Comparison of LDL-Cholesterol Enzymatic Method with Friedewald’s Formula

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This paper should be cited as: Yazdi HR, Piran F, Royani S, Nejabat M, Roshandel G, Taherizadeh M, Joshaghani HR [Comparison of LDL-Cholesterol Enzymatic Method with Friedewald's Formula]. mljgoums. 2015; 9(4):54-56
INTRODUCTION

The cause many deaths worldwide is coronary artery disease and many studies have shown that the risk of these diseases are related to increase of level low-density lipoprotein (LDL) (1). Lipoproteins carry cholesterol in plasma is five classes, including HDL, LDL, VLDL, IDL and LDL (2). LDL concentration is a marker in the diagnosis and prediction of cardiovascular disease. Also, studies have shown an association between ischemic and LDL is more than total cholesterol (3-6). LDL measurement accuracy is very important. Usually, total cholesterol and HDL-C measured directly and LDL-C calculated by Friedewald's formula (FF). The accuracy of the results of FF depends on parameters such as total cholesterol, TG and HDL. In this method, the TG exceeds from 4.5 mmol/L or presence of abnormal lipoprotein, differences can be seen between calculated LDL with reference method. Reference method for measuring LDL-C is a beta quantification method that's based on a two-step, ultracentrifugation and precipitation. LDL can be measured by chromatography, electrophoresis, immunological and nephelometry methods or can be calculated by FF that is a simple, convenient and low cost method (1, 4, 9,10). In this method (FF), LDL is calculated by this equation, LDL= total cholesterol _ (VLDL-C + HDL-C) [5], but this formula when used for the TG less than 400 mg/dl. But this formula is used when the TG < 400mg/dl, so factors such as concentrations of TG more than 400 mg/dl and initial hyperlipidemia type IV is the main limiting uses of FF (5,11). This formula for patients with diabetes, hepatopathy and nephropathy even when the TG is less than 400 mg/dl is not used (1). Despite its limitations, this method typically used to estimate the concentration of LDL-C in the laboratory (12) some studies have shown that LDL calculated using the FF in patients with TG between 4.51 and 8.82 mmol / l did not show any significant error. LDL-C was calculated using the FF even in patients with high TG 4.5mmol/l can be valid (5). In most studies, calculated LDL-C was slightly lower than the direct method (4). The aim of this study was to evaluate the correlation coefficient of LDL measured enzymatic with the FF.

MATERIAL AND METHODS

This cross-sectional descriptive-analytical study performed on 1411 patients. Cholesterol, triglycerides, HDL, LDL (Pars Azmoon) all patients assayed by enzymatic method by autoanalyzer (Mindray BS-200). For patients with triglyceride levels of less than 400 mg/dl had LDL levels were calculated by Friedewald's formula. Normal levels of LDL/HDL ratio of less than 3.5 were considered. A significance level of 95% for all tests was considered.

RESULTS

People participated in this study were 38.3% male and 61.7% female. The mean (SD) age was 46.3 (16.1) years. The mean (SD) serum cholesterol, triglycerides and HDL were 177.9 (41.1), 132.9 (73.2) and 45.8 (13.2) mg/dl, respectively. The mean (SD) LDL concentration was 82.1 (23.1) and 105.5 (35.8) mg /dl with direct assay method and calculation methods, respectively (p = 0.001). The LDL levels in the direct measurement were normal in 89.3 %, and in the calculation method 99.8 % of the cases were normal (P = 0.001). Serum LDL levels are measured using a direct method significant positive correlation with serum LDL levels were measured using a calculation method (Figure 1).

Figure 1- Correlation coefficient between the two methods of measuring LDL
DISCUSSION

Since heart disease is the most common cause of mortality worldwide and its increase is proportional to the increase in LDL, the measurement accuracy of LDL is essential. There are various methods for measuring LDL, which routinely in laboratories LDL is calculated from the Friedewald's method, but this method has limitations that the more important it is level of TG above 400 mg /dl. Some studies have shown when the TG higher than 400 mg/dl can use from FF (4). Also Türkalp study on 47 patients showed a high correlation between the measurement of LDL and FF a direct method (12,13). Another study by Cordova and co-workers with the aim of comparing the two methods of measuring LDL (direct method and FF) on 10664 patients was performed. They showed that the samples with different concentration TG two methods were not similar functional. When TG less than 150 mg/dl, may be LDL calculated by the FF can be invoked, but if TG between 301-400 mg /dl cannot be recommended to use of the FF (1). Another study by the same objective in 2005 reported even when TG less than 200 mg/dl is between LDL obtained from both direct and FF significant differences exist and the correlation coefficient between the two methods was 0.88, but some of the results inconsistent achieved and demonstrated that even in patients with TG more than 4.5 mmol /l FF method is valid and usable (14). A study by Mora and colleagues in 2009 of 27,331 healthy women with TG less than 400 mg/dl was performed. Baseline LDL determined by FF and direct measurement of fasting and non-fasting were compared homogenous risk (CVD) CVD in during an 11 year period. While low concentrations of LDL indirect method may be causing a lot of people be classified wrong in lower NCEP (Cholesterol Education Program International). In addition, the lack of association between non-fasting direct LDL with CVD (P <0.0001), causes a lot of questions regarding the clinical utility of direct methods for LDL with non-fasting blood samples (15-17). Can and colleagues' study was conducted on 1001 patients, cholesterol and TG levels were measured by enzymatic methods, and HDL and LDL levels were measured using the direct method was performed and showed LDL was estimated by the Friedewald's method significantly correlated with the direct method (P <0.01) (18). Study Timon-Zapata et al to study the effects of high levels, HDL in the calculation LDL use FF and other formulas that have recently been proposed, in the 2603 samples the HDL less than 20 mg/dl and 1953 samples with HDL more than 100 mg/dl done, showed significant differences encirclement between LDL was calculated with the formula and the direct method with two levels of HDL-C there. The results of the analysis suggest that none of the formulas should be used to calculate LDL in samples with high concentrations of HDL because there is no direct correlation with LDL may be used (19).

CONCLUSION

Despite the favorable correlation between measurement methods of LDL, the results of a calculation method is more than direct measure, this can have a negative impact on the judgment of the treating physician. The lack of standardized methods for direct measurement and comparison of results in different levels of TG, cholesterol, and HDL seems to require a comprehensive study and grouped based on different values of TG and HDL.

ACKNOWLEDGMENT

We thank the Kavosh medical laboratory that helped us in this project.

CONFLICT OF INTEREST

The authors declare no conflict of interest between them.

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