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Performance of commonly used risk triage tools in predicting clinical deterioration among hospitalized hematopoietic stem cell transplant recipients

Short Title: Predicting deterioration in stem cell recipients

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Although hospitalized hematopoietic cell transplantation (HCT) recipients frequently experience critical illness, no prediction tools for clinical deterioration have been validated in this population. We examined the discrimination of five commonly-used risk triage tools among hospitalized HCT recipients at an academic hospital (1/1/2019-12/31/2022): systemic inflammatory response syndrome (SIRS) criteria, the quick sepsis-related organ failure assessment (qSOFA), the Modified and National Early Warning Scores (MEWS, NEWS), and the Epic Deterioration Index (EDI). We calculated each tool's area under the receiver-operating characteristic curve (AUROC) for a composite of ward death, hospice discharge, or intensive care unit transfer. This outcome occurred in 137 of 1298 hospitalizations (11%). Hospitalization-level discrimination was lowest for SIRS (AUROC 0.62, 95% CI 0.57-0.66) and highest for the EDI (0.82, 0.78-0.86). However, the EDI's clinical utility may be limited by long lead times and low positive predictive values.

While outcomes for recipients of hematopoietic cell transplantation (HCT) have improved in recent years, these patients remain at high risk of clinical deterioration (i.e., physiologic worsening resulting in critical illness or death) during hospitalization.¹⁻³ Because many instances of clinical deterioration represent failure to recognize or act upon early indicators of instability, Early Warning Systems (EWS) have been developed to identify patients at risk of worsening before critical events occur, enabling timely intervention.⁴ Critically, these tools have not been well-evaluated in larger series of HCT recipients,⁵ who often display different physiology than general inpatients (e.g., fevers, cytopenias, immunocompromise) that could limit performance of risk tools including these variables.⁵⁻⁷ An inaccurate EWS could drive worse patient outcomes through both false-positive (e.g., alert fatigue, potential for inappropriate testing and/or unnecessary interventions with associated risks) and false-negative (i.e., failure to rescue) results. This study therefore evaluated the performance of several commonly-used risk triage tools for predicting clinical deterioration among hospitalized HCT recipients.

This retrospective cohort study was conducted at the Knight Cancer Institute, Oregon Health & Science University's (OHSU's) NCI-designated Comprehensive Cancer Center. All ward hospitalizations for adult HCT recipients between 1/1/2019 and 12/31/2022 were included. OHSU's institutional review board approved this study (#25188). Due to the retrospective and minimal risk nature of this study, informed consent was not required by the institutional review board. Brenna Park-Egan, MS, and Patrick Lyons, MD, MSc, analyzed the data for this study, and all authors had access to the primary data. Some of these results have previously been presented in the form of an abstract.⁸ Electronic health record (EHR) data were obtained from OHSU's Research Data Warehouse. From these, several scores were calculated hourly based on vital sign and laboratory data from the wards: Systemic Inflammatory Response Syndrome (SIRS), quick Sepsis Related Organ Failure Assessment, (qSOFA), the Modified Early Warning Score (MEWS) and the National Early Warning Score (NEWS) (Supplement Table 1).⁹ Additionally, the Epic Deterioration Index (EDI, a vendor-supplied model predicting deterioration based on vital signs, demographics, nursing assessments, and lab values) was extracted.^{10,11} At OHSU, the EDI is passively displayed within the EHR and used actively to trigger rapid response team evaluation. The other tools in this study are not used by any local clinical

protocols. The primary outcome was the composite of ward to intensive care unit (ICU) transfer, discharge to hospice, or death on the wards. Data were censored after the first of these events.

In the primary analysis, each tool's hospitalization-level area under the receiver-operating characteristic (AUROC) was calculated based on the maximum score from ward admission until deterioration or discharge. Secondly, discrete time survival analysis was used to predict, every 4 hours, whether the composite outcome would occur in the subsequent 24 hours.¹² For each tool, commonly used thresholds were used to determine the cumulative incidence of "positive" scores (i.e., hypothetical alerts), as well as the time between score-positivity and the primary outcome.^{9,10} Subgroup analyses compared tool performance between autologous and allogeneic transplant groups, and index and subsequent admissions. Statistical analyses were performed in R version 4.4.0, using packages caret (v.6.0-94) and pROC (v.1.18.5).^{14, 15}

In total, 1298 hospitalizations from 800 patients were included: 732 hospitalizations for allogeneic HCT recipients and 566 hospitalizations for autologous recipients. Of these, 789 encounters were index admissions, and 509 were subsequent hospitalizations. The composite outcome occurred in 137 hospitalizations (11%), 111 of which involved ICU transfers (Table 1). Among the ICU transfers, 49 (44%) ultimately died and 3 were discharged to hospice (3%). In overall hospitalizations, allogeneic HCT recipients were significantly more likely to experience the primary outcome than recipients of autologous HCTs ($n = 94$, 13% vs $n = 43$, 8%, $p = 0.02$). This difference was primarily driven by index transplant hospitalizations (allo-HCT $n = 41$, 12%, vs auto-HCT $n = 26$, 8%, $p = 0.003$); no significant difference was observed in outcome rates across the two groups in subsequent hospitalizations. For all scores, median values were significantly higher ($p < 0.01$) among patients who deteriorated compared to those who did not (Figure 1A).

The EDI had the greatest hospitalization-level discrimination (AUROC 0.82, 95% CI 0.78-0.86), followed by NEWS (0.69, 0.64-0.74), MEWS (0.66, 0.61-0.70), qSOFA (0.64, 0.59-0.69), and SIRS (0.62, 0.57-0.66). Tool performance using discrete time survival analysis was significantly higher for NEWS and MEWS ($p < 0.001$) while no significant differences were seen in the other tools (EDI AUROC 0.84, 95% CI 0.83-0.86; NEWS AUROC 0.80, 95% CI 0.78-.82; MEWS AUROC 0.77, 95% CI 0.75-0.79; qSOFA AUROC 0.68, 95% CI 0.67- 0.70; SIRS AUROC 0.67, 95% CI 0.65-0.69). MEWS and SIRS had higher AUROCs among allogeneic recipients ($p = 0.01$ and $p < 0.01$, respectively) than autologous recipients, but no significant differences were seen between transplant groups for the other tools. While performance of every risk prediction tool was observed to be higher in overall subsequent admissions than index admissions, none of these differences were statistically significant (Figure 2). Overall, 1,149 (89%) encounters were SIRS-positive, 971 (75%) were NEWS-positive, 506 (39%) were MEWS-positive, and 225 (17%) were qSOFA-positive at some point during ward hospitalization. A total of 638 (49%) encounters reached the EDI's "moderate" threshold at least once, while 52 (4%) reached its "high" threshold. In 5 encounters, patients deteriorated without reaching a positive threshold for any tool (these involved facilitating urgent dialysis [$n=1$], hyponatremia management [$n=2$], and increased neurological monitoring [$n=2$]). The time between first reaching a score's positive threshold and deterioration was shortest for qSOFA (3 days, range 1-9 days), followed by

MEWS (4 days, 1-10 days), the EDI (5 days, 2-11 days), NEWS (9 days, 3-16 days), and SIRS (9 days, 4-15 days).

In this 3-year cohort of hospitalized HCT recipients, commonly used risk prediction tools generally had high discrimination for clinical deterioration. However, values for all scores were frequently at or above typical thresholds for clinical alerting; for instance, only one in ten patients would have avoided meeting SIRS criteria during their hospitalization (likely due to high rates of febrile neutropenia and cytokine release syndrome). Further, hypothetical “lead times” between alerts and clinical deterioration were consistently on the order of days rather than hours. Taken together, these observations suggest limited clinical utility for these scores without substantial modification to how and when they are delivered to, and used by, clinicians. For example, temporarily suppressing subsequent alerts after an initial alert could reduce alarm fatigue. False positives would also be reduced with higher thresholds for positivity; such a strategy could either be applied overall (because the HCT population faces higher baseline risk) or dynamically during periods where risk is expected to be higher or lower based on transplant type and timing. Interestingly, while other scores had similar discrimination to what has been reported among general inpatients, the EDI showed greater discrimination than these comparators.^{9,10,13} This finding may reflect the fact that HCT recipients are typically selected based on suitable performance status and fewer comorbidities; for example, supplemental oxygen use (one parameter in the EDI) could be acute or chronic for many patients on the general wards, but is likely representative of acute pathology (and, thus, risk) among HCT recipients. Future work should focus on identifying and validating routinely-available clinical data points where the signal:noise ratio may differ for HCT patients as compared to more general populations; such variables might provide a path to more informative – and therefore more useful – predictive tools.

A limitation of this retrospective study is that some deterioration events may have been prevented or delayed by timely and effective clinical care in response to high EDI scores;⁹ our evaluation did not include data on interventions that could have contextualized these results.

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Table 1. Patient characteristics and outcomes across index and subsequent admissions.

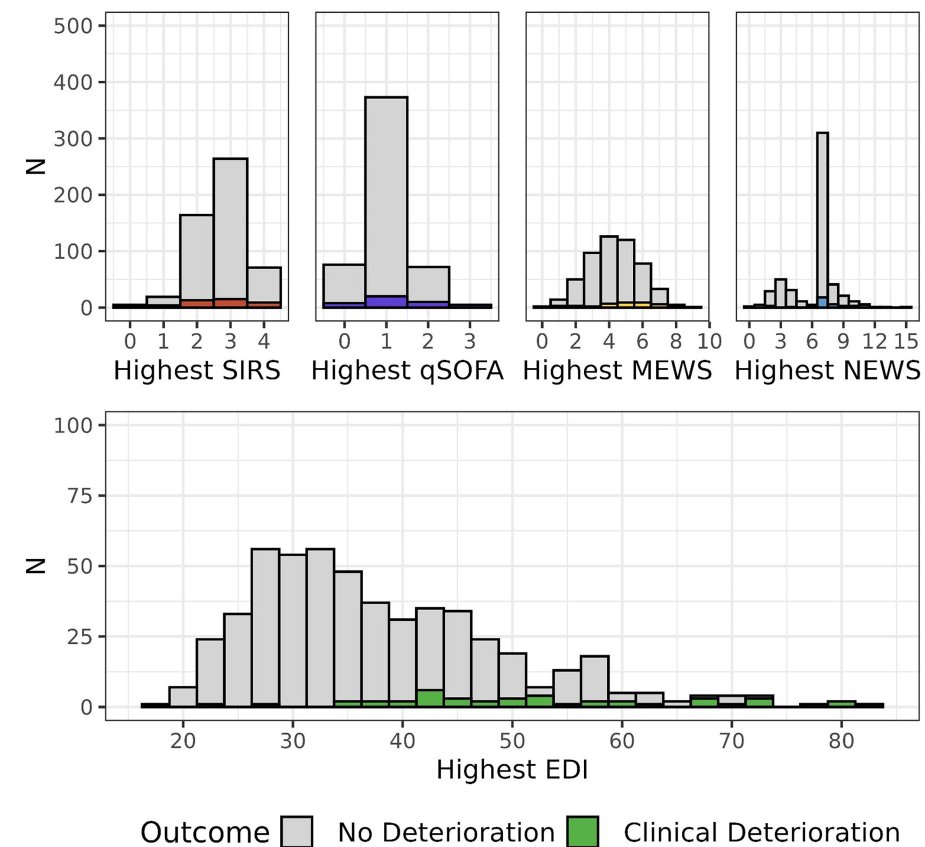
Index Admissions			
	Allogeneic Hospitalizations (n = 347)	Autologous Hospitalizations (n = 442)	p-value
<u>Characteristics</u>			
Age, median (IQR)	57 (45, 65)	62 (52, 68)	<0.001
Female, n (%)	161 (46%)	169 (38%)	0.023
Min White Blood Count, median (IQR)	0.12 (0.10, 0.15)	0.12 (0.11, 0.16)	0.2
Max Temperature, median (IQR)	100.90 (100.00, 102.40)	101.10 (100.20, 102.20)	0.6
<u>Outcomes</u>			
Composite Outcome, n (%)	41 (12%)	26 (5.9%)	0.003
Intensive Care Unit Transfer, n (%)	39 (11%)	25 (5.7%)	0.004
Length of Stay, median (IQR)	23 (20, 29)	16 (14, 19)	<0.001
100 Day All-Cause Mortality, n (%)	45 (13%)	15 (3.4%)	<0.001
Subsequent Admissions			
	Allogeneic Hospitalizations (n = 385)	Autologous Hospitalizations (n = 124)	p-value
<u>Characteristics</u>			
Age, median (IQR)	56 (41, 65)	58 (42, 67)	0.4

Female, n (%)	180 (47%)	50 (40%)	0.2
Min White Blood Count, median (IQR)	3.39 (1.59, 5.62)	2.28 (0.96, 3.94)	<0.001
Max Temperature, median (IQR)	99.10 (98.70, 100.40)	99.90 (98.80, 101.60)	0.003
<u>Outcomes</u>			
Composite Outcome, n (%)	53 (14%)	17 (14%)	>0.9
Intensive Care Unit Transfer, n (%)	33 (8.6%)	14 (11%)	0.4
Length of Stay, median (IQR)	6 (4, 15)	5 (3, 12)	0.017

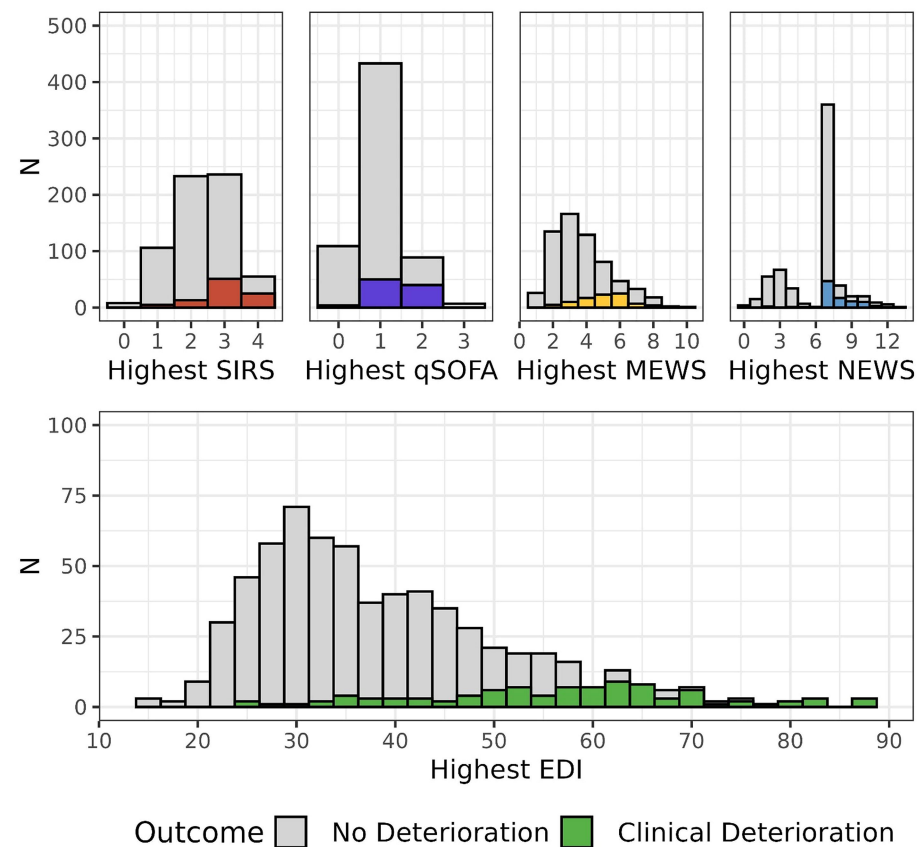
Figure 1. Hospitalization-level score distribution and cumulative incidence of positive scores across autologous and allogeneic admissions. Panels (a) and (b) depict the hospitalization-level highest score distribution of the systemic inflammatory response syndrome (SIRS) criteria, the quick sepsis-related organ failure assessment (qSOFA), the Modified and National Early Warning Scores (MEWS, NEWS), and the Epic Deterioration Index (EDI) across outcome status for autologous and allogeneic hospitalizations, respectively. The x-axis depicts the range of possible scores for each tool, while the y-axis depicts the number of hospital encounters with a maximum of each score. In (c) and (d), SIRS showed the overall highest cumulative incidence of positive scores, followed by NEWS, EDI, MEWS, and qSOFA, for autologous (c) and allogeneic (d) hospitalizations. The x-axis depicts the hospital day and the y axis depicts the proportion of hospitalizations reaching positivity for each score.

Figure 2. Hospitalization-level EWS performance across transplant and admission types. Panels (a) and (b) depict the hospitalization-level performance of the systemic inflammatory response syndrome (SIRS) criteria, the quick sepsis-related organ failure assessment (qSOFA), the Modified and National Early Warning Scores (MEWS, NEWS), and the Epic Deterioration Index (EDI) across outcome status for index and subsequent hospitalizations, respectively, for autologous and allogeneic transplant recipients. The x-axis depicts the area under the receiver operating characteristic for each tool, while the y-axis depicts the selected tool.

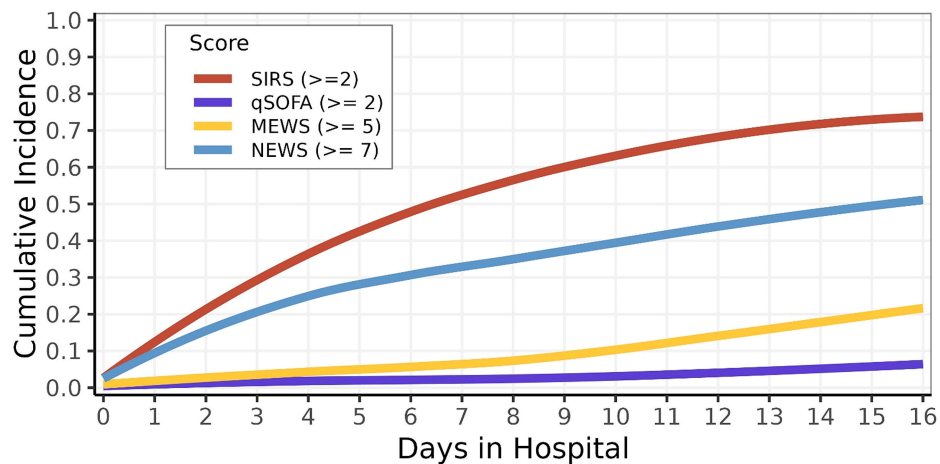
A. Autologous admissions



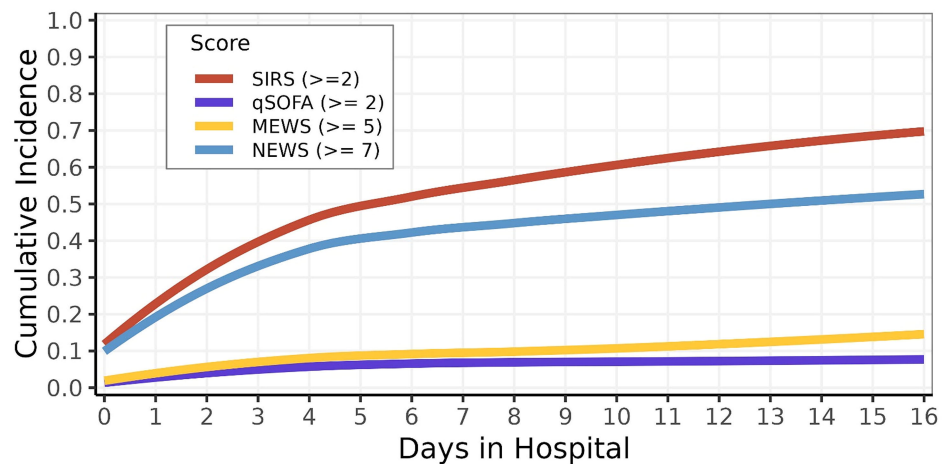
B. Allogeneic admissions



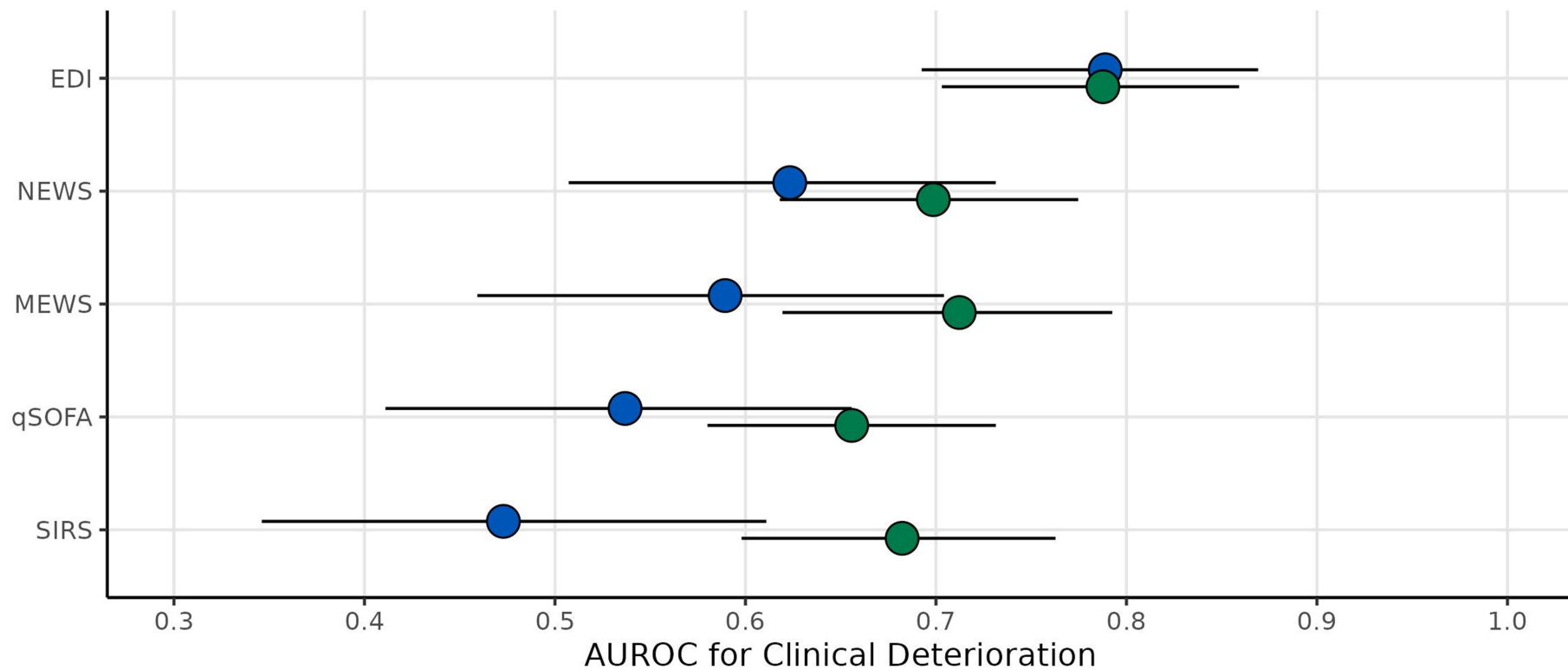
C.



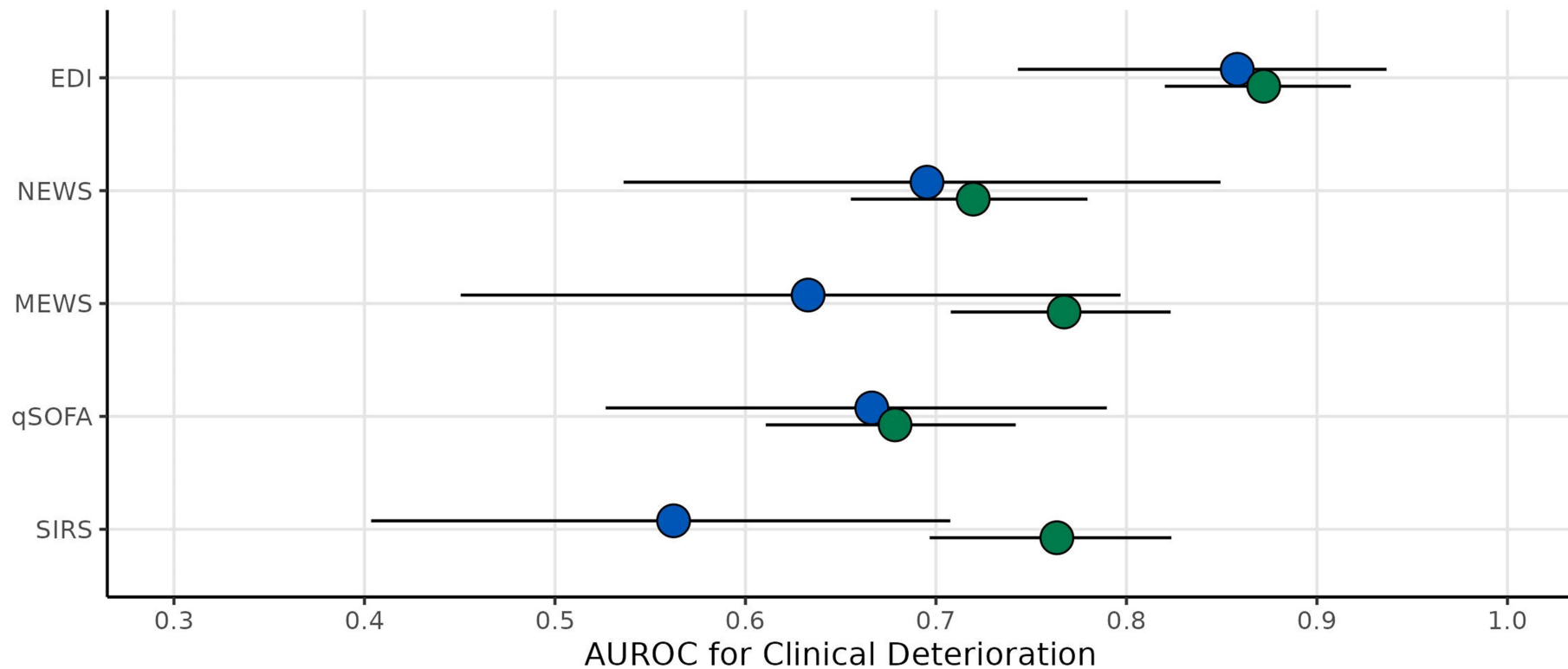
D.



A. Index Admissions



B. Subsequent Admissions



Transplant Type ● Allogeneic ● Autologous

Supplement Table 1. Triage tool score components

Triage Tool	EDI	SIRS	qSOFA	MEWS	NEWS
Score Range	0 - 100	0 - 4	0 - 3	0 - 14	0 - 20
Alert Threshold	37.4, 68.8	≥ 2	≥ 2	≥ 5	≥ 7
Temperature (F)	x*	< 96.8 or $> 100.4 = 1$		> 101.2 or $< 95 = 2$	$\geq 102.3 = 2$, $100.5 - 102.2 = 1$, $96.9 - 100.4 = 0$, $95.1 - 96.8 = 1$, $\leq 95 = 3$
Mental status (GCS)***	x*		$< 15 = 1$	$0 - 7 = 3$, $8 - 12 = 2$, $13 - 14 = 1$, $15 = 0$	$< 15 = 3$
Pulse	x*	$> 90 = 1$		$\geq 130 = 3$, $111 - 129 = 2$, $101 - 110 = 1$, $51 - 100 = 0$, $41 - 50 = 1$, $\leq 40 = 2$	$\geq 131 = 3$, $111 - 130 = 2$, $91 - 110 = 1$, $51 - 90 = 0$, $41 - 50 = 1$, $\leq 40 = 3$
SBP	x*		$< 100 = 1$	$\geq 200 = 2$, $101 - 199 = 0$, $81 - 100 = 1$, $71 - 80 = 2$, $\leq 70 = 3$	$\geq 220 = 3$, $111 - 219 = 0$, $101 - 110 = 1$, $91 - 100 = 2$, $\leq 90 = 3$
RR	x*	$> 20 = 1$	$\geq 22 = 1$	$\geq 30 = 3$, $21 - 29 = 2$, $15 - 20 = 1$, $9 - 14 = 0$, $\leq 8 = 2$	$\geq 25 = 3$, $21 - 24 = 2$, $12 - 20 = 0$, $9 - 11 = 1$, $\leq 8 = 3$
SpO2	x*				$\geq 96 = 0$, $94 - 95.9 = 1$, $92 - 93.9 = 2$, $\leq 91 = 3$
Suppl O2	x*				Any O2 support = 2
Blood counts	x*	WBC > 12 or $< 4 = 1$			
Blood chemistry**	x*				
Age	x*				

*Values not publicly available, **[EDI = Na, K, BUN, pH], ***Glasgow Coma Scale