In the 14th-century alliterative poem *Sir Gawain and the Green Knight*, the Green Knight seeks to best King Arthur and his court in a game. The Green Knight says that he will allow whomever accepts the challenge to strike him with his own axe, on the condition that the challenger find him in exactly 1 year to receive a blow in return. Sir Gawain, the ostensibly least fitting of the Knights of the Round Table, leaps to take on the challenge himself and provide honor to his King. In many respects, the valiancy of emerging diagnostics is put to a similar test as reported in the present issue by Schöchl and colleagues [1].

Trauma-induced coagulopathy (TIC) has been identified early after patient presentation [2], is associated with shock and massive transfusion (MT) requirements, may occur in the absence of coagulation factor depletion [3], and has been reported to complicate up to 30% of major trauma admissions [4]. Accordingly, early and accurate diagnosis of TIC is essential to facilitate and optimize rapid provision of appropriate blood products in order to decrease morbidity and mortality, to promote efficient utilization of valuable resources, and to avoid wastage of nonessential therapy.

Conventional coagulation testing (CCT) remains the predominant method of assessing coagulation status worldwide, including the prothrombin time, activated partial thromboplastin time, platelet tests, and fibrinogen concentration. Despite this predominance, these tests have never been proven accurate in the hemorrhaging trauma patient [5] and suffer from a prolonged laboratory turnaround time of 1 hour or more in certain settings [6]. Finally, since they are plasma based, the assays do not accurately reflect the physiological milieu of whole blood, where multiple elements – including endothelium, platelet interactions, and subsequent thrombin generation – contribute to ultimate clot strength, leading to a meticulous balance of hemostasis and lysis. Accordingly, there has been recent growing enthusiasm for alternate methods of coagulation assessment.

Thromboelastography (TEG®; the name is a trademark of Haemoscope Corp. USA) and thromboelastometry (ROTEM®; the name is a trademark of Tem Innovations GmbH, Munich, Germany) tests that assess the viscoelastic properties of whole blood are relatively newer members of the diagnostic armamentarium currently under investigation for the assessment of patients at risk for TIC [7,8]. Both tests provide the potential to supersede CCT in the diagnosis of TIC because they provide real-time assessment of clot physiology from initial activation, subsequent achievement of tensile strength, and eventual resolution/degradation via a graphical tracing of fibrin polymerization and clot strength. Beyond the early diagnosis of TIC via point-of-care assessment, perhaps the most promising future for TEG®/ROTEM® testing is the ability to evalu...
ROTEM® technology is the additional potential benefit of early goal-directed therapy, with the objective to optimize and possibly reduce blood product utilization [9].

In the present issue of Critical Care, Schöchl and colleagues evaluated whole blood ROTEM® assays and the test’s ability to predict those seriously injured patients (all with Injury Severity Score >16) likely to require MT (>10 units red blood cells/24 hours) during the subsequent hospitalization [1]. Utilizing a variety of assays (extrinsically activated assay with tissue factor, intrinsically activated assay using ellagic acid, and fibrin-based extrinsically activated assay with tissue factor and the platelet inhibitor cytochalasin D), the authors assessed the clotting time, clot formation time, clot amplitude at the end of the clotting time, and maximum clotting firmness, and compared the predictive value of these tests with CCT and other physiologic parameters. Although Schöchl and colleagues noted that the best predictive tests for MT were a simple hemoglobin or Quick value, they observed that the predictive values of the clot amplitude at the end of the clotting time and the maximum clotting firmness for the fibrin-based extrinsically activated assay with tissue factor and the platelet inhibitor cytochalasin D were similar, with the chief advantage being the availability of results within 10 minutes.

At first glance, the astute clinician may argue that severely injured patients at risk for TIC can already be identified via CCT or clinical factors, such as a variety of scoring systems [10,11], and hence why the need for another potentially expensive test to provide the same information albeit somewhat earlier? Accordingly, as with many excellent studies, the fine details raise more questions than provide answers. As pointed out by the authors, the rapid availability of point-of-care whole blood assays affords a much more comprehensive assessment of the true relational hemostatic state of the individual patient. For instance, hyperfibrinolysis – which has recently been described by several investigators in the trauma setting [12-14] and can only be rapidly identified via viscoelastic tests – was clearly noted in 19 patients. Furthermore, the mean fibrinogen level in the MT group was 95 mg/dl. Given the emerging significance of hyperfibrinolysis in the pathophysiology of TIC [12], and the current use of fibrinogen concentrates for resuscitation in European centers, it would be interesting for the authors to correlate the frequency of this phenomenon in the MT group and its association with the fibrinogen level and clot strength. Furthermore, recent evidence supporting early use of antifibrinolytics [15] has prompted many centers to now include such therapy as a first-line adjunct to their MT protocols.

Schöchl and colleagues are to be congratulated for this important work, and we agree that current evidence continues to grow in support of point-of-care viscoelastic analysis of coagulation status. Of note, however, the application and adaption of this technology present significant challenges. For instance, it should be noted that the authors’ group have been working with these techniques for some time, and have no doubt moved beyond a significant learning curve for use of these techniques. Furthermore, recognizing the complex interactions of the hemostatic system, we posit that that TEG®/ROTEM® should be considered as a tool to better appreciate relational hemostatic mechanisms that probably coexist in TIC. While we agree with the authors that ROTEM® can affirm the clinician/surgeon a rapid diagnosis of TIC, it must be noted that ROTEM®-based assessments were similar to CCT when only simply applied for predicting MT.

In sum, the authors have clearly shown that TEG®/ROTEM® holds promise as an important tool for the rapid diagnosis of TIC. We encourage the authors to continue their investigations, moving beyond observational trials to therapeutic interventions, utilizing this technology to improve our understanding of the complex physiologic changes to the coagulation system induced by significant injury. Such progress will firmly establish viscoelastic analyses as a notable and valiant member of the diagnostic roundtable.

Abbreviations
CCT, conventional coagulation testing; MT, massive transfusion; TEG, thromboelastography; TIC, trauma-induced coagulopathy.

Competing interests
The authors declare that they have no competing interests.

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