A global multicenter study on reference values: 1. Assessment of methods for derivation and comparison of reference intervals

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ABSTRACT

Objectives: The IFCC Committee on Reference Intervals and Decision Limits coordinated a global multicenter study on reference values (RVs) to explore rational and harmonizable procedures for derivation of reference intervals (RIs) and investigate the feasibility of sharing RIs through evaluation of sources of variation of RVs on a global scale.

Methods: For the common protocol, rather lenient criteria for reference individuals were adopted to facilitate harmonized recruitment with planned use of the latent abnormal values exclusion (LAVE) method. As of July 2015, 12 countries had completed their study with total recruitment of 13,386 healthy adults. 25 analytes were measured chemically and 25 immunologically. A serum panel with assigned values was measured by all laboratories. RIs were derived by parametric and nonparametric methods.

Results: The effect of LAVE methods is prominent in analytes which are reported in the second part of this paper.

Conclusion: The collaborative study enabled us to evaluate rational methods for deriving RIs and comparing the RVs on a global scale.

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1. Introduction

The reference interval (RI) is defined simply as the prediction interval which includes the central 95% of reference values (RVs), or test results from well-defined healthy individuals (reference individuals). Establishment of well-controlled, reliable RIs is an important mission for all clinical laboratories. In reality, it is very challenging, because it is not easy to recruit a sufficient number of reference individuals, to control pre-analytical variables, and to apply all statistical methods in appropriate manners. The international guideline entitled “Defining, establishing, and verifying reference intervals in the clinical laboratory” was first published as a possible solution in 1996 by collaboration between the Clinical and Laboratory Standards Institute (CLSI) and the International Federation of Clinical Chemistry and Laboratory Medicine (IFCC) (the latest 2010 version is designated as CLSI/IFCC EP28-A3c [formerly, C28-A3] [1]. However, the descriptions are generally theoretical in nature, and the rationales of the recommendations have not been well evaluated by implementing actual, down-to-earth RI studies. In fact, there have been controversies over its pragmatic aspects, such as the rationale for secondary exclusion, the use of parametric vs. nonparametric derivation, and determining how to judge the need for partitioning RVs [2–5].

In the interim period, for the achievement of global standardization for major laboratory tests, the current consensus is to derive RIs in a reproducible manner based on sufficient sample size by collaboration between multiple laboratories and to use the derived RIs in common or by transference [6–8]. The Scandinavian groups conducted a multicenter study for derivation of common RIs based on standardized test values and found virtually no between-country differences in the Nordic countries [9]. However, the appropriateness of the study protocol and the method used for evaluating between-laboratory differences need to be evaluated by use of newer statistical methods. In addition, the IFCC Committee on Plasma Proteins concurrently conducted two RI studies mainly aimed at deriving common RIs for major serum proteins in East and Southeast Asian countries in 2000 and 2004; it revealed apparent between-country differences in many of the analytes, especially those of inflammatory markers [10,11].

With this background and its mission of promoting proper implementation of multicenter RI studies, the Committee on Reference Intervals and Decision Limits (C-RIDL) was established by the IFCC in 2005. The primary project of the C-RIDL, as planned in early 2010, was to clarify between-country differences in RIs on a global scale and to seek the most practical and harmonizable methodologies for conducting the RI studies. The key strategy of the global study was to make RVs comparable among the countries through measurement of a common panel of serum samples. After conducting a feasibility study to confirm the validity of cross comparison of test results based on the panel test results [12] and the elaboration of the common protocol [13], the global multicenter study was launched at the end of 2011 on a trial basis. The more
concrete objectives were to address the issues listed below and to develop a comprehensible picture of the most rational way of conducting the RI study in an internationally harmonized way.

1) How strictly should we set-up exclusion criteria in recruitment of candidate reference individuals? It may differ from country to country.
2) In what situation is it necessary to exclude individuals secondarily? What is the rational method to do so?
3) Why not just use a nonparametric method for statistical derivation of the RI? Does the parametric method give the RI reproducibility for each analyte?
4) How comparable are the RIs if the analytes are supposed to be globally standardized?
5) How reliable is the scheme of aligning RIs based on panel test results measured in common?
6) Are there ethnic or regional effects in the RIs?
7) Are major sources of variations (sex, age, BMI, etc.) of RIs the same among the countries?

As of August 2015, the global study involved 17 countries from 5 continents, the majority of which sought to establish country-specific RIs. In this intermediary report, as the first part of the key note article for the study, the methodologies in common use are briefly described, and the analytical results from 12 countries that had completed their studies were evaluated collectively for addressing the first five of the above issues. Evaluation for the issues (6) and (7) will be presented in a second report [14].

2. Methods

2.1. The common protocol for the study

The study protocol [13] commonly used in each country is briefly described as follows:

2.1.1. Recruitment

The target population was apparently healthy individuals 18–65 years of age with equal gender and age distributions. Those above 65 years were also included, even with small sample sizes. The test results of that age range were not used for any data analysis except in making profiles for sex- and age-related changes of reference values for the extended age range.

All potential participants should be feeling subjectively well. Any subject taking medications or vitamin supplements had them recorded at the time of recruitment. The following medications were prescribed as follows:

- Contraceptive pills or estrogen, plus thyroxine if the subject was maintained in a euthyroid state (TSH < upper reference limit).

Exclusion criteria were known diabetes on oral therapy or insulin (diet alone is acceptable); a history of chronic liver or kidney disease; blood test results that clearly point to a severe disease; a history of being a hospital inpatient or otherwise seriously ill during the previous 4 weeks; blood donation in the previous 3 months; known carrier state for HBV, HCV, or HIV; female participants who were pregnant, breast feeding, or within one year after childbirth; any other significant disease or disorder which, in the opinion of the investigator, might either put the participants at risk because of participation in the study or might influence the results of the study; or participation in another research study involving an investigational product in the past 12 weeks.

2.1.2. Sampling, storage and measurements

Blood sampling from the volunteers was done under basal conditions, including overnight fasting of >10 h, sitting for at least 20 min prior to sampling, and avoidance of strenuous muscular exertion for three days or working a night shift before the sampling. Sera were separated soon after sampling and stored immediately at −80 °C. This was because all specimens were tested collectively in batches in each country's central laboratory regardless of the time of sampling.

2.1.3. Target analytes and measurements

The 50 target analytes for the study were as described in the common protocol [13] and were composed of standardized analytes (lipids, electrolytes, enzymes, major serum proteins) and non-standardized but harmonizable analytes (inflammatory and tumor markers, hormones and vitamins). All were measured in the serum samples by use of automated analyzers employed in each central laboratory for routine use. Quality control of the assays was performed by use of commercial QC sera of routine use and the mini panel composed of 5–6 sera from healthy individuals [5].

2.2. The serum panel and assignment of traceable values for standardization

Two serum panels were produced by the C-RIDL in 2011 and in 2014, respectively composed of 80 (all Japanese) and 100 (70 Japanese; 30 from other ethnicities) sera from healthy individuals with nearly equal gender and age (20–65 years) proportions. The scheme of the value assignment for the first panel for standardized analytes was described previously [5,13]. The second panel was divided into two parts, one for chemistry analytes including those measured by immunoturbidimetry, and the other for analytes measured by other immunoassays. Assignment of values traceable to the reference measurement procedures (RMPs) for five chemistry analytes (TC, TG, urea, UA, Cre), and for seven enzymes (AST, ALT, LDH, GGT, ALP, CK, AMY) was performed by reference laboratories associated with the Committee on Traceability of Laboratory Medicine (C-TLM) of IFCC: the University of Bonn Faculty of Medicine (led by Dr. Anja Kessler and Prof. Lother Siekmann) and the Hanover University School of Medicine (Prof. Gerhard Schumann).

More specifically, for the first five analytes, six sera out of the 50 chemistry part of the panel were chosen for each analyte so that their values covered a wide portion of the presumptive RIs: i.e., a total of 25 sera (3–6 levels × 5 analytes) were directly value assigned. The indirect value assignment was accomplished by collaborative measurement of all 50 sera in five leading laboratories in China (Beijing Union Medical School Hospital, Beijing), South Africa (PathCare Laboratories, Cape Town), Japan (Beckman-Coulter Laboratory, Mishima, Shizuoka), and Canada (Hamilton General Hospital, Hamilton, Ontario). Based on the linear relationship between the assigned and measured values for the 3–6 sera, all 50 test results were recalibrated. Finally, the average recalibrated values for each of the 50 sera were adopted as the indirectly assigned values.

For the 7 enzymes, three pooled sera with different enzyme activities covering the presumptive RIs were first prepared for each enzyme. The 3 × 7 pooled sera were then value assigned directly for use as the calibrators. The 50 sera in the chemistry part of the panel were assigned values of their enzyme activities indirectly by use of the three sets of calibrators for each enzyme.

The list of assigned values for the 50 chemistry samples of the panel is shown in Suppl. Table 1, which also includes tables of assigned values for the first serum panel both in SI and conventional units [5].

2.3. Alignment/harmonization of reference values among the countries

The basic scheme for conducting the global study was to make test results comparable among the countries based on the panel test results measured in common. All pairwise method comparisons were performed by use of major-axis linear regression analysis (MALRA) [12]. As a result, even if values from a laboratory were biased despite the assumption of standardization, accurate comparison across the countries was still possible by recalibration of results from all the laboratories into those matched to assigned values of the panel in the case of
standardized analytes. In the case of non-standardized analytes, test results from any country were aligned to those of a specified country for harmonized comparison of reference values.

In order to perform this recalibration or alignment procedure appropriately, the allowable limits of errors in converting values were formulated based on the theory of allowable analytic errors [5,12]. The theory in brief is as follows. The precision of RI limits (LL, UL), converted from one laboratory to another, based on the major-axis regression can be expressed in two ways. The first is by the SD ratio (SDR) of the standard error (SE) of LL or UL to the SD comprising RI [(UL-LL)/3.92, SDRLL (= SELL/SDRI) or SDRUL (= SEUL/SDRI). The other is by the CV of the regression slope b, CV(b). Analogous with the theory of allowable analytical bias [15], SDR ≤ 0.125 is regarded as optimal, and SDR ≤ 0.25 as desirable. We demonstrated that there was a close linear relationship between CV(b) and SDRLL or SDRUL [12], and SDR = 0.125 corresponded to CV(b) = 5.5%, and SDR = 0.25 to CV(b) = 11.0%. In this study, for aligning test results of any given analyte by the regression, we regarded CV(b) ≤ 5.5% as indicating the conversion error is at the “optimal” level, and 5.5 < CV(b) ≤ 8.0% as indicating “acceptable”. No alignment procedure was performed if CV(b) ≥ 8.0%.

2.4. Statistical analysis

2.4.1. Expression of between-group differences based on ANOVA

The magnitude of between-country or between-region differences was expressed as SDR. By use of one-way ANOVA, between-country SD (SDcent) and net between-individual SD (SDin) (after adjusting for the between-country variations) were calculated and their ratio was set as the SDR [5]. Analogous to the theory of allowable analytical bias, the critical limit for SDR (=SDcent/SDin) can be set to 0.25. However, the denominator, SDin, tends to be narrower when it is computed by ANOVA, especially by use of two or more-level nested ANOVA with successive subtraction of factor-dependent variations. This implies that the SDR is increased in proportion to a decrease of its denominator; therefore, we chose 0.3 rather than 0.25 as a guide for judging the practical implication of between-country differences. In this calculation of SDR, reference values of analytes with obviously skewed distributions, such as TG, ALT, GGT, and CRP (their names are marked by asterisks in Fig. 3), were logarithmically transformed before applying the ANOVA, and then, SDs in untransformed scales were computed through a method of reverse transformation as described previously [16].

2.4.2. Latent abnormal values exclusion (LAVE) method

Inevitably some individuals who had commonly-occurring latent diseases or who failed to observe the basal conditions required before sampling were included in the study population. To overcome this problem, the LAVE method was developed, which makes use of the association between abnormal values found in different analytes which are caused by common disorders such as the metabolic syndrome, muscular damage and latent inflammation. LAVE is an iterative optimization method for refining reference individuals by excluding subjects possessing abnormal values in related analytes [2,4,5]. Its advantageous features are (1) truncation of reference distributions does not occur, unlike conventional methods of outlier exclusion, because it excludes individuals who have results outside RIs of analytes other than the one for which the RI is being derived, and (2) there is no effect on analytes whose values are rarely outside the RI in healthy individuals. The disadvantage is a reduction in the number of subjects included in the statistics, but this reduction should be minimal, depending on how strictly the exclusion criteria have been set and how many extreme values are allowed in the selection process. In this study, 10 analytes (Alb, UA, Glu, TG, AST, ALT, LDH, GGT, CK, CRP) were set as reference tests (RTs), and one value outside the RI was allowed among them. In the procedure, the RI (LL–UL) of RTs was extended on both ends by 5% of the width of the RI, or (UL–LL) × 0.05. Therefore, assuming a Gaussian distribution of RVs, the extended RI (RI') corresponded to the mean ± 2.156 SD or the central 97% ranges of RVs [2]. The probability (Pr) of having two or more abnormal results in “fully normal” individuals is computed by use of 10 RTs with assumption of no association of test values among the RTs.

Pr(two or more values outside RI')

1 − Pr (no value outside RI') − Pr(one outside RI')

1.0 − [0.97]0.15 × [0.97]0.03

Therefore, the influence of the LAVE procedure is not great in deriving the RI as a central 95% range. On the other hand, if test results of RTs are associated with each other, and there exist individuals with latent abnormal results among RTs, reduction in the data size remaining as “full normal” will be greatly reduced. In this study, the performance of the LAVE methods was evaluated by comparing the RIs determined with or without applying it in various ways.

2.4.3. Derivation of the RI and 90% confidence interval (CI) of the RI limits

The RIs were derived using both parametric and nonparametric methods.

For the parametric method, reference values were first transformed to Gaussian by use of a modified Box-Cox equation [2,4]:

\[
X = \frac{(x-\bar{a})^p - 1}{p}.
\]

From the mean and SD in the transformed scale (mT, SDT), the central 95% interval was given as mT ± 1.96 SDT. Then, the lower limit (LLT = mT – 1.96 SDT) and upper limit (ULT = mT + 1.96 SDT) were back-transformed to obtain the RI (LL – UL) in the original scale as follow:

\[
LL = \left(\text{pULT} + 1\right)^1 + a \quad \text{UL} = \left(\text{pULT} + 1\right)^1 + a
\]

For the nonparametric method, reference values were first sorted, and the 2.5 and 97.5 percentile points were determined. The 90% CI for the LL and UL were predicted by the bootstrap method with repetitive resampling of 100 times both for the parametric and nonparametric methods. For smoothing the RI, average values for LL, Me, and UL were adopted as the final RI.
Table 1: Demographic profiles of each country's study.

<table>
<thead>
<tr>
<th>Area</th>
<th>N</th>
<th>18−65</th>
<th>66−</th>
<th>Male N by age</th>
<th>BMI</th>
<th>Drk (%)</th>
<th>Smk (%)</th>
<th>drug Tx%</th>
<th>Female N by age</th>
<th>BMI</th>
<th>Drk (%)</th>
<th>Smk (%)</th>
<th>drug Tx%</th>
</tr>
</thead>
<tbody>
<tr>
<td>China</td>
<td>2621</td>
<td>1249</td>
<td>1163</td>
<td>86</td>
<td>23.6 ± 3.0</td>
<td>23.5</td>
<td>29.1</td>
<td>3.2</td>
<td>0.2</td>
<td>0.1</td>
<td>1372</td>
<td>1279</td>
<td>93</td>
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<tr>
<td>Japan</td>
<td>655</td>
<td>295</td>
<td>264</td>
<td>31</td>
<td>22.9 ± 2.6</td>
<td>53.6</td>
<td>19.3</td>
<td>8.8</td>
<td>2.7</td>
<td>0.3</td>
<td>360</td>
<td>324</td>
<td>36</td>
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<tr>
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<td>512</td>
<td>257</td>
<td>244</td>
<td>13</td>
<td>24.6 ± 3.5</td>
<td>24.5</td>
<td>7.8</td>
<td>4.3</td>
<td>0.0</td>
<td>1.2</td>
<td>255</td>
<td>247</td>
<td>8</td>
</tr>
<tr>
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<td>757</td>
<td>246</td>
<td>245</td>
<td>1</td>
<td>23.0 ± 3.8</td>
<td>20.3</td>
<td>24.0</td>
<td>8.1</td>
<td>0.0</td>
<td>0.0</td>
<td>511</td>
<td>508</td>
<td>3</td>
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<tr>
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<td>311</td>
<td>303</td>
<td>8</td>
<td>24.1 ± 3.6</td>
<td>3.2</td>
<td>2.9</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>250</td>
<td>246</td>
<td>4</td>
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<tr>
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<td>396</td>
<td>389</td>
<td>7</td>
<td>28.5 ± 5.6</td>
<td>0.0</td>
<td>16.9</td>
<td>1.0</td>
<td>4.5</td>
<td>0.0</td>
<td>426</td>
<td>418</td>
<td>8</td>
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<tr>
<td>Turkey</td>
<td>3172</td>
<td>1589</td>
<td>1450</td>
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<td>26.6 ± 3.6</td>
<td>10.2</td>
<td>24.6</td>
<td>3.7</td>
<td>0.9</td>
<td>0.3</td>
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<tr>
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<td>114</td>
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<td>26.0 ± 3.3</td>
<td>78.1</td>
<td>8.8</td>
<td>2.6</td>
<td>0.0</td>
<td>0.9</td>
<td>187</td>
<td>179</td>
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<tr>
<td>South Africa</td>
<td>651</td>
<td>340</td>
<td>333</td>
<td>7</td>
<td>26.0 ± 5.8</td>
<td>5.6</td>
<td>1.8</td>
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<td>0.0</td>
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<td>311</td>
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<tr>
<td>South Africa</td>
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<td>194</td>
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<td>25.9 ± 3.7</td>
<td>56.3</td>
<td>13.6</td>
<td>1.0</td>
<td>2.0</td>
<td>0.0</td>
<td>395</td>
<td>379</td>
<td>16</td>
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<td>8</td>
<td>27.2 ± 5.1</td>
<td>65.2</td>
<td>5.4</td>
<td>5.4</td>
<td>8.7</td>
<td>4.3</td>
<td>152</td>
<td>135</td>
<td>17</td>
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<tr>
<td>Argentina</td>
<td>528</td>
<td>313</td>
<td>313</td>
<td>0</td>
<td>27.3 ± 4.0</td>
<td>47.0</td>
<td>22.7</td>
<td>3.8</td>
<td>1.0</td>
<td>0.0</td>
<td>215</td>
<td>213</td>
<td>2</td>
</tr>
</tbody>
</table>

BMI: body mass index (mean ± SD); Drk: alcohol intake; Smk: smoking; drug Tx%: % under drug therapy; dHT: drug for hypertension; dDL: drug for dyslipidemia; dHU: drug for hyperuricemia; HR Thy: thyroid hormone replacement; OC: oral contraceptive or female hormones
shown in Suppl. Figs. 1-1 and 1-2. Distributions of BMI differed greatly among the countries. The influence of these factors on the analysis of RVs will be described in part 2 of this report [14]. The proportions of those who smoked cigarettes were quite comparable among the countries. In contrast, the proportions of individuals who drank occasionally or regularly differed greatly between countries, partly for religious reasons.

For medications, we focused on drugs for hypertension (dHT) and drugs for dyslipidemia and hyperuricemia (dDL and dHU), all of which are known to affect test results for some analytes. Although the proportions of individuals taking either or both categories of drugs were low, those taking dHT were excluded from the analysis of UA, urea, Cre, and K, those taking dDL from the analysis of lipids, and those taking dHU from the analysis of UA. Oral contraceptives (OC) are reported to be associated with changes in serum levels of some chemistry analytes [21]. From a preliminary analysis, we found TG were higher and Alb and ALP were lower in users of OCs with their between users and non-users SDR ranging from 0.3 to 0.65. Therefore, in the analysis of the three analytes, we excluded individuals on OC (or receiving female hormone replacement therapy) in countries where the proportions of OC users were >5.0% (US 12.4%, Non-African in South Africa 12.9%, PHL 5.9%) of females, while those in other countries with the proportion between 0–3.1% were neglected (Table 1).

3.1.2. Analytes and measurements

Analytes measured in each country are listed in Suppl. Table 1. Serum proteins (CRP, IgG, IgA, IgM, C3, C4, TTR, and Tf) were measured by immunoturbidimetry and grouped with the general chemistry analytes. Most of the analytes except these specific serum proteins were measured in all countries; the analytes measured by so-called labeled immunoassays were less frequently measured because of the cost of reagents.

3.2. Correlations of panel test results among countries

The serum panel of the first lot (Panel-I) was assayed by all countries except Pakistan and the Philippines, which used the panel of the second lot (Panel-II). South Africa measured both panels on two occasions. The relationships between Panel I and II were evaluated in the Beckman Coulter Laboratory (Y) in Japan by parallel measurements of both panels. Then, the relationships between the new and past measurements of Panel-I (Y and Y′) were evaluated. Test results of almost all analytes were nearly identical (Y ≈ Y′), with only a few analytes showing bias that required recalibration by use of the linear equation Y′ = a + bY. Using these linear relationships, it was possible to convert test result X from Lab-X (Pakistan, South Africa) which measured Panel II to test result Z of Lab-Z which measured Panel I. As illustrated in Fig. 1, this cross-conversion becomes possible by first converting X to value Y of Lab-Y, which measured both Panel I and Panel II simultaneously. This conversion of X to Y is done by use of the regression equation for Panel-II test results: Y = a + bX. Then, from the relationship between the new and past values (Y and Y′) for Panel I in Lab-Y: Y′ = a′ + b′Y, value X is further converted to value Y′: Y′ = a′ + b′(a + bX). Finally, from the linear relationship between Y′ and Z for Panel I: Z = a″ + b″Y′, X was further converted to Z by use of the formula: Z = a″ + b″(a′ + b′(a + bX)).

By use of this conversion scheme in comparing test results among the countries, all the test results of countries which measured Panel II were aligned first to the values of Japan at the time of measuring Panel I in 2012. Then, further conversions were done if necessary depending on to which country’s values the alignment was to be made. Since the above procedure results in additive errors with successive conversions, we applied CV(b) ≤ 5.5, not 8.0%, as the allowable limit of conversion for the analysis of data from Pakistan, the Philippines, and South Africa.

Actual test results of each participating laboratory, and assigned values, if any, for the first and second lots of the serum panel are shown in Suppl. Fig. 2 by use of a correlation matrix graph.

3.3. Comparison of reference values among the countries before and after alignment

In order to compare reference values among the countries, reference values were all aligned to the assigned values for the analytes listed in Suppl. Table 1 (TP, Cre, UA, urea, TBil, TC, TG, HDL-C, LDL-C, Ca, IP, AST, ALT, GGT, CK, GGT, CRP, IgG, IgA, IgM, C3, C4, Tf, and TTR); they were otherwise aligned to the values measured in Japan based on pairwise linear regression analysis of the panel test results between assigned/Japanese values and each country’s values. Typical results are shown for UA and Cre in Fig. 2, with full results posted as Suppl. Figs. 1-1 and 1-2, for males and females, respectively. SDRs for between-country differences (SDRcnt) before and after alignment are shown at the top of each graph. The full list of SDRcnt computed separately for each sex is shown for 42 analytes in Suppl. Fig. 3. Not included in the list are those analytes which were measured by <6 countries (GH and five sex hormones: testosterone, estradiol, progesterone, LH, and FSH). For two analytes (Na

Fig. 1. Between-laboratory value conversion scheme based on test results of the serum panels. RVs of serum panel measured in common by two laboratories were converted from one laboratory to another by use of major-axis linear regression analysis (MALRA). Since there were two different serum panels (Panel I and Panel II), it was necessary to make a nested procedure for converting RVs from Lab-X (which measured Panel II in 2015) to those of Lab-Z (which measured Panel I in 2012). It was achieved by use of panel test results of Lab-Y, which measured both Panel I and II simultaneously in 2015 and also measured Panel I since 2012. The RV of Lab-X (X) was first converted to that of Lab-Y (Y’ measured by Panel II) by use of MALRA equation (right graph). The value Y based on Panel I was identical with the value Y based on Panel II because both panels were measured simultaneously. Therefore, the Y measured in 2015 could be converted to the Y of 2012 (Y′) by use of the MALRA for the relationship between Panel I values of 2015 and 2012 in Lab-Y (middle graph). Then, Y′ was further converted to the value of Lab-Z (Z) by use of the relationship between Lab-Y and Lab-Z for the Panel I test results (left graph). This serial value conversion from X to Z was achieved by use of the formula shown in the bottom.
Fig. 2. Comparison of RVs across the countries before and after alignments. The distribution of RVs for males were compared before and after alignment based on the panel test results. The box and center line in each scattergram represents the mid 50% range and median of RVs, respectively. The data size are shown to the right bottom of the country name. Af and NAf for ZAF represent African and Non-African population. SDR for between-country differences is shown on the right upper corner. The graphs are shown for two typical analytes (UA and Cre). Full figures are available in Suppl Figs. 1-1 and 1-2.
and Cl), it was not possible to align values across the countries because of the large CV(b).

Using SDRcnt ≥ 0.3 as a guide, prominent reduction in SDRcnt after alignment was observed for urea, LDH, Mg, cortisol, and PTH. Apparent between-county differences even after alignment were observed in many analytes in males (M) or in females (F): Those with SDRcnt > 0.5 were Alb, Mg, TBil (F), HDL-C (F), ALT(F), CRP(F), IgG, C3(F), VitB12, and Folate (F). However, the between-country differences observed were crude ones not adjusted for any possible influences on test results.

The most important factor which may confound the between-country differences is BMI, which differed widely from one country to another. The issues of adjusted comparison and multivariate evaluation of sources of variation (SV) are presented in the second part of this report [14].

3.4. Assessment of need for secondary exclusions

In this global study, we derived RIs in various ways in order to determine an optimal method for each analyte. Therefore, in this report,
comprehensible comparisons among the countries were made regarding differences of RIs between parametric and nonparametric methods, and with or without application of LAVE methods.

Three methods of deriving the RIs were compared: (1) nonparametric method without LAVE [NP-LAVE (—); CLSI method], (2) parametric method without LAVE [P-LAVE (—)] and (3) parametric method with LAVE allowing one abnormal result (value outside the slightly extended RI) in the reference analytes [P-LAVE (+)]. Since the target disorders or conditions which have high prevalence among apparently healthy individuals were those related to nutritional status or muscular damage, we chose the 10 analytes listed in the Methods section as the standard reference analytes to detect individuals with multiple abnormal values.

As typical examples, in Fig. 4, nine clinical chemistry analytes were chosen for evaluation: three for LAVE insensitive analytes (urea, Cre, K) and six LAVE sensitive ones (AST, ALT, LDH, GGT, TG, and CK). Data were limited to those of males because of their higher prevalence of the metabolic syndrome. The minimum number of subjects included in a data set was 300 males. This criterion led to inclusion of the following 7 countries: CHN, SAU, TUR, RUS, ZAF, USA, and ARG.

As expected, with regard to urea, Cre, and K, there were no changes in RI regardless of LAVE method, while the nonparametric method generally gave slightly wider RIs (higher ULs). For TG, AST, ALT, LDH, GGT and CK, there was a consistent tendency for reduction of the UL of RIs regardless of the country. ULs of RIs by the nonparametric method always were higher than those of the parametric method.

With application of the LAVE method, the reduction in data size for calculation of the RI was between 15 and 22%, pointing to worldwide high prevalence of latent abnormal values attributable to nutritional disorders or failure in complying with the pre-sampling requirements (muscular exertion, insufficient resting/fasting). Please note that the slightly smaller data size analyzed by the parametric method compared to the nonparametric method despite non-use of the LAVE method was due to the exclusion of values outside mean ± 2.81 SD once after Gaussian transformation in applying the parametric method. This procedure of removing influential data points in the periphery is required for an improved result in the parameter fitting. This fact partly explains why the nonparametric method gave higher ULs than the parametric method for almost all the analytes examined.

3.5. Consistency of the reference distributions and appropriateness of Gaussian transformations

RIs for 24 chemistry analytes were derived by the parametric method separately for males and females using the datasets from 10 countries (the UK was not included because of insufficient data size). The modified Box-Cox power transformation formula was used to make each reference distribution Gaussian (normal). The maximum likelihood method was used to predict power (p), and then the origin of transformation ‘a’ was optimized to the predictive p, iteratively. The value p represents the distribution pattern; i.e., p < 1.0 implies the distribution has a longer tail in its higher side; p = 1.0, Gaussian (normal) pattern or symmetrical; p > 1.0, the distribution has a longer tail in its lower side; p ≈ 0.0, log-normal.

In the left panel of Fig. 5, the distributions of the ‘p’ consisting of 20 points (10 countries × 2 sexes) were depicted for each analyte as box and whisker plots with their spans representing the central 50% and 90% ranges, respectively; 90% was chosen rather than 95% because of the small sample size. It is notable that almost all of the p values are below 1.0, implying that the reference distributions have a tendency of more or less larger tail toward the higher side of values. It is also of note that each analyte has its own distribution pattern regardless of the country. We found that the distribution pattern expressed as ‘p’ does not differ between the two sexes. From the figure, we can roughly categorize the analytes from the p value as follows: highly skewed (p < 0.25: very close to the logarithmic normal distribution), CRP, GGT, CK, ALT, and TG; moderately skewed (0.25 s p < 0.4: cube-root normal): TBil, AST, and AMY; slightly skewed (0.4 s p < 0.7: square-root normal): ALP, Urea, LDH, Fe, HDL-C, TC, LDL-C, UA, Cre, and K; almost normal (0.7 s p): Cl, Na, Alb, IP, Ca, and TP.

The success of the Box-Cox formula in transforming the reference distribution into the Gaussian pattern was evaluated by computing the skewness (Sk) [22], which reflects the degree of symmetry of the distribution. In the right panel of Fig. 5, the distributions of Sk computed after Gaussian transformation are depicted for each analyte using the same drawing scheme as the left panel. As shown on the top right, the Sk values for Gaussian (p = 1.0) and log-Gaussian (p = 0.0) distributions are theoretically 0.0 and 3.8, respectively, while the distributions with |Sk| < 0.2 are almost indistinguishable from the Gaussian distribution by inspection. It is obvious from the figure that the modified Box-Cox formula was very successful in transforming the reference distributions into Gaussian form in the vast majority of the cases. The conformity of power-transformed values to the Gaussian distribution was also evaluated by performing three statistical tests (test of skewness, χ² test of goodness-of-fit, and Kolmogorov-Smirnov test) and by visual inspection of probability plots (cumulative frequency on Y-axis and a power-transformed value on X-axis). Used for those evaluations were two examples of male datasets from Saudi Arabia and Japan that remained after the application of the LAVE method. The final data sizes were approximately 320 and 215, respectively.

Although the statistical testing of the datasets with those data sizes is generally very sensitive to a slight deviation from the Gaussian form, power-transformed values of any analytes were almost invariably judged as not significant (Suppl. Table 3). The probability plot (Suppl. Fig. 3) also showed linearity for its main segment (cumulative frequencies of 10 – 90%), except for the CRP dataset from Japan which required non-parametric derivation of its RI.

These findings support the reproducibility and accuracy of the parametric method in estimating the central 95% range of the reference distribution.

4. Discussion

The choice of appropriate reference individuals for the study of RIs (RI study) is a challenging problem. It is expected that reference individuals are healthy without any disease or condition which affects test results of the analyte for which the RI is being determined. It is usually essential to determine RIs for as many analytes as practical in the same samples for the sake of efficiency. However, it is difficult to find individuals who fulfill “normal” requirements for all the analytes. Furthermore, it is not easy to identify individuals who have latent disorders beforehand without the actual tests being evaluated. If the prevalence of a disorder is low, this does not pose a problem in determining the RIs. In contrast, if the prevalence is relatively high, it is crucial either to apply very strict exclusion criteria in recruitment or to apply secondary exclusion after testing.

Fig. 4. Comparison of RIs and their 90%CI derived in three ways. RIs were derived in three ways by nonparametric (NP) or parametric (P) method with/without LAVE method. Each RI was depicted by the horizontal bar with shades on both ends corresponding to the 90%CI derived by the bootstrap method. LAVE method was applied by use of the following test items as reference for exclusion: Alb, UA, Glu, TG, AST, ALT, LDH, GGT, CK, and CRP. They are reliable markers of nutritional derangement, muscular exertion or inflammation which constitute latent disorders commonly seen in apparent healthy individuals. Data sizes used for computation of the RIs after the selection process are shown next to the Y-axis for each subgroup. Nine analytes were chosen for contrasting the effect of LAVE: top-three as LAVE-insensitive, and other six as LAVE-sensitive analytes. Since the RIs were derived for male subgroup with data size of ≥ 300, the number of countries fulfilling the condition were 7: China (CHN), Saudi Arabia (SAU), Turkey (TUR), Russian (RUS), African of South Africa (ZAF) not including non-African, USA, and Argentina (ARG).
In the global study, the objective was to have a harmonizable, pragmatic protocol for conducting the RI study which could be applicable to all participating countries. Therefore, we had to apply a lenient policy for recruitment without setting any limitation to BMI, habits of alcohol drink or smoking, and even use of common medications, as long as the individuals were feeling healthy and did not have any chronic renal or hepatic disorder or a past history of significant diseases.

This policy inevitably required us to apply secondary exclusions criteria customized to each analyte and to some countries. The exclusion was done on an analyte by analyte basis for those on medications. For example, for subjects who were on medication or supplements which may affect lipids, Fe, Ca, vitamin D, B12, and folate were excluded. A similar approach was taken for subjects on antihypertensive drugs which may affect urea, UA, Cre, and K values. However, the percentage of individuals who were on these drugs was low (<10% for individuals over 50 years of age), indicating reasonable self-restraint on the part of volunteers.

Aside from medications, it is most important to remove the influence of latent diseases or commonly-occurring conditions as much as possible. In the CLSI/IFCC document EP28-A3c, this problem is treated as the power (p) by regression to the modified Box-Cox formula for Gaussian transformation. The success of the transformation was assessed by the skewness (Sk) of the distribution: i.e., |Sk| ≤ 0.2 indistinguishable from Gaussian pattern and |Sk| ≤ 0.3 very close to Gaussian pattern. Distribution of 20 p and Sk values, which were obtained from 10 countries in the derivation of RIs separately for males and females, were compared for 24 chemistry analytes. The spans of the box and whisker drawn for each analyte represent central 50% and 90% range of the values.

Fig. 5. The power (p) required for Gaussian transformation of reference values and the skewness after the transformation. In the process of deriving the RIs by use of parametric method, the distribution pattern of RVs were predicted as the power (p) by regression to the modified Box-Cox formula for Gaussian transformation. The success of the transformation was assessed by the skewness (Sk) of the distribution: i.e., |Sk| ≤ 0.2 indistinguishable from Gaussian pattern and |Sk| ≤ 0.3 very close to Gaussian pattern. Distribution of 20 p and Sk values, which were obtained from 10 countries in the derivation of RIs separately for males and females, were compared for 24 chemistry analytes. The spans of the box and whisker drawn for each analyte represent central 50% and 90% range of the values.

The possibility of too great a reduction in data size by applying the LAVE method can be a concern. The relative decrease in data size can be reduced if we apply stricter exclusion criteria at the time of recruitment. In reality, however, that policy makes it difficult to recruit a sufficient number of volunteers. On the other hand, the LAVE method can be applied in a looser way either by allowing up to two abnormal results among the reference analytes, or by a 2 to 5% expansion of the RI limits (LL, UL) of each reference analyte in judging abnormality. In fact, the LAVE method is of no use and leads only to reduction of the sample size and expansion of the 90% CI of the RI.
The reason why the parametric method was not recommended in the CLSI/IFCC guideline was the failure of Gaussian transformation by the Box-Cox transformation [1,27]. Therefore, it has been thought that the pattern of reference distribution is uncertain and cannot be expressed by the known mathematical formula on which the parametric method relied. As expected, the nonparametric method was recommended instead. However, it has been shown that the parametric method works appropriately when the modified Box-Cox power transformation formula is used, which includes the origin of transformation ‘a’ as an additional parameter to be fitted [2,4,5]. In this study, by compiling the intermediary data obtained in the process of computing RIs for each country, it now became clear that the reference distribution can be successfully transformed into the Gaussian pattern almost invariably, judged from the virtual lack of skewness after the transformation (Fig. 5, right panel; Suppl. Table 3; Suppl. Fig. 3). As for the controversial view of uncertainty about the distribution pattern of RVs, we have demonstrated that, viewed across countries, there is consistent pattern of distributions specific to each analyte (Fig. 5, left panel). This implies that the power required for Gaussian transformation is not determined by chance, but it is predetermined for a given analyte regardless of the countries. In fact, the p values (power) of the reference distributions for both males and female were very similar.

These findings imply that the parametric method based on the modified Box-Cox formula is the method of choice for deriving RIs in almost all situations. A known exception is a situation where the reference distribution is truncated on its lower side due to a limited level of assay quantitation. We encountered this problem in the derivation of the RIs for CRP (measured by routine and not high sensitivity methods), PSA in females, rheumatoid factor, and anti-thyroglobulin antibody. It was necessary to use the nonparametric method or a probability plot method for estimating the upper 2.5 or 5% limit. Another situation where the parametric method is not appropriate is when there are only an insufficient number of RVs for computing the RI. The method is sensitive to outlying points with data size below 150. However, this problem is easily overcome by applying the bootstrap resampling method to the same dataset and adopting the average LL and UL as the final limits of the RI. We used this smoothing technique both for the parametric and nonparametric methods.

So far we have discussed the C-RIDL task of establishing optimal methodologies for derivation of RIs, which was made possible through the comparison of analytical results gathered from the RI studies around the world. Another important task of C-RIDL was to put together RVs from around the world, overcoming the problems of insufficient standardization or non-harmonized status of assay methods. Our objective was to compare the RVs through harmonization and to investigate the regionality and ethnicity of RVs on a global scale. If successful, the information we collected is expected to contribute to globalized medical practice.

In the history of laboratory medicine, there have been no attempts to overcome the difficulties in establishing collaborative teams needed to recruit a large number of reference individuals in a harmonized way. Fortunately, toward the end of 2011, several laboratories from 17 countries willingly joined the study in response to the C-RIDL’s call for participation.

C-RIDL proposed that to achieve the goal, the key strategy was to produce a serum panel composed of many individual commutable specimens and to ask for its measurement by all collaborating laboratories for cross-comparison of the results. Conventionally, a panel of sera has been used mainly as a reference set of specimens for the validation of an assay under evaluation. It was necessary to confirm the feasibility of using it for aligning test results among the countries. In early 2011, 60 sets of the serum panel (the first lot), composed of 80 specimens from healthy individuals, were produced, and the feasibility study was carried out in spring of 2011 through collaboration among 4 laboratories from the US, Turkey and Japan [12]. The results clearly revealed the usefulness of the panel in aligning test results by pair-wise comparison of test results among collaborating laboratories, and the concept of allowable limit of error using CV(b) was also elaborated.

From this study, the practicality of the scheme of using the panel for cross-comparison and alignment of RVs has now become obvious. It is of note that some bias existed even for standardized analytes, and it was necessary to make adjustment for proper comparison of the results. Furthermore, the assignment of serum panel values traceable to the RMPs proved to be very useful for recalibration of the RVs and RIs. As an important outcome of the scheme, we showed an absence of between-country differences (SDRcnt < 0.3) regardless of sex in RVs for TG, Fe, LDL, ALP, IgM, TF, AFP, CEA, CA125, PSA, cortisol, PRL, and TSH. In contrast, prominent differences in both sexes (SDRcnt > 0.4) were revealed in RVs for TP, Tbili, HDL-C, Mg, CRP, IgG, C3, and Vit-B12. It is important to note, however, that these findings need to be confirmed by multivariate analysis to adjust for possible confounding effects of age, BMI and other sources of variation of RVs. The results of the detailed analyses are to be described in the second part of this report [14].

C-RIDL is now considering another utility of the panel: to provide a service of transferring the RIs determined country by country to any party (clinical laboratories or IVD manufacturers) that is interested in confirming, comparing or introducing the RVs without actually conducting an RI study. If some bias is found from the panel test results, the RI can be recalibrated easily based on the regression analysis. If the purpose is just to verify the RVs before transference or just to confirm the standardized status of the assay, it is not necessary to measure all of the sera in the panel. Only 5 to 10 sera are enough if their concentrations are widely apart. Exceptions are some analytes measured by immunoassays showing a high degree of molecular heterogeneity or assay imprecision [12]. It is notable that certified reference materials (CRMs) currently available are composed of just one to three specimens. Recalibration of any assay based on a few data points is not accurate, no matter how precisely CRMs were measured. In contrast, the specimens in the serum panel are value assigned for many of the standardized analytes. Therefore, only a small fraction of the panel is sufficient for use either to confirm the state of standardization or to recalibrate the RIs more accurately than CRMs. However, to make use of the panel for these purposes, it is necessary to produce a large number of aliquots of a smaller sub-set of a panel to make them widely available. With this perspective in mind, the second panel, fully value-assigned for the 12 analytes, was produced in 2014. It is now possible to produce multiple “secondary” serum panels inheriting the values from the second panel, each of which is composed of a smaller number of sera with traceable values. C-RIDL has now set this as one of the most important activities for the coming years.

5. Conclusion

There were two main objectives in the C-RIDL’s global project. One was to establish optimal methodologies for derivation of RIs through analysis of real-world datasets gathered from studies around the world. The other was to put together RVs on a global scale for comparison after harmonization and for exploration of the regionality and ethnicity of these RVs among all countries. Both objectives were achieved as planned, and the controversial issues posted at the beginning of this article were answered as below:

1) The strict criteria for recruitment tailored to each country may not be necessary if investigators in each country make efforts by their own standards of “healthiness”, and the measures below are applied.

2) In the situation where test results from healthy volunteers are expected to contain a sizable number of abnormal results due to the presence of common diseases or conditions influencing test results of the analyte concerned, it is definitely necessary to apply some measures for secondary exclusion regardless of the country. The more rational one is not outlier exclusion methods (univariate
approach, but rather the LAVE method (multivariate approach) despite the reduction in the data size.

3) The parametric method is applicable for derivation of the RIs almost invariably. Its validity was suggested from the consistent pattern of reference distributions specific for each analyte. On the other hand, the nonparametric method is applicable but not recommended when abnormal results are expected in test results from healthy individuals because there is no way to fully exclude them non-parametrically.

4) Some bias in the RVs existed even for the standardized analytes. Therefore, it is essential to make cross-comparison of them by use of multiple commutable specimens (serum panel) in conducting the multicenter study for derivation of the RVs.

5) Unbiased comparison of the RVs for most analytes is possible by use of the serum panel regardless of the standardized status of the assay.

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Conflict of interest statement

The authors state that there are no conflicts of interest with regard to publication of this article.

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