Lipoprotein(a) as a Risk Factor for Predicting Coronary Artery Disease Events: A Meta-analysis

Abstract

Backgrounds: Lipoprotein(a) [Lp(a)] are causally associated with risk of cardiovascular risk. Previous studies have proved that Lp(a) is an independent risk factor, which positively associated with stroke; but its role as a risk factor for coronary artery disease (CAD) remains controversial. We conduct this meta-analysis to determine the relationship between Lp(a) and CAD.

Methods: We retrieved articles in two electronic databases including PubMed and ScienceDirect, using the MeSH terms “lipoprotein a” OR “lipoproteins” AND “coronary artery diseases”. These articles were prospective studies, ranged from 2006 to August 2018.

Results: A total of 8 studies with 86808 participants and 8180 CAD events were incorporated in the meta-analysis. Compared with the low Lp(a) levels, the pooled HR was 1.132 (95% CI, 1.063-1.204, p=0.000). However, the eligible studies were heterogeneous (Q=25.74; I2=68.9%; p=0.001). Subgroup analysis for Lp(a) on CAD overall survival based on age and Lp(a) measurement method were also conducted. The results showed that Lp(a) is a risk factor for future CAD events, especially in the young.

Conclusion: Lp(a) level is an independent risk factor for CAD and may be especially relevant for young patients. Different detection methods for Lp(a) may modified the predictive value of lipoprotein (a) for coronary heart disease.

Keywords: Coronary artery disease; Lipoprotein(a); Meta-analysis; Prognostic value; Cardiovascular risk factors

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Abbreviations: ASCVD: Arteriosclerosis Cardiovascular Disease; AIS: Acute Ischemic Stroke; CAD: Coronary Artery Disease; CI: Confidence Interval; CVD: Cardiovascular Diseases; HDL-C: Low Level High-density Lipoprotein Cholesterol; HR: Hazard Ratio; LDL-C: Low-density Lipoprotein Cholesterol; Lp(a): Lipoprotein (a); OxPL: Oxidized Phospholipids; SE: Standard Error; TC: Triglycerides

Introduction

Lipid disorders are always divided into 4 categories: elevated level low-density lipoprotein cholesterol (LDL-C), low level high-density lipoprotein cholesterol (HDL-C), elevated triglycerides (TC), and elevated lipoprotein(a) [Lp(a)]. Elevated Lp(a) in the 4 types’ lipid disorders was rarely studied in the past decades. However, Lp(a) was an emerging marker for assessing the residual risk of cardiovascular diseases in recent year. Precious studies had proved Lp(a) was a risk factor for ischemic stroke [1-3], but the predictive value remained controversial in CAD prognosis.

In 1963, Berg firstly found Lp(a) in human plasma. Lp(a) is composed of an LDL-like particle in which Apo(B) is covalently bound by a single disulfide bond to apolipoprotein(a), and is highly homologous to plasminogen. There is extensive Apo(A) protein size heterogeneity, with >40 different isoforms, and >40 different sizes of Lp(a) particles, which is a unique occurrence
unlike other circulating proteins [4].

Circulating Lp(a) levels primarily depend on the LPA gene locus, and is not affected by diet and environment. The site of Lp(a) production has not been confirmed, and may be synthesized in hepatocytes, disse space or plasma compartment [5]. The steps contain that Apo(A) dock to LDL, and then form a covalent disulfide bond between KIV-9 of Apo(A) and Apo(B) of LDL. Lp(a) metabolism is different from LDL which stays longer in plasma than LDL. But the mechanisms which Lp(a) is cleared from plasma remain controversial, maybe relate to the numbers of LDLR.

Lp(a) lacks of sufficient evidence for physiological effects, presumable physiological effects including: repairing damaged vascular system and negative regulation of angiogenesis. In pathological condition, Lp(a) carries LDL particles with all of risk of atherosclerosis, including their propensity to oxidize after entry into the vessel wall, creating highly immunogenic and oxidized LDL. Apo(A) potentiates athero-thrombosis through other mechanisms, and potential anti-fibrinolytic effects by inhibiting plasminogen activation [6].

Elevated Lp(a) mediates myocardial infarction (MI), stroke, and peripheral arterial disease (PAD). In the last decade, the role in the underlying pathophysiology of atherosclerosis cardiovascular disease (ASCVD) has been fully studied. Zhang et al. [7] conducted a clinical trial and aimed to analysis the relationship between acute ischemic stroke (AIS) and Lp(a) levels, and to determine the prognosis value for predicting the functional outcome. The results showed that there was an increased risk of unfavorable outcome associated with Lp(a) levels 300 mg/L, and Lp(a) could be considered as an independent short-term prognostic marker. It also found that there was a relatively relationship between Lp(a) and ischemic stroke in children [8]. But the role of Lp(a) in CAD remained uncertain [9,10]. Therefore, this meta-analysis summarized current evidence on Lp(a) and CAD, and further analyzed the effect of age and detection methods.

Materials and Methods

Literature search and selection of studies

The electronic databases of PubMed and ScienceDirect were searched systematically in August 2018 without time limitation. The search strategy was performed by two independent readers (Yang Lan and Xiaoxiao Zhao), using the following keywords (“lipoprotein a” OR “lipoproteins” AND “coronary artery disease”).

Studies were eligible if they met the following criteria: (1) The study determined the relationship between Lp(a) levels and CAD events; (2) All studies must be prospective study and provide hazard ratio (HR) and 95% confidence interval (CI); (3) The sample size of each study was greater than 100; (4) Including either subjects chosen from the general population, with established cardiovascular disease (CHD, cerebrovascular or carotid disease, or peripheral artery disease) or hyperlipidemia; (5) The studies must be published in English with full-text paper available.

Data extraction

The papers selection and data extraction were accomplished independently by two reviewers. If the results provided are analyzed by both univariate and multivariate methods, we chose the latter. The extracted data included: Year of publication; first author; the journal of publication; follow-up periods; sample size; number of CHD events; age and sex; technique of Lp(a) measurement; hazard ratio;95% confidence interval.

Assessment of study quality

We used the standard Newcastle-Ottawa quality assessment scale to assess the Quality of all the studies. Newcastle-Ottawa scale [11] consisted of 3 factors: patient selection, comparability of the study groups, and assessment of outcome. Numbers from 0 to 9 (labeled as stars) represent different quality of each article. Cohort studies achieving 6 or more stars were considered to be of high quality.

Statistical analysis

In the study, the data was extracted into the EXCEL form, and then we used the stata version 12.0 (12.0 for Win, Stata Corp, College Station, TX, USA) to analyze. We extracted HR and 95% CI for each study, and HR was logarithmically transformed. Log HR was used to evaluate the relationship between Lp (a) and CAD events. We hypothesized that the heterogeneous maybe represent, and used random effects model for analysis. Random effects model was better suitable for a higher degree of heterogeneity. The prediction interval helped to estimate the range of real therapeutic effects in heterogeneous environments [12]. According to the Cochrane Handbook, Q and I^2 statistical indicators were used to analyze literature heterogeneity [13]. When p >0.05 or I^2 <50%, the fixed effect model could be performed, otherwise, using the random effects model. If there was significant heterogeneity in the given analysis, we adjusted for the type of literature and sample size to research for the source of heterogeneity. Finally, Egger test was used to analyze the potential publication bias.

Results

Search results

The selected studies were researched in two electronic databases (Figure 1). According to the prior search strategy, 1077 literatures were researched on PubMed, and 1202 literatures were searched in ScienceDirect. After reading the literature titles and abstracts, a total of 51 articles were selected for further study. We excluded 43 articles after reviewing the full text, the reasons including: duplicates; case report or cross-sectional study; the results without hazard ratios; the small sample size; study population repeated. Finally, 8 studies with 86808 patients met our inclusion criteria, and were performed for meta-analysis [4,14-20]. The mean follow-up period of all eligible studies ranged from 2 years to 20 years, and the quality of the studies assessed by the Newcastle-Ottawa quality assessment scale was about 6 scores, the detailed characteristics of all studies were summarized in Table 1.
Table 1: Clinical basic characteristics of the eligible articles.

<table>
<thead>
<tr>
<th>Time</th>
<th>First Author</th>
<th>Follow-up Period (year)</th>
<th>Numbers (CHD/total)</th>
<th>Male/female</th>
<th>Age (year)</th>
<th>Measurement of Lp(a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2006</td>
<td>Danik</td>
<td>10</td>
<td>934/27791</td>
<td>0/27791</td>
<td>54.2 ± 7.1</td>
<td>immunoturbidimetric assay</td>
</tr>
<tr>
<td>2009</td>
<td>Kamstrup</td>
<td>16</td>
<td>599/8637</td>
<td>3335/5302</td>
<td>55 ± 17</td>
<td></td>
</tr>
<tr>
<td>2010</td>
<td>Florance</td>
<td>10</td>
<td>635/9711</td>
<td>-------</td>
<td>55.3 ± 2.93</td>
<td>double-antibody enzyme-linked immunosorbent assay technique</td>
</tr>
<tr>
<td>2011</td>
<td>Lamon-Fava</td>
<td>12.3</td>
<td>145/2890</td>
<td>1328/1562</td>
<td>-------</td>
<td>immunoturbidimetric assay</td>
</tr>
<tr>
<td>2012</td>
<td>Viranic</td>
<td>20</td>
<td>1292/5777</td>
<td>5777/13318</td>
<td>53</td>
<td>double-antibody enzyme-linked immunosorbent assay technique</td>
</tr>
<tr>
<td>2012</td>
<td>Viranic</td>
<td>20</td>
<td>753/7541</td>
<td>7541/13318</td>
<td>54</td>
<td>double-antibody enzyme-linked immunosorbent assay technique</td>
</tr>
<tr>
<td>2012</td>
<td>Gurdasani</td>
<td>15</td>
<td>2365/18720</td>
<td>8381/10339</td>
<td>59.2 ± 9.2</td>
<td>immunoturbidimetric assay</td>
</tr>
<tr>
<td>2018</td>
<td>Dai</td>
<td>3.3</td>
<td>166/1602</td>
<td>1077/522</td>
<td>62.4 ± 10.6</td>
<td>Immunoturbidimetric assay</td>
</tr>
<tr>
<td>2018</td>
<td>Gregory</td>
<td>2.4</td>
<td>1146/4139</td>
<td>3328/811</td>
<td>63 ± 10/60 ± 9</td>
<td>Immunoturbidimetric assay</td>
</tr>
</tbody>
</table>

Prospective cohort studies

A total of 86808 participants and 8180 subjects with CAD in 8 prospective cohort studies were eligible for final analysis. These studies included different large scale populations, and the research methods were different. Six in eight studies reported that Lp(a) levels are closely related to CAD events. The meta-analysis results showed that the elevated Lp(a), the greater the CAD risk, and the pooled adjusted HR was 1.132 (95% CI, 1.063-1.204, p=0.000; (Figure 2). We created a funnel plot Figure 3 and performed an Egger test with a regression intercept of 3.19 (p=0.015), indicating potential publication bias in the meta-analysis. High heterogeneity was also observed (Q=25.74; I²=68.9%; p=0.001). Then we conducted a subgroup analysis to explore factors that affect heterogeneity.

Subgroup Analysis

Age

These prospective studies with mean age <55 years followed
Forest plot of meta-analysis for analyzing the relationship between Lp(a) and CAD. The pooled HR; CAD: Coronary Artery Disease.

Egger test and funnel plot for assessing publication bias. HR: Hazard Ratio; SE: Standard Error.

Detection measurement for Lp(a)

We screened the detection measurement for Lp(a) in the eligible literatures. In our meta-analysis, immunoturbidimetric assay were used in 4 studies, double-antibody enzyme-linked immunosorbent assay were used in 3 studies, and the remaining one did not provide the measurement. The different detection measurements didn’t modify the relationship between Lp(a) and CAD (immunoturbidimetric assay: the pooled HR: 1.191; 95%CI 1.039-1.365; double-antibody enzyme-linked immunosorbent assay technique: the pooled HR:1.150; 95%CI 1.074-1.231; (Figure 5).

Discussion

Cardiovascular diseases (CVD) are the leading killer in chronic diseases. Early detection of risk factors and prevention could reduce future CVD events. In the past decade, researchers were not optimistic about the emerging maker due to the detection measurements and medications. In recent years, researchers have gained a new understanding of Lp(a). Although Lp(a) lacks of sufficient evidence for physiological effects, it has widely proved that Lp(a) was closely related to the atherosclerotic cardiovascular disease (ASCVD) and may be related to its ability to transport oxidized phospholipids (OxPL). Covalent binding of OxPL to Apo(A) mediated the development inflammation [21]. In addition, Lp(a) could directly enter and accumulate, causing inflammation and calcification in the intima of artery or aortic valve leaflets [6]. It also reported that Lp(a) was considered to be an independent
and genetic risk factor, which causing cardiovascular events. Supporting evidence for the atherosclerosis of Lp(a) comes from Mendelian randomized studies [22], which showed that lifelong exposure to altered Lp(a) levels due to LPA gene variants could lead to changes in cardiovascular risk. It also proved that Apo(a) could promote atherosclerosis. The inflammatory process is now extensively studied in arteries.

In the large-scale prospective population study with 18,720 participants [17], it found that Lp(a) levels were associated with future risk of hospitalization and death in peripheral arterial disease (PAD). It confirmed the positive correlation between Lp(a) and CAD risk. Multiple studies in different populations and ethnic groups also confirmed the value of Lp(a) in the occurrence and prognosis of stroke [23-25]. These findings supported our conclusions and suggested that Lp(a) was a risk factor for vascular diseases. A previous meta-analysis has confirmed that elevated Lp(a) is an independent risk factor for ischemic stroke and may be particularly relevant for young stroke patients [26]. In the early family history of CVD, Lp(a) should be detected. The 2016 ESC/EAS guidelines recommended measurement of Lp(a) in selected cases at high risk or in patients with a family history of premature CVD [27]. However, a recent study showed that Lp(a) concentration was not associated with adverse cardiovascular outcomes in patients with recent ACS who are treated with statins. These findings called into question whether treatment specifically targeted to reduce Lp(a) levels would thereby lower the risk for ischemic cardiovascular events after ACS [19].

We conducted this meta-analysis with 8 prospective studies containing a total of 86,808 subjects with a follow-up period of 2 to 20 years, and 8180 subjects eventually developed CAD. The results showed that Lp(a) was significantly associated with CAD (the pooled HR: 1.132; 95% CI, 1.063-1.204; p=0.000), but the heterogeneity was high enough (Q=25.74; I²=68.9%; p=0.001). We found that the two studies mainly caused heterogeneity [19,20]. We removed the two studies and further analyzed. The results were shown in Figure 6. Lp(a) still independently predicted CAD prognosis, and the pooled HR was: 1.161 (95% CI: 1.106-1.219; p=0.000).

We performed a subgroup analysis to explore whether age and measurement methods could modify the relationship between Lp(a) and CAD. Subgroup analysis found that Lp(a) was more valuable in the young population than in the elderly, so we advocated early detection of Lp(a) in high risk population. The Lp(a) detection measurements had little effect on the relationship between Lp(a) and CAD. In recent years, it has been proposed that the amount-of-substance concentration of Lp(a) (nmol/L) is superior to the mass concentration (mg/L). Therefore, we still advocated the selection of more accurate measurement methods.

There are a number of limitations in this meta-analysis which should be discussed seriously. The 8 articles are from large-scale clinical trials. Newcastle-Ottawa scale were used to assess the qualities of literatures, and the results show that the score for each article≥6. However, we only research from two electronic databases, and the literatures are not comprehensive enough. There is also a high level of heterogeneity in the literature. We speculate that analytical data with high contrast and low contrast may be a source of heterogeneity, and we have not further analyzed it.

**Conclusion**

In conclusion, our meta-analysis emphasized the importance of high Lp(a) as an independent but modest risk factor in coronary artery diseases, and it became especially evident in younger populations.

**Acknowledgement**

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**Statement of Ethics**

The authors have no ethical conflicts to disclose.

**Disclosure Statement**

The authors have no conflicts of interest to declare.

**Author Contributions**

Yang Lan and Xiaoxiao Zhao respectively screen the literature from two electronic databases based on inclusion and exclusion criteria. When the two reviewers have disagreement, Yingyi Zhang, a third person, makes further judgments; Xiaofei Wang and Zhuangzhuang Zhang discuss statistical methods; Xincheng Song and Jia Chen revises the article.
References


