Analytical validation of therapeutic drug monitoring (TDM) on AxSYM Abbott analyzer

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Abstract

**Introduction:** Careful monitoring of drug concentration (therapeutic drug monitoring, TDM) is essential for a large number of drugs. The aim of this study was to evaluate analytical performance of Abbott AxSYM analyzer for therapeutic drug monitoring of theophylline, carbamazepine, phenobarbitol and valproic acid.

**Materials and methods:** For the purpose of analytical validation following parameters were determined for all analytes: inaccuracy (bias), within-run and between-run imprecision and measurement uncertainty. Additionally, concentration of valproic acid was compared with the previously used analytical system (TDx FLx Abbott analyzer) for 30 patients’ samples.

**Results:** Inaccuracy results (bias) were as follows: for theophylline from −3.66% to −5.84%; for carbamazepine −0.46% to 1.00%; for phenobarbitol −1.83% to −8.08% and for valproic acid from −1.01% to −5.65%. The highest coefficient of variation (CV) for within run imprecision was observed for phenobarbitol (7.07%) and the lowest for theophylline (2.71%). The highest CV for between run imprecision was observed for carbamazepine (4.73%) and the lowest for theophylline (2.94%). The highest measurement uncertainty was observed for phenobarbitol assay (21.7%) and the lowest for carbamazepine (10.7%).

Passing-Bablok regression analysis of valproic acid comparison on two analyzers showed statistically significant, but clinically insignificant deviation in slope of the regression equation (b = 1.121; 95% CI = 1.028–1.197); however the Cusum linearity test proved that there was a linear relationship between two methods.

**Conclusion:** In conclusion, analytical validation fulfilled all previously established criteria and could be implemented in a routine laboratory work.

**Key words:** analytical validation; measurement uncertainty; valproic acid; phenobarbitol; carbamazepine; theophylline

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**Introduction**

Careful monitoring of drug concentration (therapeutic drug monitoring, TDM) is essential for a large number of drugs. Drugs with narrow therapeutic range can easily achieve either too high or too low serum concentration. Elevated drug concentration in the bloodstream can cause side effects with serious and sometimes even fatal consequences. On the contrary, drug concentration below the therapeutic range can lead to inadequate response to therapy.

Valproic acid, phenobarbitol and carbamazepine are antiepileptic drugs (AEDs) that have been used for treatment of seizures in epilepsy and other disorders (1–3). These drugs require constant monitoring to maintain the concentration within the therapeutic range. Although, novel second- (4) and third-generation (5) antiepileptics have now days been prescribed more frequently, these drugs still remain treatment of choice for some types of pediatric epilepsy (6).

Theophylline is a bronchodilator with a narrow therapeutic range and large interindividual variability in pharmacokinetics. High serum concentrations of theophylline are associated with serious adverse reactions. Although its use has been declining in the past two decades, it is still used as an additional treatment drug in the asthmatic acute phase in children (7,8).
In our laboratory, TDM is performed using fluorescence polarization immunoassay (FPIA) technology on TDX FLx Abbott analyzer (Abbott Laboratories, Abbott Park, IL, USA). For methods based on antigen-antibody reaction a large variability between different types/generations of reagents can be observed (9). Our experiences have shown that even interchangeable analytic systems of the same kind of analyzers can show discrepancies in a result and should be carefully monitored and compared (10). Furthermore, as a laboratory accredited according to ISO 15189 standard, we are obliged to perform an analytical validation of each new reagent/analyzer prior to method verification and introduction to routine laboratory work (11).

Aim of this paper was to describe the results of analytical validation of Abbott AxSYM analyzer for therapeutic drug monitoring of valproic acid, phenobarbital, carbamazepine and theophylline.

Materials and methods

Methods

This study was performed at University Department of Chemistry, Sestre milosrdnice University Hospital in November 2009. Concentration of therapeutic drugs theophylline, carbamazepine, phenobarbital and valproic acid was determined on AxSYM analyzer (Abbott Laboratories, Abbott Park, IL, USA) using original manufacturer’s reagents and procedures. For the comparability study, valproic acid was determined on Abbott TDX FLx analyzer (Abbott Laboratories, Abbott Park, IL, USA). All analytes were determined by fluorescence polarization immunoassay (FPIA) technology.

Therapeutic intervals used in our laboratory were as follows: for theophylline 28–83 µmol/L; for carbamazepine 17–42 µmol/L; for phenobarbital 65–172 µmol/L and for valproic acid 347–693 µmol/L.

Analytic evaluation

Inaccuracy

Inaccuracy was determined using three concentration levels of the commercial control samples. Lot numbers for control samples were: 69072Q100 for Theophylline II Control; 76484Q100 for Carbamazepine Control; 69641Q100 for Phenobarbital Control and 76480Q100 Valproic Acid Control (Abbott Laboratories, Abbott Park, IL, USA). Each sample was analyzed three times; average result was compared to manufacturers declared value and the bias was calculated. Acceptance criteria for each analyte were established according to the CLIA Proficiency Testing criteria (12).

Within run imprecision (repeatability)

Repeatability was determined using sera of two patients for each analyte (samples A and B). Each sample was analyzed ten times in a batch analysis. Coefficient of variation (CV) was calculated and compared to the acceptance criteria declared by the manufacturer for appropriate concentration levels, as well as to criteria proposed by Guidelines of the German Federal Medical Council (13).

Between run imprecision (intermediate precision)

Intermediate precision was determined using pool of three patients’ samples for each analyte. These samples were remaining sera from the routine drug monitoring analysis. Samples were mixed and aliquoted in ten separate tubes and frozen at –20 °C. During the 15-days period, once a day at different time of the routine work one sample was thawed and analyzed. Coefficient of variation was calculated and compared to the acceptance criteria declared by the manufacturer, as well as to criteria proposed by Guidelines of the German Federal Medical Council (13).

Measurement uncertainty

Measurement uncertainty (U) was calculated for all tests according to the formula (14):

$$U = 2 \times \sqrt{u^2_{\text{calibrator}} + u^2_{\text{imprecision}} + u^2_{\text{bias}} + u^2_{\text{bias inaccuracy}}}$$

Where:

- \(u_{\text{calibrator}}\) is a calibrator measurement uncertainty as declared by the manufacturer.
- \(u_{\text{imprecision}}\) is an uncertainty caused by imprecision. In our case, coefficient of variation for between run imprecision was used.
$U_{\text{bias}}$ is an uncertainty caused by inaccuracy. In our case, bias of the commercial control sample with the value closest to the sample for the between run imprecision was used.

$U_{\text{bias inaccuracy}}$ is an uncertainty caused by inaccuracy when calculating bias. It is calculated using following formulas:

$$u_x = \frac{SD}{\sqrt{n}}$$

$$U_{\text{bias inaccuracy}} = \frac{ux}{X} \times 100$$

**Method comparison**

Method comparison was done using remaining sera for the patients referred to laboratory for valproic acid determination. For 30 patients, valproic acid was determined on the Abbott TDx FLX analyzer as a part of a routine workup and than immediately analyzed at AxSYM Abbott analyzer.

**Statistical analysis**

Data on the commercial control samples are presented as arithmetic mean and the bias to the manufacturers’ declared value was calculated. For the imprecision analysis, minimum, maximum, arithmetic mean, standard deviation and coefficient of variation were calculated. For the comparison analysis, valproic acid concentration is presented as mean ± standard deviation and Passing-Bablock regression analysis (along with Cusum linearity test) was performed. Statistical analysis was performed with MedCalc® software (MedCalc 10.1.3.1., Frank Schoonjans, Mariakerke, Belgium).

**Results**

Inaccuracy for all analytes was in accordance with the previously established criteria (Table 1). The highest bias was observed for the Phenobarbital Medium Control (−8.08%), but it was also within the acceptance criteria.

Data on within run imprecision (repeatability) are presented in Table 2. The highest coefficient of variation was observed for the phenobarbital assay (7.07%) and the lowest for the theophylline (2.71%). Table 3 shows the data on between run imprecision (intermediate precision). The highest coefficient of variation was observed for the carbamazepine (4.73%) and the lowest for the theophylline (2.94%). None of the obtained coefficients of variation met the criteria declared by the manufacturer. However, all values fulfilled the requirements as proposed by the German Federal Medical Council (13).

**Table 1.** Results of inaccuracy analysis.

<table>
<thead>
<tr>
<th>Commercial control sample</th>
<th>Declared value (μmol/L)</th>
<th>Mean measured value (μmol/L)</th>
<th>Bias</th>
<th>Acceptance criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Theophylline II Control</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td>38.85</td>
<td>37.00</td>
<td>−4.75%</td>
<td>± 25%</td>
</tr>
<tr>
<td>Medium</td>
<td>66.60</td>
<td>62.71</td>
<td>−5.84%</td>
<td></td>
</tr>
<tr>
<td>High</td>
<td>144.30</td>
<td>139.02</td>
<td>−3.66%</td>
<td></td>
</tr>
<tr>
<td>Carbamazepine Control</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td>12.69</td>
<td>12.82</td>
<td>1.00%</td>
<td>± 25%</td>
</tr>
<tr>
<td>Medium</td>
<td>25.38</td>
<td>25.23</td>
<td>−0.58%</td>
<td></td>
</tr>
<tr>
<td>High</td>
<td>67.68</td>
<td>67.37</td>
<td>−0.46%</td>
<td></td>
</tr>
<tr>
<td>Phenobarbital Control</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td>64.65</td>
<td>62.45</td>
<td>−3.40%</td>
<td>± 20%</td>
</tr>
<tr>
<td>Medium</td>
<td>129.30</td>
<td>118.85</td>
<td>−8.08%</td>
<td></td>
</tr>
<tr>
<td>High</td>
<td>215.50</td>
<td>211.57</td>
<td>−1.83%</td>
<td></td>
</tr>
<tr>
<td>Valproic Acid Control</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td>259.88</td>
<td>245.19</td>
<td>−5.65%</td>
<td>± 25%</td>
</tr>
<tr>
<td>Medium</td>
<td>519.75</td>
<td>493.33</td>
<td>−5.08%</td>
<td></td>
</tr>
<tr>
<td>High</td>
<td>866.25</td>
<td>857.53</td>
<td>−1.01%</td>
<td></td>
</tr>
</tbody>
</table>

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 Measurement uncertainty was calculated for all analytes according to the previously described formula (14) (Table 4). The highest uncertainty was observed for phenobarbital (21.7%), which was expected, since the highest inaccuracy and coefficient of variation were also observed for the same analyte.
Comparison of valproic acid concentration determination on two analyzers is presented in table 5 and figure 1. Passing-Bablock regression analysis showed statistically significant, but clinically insignificant deviation in slope of the regression equation \(b = 1.121; 95\% \text{ CI} = 1.028–1.197\); however the Cusum linearity test proved that there was a linear relationship between two methods.

**Discussion**

The aim of this study was to evaluate TDM assays on Abbott AxSYM analyzer. Overall, all obtained parameters were satisfactory and fulfilled previously established criteria.

Inaccuracy study showed the highest bias for phenobarbital assay \((-8.08\%)\), for the medium control concentration level of 129.30 \((99.56–159.04) \mu mol/L\), surprisingly, since one could expect the highest error for the highest concentration values. However, this concentration range is within the reference range of phenobarbital monitoring \((65–172 \mu mol/L)\) and not near the critical value that is associated with the increased risk of side effects \((> 220 \mu mol/L)\). Due to that, we believe that inaccuracy couldn’t have high clinical impact. Nevertheless, the proposed acceptance criteria for toxicology TDM by CLIA Proficiency Testing criteria are much higher than those obtained in our evaluation \((12)\).

Imprecision analysis showed similar CVs in within run and between run studies. Moreover, for some assays (valproic acid and phenobarbital) CVs obtained in between run analysis were even smaller than those in the batch analysis, suggesting a great stability of performance for these assays. However, neither of obtained coefficients of variation fulfilled the criteria established by the manufacturer. It could be expected, since the manufacturer defined those CVs using commercial control samples as the material of the analysis. In our study, we

**Table 5.** Passing-Bablock regression analysis.

<table>
<thead>
<tr>
<th>Valproic Acid TDx (μmol/L)</th>
<th>Valproic Acid AxSYM (μmol/L)</th>
<th>regression equation</th>
<th>95% confidence interval</th>
<th>P (Cusum linearity test)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N=30</td>
<td></td>
<td>(y = -25.161 + 1.121x)</td>
<td>(-48.318–1.535)</td>
<td>(1.028–1.197)</td>
</tr>
</tbody>
</table>

![Figure 1](image_url). Comparison of valproic acid concentration determination on two analyzers (Abbott TDx and AxSYM).
wanted to simulate the conditions of routine laboratory work and we have determined our imprecision using patients’ samples. Our obtained CVs also fulfilled acceptance criteria established by the German Federal Medical Council (13).

We have decided to expand our analytical validation by presenting results of measurement uncertainty. Measurement uncertainty encompasses all components that can influence on the result: systematic and random error parameters of the assay and errors obtained by uncertainty of calibrators and it’s estimate (15). Measurement uncertainty is still not fully implemented in a standard laboratory evaluations, since one could be puzzled by “high” numbers usually obtained in this analysis. Since the measurement uncertainty is a property of an each measurement result, we are obliged to give an estimate of range where our result is positioned in order to ensure to the users of our analysis a true interpretation of obtained value. Measurement uncertainties for our validation were also satisfactory (~10–20%), since we could expect these values for immunochemistry analysis (16).

Since all tested analytes utilize the same technology (FPIA), we have decided to make the comparison between two analytic systems only for one representative analyte – valproic acid. Comparability study also showed a satisfactory concordance between valproic acid determinations on two analyzers. When we have to replace an analytic system in a routine laboratory practice, it is very important not to have a constant deviation from the result. In that case, a reference range should be modified in order to maintain a correct interpretation of laboratory result and ensure proper TDM for the patient. The small deviation in slope of the regression line that we have observed in our analysis was in the high concentration range (Figure 1) and that small difference has no clinical implication on interpretation of the result. Our conclusions should be confirmed on larger number of samples, since the low number of valproic acid determinations used in our comparability study is a limitation to our work.

In conclusion, our analytical validation fulfilled all previously established criteria and could be implemented in a routine laboratory work. We could replace our existing system with a new analyzer without having influence on the result of therapeutic drug monitoring for patients.

Acknowledgments

The authors wish to thank Abbott Laboratories Ltd., Croatia for providing reagents for this evaluation.

References

Analitička validacija terapijskog praćenja koncentracije lijeka na analizatoru AxSYM tvrtke Abbott

Sažetak

Uvod: Pažljivo terapijsko praćenje koncentracije lijeka (engl. therapeutic drug monitoring, TDM) od ključne je važnosti za velik broj lijekova. Cilj ovog istraživanja bio je odrediti analitičke značajke analizatora AxSYM tvrtke Abbott u terapijskom praćenju koncentracije teofilina, karbamazepina, fenobarbitala i valproične kiseline.

Materijali i metode: U svrhu analitičke validacije za sve analite određeni su sljedeći parametri: netočnost (engl. bias), nepreciznost u seriji i iz dana u dan (engl. within-run and between-run imprecision) i mjerna nesigurnost (engl. measurement uncertainty). Dodatno je za 30 uzoraka načinjena usporedba određivanja koncentracije valproične kiseline s prethodno korištenim sustavom (analizator TDx FLx tvrtke Abbott).

Rezultati: Rezultati za netočnost (pogreška) bili su za teofilin od −3,66% do −5,84%; za karbamazepin −0,46%−1,00%; za fenobarbital −1,83%−(−8,08%) i za valproičnu kiselinu od −1,01% do −5,65%. Najviši koeficijent varijacije (engl. coefficient of variation, CV) za nepreciznost unutar serije dobiven je za fenobarbital (7,07%), a najniži za teofilin (2,71%). Najviši CV za nepreciznost iz serije u seriju izračunat je za karbamazepin (4,73%), a najniži za teofilin (2,94%). Najviša mjerna nesigurnost dobivena je za fenobarbital (21,7%), a najniža za karbamazepin (10,7%). Rezultat uspoređene rezultata koncentracije valproične kiseline na dva analizatora regresijskom analizom po Passing-Babloku pokazao je statistički značajno, ali klinički neznatno odstupanje u nagibu regresijske jednadžbe (b = 1,121; 95% CI = 1,028–1,197); međutim Cusum test linearnosti pokazao je da postoji linearna povezanost između dviju metoda.

Zaključak: Možemo zaključiti da je analitička validacija ispunila sve prethodno utvrđene kriterije te da se analizator može uključiti u rutinski rad laboratorija.

Ključne riječi: analitička validacija; mjerna nesigurnost; valproična kiselina; fenobarbital; karbamazepin; teofilin