Comparative Accuracy Evaluation of a Blood Glucose Meter With Novel Hematocrit Correction Technology, With Three Currently Used Commercially Available Blood Glucose Monitoring Systems

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Abstract
Hematocrit is known to influence glucose values obtained on some blood glucose meters, with bias observed especially at low and high hematocrit levels. We evaluated the performance of a meter with hematocrit correction technology alongside 3 other commercially available meters. Capillary blood samples from 100 subjects were analyzed in duplicate and compared to the plasma values obtained by reference laboratory analyzer. Bias, error grid, and sensitivity to hematocrit analyses were performed for each meter. Average percentage bias was similar for all meters, however the evaluated meter performed best with respect to error grid analysis, with 100% of values falling within the “no effect on clinical action” and “no risk” categories and did not display any hematocrit associated bias.

Keywords
accuracy, blood glucose meter, hematocrit

Diabetes is a chronic condition, affecting approximately 425 million people globally.¹ Achieving tight glucose control in people with diabetes has been shown to reduce the development of a number of diabetes-associated complications.² Self-monitoring of blood glucose (SMBG) provides a means to improve the glycemic control of a person with diabetes, in addition to identifying hypoglycemic events in those on insulin therapy.³

The guidelines for determining system accuracy on blood glucose meters (BS EN ISO 15197:2015)³ state that accuracy should be determined from at least 100 different subjects, and evaluated under actual conditions of use to include the effects of both systematic error (measurement bias) and random error (measurement imprecision). The system accuracy evaluation should be performed on fresh blood samples, at least in duplicate and compared to reference laboratory glucose concentrations, both pre- and post-measurement with the glucose meter.

Hematocrit has previously been shown to demonstrate considerable influence on the glucose readings obtained from certain blood glucose monitors, even when tested within the manufacturer’s recommended hematocrit range.⁵,⁶ Typically, a positive bias is observed at low hematocrit levels and a negative bias at higher hematocrit levels.⁷,⁹ In this study, the system accuracy of the evaluated blood glucose meter, the GlucoRx HCT meter (GlucoRx, Surrey, UK), which utilizes a novel hematocrit correction technology, was compared to 3 other blood glucose meters used in the United Kingdom across a wide range of blood glucose concentrations, similar to that specified in Section 8 BS ISO 15197:2015.⁴

Methods
This study was performed between March and April 2018 at the Joint Clinical Research Facility, ILS2, Swansea University, Swansea, UK, in compliance with Good Clinical Practice and was approved by Wales REC 6 (18/WA/0023) prior to commencement of the study.

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Subjects
A total of 114 participants were recruited to achieve 100 full, evaluable datasets. Subjects with diabetes mellitus (type 1 and type 2) and without diabetes were included in the study, and on entry were all aged ≥18 years old. Informed, signed consent was received from all participants.

Blood Glucose Monitoring Systems
The evaluated meter (meter A) is a multiparameter meter, capable of measuring blood glucose, ketones and hematocrit. The meter uses advanced GDH-FAD enzyme technology with an AC signal which provides the hematocrit result, and DC signal which calculates glucose. The meter then modulates the glucose level according to the hematocrit value allowing enhanced accuracy in measurement of glucose. Meter A was compared to three other meters (meters B, C, and D—in no particular order); these included the Roche Accu-Chek Aviva (Roche Diagnostics Ltd, West Sussex, UK), which measures blood glucose only, and two dual glucose and ketone meters; Caresens Dual BGM System (distributed by Spirit Healthcare, Leicester, UK); and Menarini GlucoMen Areo 2K (A Menarini Diagnostics, Berkshire, UK). Meters B, C, and D were selected for this study due to their availability and current use within the United Kingdom. For this study, the ketone function was not used on the dual systems. All meters and strips were purchased directly by the research team and were not provided by the manufacturers.

Study Procedure
Procedures performed were similar to those outlined in EN ISO 15197:2015. Briefly, eligible subjects had a fingerprick performed using a high flow lancet. Capillary blood (approximately 100 ul) was collected into lithium heparin anticoagulant (Microvette, Sarstedt, Leicester, UK), with the sample used for determination of plasma glucose on the reference laboratory glucose analyzer. Following this, duplicate blood glucose measurements were performed on the 4 blood glucose meters in a random order, before a second reference glucose sample was collected. Finally, hematocrit was determined. For the reference YSI measurement, the Microvette tubes were centrifuged within 5 minutes of collection and the plasma component used for glucose determination.

Laboratory Measurements
The reference laboratory plasma glucose was determined using a YSI 2300 Stat Plus (Yellow Springs Instruments, Fleet, UK). Daily internal quality control was performed before any study samples were run (Assayed Chemistry Control Plus, levels 2 and 3, Randox, UK). Microvette tubes were centrifuged within 5 minutes of collection and the plasma component used for glucose determination.

Hematocrit was determined using a HemoControl analyzer (EKF Diagnostics, Cardiff, UK)

Data Analysis
Data were excluded from analysis if the reference laboratory measurements differed by >4% at glucose concentrations <100 mg/dL or >4 mg/dL at glucose concentrations ≥100 mg/dL, if valid glucose readings were not obtained on all meters or if insufficient sample volume was available for all measurements to be performed.

Accuracy for each meter was assessed according to the number of readings within 5, 10, and 15 mg/dL (glucose <100 mg/dL) or 5, 10, and 15% (glucose ≥100 mg/dL) of the reference glucose value and the accuracy for each individual meter compared using the Cochran-Mantel-Haenszel test. Bias plots, Parkes consensus error grids (CEG),10 and surveillance error grids (SEG)11 were performed for each meter. In addition, sensitivity of the individual meters to hematocrit was assessed by regression analysis of the relative glucose differences (meter and reference glucose) versus hematocrit.

Results
The batches of meters and strips used were:

- Meters 4279317350041211 / 4279317350041299; Strips TD17 G110-BHF
- Meters 45900007291 / 45900013602 / 45900019971; Strips 497275
- Meters F024073F0910 / F024073F0906; Strips PJ25CAQ1B
- Meters GT168145 / GT168206; Strips HS170705

For the 100 evaluated samples, the observed glucose range determined on the reference analyzer was 33 to 581 mg/dL and hematocrit was 28 to 50%.

The average percentage bias between the meters and reference glucose measurement were 5.3, 5.0, 5.9, and 6.6%, meters A to D respectively (Figure 1).

For meters A to D, respectively, 99.5, 98.0, 96.0, and 96.0% of the meter values were within ±15 mg/dL (glucose <100 mg/dL) or ±15% (glucose ≥100 mg/dL) of the reference glucose value (Table 1), with meter A displaying significantly greater accuracy when compared to both meters C and D (P = .044 and .043, respectively).
Error Grid Analysis

For meter A, 100% of glucose values fell within Zone A (no effect on clinical action) and were classed as “no risk” on the CEG and SEG plots, respectively. For the remainder of the meters, the corresponding values were 100% and 97.5% (meter B), 99.5% and 98.5% (meter C) and 98.0% and 96.5% (meter D) (Table 2 and Figures 2 and 3).

Sensitivity to Hematocrit

The hematocrit values of the samples tested were within the accepted ranges for all meters. Across an increasing hematocrit range, the magnitude of spread of relative glucose differences (reference and meter) remained relatively constant for meters A, B, and C, however greater variation was observed for meter D (Figures 4 and 5). Neither the slope nor intercept of the calculated regression lines differed from zero for meters A or D but were significantly different for both meters B and C (all \( P < .01 \)).

Discussion

In this study, the performance of a blood glucose meter with novel hematocrit correction technology (meter A) was evaluated and compared to the performance of three
commercially available blood glucose monitoring systems across a wide glucose concentration range, based on the guidelines stated in Section 8 BS ISO 15197:2015. One of the limitations of this study is that while the latest ISO guidelines were followed in terms of number of samples and glucose concentration range tested, only a single batch of test strips was employed. The three comparator meters (B to D) were selected due to their availability and usage within the United Kingdom. The accuracy of readings generated by blood glucose meters is essential to ensure both tight glucose control and patient safety; as such, the International Organization for

<table>
<thead>
<tr>
<th>CEG zone</th>
<th>Slight, lower</th>
<th>Slight, higher</th>
<th>Moderate, lower</th>
<th>Moderate, higher</th>
<th>Great, lower</th>
<th>Great, higher</th>
<th>Extreme</th>
</tr>
</thead>
<tbody>
<tr>
<td>A B C D E</td>
<td>None</td>
<td>Slight, lower</td>
<td>Slight, higher</td>
<td>Moderate, lower</td>
<td>Moderate, higher</td>
<td>Great, lower</td>
<td>Great, higher</td>
</tr>
<tr>
<td>Meter A</td>
<td>200</td>
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<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>200</td>
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<td>100.0%</td>
<td>—</td>
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<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>100.0%</td>
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<tr>
<td>Meter B</td>
<td>200</td>
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<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>195</td>
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<tr>
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<td>—</td>
<td>—</td>
<td>—</td>
<td>97.5%</td>
</tr>
<tr>
<td>Meter C</td>
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<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>197</td>
</tr>
<tr>
<td>99.5%</td>
<td>0.5%</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>98.5%</td>
</tr>
<tr>
<td>Meter D</td>
<td>194</td>
<td>2</td>
<td>2</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>193</td>
</tr>
<tr>
<td>98.0%</td>
<td>1.0%</td>
<td>1.0%</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>96.5%</td>
</tr>
</tbody>
</table>

Figure 2. Consensus error grids, meters A, B, C, and D.
Figure 3. Surveillance error grids, meters A, B, C, and D.

Figure 4. Relative difference between meter and YSI according to hematocrit, meters A, B, C, and D.
Standardization (ISO) has published guidelines to evaluate blood glucose meters. According to the most recent recommendations (ISO 15197:2015), 95% of meter readings should be within ±15 mg/dL (for glucose concentrations <100 mg/dL) or ±15% (glucose concentrations ≥100 mg/dL) of the reference glucose concentrations. In this study, all meters were found to achieve these targets, ranging from 99.5% (meter A) to 96.0% (meters C and D).

With respect to the CEG and SEG scores the evaluated meter (meter A) performed the best with all results in zone A (no effect on clinical outcome—CEG) or “no risk” (SEG). Meter D was more variable with 1% of values falling in Zone C (leading to unnecessary treatment; CEG) and “moderate, lower risk” (SEG).

The magnitude of the relative glucose differences between meter and reference glucose was similar across the tested hematocrit range for meters A, B, and C, but showed greater variation for meter D. However, despite this similarity in magnitude of spread for meters A, B, and C, meter A was the only one that did not display hematocrit associated bias.

**Conclusion**

In this study, the performance of all the blood glucose meters was within ISO 15197:2015 guidelines; meter A had the best performance due to the hematocrit correction technology employed within this meter.

**Abbreviations**

BGM, blood glucose meter; CEG, consensus error grid; ISO, International Organization for Standardization; SEG, surveillance error grid; SMBG, self-monitoring of blood glucose.

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