The Association Between an Ultrabrief Cognitive Screening in Older Adults and Hospital Outcomes

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BACKGROUND: Though often recommended, hospital cognitive assessment is infrequently completed due to clinical and time constraints.

OBJECTIVE: This analysis aimed to evaluate the relationship between performance on ultrabrief cognitive screening instruments and hospital outcomes.

DESIGN: This is a secondary data analysis of a quality improvement project.


PATIENTS: Patients ≥60 years old, admitted to the hospital.

INTERVENTION: None.

MEASUREMENTS: Upon admission, patients were administered 2 cognitive screening tools. The modified Richmond Agitation and Sedation Scale (mRASS) is a measure of arousal that can be completed in 15 seconds. The months of the year backward (MOYB) is a measure of attention that can be administered in ≤1 minute. In-hospital outcomes included restraints and mortality, whereas discharge outcomes included length of stay, discharge not home, and variable direct costs. Risk ratios were calculated for dichotomous outcomes and unadjusted Poisson regression for continuous outcomes.

RESULTS: Patients (n = 3232) were screened. Altered arousal occurred in 15% of patients (n = 495); incorrect MOYB was recorded in 45% (n = 1457). Relative to those with normal arousal and attention, those with abnormal mRASS and incorrect MOYB had increased length of stay (incident rate ratio [IRR]: 1.08, 95% confidence interval [CI]: 1.02–1.14), restraint use (risk ratio [RR]: 5.05, 95% CI: 3.29–7.75), in-hospital mortality (RR: 3.46, 95% CI: 1.24–9.63), and decreased discharge home (RR: 2.97, 95% CI: 2.42–3.64). Hospital variable direct costs were slightly, but not significantly, higher (IRR: 1.02, 95% CI: 0.88–1.17).


Hospitalization is a critical time for older patients with cognitive impairment. Past research has found that hospitalized older adults with cognitive dysfunction have more rapid cognitive decline, increased morbidity and mortality, and higher costs of healthcare utilization.1–3 Those with preexisting cognitive dysfunction, such as dementia, are most susceptible to the negative impacts of hospitalization.4–8 Identification of cognitive deficits upon admission is important for risk stratification of patients and prevention of negative hospital health events.

Frontline healthcare providers are underequipped to detect acute cognitive dysfunction.9,10 Current practice and research for the detection of cognitive dysfunction in the acute care setting utilizes instruments that require training11 and are relatively lengthy (>5 minutes).12 Although these cognitive screening tests are accurate and reliable, the time requirement is not feasible in a fast-paced clinical setting. A possible alternative is the use of ultra-brief cognitive screening instruments (<1 minute) that have the potential to identify those individuals requiring additional evaluation and follow-up. These brief instruments are composed of screening tools that emphasize core features of acute cognitive dysfunction such as level of arousal or attention.13–16 Arousal, the ability to respond to or interact with the environment,17 is an important component of cognition because it is generally preserved in chronic cognitive disorders (eg, dementia). Thus, an alteration in arousal may be a harbinger of more acute impairment17 in need of evaluation, and in these...
lowered states of arousal it may be difficult to test for attention. Attention is a broadly defined cognitive domain indicating focus. Older adults, regardless of preexisting cognitive dysfunction, warrant additional cognitive testing if levels of arousal or attention are altered due to the significant relationship to delirium, which is associated with adverse events in this population. Recent research has demonstrated that these brief cognitive screening instruments provide information about the risk for delirium and are a strong test for clinical characteristics of delirium.

The purpose of this analysis was to demonstrate the clinical outcomes of poor performance on ultrabrief assessments arousal and attention by frontline staff using a quality improvement database. Specific objectives include determining (1) the association of poor performance on brief cognitive assessments and hospital outcomes and (2) the inter-relationship between alterations in the levels of arousal and attention on in-hospital and discharge outcomes.

METHODS
Setting and Study Design
This is a secondary analysis of data collected from a quality improvement program for delirium risk modification. This program collected data from October 2010 until September 2012 at a Veterans Affairs (VA) tertiary referral center for the New England region. Patients aged 60 years or older and admitted to medical wards were screened upon admission or transfer to VA Boston Healthcare System and provided appropriate interventions to modify delirium risk. Excluded were individuals admitted as observational status, or those readmitted within 30 days of initial screening, and those screened more than 72 hours after admission. Age and sex were abstracted from the medical record. Outcome data were collected from the medical record for the purpose of operating and sustaining the program. In a previous article, the length of stay (LOS) outcome was reported in a subset of this population. The analysis presented here includes the full cohort, presents the interaction with month of the year backward (MOYB), and provides additional outcomes not included in the other article. The VA institutional review board (IRB) reviewed and approved the secondary data analysis of the quality improvement project.

Measures
Brief Cognitive Screening
The baseline assessments of levels of arousal and attention were collected within 72 hours of admission to identify delirium risk. Trained study staff, not involved in the clinical care of patients, administered these assessments as part of the quality improvement project. It is estimated that these assessments took less than 1 minute to complete per individual, but actual administration time was not measured. Assessments were documented within the electronic health record as part of a delirium risk stratification system.

Arousal
The arousal level assessment was the modified Richmond Agitation and Sedation Scale (mRASS). The mRASS is a brief, reliable, observational tool used to determine arousal level. It is a text modification of the RASS for less acutely ill patients, capturing hyperactive and hypoactive altered levels of arousal. The mRASS asks an open-ended question followed by observation for 10 seconds and completion of a −5 to +4 rating scale. Alert and calm (score = 0) is considered normal, with positive numbers related to an increased level of arousal and attention, whereas negative numbers denote decreased levels. For the analyses, an mRASS of 0 is utilized as the reference. Categories were collapsed into ≥2 and ≤−2 due to few patients on the extremes of the mRASS.

Attention
The MOYB is a brief measure of attention that is included in several instruments for delirium. For this study, the patient was asked to recite the 12 months backward beginning with December. A correct score was given if the individual was able to recite all 12 months to January without any error. An incorrect score was given if any mistake was made. Scoring for the MOYB is not standardized by age, preexisting medical diagnosis, or any other rational. Others have used July or June as a cutoff for a correct score on the MOYB, but a more conservative score of correct to January was used in this study, which has been previously used. A score of not completed was given when the patient was unable to participate or declined to complete the assessment. For the analysis, a correct score on the MOYB is the referent group.

Outcomes
In-hospital outcomes included (1) restraint use and (2) in-hospital mortality. Physical restraint use was identified by focused medical record review and identification of required restraint documentation, which, by center policy requires daily review and documentation. Any restraint use during the hospitalization was included.

Discharge outcomes included (1) LOS, (2) discharge other than a location to home, and (3) variable direct costs. LOS was calculated from date of admission until date of discharge. Discharge disposition was identified in the electronic medical record discharge documentation and categorized into discharge to the prehospital residence (home) or not. Hospital variable direct costs were collected from the VA decision support system, a centrally maintained administrative database. The VA decision support system is challenged with accounting for costs of a single-day...
admission and patients who are hospitalized from VA long-term care. To address the missing data from these cases, multiple imputations (n = 20) of the missing data were performed. Sensitivity analyses were performed to determine the impact of the imputation and the cost analysis strategy (see Supporting Information, Appendix 1, in the online version of this article).

**Statistical Analyses**

For this analysis, outcomes are reported at each level of performance on the mRASS (≤-1 to ≥1) and MOYB (correct, incorrect, not completed). For each analysis, the performance with a mean and standard deviation (SD) are reported for continuous outcomes and a percentage for dichotomous outcomes. For dichotomous outcomes, including restraint use, in-hospital mortality, and discharge disposition, a risk ratio (RR) with 95% confidence interval (CI) is presented. The median is presented for the cost data because variable direct cost is highly skewed. For LOS and cost outcomes, unadjusted incident rate ratio (IRR) from a Poisson regression relative to the referent is presented to compare the categories. A Poisson regression was selected because LOS (a count of days) and variable direct costs (a count of dollars) are highly skewed. The output of Poisson regression produces an IRR and 95% CI relative to the referent group. The Poisson regression could not be adjusted because the quality improvement nature of these data limited the number of covariates collected. Sensitivity analyses did not identify significant interactions of age and sex (results not shown).

MOYB was also compared by level of arousal (mRASS = 0 vs all others). Due to the relatively few patients with positive mRASS, it was compressed into a category of abnormal mRASS relative to alert and calm. Similar to the previous analyses, Poisson regression was performed to calculate the IRR (95% CI) relative to correct MOYB for the continuous variables. An RR was calculated for the dichotomous variables. All statistical analyses were performed using Stata version 11.0 (StataCorp, College Station, TX).

**RESULTS**

**Population Description**

Over the 2-year project timeline, a total of 3232 unique individual records were analyzed (Table 1). Patients admitted and screened within the prior 30 days (n = 501) and patients screened more than 3 days after admission (n = 664) were not included in the analysis. Older adults were on average 74.7 years old (SD = 9.8), and 98.2% were male, consistent with the veteran population. Altered level of arousal, as defined by an abnormal mRASS score, was found in 15.3% of the population. Average LOS was 5.2 days (SD = 5.6), restraint use occurred in 5.5% during the hospital stay, patients were likely to be discharged home (71.7%), and a small portion died during hospitalization (1.3%). Mean variable direct costs were $11,084 with expected variability (SD = $15,682, median $6,614). Patients who died during the hospitalization were excluded from this analysis.

**TABLE 1. Baseline Characteristics**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Result, N = 3,232, Mean (SD) or % (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, y</td>
<td>74.7 (9.8)</td>
</tr>
<tr>
<td>Male</td>
<td>98.2 (1,174)</td>
</tr>
<tr>
<td>mRASS</td>
<td></td>
</tr>
<tr>
<td>≤-2</td>
<td>2.0% (64)</td>
</tr>
<tr>
<td>-1</td>
<td>8.5% (273)</td>
</tr>
<tr>
<td>0</td>
<td>84.7% (2,737)</td>
</tr>
<tr>
<td>1</td>
<td>4.0% (131)</td>
</tr>
<tr>
<td>≥2</td>
<td>0.8% (27)</td>
</tr>
<tr>
<td>MOYB</td>
<td></td>
</tr>
<tr>
<td>Correct</td>
<td>48.8% (1,578)</td>
</tr>
<tr>
<td>Incorrect</td>
<td>45.1% (1,457)</td>
</tr>
<tr>
<td>Not completed</td>
<td>6.1% (197)</td>
</tr>
<tr>
<td>Restraint use</td>
<td></td>
</tr>
<tr>
<td>In-hospital mortality</td>
<td>1.3% (41)</td>
</tr>
<tr>
<td>Length of stay, d*</td>
<td>5.1 (5.4)</td>
</tr>
<tr>
<td>Discharge other than home*</td>
<td>71.7% (2,292)</td>
</tr>
<tr>
<td>Variable direct hospital cost, $*</td>
<td>11,084 (15,682)</td>
</tr>
<tr>
<td>Median cost, $</td>
<td>6,614</td>
</tr>
</tbody>
</table>

*Length of stay, discharge to location other than home, and variable direct cost exclude patients who died during the hospitalization.

**TABLE 2. Association of Arousal and Outcomes**

<table>
<thead>
<tr>
<th>mRASS Alert and Calm, n = 2,737</th>
<th>mRASS Negative, n = 337</th>
<th>mRASS Positive, n = 158</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Value (SD)*</td>
<td>IRR/RR (95% CI)</td>
</tr>
<tr>
<td>Restraint use % (n)</td>
<td>4.2% (114)</td>
<td>Referent</td>
</tr>
<tr>
<td>In-hospital mortality % (n)</td>
<td>1.0% (28)</td>
<td>Referent</td>
</tr>
<tr>
<td>Length of stay, d (SD)*</td>
<td>4.9 (6.2)</td>
<td>Referent</td>
</tr>
<tr>
<td>Discharge other than home, % (n)*</td>
<td>24.9% (675)</td>
<td>Referent</td>
</tr>
<tr>
<td>Variable direct cost, $ (SD)*</td>
<td>10,581 (14,905)</td>
<td>Referent</td>
</tr>
<tr>
<td>Median cost, $</td>
<td>6,318</td>
<td></td>
</tr>
</tbody>
</table>

NOTE: IRR is calculated for length of stay and cost. RR is calculated for restraint use, discharge home, and in-hospital mortality. Abbreviations: CI, confidence interval; IRR, incident rate ratio; mRASS, Modified Richmond Agitation and Sedation Scale; SD, standard deviation.

NOTE: Missing variable direct costs were imputed. Abbreviations: MOYB, months of the year backward; mRASS, Modified Richmond Agitation and Sedation Scale; SD, standard deviation.

*Length of stay, discharge to location other than home, and variable direct cost exclude patients who died during the hospitalization.

Variable direct cost is calculated with imputation of missing cost data.
Impact of Altered Level of Arousal on Outcomes

There is an association between a deviation from a normal level of arousal (mRASS not equal to 0) and worsening outcomes (Table 2). Relative to a normal level of arousal (4.9 ± SD 5.2 days), decreased level of arousal (negative mRASS), and increased arousal (positive mRASS) resulted in longer LOS (6.0 ± SD 5.6 days, 5.7 ± SD 6.8 days, respectively). Similarly, increased or decreased arousal was associated with heightened risk of restraints and less frequent discharge to home. In-hospital mortality and hospital variable direct costs were significantly higher in those with decreased levels of arousal (IRR: 2.8, 95% CI: 1.3–6.0; IRR: 1.10, 95% CI: 0.95–1.26, respectively). The pattern does not hold for increased arousal with respect to in-hospital mortality and variable direct hospital cost outcomes. The unadjusted analysis found that, relative to normal arousal, there is a significant change in outcomes with decreased levels of arousal. Increased arousal is associated with worsened IRR in LOS, restraint use, and discharge home, but not in-hospital mortality and variable direct cost.

Impact of Altered Attention on Outcomes

Patients who completed the MOYB incorrectly had increased restraint use (RR: 2.11, 95% CI 1.44–3.11) and LOS (IRR: 1.06, 95% CI: 1.02–1.10), but no difference in in-hospital mortality, discharge home (RR: 0.78, 95% CI: 0.75–0.82), and variable direct costs, relative to those who completed the MOYB correctly (Table 3). Importantly, patients who did not complete the MOYB assessment had a 2-fold increase in restraint use (RR: 2.05, 95% CI: 0.94–4.50), in-hospital mortality was nearly 6-fold higher (RR: 6.36, 95% CI: 2.16–18.69), longer LOS (IRR: 1.12, 95% CI: 1.03–1.21), and returned home less frequently (RR: 1.77, 95% CI: 1.26–2.48).

Inter-relationship of Altered Level of Arousal and Attention on Outcomes

The inter-relationship of altered level of arousal and attention is presented in Table 3. Of patients with a normal mRASS, 52% had correct MOYB. The percentage of correct MOYB declined with the level of arousal, such that 38% had normal MOYB and a mRASS of –1 and 9% had normal MOYB with mRASS of ≤–2. In general, in-hospital outcomes (restraints and mortality) are associated with MOYB performance, and discharge outcomes (LOS, discharge location, and variable direct costs) are associated with mRASS. Those patients who did not complete the MOYB demonstrated worse outcomes, regardless of

<table>
<thead>
<tr>
<th>Table 3. Performance on mRASS, MOYB, and Outcomes</th>
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<tbody>
<tr>
<td>mRASS Abnormal</td>
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<tr>
<td>-----------------</td>
</tr>
<tr>
<td>IRR/RR (95% CI)</td>
</tr>
<tr>
<td>Restraint use, % (n)</td>
</tr>
<tr>
<td>In-hospital mortality, % (n)</td>
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<tr>
<td>Length of stay, d (SD)*</td>
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<td>Median cost, $</td>
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NOTE: IRR is calculated for length of stay and cost. RR is calculated for restraint use, discharge home, and in-hospital mortality. MOYB, months of the year backward; mRASS, Modified Richmond Agitation and Sedation Scale; RR, risk ratio; IRR, incident rate ratio; SD, standard deviation.

\*Length of stay, discharge other than home, and variable direct cost calculated with imputation of missing cost data (n = 168).
mRASS performance, including a 6-fold increase in mortality and significant increases in LOS and discharge location.

**DISCUSSION**

Impaired performance on a one-time assessment of arousal or attention during hospitalization demonstrated a relationship with in-hospital and discharge outcomes. Relative to normal levels of arousal and attention, alterations in attention, level of arousal, or both were associated with progressively negative consequences. Combined with the prognostic value, the administration of ultra-brief cognitive screening measures may have value in the identification of patients who would benefit from additional screening, supporting prior work in this area. The brevity of the assessments enhances clinical utility and implementation potential.

Cognitive function during hospitalization has been associated with many negative outcomes including delirium, falls, pressure ulcers, and functional decline. The findings of this analysis are consistent with previous studies and provide important clinical implications. First, prior work in cognitive screening has focused on more time-consuming instruments. By focusing on brief instruments, particularly those under 1 minute that do not require paper or props, a user-friendly tool that is associated with health outcomes is provided.

In addition, this analysis demonstrates that each assessment, when used individually, has some prognostic significance associated with the identification of delirium or other types of cognitive impairment. When used alone, abnormal scores on the mRASS or MOYB may be indicative of individuals requiring further cognitive assessment, supporting previous research. Individuals with abnormal scores on both the mRASS and MOYB identify a high-risk group in need of further clinical assessment for delirium (Figure 1). Neither of these assessments are meant to be used as the only means to diagnosis delirium, but together they identify key clinical characteristics of delirium (arousal and attention). Considering the significant negative consequences associated with delirium, it is not surprising that tools identifying core features of delirium, such as those presented here, would also be associated with in-hospital and discharge outcomes.

The quality improvement design of this project allowed the recording of outcomes in those who were unable or refused to complete the screening. This may be a potentially high-risk group who would otherwise go unnoticed. A recent editorial from the American and European Delirium Societies highlights that individuals who are unable or refuse to complete testing due to impaired arousal are neglected in the most recent American Psychiatric Association Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition definition of delirium. Further work to identify
and intervene on behalf of individuals who are unable to complete testing will aid in understanding arousal and its relationship to delirium and other disorders.

This analysis provides additional insight in the selection of measures of arousal and attention. Level of arousal is a complex concept that involves components of awareness and alertness, including external stimuli and self-awareness. As an ultra-brief measure of arousal level, the mRASS incorporates both external stimuli (asking an open-ended question) and self-awareness (describing current state) to determine basic cognitive function. Attention can be defined as the selection of stimuli for further cognitive processing. Administration of both the mRASS and MOYB is a useful bedside tool for clinicians to examine both basic cognitive function and more complex tasks of attention.

The quality improvement nature of this work has limitations and strengths that deserve mention. The significant strength of this work is the robust sample size. Also, trained staff not involved in the direct clinical care of patients administered the cognitive screens, suggesting that non–clinically trained personnel could be utilized for risk assessment. The major limitation is the restricted amount of covariate data that were collected. Data for this project were collected to operationalize and demonstrate the impact and business case of a delirium risk modification program, limiting the ability to perform adjustment for other covariates such as comorbidity and reason for admission. Also, due to the nature of this project, a diagnosis of delirium was not determined. A limitation of excluding in-hospital deaths from the cost analysis was that some individuals at high risk died early, thus costing less overall. Generalizability is limited by an over-representation of males within a single setting. Further use and understanding of the mRASS and MOYB in other populations is warranted and welcomed. Use of MOYB is also a limitation considering that scores are not standardized across patients or settings. Data regarding administration time of either of these tools were not collected; therefore, determining that these are ultra-brief assessments (<1 minute) is based on estimates. As such, these measures should not be the sole source of information for clinical evaluation and diagnosis.

CONCLUSION

This work found that impaired performance on brief cognitive assessments of arousal and attention in hospitalized patients were associated with restraint use, in-hospital mortality, longer LOS, less discharge home, and hospital costs. Routine screening of older patients with brief, user-friendly cognitive assessments upon admission can identify those who would benefit from additional assessment and intervention to alleviate individual and economic burdens.

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