Lipoprotein (a) and cardiovascular risk factors in children and adolescents

ABSTRACT

Objective: To review the relationship between lipoprotein (a) [Lp(a)] and other risk factors for cardiovascular disease (CVD) in children and adolescents.

Data sources: This systematic review included studies from 2001 to 2011, a ten-year time period. Epidemiological studies with children and/or adolescents published in English, Portuguese or Spanish and fully available online were included. The searches were performed in Science Direct, PubMed/Medline, BVS (Biblioteca Virtual em Saúde) and Cochrane Library databases, using the following combination of key-words: “lipoprotein a” and “cardiovascular diseases” and “obesity”.

Data synthesis: Overall, 672 studies were obtained but only seven were included. Some studies assessed the family history for CVD. In all of them, Lp(a) levels were increased in patients with family history for CVD. There was also a positive correlation between Lp(a) and LDL-cholesterol, total cholesterol, and apolipoprotein B levels, suggesting an association between Lp(a) levels and the lipid profile.

Conclusions: The evidence that CVD may originate in childhood and adolescence leads to the need for investigating the risk factors during this period in order to propose earlier and possibly more effective interventions to reduce morbidity and mortality rates.

Key-words: cardiovascular diseases; obesity; child; adolescent; lipoprotein(a).

RESUMO

Objetivo: Revisar a relação da lipoproteína (a) [Lp(a)] com outros fatores de risco para doenças cardiovasculares (DCV) em crianças e adolescentes.

Fontes de dados: Revisão sistemática, com estudos do período de 2001 a 2011, caracterizando um recorte temporal de dez anos. Incluíram-se estudos epidemiológicos realizados com crianças e/ou adolescentes, publicados em inglês, português ou espanhol, disponíveis integralmente on-line. Realizou-se a busca nas bases de dados Science Direct, PubMed/Medline, Biblioteca Virtual em Saúde e Biblioteca Cochrane, utilizando-se a combinação dos descritores “lipoproteína a” e “doenças cardiovasculares” e “obesidade”.

Síntese dos dados: Encontraram-se 672 estudos, porém apenas sete foram incluídos na revisão. Alguns trabalhos avaliaram o histórico familiar para DCV. Em todos, os níveis de Lp(a) eram aumentados nos pacientes com esse histórico. Observou-se também correlação positiva entre Lp(a) e colesterol LDL, colesterol total e apolipoproteína B, sugerindo uma associação entre concentrações de Lp(a) e perfil lipídico.

Conclusões: A evidência de que as DCV podem ter sua origem na infância e na adolescência leva à necessidade de se...
investigaremos os fatores de risco nesse período, para planejar intervenções cada vez mais precoces e, possivelmente, mais efetivas, reduzindo a morbimortalidade.

**Palavras-chave:** doenças cardiovasculares; obesidade; criança; adolescente; lipoproteína(a).

**RESUMEN**

**Objetivo:** Revisar la relación de la lipoproteína (a) \( \text{Lp(a)} \) con otros factores de riesgo para enfermedades cardiovasculares (ECV) en niños y adolescentes.

**Fuentes de datos:** Se trata de una revisión sistemática, realizada de julio a agosto de 2011, con estudios del período de 2001 a 2011, caracterizando un recorte temporal de diez años. Se incluyeron estudios epidemiológicos realizados en niños y/o adolescentes, publicados en inglés, portugués o español, disponibles integralmente en línea. Se realizó la búsqueda en las bases de datos **Science Direct**, PubMed/Medline, Biblioteca Virtual salud y Biblioteca Cochrane, utilizando la combinación de los descriptores «lipoproteína a», «enfermedades cardiovasculares» y «obesidad».

**Síntesis de los datos:** Se encontraron 672 estudios, pero solamente siete fueron incluidos en la revisión. Algunos trabajos evaluaron el histórico familiar para ECV. En todos, los niveles de Lp (a) eran aumentados en los pacientes con ese histórico. Se observó además la correlación positiva entre \( \text{Lp(a)} \) y colesterol LDL, colesterol total y apolipoproteína B, sugiriendo una asociación entre concentraciones de \( \text{Lp(a)} \) y perfil lipídico.

**Conclusiones:** La evidencia de que las ECV pueden tener su origen en la infancia y adolescencia lleva a la necesidad de investigar los factores de riesgo en ese periodo, para planear intervenciones cada vez más tempranas y, posiblemente, más efectivas, reduciendo la morbimortalidad.

**Palabras clave:** enfermedades cardiovasculares; obesidad; niño; adolescente; lipoproteína (a).

**Introduction**

Cardiovascular diseases (CVD) are characterized as the main cause of death worldwide. These diseases develop slowly and progressively over life since early childhood. Among the variables associated with CVD, excess of weight (overweight and obesity) is one of the most important factors\(^1\). Therefore, it is believed that the prevention of CVD should begin in early childhood, with a special focus on the education process for cardiovascular health promotion, emphasizing the importance of the diet and the regular practice of physical activities over life\(^2\).

Accordingly, new risk markers related to lipid metabolism have been identified and studied, among which lipoprotein (a) or \( \text{Lp(a)} \) stands out because its persistently high plasma levels appear to be strong and independently associated with atherosclerosis\(^3,4\). Although the mechanisms of action associated with \( \text{Lp(a)} \) are still unclear, their involvement in atherosclerotic processes is active and silent\(^5\). Recent studies have reported that \( \text{Lp(a)} \) is a stable risk marker of the major forms of vascular diseases, with atherogenic and thrombotic properties\(^6\).

In this context, studies should be carried out to identify the cardiovascular risk factors in this age group and to clearly determine the problem magnitude, guiding actions to combat and prevent these risks. Thus, assuming that \( \text{Lp(a)} \) is related to other cardiovascular risk factors, this review summarized the main results of studies performed with this protein and verified its relationship with other risk factors for CVD in children and adolescents.

**Method**

To screen the most recent publications on this topic, a systematic and descriptive review was carried out, from July to August, 2011. Studies from 2001 to 2011 were included, featuring a ten-year time period. Epidemiological studies with children and/or adolescents, published in English, Portuguese or Spanish and fully available online were included.

The databases used for screening the articles were Science Direct, PubMed/Medline, Biblioteca Virtual en Salud and Biblioteca Cochrane, applying the following key-words combination: “lipoprotein a” and “cardiovascular diseases” and “obesity”. The first screening included the combination of listed key-words in order to identify abstracts in duplicate, and to exclude references related to abstracts without full article available, books and book indexes, review articles and those not specifically related to the topic. The considered articles were fully read, excluding those focusing on other outcomes. A new stage of research was conducted, in which the references of eligible articles were reviewed to capture manuscripts not found in the first search.
Design
To evaluate the Lp(a) sample
To assess levels of
To investigate, in children and adolescents with familial dyslipidemia, the association between Lp (a) and family history for CVD, and whether this association is independent of the altered lipid profile
Methodology
Main results

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Table 1 - Characteristics and main results of the included studies

| Author, year and country | Design | Objective | Sample | Methodology | Main results |
|--------------------------|--------|-----------|--------|-------------|--------------|
| Guardamagna et al(7), 2011, Italy | Cross-sectional | To investigate, in children and adolescents with familial dyslipidemia, the association between Lp (a) and family history for CVD, and whether this association is independent of the altered lipid profile | 231 children and adolescents (2–18 years) | Family history for CVD, anthropometric measurements (weight and height), BMI calculation; lipid profile measurements (TC, HDL-c, LDL-c and TG) and Lp(a) | % of patients with elevated Lp(a) levels was higher in those with family history for CVD. No significant correlation was found between Lp(a), age, BMI and lipid profile. |
| Saez de Lafuente et al(8), 2006, Spain | Cross-sectional | To determine the Lp(a) distribution in children and its relationship with anthropometric variables, lipid and thrombogenic factors | 98 healthy children (6–7 years) | Anthropometric measurements (weight and height), BMI calculation; Family history for CVD, lipid profile measurements (TC, HDL-c, LDL-c and TG), Lp(a), fibrinogen, D-dimer and PAI-1 | + correlation between Lp(a) and LDL-c; high Lp(a) levels in children with family history for CVD. Lp(a) without correlation with thrombogenic factors; |
| Obisesan et al(9), 2006, Spain | Cross-sectional | To evaluate the Lp(a) distribution in children and assess its association with lipid profile and anthropometric variables | 98 healthy children (6–7 years) | Anthropometric measurements (weight and height), BMI calculation; Family history for CVD, lipid profile measurements (TC, HDL-c, LDL-c and TG), Lp(a), Apo A1 and Apo B | High Lp(a) levels associated with LDL-c and Apo B |
| Wang et al(10), 2005, United States | Cross-sectional | To assess the effects of age and sex in the Lp (a) distribution and its relationship to other risk factors for CVD and diabetes among patients participating in the Study of Diabetes in Cherokee | 2,182 participants (5–40 years) | Calculation of BMI, weight, body fat percentage, SBP measurement, lipid profile measurements (TC, HDL-c, LDL-c and TG), glucose, insulin, OGTT, Lp(a), Apo and Apo B | In girls 5 to 19 years old, Lp(a) levels were associated with TC, LDL-c and Apo. |
| Obisesan et al(11), 2004, United States | Cross-sectional | To determine the correlation of Lp (a) in children and adolescents in the United States | 3,585 children and adolescents (4–19 years) | Calculation of BMI, height and WC, family history for CVD, birthweight, lipid profile measurements (TC, HDL-c, LDL-c and TG) | Ethnicity is associated with Lp(a) (higher in blacks). TC and family history associated with elevated Lp(a) levels. |
| Glowinska et al(12), 2003, Poland | Cross-sectional | To assess levels of novel risk factors for atherosclerosis in children and adolescents with obesity, hypertension and diabetes | 285 children and adolescents (6–20 years) | Anthropometric measurements (weight and height), BMI calculation, SBP measurement, lipid profile measurement (TC, HDL-c, LDL-c and TG), Lp(a), Apo A1, Apo B, homocysteine, fibrinogen, t-PA, PAI-1 | Obese, hypertensive and diabetic patients had significant changes in lipid metabolism, especially in TC, LDL-c, TG, Lp(a), Apo A1 and Apo B levels. |
| Gillumi et al(13), 2001, United States | Cross-sectional | To examine the association between body fat distribution and apolipoproteins, Lp (a) and TG in a representative sample of black, white and Hispanic children in the United States | 5,056 children (4–11 years) | Anthropometric measures (WC, HC, WHR, WTR, subcapsular and triceps skinfolds), calculation of BMI, TG, Lp(a), Apo A1, Apo B | Lp(a) was not associated with body fat distribution, regardless of age and BMI. |

BMI: body mass index; TC: total cholesterol; HDL-c: HDL cholesterol; LDL-c: LDL cholesterol; TG: triglycerides; SBP: systemic blood pressure; Hcy: homocysteine; PAI-1: plasminogen activation inhibition; Apo A1: apolipoprotein A1; Apo B: apolipoprotein B; OGTT: oral glucose tolerance test; t-PA: tissue plasminogen activator; WC: waist circumference; HC: hip circumference; WHR: waist-hip ratio; WTR: waist-thigh ratio.
The data were independently extracted by three authors, and disagreements were consensually resolved. The selected articles were classified according to year of publication, survey location, study type, population’s characteristics (subjects, number and age group), objectives, methodology and major findings.

Results

Initially, 672 papers were screened in the mentioned databases, but 667 were excluded (257 were revisions, 195 were not related to the topic, 140 were books and book indexes, 52 were not available in the full online version, 12 were only references, ten were mostly focused on other outcomes and one was found in two different databases). Thus, only five met the established inclusion criteria for this review. After reviewing the references of the selected articles, two new ones were added, totaling seven articles described in Table 1.

Regarding the population, four studies included children and adolescents, and only one included also adults. All investigations were conducted with boys and girls, aged from two to 19 years old.

The studies were conducted with healthy, dyslipidemic, obese, hypertensive or diabetic children and adolescents. The sample of most studies was obtained from other ones performed with larger numbers of participants.

All the selected papers were cross-sectional and carried out in Italy, Spain, United States and Poland. None of them was conducted in Brazil. Of the three studies carried out in the United States, two worked with different ethnic groups (white, black and Hispanic).

Some of the investigations assessed the family history for CVD. In all of them, Lp(a) levels were increased in patients with family history for the diseases. The Lp(a) dosage was performed with the patient’s serum using two different techniques (immunoturbidimetry and ELISA). For the Lp(a) levels interpretation, most studies considered the values above 30mg/dL as high, as recommended in the III Brazilian Guidelines on Dyslipidemias and Guideline of Atherosclerosis Prevention – Atherosclerosis Department of the Brazilian Society of Cardiology.

The goal of most studies was to investigate the relationship between Lp(a) and other cardiovascular risk factors, mainly the lipid profile, as well as anthropometric measurements and thrombogenic factors.

Discussion

The fundamental aim of this systematic review was to evaluate whether Lp(a) is associated with other cardiovascular risk factors. Although the action mechanisms associated with Lp(a) are still unclear, its involvement in atherosclerotic processes is active and silent. Recent studies have reported that Lp(a) is a stable risk marker of the vascular diseases major forms, with atherogenic and thrombotic properties.

Lp(a) shows a double-atherogenic character due to the fact that its lipid composition is similar to LDL and to the presence of apolipoprotein (a) in its structure, a protein with high degree of homology with plasminogen. In the study by Guardamagna et al., patients with family history of three or more cardiovascular events were more likely to have high Lp(a) levels, regardless of other lipid fractions. This fact could be observed in two other studies, Sáez de Lafuente et al. and Obisesan et al., in which the family history was significantly associated with elevated Lp(a) levels. These results indicate the need for pediatricians and cardiologists’ attention in order to identify and early intervene in the risk factors in children and adolescents with proven family history for CVD.

Family history is a strong and independent risk factor for CVD. At genetic level, a gene involved in the risk of developing a certain disease may predispose individuals of the following generations to the same risk. Positive family history for several cardiovascular events or stroke can be a simple criterion to assess pediatric patients who could benefit from Lp(a) measurements, thus contributing to identify patients at higher risk for CVD.

A positive correlation was observed between Lp(a) and LDL cholesterol (LDL-c), Lp(a) and total cholesterol (TC), and Lp(a) and apolipoprotein B (Apo B), which suggests an association between Lp(a) levels and lipid profile. According to Meabe et al., the fact that high Lp(a) levels are associated with high LDL-c levels proposes that the LDL metabolism may be involved in the Lp(a) synthesis. In patients with high LDL-c levels, Lp(a) is an important factor to determine atherosclerotic disease, as well as its severity and progression rate.

According to Giuliano et al., serum lipids and lipoproteins levels undergo profound changes during growth and development, with two phases of significant increase: up to the second year of life and during sexual maturation. There
is also a significant increase in LDL-c in puberty, especially among white boys.

In the study by Wang et al\textsuperscript{(9)}, girls showed Lp(a) levels associated with TC, LDL-c and Apo B. Epidemiological studies on the lipid distribution in children and adolescents have shown high levels of all lipoproteins and lipids in girls, regardless of age or skin color\textsuperscript{(16)}. The general consensus in literature confirms that Lp(a) concentrations are largely genetically determined\textsuperscript{(9)}, being therefore considerably variable between populations\textsuperscript{(17)}.

In the study by Obisesan et al\textsuperscript{(10)}, high Lp(a) levels were observed in black individuals. They have higher levels compared to other populations and the distribution is less distorted than in white individuals. Moreover, Lp(a) provides less risk in blacks than in whites, Asians or Indians. This risk is reduced due to its lower anti-atherogenic lipid profile (low LDL-c and triglycerides and high HDL-c levels compared to whites), which can partly offset the atherogenic potential of Lp(a)\textsuperscript{(15)}.

In the study by Glowinska et al\textsuperscript{(11)}, young obese, hypertensive and diabetic patients showed significant changes in lipid metabolism, particularly in relation to TC, LDL-c, triglycerides (TG), as well as Lp(a), apolipoprotein A (Apo A) and Apo B levels. This changed lipid profile was characteristic in children with obesity associated to hypertension. Several studies have demonstrated association between obesity and hypertension in children of various racial and ethnic groups, regardless of gender and age, synergistically influencing the cardiovascular risk. There are higher levels of blood pressure and/or higher prevalence of hypertension in obese children and, according to some authors, their risk of developing hypertension is two to three times higher than in non-obese individuals\textsuperscript{(18)}.

It has been well documented in literature that there is a direct correlation between plasma TC levels and LDL-c with CVD. Moreover, there is an inverse correlation between plasma HDL cholesterol (HDL-c) levels and the risk for these diseases. However, unlike the available literature on the CVD risk, the correlation between apolipoprotein abnormalities is not well established in diabetic and hypertensive patients\textsuperscript{(14)}.

According to some authors, apolipoprotein A1 (Apo A1) and Apo B concentrations have a stronger correlation with the atheroma development when compared to their equivalent lipoproteins LDL and HDL; in children, they are associated with coronary artery disease in their parents\textsuperscript{(11)}. Plasma Apo B concentrations have been considered a better representative of atherogenic particles, and some studies have shown that elevated plasma Apo B levels may be a valuable predictor of coronary artery diseases\textsuperscript{(14)}.

According to Sáez de Lafuente et al\textsuperscript{(8)}, obesity is another risk factor for CVD, significantly associated with lipids and lipoprotein metabolism. Studies have shown that obese children and adolescents have, even before reaching maturity, risk factors for CVD development, such as dyslipidemia, hyperinsulinemia, increased blood pressure and autonomic dysfunction\textsuperscript{(19)}. According to Silva and Zurita\textsuperscript{(20)}, childhood obesity is a risk factor for increased morbidity and mortality in adults, causing the later onset of CVD such as hyperlipidemic, colorectal cancer, type 2 diabetes, gout and arthritis. Obese children are also submitted to severe psychological stress due to social stigma. There are also frequent respiratory, orthopedic, dermatologic and immunologic complications, as well as hormonal disorders.

Pinhas-Hamiel et al\textsuperscript{(21)} reported that, in normal weight children, serum Lp(a) levels remain relatively constant throughout puberty. Glowinska et al\textsuperscript{(11)} found that Lp(a) levels are higher in diabetic patients, a fact that can be ambiguous in literature, because some studies claim that higher Lp(a) concentration is present in all diabetic children and others have reported increased Lp(a) concentrations only in patients with inadequate metabolic control. That is, further studies should be carried out in order to obtain a better understanding about the involvement of this marker in relation to diabetes. Nawawi et al\textsuperscript{(22)} reported that it is well established that diabetic patients have an increased risk of developing atherosclerotic vascular diseases compared with non-diabetic subjects. However, the contribution of Lp(a) in atherosclerotic complications in patients with type 1 and type 2 diabetes is still unclear. There are conflicting results as to whether Lp(a) concentrations are elevated in type 1 and type 2 diabetes. Several studies have shown that high Lp(a) concentrations in type 2 diabetes patients are an independent risk factor for coronary diseases, but, in relation to type 1 diabetes, the results are conflicting\textsuperscript{(22)}.

In the study by Sharma et al\textsuperscript{(23)}, conducted with obese or overweight African-American children and adolescents, it was found that Lp(a) is not considered an independent risk factor for CVD. There are some possible reasons for the controversial results found by studies on Lp(a) and cardiovascular risk factors. The factors affecting the plasma levels of this lipoprotein are different in each population. Dietary habits,
genetic factors, lifestyle and race are important factors that affect diseases of multifactorial origin, such as CVD.

In this context, some limitations of the present review should be noted. A small number of scientific studies met the established inclusion criteria. Despite attempts to search the references of the selected articles, only two additional studies were found. Moreover, the number of studies excluded from the review, mainly after the first screening, reflects the literature on this subject. Several studies were actually focused on the association between Lp(a) and cardiovascular risk in children and adolescents. It is noteworthy that all studies showed cross-sectional design. Additionally, long-term studies on cardiovascular risk factors are needed, because there are several metabolic and anthropometric changes in the age group under study, especially during puberty (21).

The association between Lp(a) and cardiovascular risk factors was studied mainly by international groups. Brazilian research may bring a more reliable and adequate panorama of our population, particularly among children and adolescents, in which Lp(a) is a possible predictor of CVD risk. According to Silva and Bittar (22), immediate interventions should be performed as soon as the first risk factors for CVD are identified in children and adolescents.

Considering this potential and the fact that there are few investigations about this new risk marker, more studies and researches on this topic should be carried out, which can provide support for intervention programs in health promotion and for actions to prevent and reduce cardiovascular risk factors in childhood and adolescence, providing greater participation of professionals involved in health promotion, and contributing to reduce morbidity and mortality.

However, the implementation of programs to change the lifestyle of risk factors carriers should be associated with environmental changes, which favors individual choices in adopting and keeping healthy habits. Moreover, encouraging cultural appreciation of health is an extremely important tool to achieve the healthy lifestyle goals.

It is expected that all these considerations help in the development of epidemiological research on these new cardiovascular risk markers in Brazil.

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