



Scorer, T. G., Fitzgibbon, L., Aungraheeta, R., Sharma, U., Peltier, G. C., McIntosh, C. S., Reddoch-Cardenas, K. M., Meyer, A., Cap, A. P., & Mumford, A. D. (2020). TEG PlateletMapping assay results may be misleading in the presence of cold stored platelets. *Transfusion*, *60*(S3), S119-S123. https://doi.org/10.1111/trf.15753

Peer reviewed version

Link to published version (if available): 10.1111/trf.15753

Link to publication record in Explore Bristol Research PDF-document

This is the author accepted manuscript (AAM). The final published version (version of record) is available online via Wiley at https://onlinelibrary.wiley.com/doi/full/10.1111/trf.15753. Please refer to any applicable terms of use of the publisher.

# University of Bristol - Explore Bristol Research General rights

This document is made available in accordance with publisher policies. Please cite only the published version using the reference above. Full terms of use are available: http://www.bristol.ac.uk/red/research-policy/pure/user-guides/ebr-terms/

## **Brief Report – Transfusion – THOR supplement 2019**

TEG PlateletMapping assay results may be misleading in the presence of cold stored platelets

TG Scorer<sup>1,2,3</sup>, L FitzGibbon<sup>2</sup>, R Aungraheeta<sup>2</sup>, U Sharma<sup>3</sup>, GC Peltier<sup>3</sup>, CS McIntosh<sup>3</sup>, KM Reddoch-Cardenas<sup>3</sup>, A Meyer<sup>3,4</sup>, AP Cap<sup>3</sup>, AD Mumford<sup>2</sup>

<sup>1</sup> Centre of Defence Pathology, Royal Centre of Defence Medicine, Birmingham, UK.

<sup>2</sup> School of Cellular and Molecular Medicine, University of Bristol, Bristol, UK.

<sup>3</sup>Coagulation and Blood research, U.S. Army Institute of Surgical Research, Fort Sam Houston, Texas, USA.

<sup>4</sup> Division of Pediatric Critical Care, Department of Pediatrics, University of Texas Health Science Center, San Antonio, TX

Word count: 1720 Abstract word count: 250 Number of figures and tables: 3

References: 24

**Corresponding author**: Dr Tom Scorer, Research Floor Level 7, University of Bristol, Bristol Royal Infirmary, Bristol, BS2 8HW United Kingdom.

Email: tom.scorer@bristol.ac.uk

The authors have no relevant conflicts of interest to declare.

### DISCLAIMER

TS was funded by Ministry of Defence. The study was supported by the U.S. Army Medical Research and Materiel Command, NIHR Biomedical Research Centre at University Hospitals Bristol NHS Foundation Trust and the University of Bristol. The opinions or assertions contained herein are the private views of the authors and are not to be construed as official or as reflecting the views of the U.S. Department of the Army, the U.S. Department of Defense, the Ministry of Defence, the National Institute for Health Research or the Department of Health.

#### ABSTRACT

**Background:** Viscoelastic tests (VETs) are used widely to monitor hemostasis in settings such as cardiac surgery. There has also been renewed interest in cold stored platelets (CSP) to manage bleeding in this setting. CSPs are reported to have altered hemostatic properties compared to room temperature platelets (RTP), including activation of GPIIb/IIIa. We investigated whether the functional differences between CSP and RTP affected the performance of the PlateletMapping VET on the TEG 5000 and 6s analyzer.

**Method:** Platelet concentrates were divided equally into CSP (stored at  $4^{\circ}C\pm 2^{\circ}C$ ) and RTP (stored at  $22^{\circ}C\pm 2^{\circ}C$ ) fractions. Whole blood was treated to induce Platelet dysfunction (WBIPD) by incubating with anti-platelet drugs (1.0µM ticagrelor and 10µM aspirin) or by simulating cardiopulmonary bypass. WBIPD samples were then mixed with 20% by volume of CSP or RTP to model platelet transfusion before analysis using the PlateletMapping VET.

**Results:** Addition of CSP to WBIPD increased the PlateletMapping MA<sub>FIBRIN</sub> and MA<sub>ADP</sub> parameters with the TEG 5000 analyzer (both p<0.0001 compared to addition of buffer alone). This effect was not observed with RTP. The differential effect of CSP on the MA<sub>FIBRIN</sub> corrected after pre-incubation with the GPIIb/IIIa antagonist tirofiban and was quantitatively less with the PlateletMapping test for the TEG 6s analyzer which contains the GPIIb/IIa antagonist abciximab.

**Discussion:** The PlateletMapping MA<sub>FIBRIN</sub> and MA<sub>ADP</sub> test results may be misleadingly high with CSP, particularly with the TEG 5000 analyzer, most likely due to constitutive activation of GPIIb/IIIa on CSP during storage. TEG PlateletMapping results should be interpreted with caution following CSP transfusion.

#### FIGURE LEGENDS

Figure 1: TEG PlateletMapping assay percentage (%) inhibition/aggregation is calculated from the maximal amplitude (MA) of three TEG assays;  $MA_{THROMBIN}$  obtained using the standard TEG Kaolin activator, which stimulates thrombin generation in the blood sample. This parameter reflects the thrombin-dependent contributions of fibrin plus platelets to clot strength.  $MA_{FIBRIN}$  obtained using ActivatorF, containing reptilase and FXIII to generate and stabilize fibrin clot independently of thrombin. This reflects the fibrin component.  $MA_{ADP/AA}$  in which ActivatorF is combined with the direct platelet activators adenosine di-phosphate (ADP) or arachidonic acid (AA). This reflects the contribution of platelet ADP or AA pathways to clot strength. The equation shown is used to calculate the '% aggregation' parameter from the three MA measurements and represents the contribution of either ADP-mediated or AAmediated platelet activation to clot strength. Percentage (%) inhibition = 100 - % aggregation

Figure 2: TEG 5000 PlateletMapping assay maximal amplitude (MA) and calculated percentage (%) aggregation of; whole blood with induced thrombocytopathy (WBIPD) and addition of 20% v/v buffer alone, WBIPD with a simulated transfusion of 20% v/v, cold stored platelets (WBIPD + CSP), and WBIPD with a simulated transfusion of 20% v/v, room temperature stored platelets (WBIPD + RTP) (n = 14). A – MA<sub>THROMBIN</sub>, B – MA<sub>FIBRIN</sub>, C – MA<sub>ADP</sub> and D – Calculated Percentage (%) aggregation. Statistical significance relative to WBIPD \* = p < 0.05, \*\* = p < 0.01, \*\*\* = p < 0.001, \*\*\*\* = p < 0.001, ns - non-significance.

Figure 3: TEG MA<sub>FIBRIN</sub> dose response curve illustrating tirofiban treatment of pooled buffy coat CSP (n = 3) with the TEG 5000 analyzer, R2 = 0.93, 95% confidence intervals, standard error of the mean (SEM).

#### INTRODUCTION

The use of viscoelastic hemostatic tests (VETs) to guide resuscitation during major hemorrhage has increased significantly in recent years and is now recommended by the National Institute for Health and Care Excellence (NICE) for cardiac surgery.<sup>1</sup> VETs are also used commonly in other clinical settings such as traumatic or obstetric hemorrhage, but are not formally recommended in practice guidelines because evidence supporting clinical benefit remains incomplete.<sup>2-5</sup>

In both of the two most widely adopted VET technologies thromboelastography (TEG; Haemonetics, Braintree, MA) and rotational thromboelastometry (ROTEM; TEM International, Munich, Germany), clot formation is initiated using specific activators in whole blood samples housed in analysis cups with a suspended pin. Clot viscoelastic strength is then monitored continuously by detecting the mechanical resistance to oscillations applied to either cup (TEG) or pin (ROTEM).

Early versions of these VETs using contact (TEG-Kaolin test) or extrinsic pathway (ROTEM-EXTEM test) activators did not distinguish the specific contributions of either fibrin or platelets to clot strength. However, subsequent refinements of VETs utilize selective coagulation activators and platelet inhibitors to more precisely phenotype clot formation.<sup>6,7</sup> One example is the TEG PlateletMapping system which enables the platelet contribution to clot strength to be inferred from three different measurements of the maximum amplitude (MA) of the test traces:<sup>8-10</sup> 1. MATHROMBIN obtained using the standard TEG Kaolin activator, which stimulates thrombin generation in the blood sample. This parameter reflects the thrombin-dependent contributions of fibrin plus platelets to clot strength. 2. MAFIBRIN obtained using ActivatorF, containing reptilase

and FXIII to generate and stabilize fibrin clot independently of thrombin. This reflects the fibrin component. 3. MAADP/AA in which ActivatorF is combined with the direct platelet activators adenosine di-phosphate (ADP) or arachidonic acid (AA). This reflects the contribution of platelet ADP or AA pathways to clot strength. The main '% inhibition/aggregation' parameter from TEG PlateletMapping system is derived from the three MA measurements (Figure 1) and represents the contribution of either ADPmediated or AA-mediated platelet activation to clot strength. This enables detection of the antiplatelet drugs aspirin (inhibits AA-mediated platelet function) and P2Y<sub>12</sub>blockers (inhibit ADP-mediated platelet function), which is critical in settings such as cardiac surgery to quantify the impact of anti-platelet drugs on clot strength and to guide platelet transfusion. Although initially developed for the TEG 5000 analyzer, the TEG PlateletMapping system has now been refined for the cartridge-based TEG 6s analyzer. For this analyzer, the ActivatorF reagent used to measure the MAFIBRIN also includes the potent GPIIb/IIIa receptor antagonist abciximab.<sup>10</sup> This prevents plateletfibrin interactions ensuring that the MAFIBRIN only reflects the fibrin component of clot strength, thereby enabling a more reliable calculation of platelet 'percentage inhibition/aggregation'.

One important trend in transfusion practice is the emergence of evidence that cold stored platelets (CSP) may be a useful alternative to conventional room temperature platelets (RTP) is settings such trauma or cardiac surgery.<sup>11-15</sup> It is established that CSP have altered expression of several surface adhesive receptors including GPIIb/IIIa, and also increased basal platelet activation compared to WSP.<sup>16-19</sup> Since these characteristics are critical for the endpoints of the TEG PlateletMapping test, we investigated whether CSP had a different effect on test results compared to RTP when

platelets were mixed with coagulopathic blood samples to simulate platelet transfusion.

### MATERIALS AND METHODS

#### **Preparation of platelet concentrates**

Apheresis platelets were collected from healthy volunteers using the Trima Accel system (TerumoBCT Lakewood, CO) as previously described.<sup>17,20-22</sup> Platelets were collected as hyper-concentrated double collections, before being split equally into two platelet storage bags (Polyolefin PL-2410, Fenwal, Lake Zurich, IL) within two hours. Pooled buffy coat platelets were manufactured by NHS Blood and Transplant (NHSBT, Bristol UK) as previously described<sup>23</sup> in compliance with Blood Safety and Quality Regulations, Statutory Instrument 2005 No.50.<sup>24</sup> Both preparations were suspended in 35% plasma and 65% platelet additive solution (T-PAS, TerumoBCT, Lakewood, CO for apheresis and SSP+, MacoPharma, Mouvaux, France for buffy-coat platelet concentrates were separated aseptically 24-36 hours after donation into two equal volumes into two TOTM bags (MacoPharma, Mouvaux, France) for storage at 22°C  $\pm 2$  °C with continuous agitation (RTP) or at 4°C  $\pm 2$  °C without agitation (CSP).

#### Preparation of Whole blood with induced platelet dysfunction (WBIPD) samples.

Whole blood (WB) was collected from healthy volunteers by peripheral venipuncture into 3.2% trisodium citrate tubes (BD Biosciences, Oxford, UK) in compliance with the Declaration of Helsinki. Two platelet dysfunction models were simulated and in this analysis are considered together (n = 14) 1.) *dual anti-platelet therapy:* incubation with 1.0µM ticagrelor and 10µM acetylsalicylic acid (ASA) for a minimum of 15 minutes (n

= 8 samples), and, 2) *extracorporeal bypass*; WB collected into a 500 mL Capiox reservoir (TerumoBCT, Lakewood, CO) containing 1 IU/mL heparin sulfate then circulated through an extracorporeal bypass circuit containing a roller pump (Cobe CV Model 43600, Sorin Biomedica, Milan, Italy), blood oxygenator (KIDS D100 Oxygenator, Sorin Biomedica, Milan, Italy) and pressure monitoring catheters (DLP catheters, 8 Fr, 12 Fr, Medtronic, Minneapolis, MN). Blood was circulated at a flow rate of 0.5 L/min with samples taken for analysis after 6 hours (n = 6 samples).

## Mixing experiments and PlateletMapping analysis

Apheresis CSP or RTP stored between three and seven days were mixed with WBIPD samples at a dose of 20% by volume. In control experiments, the same volume of PlasmaLyte A pH 7.4 (Baxter, Deerfield, IL) was mixed with the WBIPD samples. The sample mixtures were analyzed using the TEG 5000 analyzer according to manufacturer's guidelines.<sup>10</sup> In additional experiments to study the effect of additional GPIIb/IIIa inhibition, buffy coat CSP were incubated with tirofiban (Correvio Pharma Corp. Vancouver, Canada) for at least 15 minutes and analyzed on both TEG 5000 and 6s platforms. Data were compared by analysis of variance (ANOVA) performed using GraphPad Prism version 8.2.1 (GraphPad Software, San Diego, CA).

#### RESULTS

Addition of both CSP and RTP increased the TEG 5000 MA<sub>THROMBIN</sub> in the WBIPD samples (both p<0.001 compared to addition of buffer alone, n = 14, figure 2A), consistent with reversal of thrombocytopathy. As expected, the TEG 5000 MA<sub>FIBRIN</sub> was not altered by the addition of RTP to WBIPD. However, CSP substantially increased the MA<sub>FIBRIN</sub> (p<0.0001 compared to addition of buffer alone, n = 14, figure

2B). There was a small, but non-significant increase in TEG 5000 MA<sub>ADP</sub> with RTP, but a significant increase with CSP (p<0.0001 compared to addition of buffer alone, n = 14, figure 2C). The net effect of the differences in MA<sub>FIBRIN</sub> and MA<sub>ADP</sub> between RTP and CSP was an apparent reduction in the derived percentage aggregation parameter with CSP. However, because of the broad variation in results, this did not reach statistical significance (figure 2D).

One explanation for the increased MAFIBRIN with CSP was that increased constitutive activation of GPIIb/IIIa could potentially cause the CSP to contribute to clot strength even though platelets are not directly activated with the ActivatorF reagent that is used to measure the MAFIBRIN. In order to test this, the CSP were incubated with the GPIIb/IIIa antagonist tirofiban before mixing with WBIPD and testing using the TEG 5000 platform. The addition of tirofiban resulted in a dose-dependent decrease in MAFIBRIN with CSP, resulting in values similar to those with RTP at tirofiban concentrations above 1mM (figure 3). In order to assess whether the inclusion of abciximab in the MAFIBRIN channel of the TEG 6s PlateletMapping cartridge reduces the CSP MAFIBRIN in the same way as tirofiban, three samples of CSP products were analyzed on both TEG 5000 and 6s analyzers simultaneously. The MAFIBRIN was 56.2  $\pm$  3.28 mm (mean $\pm$  SEM) and 45.2  $\pm$  3.0 mm for TEG 5000 and TEG 6s respectively a reduction of 11 mm (19.6%) with TEG 6s.

Together, these observations support that the increased MA<sub>FIBRIN</sub> with CSP compared to RTP was mostly mediated by activated GPIIb/IIIa on CSP and that this particularly impacts on the unmodified TEG 5000 PlateletMapping test because a GPIIb/IIIa antagonist is absent.

#### DISCUSSION

The TEG PlateletMapping assay has proved valuable in the management of patients, especially cardiac surgery patients in which there is frequently a complex thrombocytopathy that includes the effects of anti-platelet and the mechanical effect of cardiopulmonary bypass.<sup>2,3,5</sup> The calculation of percentage inhibition/aggregation with ADP or AA is essential to resolve the degree of platelet dysfunction. However, this derived parameter relies on the MA<sub>FIBRIN</sub> being an accurate representation of the contribution of fibrin to clot strength which can then be subtracted from the MA<sub>THROMBIN</sub> and MA<sub>ADP/AA</sub> to yield the respective platelet contributions. Our results demonstrate that particularly with the TEG 5000 PlateletMapping test, this assumption may not be valid in the presence of CSP with which there is a dramatic increase in MA<sub>FIBRIN</sub> compared with equivalent quantities of RTP. This differential effect was also observed with the MA<sub>ADP</sub> result, which together resulted in a trend towards apparent underestimation of the percentage aggregation parameter in the presence of CSP.

We show further that the differential effect of CSP on the TEG 5000 PlateletMapping MA<sub>FIBRIN</sub> can be ameliorated by the addition of the GPIIb/IIIa antagonist tirofiban. This observation is consistent with previous observations that cold storage of platelets results in constitutive activation of GPIIb/IIIa enabling binding of free fibrinogen in plasma, a process responsible for the formation of platelet clumps during cold storage in fibrinogen rich medium.<sup>19</sup> The bound fibrinogen may also complex for FXIII, providing a nucleation site for fibrin polymerisation.<sup>17</sup> We propose that addition of ActivatorF containing activated FXIII and reptilase for measurement of the PlateletMapping MA<sub>FIBRIN</sub> and MA<sub>ADP</sub>, results in crosslinked fibrin formation, but also recruitment of CSP to contribute to clot strength, even though they are not directly

activated by the test reagents. This effect is not observed with RTP because GPIIb/IIIa activation is minimal at room temperature storage. Consistent with this, the impact of CSP on MA<sub>FIBRIN</sub> is less in the PlateletMapping test developed for the TEG 6s analyzer, because of the inclusion of the GPIIb/IIIa antagonist abciximab, although still present. The partial reduction in MA<sub>FIBRIN</sub> in the TEG 6s with CSP may indicate that the dose of abciximab in the cartridge is insufficient. These findings indicate that the TEG 6s assay has the potential to be optimised further to mitigate against misleading results. In the meantime, our data suggest that caution should be exercised in interpreting TEG PlateletMapping results in the presence of CSP. Further validation, ideally in the *in vivo* setting is required to ensure patients are not put at risk.

#### ACKNOWLEDGMENTS

Thank you to Mr Chris Taswell (Bristol Royal Infirmary, cardiac theatres) for facilitating the use of TEG 5000 and 6s analyzers. Haemonetics for kindly supplying TEG PlateletMapping reagents for part of this study.

#### REFERENCES

- National Institute for Health and Care Excellence (NICE) Diagnostics Guidance 13: Detecting, managing and monitoring haemostasis: viscoelastometric point-of-care testing (ROTEM, TEG and Sonoclot systems) 2014. Available from: <u>https://www.nice.org.uk/guidance/dg13. Accessed 21st September 2019.</u>
- 2. Whiting P, Al M, Westwood M, et al. Viscoelastic point-of-care testing to assist with the diagnosis, management and monitoring of haemostasis: a systematic review and cost-effectiveness analysis. Health Technol Assess 2015;**19**: 1-228, v-vi.
- Serraino GF, Murphy GJ. Routine use of viscoelastic blood tests for diagnosis and treatment of coagulopathic bleeding in cardiac surgery: updated systematic review and meta-analysis. Br J Anaesth 2017;118: 823-33.
- 4. Hunt BJ, Allard S, Keeling D, et al. A practical guideline for the haematological management of major haemorrhage. Br J Haematol 2015;**170**: 788-803.
- Curry NS, Davenport R, Pavord S, et al. The use of viscoelastic haemostatic assays in the management of major bleeding: A British Society for Haematology Guideline. Br J Haematol 2018;182: 789-806.
- Peng HT, Nascimento B, Beckett A. Thromboelastography and Thromboelastometry in Assessment of Fibrinogen Deficiency and Prediction for Transfusion Requirement: A Descriptive Review. Biomed Res Int 2018;2018: 7020539.
- Stettler GR, Sumislawski JJ, Moore EE, et al. Citrated kaolin thrombelastography (TEG) thresholds for goal-directed therapy in injured patients receiving massive transfusion. J Trauma Acute Care Surg 2018;85: 734-40.
- Sivapalan P, Back AC, Ostrowski SR, et al. Transfusion requirements in elective cardiopulmonary bypass surgery patients: predictive value of Multiplate and Thromboelastography (TEG) Platelet Mapping Assay. Scand J Clin Lab Invest 2017;77: 345-51.

- 9. Bochsen L, Wiinberg B, Kjelgaard-Hansen M, et al. Evaluation of the TEG platelet mapping assay in blood donors. Thromb J 2007;**5**: 3.
- *TEG system.* Haemonetics. Available from: <u>https://teg.haemonetics.com/en. Accessed</u>
  21<sup>st</sup> September 2019.
- Reddoch-Cardenas KM, Bynum JA, Meledeo MA, et al. Cold-stored platelets: A product with function optimized for hemorrhage control. Transfus Apher Sci 2019;58: 16-22.
- 12. Berzuini A, Spreafico M, Prati D. One size doesn't fit all: Should we reconsider the introduction of cold-stored platelets in blood bank inventories? F1000Research 2017;6: 95.
- Apelseth TO, Cap AP, Spinella PC, et al. Cold stored platelets in treatment of bleeding.
  ISBT Science Series 2017;12: 488-95.
- Cap AP. Targeting hemorrhage: Alternative storage of platelets for hemostatic transfusion. Blood 2017;130.
- 15. Milford CEM, Reade CMC. Comprehensive review of platelet storage methods for use in the treatment of active hemorrhage. Transfusion 2016;**56**: S140-S8.
- Reddoch KM, Pidcoke HF, Montgomery RK, et al. Hemostatic Function of Apheresis
  Platelets Stored at 4°C and 22°C. Shock (Augusta, Ga.) 2014;41: 54-61.
- Nair PM, Pandya SG, Dallo SF, et al. Platelets stored at 4°C contribute to superior clot properties compared to current standard-of-care through fibrin-crosslinking. Br J Haematol 2017;**178**: 119-29.
- Wood B, Padula MP, Marks DC, et al. Refrigerated storage of platelets initiates changes in platelet surface marker expression and localization of intracellular proteins. Transfusion 2016;56: 2548-59.
- Getz TM, Montgomery RK, Bynum JA, et al. Storage of platelets at 4degreeC in platelet additive solutions prevents aggregate formation and preserves platelet functional responses. Transfusion 2016;56: 1320-8.

- 20. Montgomery RK, Reddoch KM, Evani SJ, et al. Enhanced shear-induced platelet aggregation due to low-temperature storage. Transfusion 2013;**53**: 1520-30.
- Reddoch KM, Montgomery RK, Rodriguez AC, et al. Endothelium-derived inhibitors efficiently attenuate the aggregation and adhesion responses of refrigerated platelets. Shock 2016;45: 220-7.
- Scorer T, Sharma U, Peltier G, et al. Ticagrelor Induced Platelet Dysfunction Can be Assessed Under Shear Conditions and Correction By Platelets Is Influenced By Storage Temperature. Blood 2018;132.
- JPAC. Guidelines for the Blood Transfusion Services in the UK: 8th Edition Red Book.
  2013. Available from: <u>https://www.transfusionguidelines.org/red-book. Accessed 21st</u>
  <u>September 2019.</u>
- 24. Blood Safety and Quality Regulations 2005 UK Statutory Instrument No.50 2005.