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Trauma-Induced Coagulopathy: The Past, Present, and Future

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Abstract

Trauma remains a leading cause of death worldwide, and most early preventable deaths in both the civilian and military settings are due to uncontrolled hemorrhage, despite paradigm advances in modern trauma care. Combined tissue injury and shock result in hemostatic failure, which has been identified as a multi-dimensional molecular, physiologic, and clinical disorder termed trauma-induced coagulopathy (TIC). Understanding the biology of TIC is of utmost importance as it is often responsible for uncontrolled bleeding, organ failure, thromboembolic complications, and death. Investigations have exposed that TIC is characterized by multiple phenotypes of impaired hemostasis due to altered biology in clot formation and breakdown. These coagulopathies are attributable to tissue injury and shock and encompass underlying endothelial, immune, and inflammatory perturbations. Despite the recognition and identification multiple mechanisms and mediators of TIC and the development of targeted treatments, the mortality rates and associated morbidities due to hemorrhage after injury remain high. The purpose of this review is to examine the past and present understanding of the multiple distinct but highly integrated pathways implicated in TIC to highlight the current knowledge gaps and future needs in this evolving field, aimed at reducing morbidity and mortality after injury.

Keywords

Blood Coagulation Disorders; Exsanguination; Hemorrhagic Shock; Hemostasis; Trauma

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Introduction

Trauma remains a leading cause of death worldwide(1), and it is expected that rates of death due to injury will continue to rise as populations age (2). In the 1970s, Trunkey described a trimodal distribution of trauma deaths (3), with immediate deaths thought to be nonpreventable and due to hemorrhage. However, since that time there have been paradigm changes in our understanding of death due to hemorrhage including the recognition of trauma-induced coagulopathy (TIC), as well as advances in resuscitation techniques designed to mitigate the effects of TIC (4). It is now clear that one-quarter to one-third of hemorrhaging trauma patients suffer from TIC, a multi-phenotypic disease state that encompasses disorders of coagulation and inflammation characterized by impairments in clot formation, breakdown, and overall vascular homeostasis. TIC is associated with increased early transfusion requirements, the development of organ failure, and mortality (5, 6). Death from hemorrhage is not the sole consequence of TIC. Due to the innate crosstalk between coagulation and inflammation, there are widespread adverse downstream inflammatory and immune consequences associated with early trauma coagulopathies, including organ dysfunction and thromboembolic complications(6).

Despite these focused investigations into the mechanisms and mediators of TIC and improvements in care related to identification and treatment, hemorrhage related mortality continues to be the most common cause of preventable deaths after injury (7). Therefore, targeted scientific focus on advancing our understanding of TIC across molecular and clinical realms remains of utmost importance to rescue a large percentage of injured patients from death due to early hemorrhage and late organ failure. Although substantial advances have been made, the links between the injury induced impairments of each of the various mechanisms that contribute to the failure of clot formation, lysis, and vascular homeostasis remain open science. In addition, the field is desperate for clear causal and intervenable pathways between coagulation, inflammation, and immune dysfunction. However, mechanistic studies related to TIC are fraught with the limitations of analyzing blood and vascular homeostasis in ex-vivo environments that often lack endothelium, flow, and local tissue injury milieu while extrapolating microvascular environments into macro measurements. The overall purpose of this review is to understand the past and present in order to highlight the needed future directions of this evolving field.

History

Injury induced disordered clotting that is associated with bleeding and death is not a modern concept. Historic battlefield descriptions of this date back to the eighteenth century. During the Korean and Vietnam Wars, patients in hemorrhagic shock were identified to have prolonged prothrombin and partial thromboplastin times early after injury and prior to interventions, hemodilution, or hypothermia, ultimately requiring more blood and having higher mortality(8). What followed was evolution of purported mechanisms of this coagulopathy including an initial focus on consumptive states and dilutional effects of resuscitation with fluids and bloods products.

These observations of coagulopathy during major wars stimulated significant military and civilian trauma research, and further critical concepts emerged including the recognition of the clear transition from a clinically hypocoagulable to hypercoagulable state and the central role that hypoperfusion due to shock played. Multiple groups simultaneously identified that hemorrhage and resuscitation induced depletion of coagulation factors and dilutional coagulopathy led to an irreversible physiologic collapse(9). These concepts were formally coalesced into the 'Bloody Vicious Cycle' by Kashuk and colleagues in the early 1980s(10). Following this, it was increasingly recognized that post-injury hemorrhage exacerbated by the 'lethal triad' of hypothermia, acidosis, and coagulopathy was a vicious cycle that led to more coagulopathy and ultimately high mortality rates(11). This increasing recognition led to changes in clinical practice in the 1980s with formalization of damage control surgical and resuscitative techniques to halt the cycle of acidosis, hypothermia, and coagulopathy(12). In the early 2000s, Brohi, Cohen, and colleagues formally described the entity of acute traumatic coagulopathy in human and animal models, that reflected an endogenous biology independent of resuscitation, required combined tissue injury and shock, and had significant associations with poor outcomes and mortality $(6, 13-16)$. Although modern study of trauma patients has identified no significant difference in the presentation core temperature or amount of prehospital crystalloid given to patients that developed TIC requiring transfusion(17), there remains evidence that resuscitation and hypothermia continue to contribute to TIC(18), but are not necessary prerequisites. However, patients who develop TIC are profoundly more acidemic, supporting the pivotal importance of a shock state in the setting of tissue injury (17).

Finally, critical in the evolution of our understanding of TIC beyond consumption of coagulation factors, was the progress made in defining coagulation outside of the classic enzymatic coagulation cascade. In 2001, Hoffman and Monroe proposed the 'cell-based model of hemostasis' based on overlapping events of initiation (extrinsic pathway on tissue factor bearing cells), amplification (positive feedback of thrombin on platelets), and propagation of clotting (intrinsic pathway on activated platelets) that were regulated by cell surfaces rather than enzymatic protein and protease reactions alone(19). This model of hemostasis magnified research focused on multiple pathways contributing to TIC, rather than solely on a dilutional failure of the enzymatic processes of clot formation, and a large body of investigation has contributed to the current multi-dimensional understanding of the biologic contributors of TIC, to be addressed specifically below and summarized in Table 1.

However, despite years of basic and translational investigations of TIC, in 2013, Gando and the Standardization Committee on Disseminated Intravascular Coagulation (DIC) of the International Society on Thrombosis and Haemostasis reported that what was termed TIC was similar to consumptive states like DIC(20). However, best evidence at this time suggests that although TIC encompasses the criteria of DIC, it remains distinct from and not sufficiently characterized by the broad less-precise major (low platelet count, prolonged prothrombin time, increased soluble fibrin or fibrin-degradation products) and specific (low antithrombin, low protein C, and increased thrombin-antithrombin complexes) DIC criteria outlined by the Standardization Committee on DIC of the International Society on Thrombosis and Haemostasis (21) (Table 2). The definition of TIC has been in evolution for decades, shifting from the perception of a resuscitation induced dilutional coagulopathy, to a

multifactorial and multi-mechanistic event. TIC is distinct and more complex than the DIC (Table 2), covering a wide range of impaired clot formation and lysis, in combination with failure of vascular homeostasis and immunoactivation resulting in multiple clinical phenotypic states that can cause pathologic bleeding, clotting, organ failure, and death.

Mechanisms and Mediators

Table 1 depicts the estimated average quantitative values (mean or median) for each listed coagulation parameter in trauma patients with TIC in the early and later periods after injury and without TIC, supported by existing literature, as cited (16, 17, 21–31).

Tissue Injury and Shock

Impaired coagulation after injury was long thought to be solely due to iatrogenic causes(9). For decades, trauma resuscitation practices guided by the luminaries of shock research were based on restoring flow and oxygen carrying capacity with large volumes of packed red blood cells (oxygen) and crystalloid (flow). The creep of resuscitation toward larger and larger volumes of cold anemic resuscitation resulted in an iatrogenic coagulopathy characterized by impaired thrombin production and platelet function due to hypothermia, acidosis, and dilutional coagulopathy $(9-11)$. Although these iatrogenic effects are still contributors to coagulopathy and can coexistent with TIC, they are termed iatrogenic coagulopathy (IC), are separate in biology from TIC, and are fortunately more infrequent with modern hemostatic and goal-directed resuscitation techniques. An enlarging body of work has now demonstrated that to induce the mechanistic and clinical phenotypes of TIC, tissue injury (thought to activate the clotting cascade, produce thrombin, and stimulate resultant anticoagulant pathways) must be combined with tissue hypoperfusion (thought to release damage-associated molecular patterns [DAMPS], activate the contact pathway, induce thrombomodulin and endothelial protein C receptor [EPCR] expression at the endothelial surface to activate protein C). It has become clear that shock is essential to development of the phenotypes of TIC via multiple mediators described below.

Activation and Depletion of Protein C

One of the primary and first described mechanisms of TIC is activation of protein C system(13, 14, 16, 32, 33). Protein C is a serine protease with dual anticoagulant functions: induction of proteolytic cleavage of factors Va and VIIIa as well as a derepression of fibrinolysis through inhibition of plasminogen activator inhibitor-1 (PAI-1) (resulting in increased tissue plasminogen activator [t-PA] activity)(13, 14, 32). Since the initial description that severe injury combined with shock resulted in increased levels of activated protein C (aPC), concomitant reductions in coagulation factor V (FV) and factor VIII (FVIII), and increased fibrinolysis, a series of human studies and mechanistic animal and invitro cell culture models have confirmed these findings(13–16, 32). The activation of this pathway starts with tissue injury driven thrombin production that combines with thrombomodulin, the EPCR. The non-activated zymogen form of protein C on the endothelial surface is then activated and results in cleavage of FVa and FVIIIa and derepression of fibrinolysis. Multiple studies have corroborated that approximately 25–33%

of severely injured patients have elevated aPC, which is associated with increased blood transfusion and mortality, and in patients who survive, higher rates of infection and single and multiple organ failure(13, 14, 16, 32).

Interestingly, aPC also has cytoprotective functions including stabilization of endothelial and epithelial junctions, anti-apoptosis, and cleavage of extracellular histones. The enhanced activation of the protein C system may therefore represent an evolutionary maladaptive response. As humans may not be evolved to survive the massive injury created by being shot, stabbed, and run over by automobiles, it is hypothesized that the human physiologic response to these events is likely activating a large amount of protein C in an attempt to activate inflammomodulatory pathways to survive after severe trauma. The unfortunate sequela of this 'too much of a good thing' response is a systemic and local anticoagulation resulting in increased bleeding with incumbent enhanced mortality and morbidity. For those who survive their initial injury there is a rapid transition to a state of protein C depletion, characterized by hypercoagulability and impaired cytoprotectivity. Based on study of circulating aPC levels in trauma patients (16), the cytoprotective level can be assumed to be somewhere in the range of 1.05–6.00 ng/mL, as these levels correspond with preservation of Factor Va, VIIIa, and t-PA, as well as lower D-dimer levels. Conversely, the maladaptive level can be assumed to be in the >6.00ng/mL range as this level was shown to be associated with markedly decreased Factor Va and VIIIa, as well as increased t-PA and D-dimer levels (16).

The transition from hypo to hypercoagulability occurs very rapidly. Somewhere between 6 and 24 hours after injury the majority of patients are undergoing this transition, with 90% or more being normal or hypercoagulable in global hemostatic measures by 24 hours after injury, regardless of TIC (34). In addition, the coagulation factor activity in patients with and without TIC begin to rebound for most factors by 12 hours with significantly higher than baseline activity by 72 hours (17). The depletion of the non-active zymogen form of protein C is associated with both a transition to a thrombotic phenotype, and importantly a loss of cytoprotective signaling and resultant organ failure and infectious complications(16). With newly engineered activated forms of protein C which have augmented cytoprotective function with little or no anticoagulant protease activity, as well as antibodies or aptamers against the anticoagulant function of protein C, one can imagine in the near future a resuscitation where we might inhibit the anticoagulant function and concurrently augment the cytoprotective function of protein C immediately after trauma. Further precision medicine modulation of the relative amount of inhibition and augmentation could be performed as the patient moves through their post-injury coagulation and inflammation phases.

Factor Depletion, Impaired Thrombin Generation, and Fibrinogen Deficiency

Factor depletion does indeed exist after trauma(17). In addition, studies have demonstrated that clotting factor deficiencies occur without significant fluid resuscitation, immediately after injury, and are associated with worse outcomes(35). While reproducible and in

correlation with severity of injury and degree of shock, the factor level nadirs do not often reach the historically reported 20–30% level where coagulation should be impaired(17). Of course, the critical values of factor depletion were derived from closed studies in nonactivated (non-injury) blood(36). Whether the reductions in factor levels do cause tissue specific impaired coagulation at the site of injury that requires levels higher than 20–30% in the setting of injury, and what circulating levels are ideal after significant injury for both tissue specific hemostasis and survival remains an open experimental and clinical question.

Clot formation requires thrombin generation on injured tissues. There are some suggestions that the measurement of thrombin generation is an optimal real-time assessment of coagulation capacity in the setting of injury because thrombin generation does not cease with fibrin deposition. However, both hypocoagulable and hypercoagulable thrombin generation profiles have been associated with trauma with variable associations to bleeding and thromboembolic outcomes(37, 38). Studies do suggest that there is sufficient thrombin generation in TIC(22, 38), yet both trauma patients with and without TIC show evidence of increased thrombin-antithrombin, and prothrombin fragments(39), and the biochemical milieu of injury and shock may increase thrombin generation due to decreased antithrombin activity(40).

Fibrinogen is the terminal substrate for clot formation and low quantity or quality of fibrinogen in TIC is associated with bleeding and death(41, 42). Animal models of TIC confirm low levels of fibrinogen(43). Fibrinogen replacement in the form of cryoprecipitate or newer fibrinogen concentrates is used in many protocols for treatment of TIC (44), and often to maintain levels of fibrinogen well above standard triggers for fibrinogen replacement despite weak supporting evidence for this(45). Whether this technique improves outcomes continues to be a moving target as goal-directed treatment of hypofibrinogenemia remains without clear evidence-based treatment cutoffs, but with ever-broadening therapies beyond blood components.

Altered Post-Injury Platelet Biology

Although the pivotal importance of platelets in the overlapping stages of clot formation is highlighted in Hoffman and Monroe's 'cell-based model of hemostasis', understanding the effect of local versus diffuse injury and shock states on platelet biology remains unclear in TIC(41, 46–48). However, platelets contribute significantly to the strength of clot formation in systemic blood during the post-injury state. Kornblith and colleagues demonstrated that the platelet contribution to clot strength is higher than that of fibrinogen at all time points after injury, using functional assessment of whole blood in trauma patients(41). In addition, thrombocytopenia is a poor prognostic marker following injury and is independently associated with transfusion, progression of brain injury, and death after injury(47). However, even with normal platelet counts, nearly half of injured patients demonstrate impaired platelet aggregation in aggregometry assays immediately after injury(24, 46), which has been replicated in animal models of injury and hemorrhagic shock(49). Excessive platelet activation(24) with subsequent exhaustion is one of the proposed mechanisms. This injury induced alteration in platelet biology has been found to have independent associations to brain injury, severe injury, and shock, and all injury patterns consistent with significant

endothelial damage(24). Multiple studies have demonstrated increased morbidity and mortality in patients that have impairments in platelet aggregation after injury(24, 46).

Numerous unresolved aspects of post-injury platelet biology remain to be addressed. For one, due to the nature of ex-vivo aggregometry assays (lack of endothelium, lack of flow, exogenous agonist stimulation of platelets), it remains unclear where on the spectrum of adaptive to maladaptive responses to injury impaired post-injury platelet aggregation lies. It should not be ignored that the majority of investigations in this area have relied on point-ofcare platelet function assays that were intended to assess the effects of anti-platelet medication on platelet inhibition, and of concern is that viscoelastic assays do not always correlate well with point-of-care assessment of platelet function in trauma patients (50). In fact, using multiple electrode aggregometry, unpublished work by the authors of this review has identified that platelets may be endogenously activated by injury, diminishing their aggregation response to exogenous agonists. This raises concern that the identification of impaired platelet aggregation after injury may detect multiple phenotypes of post-injury platelet biology including a physiologic response (platelets activated by injury and unable to respond further to platelet activating agonists) and a maladaptive impairment in platelet aggregation. Therefore, future investigations need to use additional methods to assess the health and function of the post-injury platelets by incorporating endothelium and flow (microfluidics), assessment of structure (microscopy), mitochondrial health (mitochondrial respiration), and improved biomarkers of platelet and endothelial function(48, 51–53).

Additionally, given platelet function extends beyond coagulation to bidirectional endothelial interaction and regulation(54), control of local fibrinolysis at injury sites($19, 55$), and their role as core effector cells in local and systemic inflammation(56, 57), a focus on multiple areas of post-injury platelet biology should remain an active area of TIC science. Finally, an expanded understanding of post-injury platelet biology is critical to improving the care of post-injury hemorrhage. The standard-of-care in TIC is platelet transfusion as part of a balanced resuscitation ratio (regardless of platelet count)(7), but investigations have identified that not only does impaired post-injury platelet aggregation fail to predict the need for platelet transfusion(58), platelet transfusion does not reverse post-injury impairment in platelet aggregation(59) and does not improve outcomes for patients on anti-platelet medications who have brain injury(60). Multiple studies have demonstrated that platelet transfusions may increase morbidity and mortality after injury(59). The explanative mechanisms for this are unknown, but it is without a doubt that further studies of TIC related post-injury platelet biology, platelet transfusions, and alternative platelet therapies are required(61).

Dysregulated fibrinolysis

Local hypercoagulability promoting hemostasis at the site of injury is proposed to activate systemic fibrinolysis to "guard against thrombosis" in remote non-injured tissue(62). Elevated fibrinolysis measured by viscoelastic assays has repeatedly been demonstrated to be associated with a high mortality rate and massive transfusion in trauma, but found in less than 20% of the most severely injured trauma patients(32, 63). While excessive fibrinolysis has pathologic consequences, low fibrinolytic activity in animals has also been demonstrated

to cause microvascular occlusion to vital organs, irreversible recovery from hemorrhagic shock(64), and be reversible by administration of a profibrinolytic agent(65).

Moore and colleagues identified that low fibrinolytic activity measured by viscoelastic assays, termed fibrinolysis shutdown, has also been associated with increased mortality in severely injured patients(66). Low fibrinolytic activity (defined by a lysis at 30 minutes by thromboelastography [LY30] <0.9%) has been associated with increased mortality compared to moderate levels of fibrinolysis (LY30 0.9–2.9%) at multiple large volume trauma canters(66–68). Fibrinolysis shutdown early after injury may be protective in some patients(68). This coagulation change could counter balance platelet inhibition and prolonged prothrombin time that has been described in patients with evidence of prior fibrinolytic activation (67, 69). Historic and recent observations in trauma support that patients can present to the hospital with a spectrum of fibrinolytic activity in which patients at the pathologic extremes are at risk of increased mortality. This concept has argued for the selective use of antifibrinolytics, supported by a reduction in mortality by use in goal directed resuscitation(70). Specifically, concerns have risen that TXA may cause harm in certain trauma patients, in particular patients with moderate (physiologic) levels of fibrinolysis(71). Patients who receive TXA are also at risk of prolonged fibrinolysis inhibition. Fibrinolysis shutdown beyond 24 hours of injury is associated with increase in mortality and ventilator requirements(72, 73), and Myers, Neal and colleagues identified in a propensity matched retrospective review of a modern population of trauma patients that TXA may be an independent risk factor for the development of VTE after injury (74). In aggregate, evidence supports the need for improved understanding of complex biologic and clinical interactions and outcomes related to TXA administration in trauma patients.

Inflammation and Immune Dysfunction: Endotheliopathy, Damage-Associated Molecular Patterns (DAMPs), and Others

While the effects of trauma on the biochemistry of coagulation are the subject of much important work, concurrent progress has been made on elucidating the effects of injury and shock on inflammatory and immune dysfunction. Emerging literature describes a conglomeration of sterile inflammatory and immune responses that center around the endothelium, which are related but unique to biochemical coagulopathies of TIC (52, 75– 78). The injury response establishes a cascade of inflammatory and immune dysfunction resulting in single and multiple organ failure (acute kidney injury, Acute Respiratory Distress Syndrome, hepatic dysfunction) and a higher susceptibility to infection(79, 80). Much has been done in septic models, but it is more recently that bridging work has been done between coagulation and inflammation in trauma.

Endotheliopathy

Dr. Kozar and others have identified degradation of the endothelial glycocalyx after trauma. Further work has suggested that catecholamine surge is associated with glycocalyx degradation and associated auto-heparinization that can be identified on viscoelastic assays(81, 82) In addition, clinical studies have identified increased levels of Syndecan-1, a degradation product of the endothelial glycocalyx, to be associated with inflammation,

coagulopathy, and mortality(27). Important to this, plasma has been identified to be

restorative to the endothelial glycocalyx(79, 83), but clinical studies of early plasma administration have had mixed results in protection from morbidity and mortality(84, 85). Other pathways may also contribute to the endotheliopathy of trauma. Multiple investigators have reported aPC's cytoprotective functions including endothelial barrier protection, antiapoptosis, and cleavage of extracellular histones. Kutcher and colleagues identified that release of extracellular histones in the setting of injury and shock are associated with organ failure and death, and the compensatory activation of protein C and subsequent clearance of histones was found to be protective(86). This remains an active area of investigations, and there are hypothesis generating investigations that endothelial leak primes neutrophil reactive oxygen species release through the alternative pathway of complement(87), which could be mediated by coagulation products. Given that cultured endothelial cells exposed to plasma from coagulopathic trauma patients have increased permeability consistent with endothelial barrier dysfunction, suggesting circulating mediators in trauma patients can directly impair the endothelium, significant further work to characterize, diagnose, and treat the endotheliopathy of trauma is needed.

DAMPs and Others

The spectrum of DAMPs following injury remains a confusing mix of individual and combined mediators of biologic interest including: cytokines, soluble receptor for advanced glycation end-products [s-RAGE], high mobility group protein B1 [HMGB1], tissue inhibitor metalloproteinases-3 [TIMP-3], and numerous others(26, 30, 88, 89). These DAMPs have been measured and implicated in the sterile inflammation of trauma and the crosstalk between coagulation and inflammation(26, 30, 88, 89). Beyond this, many mechanisms abound; myosin may be implicated as a link between coagulation and inflammation biology and effects on TIC and fibrinolysis(90); metabolites are likely underappreciated in their contribution to TIC and post-injury hemorrhage(91); the relative production of alpha and meizo-thrombin is being investigated as a switch between coagulation and sterile inflammation after trauma(92). Indeed, whether each of these mediators has true mechanistic effect or is just a correlative measure of injured patients remains an important question which will be answered only by a combination of both reductionist fundamental mechanistic biology and multivariate big data approaches to clinical data. As highlighted by Dr. Timothy Billiar in PLOS Medicine in 2017 (93) and at his 2019 Western Trauma Association Founders Basic Science Lecture, both DAMP and pathogen-associated molecular pattern (PAMP) signaling are the scientific frontier of TIC and an expanded focus on trauma-induced immune activation in response to both sterile signal and microbial signal should guide the future of this field.

The Future

Improved assays

The evolution of identification and measurement of TIC has followed the evolution in mechanistic understanding. Originally TIC was identified by focusing on single, static, exvivo assessments of enzymatic coagulation cascade by measuring prolonged conventional

coagulation assays(6, 94), however functional assessment of whole-blood clot formation and degradation in real time via viscoelastic assays for both the identification and the management of TIC has become standard-of-care (70, 95). With the knowledge gained of the cellular and biochemical milieu created by combined tissue injury and shock, concerns have emerged regarding the ability to use ex-vivo conventional coagulation and viscoelastic assays for TIC interpretation, identification of 'normal ranges' of assays, and guidance of therapy.

Furthermore, there exists little ability to characterize and dynamically track the multiple competing and overlapping phenotypes which comprise TIC. Emerging evidence has highlighted that there are both clinical and biologic phenotypes of TIC and single assays are therefore ineffective for the purpose of identification and treatment of TIC(96). In addition, the optimal combination of assays that completely and effectively measure the complex and integrated pathways of all aspects of TIC related failure in coagulation, inflammation, and vascular homeostasis are unknown and perhaps undeveloped. Current best evidence supports that TIC can be identified by abnormalities in either conventional coagulation or viscoelastic assays, but not always both(96). Given this, combining multiple assays for diagnosis and therapy in TIC is likely the optimal strategy at this time. Future development of improved methods including in vivo biosensors of the biochemical milieu targeting assessments of circulating cells in flow and shear environments with the contribution of hematocrit and endothelium(97) should be a focus in the evolution of TIC science.

In-Silico Modeling

Ultimately however, we envision that biologic assays will be rapidly replaced by multivariate modeling of the complex coagulation milieu after trauma. This will be augmented by computational models which can target reductionist precision medicine approaches to resuscitation(98), while also predict and characterize dynamic phenotypes. This may allow for targeted predication and optimal personalized treatment rules for clinicians to use on their individual patients(99, 100). It is clear that we still have much to accomplish in the open science of TIC toward reducing morbidity and mortality for trauma patients.

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References:

- 1. Norton R, Kobusingye O. Injuries. N Engl J Med. 2013;368:1723–30. [PubMed: 23635052]
- 2. Rhee P, Joseph B, Pandit V, Aziz H, Vercruysse G, Kulvatunyou N, Friese RS. Increasing trauma deaths in the United States. Ann Surg. 2014;260:13–21. [PubMed: 24651132]
- 3. Baker CC, Oppenheimer L, Stephens B, Lewis FR, Trunkey DD. Epidemiology of trauma deaths. Am J Surg. 1980;140:144–50. [PubMed: 7396078]
- 4. Holcomb JB, Tilley BC, Baraniuk S, Fox EE, Wade CE, Podbielski JM, del Junco DJ, Brasel KJ, Bulger EM, Callcut RA, Cohen MJ, Cotton BA, Fabian TC, Inaba K, Kerby JD, Muskat P, O'Keeffe

T, Rizoli S, Robinson BR, Scalea TM, et al. Transfusion of plasma, platelets, and red blood cells in a 1:1:1 vs a 1:1:2 ratio and mortality in patients with severe trauma: the PROPPR randomized clinical trial. JAMA. 2015;313:471–82. [PubMed: 25647203]

- 5. Cohen MJ, Christie SA. New understandings of post injury coagulation and resuscitation. International journal of surgery. 2016.
- 6. Brohi K, Singh J, Heron M, Coats T. Acute traumatic coagulopathy. J Trauma. 2003;54:1127–30. [PubMed: 12813333]
- 7. Cannon JW. Hemorrhagic Shock. N Engl J Med. 2018;378:370–9. [PubMed: 29365303]
- 8. Simmons RL, Collins JA, Heisterkamp CA, Mills DE, Andren R, Phillips LL. Coagulation disorders in combat casualties. I. Acute changes after wounding. II. Effects of massive transfusion. 3. Postresuscitative changes. Ann Surg. 1969;169:455–82. [PubMed: 5774736]
- 9. Dunn EL, Moore EE, Breslich DJ, Galloway WB. Acidosis-induced coagulopathy. Surg Forum. 1979;30:471–3. [PubMed: 538668]
- 10. Kashuk JL, Moore EE, Millikan JS, Moore JB. Major abdominal vascular trauma--a unified approach. J Trauma. 1982;22:672–9. [PubMed: 6980992]
- 11. Rossaint R, Cerny V, Coats TJ, Duranteau J, Fernandez-Mondejar E, Gordini G, Stahel PF, Hunt BJ, Neugebauer E, Spahn DR. Key issues in advanced bleeding care in trauma. Shock. 2006;26:322–31. [PubMed: 16980877]
- 12. Stone HH, Strom PR, Mullins RJ. Management of the major coagulopathy with onset during laparotomy. Ann Surg. 1983;197:532–5. [PubMed: 6847272]
- 13. Brohi K, Cohen MJ, Ganter MT, Matthay MA, Mackersie RC, Pittet JF. Acute traumatic coagulopathy: initiated by hypoperfusion: modulated through the protein C pathway? Ann Surg. 2007;245:812–8. [PubMed: 17457176]
- 14. Cohen MJ, Brohi K, Ganter MT, Manley GT, Mackersie RC, Pittet JF. Early coagulopathy after traumatic brain injury: the role of hypoperfusion and the protein C pathway. J Trauma. 2007;63:1254–61; discussion 61–2. [PubMed: 18212647]
- 15. Chesebro BB, Rahn P, Carles M, Esmon CT, Xu J, Brohi K, Frith D, Pittet JF, Cohen MJ. Increase in activated protein C mediates acute traumatic coagulopathy in mice. Shock. 2009;32:659–65. [PubMed: 19333141]
- 16. Cohen MJ, Call M, Nelson M, Calfee CS, Esmon CT, Brohi K, Pittet JF. Critical role of activated protein C in early coagulopathy and later organ failure, infection and death in trauma patients. Ann Surg. 2012;255:379–85. [PubMed: 22133894]
- 17. Kutcher ME, Kornblith LZ, Vilardi RF, Redick BJ, Nelson MF, Cohen MJ. The natural history and effect of resuscitation ratio on coagulation after trauma: a prospective cohort study. Ann Surg. 2014;260:1103–11. [PubMed: 24846092]
- 18. Lester ELW, Fox EE, Holcomb JB, Brasel KJ, Bulger EM, Cohen MJ, Cotton BA, Fabian TC, Kerby JD, O'Keefe T, Rizoli SB, Scalea TM, Schreiber MA, Inaba K, group Ps. The impact of hypothermia on outcomes in massively transfused patients. J Trauma Acute Care Surg. 2019;86:458–63. [PubMed: 30444856]
- 19. Hoffman M, Monroe DM 3rd. A cell-based model of hemostasis. Thromb Haemost. 2001;85:958– 65. [PubMed: 11434702]
- 20. Gando S, Wada H, Thachil J, Scientific, Standardization Committee on DICotISoT, Haemostasis. Differentiating disseminated intravascular coagulation (DIC) with the fibrinolytic phenotype from coagulopathy of trauma and acute coagulopathy of trauma-shock (COT/ACOTS). J Thromb Haemost. 2013;11:826–35. [PubMed: 23522358]
- 21. Taylor FB Jr., Toh CH, Hoots WK, Wada H, Levi M, Scientific Subcommittee on Disseminated Intravascular Coagulation of the International Society on T, Haemostasis. Towards definition, clinical and laboratory criteria, and a scoring system for disseminated intravascular coagulation. Thromb Haemost. 2001;86:1327–30. [PubMed: 11816725]
- 22. Dunbar NM, Chandler WL. Thrombin generation in trauma patients. Transfusion. 2009;49:2652– 60. [PubMed: 19682336]
- 23. Scherer RU, Spangenberg P. Procoagulant activity in patients with isolated severe head trauma. Crit Care Med. 1998;26:149–56. [PubMed: 9428558]

- 24. Jacoby RC, Owings JT, Holmes J, Battistella FD, Gosselin RC, Paglieroni TG. Platelet activation and function after trauma. J Trauma. 2001;51:639–47. [PubMed: 11586152]
- 25. Kornblith LZ, Robles AJ, Conroy AS, Hendrickson CM, Calfee CS, Fields AT, Callcut RA, Cohen MJ. Perhaps it's not the platelet: Ristocetin uncovers the potential role of von Willebrand factor in impaired platelet aggregation following traumatic brain injury. J Trauma Acute Care Surg. 2018;85:873–80. [PubMed: 29985231]
- 26. Vogel S, Bodenstein R, Chen Q, Feil S, Feil R, Rheinlaender J, Schaffer TE, Bohn E, Frick JS, Borst O, Munzer P, Walker B, Markel J, Csanyi G, Pagano PJ, Loughran P, Jessup ME, Watkins SC, Bullock GC, Sperry JL, et al. Platelet-derived HMGB1 is a critical mediator of thrombosis. J Clin Invest. 2015;125:4638–54. [PubMed: 26551681]
- 27. Gonzalez Rodriguez E, Ostrowski SR, Cardenas JC, Baer LA, Tomasek JS, Henriksen HH, Stensballe J, Cotton BA, Holcomb JB, Johansson PI, Wade CE. Syndecan-1: A Quantitative Marker for the Endotheliopathy of Trauma. J Am Coll Surg. 2017;225:419–27. [PubMed: 28579548]
- 28. Ganter MT, Cohen MJ, Brohi K, Chesebro BB, Staudenmayer KL, Rahn P, Christiaans SC, Bir ND, Pittet JF. Angiopoietin-2, marker and mediator of endothelial activation with prognostic significance early after trauma? Ann Surg. 2008;247:320–6. [PubMed: 18216540]
- 29. Burk AM, Martin M, Flierl MA, Rittirsch D, Helm M, Lampl L, Bruckner U, Stahl GL, Blom AM, Perl M, Gebhard F, Huber-Lang M. Early complementopathy after multiple injuries in humans. Shock. 2012;37:348–54. [PubMed: 22258234]
- 30. Cohen MJ, Carles M, Brohi K, Calfee CS, Rahn P, Call MS, Chesebro BB, West MA, Pittet JF. Early release of soluble receptor for advanced glycation endproducts after severe trauma in humans. J Trauma. 2010;68:1273–8. [PubMed: 20539169]
- 31. Roberts DJ, Kalkwarf KJ, Moore HB, Cohen MJ, Fox EE, Wade CE, Cotton BA. Time course and outcomes associated with transient versus persistent fibrinolytic phenotypes after injury: A nested, prospective, multicenter cohort study. J Trauma Acute Care Surg. 2019;86:206–13. [PubMed: 30376538]
- 32. Davenport RA, Guerreiro M, Frith D, Rourke C, Platton S, Cohen M, Pearse R, Thiemermann C, Brohi K. Activated Protein C Drives the Hyperfibrinolysis of Acute Traumatic Coagulopathy. Anesthesiology. 2017;126:115–27. [PubMed: 27841821]
- 33. Howard BM, Kornblith LZ, Cheung CK, Kutcher ME, Miyazawa BY, Vilardi RF, Cohen MJ. Inducing Acute Traumatic Coagulopathy In Vitro: The Effects of Activated Protein C on Healthy Human Whole Blood. PLoS One. 2016;11:e0150930. [PubMed: 27008408]
- 34. Sumislawski JJ, Kornblith LZ, Conroy AS, Callcut RA, Cohen MJ. Dynamic coagulability after injury: Is delaying venous thromboembolism chemoprophylaxis worth the wait? J Trauma Acute Care Surg. 2018;85:907–14. [PubMed: 30124623]
- 35. Rizoli SB, Scarpelini S, Callum J, Nascimento B, Mann KG, Pinto R, Jansen J, Tien HC. Clotting factor deficiency in early trauma-associated coagulopathy. J Trauma. 2011;71:S427–34. [PubMed: 22071999]
- 36. Dzik WH. The James Blundell Award Lecture 2006: transfusion and the treatment of haemorrhage: past, present and future. Transfus Med. 2007;17:367–74. [PubMed: 17903138]
- 37. Park MS, Xue A, Spears GM, Halling TM, Ferrara MJ, Kuntz MM, Dhillon SK, Jenkins DH, Harmsen WS, Ballman KV, Harrison P, Heit JA. Thrombin generation and procoagulant microparticle profiles after acute trauma: A prospective cohort study. J Trauma Acute Care Surg. 2015;79:726–31. [PubMed: 26496097]
- 38. Cardenas JC, Rahbar E, Pommerening MJ, Baer LA, Matijevic N, Cotton BA, Holcomb JB, Wade CE. Measuring thrombin generation as a tool for predicting hemostatic potential and transfusion requirements following trauma. J Trauma Acute Care Surg. 2014;77:839–45. [PubMed: 25099452]
- 39. Shaz BH, Winkler AM, James AB, Hillyer CD, MacLeod JB. Pathophysiology of early traumainduced coagulopathy: emerging evidence for hemodilution and coagulation factor depletion. J Trauma. 2011;70:1401–7. [PubMed: 21460741]
- 40. Gissel M, Brummel-Ziedins KE, Butenas S, Pusateri AE, Mann KG, Orfeo T. Effects of an acidic environment on coagulation dynamics. J Thromb Haemost. 2016;14:2001–10. [PubMed: 27431334]

- 41. Kornblith LZ, Kutcher ME, Redick BJ, Calfee CS, Vilardi RF, Cohen MJ. Fibrinogen and platelet contributions to clot formation: implications for trauma resuscitation and thromboprophylaxis. J Trauma Acute Care Surg. 2014;76:255–6; discussion 62–3. [PubMed: 24458031]
- 42. Rourke C, Curry N, Khan S, Taylor R, Raza I, Davenport R, Stanworth S, Brohi K. Fibrinogen levels during trauma hemorrhage, response to replacement therapy, and association with patient outcomes. J Thromb Haemost. 2012;10:1342–51. [PubMed: 22519961]
- 43. Martini WZ, Holcomb JB. Acidosis and coagulopathy: the differential effects on fibrinogen synthesis and breakdown in pigs. Ann Surg. 2007;246:831–5. [PubMed: 17968176]
- 44. Schochl H, Nienaber U, Hofer G, Voelckel W, Jambor C, Scharbert G, Kozek-Langenecker S, Solomon C. Goal-directed coagulation management of major trauma patients using thromboelastometry (ROTEM)-guided administration of fibrinogen concentrate and prothrombin complex concentrate. Crit Care. 2010;14:R55. [PubMed: 20374650]
- 45. Rossaint R, Bouillon B, Cerny V, Coats TJ, Duranteau J, Fernandez-Mondejar E, Hunt BJ, Komadina R, Nardi G, Neugebauer E, Ozier Y, Riddez L, Schultz A, Stahel PF, Vincent JL, Spahn DR, Task Force for Advanced Bleeding Care in T. Management of bleeding following major trauma: an updated European guideline. Crit Care. 2010;14:R52. [PubMed: 20370902]
- 46. Kutcher ME, Redick BJ, McCreery RC, Crane IM, Greenberg MD, Cachola LM, Nelson MF, Cohen MJ. Characterization of platelet dysfunction after trauma. J Trauma Acute Care Surg. 2012;73:13–9. [PubMed: 22743367]
- 47. Brown LM, Call MS, Margaret Knudson M, Cohen MJ, Trauma Outcomes G, Holcomb JB, Wade CE, Brasel KJ, Vercruysse G, MacLeod J, Dutton RP, Hess JR, Duchesne JC, McSwain NE, Muskat P, Johannigamn J, Cryer HM, Tillou A, Pittet JF, De Moya MA, et al. A normal platelet count may not be enough: the impact of admission platelet count on mortality and transfusion in severely injured trauma patients. J Trauma. 2011;71:S337–42. [PubMed: 21814101]
- 48. Li R, Elmongy H, Sims C, Diamond SL. Ex vivo recapitulation of trauma-induced coagulopathy and preliminary assessment of trauma patient platelet function under flow using microfluidic technology. J Trauma Acute Care Surg. 2016;80:440–9. [PubMed: 27082706]
- 49. Sillesen M, Johansson PI, Rasmussen LS, Jin G, Jepsen CH, Imam AM, Hwabejire J, Lu J, Duggan M, Velmahos G, deMoya M, Alam HB. Platelet activation and dysfunction in a large-animal model of traumatic brain injury and hemorrhage. J Trauma Acute Care Surg. 2013;74:1252–9. [PubMed: 23609275]
- 50. Martin G, Shah D, Elson N, Boudreau R, Hanseman D, Pritts TA, Makley AT, Foreman B, Goodman MD. Relationship of Coagulopathy and Platelet Dysfunction to Transfusion Needs After Traumatic Brain Injury. Neurocrit Care. 2018.
- 51. Baimukanova G, Miyazawa B, Potter DR, Muench MO, Bruhn R, Gibb SL, Spinella PC, Cap AP, Cohen MJ, Pati S. Platelets regulate vascular endothelial stability: assessing the storage lesion and donor variability of apheresis platelets. Transfusion. 2016;56 Suppl 1:S65–75. [PubMed: 27001364]
- 52. Johansson PI, Henriksen HH, Stensballe J, Gybel-Brask M, Cardenas JC, Baer LA, Cotton BA, Holcomb JB, Wade CE, Ostrowski SR. Traumatic Endotheliopathy: A Prospective Observational Study of 424 Severely Injured Patients. Ann Surg. 2017;265:597–603. [PubMed: 27144442]
- 53. Brouns SLN, van Geffen JP, Heemskerk JWM. High-throughput measurement of human platelet aggregation under flow: application in hemostasis and beyond. Platelets. 2018:1–8.
- 54. Kornblith LZ, Robles AJ, Conroy AS, Hendrickson CM, Calfee CS, Fields AT, Callcut RA, Cohen MJ. Perhaps It's Not the Platelet: Ristocetin Uncovers the Potential Role of von Willebrand Factor in Impaired Platelet Aggregation Following Traumatic Brain Injury. J Trauma Acute Care Surg. 2018.
- 55. Mazepa M, Hoffman M, Monroe D. Superactivated platelets: thrombus regulators, thrombin generators, and potential clinical targets. Arteriosclerosis, thrombosis, and vascular biology. 2013;33:1747–52.
- 56. Tweardy DJ, Khoshnevis MR, Yu B, Mastrangelo MA, Hardison EG, Lopez JA. Essential role for platelets in organ injury and inflammation in resuscitated hemorrhagic shock. Shock. 2006;26:386–90. [PubMed: 16980886]

- 57. Ding N, Chen G, Hoffman R, Loughran PA, Sodhi CP, Hackam DJ, Billiar TR, Neal MD. Toll-like receptor 4 regulates platelet function and contributes to coagulation abnormality and organ injury in hemorrhagic shock and resuscitation. Circ Cardiovasc Genet. 2014;7:615–24. [PubMed: 25049041]
- 58. Stettler GR, Moore EE, Moore HB, Nunns GR, Huebner BR, Einersen P, Ghasabyan A, Silliman CC, Banerjee A, Sauaia A. Platelet adenosine diphosphate receptor inhibition provides no advantage in predicting need for platelet transfusion or massive transfusion. Surgery. 2017;162:1286–94. [PubMed: 28964508]
- 59. Henriksen HH, Grand AG, Viggers S, Baer LA, Solbeck S, Cotton BA, Matijevic N, Ostrowski SR, Stensballe J, Fox EE, Chen TA, Holcomb JB, Johansson PI, Cardenas JC, Wade CE. Impact of blood products on platelet function in patients with traumatic injuries: a translational study. J Surg Res. 2017;214:154–61. [PubMed: 28624038]
- 60. Holzmacher JL, Reynolds C, Patel M, Maluso P, Holland S, Gamsky N, Moore H, Acquista E, Carrick M, Amdur R, Hancock H, Metzler M, Dunn J, Sarani B. Platelet transfusion does not improve outcomes in patients with brain injury on antiplatelet therapy. Brain Inj. 2018;32:325–30. [PubMed: 29341793]
- 61. Shukla M, Sekhon UD, Betapudi V, Li W, Hickman DA, Pawlowski CL, Dyer MR, Neal MD, McCrae KR, Sen Gupta A. In vitro characterization of SynthoPlate (synthetic platelet) technology and its in vivo evaluation in severely thrombocytopenic mice. J Thromb Haemost. 2017;15:375– 87. [PubMed: 27925685]
- 62. Innes D, Sevitt S. Coagulation and Fibrinolysis in Injured Patients. Journal of clinical pathology. 1964;17:1–13. [PubMed: 14103515]
- 63. Cotton BA, Harvin JA, Kostousouv V, Minei KM, Radwan ZA, Schochl H, Wade CE, Holcomb JB, Matijevic N. Hyperfibrinolysis at admission is an uncommon but highly lethal event associated with shock and prehospital fluid administration. J Trauma Acute Care Surg. 2012;73:365–70; discussion 70. [PubMed: 22846941]
- 64. Hardaway RM, Chun B, Rutherford RB. Histologic evidence of disseminated intravascular coagulation in clinical shock. Vascular diseases. 1965;2:254–65. [PubMed: 4160038]
- 65. Hardaway RM, Drake DC. Prevention of "irreversible" hemorrhagic shock with fibrinolysin. Annals of surgery. 1963;157:39–47. [PubMed: 13952730]
- 66. Moore HB, Moore EE, Gonzalez E, Chapman MP, Chin TL, Silliman CC, Banerjee A, Sauaia A. Hyperfibrinolysis, physiologic fibrinolysis, and fibrinolysis shutdown: the spectrum of postinjury fibrinolysis and relevance to antifibrinolytic therapy. The journal of trauma and acute care surgery. 2014;77:811–7; discussion 7. [PubMed: 25051384]
- 67. Cardenas JC, Wade CE, Cotton BA, George MJ, Holcomb JB, Schreiber MA, White NJ, Group PS. Teg Lysis Shutdown Represents Coagulopathy in Bleeding Trauma Patients: Analysis of the Proppr Cohort. Shock. 2018.
- 68. Gomez-Builes JC, Acuna SA, Nascimento B, Madotto F, Rizoli SB. Harmful or Physiologic: Diagnosing Fibrinolysis Shutdown in a Trauma Cohort With Rotational Thromboelastometry. Anesthesia and analgesia. 2018.
- 69. Moore HB, Moore EE, Huebner BR, Dzieciatkowska M, Stettler GR, Nunns GR, Lawson PJ, Ghasabyan A, Chandler J, Banerjee A, Silliman C, Sauaia A, Hansen KC. Fibrinolysis shutdown is associated with a fivefold increase in mortality in trauma patients lacking hypersensitivity to tissue plasminogen activator. J Trauma Acute Care Surg. 2017;83:1014–22. [PubMed: 29190254]
- 70. Gonzalez E, Moore EE, Moore HB, Chapman MP, Chin TL, Ghasabyan A, Wohlauer MV, Barnett CC, Bensard DD, Biffl WL, Burlew CC, Johnson JL, Pieracci FM, Jurkovich GJ, Banerjee A, Silliman CC, Sauaia A. Goal-directed Hemostatic Resuscitation of Trauma-induced Coagulopathy: A Pragmatic Randomized Clinical Trial Comparing a Viscoelastic Assay to Conventional Coagulation Assays. Annals of surgery. 2016;263:1051–9. [PubMed: 26720428]
- 71. Moore HB, Moore EE, Huebner BR, Stettler GR, Nunns GR, Einersen PM, Silliman CC, Sauaia A. Tranexamic acid is associated with increased mortality in patients with physiological fibrinolysis. The Journal of surgical research. 2017.
- 72. Meizoso JP, Karcutskie CA, Ray JJ, Namias N, Schulman CI, Proctor KG. Persistent Fibrinolysis Shutdown Is Associated with Increased Mortality in Severely Injured Trauma Patients. Journal of the American College of Surgeons. 2017;224:575–82. [PubMed: 28017804]

- 73. Moore HB ME, Gonzalez E, Huebner BJ, Sheppard F, Banerjee A, Sauaia A, Silliman CC. Reperfusion Shutdown: Delayed Onset of Fibrinolysis Resistance after Resuscitation from Hemorrhagic Shock Is Associated with Increased Circulating Levels of Plasminogen Activator Inhibitor-1 and Postinjury Complications. Blood. 2016;128:206.
- 74. Myers SP, Kutcher ME, Rosengart MR, Sperry JL, Peitzman AB, Brown JB, Neal MD. Tranexamic acid administration is associated with an increased risk of posttraumatic venous thromboembolism. J Trauma Acute Care Surg. 2019;86:20–7. [PubMed: 30239375]
- 75. Naumann DN, Hazeldine J, Davies DJ, Bishop J, Midwinter MJ, Belli A, Harrison P, Lord JM. Endotheliopathy of Trauma is an On-Scene Phenomenon, and is Associated with Multiple Organ Dysfunction Syndrome: A Prospective Observational Study. Shock. 2017.
- 76. Ostrowski SR, Henriksen HH, Stensballe J, Gybel-Brask M, Cardenas JC, Baer LA, Cotton BA, Holcomb JB, Wade CE, Johansson PI. Sympathoadrenal activation and endotheliopathy are drivers of hypocoagulability and hyperfibrinolysis in trauma: A prospective observational study of 404 severely injured patients. J Trauma Acute Care Surg. 2017;82:293–301. [PubMed: 27779595]
- 77. Pati S, Potter DR, Baimukanova G, Farrel DH, Holcomb JB, Schreiber MA. Modulating the endotheliopathy of trauma: Factor concentrate versus fresh frozen plasma. J Trauma Acute Care Surg. 2016;80:576–84; discussion 84–5. [PubMed: 26808040]
- 78. Rodriguez EG, Cardenas JC, Lopez E, Cotton BA, Tomasek JS, Ostrowski SR, Baer LA, Stensballe J, Holcomb JB, Johansson PI, Wade CE. Early Identification of the Patient with Endotheliopathy of Trauma by Arrival Serum Albumin. Shock. 2017.
- 79. Kozar RA, Pati S. Syndecan-1 restitution by plasma after hemorrhagic shock. J Trauma Acute Care Surg. 2015;78:S83–6. [PubMed: 26002270]
- 80. Wu F, Peng Z, Park PW, Kozar RA. Loss of Syndecan-1 Abrogates the Pulmonary Protective Phenotype Induced by Plasma After Hemorrhagic Shock. Shock. 2017;48:340–5. [PubMed: 28107214]
- 81. Johansson PI, Stensballe J, Rasmussen LS, Ostrowski SR. High circulating adrenaline levels at admission predict increased mortality after trauma. J Trauma Acute Care Surg. 2012;72:428–36. [PubMed: 22439205]
- 82. Ostrowski SR, Johansson PI. Endothelial glycocalyx degradation induces endogenous heparinization in patients with severe injury and early traumatic coagulopathy. J Trauma Acute Care Surg. 2012;73:60–6. [PubMed: 22743373]
- 83. Pati S, Matijevic N, Doursout MF, Ko T, Cao Y, Deng X, Kozar RA, Hartwell E, Conyers J, Holcomb JB. Protective effects of fresh frozen plasma on vascular endothelial permeability, coagulation, and resuscitation after hemorrhagic shock are time dependent and diminish between days 0 and 5 after thaw. J Trauma. 2010;69 Suppl 1:S55–63. [PubMed: 20622621]
- 84. Moore HB, Moore EE, Chapman MP, McVaney K, Bryskiewicz G, Blechar R, Chin T, Burlew CC, Pieracci F, West FB, Fleming CD, Ghasabyan A, Chandler J, Silliman CC, Banerjee A, Sauaia A. Plasma-first resuscitation to treat haemorrhagic shock during emergency ground transportation in an urban area: a randomised trial. Lancet. 2018;392:283–91. [PubMed: 30032977]
- 85. Sperry JL, Guyette FX, Brown JB, Yazer MH, Triulzi DJ, Early-Young BJ, Adams PW, Daley BJ, Miller RS, Harbrecht BG, Claridge JA, Phelan HA, Witham WR, Putnam AT, Duane TM, Alarcon LH, Callaway CW, Zuckerbraun BS, Neal MD, Rosengart MR, et al. Prehospital Plasma during Air Medical Transport in Trauma Patients at Risk for Hemorrhagic Shock. N Engl J Med. 2018;379:315–26. [PubMed: 30044935]
- 86. Kutcher ME, Xu J, Vilardi RF, Ho C, Esmon CT, Cohen MJ. Extracellular histone release in response to traumatic injury: implications for a compensatory role of activated protein C. J Trauma Acute Care Surg. 2012;73:1389–94. [PubMed: 23188230]
- 87. Boueiz A, Hassoun PM. Regulation of endothelial barrier function by reactive oxygen and nitrogen species. Microvasc Res. 2009;77:26–34. [PubMed: 19041330]
- 88. Ganter MT, Brohi K, Cohen MJ, Shaffer LA, Walsh MC, Stahl GL, Pittet JF. Role of the alternative pathway in the early complement activation following major trauma. Shock. 2007;28:29–34. [PubMed: 17510601]

- 89. Cohen MJ, Brohi K, Calfee CS, Rahn P, Chesebro BB, Christiaans SC, Carles M, Howard M, Pittet JF. Early release of high mobility group box nuclear protein 1 after severe trauma in humans: role of injury severity and tissue hypoperfusion. Crit Care. 2009;13:R174. [PubMed: 19887013]
- 90. Deguchi H, Sinha RK, Marchese P, Ruggeri ZM, Zilberman-Rudenko J, McCarty OJT, Cohen MJ, Griffin JH. Prothrombotic skeletal muscle myosin directly enhances prothrombin activation by binding factors Xa and Va. Blood. 2016;128:1870–8. [PubMed: 27421960]
- 91. Wiener G, Moore HB, Moore EE, Gonzalez E, Diamond S, Zhu S, D'Alessandro A, Banerjee A. Shock releases bile acid inducing platelet inhibition and fibrinolysis. J Surg Res. 2015;195:390–5. [PubMed: 25777826]
- 92. Whelihan MF, Mann KG. The role of the red cell membrane in thrombin generation. Thromb Res. 2013;131:377–82. [PubMed: 23402970]
- 93. Billiar TR, Vodovotz Y. Time for trauma immunology. PLoS Med. 2017;14:e1002342. [PubMed: 28700602]
- 94. MacLeod JB, Lynn M, McKenney MG, Cohn SM, Murtha M. Early coagulopathy predicts mortality in trauma. J Trauma. 2003;55:39–44. [PubMed: 12855879]
- 95. Prat NJ, Meyer AD, Ingalls NK, Trichereau J, DuBose JJ, Cap AP. Rotational thromboelastometry significantly optimizes transfusion practices for damage control resuscitation in combat casualties. J Trauma Acute Care Surg. 2017;83:373–80. [PubMed: 28846577]
- 96. Christie SA, Kornblith LZ, Howard BM, Conroy AS, Kunitake RC, Nelson MF, Hendrickson CM, Calfee CS, Callcut RA, Cohen MJ. Characterization of distinct coagulopathic phenotypes in injury: Pathway-specific drivers and implications for individualized treatment. J Trauma Acute Care Surg. 2017;82:1055–62. [PubMed: 28338598]
- 97. Rong G, Corrie SR, Clark HA. In Vivo Biosensing: Progress and Perspectives. ACS Sens. 2017;2:327–38. [PubMed: 28723197]
- 98. Menezes AA, Vilardi RF, Arkin AP, Cohen MJ. Targeted clinical control of trauma patient coagulation through a thrombin dynamics model. Sci Transl Med. 2017;9.
- 99. Moore SE, Decker A, Hubbard A, Callcut RA, Fox EE, Del Junco DJ, Holcomb JB, Rahbar MH, Wade CE, Schreiber MA, Alarcon LH, Brasel KJ, Bulger EM, Cotton BA, Muskat P, Myers JG, Phelan HA, Cohen MJ, Group PS. Statistical Machines for Trauma Hospital Outcomes Research: Application to the PRospective, Observational, Multi-Center Major Trauma Transfusion (PROMMTT) Study. PLoS One. 2015;10:e0136438. [PubMed: 26296088]
- 100. Hubbard A, Munoz ID, Decker A, Holcomb JB, Schreiber MA, Bulger EM, Brasel KJ, Fox EE, del Junco DJ, Wade CE, Rahbar MH, Cotton BA, Phelan HA, Myers JG, Alarcon LH, Muskat P, Cohen MJ, Group PS. Time-dependent prediction and evaluation of variable importance using superlearning in high-dimensional clinical data. J Trauma Acute Care Surg. 2013;75:S53–60. [PubMed: 23778512]

Table 1.

Estimated Average Coagulation Parameters in Trauma Patients With Trauma-Induced Coagulopathy Early (0h) and Later (beyond 0h) After Injury vs. Without Trauma-Induced Coagulopathy

* Estimated average quantitative values (mean or median) for each listed coagulation parameter in trauma patients with trauma-induced coagulopathy (TIC) at early and later timepoints after injury vs. without TIC. Quantitative values are supported by existing literature, as cited. Abbreviations: trauma induced coagulopathy (TIC), adenosine diphosphate (ADP), thrombin receptor activating peptide-6 (TRAP), thromboelastography (TEG), activated coagulation time (ACT), percent lysis at 30 minutes (LY30), maximum amplitude (MA), high mobility group box 1 protein (HMGB-1), soluble (s), receptor for advanced glycation endproducts (RAGE), under-defined in previous studies (und).

Table 2.

Qualitative Changes in Coagulation Parameters in Trauma-Induced Coagulopathy Compared to Major and Specific Criteria for Disseminated Intravascular Coagulation from the Scientific Subcommittee on Disseminated Intravascular Coagulation of the International Society on Thrombosis and Haemostasis (21)

* Qualitative changes for each listed coagulation parameter in trauma patients with TIC vs. major (**bolded**) and specific (grey) criteria for Disseminated Intravascular Coagulation (DIC) by the Scientific Subcommittee on DIC of the International Society on Thrombosis and Haemostasis. Abbreviations: trauma-induced coagulopathy (TIC), disseminated intravascular coagulation (DIC), adenosine diphosphate (ADP), thrombin receptor activating peptide-6 (TRAP), thromboelastography (TEG), activated coagulation time (ACT), percent lysis at 30 minutes (LY30), maximum amplitude (MA), high mobility group box 1 protein (HMGB-1), soluble (s), receptor for advanced glycation endproducts (RAGE), not included (n/i).