Performance Evaluation and Local International Sensitivity Index Verification Using Automated Coagulation Analyzer Coapresta 2000

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Background: The Coapresta 2000 (CP2000; Seikisui, Japan) system is a fully-automated random-access multiparameter coagulation analyzer equipped with a photo-optical clot detection unit. It can perform clotting time assays as well as colorimetric assays.

Methods: We evaluated the analytical performance of CP2000 for several coagulation test parameters and compared its performance with that of the CA-7000 (Sysmex, Japan) system. Fresh and frozen plasma samples were used to evaluate the performance of CP2000 with respect to four routine coagulation test parameters: prothrombin time (PT), activated partial thromboplastin time, fibrinogen, and D-dimer. On-board stability of the liquid reagents was confirmed. Additionally, local international sensitivity index (ISI) verification was performed with four levels of calibrants and direct PT/international normalized ratio (INR) line.

Results: The intra- and inter-assay coefficients of variation were below 5% for every parameter in both normal and pathological ranges. Carryover was not detected. The results obtained using CP2000 showed good correlation ($r^2$ over 0.95) with those obtained by the CA-7000 analyzer. On-board stability in open-vial state, which was expected to be much longer than that of other reagents, was confirmed. Local verification of ISI showed an acceptable bias range of INR, compared with the values using calibrants.

Conclusions: The high-throughput, CP2000 analyzer is a fast, user-friendly system with long on-board reagent stability. Its results were concordant with the CA-7000 analyzer, for analysis of the routine coagulation test parameters. Furthermore, this system would add greater confidence to the reporting of INR data.

Key Words: Coapresta 2000, Coagulation test, Local international sensitivity index verification, Direct prothrombin time/international normalized ratio line

INTRODUCTION

Blood coagulation tests are used in a variety of ways in the medical field, from screening to confirmation and follow-up of bleeding and thrombotic disorders. Although previously these tests were performed manually, automated machines are now commonly used. Recently, numerous systems have been developed that can simultaneously perform various methods to obtain results, and different types of blood coagulation tests are conveniently carried out in real time. However, obtaining accurate coagulation test results has become increasingly important, and different assay methods and reagents that can alter results can have a significant effect. In addition, with the increase in inspection requirements for laboratory tests, the evaluation of the correlation of existing devices and the perfor-
Performance of new equipment has become essential whenever a new blood coagulation device is introduced.

Coapresta 2000 (CP2000; Sekisui Medical Co., Tokyo, Japan), which has a fully automated and high-throughput system, is a random-access multiparameter coagulation analyzer that is equipped with a photo-optical clot detection unit. It can perform clotting time and colorimetric assays. Efficient space utilization is possible because of its small size, and a large number of specimens can be quickly and accurately processed because of its speed (clotting assay: 400 tests/hr and photometric assay: 200 tests/hr), thus making emergency inspection possible [1,2]. Herein, the performance of the CP2000 system was evaluated, and the results were compared with that of the CA-7000 (Sysmex Co., Tokyo, Japan) system, which is now routinely used in Kyungpook National University Hospital. In addition, for more accurate international normalized ratio (INR) data, the local international sensitivity index (ISI) value was verified using calibrated plasmas.

MATERIALS AND METHODS

1. Instruments and Reagents

Quality control materials in low and high concentrations (Coagpia control set; Sekisui Virotech, Tokyo, Japan) were used to evaluate prothrombin time (PT; including the results in seconds, %, and INR value), activated partial thromboplastin time (aPTT), fibrinogen, fibrin degradation product (FDP), and D-dimer tests using the CP2000 system. For PT analysis, Coagpia PT-N (including thromboplastin of rabbit brain origin, Sekisui Virotech) material was used instead of Thromborel-S (Siemens Healthcare Diagnostics, Marburg, Germany), which is currently used in the CA-7000 system. All tests were performed according to the manufacturer’s instructions.

2. Sample and Storage

Every sample was added to a 3.2% sodium citrate tube (Becton Dickinson Ltd., Franklin Lakes, NJ, USA) at a precise sample:anticoagulant ratio of 9:1 and centrifuged at 1,500 g for 15 minutes to obtain plasma. A random selection of ordered coagulation test specimens, excluding inadequate specimens (insufficient volume, hemolysis, clot), was used for the correlation evaluation, using CA-7000 as the routine testing system. Moreover, random samples containing low or high levels of fibrinogen and D-dimer were used for the carryover evaluation. When the onboard stability of the reagents was examined, pooled plasma samples were frozen at $-70^\circ$ C as aliquots and then used for the evaluation of reagents until the expiry date as proposed by the manufacturer. After testing with CA-7000, samples were immediately run in CP2000. If immediate examination after a routine analysis was not possible, remnant samples were refrigerated ($2^\circ$C-$4^\circ$C) and tested within 4 hours.

3. Evaluation

1) Precision

The results of PT (including the results in seconds, %, and INR value), aPTT, fibrinogen, and D-dimer were calibrated using low and high concentrations of quality control materials for 10 repetitive sessions in one day to assess the intra-assay (within-run) precision. To estimate the inter-assay (between-run) precision for similar test items, two independent sessions were examined per day for 10 days. Every session was tested in duplicate according to appropriate Clinical and Laboratory Standards Institute (CLSI) guidelines [3]. The mean, standard deviation (SD), and coefficient of variation (CV) for each test item were calculated.

2) Correlation with CA-7000

PT, aPTT, fibrinogen, and D-dimer results obtained by CP2000 were compared with those of CA-7000 using the proper number of normal and abnormal patient plasma cells. CA-7000 results were treated as the standard, and the correlation coefficient ($r$) was calculated. The correlation was evaluated based on the CLSI guidelines [3,4].

3) Carryover

High and low fibrinogen and D-dimer patient samples, which were obtained during routine analysis of fibrinogen and D-dimer test carryover rates, were calculated using the following equation: $[L1-(L3-L4)/2]/(H2+H3)/2$.
(L3+L4)/2}×100. The acceptable range of the carryover rate was less than 1.0%.

4) On-board stability of reagents
As much longer sustained stability is claimed by the manufacturer for the reagents used in CP2000 than other coagulation analyzers, the confirmation process was performed during the proposed stable periods of each test. Two-level quality control materials and frozen (−70°C) pooled plasma were tested, and these results were used to evaluate the onboard stability of five reagents, including PT, aPTT, fibrinogen, FDP, and D-dimer, using CP2000. Every sample was tested in duplicate, and the mean, SD, and CV for each test item were calculated.

5) Local international sensitivity index verification
The four-level (one normal level and three abnormal levels: with an INR range of 1.5–4.5) calibrated plasma samples (AK-Calibrant: Technoclone, Vienna, Austria) were used for local ISI verification. The AK-Calibrant is a lyophilized normal plasma pool, which is known to be produced from the selected citrated plasma of at least 100 blood samples of healthy donors. The assigned value of local ISI 1.04 was verified according to the suggested guidelines (5). The INRs of the calibrated plasmas were calculated from local PTs and ISI (measured in triplicate) and compared with the calibrated values of the INR. It is considered appropriate if the difference between the two values is less than 15%. When the differences between the two values were over 15%, the local ISI was calibrated to obtain an accurate INR. After the linear regression of log local PT results that correspond to log certified INR values, a valid calibration curve (r²) was required to be greater than 0.95. A simple and direct PT/INR line, which was recently discovered, does not require a local ISI, and mean normal prothrombin time (MNPT) was used to determine if the bias between the local INR and INR that was derived from the direct line is less than 10% (6).

6) Statistics
Statistical analysis was performed using the Analyse-it Software (Analyse-it Software Ltd., Leeds, UK). The Pearson correlation coefficient (r) of the two devices was calculated.

RESULTS

1. Precision
For every item that was tested, the precision results were significant and within 3% of the CV. The intra-assay precision ranged from 0.56% to 2.08%, while the inter-assay precision ranged from 0.48% to 2.22% (Table 1).

2. Correlation with CA-7000
The results of PT (including the results in seconds, %, and INR value), aPTT, fibrinogen, and D-dimer using CP2000 showed a high and statistically significant correlation with those of CA-7000. The correlation coefficient (r) was higher than 0.95 for every item except aPTT (0.9445) (Table 2, Fig. 1).

Table 1. Intra-assay and inter-assay precisions of the CP2000 analyzer

| Test                      | Coagpia control low | Coagpia control high |
|---------------------------|---------------------|----------------------|
|                           | Mean    | SD     | CV (%)   | Mean    | SD     | CV (%)   |
| Activated partial thromboplastin time (sec) | 28.30/28.45 | 0.22/0.15 | 0.78/0.51 | 56.84/57.10 | 0.50/0.58 | 0.87/1.01 |
| PT (sec)                  | 12.50/12.50 | 0.07/0.06 | 0.56/0.48 | 20.90/20.80 | 0.21/0.24 | 1.02/1.15 |
| PT (%)                    | 91.80/91.60 | 1.31/1.11 | 1.42/1.21 | 33.80/34.00 | 0.54/0.62 | 1.60/1.83 |
| PT (international normalized ratio) | 1.00/1.04 | 0.01/0.01 | 0.67/0.57 | 1.80/1.76 | 0.02/0.02 | 1.08/1.11 |
| Fibrinogen (mg/dL)        | 273.60/268.94 | 5.64/5.48 | 2.06/2.04 | 88.90/89.44 | 1.85/1.98 | 2.08/2.22 |
| D-dimer (µg/mL)           | 3.58/3.67  | 0.04/0.05 | 1.16/1.35 | 12.52/12.50 | 0.08/0.14 | 0.65/1.12 |

Abbreviations: SD, standard deviation; CV, coefficient of variation; PT, prothrombin time.
3. Carryover
Carryover rates that were below 1% (fibrinogen, 0.2%; D-dimer, 0.2%) between high and low concentration samples were acceptable.

4. Onboard Stability of the Reagents
For every item that was tested during the manufacturer-proposed shelf life, the reagents’ onboard stability in the open-vial state was sustained, with every test result falling within 5% of the CV, ranging from 0.50% to 4.53% (Table 3).

5. Local ISI Verification
The differences between routine INR and certified INR were 15% or less (Table 4). The linear regression of log local PT results that correspond to the log certified INR values revealed that $r^2$ was greater than 0.95 (Fig. 2). In addition, the difference of less than 10% between the

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**Table 2. Correlation between CA-7000 and CP2000**

| Test                        | Number | CA-7000 | CP2000 | $r^*$   |
|-----------------------------|--------|---------|--------|---------|
| Activated partial thromboplastin time (sec) | 209    | 35.6    | 40.7   | 0.9445  |
| PT (sec)                    | 367    | 22.6    | 26.6   | 0.9740  |
| PT (%)                      | 367    | 49.4    | 46.0   | 0.9848  |
| PT (international normalized ratio) | 367    | 2.1     | 2.3    | 0.9730  |
| Fibrinogen (mg/dL)          | 95     | 350.1   | 379.0  | 0.9890  |
| D-dimer (µg/mL)             | 169    | 3.1     | 3.8    | 0.9953  |

Values are presented as mean. Abbreviation: PT, prothrombin time. *Correlation coefficient.

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**Fig. 1.** Correlation of the test results between CA-7000 (Sysmex Co., Tokyo, Japan) and CP2000 (Sekisui Medical Co., Tokyo, Japan) for five coagulation test items. (A) Correlation PT (sec); (B) Correlation PT (%); (C) Correlation PT (international normalized ratio); (D) Correlation activated partial thromboplastin time (sec); (E) Correlation fibrinogen; and (F) Correlation D-dimer. Abbreviations: CP2000, Coapresta 2000; PT, prothrombin time; DDU, D-dimer units; FEU, fibrinogen-equivalent units.
calculated INR, which uses direct PT/INR line, and the routine INR, which utilizes ISI, was acceptable (data not shown).

DISCUSSION

Various devices and reagents for blood coagulation tests are used in laboratories. However, blood coagulation tests are highly influenced by analytical instruments and the reagents used. Therefore, when introducing a new blood coagulation system into a laboratory, an evaluation of not only the correlation with existing devices, but also of its function, effectiveness, simplicity, and speed should be conducted [7,8]. Since CP2000 has a fast and easy-to-operate coagulation test system, its use for routine analysis was evaluated.

The intra- and inter-assay precisions of the four routine coagulation test items showed significantly low CV values (below 3%), whereas the proposed allowable CV value was <5% in the CLSI guideline [4]. A good correlation with the existing CA-7000 system was observed, and this result is similar with those of other reports [1,2]. The precision of aPTT can be lower because of more complex testing procedures and different sensitivities to different reagents and testing methods, compared to PT results, as reported by the authors [9]. Thus, the lowest correlation coefficient (r=0.9445) was obtained. However, the results using CP2000 showed a bias over those of CA-7000 (PT, 14.716%; INR value, 7.605%; aPTT, 17.784%), and this result is similar to those of other reports [2]. The sustained onboard stability of the CP2000 reagents, even in the open-vial state, could be an important advantage for practical procedures.

INR values are used to monitor patients who are on warfarin therapy. Thus, an accurate report of the INR value that minimizes the variation effect between different laboratories (by instruments and reagents) is necessary for the exact drug dosage arrangement. INR results are mathematically converted based on the instrument-specific reagent combination, using proposed ISI values from the manufacturer. However, in spite of the fact that INR theoretically eliminates the bias of particular instrument/
reagent combinations, the values still vary [6]. Thus, INR should be verified using calibrated plasmas. Commercially available plasmas with a range of certified INR values can be used to verify local ISI, and if the difference is greater than 15%, ISI should be recalculated for more accurate INR data. Recently, a simple and direct PT/INR method was discovered that is different from the previously used local calibration method, which was more labor intensive and impractical in routine analysis. Moreover, it can be used by laboratory workers with ease and does not require instrument-specific ISI or MNPT [6,10,11]. The direct PT/INR line method acquired almost identical results to conventional calibration in a multicenter analysis [12]. Acceptable differences were all obtained through one normal and three abnormal levels of certified INR values using CP2000 and an instrument-specific reagent. Although further ISI calibration was not necessary, the results could also be verified with calibrated values using the direct PT/INR line by linear regression. Locally calibrated INR values can simply be obtained using the website spreadsheet of the direct PT/INR line [11].

In conclusion, the high-throughput CP 2000 analyzer is a user-friendly system with sustained onboard reagent stability. This analyzer correlated well with CA-7000 for the analysis of routine coagulation test parameters. This instrument is expected to be used successfully in routine coagulation analysis.

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Coapresta 2000을 이용한 성능 평가 및 자체 International Sensitivity Index 검증

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배경: 높은 검체 처리량의 Coapresta 2000 (CP 2000; Sekisui, Japan) 시스템은 광-광학 응고 탐지장치가 장착된 완전 자동 랜덤액세스 다중응고분석기가다. 이 시스템은 비색 분석 및 응고시간 분석을 수행할 수 있다. 본 연구에서 저자들은 몇 가지 응고검사결과에 대한 CP 2000의 분석성능을 평가하고 현재 병원의 일상적인 응고검사에 사용되고 있는 CA–7000 (Sysmex, Japan) 시스템의 성능과 비교하였다. 또한 보다 정확한 international normalized ratio (INR) 결과보고를 위해 인증된 혈장을 사용하여 local international sensitivity index (ISI) 검증을 수행하였다.

방법: Prothrombin time (PT), activated partial thromboplastin time, fibrinogen과 D-dimer 4가지 일상적인 응고검사와 관련하여 신선하거나 동결된 환자 혈청으로 CA–7000과 성능을 비교 평가하였다. 다른 시약에 비해 안정성이 길다고 알려진 액체 시약의 검사성능과 안정성도 확인하였다. 로컬 ISI 검증은 4단계의 AK-Calibrant (Technoclone, Austria)로 수행하였으며, 기존 ISI를 사용한 계산 INR 값과 직접 PT/INR 직선을 이용한 값과 비교하였다.

결과: 분석범위 내 및 분석 간 변동계수는 정상 및 병적 범위 모두에서 모든 검사에 대해 5% 미만의 좋은 정밀도를 보였다. CP 2000을 사용하여 얻은 결과는 CA–7000에서 얻은 결과와 좋은 상관관계(0.96를 초과하는 ²)를 보였다. 장비 내 개봉된 상태에서의 평가된 시약 안정성도 제조사에서 제공한 기간 동안 잘 유지되는 것으로 확인되었다. ISI의 local 검증에서의 인증 헐리안과 차이는 헐리안 가능한 바이어스 범위(15 % 미만)로 나타났다.

결론: CP 2000 자동응고분석기는 좋은 시약 안정성과 용이성을 갖춘 시스템으로, 평가된 응고검사결과는 기존 사용하고 있는 CA–7000과 높은 상관관계를 보여 동상적인 응고검사에 충분히 유용하게 사용될 것으로 생각된다.

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