
≥300	1712	403 (23.5)	3050	946 (31.0)	

made to standardise the care delivered in the two time periods analysed and no co-interventions relating to ventilation practice changed during the study. The reduction of tidal volume in the intervention group in this study suggests the need for a larger randomised, multicentre trial, which will now be undertaken.

Compliance with low tidal volume ventilation was low in this study, but it compares reasonably with previous studies. 12 13 The average tidal volume in both groups is substantially lower than that reported in the conventional ventilation group of the OSCAR trial¹⁷ and, as such, the practice observed in this trial is likely to represent that of a well-performing UK ICU. During the period of data collection our ICU used predominantly pressurecontrolled ventilation in which tidal volumes can vary according to changes in lung compliance. It may be easier to control tidal volumes using a volume-controlled mode, but we are aware of no evidence that suggests that volume-controlled ventilation results in better adoption of LPV. The choice of the mode of ventilation deserves further study, and we plan to do this.

Computerized decision support systems have been shown in several studies to improve adherence to LPV protocols when used in conjunction with a CIS. In one study, a ventilator-induced lung injury 'sniffer' was employed to screen the medical record for patients at risk and alert staff via a page when ventilator settings were potentially injurious.²⁵ This led to a significant reduction in the time spent with excessive tidal volumes. A non-rulebased alert within the medical record that reminded staff of the target tidal volume in ml kg⁻¹ PBW was effective at reducing excessive tidal volumes in all patients ventilated for >24 h.26 Retrospective feedback on compliance with LPV in conjunction with an education programme successfully improved compliance with LPV in one study.27

We displayed information regarding tidal volume in ml kg⁻¹ PBW in all patients receiving controlled mechanical ventilation

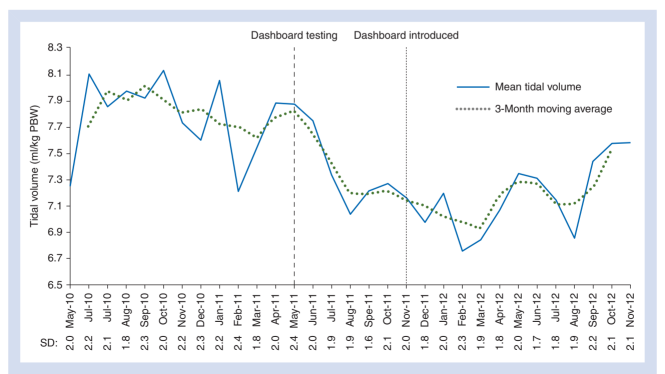


Fig 2 Mean tidal volume in each month and 3-month rolling averages during the periods before dashboards, during testing (May-November 2011), and after the full introduction of the display screens in November 2011.

in real time. This is simpler to configure than systems that attempt to filter for patients with criteria for ARDS, which require free text searches of radiology reports. 25 28 29 The dashboard display is constantly visible, with the advantage that staff don't need to enter individual patient records to ascertain compliance. No education programme was associated with our intervention and our analysis demonstrates the effectiveness of real-time feedback alone in changing practice.

It could be argued that a formal quality improvement process with repeated education, use of champions, repeated meetings, and strategies to share and feed back data would have enhanced compliance with LPV in this study. We deliberately avoided this approach in order to demonstrate the effect of simple changes to the display of information on clinical practice. As such, our study highlights a useful additional strategy to be deployed alongside established quality improvement methodologies. Our qualitative work demonstrated that bedside nurses may have found an education programme useful alongside introduction of the screens, and we will include this in future work.

Several studies have suggested that difficulties with the identification of the subgroup of patients with ARDS is a barrier to implementation of LPV. 14 18 19 We alerted for high tidal volumes in all patients, not just those with ARDS, and this is likely to have contributed to the effect in the intervention group. Interestingly, the severity of lung injury as evidenced by Pao, /FIo, at the start of controlled ventilation was not associated with the increased use of low tidal volumes in our study, although all patients with a Pa_{O2}/FI_{O2}<300 mm Hg benefited from an increased use of low tidal volume ventilation in the intervention group.

Improving the implementation of evidence-based interventions involves changing behaviour and influencing the decisions that clinicians make. Research in the fields of cognitive psychology, neuroscience, and behavioural economics has converged on a dual process model of decision-making.30 Many decisions are rapid, often subconscious, and are heavily influenced by environmental and emotional factors. A minority of decisions are made more deliberately but require a high degree of cognitive effort. Influencing the environment in which decisions are made has the potential to deliver significant behaviour change independent of changing people's intentions.31 32

Simple changes to the choices presented to clinicians can have profound effects on their subsequent decisions. One study found a highly significant difference in the dose of sedation delivered to endoscopy patients, which depended on the volume of pre-filled syringes available to the endoscopist.³³ The screens in our study improved the availability of salient information regarding tidal volume. Calculation of the 'correct' tidal volume for any particular patient requires knowledge of their height, the use of a formula to calculate predicted body weight, and the division of the actual tidal volume by the predicted body weight. This kind of complex cognitive activity is difficult to incorporate into busy clinical practice.

The screens in this study changed the clinical environment by presenting information from multiple sources in a format that is more likely to result in the reliable delivery of ventilation with lower tidal volumes. The configuration of the clinical environment is often ad hoc and based on historic preferences that may not reflect changes in desired care processes as new evidence emerges. This is one of the first studies to demonstrate the impact of environmental influences on clinician behaviour in the ICU and, as such, provides insight into how this approach might be developed in the ICU and beyond.

The control of tidal volume must occur at all times of the day in order to be effective. In our unit, ventilator settings are often only adjusted when clinicians pay attention to individual patients during rounds or when prompted by nursing staff on the basis of clinical concern. Adherence to low tidal volume ventilation is one consideration among many that clinicians must bear in mind when adjusting the ventilator. The intervention in this study appears to have stimulated greater interaction between clinicians with oversight of the entire unit and staff at the bedside with particular reference to the desired tidal volume. This resulted in a highly significant reduction in delivered tidal volumes. Reducing tidal volume is important because the consistent implementation of low tidal volumes is likely to improve outcomes and be cost effective in patients with acute lung injury/ARDS.34 The principle of changing the format of displayed information from a CIS is highly applicable to all ICUs with a CIS and may improve the delivery of many evidence-based interventions unrelated to ventilation.

Conclusions

The implementation of a large screen configured to display information routinely collected from a CIS in a format more likely to promote the implementation of low tidal volume ventilation resulted in a significant and sustained improvement in the use of evidence-based ventilation practice and was acceptable to the staff. The principle of configuring the clinical environment in which decisions are made in order to make it easier to comply with desired goals is widely applicable to health care. It has the potential to reduce unwanted variation in clinical practice and improve the implementation of evidence-based interventions.

Authors' contributions

C.B., M.T. and T.G. designed the study. C.B. and K.B. wrote the manuscript. K.B., J.S., A.T. and J.D. undertook the statistical analysis. J.B. and J.B. designed and undertook the qualitative evaluation.

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Declaration of interest

None declared.

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