Changes in blood gas samples produced by a pneumatic tube system

P O Collinson, C M John, D C Gaze, L F Ferrigan, D G Cramp

Aims: To investigate the effect of a pneumatic tube system (PTS) on the results of samples sent for blood gas analysis to a central laboratory.

Methods: Blood gas samples were analysed immediately or sent via the PTS to the laboratory for analysis. In addition, samples sent via the PTS in a pressure sealed container were compared with those sent non-pressure sealed to the laboratory.

Results: Samples sent via the PTS had significant alterations in their pO₂ values, which were not seen when samples were carried by hand to the laboratory. There was no effect on pCO₂ and pH values.

Conclusions: Samples for blood gas analysis should be transported via a PTS using a pressure sealed container to avoid artefacts in the pO₂.

METHODS

Arterial blood samples were drawn from patients in the intensive care unit (ICU) of a UK district general hospital. The ICU is located in a separate building from the pathology laboratory, but on the same geographical site. All of the hospital wards including the ICU are connected to the central pathology laboratory via a pneumatic tube delivery system. The ICU is 520 metres from the laboratory, with a median delay from sampling to arrival at the laboratory of 19 minutes (interquartile range, 13 to 23). Ethical permission for our study was obtained from the local research ethics committee. Samples were taken into commercially supplied preheparinised syringes (Mieno Corp, Tokyo, Japan). Visible air bubbles were expelled and the samples capped and processed immediately.

A three stage study was performed, as follows:

Phase 1. Consecutive samples were drawn in duplicate from patients over a two week period. One sample was analysed immediately on the ICU by a member of the ICU staff. The second sample was capped, sealed in a gas tight plastic envelope, and sent to the laboratory via the PTS. Immediately on receipt by the laboratory the sample was analysed by a member of the laboratory staff. The time of sample draw and sample analysis was recorded for each sample.

Phase 2. Consecutive samples were drawn in duplicate from patients over a two week period. One sample was analysed immediately on the ICU by a member of the ICU staff. The second sample was capped, sealed in a gas tight plastic envelope, and sent to the laboratory via the PTS. Immediately on receipt by the laboratory the sample was analysed by a member of the laboratory staff. The time of sample draw and sample analysis was recorded for each sample.

Phase 3. Consecutive samples were drawn in triplicate from patients over a two week period. One sample was analysed immediately on the ICU by a member of the ICU staff. The remaining two samples were then capped and each sealed in a separate gas tight plastic envelope. One sample was then sent to the laboratory via the pneumatic tube system using the conventional canister. The second was sealed inside a pressure tight container and sent to the laboratory via the PTS inside the conventional canister. Immediately on receipt by the laboratory the samples were analysed by a member of the laboratory staff. The time of sample draw and sample analysis was recorded for each sample.

Blood gas analysis in the ICU was performed using ion sensitive electrodes on a Corning 850 blood gas analyser (Corning Instruments, Halstead, Essex, UK). Coefficients of variation (CVs) were—pH: 0.1% at 7.152, 0.3% at 7.429, 0.1% at 7.587; pCO₂: 1.5% at 10.30 kPa, 3.1% at 5.56 kPa, 2.9% at 3.09 kPa; pO₂: 1.1% at 8.05 kPa, 7.9% at 12.66 kPa, 2.3% at 17.46 kPa. Blood gas analysis in the laboratory was performed using ion sensitive electrodes on an ABL 50 blood gas analyser (Radiometer UK, Crawley, Sussex). CVs were—pH: 0.1% at 7.128, 0.1% at 7.384, 0.1% at 7.625; pCO₂: 8.3% at 8.53 kPa, 1.9% at 5.19 kPa, 4.1% at 2.36 kPa; pO₂: 2.5% at 8.50 kPa, 8.8% at 14.9 kPa, 2.5% at 23.9 kPa. The laboratory maintained both blood gas analysers with daily quality control measurements performed by a qualified laboratory technician. Results of pH, pO₂ and pCO₂ measurements were compared by non-parametric statistical analysis, with calculation of the
RESULTS

Twenty samples were analysed on the ICU and after being immediately transferred by hand carriage to the laboratory. There was no difference between the values for pH (median difference, 0.001; IQR, 0.015; \( p = 0.9461 \)), \( pCO_2 \) (median difference, 0.230; IQR, 0.408; \( p = 0.4487 \)), or \( pO_2 \) (median difference, 0.280; IQR, 1.27; \( p = 0.7150 \)) by Mann Whitney U test. pH ICU = 0.942 pH lab + 0.432; \( pCO_2 \) ICU = 1.079 \( pCO_2 \) lab – 0.018; median difference and interquartile range (IQR) of differences. Passing and Bablock regression plots and Bland-Altman difference plots were constructed for each analyte and the results compared.

Figure 1 Altman and Bland plot for \( pO_2 \) determined in samples carried by hand to the laboratory and assayed on the intensive care unit (ICU).

Figure 2 Altman and Bland plot, for \( pO_2 \), determined in samples assayed in the intensive care unit (ICU) and carried to the laboratory via the pneumatic tube.

Median difference and interquartile range of differences. Passing and Bablock regression plots and Bland-Altman difference plots were constructed for each analyte and the results compared.

DISCUSSION

The determinants of TAT are: the time interval from blood draw to the time of delivery of the sample to the point of analysis, the analytical TAT, and the time taken to return results. Whole blood analytical systems and modern fast analytical systems can result in a reduction in analytical TAT to as little as five minutes. Electronic data links mean that results can be viewed remotely within seconds of analysis. Therefore, the limitation remains the time taken to deliver the sample to the point of analysis. The options therefore are to move analysis nearer to the patient, either by the use of satellite laboratories or POCT, or speed up the transport of samples. This is the rationale for the use of PTS for sample delivery. The case for the existence of satellite laboratories is the maintenance of laboratory quality. The use of POCT is based on the premise that the quality of results produced is clinically acceptable. The PTS option assumes that samples are unaffected by transport within the system. Most analytes would be expected to be unaffected by transport and most studies support this, although increased concentrations of lactate dehydrogenase were reported in samples allowed to clot before analysis. One recent study of 291 samples in routine clinical use in particular showed no increased incidence of haemolysis in the samples transported via PTS. However, samples transported by PTS may not be suitable for all applications. The samples most sensitive to transport changes would be expected to be those taken for blood gas analysis. Recently, a report suggested that the presence of air contamination resulted in significant interference in PTS transported samples.

We confirmed that the presence of air contamination resulted in significant interference in PTS transported samples.

Mean of \( pO_2 \) (kPa)

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<tr>
<th>Mean of ( pO_2 ) (kPa)</th>
<th>Difference between methods (kPa)</th>
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<tbody>
<tr>
<td>6</td>
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<tr>
<td>8</td>
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<td>10</td>
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<td>-1.0</td>
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<td>14</td>
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Pressure changes as a possible cause of interference have not been investigated previously. It seems that the previous
studies did not use pressure sealed containers. The lack of correlation between delay time and pO₂ difference is in accordance with other studies. The changes in pO₂ but not CO₂ and pH are at first surprising, but are readily explained. Blood pH and pCO₂ are tightly controlled and heavily buffered. The determination of pO₂ is made solely from the oxygen dissolved in solution, in excess of that bound by the haemoglobin in erythrocytes. Thus, any minor changes in pressure will affect the dissolved oxygen content of the sample, especially if there are microbubbles, and hence change the pO₂.

In routine clinical use, samples sent via a pneumatic delivery system may show a large discrepancy when compared with samples analysed immediately. This most likely results from air bubbles in the samples, despite care being taken to expel any air bubbles, combined with pressure effects from the delivery system. Unless samples sent for blood gas analysis to the laboratory can be transported in a pressure tight system that can be conveniently used, samples should be hand carried or analysed by POCT. This needs to be incorporated into recommendations for the implementation of such systems.²

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