Safety Huddle Intervention for Reducing Physiologic Monitor Alarms: A Hybrid Effectiveness-Implementation Cluster Randomized Trial

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BACKGROUND: Monitor alarms occur frequently but rarely warrant intervention.

OBJECTIVE: This study aimed to determine if a safety huddle-based intervention reduces unit-level alarm rates or alarm rates of individual patients whose alarms are discussed, as well as evaluate implementation outcomes.

DESIGN: Unit-level, cluster randomized, hybrid effectiveness-implementation trial with a secondary patient-level analysis.

SETTING: Children’s hospital.

PATIENTS: Unit-level: all patients hospitalized on four control (n = 4177) and four intervention (n = 7131) units between June 15, 2015 and May 8, 2016. Patient-level: 425 patients on randomly selected dates postimplementation.

INTERVENTION: Structured safety huddle review of alarm data from the patients on each unit with the most alarms, with a discussion of ways to reduce alarms.

MEASUREMENTS: Unit-level: change in unit-level alarm rates between baseline and postimplementation periods in intervention versus control units. Patient-level: change in individual patients’ alarm rates between the 24 hours leading up to huddles and the 24 hours after huddles in patients who were discussed versus not discussed in huddles.

RESULTS: Alarm data informed 580 huddle discussions. In unit-level analysis, intervention units had 2 fewer alarms/patient-day (95% CI: 7 fewer to 6 more, P = .50) compared with control units. In patient-level analysis, patients discussed in huddles had 97 fewer alarms/patient-day (95% CI: 52–138 fewer, P < .001) in the posthuddle period compared with patients not discussed in huddles. Implementation outcome analysis revealed a low intervention dose of 0.85 patients/unit/day.

CONCLUSIONS: Safety huddle-based alarm discussions did not influence unit-level alarm rates due to low intervention dose but were effective in reducing alarms for individual children. Journal of Hospital Medicine 2018:13:609-615. Published online first February 27, 2018. © 2018 Society of Hospital Medicine

Physiologic monitor alarms occur frequently in the hospital environment, with average rates on pediatric wards between 42 and 155 alarms per monitored patient-day. However, average rates do not depict the full story, because only 9%-25% of patients are responsible for most alarms on inpatient wards. In addition, only 0.5%-1% of alarms on pediatric wards warrant action. Downstream consequences of high alarm rates include interruptions and alarm fatigue.

Alarm customization, the process of reviewing individual patients’ alarm data and using that data to implement patient-specific alarm reduction interventions, has emerged as a potential approach to unit-wide alarm management. Potential customizations include broadening alarm thresholds, instituting delays between the time the alarm condition is met and the time the alarm sounds, and changing electrodes. However, the workflows within which to identify the patients who will benefit from customization, make decisions about how to customize, and implement customizations have not been delineated.

Safety huddles are brief structured discussions among physicians, nurses, and other staff aiming to identify and mitigate threats to patient safety. In this study, we aimed to evaluate...
the influence of a safety huddle-based alarm intervention strategy targeting high alarm pediatric ward patients on (a) unit-level alarm rates and (b) patient-level alarm rates, as well as to (c) evaluate implementation outcomes. We hypothesized that patients discussed in huddles would have greater reductions in alarm rates in the 24 hours following their huddle than patients who were not discussed. Given that most alarms are generated by a small fraction of patients,\(^1,2\) we hypothesized that patient-level reductions would translate to unit-level reductions.

**METHODS**

**Human Subject Protection**

The Institutional Review Board of Children’s Hospital of Philadelphia approved this study with a waiver of informed consent. We registered the study at ClinicalTrials.gov (identifier NCT02458872). The original protocol is available as an Online Supplement.

**Design and Framework**

We performed a hybrid effectiveness-implementation trial at a single hospital with cluster randomization at the unit level (CONSORT flow diagram in Figure 1). Hybrid trials aim to determine the effectiveness of a clinical intervention (alarm customization) and the feasibility and potential utility of an implementation strategy (safety huddles).\(^14\) We used the Consolidated Framework for Implementation Research\(^15\) to theoretically ground and frame our implementation and drew upon the work of Proctor and colleagues\(^6\) to guide implementation outcome selection.

For our secondary effectiveness outcome evaluating the effect of the intervention on the alarm rates of the individual patients discussed in huddles, we used a cohort design embedded within the trial to analyze patient-specific alarm data collected only on randomly selected “intensive data collection days,” described below and in Figure 1.

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**FIG 1. CONSORT flow diagram.**

8 non-ICU medical units invited to participate

8 Units randomized (prior to randomization, units paired based on shared characteristics*)

4 Units received the intervention (22,231 patient-days)

- 580 Patient-customized alarm discussions occurred during huddles

4 Units served as controls (22,102 patient-days)

- 0 Patient-customized alarm discussions occurred during huddles

Restricted patient-level analysis to randomly selected intensive data collection days during postimplementation period only

Evaluated the 24 hours leading up to, and the 24 hours following each huddle in an embedded cohort of patients

Intervention unit patients 512 patient-days

Control unit patients 512 patient-days

*Shared characteristics: participation in hospital-wide Joint Commission alarm management preparation activities, use of alarm middleware that relays detailed alarm information to nurses’ mobile phones, and baseline alarm rates.

ICU indicates intensive care unit.
Setting and Subjects
All patients hospitalized on eight units that admit general pediatric and medical subspecialty patients at Children’s Hospital of Philadelphia between June 15, 2015 and May 8, 2016 were included in the primary (unit-level) analysis. Every patient’s bedside included a General Electric Dash 3000 physiologic monitor. Decisions to monitor patients were made by physicians and required orders. Default alarm settings are available in Supplementary Table 1; these settings required orders to change.

All eight units were already convening scheduled safety huddles led by the charge nurse each day. All nurses and at least one resident were expected to attend; attending physicians and fellows were welcome but not expected to attend. Huddles focused on discussing safety concerns and patient flow. None of the preexisting huddles included alarm discussion.

Intervention
For each nonholiday weekday, we generated customized paper-based alarm huddle data “dashboards” (Supplementary Figure 1) displaying data from the patients (up to a maximum of four) on each intervention unit with the highest numbers of high-acuity alarms (“crisis” and “warning” audible alarms, see Supplementary Table 2 for detailed listing of alarm types) in the preceding four hours by reviewing data from the monitor network using BedMasterEx v4.2 (Excel Medical Electronics, Jupiter, Florida). Dashboards listed the most frequent types of alarms, alarm settings, and included a script for discussing the alarms with checkboxes to indicate changes agreed upon by the team during the huddle. Patients with fewer than 20 alarms in the preceding four hours were not included; thus, sometimes fewer than four patients’ data were available for discussion. We hand-delivered dashboards to the charge nurses leading huddles, and they facilitated the multidisciplinary alarm discussions focused on reviewing alarm data and customizing settings to reduce unnecessary alarms.

Study Periods
The study had 3 periods as shown in Supplementary Figure 2: (1) 16-week baseline data collection, (2) phased intervention implementation during which we serially spent 2-8 weeks on each of the four intervention units implementing the intervention, and (3) 16-week postimplementation data collection.

Outcomes
The primary effectiveness outcome was the change in unit-level alarms per patient day between the baseline and postimplementation periods in intervention versus control units, with all patients on the units included. The secondary effectiveness outcome (analyzed using the embedded cohort design) was the change in individual patient-level alarms between the 24 hours leading up to a huddle and the 24 hours following huddles in patients who were versus patients who were not discussed in huddles.

Implementation outcomes included adoption and fidelity measures. To measure adoption (defined as “intention to try” the intervention),16 we measured the frequency of discussions attended by patients’ nurses and physicians. We evaluated three elements of fidelity: adherence, dose, and quality of delivery.17 We measured adherence as the incorporation of alarm discussion into huddles when there were eligible patients to discuss. We measured dose as the average number of patients discussed on each unit per calendar day during the postimplementation period. We measured quality of delivery as the extent to which changes to monitoring that were agreed upon in the huddles were made at the bedside.

Safety Measures
To surveil for unintended consequences of reduced monitoring, we screened the hospital’s rapid response and code blue team database weekly for any events in patients previously discussed in huddles that occurred between huddle and hospital discharge. We reviewed charts to determine if the events were related to the intervention.

Randomization
Prior to randomization, the eight units were divided into pairs based on participation in hospital-wide Joint Commission alarm management activities, use of alarm middleware that relayed detailed alarm information to nurses’ mobile phones, and baseline alarm rates. One unit in each pair was randomized to intervention and the other to control by coin flip.

Data Collection
We used Research Electronic Data Capture (REDCap)18 database tools.

Data for Unit-Level Analyses
We captured all alarms occurring on the study units during the study period using data from BedMasterEx. We obtained census data accurate to the hour from the Clinical Data Warehouse.

Data Captured in All Huddles
During each huddle, we collected the number of patients whose alarms were discussed, patient characteristics, presence of nurses and physicians, and monitoring changes agreed upon. We then followed up four hours later to determine if changes were made at the bedside by examining monitor settings.

Data Captured Only During Intensive Data Collection Days
We randomly selected one day during each of the 16 weeks of the postimplementation period to obtain additional patient-level data. On each intensive data collection day, the four monitored patients on each intervention and control unit with the most high-acuity alarms in the four hours prior to huddles occurring – regardless of whether or not these patients were later discussed in huddles – were identified for data collection. On these dates, a member of the research team reviewed each patient’s alarm counts in four-hour blocks during the 24 hours before and after the huddle. Given that the huddles were not always at the same time every day (ranging between 10:00 AM and 1:00 PM), we operationally set the huddle time as 12:00 PM for all units.
Data Analysis
We used Stata/SE 14.2 for all analyses.

Unit-Level Alarm Rates
To compare unit-level rates, we performed an interrupted time series analysis using segmented (piecewise) regression to evaluate the impact of the intervention. We used a multivariable generalized estimating equation model with the negative binomial distribution and clustering by unit. We bootstrapped the model and generated percentile-based 95% confidence intervals. We then used the model to estimate the alarm rate difference in differences between the baseline data collection period and the postimplementation data collection period for intervention versus control units.

Patient-Level Alarm Rates
In contrast to unit-level analysis, we used an embedded cohort design to model the change in individual patients’ alarms between the 24 hours leading up to huddles and the 24 hours following huddles in patients who were versus patients who were not discussed in huddles. The analysis was restricted to the patients included in intensive data collection days. We performed bootstrapped linear regression and generated percentile-based 95% confidence intervals using the difference in four-hour block alarm rates between pre- and posthuddle as the outcome. We clustered within patients. We stratified by unit and preceding alarm rate. We modeled the alarm rate difference between the 24-hour prehuddle and the 24-hour posthuddle for huddled and nonhuddled patients and the difference in differences between exposure groups.

Implementation Outcomes
We summarized adoption and fidelity using proportions.

RESULTS
Alarm dashboards informed 580 structured alarm discussions during 353 safety huddles (huddles often included discussion of more than one patient).

Unit-Level Alarm Rates
A total of 2,874,972 alarms occurred on the eight units during the study period. We excluded 15,548 alarms that occurred during the same second as another alarm for the same patient because they generated a single alarm. We excluded 24,700 alarms that occurred during 4 days with alarm database downtimes that affected data integrity. Supplementary Table 2 summarizes the characteristics of the remaining 2,834,724 alarms used in the analysis.

Visually, alarm rates over time on each individual unit appeared flat despite the intervention (Supplementary Figure 3). Using piecewise regression, we found that intervention and control units had small increases in alarm rates between the baseline and postimplementation periods with a nonsignificant difference in these differences between the control and intervention groups (Table 1).

Patient-Level Alarm Rates
We then restricted the analysis to the patients whose data were collected during intensive data collection days. We obtained data from 1974 pre-post pairs of four-hour time periods. Patients on intervention and control units who were not discussed in huddles had 38 fewer alarms/patient-day (95% CI: 23–54 fewer, P < .001) in the posthuddle period than in the prehuddle period. Patients discussed in huddles had 135 fewer alarms/patient-day (95% CI: 93–178 fewer, P < .001) in the posthuddle 24-hour period than in the prehuddle period. The pairwise comparison reflecting the difference in differences showed that huddled patients had a rate of 97 fewer alarms/patient-day (95% CI: 52–138 fewer, P < .001) in the posthuddle period compared with patients not discussed in huddles.

To better understand the mechanism of reduction, we analyzed alarm rates for the patient categories shown in Table 2 and
visually evaluated how average alarm rates changed over time (Figure 2). When analyzing the six potential pairwise comparisons between each of the four categories separately, we found that the following two comparisons were statistically significant: (1) patients whose alarms were discussed in huddles and had changes made to monitoring had greater alarm reductions than patients on control units, and (2) patients whose alarms were discussed in huddles and had changes made to monitoring had greater alarm reductions than patients who were also on intervention units but whose alarms were not discussed (Table 2).

**Implementation Outcomes**

**Adoption**
The patient's nurse attended 482 of the 580 huddle discussions (83.1%), and at least one of the patient's physicians (resident, fellow, or attending) attended 394 (67.9%).

**Fidelity: Adherence**
In addition to the 353 huddles that included alarm discussion, 123 instances had no patients with ≥20 high acuity alarms in the preceding 4 hours therefore, no data were brought to the huddle. There were an additional 30 instances when a huddle did not occur or there was no discussion in the huddle despite data being available. Thus, adherence occurred in 353 of 383 huddles (92.2%).

**Fidelity: Dose**
During the 112 calendar day postimplementation period, 379 patients’ alarms were discussed in huddles for an average intervention dose of 0.85 discussions per unit per calendar day.

**Fidelity: Quality of Delivery**
In 362 of the 580 huddle discussions (62.4%), changes were agreed upon. The most frequently agreed upon changes were discontinuing monitoring (32.0%), monitoring only when asleep or unsupervised (23.8%), widening heart rate parameters (12.7%), changing electrocardiographic leads/wires (8.6%), changing the pulse oximetry probe (8.0%), and increasing the delay time between when oxygen desaturation was detected and when the alarm was generated (4.7%). Of the huddle discussions with changes agreed upon, 346 (95.6%) changes were enacted at the bedside.

**Safety Measures**
There were zero code blue events and 26 rapid response team activations for patients discussed in huddles. None were related to the intervention.

**DISCUSSION**
Our main finding was that the huddle strategy was effective in safely reducing the burden of alarms for the high alarm pedi-
TABLE 2. Alarm Rate Differences Based on Patient Category

<table>
<thead>
<tr>
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<tbody>
<tr>
<td>Control Unit, Not Discussed in Huddle</td>
<td>Intervention Unit, Not Discussed in Huddle</td>
<td>Intervention Unit, Discussed in Huddle, But No Changes Made in 4 h After Huddle</td>
<td>Intervention Unit, Discussed in Huddle, Monitor Changes Made in 4 h After Huddle</td>
</tr>
<tr>
<td>Huddles or huddle opportunities, n</td>
<td>256</td>
<td>135</td>
<td>34</td>
</tr>
<tr>
<td>Unique patients, n</td>
<td>201</td>
<td>126</td>
<td>27</td>
</tr>
<tr>
<td>Patient age in years, median (IQR)</td>
<td>4.1 (0.5–14.3)</td>
<td>4.1 (1.0–12.4)</td>
<td>0.6 (0.3–4.9)</td>
</tr>
<tr>
<td>Pre/post-huddle difference in alarms/patient-day (95% CI, P)</td>
<td>49 fewer (29 to 70 fewer)</td>
<td>14 fewer (35 fewer to 11 more)</td>
<td>54 fewer (155 fewer to 31 more)</td>
</tr>
<tr>
<td>Versus Category 1:</td>
<td></td>
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<tr>
<td>Difference in differences contrast in alarms/patient-day (95% CI, P)</td>
<td>35 more (7 fewer to 78 more, P = .17)</td>
<td>5 fewer (130 fewer to 121 more, P = .99)</td>
<td>119 fewer (186 fewer to 52 fewer, P &lt; .001)</td>
</tr>
<tr>
<td>Versus Category 2:</td>
<td></td>
<td></td>
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<tr>
<td>Difference in differences contrast in alarms/patient-day (95% CI, P)</td>
<td>40 fewer (165 fewer to 85 more, P = .99)</td>
<td>154 fewer (220 fewer to 89 fewer, P &lt; .001)</td>
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<tr>
<td>Versus Category 3:</td>
<td></td>
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<tr>
<td>Difference in differences contrast in alarms/patient-day (95% CI, P)</td>
<td></td>
<td>114 fewer (253 fewer to 24 more, P = .17)</td>
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</tbody>
</table>

*Patients who were not discussed in huddles (Groups 1 and 2) but whose data we obtained for comparison on intensive data collection days are enumerated here as “huddle opportunities.”

Abbreviations: CI, confidence interval; IQR, interquartile range.

atrial ward patients whose alarms were discussed, but it did not reduce unit-level alarm rates. Implementation outcomes explained this finding. Although adoption and adherence were high, the overall dose of the intervention was low.

We also found that 36% of alarms had technical causes, the majority of which were related to the pulse oximetry probe detecting that it was off the patient or searching for a pulse. Although these alarms are likely perceived differently by clinical staff (most monitors generate different sounds for technical alarms), they still represent a substantial contribution to the alarm environment. Minimizing them in patients who must remain continuously monitored requires more intensive effort to implement other types of interventions than the main focus of this study, such as changing pulse oximetry probes and electrocardiographic leads/wires.

In one-third of huddles, monitoring was simply discontinued. We observed in many cases that, while these patients may have had legitimate indications for monitoring upon admission, their conditions had improved; after brief multidisciplinary discussion, the team concluded that monitoring was no longer indicated. This observation may suggest interventions at the ordering phase, such as pre-specifying a monitoring duration.22,23

This study’s findings were consistent with a quasi-experimental study of safety huddle-based alarm discussions in a pediatric intensive care unit that showed a patient-level reduction of 116 alarms per patient-day in those discussed in huddles relative to controls.11 A smaller quasi-experimental study of implementing a nighttime alarm “ward round” in an adult intensive care unit showed a significant reduction in unit-level alarms/patient-day from 168 to 84.9 In a quality improvement report, a monitoring care process bundle that included discussion of alarm settings showed a reduction in unit-level alarms/patient-day from 180 to 40.10 Our study strengthens the body of literature using a cluster-randomized design, measuring patient- and unit-level outcomes, and including implementation outcomes that explain effectiveness findings.

On a hypothetical unit similar to the ones we studied with 20 occupied beds and 60 alarms/patient-day, an average of 1,200 alarms would occur each day. We delivered the intervention to 0.85 patients per day. Changes were made at the bedside in 60% of those with the intervention delivered, and those patients had a difference in differences of 119 fewer alarms compared with the comparison patients on control units. In this scenario, we could expect a relative reduction of 0.85 × 0.60 × 119 = 61 fewer alarms/day total on the unit or a 5% reduction. However, that estimated reduction did not account for the arrival of new patients with high alarm rates, which certainly occurred in this study and explained the lack of effect at the unit level.

As described above, the intervention dose was low, which translated into a lack of effect at the unit level despite a strong effect at the patient level. This result was partly due to the manual process required to produce the alarm dashboards that restricted their availability to nonholiday weekdays. The study was performed at one hospital, which limited generalizability. The study hospital was already convening daily safety huddles relative to controls. The study hospital was already convening daily safety huddles relative to controls.11 A smaller quasi-experimental study of implementing a nighttime alarm “ward round” in an adult intensive care unit showed a significant reduction in unit-level alarms/patient-day from 168 to 84.9 In a quality improvement report, a monitoring care process bundle that included discussion of alarm settings showed a reduction in unit-level alarms/patient-day from 180 to 40.10 Our study strengthens the body of literature using a cluster-randomized design, measuring patient- and unit-level outcomes, and including implementation outcomes that explain effectiveness findings.

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pitals without existing huddle structures may face challenges in implementing similar multidisciplinary alarm discussions. In addition, the study design was randomized at the unit (rather than patient) level, which limited our ability to balance potential confounders at the patient level.

CONCLUSION

A safety huddle intervention strategy to drive alarm customization was effective in safely reducing alarms for individual children discussed. However, unit-level alarm rates were not affected by the intervention due to a low dose. Leaders of efforts to reduce alarms should consider beginning with passive interventions (such as changes to default settings and alarm delays) and use huddle-based discussion as a second-line intervention to address remaining patients with high alarm rates.

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References