Is the BD Barricor Tube the New Standard for Lithium Heparin Plasma?

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Abstract

Background: Lithium heparin plasma from gel separator tubes has one major advantage over serum it can be centrifuged without delay. The major disadvantage is it contains a large number of cells, cell stroma and fibrin which renders it unsuitable for delayed testing of some analytes e.g. potassium.

Methods: BD tubes, plasma separator tube - PST, Barricor tube, rapid serum tube - RST and serum separator tube-SST were evaluated for stability of potassium, glucose, LDH and phosphate at different storage time points and centrifugation settings from 22 participants. Additionally tubes from a participant were centrifuged at 3000g, and the buffy coat containing sample was then transferred into a Cytospin centrifuge to prepare slides for staining and content examination.

Results: The Barricor tube showed decreased changes in analyte concentrations compared with the PST tube. The stability of analytes was best in the RST and at 3000g across all tubes. The buffy coat study showed the PST tube had the largest number cells present.

Conclusion: The Barricor tube offers improvement in lithium heparin plasma quality due to reduced cell numbers leading to decreasing the change in potassium, LDH and phosphate results from cellular lysis and glucose results from cellular consumption upon sample storage. Based on the data from this study the Barricor tube should become the new standard for lithium heparin plasma.

Keywords: Barricor; Tubes; Plasma; Serum; Gel separator

Introduction

Biochemical analysis primarily uses lithium heparin plasma or serum specimens, with serum being the more commonly used specimen type. It is well documented the lithium heparin and serum tubes do not always produce the optimal quality sample to meet timeliness and accuracy requirements, and the deficiency of these tubes are well known [1-3]. Obtaining the highest quality sample is in large part dependent on the tube components (procoagulant/anticoagulant, surfactant and gel separator) [4] and centrifugation settings [5]. Gel separators offer greater stability of analytes, by separating cells from plasma or serum and almost eliminating subsequent changes in analytes due to cellular activity or release from cellular rupture post centrifugation. The presence of “white particulate matter” on top of the gel consisting of cells and fibrinogen in lithium heparin impacts on LDH, and there is presence of oil droplet which was first reported from our laboratory in 2004 [6]. Cadamuro et al. [7] observed the presence of fat droplets in the “white particulate matter” that most probably originated from the separator gel in the tubes. Gel separator tubes have been reported to affect measurements of therapeutic drugs [8], red blood cells surpassing the separator gel barrier in both plasma and serum tubes increasing the potassium concentration [9], and the separator gel components in tubes has been the source of interference in their liquid chromatography-MS testosterone assay [10].

BD released the next-generation blood separation technology lithium heparin tube, Barricor designed to enhance plasma quality by lowering cell count in plasma. Rather than using gel as the separator it uses a mechanical device, high density plastic to separate cells from plasma. There are only a few publications on the Barricor tube and one is by Füzéry et al. [11] for suitability with the Beckman TnI assay. In this study we examined the effect on cell count in the Barricor tubes by using four biochemical analytes (potassium, glucose, phosphate, and LDH) and examining the composition of the buffy coat.

Materials and Methods

The study was in two parts. The study compared the BD Barricor tube (#365032, Lot 6011667) against the current commercial tubes from BD lithium heparin tube or plasma separator tube (PST) (#367375, Lot 6179937), rapid serum tube (RST) (#368774, Lot 160308), and the serum separator tube (SST) #367954, Lot 6144963). Part one was a comparison of the most significantly affected common analytes (potassium, glucose, phosphate, and LDH) by the presence of cells in the plasma or serum component post centrifugation and gel/mechanical device separation at four time points. In total 22 healthy participants were recruited with appropriate ethical approval and informed consent. The blood was collected by an experienced phlebotomist into serum tubes.
first (RST or/and SST) then into the lithium heparin tubes (PST or/and Barricor). The RST, PST and Barricor tubes were mixed as recommended by BD and centrifuged without delay while the SST tube was allowed to clot for 30 minutes then centrifuged. All tubes were centrifuged at 4000, 3000 and 2050g for 10 minutes at 20 °C. The 2050g was selected as per BD recommendations [12]. The samples were analysed at 0 time, 24, 48 and 168 hours post storage at 2-8 °C. Analysis was performed within ~30 minutes post centrifugation on a DxC800 general chemistry analyser (Beckman Coulter, Brea, CA, USA). Results were significant and analytically important when the percent difference was greater than the upper limit of the cumulative imprecision of the between-run coefficient of variations (CVs) from the internal quality control concentrations on the Beckman analysers. The acceptable cut-off CVs used was: <2% for potassium at 3.84mmol/L, <3% for glucose at 4.5mmol/L, <2.5% for LD at 182 U/L and <2% for phosphate at 1.01mmol/L. These acceptable limits are significantly lower than the within-subject coefficient of variation published by Ricos et al. [13] (<4.8% for potassium, <6.5% for glucose, <6.6% for LD, and <8.5% for phosphate in serum). The second part was to examine the buffy coat in each of the four tubes and demonstrate the cell presence and how this aligns to analytical changes. This was performed from the blood of a single participant. The blood was collected in each of the four tubes and then the tubes were centrifuged at 3000g post completion of part one of the studies where 3000 g showed the least changes and aligns with previous findings [5]. Post centrifugation, all but the last ~0.5mL of plasma or serum was left in each tube. The 0.5mL sample in each tube was well mixed to obtain a homogenous mixture, and then a 0.1mL aliquot was transferred into the cytospin cups with slides (Cytospin 4, Thermo Fisher Scientific, Sydney, Australia) to concentrate the cells. The samples were centrifuged at 1260 rpm for 5 minutes. The slides were stained with May-Grünwald-Giemsa stain and processed as per normal blood film. The slides were examined for composition at x20 and x400 magnification. The cell count was performed on grid slides with counting chambers using a phase contrast microscope (x40 magnification). The concentration of all cell is x10⁶. The samples were diluted 1 in 20 with normal saline prior to loading on the counting chamber.

**Results**

![Barricor tube x20 magnification.](image)

![PST x20 magnification.](image)

![RST x20 magnification.](image)

![SST x20 magnification.](image)

![PST x400 magnification.](image)

![RST x400 magnification.](image)

![SST x 400 magnification.](image)

**Figure 1:** Stained slides from each of the BD tubes buffy coat layers to demonstrate visually the cell type and amount of cells present at 20x and 400x magnification.
Table 1: Potassium, glucose, LDH and phosphate results using the different BD tubes at different centrifugation speeds and storage times. (# clinically significant difference).

<table>
<thead>
<tr>
<th>Tube</th>
<th>Potassium (mmol/L)</th>
<th>Glucose (mmol/L)</th>
<th>LDH (U/L)</th>
<th>Phosphate (mmol/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Time (hours)</td>
<td>Difference (%)</td>
<td>Time (hours)</td>
<td>Difference (%)</td>
</tr>
<tr>
<td>PST 4000g</td>
<td>0 24 48 168</td>
<td></td>
<td>0 24 48 168</td>
<td></td>
</tr>
<tr>
<td>3.84</td>
<td>4.02 4.1 4.1 4.6 7.4</td>
<td>-19.98</td>
<td>5.22 4.89 4.7 4.13 -6.4 -10.2 -21.3</td>
<td>211 226 230 265 7.4 9.5 # 26.3</td>
</tr>
<tr>
<td>3.86</td>
<td>4.01 4.1 3.9 5.8 10.4</td>
<td>5.23 4.98 4.82 3.44 -4.8 -8.2 # -18.1</td>
<td>210 228 234 268 8.8 11.7 # 27.9</td>
<td>1.2 1.2 1.25 1.37 2.2 3.7 13.2</td>
</tr>
<tr>
<td>3.8</td>
<td>3.98 4.1 4.8 8.1 26.4</td>
<td>5.23 4.94 4.73 4.16 -5.7 -9.6 # -20.0</td>
<td>284 230 236 271 13 16.1 61.6 # 32.8</td>
<td>1.2 1.2 1.25 1.38 2.1 3.7 14.5</td>
</tr>
<tr>
<td>SST 2050g</td>
<td>4.02 4.1 4.1 1.9 7.3</td>
<td>5.04 4.92 4.91 4.87 -2.4 -2.7 -3.4</td>
<td>217 222 222 213 2.3 4.3 # 1.7</td>
<td>1.3 1.3 1.27 1.27 1.3 2.7 4.2</td>
</tr>
<tr>
<td>3000g</td>
<td>4 4.07 4.1 1.7 1.8 7.5</td>
<td>5.03 4.93 4.92 4.8 -2.1 -2.2 -4.4</td>
<td>214 221 220 212 1.1 2.9 0.5</td>
<td>1.3 1.3 1.27 1.3 1.6 1.6 4.5</td>
</tr>
<tr>
<td>2050g</td>
<td>4.04 4.15 4.2 2.7 3.7 13.1 #</td>
<td>5.06 4.93 4.86 4.68 -2.6 -4 -7.1</td>
<td>286 215 217 212 4.9 5.7 # 3.3</td>
<td>1.3 1.3 1.27 1.31 1 0.5 4.2</td>
</tr>
<tr>
<td>RST 4000g</td>
<td>4.04 4.08 4.1 1.1 0.9 4.7</td>
<td>5.12 5.02 5.03 5 -2.1 -1.9 -2.4</td>
<td>282 202 198 189 0 1.8 6.3</td>
<td>1.3 1.3 1.31 1.34 1.5 0.8 3.5</td>
</tr>
<tr>
<td>3000g</td>
<td>4.02 4.08 4.1 1.5 1.3 5.5</td>
<td>5.1 5.02 5 4.85 -1.7 -2 -5.8</td>
<td>282 200 198 190 0.6 1.7 5.2</td>
<td>1.3 1.3 1.34 1 0.4 3.2</td>
</tr>
<tr>
<td>2050g</td>
<td>3.99 4.11 4.2 3.3 4.6 8.9 #</td>
<td>5.14 5.04 5.03 4.95 -1.9 -2.2 -4</td>
<td>281 200 198 190 0.3 1.3 5.1</td>
<td>1.3 1.3 1.31 1.35 1.7 1.2 4.4</td>
</tr>
<tr>
<td>Barricor 4000g</td>
<td>3.8 3.89 3.9 2.4 2.7 6.3</td>
<td>5.4 5.08 4.95 4.7 -5.9 -8.4 # -13.2 #</td>
<td>284 209 209 216 2.6 2.7 6.1</td>
<td>1.2 1.2 1.22 1.27 1.1 1.5 5.5</td>
</tr>
<tr>
<td>3000g</td>
<td>3.8 3.87 3.9 2.1 2.2 6.6</td>
<td>5.27 5.09 5 4.7 -2.9 -5.1 -9.8 #</td>
<td>284 207 208 219 1.9 2.2 7.6</td>
<td>1.2 1.2 1.21 1.27 1.3 1.2 6</td>
</tr>
<tr>
<td>2050g</td>
<td>3.8 3.87 3.9 2.6 4.2 8.0 #</td>
<td>5.4 5.02 4.89 4.6 -6.6 -8.6 # -14.2 #</td>
<td>284 214 214 222 4.8 4.9 9.5</td>
<td>1.2 1.2 1.21 1.27 1.7 1.8 6.5</td>
</tr>
</tbody>
</table>

Table 2: Cell count of the buffy coat in each of the four tubes performed at x400 magnification with phase contrast microscope.

<table>
<thead>
<tr>
<th>Tube</th>
<th>White Cell Count x10^6</th>
<th>Red Cell Count x10^6</th>
<th>Platelet Count x10^6</th>
</tr>
</thead>
<tbody>
<tr>
<td>LH</td>
<td>405</td>
<td>12150</td>
<td>4050</td>
</tr>
<tr>
<td>Barricor</td>
<td>36</td>
<td>1197</td>
<td>7290</td>
</tr>
<tr>
<td>RST</td>
<td>24</td>
<td>2835</td>
<td>153</td>
</tr>
<tr>
<td>SST</td>
<td>24</td>
<td>4293</td>
<td>144</td>
</tr>
</tbody>
</table>

The biochemical analytes comparative stability results (means, percentage differences and if the difference is clinically significant) are shown in (Table 1). The Barricor tube plasma results clearly showed decreased changes in the biochemical analytes concentration over time when compared with the PST plasma. The PST plasma for all four analytes showed the most significant changes across all tubes, centrifugation settings and storage time points. It exceeded the allowed technical limits for each time point and centrifugation setting. The most significant changes across all tubes were observed with the 2050g centrifugation. The least changes observed overall were with the 3000g centrifugation, and with the serum from the RST. The slides, Figure 1 showed significant higher number of cells were present in the lithium heparin tubes, PST having higher red cells and white cells numbers than the Barricor tube which had slightly higher platelet numbers. The least number of cells were present in the SST serum sample. The breakdown of cell types from counts under microscope in each tube is shown in (Table 2). The haemolysis index levels measured on the Beckman Dx800 analysers were insignificant in all tubes, below levels that affect any analytes.

Discussion

Sample quality can influence accuracy of results of any test. The data from part one showed the mechanical barrier in the Barricor tube is significantly more effective in separating cells from the plasma than the gel in the PST. Hence analytical changes in the four analytes are minimised across the different storage time points and centrifugation settings. It is well known blood cells (red, white and platelets) are progressively lysed post storage, leading to release of potassium, LDH, and phosphate while viable cells continue to consume glucose[7]. The LDH activity decrease in serum tubes is due to fraction F4(F4) instability on storage at 0-4 ºC. The concentration of F4 falls rapidly in the first week, and disappears within 3 weeks [14]. Yet IFCC states serum is the preferred sample for LDH [15]. The data in this study showed heparin inhibits this degradation of the F4, hence LDH activity follows the same path as the other cell present analytes (potassium and phosphate), increasing with time. The 3000g centrifugation setting was the most effective in minimising cell numbers presence in both plasma and serum, and provided the best stability in concentration of analytes. The slides in part two data clearly showed the amount of buffy coat varies from tube to tube. With the x20 magnification the red and white cell numbers were highest in the PST, followed by the Barricortube, the SST and the least in the RST tube. The Barricor tube had the highest platelet cell count. The x400 magnification provided clear distinction on the cell types present, and that the platelets were removed during the clotting process in the RST and SST tubes. WHO report that a buffy coat presents with all three cell types, cell debris, fibrin, decreasing the specimen integrity.

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The "white particulate matter" formation of insoluble fibrin is enhanced when heparin plasma is stored at low temperature [17]. It has also been suggested that individual molecules within pharmaceutical heparin preparations may have procoagulant activity particularly in the absence of antithrombin III [18]. Stimulation of platelet factor 4 (PL4), which is a protein stored in the alpha granules of platelets that is released during platelet aggregation, can neutralize heparin [19,20]. PL4 may be released at the site of collection and into the sample or from platelets trapped in microdots which in turn can aid the formation of fibrin strands in the specimen. If all of these physico-chemical factors are present in the right balance, fibrin formation is likely to be enhanced in heparinised specimens. Over time heparin activity in stored lithium heparin blood specimens is decreased, mainly due to PL4 activity. In our tertiary level laboratory up to 5% of the lithium heparin specimens stored for two days at 4 °C showed visible fibrin formation which would render them unsuitable for re-analysis. Such stored plasma specimens require aliquoting and re-centrifugation for re-testing or performing additional tests and due to the presence of cells renders the plasma samples unsuitable for determination of some analytes such as potassium and glucose. The only other notable factor is the Barricor tube with the mechanical separator leads to slightly decreased plasma volume. The limitations of this study are therapeutic drugs and broader range of analytes was not studied for stability due to the mechanical separator, if it does bind to drugs leading to concentration changes.

Conclusion

The Barricor tube offers improvement in lithium heparin plasma quality as shown by the decreased changes in potassium, LDH and phosphate concentrations due to cellular lysis and glucose results from cellular consumption upon sample storage. The least differences in results were observed across all four tubes with 3000 g centrifugation speed. Based on the data from this study the Barricor tube should become the new standard for lithium heparin plasma.

Acknowledgement

BD provided the Barricor and RST tubes but played no part in study design or write up.

References
