Evaluation of Cobas b 101 HbA1c Analyzer Performance for Point-of-Care Testing

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Background: The use of point-of-care (POC) devices for evaluating HbA1c is increasing; accordingly, comparisons between these devices and central laboratory methods are important. In the present study, we evaluated the analytical performance of the cobas b 101 analyzer for POC HbA1c testing.

Methods: The analytical quality of the cobas b 101 system was assessed based on repeatability, within-laboratory precision, linearity, and lot-to-lot reproducibility. Two specimen types, i.e., EDTA whole blood and capillary blood, were examined using the cobas b 101 system and the Variant II Turbo instrument.

Results: The method showed good linearity, with a coefficient of correlation of 0.990. In a comparison of two different HbA1c disk lots, a strong correlation ($r = 0.986$) and a mean % difference of -2.9% were observed. The cobas b 101 results using EDTA whole blood were strongly correlated with the Variant II Turbo results ($r = 0.958$), with a mean % difference of 0.8%; the cobas b 101 results using capillary blood were strongly correlated with the Variant II Turbo results, using EDTA whole blood ($r = 0.976$), with a mean % difference of 2.0%. A comparison between HbA1c levels in EDTA whole blood and capillary blood obtained using the cobas b 101 showed a strong correlation ($r = 0.985$) and a mean % difference of 1.3%.

Conclusions: The cobas b 101 analyzer is convenient for the measurement of HbA1c levels for diabetes management.

Key Words: HbA1c, Point-of-care testing, Diabetes

INTRODUCTION

The HbA1c level indicates a patient’s average glucose levels for the past 3 months and facilitates the long-term management of blood glucose levels by clinicians [1]. HbA1c levels are highly standardized worldwide owing to the development of reference measurement procedures and primary reference materials. In addition, test results are standardized according to the International Federation of Clinical Chemistry (IFCC) Working Group on HbA1c Standardization and the National Glycohemoglobin Standardization Program (NGSP) HbA1c Harmonization Program [2, 3]. Various studies have reported that immediate feedback on HbA1c levels improves glycemic control in patients with diabetes mellitus (DM) [4-6]. Furthermore, studies have shown that HbA1c levels and intensive care based on HbA1c monitoring are correlated with the risk of developing DM-associated complications [7-9].

Among laboratory examination methods, point-of-care (POC) testing is the quickest and most convenient method used by doctors to make clinical decisions [10]. Recently, various HbA1c testing devices for capillary blood or EDTA whole blood have shown positive results [10-14]. Furthermore, several studies have established the effectiveness of HbA1c monitoring by POC testing for glycemic control and the prevention of complications in patients with DM [15, 16].

In this study, we evaluated the analytical performance of the
cobas b 101 (Roche Diagnostics, Mannheim, Germany) analyzer for estimating HbA1c levels. The cobas b 101 analyzer was compared with the standard central laboratory method, which uses the Variant II Turbo instrument.

**MATERIALS AND METHODS**

1. **Study design**
   This study was performed from July to November 2015 in the Department of Laboratory Medicine at our institute. All analyses were reviewed and approved by the Institutional Review Board (IRB) at our institute (KBSMC 2015-01-049). Patient samples were obtained from residual EDTA samples or from EDTA and capillary samples collected immediately before the test. All participants provided written informed consent prior to blood collection. EDTA whole blood specimens were taken from the antecubital vein after at least 8 hours of fasting and analyzed promptly within 1 hour. Simultaneously, capillary blood collection was performed by finger pricking to obtain 2 µL of whole blood. Participant data were subjected to a de-identification process and coded with serial numbers for the test.

2. **Precision and linearity**
   Repeatability and within-laboratory precision were determined using two quality control (QC) materials with different HbA1c levels, supplied by the manufacturer of the cobas HbA1c control. Both QC materials were assayed in duplicate twice daily at 9 am and 3 pm for a total of 20 days according to the 2014 Clinical and Laboratory Standards Institute (CLSI) guideline EP05-A3 [17]. The linearity of the assay was evaluated for specimens with five different HbA1c levels with duplication by mixing venous EDTA blood samples from two patients according to the CLSI guideline EP06-A [18].

3. **Lot-to-lot reproducibility**
   The lot-to-lot reproducibility was evaluated using two different HbA1c disk lots (#434022-01 and #435021-01), and 20 residual venous EDTA blood samples treated in the same manner by the same user were evaluated for 5 days, with 4 samples evaluated in duplicate each day. The test results for 20 samples using the first lot were in the range of 5.2-8.9% HbA1c.

4. **Comparative analysis**
   Linear regression and %differences in estimated HbA1c levels between the cobas b 101 and Variant II Turbo were determined according to the 2013 CLSI guideline EP09-A3 protocol [19]. Two types of blood specimens were obtained immediately before the tests from the 40 enrolled participants: capillary blood and EDTA whole blood. The capillary finger prick samples and EDTA whole blood samples were analyzed using the cobas b 101 POC system in the phlebotomy room and the EDTA whole blood samples were analyzed using the Variant II Turbo in the central laboratory. The HbA1c levels of samples spanned the clinically relevant range of 4.8% to 8.6%.

5. **Laboratory method**
   HbA1c was measured at the central laboratory of our hospital using the Variant II Turbo (Bio-Rad Laboratories, Hercules, CA, USA), which is based on high-performance liquid chromatography (HPLC). The HbA1c program of the reference device was certified by the NGSP with documented traceability to the Diabetes Control and Complications Trial (DCCT) reference method. This test participated in the quality assurance survey from the College of American Pathologists (CAP), and all five of the challenges were within the acceptable range, with a mean bias of 0.06% for HbA1c levels. QC materials with low (5.10 ± 0.2%) and high (9.80 ± 0.2%) levels were used. The coefficient of variation was 0.96-3.03% for the low-HbA1c QC materials and 1.18-2.90% for the high-HbA1c QC materials during the study period.

6. **POC analyzer methods**
   The cobas b 101 POC analyzer (Roche Diagnostics) is based on a photometric transmission measurement method using a latex agglutination inhibition immunoassay. The test uses 2 µL of EDTA venous whole blood or 2 µL of capillary whole blood. The POC devices were designed to operate with ready-to-use HbA1c disks and provide results in 340 seconds. This device can measure HbA1c levels in the 4-14% range. The test method was certified by the NGSP and standardized or traceable to the DCCT reference assay. HbA1c values obtained using both devices are expressed in NGSP units as % HbA1c.

7. **Statistical analysis**
   Test imprecision was analyzed based on repeatability and within-
laboratory precision. Each formula was obtained from EP05-A3 [17]. The lot-to-lot reproducibility and comparative results were evaluated based on Pearson’s correlation coefficients (r). The results are displayed in scatter plots. %Differences between each POC test result and the reference method were analyzed by calculating the percentage of HbA1c reporting unit differences and a Bland-Altman plot was generated. The acceptable standard was a difference of ±6% according to the HbA1c acceptable limit established by NGSP, which was applied in the CAP GH2 survey [20]. All statistical analyses were implemented in Microsoft Excel 2010 and IBM SPSS version 18.0 (IBM, New York, NY, USA) and statistical significance was defined as P<0.05.

RESULTS

1. Precision and linearity

Imprecision was analyzed using QC materials with two levels of HbA1c for 20 days; the mean levels were 5.1% (normal) and 9.5% (pathological). The standard deviation (SD) and coefficient of variation (CV) were 0.25% and 4.83% for repeatability and 0.27% and 5.22% for within-laboratory precision tests for the normal level, and 0.21% and 2.22% for repeatability and 0.24% and 2.56% for within-laboratory precision for the pathological level, respectively (Table 1). This method showed good linearity between HbA1c levels of 4.6% and 13.3%. The estimated slope and intercept of the regression were 1.014 and 0.392, respectively with a coefficient of correlation of 0.990.

Table 1. Imprecision with 95% confidence intervals (95% CI) for the cobas b 101 HbA1c analyzer based on EP05

<table>
<thead>
<tr>
<th>Quality control materials</th>
<th>Normal level (Mean 5.1% HbA1c)</th>
<th>Pathological level (Mean 9.5% HbA1c)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SD (95% CI)</td>
<td>%CV (95% CI)</td>
</tr>
<tr>
<td>Repeatability</td>
<td>0.25 (0.20–0.31)</td>
<td>4.83 (3.97–6.18)</td>
</tr>
<tr>
<td>Within-Laboratory Precision</td>
<td>0.27 (0.23–0.32)</td>
<td>5.22 (4.48–6.25)</td>
</tr>
<tr>
<td></td>
<td>SD (95% CI)</td>
<td>%CV (95% CI)</td>
</tr>
<tr>
<td></td>
<td>0.21 (0.17–0.27)</td>
<td>2.22 (1.83–2.85)</td>
</tr>
</tbody>
</table>

Fig. 1. Lot-to-lot reproducibility of the cobas b 101 HbA1c test. (A) Regression analysis of the cobas b 101 using EDTA venous blood from two different lots. The linear curve and 95% confidence interval [CI] are represented as red and blue lines, respectively. (B) Bland-Altman difference plot summarizing lot-to-lot comparisons.

2. Lot-to-lot reproducibility

The lot-to-lot reproducibility for cobas b 101 HbA1c test results using two different lots is shown in Fig. 1A (r = 0.986, P < 0.001). The mean %difference was -2.9% (range -8.7% to 0.0%) (Fig. 1B). For the first lot, there was a strong correlation between the results obtained using the cobas b 101 and Variant II Turbo; the estimated coefficient of correlation and %difference were 0.983 (P < 0.001) and -1.5% (range -8.6% to 2.6%), respectively. For the second lot, there was also a strong correlation between the results obtained using the cobas b 101 and the Variant II Turbo, with a correlation coefficient of 0.985 (P < 0.001) and a %difference of 1.7% (range -5.0% to 5.8%). The slope and the intercept of the regression for
the two lots using the central laboratory method were 0.905 and 0.525 for the first lot and 0.834 and 0.539 for the second lot.

3. Comparative analysis

The regression of the estimates obtained using the cobas b 101 HbA1c and Variant II Turbo using EDTA whole blood showed a slope of 0.946 and an intercept of 0.374. On the scatter plot, the graph showed a strong correlation, with a correlation coefficient of 0.958 ($P<0.001$; Fig. 2A). The mean %difference was 0.8% (range -5.4% to 5.9%) and tended to be more highly dispersed at the higher HbA1c level (Fig. 2B). No specimen exceeded a 6% %difference.

When comparing the cobas b 101 analyzer results using capillary blood and the Variant II Turbo results using EDTA whole blood, a regression line with a slope of 0.991 and intercept of 0.179 was obtained. On the scatter plot, the graph showed a strong correlation, with a correlation coefficient of 0.976 ($P<0.001$; Fig. 2C). The mean %difference was 2.0% (range -4.8% to 6.8%) (Fig. 2D). Moreover, a strong correlation was observed between EDTA whole blood and capillary HbA1c levels, with a correlation coefficient of 0.985, using the cobas b 101 system ($P<0.001$; Fig. 3A). The mean %difference was 1.3% (range -5.5% to 7.7%) (Fig. 3B). Table 2 contains all results for the reproducibility and comparative analyses.
indicating the expected device or lot value (Y) when the other
device or lot (X) had a 6.5% HbA1c level.

**DISCUSSION**

Venous blood sampling is currently the gold standard for the
assessment of blood glucose levels. However, this sampling method
cannot be applied to home-based glucose self-monitoring. Capil-
lary blood glucose testing using portable POC devices has been
hailed as an alternative method to venous blood sampling owing
to its better compliance, rapid reporting of test results, low cost,
and potential for self-monitoring [21]. Nonetheless, there are some
doubts regarding its diagnostic accuracy compared to that of ve-
 nous blood sampling, and a recent study has shown that many
current POC HbA1c devices do not meet the analytical require-
ments stipulated by the NGSP [2, 22].

Current HbA1c POC devices have shown good performance in
some recent studies designed to compare the validity of capillary
blood glucose and venous blood glucose testing [10-14]. In our
study, we not only evaluated POC device performance for EDTA
whole blood glucose, but also compared HbA1c levels using EDTA
whole blood and capillary blood as samples determined by the
cobas b 101 assay to evaluate its clinical efficiency.

The NGSP has tightened the criteria several times for the certifi-
cation of manufacturer methods, with the goal of improving the
quality of HbA1c testing. The performance threshold for manufac-
turers is ±6% with respect to the relative bias of the Secondary
Reference Laboratory measurements [23]. Similarly, the CAP re-
placed peer-group grading of the HbA1c level for the GH-2 HbA1c
survey with accuracy-based grading, and has since tightened the
acceptable performance limits from ±15% to ±7% in 2011-2012
and ±6% in 2013-2016 [24, 25]. The performance of POC tests for
the determination of HbA1c levels is generally assessed based on
precision; it is recommended that an imprecision of less than 3%
is a desirable analytical goal for laboratory HbA1c methods based
on clinical requirements, and an optimal imprecision of 2% NGSP
units is now recommended by leading professional groups [2, 26]. In this study, the cobas b 101 analyzer showed good precision at the pathological level according to the desired analytical goal, but did not meet the optimal imprecision goal. At the normal level, the within-laboratory precision did not meet the goal. Therefore, this test was considered acceptable for follow-up monitoring of HbA1c at the 7.0% HbA1c treatment goal recommended by the American Diabetes Association Standards of Medical Care in Diabetes 2017 [27]. Overall comparative analyses showed a strong correlation and all of the mean %differences were within 6%.

In a previous study, the cobas b 101 showed an acceptable imprecision result in an evaluation of seven HbA1c POC devices using patient venous blood samples [14]. Two different reagent lot numbers for the cobas b 101 met the NGSP criteria, with an intra-laboratory precision of 2.4% and 1.2% (CVs) and -0.05%–0.23% bias with three certified secondary reference measurement procedures, indicating good performance compared to that of the other six POC instruments.

Although the cobas b 101 showed promising results, this study had a few limitations. We could not evaluate a broad range of HbA1c levels owing to a lack of patient samples with extremely low or high levels. In the repeatability and within-laboratory precision studies for the cobas b 101 POC device, the 5.1% level for QC materials did not fully represent the diagnostic performance for a diabetes cutoff HbA1c level of 6.5%. We did not exclude interference from hemoglobin variants because we could not obtain fresh specimens with hemoglobinopathies, which have the potential to impact the HbA1c results.

In conclusion, comparative analyses with the reference method using the Variant II Turbo and the POC test using the cobas b 101 showed strong correlations using EDTA whole blood samples in our study. Moreover, our findings demonstrated a strong correlation between HbA1c levels obtained using EDTA whole blood and capillary samples in the cobas b 101 assay. However, exhaustive precision analyses are necessary before clinical use. Therefore, the cobas b 101 analyzer is a convenient assay for HbA1c levels and may be useful for diabetes management.

요 약

배경: 당화혈색소 측정을 위한 현장검사 장비의 사용이 점차 늘고 있으므로 현장검사 장비와 중앙 검사실에서 사용하는 검사법 간의 비교 분석은 매우 중요하다. 본 연구에서 당화혈색소 현장검사 장비인 cobas b 101의 분석능을 중앙검사실 장비와 비교하여 평가하였다.

방법: 반복성, 검사실 내 정밀도, 적정성, 로트 간 재현성을 분석하여 cobas b 101 system의 분석적 질을 평가하였다. EDTA 전혈과 모세혈관에서 채취한 혈전이 사용되었으며 검사는 cobas b 101 system과 Variant II Turbo 장비로 시행되었다.

결과: 정상 범위의 정도관리 물질에서 검사실 내 정밀도는 5.22% 변동계수를 나타내었고, 높은 값의 정도관리 물질에서 2.56% 변동 계수를 나타내었다. 적정성 평가에서 상관계수 0.990으로 좋은 결과를 나타내었다. 서로 다른 로트 간 재현성 분석에서 상관 계수 0.986으로 좋은 상관성을 보였으며 당화혈색소 값 차이의 백분율 평균은 -2.9%였다. Cobas b 101과 Variant II turbo를 이용한 당화혈색소 검사 결과의 비교는 EDTA 전혈을 사용하였을 때 0.958의 상관계수와 0.8%의 차이의 백분율 평균값을 보였으며 모세혈관 전혈을 사용하였을 때 0.976의 상관계수와 2.0%의 차이의 백분율 평균값을 보여 전체적으로 좋은 상관성을 보였다. Cobas b 101을 이용한 EDTA 전혈과 모세혈관 검체 결과의 비교분석 부분에서 상관 계수 0.985로 좋은 상관관계를 나타내었으며 차이의 평균 백분율은 1.3%로 나타났다.

결론: Cobas b 101은 전반적인 성능평가에서 좋은 결과를 나타내어 당뇨 환자의 추적검사를 위한 당화혈색소 측정 장비로서 편리하고 유용하게 쓰일 것으로 기대되나 사용 전주의 값은 정밀도 평가가 선행되어야 할 것이다.

AUTHORS’ DISCLOSURES OF POTENTIAL CONFLICTS OF INTEREST

No potential conflicts of interest relevant to this article were reported.

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