

Single step separation of plasma from whole blood without the need for centrifugation applied to the quantitative analysis of warfarin

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1. Introduction

Dried plasma sample collection and storage from whole blood without the need for centrifugation separation and refrigeration opens new opportunities in blood sampling strategies for quantitative LC/MS/MS bioanalysis. Plasma samples were generated by gravity filtration of a whole blood sample through a laminated membrane stack allowing plasma to be collected, dried, transported and analysed by LC/MS/MS. This novel plasma separation card (PSC) technology was applied to the quantitative LC/MS/MS analysis of warfarin, in blood samples. Warfarin is a coumarin anticoagulant vitamin-K antagonist used for the treatment of thrombosis and thromboembolism. As a result of vitamin-K recycling being inhibited, hepatic synthesis is in-turn inhibited for blood clotting factors as well as anticoagulant proteins. Whilst the measurement of warfarin activity in patients is normally measured by prothrombin time by international normalized ratio (INR) in some cases the quantitation of plasma warfarin concentration is needed to confirm patient compliance, resistance to the anticoagulant drug, or diet related issues. In this preliminary evaluation, warfarin concentration was measured by LC/MS/MS to evaluate if PSC technology could complement INR when sampling patient blood.

2. Materials and Methods

2-1. Sample preparation

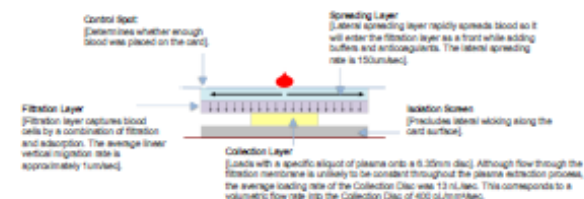
Warfarin standard was dissolved in water containing 50% ethanol + 0.1% formic acid, spiked (60uL) to whole human blood (1mL) and mixed gently. 50uL of spiked blood was deposited onto the PSC. After 3 minutes, the primary filtration overlay was removed followed by 15 minutes air drying at room temperature. The plasma sample disc was prepared directly for analysis after drying. LC/MS/MS sample preparation involved vortexing the sample disk in 40uL methanol, followed by centrifugation 16,000g 5 min. 20uL supernatant was added directly to the LC/MS/MS sample vial already containing 80uL water (2uL analysed). Control plasma comparison was prepared by centrifuging remaining blood at 1000g for 10min. 2.5uL supernatant plasma was taken, 40uL methanol added, and prepared as PSC samples. LC/MS/MS sample injection volume, 2uL.

2-2. LC-MS/MS analysis

Warfarin was measured by MRM, positive negative switching mode (15msec).

LC/MS/MS System	Nexera UHPLC system + LCMS-8040 Shimadzu Corporation
Flow rate	0.4mL/min (0-7.75min), 0.5mL/min (7.5-14min), 0.4mL/min (15min)
Mobile phase	A= Water + 0.1% formic acid B= Methanol + 0.1% formic acid
Gradient	20% B (0-0.5 min), 100% B (8-12 min), 20% B (12.01-15 min)
Analytical column	Phenomenex Kinetax XB C18 100 x 2.1mm 1.7um 100A
Column temperature	50°C
Ionisation	Electrospray, positive, negative switching mode
Dissolution line	250°C
Drying/Nebulising gas	10L/min, 2L/min
Heating block	400°C

Design of plasma separator technology



Plasma separation workflow



Figure 1. Noviplex workflow.

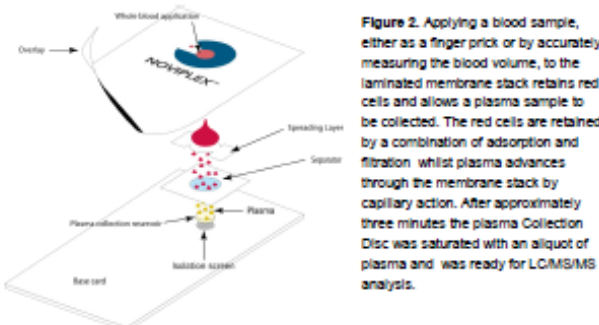


Figure 2. Applying a blood sample, either as a finger prick or by accurately measuring the blood volume, to the laminated membrane stack retains red cells and allows a plasma sample to be collected. The red cells are retained by a combination of adsorption and filtration whilst plasma advances through the membrane stack by capillary action. After approximately three minutes the plasma Collection Disc was saturated with an aliquot of plasma and was ready for LC/MS/MS analysis.

3 Results

Comparison between plasma separation cards (PSC) and plasma

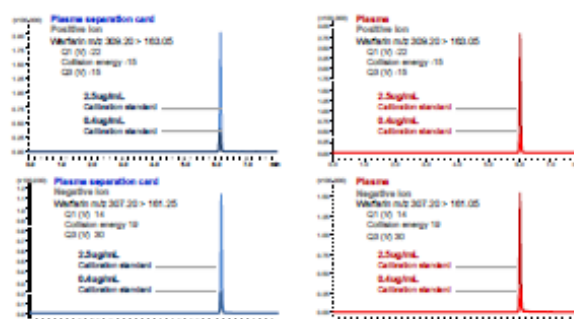


Figure 3. Comparison between the warfarin response in both positive and negative ion modes for warfarin calibration standards at 2.5ug/mL and 0.4ug/mL extracted from the plasma separation cards and a conventional plasma sample. There is a broad agreement in ion signal intensity between the 2 sample preparation techniques.

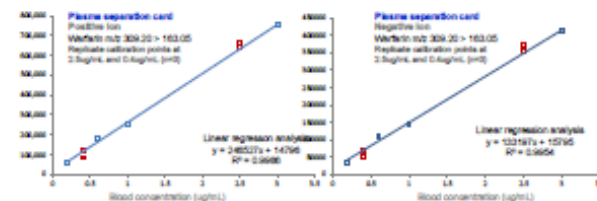


Figure 4. In both ion modes, the calibration curve was linear over the therapeutic range studied for warfarin extracted from PSC's (calibration range 0-3ug/mL, single point calibration standards at each level with the exception of replicate calibration points at 2.5ug/mL and 0.4ug/mL (n=3); $r^2 > 0.99$ for PSC analysis [$r^2 > 0.99$ for a conventional plasma extraction]).

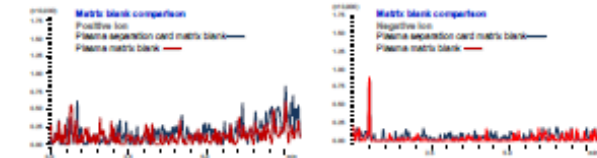


Figure 5. Matrix blank comparison. In both ion modes, the MRM chromatograms for PSC and plasma are comparable. Warfarin ion signals were not detected in the any PSC or plasma matrix blank.

3.1 Plasma separation card comparison

The drive to work with smaller sample volumes offers significant ethical and economical advantages in pharmaceutical and clinical workflows and dried blood spot sampling techniques have enabled a step change approach for many toxicokinetic and pharmacokinetic studies. However, the impressive growth of this technique in the quantitative analysis of small molecules has also discovered several limitations in the case of sample instability (some enzyme labile compounds, particularly prodrugs, analyte stability can be problematic), hematocrit effect and background interferences of DBS. DBS also shows noticeable effects on many lipids dependent on the sample collection process. To compare PSC to plasma lipid profiles the same blood sample extraction procedure applied for warfarin analysis was measured by a high mass accuracy system optimized for lipid profiling.

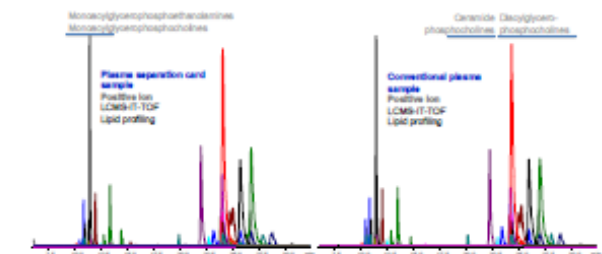


Figure 6. Lipid profiles from the same human blood sample extracted using a plasma separation card (left hand profile) compared to a conventional plasma samples (centrifugation). Both lipid profiles are comparable in terms of distribution and the number of lipids detected (the scaling has been normalized to the most intense lipid signal).

4. Conclusions.

- In this limited study, plasma separation card (PSC) sampling delivered a quantitative analysis of warfarin spiked into human blood.
- PSC generated a linear calibration curve in both positive and negative ion modes ($r^2 > 0.99$; $n=5$).
- The warfarin plasma results achieved by using the PSC technique were in broad agreement with conventional plasma sampling data.
- The plasma generated by the filtration process appears broadly similar to plasma derived from conventional centrifugation.
- Further work is required to consider the robustness and validation in a routine analysis.

5. References

- Jensen, B.P., Chin, P.K.L., Begg, E.J. (2011) Quantification of total and free concentrations of R- and S-warfarin in human plasma by ultrafiltration and LC-MS/MS. *Anal Bioanal Chem*, 401, 2187-2193
- Radiwan, M.A., Bawazeer, G.A., Aloudah, N.M., Alusdeib, B.T., Abou-Enein, H.Y. (2012) Determination of free and total warfarin concentrations in plasma using UPLC MS/MS and its application to patient samples. *Biochemical Chromatography*, 28, 6-11