Comparison and Validation of 10 Equations Including a Novel Method for Estimation of LDL-cholesterol in a 168,212 Asian Population

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Abstract: Low-density lipoprotein cholesterol (LDL-C) is frequently estimated using the empirical Friedewald equation. We compared the accuracy of the novel equation named as the 180-c method (180-c), which estimates LDL-C using a stratification approach, to those of 9 previously suggested formulas, including the Friedewald equation.

We compared the 9 previously suggested formulas by calculating intraclass correlation coefficient (ICC) and weighted kappa index in relation to direct LDL-C measurement values. Two independent populations used in the validation were the Severance Hospital LDL-C (SHL) registry (n = 164,358) and the Korea National Health and Nutrition Examination Survey (KNHANES) 2009 to 2010 (n = 3,854), each representing the hospital patient population and the general Korean population, respectively.

The 180-c and DeLong equations showed the highest ICCs, indicating the best agreement with direct LDL-C measurement. The 180-c and Chen equations showed the highest kappa indices. For the hypertriglyceridemic subpopulation from SHL, the 180-c equation showed the best agreement with direct LDL-C measurement in terms of ICC. The 180-c and DeLong equations showed the highest ICCs, indicating the best agreement with direct LDL-C measurement. The 180-c equation performed appropriately in Asian population.

INTRODUCTION

Cardiovascular disease (CVD) is one of the most common causes of morbidity and mortality worldwide. Elevated low-density lipoprotein cholesterol (LDL-C) has been recognized as an important risk factor for CVD, with clinical trials conclusively showing that LDL-C lowering therapy can reduce the risk of CVD. For this reason, numerous clinical practice guidelines have continuously identified LDL-C as the primary target for CVD prevention and therapy. Understandably, accurate measurement of LDL-C is essential to proper utilization of clinical practice guidelines for patient care. Among various principles of LDL-C measurement, the ultracentrifugal method, which is also called as β-quantification, remains as the reference procedure because LDL-C is unaffected by the presence of chylomicrons or other triglyceride-rich lipoproteins in this method. Nevertheless, β-quantification is inadequate for routine clinical laboratory use as it is labor-intensive, time-consuming and required impractically large volume of plasma. Therefore, LDL-C is alternatively estimated using the empirical Friedewald equation in most clinical environments: LDL-C = (total cholesterol [TC]) – (high-density lipoprotein cholesterol [HDL-C]) – (triglyceride [TG/5]). The final term, which refers to the estimate of very low-density lipoprotein cholesterol (VLDL-C), is calculated using a fixed denominator of 5 in this equation. However, this uniformly fixed ratio has been suspected to cause incorrect results because VLDL-C is a group of various lipoproteins containing individually different proportions of TG to TC.

Many groups have consistently evaluated the accuracy of the Friedewald equation in different ethnicities or various disease entities, and proposed alternative formulas for more precise LDL-C estimation until today. For instance, DeLong et al proposed the expression (0.16 × TG) as a more accurate estimate of VLDL-C, suggesting a fixed factor of 6 rather than 5. Other factor values have also been suggested in specific populations, but no fixed factor appears to be accurate under all circumstances due to high inter-individual variance in the TG/VLDL-C ratio. To date, none of proposed alternative
methods have replaced the Friedewald equation in routine clinical practice, and the National Cholesterol Education Program (NCEP) still recommends the use of the original factor of 5 for estimating LDL-C.\textsuperscript{11}

Recently, Martin et al. have recommended a novel equation to estimate LDL-C by applying an adjustable factor for the TG:VLDL-C ratio, which was developed using a stratification approach based on the levels of TG and non-HDL-C.\textsuperscript{5} This formula was derived from the United States (US) population. However, a subsequent validation study reported that this novel equation has no clear benefit over the Friedewald calculation to make changes in medical decisions.\textsuperscript{12}

Although previous studies have already compared the accuracy of several formulas to estimate LDL-C in relation to the Friedewald equation, we compared a total of 10 formulas for LDL-C calculation, including the latest novel method,\textsuperscript{9} to the direct measurement of LDL-C. Therefore, the aim of this study was to compare the 10 equations and validate the most powerful method for LDL-C estimation, using the largest cohort sample size to date. We validated the clinical utility and application of different formulas for LDL-C calculation not only in the hospitalized patients, but also in the general population.

**METHODS**

**Study Subjects**

To compare the accuracy of LDL-C estimating equations, we used two independent population datasets; the Severance Hospital LDL-C (SHL) registry and the Korea National Health and Nutrition Examination Survey (KNHANES) 2009–2010.

The SHL registry was established from the electronic database of hospital patient records in Severance hospital for this study. Severance hospital is a 2100-bed university-affiliated teaching tertiary-level hospital in Seoul, Korea, with more than 10,000 outpatients visiting daily. From January 2008 to December 2013, we collected subjects who were \( \geq 18 \) years and whose lipid profiles (i.e., TC, HDL-C, LDL-C, and TG) were measured by the central laboratory (Neodin Medical Institute, Seoul, Korea) and analyzed within 24 hours after transportation.\textsuperscript{14} TC and TG levels were determined by enzymatic method using reagents from Sekisui Medical Corporation. HDL-C and LDL-C levels were measured by a homogenous direct assay using reagents from Sekisui Medical Corporation. All analytes were analyzed on a Hitachi 7600 automated analyzer.

**Measurements of Lipid Profiles**

In patients from the SHL, serum TC and TG levels were determined by enzymatic method using reagents from Sekisui Medical Corporation (Tokyo, Japan) and Roche Diagnostic (Indianapolis, IN), respectively, on a Hitachi 7600 automated chemistry analyzer (Hitachi, Tokyo, Japan). HDL-C and LDL-C levels were measured by a homogenous direct assay using reagents from Sekisui Medical Corporation on a Hitachi 7600 automated analyzer.

In the KNHANES 2009–2010, blood samples were collected from each subject after overnight fasting for more than 8 hours, refrigerated immediately, transported in cold storage to the central laboratory (Neodin Medical Institute, Seoul, Korea) and analyzed within 24 hours after transportation.\textsuperscript{14} TC and TG levels were determined by enzymatic method using reagents from Sekisui Medical Corporation. HDL-C and LDL-C levels were measured by a homogenous direct assay using reagents from Sekisui Medical Corporation. All analytes were analyzed on a Hitachi 7600 automated analyzer.

**Equations for Estimating LDL-C**

A total of 10 formulas for estimating LDL-C in units of mg/dL are summarized in Supplementary Table 1, http://links.lww.com/MD/A865. These include the novel 180-cell (180-c) method\textsuperscript{9} along with equations suggested by Friedewald et al.,\textsuperscript{8} Hattori et al.,\textsuperscript{15} Anandaraja et al.,\textsuperscript{16} Chen et al.,\textsuperscript{17} Cordova,\textsuperscript{18} Teerakanchana et al.,\textsuperscript{19} Ahmadi et al.,\textsuperscript{20} DeLong et al.,\textsuperscript{10} and Rao et al.\textsuperscript{21}

**Statistical Analyses**

We assessed the agreement and accuracy of the 10 equations for estimating LDL-C using the direct LDL-C measurement as the reference value. Two statistical concepts were utilized for comparison of the accuracy of the 10 equations compared to the direct LDL-C measurement.

Firstly, the intraclass correlation coefficient (ICC) was calculated to compare the degrees of agreement between the 10 formulas and the direct LDL-C measurement. The level of agreement was defined according to ICC value; good agreement when ICC >0.75, and moderate agreement when 0.5 < ICC <0.75.\textsuperscript{22}

Secondly, we calculated a weighted kappa (\( \kappa \)) index to compare the concordance for estimating LDL-C in relation to the direct LDL-C measurement according to the LDL-C level classification guideline. Similar to the ICC values, the \( \kappa \) index was used to define levels of agreement; good concordance when \( \kappa \) index >0.8, and substantial concordance when 0.6 < \( \kappa \) index <0.8.\textsuperscript{22} To induce the \( \kappa \) index from the datasets, two most commonly used clinical practice guidelines to classify LDL-C values were applied; the US guideline (<70, 70–99, 100–129, 130–159, 160–189, and \( \geq 190 \) mg/dL) and the European (EU) guideline (<70, 70–99, 100–154, 155–189, and \( \geq 190 \) mg/dL).\textsuperscript{6,7}

Bland–Altman plots were also expressed to compare the direct measurement of LDL-C and other estimates calculated using the 10 equations.\textsuperscript{22} Categorical variables regarding concordance were compared using the \( \chi^2 \) test, whereas numerical variables were compared using the \( t \)-test. A \( P <0.05 \) was considered statistically significant. All statistical analyses were performed using SAS version 9.2 (SAS Institute, Cary, NC), MedCalc version 12.7 (MedCalc software, Ostend, Belgium), R package version 3.0.2 (http://www.R-project.org), and SPSS version 20.0 for Windows (IBM Corp., Armonk, NY).
RESULTS

Characteristics of Two Independent Population Datasets—the SHL and the KNHANES 2009–2010

Table 1 summarizes the general characteristics and lipid profiles of subjects in two independent datasets; the SHL and the KNHANES 2009–2010. Among a total of 164,358 subjects in the SHL, 1,481 (0.9%) individuals showed TG level higher than 400 mg/dL. The mean age was older in the SHL than in the KNHANES 2009–2010, possibly due to the difference between the hospital registry and the general population.

Comparison of the Performance of the 10 Equations

The quantitative agreements according to ICC value between the direct LDL-C measurement and other LDL-C estimates using the 10 equations are presented in Table 2. Most equations except the Ahmadi and Anandaraja formulas showed ICCs >0.90 in both datasets, indicating excellent agreement with the direct LDL-C measurement. ICCs of the 180-c (0.980), DeLong (0.980), and Chen (0.977) equations were superior to that of the Friedewald equation (0.975) in the SHL. Similar patterns were observed in the KNHANES 2009–2010. Mean differences between direct LDL-C measurement and estimates of each method were less than 10 mg/dL in all equations except the Ahmadi equation, indicating that most formulas investigated in this study were comparable with the direct LDL-C measurement in terms of ICC validation. The Bland–Altman plots between LDL-C values determined by direct measurement and other methods in the SHL are depicted in Supplementary Figure 1, http://links.lww.com/MD/A865.

Concordance According to Clinical LDL-C Classification Guidelines

The weighted k index was used to assess the concordance between direct LDL-C measurement and other LDL-C estimates calculated using equations according to the clinical classification guidelines from the US and EU (Table 3). Only two methods showed superior concordance with direct LDL-C measurement when compared to the Friedewald equation; 180-c and Chen equations [k index: 0.866 (0.865) and 0.863 (0.866) according to US (EU) classification, respectively]. Other equations showed inferior concordance when compared to the Friedewald equation. Similar patterns were observed in the KNHANES 2009–2010.

Comparison Between the Friedewald Equation and the 180-c Method According to LDL-C Level

Since 180-c method showed the best agreement with direct measurement among the 10 equations based on ICC and k index, concordance of direct LDL-C measurement with LDL-C estimates using the 180-c method was evaluated in comparison with the Friedewald equation (Figure 1). Interestingly, the 180-c method appeared to overestimate LDL-C values significantly more often than the Friedewald equation (12.3% vs. 9.8%; \( P<0.001 \)); however, the concordant percentage was also significantly higher in the 180-c method (84.0% vs. 82.6%; \( P<0.001 \)) (Figure 1A). After stratification by LDL-C level, both Friedewald and 180-c equations tended to overestimate LDL-C compared to direct measurement, especially in the low ranges of LDL-C (<100 mg/dL for Friedewald and <130 mg/dL for 180-c), whereas both equations underestimated LDL-C in the high ranges (Figure 1B and C).

Subgroup Analyses Based on LDL-C and TG Levels Using the Friedewald Equation and the 180-cell Method

Subgroup analyses were performed to measure quantitative agreement using ICC values according to the strata of LDL-C (<69, 70–99, 100–129, 130–159, and ≥160 mg/dL) and TG (<150, 100–149, 150–199, and 200–399 mg/dL) levels. In other words, direct LDL-C and TG levels were divided into 5 and 4 categories, respectively, to conduct a total of 20 comparisons in different combinations. The quantitative agreement between direct LDL-C measurement and the Friedewald or 180-cell equation were compared by means of ICC values).

Overall, higher ICC values were observed in the 180-c equation than the Friedewald equation for all subgroups (Figure 2 and Supplementary Table 2, http://links.lww.com/MD/A865). However, the differences between the two equations were not significant. Subjects with high LDL-C (≥160 mg/dL) and low TG (<150 mg/dL) levels showed the
best agreement by the 180-c equation (ICC: 0.93), and the Friedewald equation (ICC: 0.92). Subjects with the lowest LDL-C (<70 mg/dL) and TG (<100 mg/dL) levels also showed comparable ICCs as determined by both equations (ICC by 180-c: 0.92, ICC by Friedewald: 0.91). Regardless of TG level, ICCs among the subjects with LDL-C <160 mg/dL, (all ICCs >0.90) (Supplementary Table 4, http://links.lww.com/MD/A865) describes the concordance levels between direct LDL-C measurement and other LDL-C estimates in terms of weighted k index in subjects with TG level >400 mg/dL, using the same classification as in previous analyses. The DeLong equation showed the best concordance with direct LDL-C measurement [k index: 0.618 (0.617) by US (EU) classification], superior to that of the Friedewald equation. The 180-c equation resulted in inferior concordance compared to the Friedewald equation.

Ancillary Analyses for the Hypertriglyceridemic Subpopulation

Quantitative and qualitative agreements between direct LDL-C measurement and other LDL-C estimates calculated using the different equations were assessed in subjects with TG level ≥400 mg/dL derived from the SHL. Among the 10 formulas, the 180-c equation showed the best agreement with direct LDL-C measurement (ICC: 0.82, P < 0.001), followed by Teerakanchana’s equation (ICC: 0.81, P < 0.001) (Supplementary Table 3, http://links.lww.com/MD/A865). The equations proposed by Chen, Anandaraja, and DeLong also showed better agreement with direct LDL-C measurement than the Friedewald equation (ICC: 0.77). Supplementary Table 4, http://links.lww.com/MD/A865 describes the concordance levels between direct LDL-C measurement and other LDL-C estimates in terms of weighted k index in subjects with TG level >400 mg/dL, using the same classification as in previous analyses. The DeLong equation showed the best concordance with direct LDL-C measurement [k index: 0.618 (0.617) by US (EU) classification], superior to that of the Friedewald equation. The 180-c equation resulted in inferior concordance compared to the Friedewald equation.

Discussion

The international guidelines for CVD treatment continuously emphasize the importance of LDL-C assay for risk assessment and patient follow-up.1–7,25 The current NCEP Adult Treatment Panel recommendations for cardiovascular risk assessment are mostly based on early epidemiologic studies that used the Friedewald equation to estimate LDL-C.4 However, the accuracy of the Friedewald formula has been called into question based on several critical limitations including inaccuracy in patients with hypertriglyceridemia, subjects with very low TG level, and patients with liver or renal disease. Although the Friedewald equation has been traditionally believed to show high accuracy compared to β-quantification when TG <4.52 mmol/L (or 400 mg/dL), this has been also

| TABLE 2. Quantitative agreement by intraclass correlation coefficient between direct LDL-C and other equations for estimating LDL-C |
|-----------------|-----------------|-----------------|-----------------|
|                  | Intracl.        | Mean Difference |
|                  | correlation     | Mean ± SD       | 95% CI          |
|                  | coefficient     |                  |                 |
|                  | Value | 95% CI     | P-value | Value | Mean ± SD | 95% CI |
| SHL (N = 162,877) |       |            |         |       |       |        |
| LDL-C, Friedewald8 | 0.975 | 0.975–0.975 | Reference | −0.35 | 0.35 | −0.38–0.31 |
| LDL-C, 180-c9 | 0.980 | 0.979–0.980 | <0.0001 | −2.32 | 2.32 | −2.35–2.28 |
| LDL-C, Hatori et al15 | 0.974 | 0.973–0.974 | <0.0001 | 6.14 | 6.14 | 6.11–6.18 |
| LDL-C, Anandaraja16 | 0.901 | 0.900–0.902 | <0.0001 | −6.32 | 6.32 | −6.39–6.25 |
| LDL-C, Chen et al17 | 0.977 | 0.977–0.977 | <0.0001 | 0.42 | 0.42 | 0.39–0.46 |
| LDL-C, Cordova and Cordova18 | 0.935 | 0.934–0.936 | <0.0001 | 7.61 | 7.61 | 7.55–7.66 |
| LDL-C, Teerakanchana et al19 | 0.973 | 0.973–0.973 | <0.0001 | −6.39 | 6.39 | −6.43–6.36 |
| LDL-C, Ahmadi et al20 | 0.584 | 0.581–0.587 | <0.0001 | −26.05 | 26.05 | −26.24–25.85 |
| LDL-C, DeLong10 | 0.980 | 0.980–0.980 | <0.0001 | −5.17 | 5.17 | −5.20–5.14 |
| LDL-C, Rao et al21 | 0.969 | 0.968–0.969 | <0.0001 | −7.37 | 7.37 | −7.42–7.33 |
| KNHANES (N = 3854) |       |            |         |       |       |        |
| LDL-C, Friedewald8 | 0.957 | 0.955–0.960 | Reference | −0.50 | 0.50 | −0.79–0.21 |
| LDL-C, 180-c9 | 0.970 | 0.968–0.972 | <0.0001 | −2.44 | 2.44 | −2.68–2.20 |
| LDL-C, Hatori et al15 | 0.956 | 0.953–0.958 | 0.4554 | 6.52 | 6.52 | 6.23–6.80 |
| LDL-C, Anandaraja et al16 | 0.867 | 0.859–0.875 | <0.0001 | −4.89 | 4.89 | −5.39–4.39 |
| LDL-C, Chen et al17 | 0.968 | 0.966–0.970 | <0.0001 | 0.91 | 0.91 | 0.67–1.16 |
| LDL-C, Cordova and Cordova18 | 0.927 | 0.923–0.932 | <0.0001 | 9.21 | 9.21 | 8.86–9.55 |
| LDL-C, Teerakanchana et al19 | 0.959 | 0.956–0.962 | 0.1253 | −5.74 | 5.74 | −6.02–5.47 |
| LDL-C, Ahmadi et al20 | 0.509 | 0.485–0.532 | <0.0001 | −26.82 | 26.82 | −28.29–25.35 |
| LDL-C, DeLong et al20 | 0.968 | 0.965–0.970 | <0.0001 | −5.44 | 5.44 | −5.69–5.18 |
| LDL-C, Rao et al21 | 0.947 | 0.943–0.950 | <0.0001 | −7.84 | 7.84 | −8.18–7.51 |

Subjects with triglyceride ≥400 mg/dL were excluded from the analysis.
Mean Difference = [LDL-C measured by the direct method] − [LDL-C estimated by specific methods indicated above].
CI = confidence interval, KNHANES = Korea National Health and Nutrition Examination Survey, LDL-C = low-density lipoprotein cholesterol, SD = standard deviation, SHL = Severance Hospital low-density lipoprotein cholesterol registry.
suspected with the development of new direct LDL-C assays. These shortcomings of the Friedewald formula have prompted the recent development of new equations for LDL-C calculation. Among the formulas developed to overcome problems of the Friedewald equation, the most up-to-date 180-cell method derived from a US population has been highlighted for possibility to be used in practice with high accuracy, which applies an adjustable factor for the TG:VLDL-C ratio using a stratification approach according to TG and non-HDL-C levels. This estimation method was shown to provide higher fidelity estimates than the Friedewald equation, resulting in more accurate guideline risk classification. However, these favorable findings are required to be externally validated in different ethnicities before being applied to other populations. Although one study validated the novel 180-cell method concluded that its improvement over the Friedewald equation is not sufficient to supplant the original formula, the results in this study were derived from another US population. To the best of our knowledge, this is the first study to validate a new stratification approach in an Asian population using two independent large populations: one hospital patient based cohort and one general population.

Among the 10 equations compared in the present study, all equations except the Ahmadi equation revealed ICC >0.90, suggesting that these equations are in good accordance with direct LDL-C measurement. The fact that the Ahmadi equation was derived only from the patients with high cholesterol (>250 mg/dL) could have caused this inferior ICC value compared to those of other formulas. When we analyzed k index to distinguish the most concordant equation, two most accurate methods showing the best concordance agreement according to NCEP category with direct LDL-C measurement were the 180-c method and the Chen equation ([k index: 0.866 (0.865) and 0.863 (0.866) by US (EU) classification, respectively]. The Friedewald and DeLong equations followed with k index: 0.856 (0.854) and 0.825 (0.823) by US (EU) classification, respectively.

Considering the results of several previous reports by objective third parties who focused on the validation and comparison of suggested equations for LDL-C estimation, our results highlight the superior performance of the novel 180-c method. Also our data support the good predictive performance of the Chen equation, which has been undervalued in other reports. Martins et al recently reported that the Hattori formula performed the best in hospitalized patients when compared to the Friedewald, Chen and Cordova equations. Although direct LDL-C measurement was used for reference value in comparisons (as in our study), the use of different reagents and instruments for lipid measurements as well as different ethnicities and populations with various health conditions might have caused the discordant findings with our results. On the contrary, Oliveira et al concluded that the

### Table 3. Weighted Kappa Index: Concordance in Guideline Classification of LDL-C Levels by Friedewald versus Other Equations for Estimating LDL-C in Relation to Direct LDL-C

| US Guideline     | EU Guideline     |
|------------------|------------------|
|                  | Weighted Kappa   | SE     | P-value | Weighted Kappa   | SE     | P-value |
| SHL (N = 162,877) |                  |        |         |                  |        |         |
| LDL-C, Friedewald et al<sup>8</sup> | 0.856 | 0.0008 | –       | 0.854 | 0.0009 | –       |
| LDL-C, 180-c<sup>9</sup> | 0.866 | 0.0008 | <0.00001 | 0.865 | 0.0009 | <0.00001 |
| LDL-C, Hattori et al<sup>15</sup> | 0.806 | 0.0009 | <0.00001 | 0.812 | 0.001 | <0.00001 |
| LDL-C, Anandaraja et al<sup>16</sup> | 0.678 | 0.0012 | <0.00001 | 0.673 | 0.0014 | <0.00001 |
| LDL-C, Chen et al<sup>17</sup> | 0.863 | 0.0008 | <0.00001 | 0.866 | 0.0009 | <0.00001 |
| LDL-C, Cordova and Cordova<sup>18</sup> | 0.695 | 0.0011 | <0.00001 | 0.724 | 0.0012 | <0.00001 |
| LDL-C, Teerakanchana et al<sup>19</sup> | 0.789 | 0.001 | <0.00001 | 0.776 | 0.0012 | <0.00001 |
| LDL-C, Ahmadi et al<sup>20</sup> | 0.395 | 0.0013 | <0.00001 | 0.394 | 0.0015 | <0.00001 |
| LDL-C, DeLong et al<sup>10</sup> | 0.825 | 0.0009 | <0.00001 | 0.823 | 0.001 | <0.00001 |
| LDL-C, Rao et al<sup>21</sup> | 0.759 | 0.001 | <0.00001 | 0.757 | 0.0012 | <0.00001 |
| KNHANES (N = 3854) |                  |        |         |                  |        |         |
| LDL-C, Friedewald et al<sup>8</sup> | 0.804 | 0.0062 | –       | 0.788 | 0.0078 | –       |
| LDL-C, 180-c<sup>9</sup> | 0.835 | 0.0058 | 0.0003 | 0.822 | 0.0073 | 0.0015 |
| LDL-C, Hattori et al<sup>15</sup> | 0.762 | 0.0068 | <0.00001 | 0.757 | 0.0081 | 0.0058 |
| LDL-C, Anandaraja et al<sup>16</sup> | 0.631 | 0.0084 | <0.00001 | 0.603 | 0.0103 | <0.00001 |
| LDL-C, Chen et al<sup>17</sup> | 0.837 | 0.0058 | 0.001 | 0.831 | 0.0071 | <0.00001 |
| LDL-C, Cordova and Cordova<sup>18</sup> | 0.65 | 0.0077 | <0.00001 | 0.666 | 0.0092 | <0.00001 |
| LDL-C, Teerakanchana et al<sup>19</sup> | 0.761 | 0.0067 | <0.00001 | 0.731 | 0.0086 | <0.00001 |
| LDL-C, Ahmadi et al<sup>20</sup> | 0.381 | 0.0086 | <0.00001 | 0.366 | 0.0094 | <0.00001 |
| LDL-C, DeLong et al<sup>10</sup> | 0.787 | 0.0063 | 0.0544 | 0.768 | 0.008 | 0.0734 |
| LDL-C, Rao et al<sup>21</sup> | 0.703 | 0.007 | <0.00001 | 0.68 | 0.0089 | <0.00001 |

Subjects with triglyceride ≥400 mg/dL were excluded from the analysis.

KNHANES = Korea National Health and Nutrition Examination Survey, LDL-C = low-density lipoprotein cholesterol, SE = standard error, SHL = Severance Hospital low-density lipoprotein cholesterol registry.
FIGURE 1. Concordance of direct LDL-C with LDL-C using the Friedewald or 180-cell equations (A), and concordance according to LDL-C stratum between direct LDL-C and Friedewald (B) or 180-cell equations (C). LDL-C = low-density lipoprotein cholesterol.
Friedewald equation showed the best accuracy when compared to the Chen, Anandaraja, and Vujovic (which was not included in this study) formulas. Even though the reference method was different from that in our study (β-quantification in Oliveira’s study), none of the equations performed adequately for hypertriglyceridemic patients, which is in line with our findings.

When we focused on the comparison of only the 180-c method and the Friedewald equation, the ICC of the 180-c method (0.980) was superior to that of the Friedewald equation (0.975) in the SHL. In all subgroups classified by different strata of TG and LDL-C levels, higher ICC values were also observed using the 180-c method compared to the Friedewald equation; however, the difference between the ICC values was small. These results suggest that the novel 180-c method can be suitably and appropriately applied in Asian populations. Furthermore, it might perform more accurately than the Friedewald equation in all ranges of TG and LDL-C.

One interesting result in this study was that the ICC values for subgroup with LDL-C >160 mg/dL consistently showed the highest values with only a slight decreasing trend as TG level increased. However, in the subgroup with LDL-C <160 mg/dL, the ICC values proportionally decreased by significant amounts as TG level increased, resulting in ICC values of 0.54 to 0.63 among the subgroup with TG level of 200 to 399 mg/dL. From the clinical perspective, this finding is important because physicians can easily use and rely on the Friedewald or 180-c equation at their discretion in order to manage patients with high level of LDL-C (>160 mg/dL). Relatively small differences between direct LDL-C measurement and LDL-C estimates calculated using the other equations might not crucially change the treatment practice among patients who are already being treated. More importantly, a relatively large bias of LDL-C estimate in individuals with a low or middle range LDL-C is of serious concern because this underestimation will delay timely prevention and appropriate treatment for dyslipidemia. This is especially true for Asian populations because large proportion of the population was reported to be unaware of high LDL-C.

From the results of subgroup analyses, we challenged to find out the best equation to estimate LDL-C level in patients with hypertriglyceridemia, which is the most commonly recognized factor to cause misleading estimated as determined by the Friedewald equation. Among the 7 equations that had a good level of agreement according to the ICC value, the 180-c method outperformed all other equations in estimating LDL-C compared to direct measurement. Meanwhile, the Friedewald equation was indeed inferior to 6 equations in hypertriglyceridemic conditions. When we compared the performances of the 10 equations in terms of k index, all equations except the DeLong formula showed poor concordance level, not satisfying the minimum value of 0.6. Since the DeLong equation was the only formula that demonstrated fairly good agreement and concordance level by the indices of ICC and k index, we propose that physicians consider using the DeLong equation for LDL-C prediction in patients with hypertriglyceridemia. However, it is important to realize that the performance of all the suggested equations in the hypertriglyceridemic subpopulation was worse than that of the subgroup with TG level <400 mg/dL. Therefore, we suggest that the critical limitation of the LDL-C formulas still remains in patients with hypertriglyceridemia despite the development of several new equations.

One limitation in this study is that we used direct homogeneous LDL-C measurement as the reference value instead of β-quantification. Due to the limited clinical availability of β-quantification, direct LDL-C assays have frequently served as the reference values in several reports validating different equations for LDL-C estimation. To avoid incorrect comparisons of formulas with direct LDL-C data obtained from different methods, we used only one direct LDL-C assay (Sekisui reagent), which was reported to perform the best among seven assays when compared to β-quantification. Another limitation of our study is the different TC and TG assays used in two independent populations. However, a recent study reported that the use of different direct HDL assays is more likely to significantly affect the LDL-C estimates than various TC and TG methods because of the better standardization of TC and TG.

In conclusion, we compared the novel 180-c method for LDL-C estimation with 9 previously reported formulas in a Korean population as the first external validation in a non-US population. The 180-c equation appeared to be more accurate than the most commonly used Friedewald equation, along with the Chen equation. Although the DeLong equation showed better performance in the hypertriglyceridemic subpopulation, the 180-c method might perform better in hospitalized Korean patients as well as a general Korean population without hypertriglyceridemia.

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