Success Of An Expedited Emergency Department Triage Evaluation System For Geriatric Trauma Patients Not Meeting Trauma Activation Criteria

Background: Geriatric patients are at increased risk of injury following low-energy mechanisms and are less tolerant of injury. Current criteria for trauma team activation (TTA) often miss these injuries. We evaluated a novel triage process for an expedited Emergency Medicine Physician evaluation protocol (T3) for at-risk geriatric sub-populations not meeting trauma team activation (TTA) criteria.

Methods: Retrospective review of injured patients (≥65 years) from a Level II Trauma Center with an Injury Severity Score (ISS < 16), prior to (Pre-T3, Jan 2007-Oct 2009), and after (Post-T3, Jan 2010-Oct 2012), implementation of T3, as well as a contemporary period (CP, Jan 2013-Oct 2015). Demographics, physiologic variables, and timeliness of care were measured. Rates of ICU admission, operative procedures and lengths of stay and in-hospital mortality were compared for all periods. Logistic regression analysis determined variables independently associated with mortality.

Results: Post-T3, 49.2% of geriatric registry patients underwent T3 with a reduction in key time intervals. Median time to evaluation (42.1 mins vs 61.7 min, p<0.001), median time to CT (161.3 mins vs 212.9 mins, p<0.001) and EDLOS (364.6 mins vs 451.5 mins, p=0.023) were all reduced compared to non-expedited evaluations. There was no change in mortality after the implementation of the protocol.

Conclusion: The T3 protocol expedited patient evaluation of at-risk geriatric patients that would not otherwise meet TTA criteria. The new process met the goals of the American College of Surgeons Trauma Quality Improvement Program while conserving resources.

Keywords: trauma, triage, geriatrics

Introduction

Older adults, defined as those aged over 65, are at increased risk of significant injury following low-energy traumatic mechanisms and are also less tolerant of physiologic insult.\(^1\),\(^2\) Traditional triage thresholds for trauma activation that rely solely on abnormal vital signs, altered sensorium, anatomic injury, or the presence of a high-energy mechanism often fail to capture significantly injured patients in this population.\(^3\),\(^4\) Prior to November 2009, a significant proportion of older adults presenting to our trauma center did not meet our trauma team activation (TTA) criteria, and were evaluated by an Emergency Medicine physician (EMP), with subsequent consultation to the Trauma Service after injuries were identified. To expedite the evaluation of this vulnerable population, we created a new EMP-led third triage tier. Criteria for patient triage to “Tier 3” (T3) were designed to identify patients with occult head, neck, and torso injury.
following low-energy mechanisms without baseline neurologic or physiological derangements meeting TTA criteria.

Our outcomes of interest were time to physician evaluation (TPE), time to computed tomography (TCT), and Emergency Department (ED) length of stay in older patients following low-energy mechanisms. We hypothesized that the introduction of T3 would have a positive impact on these metrics. Although initiated in 2009, the addition of T3 aligns with the goals of the 2013 American College of Surgeons Trauma Quality Improvement Program (ACS TQIP) Geriatric Trauma Management guidelines.5

Methods
The study took place at a Level II state-accredited trauma center serving a broad urban and rural catchment area of approximately 450,000 people. Blunt mechanisms of injury predominate (93%). The center averages 1,500 TTAs, and 2000 trauma service admissions annually. Patients ≥ 65 years of age make up approximately 40% of our registry and 15% of the registry is aged ≥ 85. Emergency Department (ED) volume exceeds 130,000 contacts annually.

Study Design
The Reading Hospital Institutional Review Board approved the study. The IRB found that the study was a review and satisfied criteria for Federal regulation 45 CFR 164.512i with waiver of consent granted. Respect for patient data confidentiality was maintained as all analysis was performed on de-identified patient information. This study was performed in compliance with the Declaration of Helsinki.

We performed a retrospective review from our trauma registry of injured patients aged ≥ 65 years with an ISS < 16 meeting Pennsylvania Trauma Outcome Study (PTOS) inclusion criteria (Box 1).6 The first period of study (Pre-T3: Jan 2007 – Oct 2009), was prior to the introduction of the T3 protocol, during which an EMP evaluated patients not meeting TTA triage criteria. A learning period of 2 months (Nov 2009–Dec 2009) was used to implement the protocol. This study compared Pre-T3 outcomes with a similar time-frame following completion of the learning period (Post-T3, Jan 2010–Oct 2012), as well as a third comparable contemporary period (CP, Jan 2013–Oct 2015). The contemporary period was included to evaluate the sustainability of the impact on outcomes with T3 implementation.

Triage
Our center’s triage criteria delineate three levels of activation (Figure 1). A “Trauma Alert” (Level 1, highest level) activates the trauma team (Trauma Surgeon, EMP, Resident/Advanced Practitioner, Trauma Nurse (TN) x 2, Paramedic, Laboratory Technician (LT), Radiology Technician (RT), ED Registrar (EDR), and Chaplain) as well as an additional Anesthesia Provider, Respiratory Therapist, and delivery of 2 type-specific units of packed red blood cells (PRBCs). A “Trauma Response” (Level 2) activates the trauma team only without the immediate mobilization of the Anesthesia Provider or blood products. The T3 protocol (Level 3) mobilizes a board-certified EMP, TN, LT, RT, and EDR for expedited evaluation. Patients not meeting entry criteria for the first 3 levels of activation were triaged to a traditional non-expedited EMP evaluation (NE). The T3 evaluation consisted of a standard history and physical with emphasis on the primary and secondary trauma survey. The EMP then decided which advanced imaging studies were needed in addition to standard trauma laboratory studies. The Radiology Department also expedited the acquisition and reading of computed tomography studies ordered under the T3 process. The institution set an a priori goal of 30 mins for TPE, ≤ 90 mins for TCT, and ≤ 180 mins for ED LOS.

Descriptive And Outcome Variables
Table 1 lists descriptive variables from the electronic medical record (EMR). Time interval metrics included time from patient arrival to the initial EMP evaluation (TPE), time to first CT acquisition (TCT), and ED length of stay (EDLOS). Variables were then compared between the three periods. Data elements were missing in 2.8% of patients and such patients were excluded from the analysis. Upgrades from T3 to TTA were included in the T3 group, consistent with an intention-to-treat principle.

Statistical Analysis
All statistical analysis was performed using SPSS v24.0 (IBM corp., Armonk NY). Data analysis was conducted using

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Box 1 Pennsylvania Trauma Outcome Study (PTOS) Inclusion Criteria

- Trauma related Intensive Care Unit admission
- Trauma related step-down unit admission
- Trauma related Dead on Arrival
- Trauma related pronounced Dead after Arrival
- Trauma patients remaining in facility over 48 hrs
- Trauma patients remaining in facility between 36 and 48 hrs with an injury severity score (ISS) ≥ 9
- All admitted transfers in or out of facility
planned comparisons between periods, Pre-T3, Post-T3 and CP, in order to maintain the sequential integrity of the analysis and control for the experiment-wise error rate. A Chi-square analysis was used for the discreet variables and group t-tests for the continuous level variables. Significance was determined by a p-value of less than 0.05. There were no corrections used for multiple comparisons because some analyses were outcomes related and some analyses were conducted as the goodness of fit measures. Logistic regression was performed using mortality as the dependent variable and assessing the impact of the discrete variables of time-period, anticoagulant use, PRBC transfusion, and male gender (Referents: Pre-T3, no anticoagulated use, no PRBC transfusion and female gender, respectively). Continuous variables were Age, ISS, SBP and HR. The model examined was the enter method which assesses all variables without exclusion, and reports the beta coefficient, p-value, odds ratio and the 95% confidence interval of each variable. Model diagnostics were the initial model prediction used as a comparator for enhancement, the final model prediction as well as the Nagelkerke R2 value for all variables and the Hosmer–Lemeshow Goodness of fit statistic.

**Results**

Compared to the pre-T3 period, the new triage level resulted in a significant reduction in TPE and overall EDLOS in the post-T3 period (42.1 mins vs 61.7 mins p < 0.001 and 364.6 mins. vs 451.5 mins, p < 0.001, respectively). All 3-time intervals, TPE, TCT, and EDLOS were significantly reduced in the CP compared to the pre-T3 period (44.6, 125.7, and 370.0 mins respectively, p < 0.001) (Table 2).

Overall ED volume increased throughout the three periods of study (Table 1) for our study population. Post-T3, 49.2% of all admissions with an ISS < 16 were evaluated via the T3 pathway (CP, 43.8%). The proportion of patients undergoing TTAs declined after implementation of the protocol. Differences in mean age, admission GCS, admission SBP, and admission HR were not clinically meaningful. There was a change in injury pattern identification with a higher percentage of PTOS patients identified with significant head (AIS Head ≥ 3) and chest (AIS Chest ≥ 2) injuries through the three periods. For ISS < 16, there was no statistically significant change of in-house mortality with the implementation of the protocol (Pre-T3: 3.7% vs Post-T3: 2.7%, and CP: 2.3%, p<0.145). There was a statistically significant reduction in overall hospital LOS post-implementation that was sustained in the CP. Utilization of Pan-CT imaging increased significantly after implementation.

Logistic regression analysis incorporating the above-mentioned plausible variables showed that advanced age, lower GCS and lower SBP were independent predictors of mortality in this population (Table 3). The post T3 period was not associated with a mortality benefit.
Discussion

The novel triage process met our primary objective of decreasing key metric times in evaluating patients older than age 65: time to physician evaluation, time to CT scan, and overall emergency department length of stay. Although we did not meet our a priori target goals, the process did result in significant improvements.

The T3 process did not meet our secondary objective of lowering mortality but it did impact overall hospital length of stay when compared to those patients who underwent the standard non-expedited evaluation. This finding is not surprising given the overall lower ISS of the T3 cohort.

In the United States, 13% of the total population is over 65 and this proportion is expected to increase to over 20% by 2030.\(^7\,8\) Although the majority of injuries sustained by this subgroup are by low-energy mechanisms, this population currently represents 25% of all trauma-related admissions\(^5\,9\) and a similar proportion of injury-related deaths.\(^1\,2,10\) Pre-existing encephalopathy and co-morbid conditions often mask clinical presentation,
complicating clinical investigation and increasing vulnerability to injury.

Multiple strategies have been proposed for meeting this triage challenge. The ACS TQIP Geriatric Guideline currently recommends the following: 1) a lowered threshold for transporting injured patients to trauma centers, and 2) Level 1 or 2 conventional activations upon arrival. 3) A lower threshold for laboratory and radiologic investigation.

In our study, we reviewed a novel triage strategy that meets the goals of the ACS TQIP Geriatric Guideline while judiciously using both Emergency Department and trauma resources. The T3 pathway was utilized for evaluation of 43.8% of all older adult PTOS patients (CP) with ISS < 16. Despite this high percent, only 21% of patients triaged to T3 ultimately had an injury requiring admission to a surgical service (CP). A similar fraction was found to have medical conditions requiring non-surgical inpatient or observational care with the remainder being discharged. This observation suggests that a majority of T3 patients would not benefit from a full TTA as suggested by the ACS-TQIP Guideline.

There was an increase in the number of patients meeting PTOS criteria after implementing the T3 process. The subsequent decrease in the injury severity for extremity/pelvis is most likely a factor of including more patients with lower acuity injuries although meeting PTOS admission criteria.

Multiple studies have been performed evaluating the impact of triage and trauma team activation on geriatric outcomes. This study confirms the low sensitivity of triage activation criteria in capturing significant injuries in patients older than 65. It adds to the current understanding of geriatric triage in demonstrating that the goals of the ACS TQIPS can be achieved in a resource responsible manner while improving the quality of care.

Post-T3, the threshold for utilization of computed tomography imaging decreased significantly. Geriatric patients underwent Pan-CT imaging, defined as CT imaging of the brain, cervical spine, chest, abdomen and pelvis, much more frequently in the Post-T3 period. This increase was intentional as the working group initially challenged EMP to maintain a threshold for diagnostic evaluation comparable to the Trauma Service. Though

| Variable       | Coefficient | p-value | Odds Ratio | 95% CI |
|----------------|-------------|---------|------------|------|
| Intercept      | 1.138       | 0.383   | 1.831      | 0.982–3.413 |
| Post-T3 Period | 0.605       | <0.001  | 0.945      | 0.916–0.974 |
| Age            | −0.057      | <0.001  | 1.09       | 1.09–1.11 |
| GCS            | 0.411       | <0.001  | 1.509      | 1.403–1.623 |
| ISS            | −0.051      | 0.106   | 0.95       | 0.993–1.011 |
| Anticoagulant Use | −0.414     | 0.109   | 0.661      | 0.399–1.096 |
| PRBC in ED     | −0.306      | 0.142   | 0.737      | 0.490–1.108 |
| SBP            | 0.018       | <0.001  | 1.018      | 1.011–1.025 |
| HR             | −0.008      | 0.149   | 0.992      | 0.982–1.003 |
| Male gender    | −0.459      | 0.065   | 0.632      | 0.389–1.028 |
| Fall           | 0.057       | 0.963   | 1.059      | 0.552–2.030 |

**Table 2** Timeliness Of Care Before And After Implementation Of T3 Protocol

| TPE | n   | Pre-T3 | Post-T3 | CP   | p-value | Post-Hoc# |
|-----|-----|--------|---------|------|---------|-----------|
| Mean (SD) | 614 | 1298   | 1443    | <0.001 | Pre vs Post (p<0.01), Pre vs CP (p<0.001) |
| TCT | n   | 534    | 1201    | 1319  | 0.001   | Pre vs CP (p=0.001) |
| Mean (SD) | 212.9 (661.5) | 161.3 (550.9) | 125.7 (217.5) | 0.001 | Pre vs CP (p=0.001) |

| EDLOS | n   | 708    | 1449    | 1632  | <0.001 | Pre vs Post (p<0.001), Pre vs CP (p<0.001) |
|       | Mean (SD) | 451.5 (376.1) | 364.6 (277.9) | 370.9 (303.5) | <0.001 | Pre vs Post (p<0.001), Pre vs CP (p<0.001) |

Notes: #Omnibus p-value test. ANOVA analysis performed for three time periods with Scheffe post hoc test.
highly sensitive for injury identification, the ultimate clinical benefit of aggressive CT imaging has been questioned by several investigators.23–25 Post T3, confirmation of previously occult injuries, such as isolated rib fractures and stable spinal or pelvic fractures may have resulted in an increased number of patients meeting criteria for admission or observation with a lower overall mortality risk. The observed decline in the prevalence of significant extremity injuries, and the increase in significant head and chest injuries post-implementation supports this hypothesis.

Limitations
Our study has many of the anticipated limitations of a retrospective study. The data analysis is derived from a single-institution registry of only injured patients. The detailed outcomes of non-injured T3 patients are not available. In addition, our results may not be translatable to smaller EDs with a low prevalence of presenting older adult patients. Additionally, the period of study spans almost 10 years, and many improvements to our institution’s infrastructure have been implemented over that time. Although a significant mortality decline occurred concurrent with the implementation of the protocol, the observed outcomes may have resulted from other enhancements in care. However, like most improvements in care, such as massive transfusion protocols and damage control surgery apply to the most severely injured, those with ISS > 16, these changes should not affect our population studied. Further study of long-term mortality and functional outcomes is needed to characterize the ultimate impact of T3 in improving outcomes.

A potential confounder in this analysis was a conversion from AIS 2005 to AIS 2008 which occurred in January 2012. ISS calculations in the Pre-T3 time period used the former system, while scores Post-T3 used the latter. Some authors have raised the possibility of induced discrepancies by transitioning between versions.26,27

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Author Contributions
FBF, AO, CWS, ARM and CFB designed the study along with writing initial manuscript and edits. ARM and RT performed pilot chart review for design phase, had input into original manuscript and edits. APM extracted data for analysis. TW, APM, ALM and APS performed the statistical analysis and composition of Methods section along with input into original manuscript and design of the Tables. FBF, AO, APM, and TW interpreted the results. CFB and APS implemented the program in ED. FBF, AO, CFB, ARM, CAB, APS, CWS, ALM and APM all contributed to critical revisions of the final document. All authors contributed to data analysis, drafting and revising the article, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

Disclosure
The authors report no conflicts of interest in this work.

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