Gaps in the Traceability Chain of Human Growth Hormone Measurements

Sébastien Boulo, Katja Hanisch, Martin Bidlingmaier, Cristian-Gabriel Arsene, Mauro Panteghini, Guy Auclair, Catharine Sturgeon, Heinz Schimmel, and Ingrid Zegers*

BACKGROUND: Human growth hormone (hGH) is measured for the diagnosis of secretion disorders. These measurements fall under the EU Directive 98/79/EC on in vitro diagnostic medical devices requiring traceability of commercial calibrator values to higher-order reference materials or procedures (Off J Eur Communities 1998 Dec 7;L 331:1–37). External quality assessment schemes show large discrepancies between results from different methods, even though most methods provide results traceable to the recommended International Standard (IS 98/574). The aim of this study was to investigate possible causes for these discrepancies.

METHODS: We investigated the commutability and recovery of hGH in reconstituted IS 98/574. We tested different reconstitution protocols and used 4 different serum matrices for spiking. These IS preparations were measured together with serum samples. We quantified hGH by 5 different methods in 4 different laboratories.

RESULTS: Results from the different methods correlated well for the serum samples. Mean discrepancies between results from different methods were ≤20%. None of the IS preparations was commutable for all the method comparisons. The recovery of hGH in preparations of IS 98/574 depended on the reconstitution protocol (>10-fold differences) and background matrix (relative differences ≤17% for different serum matrices).

CONCLUSIONS: The use of different protocols for reconstitution and spiking of hGH reference preparations affects quantification by immunoassays, potentially leading to a bias between commercial methods, despite the use of calibrators with values claimed to be traceable to the same higher-order reference material. © 2013 American Association for Clinical Chemistry

Human growth hormone (hGH) is a protein secreted by the pituitary gland. It regulates the increase in height during childhood and has other anabolic effects. An excess of hGH leads to acromegaly, whereas a deficiency causes growth failure and short stature. The biochemical diagnosis of GH deficiency is based on peak GH concentrations obtained during different stimulation tests. A nadir value after oral glucose load (<1 μg/L) is commonly used as decision limit to diagnose acromegaly, although this decision limit has been criticized for being too high (1–3).

Circulating hGH shows large heterogeneity due to alternative splicing, different posttranslational modifications, oligomerization, complex formation, and proteolytic processing. The 22-kDa hGH form represents ≤90% of total circulating hGH (4, 5). The 20-kDa hGH (approximately 5%–10% of total circulating hGH) lacks residues 32–46 of the 22-kDa hGH form. These 2 forms are co-secreted in a fixed ratio (6–8). Dimers represent ≤10% of circulating hGH. Fragments of hGH with 16 and 17 kDa and other forms are also present (approximately 2%) (9).

In vitro diagnostic (IVD) measurements of hGH are within the scope of EU Directive 98/79/EC on IVDs, which requires traceability of values assigned to commercial calibrators to higher-order reference materials or methods (10). Calibration of hGH immunoassays in terms of International Standard (IS) 98/574 has been recommended by the International Collaborative on Standardisation of Growth Hormone methods.
Commutability of hGH Calibrators

Materials and Methods

hGH

The hGH material used was the second WHO IS 98/574, containing the lyophilized 22-kDa form of hGH (somatropin) plus excipients (16).

SAMPLES

The serum matrices used in this study were hGH-depleted serum (ref. SF220-2, SCIPAC), human charcoal-stripped serum (ref. IPLA-SER7, Dunn Labortechnik), NIST Standard Reference Material (SRM) 971 male serum, and sheep serum (ref. S2263, Sigma-Aldrich). The density of each of these sera and of the different buffers used in this study were all measured by the American Society for Testing and Materials (ASTM) D 4052 method at 20 °C (17).

We selected serum samples from a large number of sets of 0.5-mL samples, stored at −70 °C. These sera were obtained from blood donations from healthy people (18). Selected serum samples had hGH concentrations between 0.1 and 12 μg/L (mean values of results from the 5 immunoassays used).

PREPARATION OF MATERIALS SPIKED WITH IS 98/574

We designed the protocols for reconstitution, dilution, and spiking while keeping in mind practices used by IVD manufacturers to produce calibrators and the UK National External Quality Assessment Service (NEQAS) for preparing samples for distribution in EQA schemes. We performed reconstitution in different diluents to assess the importance of the buffer and the protein background separately. The final matrices used for the preparation of calibrators reflect the choices available for materials that are free (or almost free) of human GH but have a matrix comparable to that of human serum. Sheep serum is used to avoid interference by (human) GH binding protein (GHB).

For all studies, we gave randomly assigned numbers to serum lots from individual patients and spiked materials. Serum samples were thawed and combined in sets with spiked samples on the day of shipment. Serum samples and spiked samples were filled in identical vials. In combination with the use of randomized numbers, this protocol ensured that all samples were treated equally during shipment and by the participating laboratories. Samples transport took <24 h and included cooling elements to ensure that the temperature remained at 2 °C–8 °C. Samples were stored at 2 °C–8 °C at each of the laboratories. hGH in serum has been shown to be stable at 4 °C for 8 days; therefore we requested that measurements be performed within 5 days of receipt of the samples.

Commutability study. For the commutability study, we made 13 preparations volumetrically to achieve a hGH concentration, after spiking, of approximately 7 μg/L (Fig. 1). IS 98/574 was first reconstituted with MilliQ water, PBS (137 mmol/L NaCl, 2.7 mmol/L KCl, and 10 mmol/L phosphate buffer solution at pH 7.4), or PBS containing 5% BSA (PBS-BSA). After reconstitution, we made several serial dilutions of the respective solutions (3 times by a factor of 100), before the final spiking of the hGH solution into the different sera at 7 μg/L (Fig. 1). We divided these preparations into 0.5-mL aliquots. All preparations were performed on ice.

Recovery study. For the recovery study, we made 10 preparations gravimetrically. IS 98/574 was reconstituted with PBS or PBS-BSA. After reconstitution, we performed 2 serial dilutions (1 by a factor of 250 and 1 by a factor of 40), spiked hGH into the different serum matrices at 5 μg/L, and then divided the samples into aliquots. To accurately determine the hGH mass fraction of each preparation, we measured the density of sera and reconstitution buffers. The results were 1.0052, 1.0174, 1.0214, 1.0196, 1.0249, and 1.0237 kg/L, respectively, for PBS, PBS-BSA, hGH-depleted serum, human charcoal-stripped serum, SRM 971...
male serum, and sheep serum (SD < 0.07 g/L for all measurements).

**hGH MEASUREMENTS**
We provided participating laboratories with detailed protocols. All measurements were performed in triplicate within 1 analytical run. The methods used were Immulite 2000 (Siemens Healthcare Diagnostics) at the Universitair Ziekenhuis Gent (Belgium); Cobas e411 (Roche Diagnostics) at the CIRME, University of Milan (Italy); Liaison (DiaSorin) and iSYS (IDS) at the Klinik der Universität München (Germany); and hGH-sensitive ELISA (Mediagnost) at Institute for Reference Materials and Measurements (IRMM) (Belgium). All the samples were measured once in 1 sequence, then measured in reverse order in a second sequence, and finally measured again as in the first sequence.

**DATA ANALYSIS**
We analyzed the equivalence of results from different methods by performing, for each pair of methods, a Passing-Bablok regression and determining the Pearson correlation coefficient with Analyse-it software. We assessed commutability by Deming regression on the measurement results obtained for the serum samples and verified whether the results for the hGH preparations were within the 95% prediction interval with Excel and R software (19).

**LC-MS MEASUREMENTS**
We performed LC-MS measurements for 3 serum samples according to the method described by Arsene et al. (20). With this isotope-dilution mass spectrometry method, hGH tryptic cleavage products YSFLQNPQTSLCFSEIPTPSNR (T6) and LEDGSPR (T12) were quantified by LC-MS. T12 is present in both the 22- and 20-kDa forms of the growth hormone, whereas T6 is present only in the 22-kDa form. Samples were measured in duplicate. Measurements were performed using either labeled peptides or the fully [15N]-labeled 22-kDa form of growth hormone as internal standard. Results were obtained in micrograms per gram and converted to micrograms per liter using a serum density of 1.02 kg/L, the density of the sera used in this study.

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**Fig. 1. Overview of the volumetric preparations.**
Each hGH ampoule (WHO standard IS 98/574) was reconstituted in water, PBS, or PBS-BSA. Each reconstitution was then diluted into the solution used for reconstitution and subsequently spiked at about 7 µg/L into the different matrices and buffers (hGH-depleted serum, human charcoal-stripped serum, NIST SRM 971 male serum, sheep serum, PBS, and PBS-BSA). Five control matrices were also tested.
IMPACT OF GHBP
GHBP purified from natural sources (ProSpec) was spiked into the samples to produce final concentrations of 0, 10, 75, 250, and 500 μg/L. We then performed hGH measurements in a single analytical sequence, in triplicate, with the hGH-sensitive ELISA (Mediagnost). GHBP content was also measured in both serum samples and hGH preparations using the Active® Human Growth Hormone Binding Protein ELISA (DLS-10-48100, Diagnostic Systems Laboratories). The measurements were performed in 1 single run, in triplicate.

Results

COMMUTABILITY STUDY
For the commutability study, 13 preparations were measured together with 29 serum samples and appropriate control samples (buffer or serum), in 1 analytical run.

For those control preparations with hGH concentrations >0.05 μg/L, the value was corrected by subtraction: specifically, for the NIST SRM 971 material, where the measured hGH concentration was 0.24 μg/L, and for the human charcoal-stripped serum, where the measured hGH concentration was 0.07 μg/L (mean values from all measurements with all methods). The mean value for the control sample, calculated per method, was subtracted from each value measured for a spiked material with that method. An example of the measurement results is presented in Fig. 2A. The data for all the different methods are shown in the Data Supplement, which accompanies the online version of this article at http://www.clinchem.org/content/vol59/issue7.

**Method comparisons.** All methods showed intraassay CVs <6.0% for the serum samples (Table 1), with the Roche method having the lowest mean CV of 1.7%. For each pair of methods, the correlation between the results (mean value calculated per serum sample) was calculated per method, was subtracted from each value measured for a spiked material with that method. An example of the measurement results is presented in Fig. 2A. The data for all the different methods are shown in the Data Supplement, which accompanies the online version of this article at http://www.clinchem.org/content/vol59/issue7.

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sample) was evaluated. The Pearson correlation coefficients were 0.99 or higher, indicating very good correlation of results for individual patient samples despite potential differences in the selectivity of the methods. The slopes of the correlation lines were between 0.78 and 1.13, indicating that a large bias existed between the results from the different methods. The intercepts were either small (<0.2 μg/L) or insignificant.

**Commutability.** For the 5 methods, the different hGH preparations behaved in a manner that seemed similar to those of the serum samples. However, when commutability was assessed according to CLSI guide C53-A, comparing the reference material preparation with the 95% prediction interval for the serum samples (15), none of the preparations was commutable for all comparisons of methods (see online Supplemental Fig. 1, comparisons A to J). Preparations with low measured hGH concentrations (those reconstituted with water or PBS and spiked into hGH-depleted serum) were more often found to be commutable. Comparing the 10 different pairs of experiments, the preparations in which hGH was reconstituted with PBS-BSA and then spiked into hGH-depleted serum (see online Supplemental Fig. 1) showed the best commutability. This preparation was commutable for 8 of the 10 combinations of methods.

**RECOVERY STUDY**

To investigate the recovery of hGH in preparations of IS 98/574, all the reconstitution and spiking steps were carefully controlled, and the amount of hGH in each preparation was determined with a low relative standard uncertainty (<0.07%). IS 98/574 was reconstituted with PBS or PBS-BSA and was spiked into the different matrices. The calculated final hGH concentrations of the gravimetric preparations were all close to the intended value, between 4.91 and 5.18 μg/L (Table 2).

The 10 preparations were measured together with 9 serum samples and 6 control samples in 1 analytical run. hGH concentrations determined in the controls (mean value 0.25 μg/L for NIST SRM 971 and 0.08 μg/L for human charcoal-stripped serum) were subtracted from the values of the respective preparations.

The results from the recovery study confirmed that there were large differences in recovery of hGH in different preparations (Fig. 2B). Preparations of IS 98/574 with PBS for reconstitution gave hGH values between 1 and 2 μg/L (Fig. 2B; see online Supplemental Fig. 2). In preparations where reconstitution had been performed with PBS-BSA, the hGH concentrations measured were between 4.5 and 7 μg/L, depending on the method and the background matrix into which the hGH was finally spiked (Fig. 2B). hGH reconstituted with PBS-BSA and spiked into the same buffer gave a high value, well above the actual concentration of approximately 5 μg/L.

The effect of using different background sera (hGH-depleted serum, charcoal-stripped serum, SRM 971, or sheep serum) was smaller. Differences between hGH concentrations measured in different back-

### Table 2. Results from the hGH recovery study.

<table>
<thead>
<tr>
<th>Background matrix</th>
<th>Recovery value, %</th>
<th>hGH, μg/L</th>
<th>Immulite</th>
<th>Cobas e411</th>
<th>hGH-sensitive ELISA</th>
<th>DiaSorin</th>
<th>IDS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reconstitution in PBS</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PBS</td>
<td>5.05</td>
<td>&lt;1</td>
<td>0.60</td>
<td>&lt;1</td>
<td>&lt;1</td>
<td>&lt;1</td>
<td></td>
</tr>
<tr>
<td>hGH-depleted serum</td>
<td>5.18</td>
<td>38</td>
<td>38</td>
<td>32</td>
<td>28</td>
<td>31</td>
<td></td>
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<tr>
<td>Charcoal-stripped serum</td>
<td>5.09</td>
<td>28</td>
<td>28</td>
<td>23</td>
<td>21</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td>SRM 971 male</td>
<td>5.03</td>
<td>25</td>
<td>22</td>
<td>17</td>
<td>16</td>
<td>20</td>
<td></td>
</tr>
<tr>
<td>Sheep serum</td>
<td>5.12</td>
<td>28</td>
<td>24</td>
<td>20</td>
<td>15</td>
<td>18</td>
<td></td>
</tr>
<tr>
<td>Reconstitution in PBS-BSA</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PBS-BSA</td>
<td>4.94</td>
<td>145</td>
<td>135</td>
<td>103</td>
<td>89</td>
<td>106</td>
<td></td>
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<tr>
<td>hGH-depleted serum</td>
<td>5.01</td>
<td>107</td>
<td>98</td>
<td>88</td>
<td>81</td>
<td>89</td>
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<tr>
<td>Charcoal-stripped serum</td>
<td>5.03</td>
<td>103</td>
<td>94</td>
<td>81</td>
<td>82</td>
<td>88</td>
<td></td>
</tr>
<tr>
<td>SRM 971 male</td>
<td>4.91</td>
<td>106</td>
<td>90</td>
<td>83</td>
<td>80</td>
<td>92</td>
<td></td>
</tr>
<tr>
<td>Sheep serum</td>
<td>4.97</td>
<td>122</td>
<td>100</td>
<td>90</td>
<td>77</td>
<td>90</td>
<td></td>
</tr>
</tbody>
</table>

*Comparison of different immunoassay measurement values to the theoretical hGH content (gravimetrically controlled). Data are the mean recovery values of triplicate measurements for each of the 5 methods.*
grounds ranged from 2% to 17%, depending on the method. However, these differences were of the same order of magnitude as the discrepancies found between results from different methods. The different recoveries in different serum backgrounds could therefore contribute substantially to the observed between-method bias.

**MASS SPECTROMETRY MEASUREMENTS**

For 3 serum samples, hGH concentrations were measured by LC-MS. For each of these samples, the means and SDs were calculated from 2 measurements with labeled peptides and 2 measurements with labeled hGH. The SDs of the measurement results were very low (Table 3). From these data, the calculated mass fraction of hGH in the 20-kDa form was 12.4%, 14.4%, and 15.8% for samples A, B, and C, respectively. The comparison with immunoassay results for the same samples showed that the mean ratio between LC-MS and immunoassay results, calculated over the different methods, was 0.88 for sample A, 0.83 for sample B, and 1.27 for sample C. There was no constant ratio between immunoassay and LC-MS results.

**GHBP MEASUREMENTS**

In plasma, 50% of hGH molecules are expected to be bound to GHBP (9). GHBP may also have an impact on the detection of hGH by immunoassay, whereas for the LC-MS method it has been shown that recombinant GHBP does not interfere with measurements (20). Different concentrations of GHBP were therefore spiked into serum samples and other preparations to investigate the effect of the presence of GHBP. The presence of larger concentrations of GHBP reduced hGH values as measured by immunoassay by up to 38%, 33%, and 31% in serum samples A, B, and C, respectively. For preparations containing IS 98/574, the measured hGH concentration was reduced by 46% in the preparation made from the IS reconstituted in PBS-BSA and spiked into PBS-BSA, and by 44% in the preparation where the IS was spiked into hGH-depleted serum (see online Supplemental Fig. 3).

**Discussion**

The present study addressed the traceability of results from hGH measurements performed with commercial immunoassays. Measurement results are made traceable to a higher-order reference material or reference method with the aim of obtaining equivalent results. However, hGH values measured with different routine methods differ significantly (14), despite the fact that the values assigned to the calibrators (true for all 5 methods used in this study) are stated to be traceable to IS 98/574. Possible causes for the lack of harmonization were investigated in this study, as well as possible solutions.

Differences in selectivity of methods intended to measure the same quantity are common in clinical chemistry, because different methods use different reagents (at least) for the same target analyte. Antibodies

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**Table 3. Comparison of LC-MS and immunoassay techniques with respect to hGH quantification.**

<table>
<thead>
<tr>
<th>Sample</th>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>[hGH] by LC-MS, μg/L</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Peptide T12</td>
<td>7.250 (0.023)</td>
<td>5.205 (0.010)</td>
<td>4.365 (0.019)</td>
</tr>
<tr>
<td>Peptide T6</td>
<td>6.348 (0.005)</td>
<td>4.458 (0.017)</td>
<td>3.675 (0.012)</td>
</tr>
<tr>
<td>[hGH]LC-MS T12/[hGH]IA</td>
<td>0.88</td>
<td>0.83</td>
<td>1.27</td>
</tr>
<tr>
<td>[GHBP], μg/L</td>
<td>42.2 (1.8)</td>
<td>21.2 (1.6)</td>
<td>24.7 (1.4)</td>
</tr>
<tr>
<td>Decrease in [hGH] upon addition of 10 μg/L GHBP, %b</td>
<td>9.7</td>
<td>10.6</td>
<td>14.8</td>
</tr>
<tr>
<td>[hGH] by immunoassay, μg/L</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Siemens</td>
<td>9.78 (0.31)</td>
<td>7.14 (0.44)</td>
<td>4.21 (0.18)</td>
</tr>
<tr>
<td>Roche</td>
<td>9.29 (0.21)</td>
<td>7.26 (0.12)</td>
<td>4.10 (0.11)</td>
</tr>
<tr>
<td>Medigast</td>
<td>7.91 (0.17)</td>
<td>5.62 (0.06)</td>
<td>3.14 (0.07)</td>
</tr>
<tr>
<td>DiaSorin</td>
<td>6.90 (0.87)</td>
<td>5.60 (0.30)</td>
<td>2.87 (0.15)</td>
</tr>
<tr>
<td>IDS</td>
<td>7.99 (0.76)</td>
<td>6.18 (0.25)</td>
<td>3.25 (0.13)</td>
</tr>
</tbody>
</table>

a Data are means (SD) unless noted otherwise. Data are the average values of quadruplicate measurements from LC-MS and triplicate measurements for each of the 5 methods.

b As measured by hGH-sensitive ELISA.
can be monoclonal or polyclonal, derived from different species (e.g., goat, mouse, chicken), and raised against different targets (e.g., recombinant hGH, a mixture of isoforms, a single peptide). Our results showed that for the 5 methods evaluated here, the correlation of results for the serum samples was very good (correlation coefficients >0.99). This means that either these methods had no major differences in selectivity or they had different selectivities for analytes present in a constant ratio in the samples. The latter is more likely, because substantial differences in isoform recognition by the different antibodies used have been described (21).

Because the results from the different methods correlated, it should be possible to reduce discrepancies in measurement results by calibrating all methods with a common commutable calibrator or by data adjustment. Muller et al. (22) showed that the between-method CV for hGH measurements could be reduced from 24.3% to <14% by data adjustment. The mean between-method CV for the serum samples in our study was 17.7%. When calculating mean correction factors per method (by averaging correction factors calculated for individual samples) and applying this to the results for the serum samples, the between-method CV was reduced to 7.5%, comparable to what was found in a study on the potential use of a common harmonization sample (23, 24).

The most common currently available calibration hierarchy, or traceability chain, for hGH measurements is shown in Fig. 3. Clinical measurements are performed in a complex matrix, on a mixture of hGH forms. IS 98/574, referred to in traceability statements, is several steps up in the traceability chain and consists of a purified single isoform without matrix. There are different problems in linking the IS to the clinical measurement results. First, the lack of a common reconstitution and spiking protocol leads to a situation where manufacturers may be using calibrators that are de facto different and produce uncontrolled bias. For example, the use of different serum backgrounds for spiking resulted in values for hGH that differed by ≲17%.

Second, none of the calibrator solutions produced from IS 98/574 according to the protocols presented here were commutable for all the combinations of methods. The distance of the results for the reconstituted IS 98/574 from the regression line for the serum samples provided a measure of the bias caused by a lack of commutability. This bias was of the same order of magnitude as the between-method bias. Therefore, it would not be sufficient to use a common reconstitution and spiking protocol for the production of a master calibrator from IS 98/574. Optimal harmonization requires that such a protocol ensures a commutable matrix preparation.

Third, since different protocols lead to different recoveries of hGH, the value obtained with an immunoassay would be protocol dependent. This could eventually be solved by use of an orthogonal method for measuring hGH, such as LC-MS. A reconstitution and spiking protocol yielding immunoassay and LC-MS results having the same ratio for the calibrator and serum samples could be selected. This would anchor the immunoassay results to the hGH concentration as determined by LC-MS, which should be independent of matrix used. The first results showed, however, that the ratio of hGH as measured by LC-MS and immunoassay was significantly different for the different samples. This indicates that a better understanding of the influence of quantities and parameters of immunoassays and LC-MS measurements—i.e., of what we are measuring—is required before LC-MS measurements can be used for anchoring the immunoassay results. A larger-scale study of samples measured both by LC-MS and immunoassay should provide more information.

One of the main influencing factors for hGH immunoassays is the presence of GHBP (5). Different serum samples have different GHBP concentrations, and different GHBP concentrations in samples change hGH values as measured by immunoassay (see online Supplemental Fig. 3). This is true not only for the method used in our study but also for other methods (25). GHBP was measured in the 3 serum samples to determine whether this could explain the large ratio of [hGH]LC-MS/[hGH]IA for sample C. However, there was no simple correlation between immunoassay values, LC-MS values, and GHBP concentrations, since the GHBP concentrations were comparable in samples B and C.
In conclusion, the commutability and recovery studies reported here show that to standardize hGH measurements, it is not sufficient to recommend traceability to a higher-order reference material such as IS 98/574. Results from different immunoassays correlate well for serum samples, so it is possible in principle to systematically improve the equivalence of hGH measurement results. It has been proposed that correction factors be used to achieve this goal, at least for the present. However, this approach has some inherent disadvantages; it would be difficult to maintain standardization of hGH measurements and avoid drift in values. The production of a matrix reference material from pooled serum, and its use for calibration, should lead to equivalence of results. Standardization might also be achieved through a protocol for producing a commutable matrix calibrator from IS 98/574, but our results show that this is challenging. LC-MS methods could provide reference values. However, the limited correlation between the data from the group of immunoassay-based procedures and data from the LC-MS-based measurement procedure may indicate differences in selectivity. Therefore, the many factors influencing commercial measurement procedures and LC-MS need to be clearly understood to allow control of all sources of calibration bias and design a stable reference system on the basis of LC-MS.

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References


