Physiologic Monitor Alarms for Children: Pushing the Limits

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Deciding when a hospitalized child’s vital signs are acceptably within range and when they should generate alerts, alarms, and escalations of care is critically important yet surprisingly complicated. Many patients in the hospital who are recovering appropriately exhibit vital signs that fall outside normal ranges for well children. In a technology-focused hospital environment, these out-of-range vital signs often generate alarms in the electronic health record (EHR) and alarms on physiologic monitors that can disrupt patients’ sleep, generate concern in parents, lead to unnecessary testing and treatment by physicians, interrupt nurses during important patient care tasks, and lead to alarm fatigue. It is this last area, the problem of alarm fatigue, that Goel and colleagues2–6 have used to frame the rationale and results of their study reported in this issue of the Journal of Hospital Medicine.

Goel and colleagues correctly point out that physiologic monitor alarm rates are high in children’s hospitals, and alarms warranting intervention or action are rare.2–6 Few studies have rigorously examined interventions to reduce unnecessary hospital physiologic monitor alarms, especially in pediatric settings. Of all the potential interventions, widening parameters has the most face validity: if you set wide enough alarm parameters, fewer alarms will be triggered. However, it comes with a potential safety tradeoff of missed actionable alarms.

Before EHR data became widely available for research, normal (or perhaps more appropriate for the hospital setting, “expected”) vital sign ranges were defined using expert opinion. The first publication describing the distribution of EHR-documented vital signs in hospitalized children was published in 2013.7 Goel and colleagues have built upon this prior work in their article, in which they present percentiles of EHR-documented heart rate (HR) and respiratory rate (RR) developed using data from more than 7000 children hospitalized at an academic children’s hospital.

In a separate validation dataset, they then compared the performance of their proposed physiologic monitor alarm parameters—the 5th and 95th percentiles for HR and RR from this study—to the 2004 National Institutes of Health (NIH) vital sign reference ranges8 that were the basis of default alarm parameters at their hospital. They also compared their percentiles to the 2013 study.7

The 2 main findings of Goel and colleagues’ study were: (1) using their separate validation dataset, 55.6% fewer HR and RR observations were out of range based on their newly developed percentiles as compared to the NIH vital sign reference ranges; and (2) the HR and RR percentiles they developed were very similar to those reported in the 2013 study,7 which used data from 2 other institutions, externally validating their findings.

The team then pushed the data a step further in a safety analysis and evaluated the sensitivity of the 5th and 95th percentiles for HR and RR from this study for detecting deterioration in 148 patients in the 12 hours before either a rapid response team activation or a cardiorespiratory arrest. The overall sensitivity for having either a HR or RR value out of range was 93% for Goel and colleagues’ percentiles and 97% for the NIH ranges. Goel and colleagues concluded that using the 5th and 95th HR and RR percentiles provides a potentially safe means by which to modify physiologic bedside monitor alarm limits.

There are 2 important limitations to this work. The first is that the study uses EHR-documented data to estimate the performance of new physiologic monitor settings. Although there are few published reports of differences between nurse-charted vital signs and monitor data, those that do exist suggest that nurse charting favors more stable vital signs,9,10 even when charting oxygen saturation in patients with true, prolonged desaturation.9 We agree with the authors of 1 report, who speculated that nurses “recognize that temporary changes in vital signs are untypical for that patient and might choose to ignore them and either await a period of stability or make an educated estimate for that hour.”9 When using Goel and colleagues’ 5th and 95th percentiles as alarm parameters, the expected scenario is that monitors will generate alarms for 10% of HR values and 10% of RR values. Because of the differences between nurse-charted vital

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signs and monitor data, the monitors will probably generate many more alarms.

The second limitation is the approach Goel and colleagues took in performing a safety analysis using chart review. Unfortunately, it is nearly impossible for a retrospective chart review to form the basis of a convincing scientific argument for the safety of different alarm parameters. It requires balancing the complex and sometimes competing nurse-level, patient-level, and alarm-level factors that determine nurse response time to alarms. It is possible to do prospectively, and we hope Goel’s team will follow up this article with a description of the implementation and safety of these parameters in clinical practice.

In addition, the clinical implications of HR and RR at the 95th percentile might be considered less immediately life threatening than HR and RR at the 5th percentile, even though statistically they are equally abnormal. When choosing percentile-based alarm parameters, statistical symmetry might be less important than the potential immediate consequences of missing bradycardia or bradypnea. It would be reasonable to consider setting high HR and RR at the 99th percentile or higher, because elevated HR or RR alone is rarely immediately actionable, and set the low HR and RR at the 5th or 10th percentile.

Despite these caveats, should the percentiles proposed by Goel and colleagues be used to inform pediatric vital sign clinical decision support throughout the world? When faced with the alternative of using vital sign parameters that are not based on data from hospitalized children, these percentiles offer a clear advantage, especially for hospitals similar to Goel’s. The most obvious immediate use for these percentiles is to improve noninterruptive\textsuperscript{11} vital sign clinical decision support in the EHR, the actual source of the data in this study.

The question of whether to implement Goel’s 5th and 95th percentiles as physiologic monitor alarm parameters is more complex. In contrast to EHR decision support, there are much clearer downstream consequences of sounding unnecessary alarms as well as failing to sound important alarms for a child in extremis. Because their percentiles are not based on monitor data, the projected number of alarms generated at different percentile thresholds cannot be accurately estimated, although using their 5th and 95th percentiles should result in fewer alarms than the NIH parameters.

In conclusion, the work by Goel and colleagues represents an important contribution to knowledge about the ranges of expected vital signs in hospitalized children. Their findings can be immediately used to guide EHR decision support. Their percentiles are also relevant to physiologic monitor alarm parameters, although the performance and safety of using the 5th and 95th percentiles remain in question. Hospitals aiming to implement these data-driven parameters should first evaluate the performance of different percentiles from this article using data obtained from their own monitor system and, if proceeding with clinical implementation, pilot the parameters to accurately gauge alarm rates and assess safety before spreading hospital wide.

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