Small Dense Low-Density Lipoprotein: Biomarker or Potential Drug Target?

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Abstract

Ischemic heart disease is currently an epidemic affecting individuals worldwide. Increased incidence along with earlier onset of disease has led to the constant search for biomarkers that will help in earlier identification and treatment of at risk individuals. Small dense low-density lipoprotein (sdLDL) is the atherogenic subtype of low-density lipoprotein (LDL). It is smaller in size and higher in density in comparison to other LDL subtypes. Higher levels of sdLDL have been found to be associated with increased incidence of ischemic heart disease and adverse outcomes. Properties including decreased resistance to oxidative stress and prolonged residence time in the circulation account for its increased atherogenic potential. Hence intervention approaches targeting sdLDL directly in at risk individuals may be beneficial.

Genetic, lifestyle, and environmental factors affect sdLDL levels. But the main determining factor is the level of triglycerides (TGs). Higher TG levels are associated with higher levels of very low density lipoprotein (VLDL) 1 and sdLDL. Various drugs have been used for targeting sdLDL with varying outcomes; drugs tried out include statins, fibrates, niacin, cholesterol ester transfer protein inhibitors and sodium-glucose co-transporter-2 inhibitors. Future prospects include modification of enzymes involved in fatty acid and TG synthesis, for example, lipoprotein lipase and acyl CoA carboxylase. However, further research is still necessary to draw clear guidelines for sdLDL reduction therapy in coronary artery disease treatment and prevention.

Keywords

► ischemic heart disease
► small dense low-density lipoprotein
► atherogenicity
► drug target

Introduction

Cardiovascular disease (CVD), including coronary heart disease (CHD) and coronary artery disease (CAD), is a major cause of morbidity and mortality worldwide. According to the 2016 Heart Disease and Stroke Statistics update of the American Heart Association, in the United States, 15.5 million people ≥20 years of age have CHD. The prevalence increases with age for both men and women. In India the prevalence of CHD has increased fourfold in the past 40 years. The Registrar General of India reported that CHD-related adult deaths have increased from 26% in 2001 to 2003 to 32% in 2010 to 2013. Gupta et al reported that cardiovascular risk factors among Indians increase exponentially in the age group of 30 to 39 years. Covering 24.8%, CVD is the leading cause of death throughout India, including both rural and urban areas.

Increasing incidence combined with earlier onset has led to the constant search for biomarkers which will help in identifying at risk individuals at an early stage. Small dense LDL (sdLDL) is the smaller and denser subtype of LDL. Higher levels of sdLDL have been found to be associated with increased incidence of cardiovascular events. Hence sdLDL estimation could be used as a means of identification of high-risk individuals. Early intervention using drug therapy for reduction of sdLDL levels could ultimately lead to a better outcome for these people.
Cardiovascular Risk Assessment

Risk factors which increase the probability of CHD include diabetes, hypertension, central obesity, physical inactivity, smoking, psychosocial factors, dyslipidemias, and genetic predisposition. The INTERHEART study reported that more than 90% of the risk factors for CHD were measurable and modifiable. Among the lipoproteins, traditionally high-density lipoprotein (HDL) is considered as “good cholesterol” and LDL as “bad cholesterol.” Longitudinal studies had established the role of LDL as a risk factor for CHD occurrence, recurrence, and fatal outcome. Clinical trials proved that control of elevated LDL in high-risk patients was a very cost effective and efficient means of cardiovascular risk reduction. Hence lipid profile, in particular LDL, is routinely used by clinicians as a tool for cardiovascular risk assessment.

Other studies, however, found that LDL is not elevated in all those patients suffering from CAD. Moreover, though lipid lowering agents are widely used in clinical practice as a treatment component in CAD, the risk reduction after lipid lowering therapy has been found to be not more than 30% in most of the clinical studies.

These findings led to the search of risk factors in addition to those previously known which could contribute to the development of atherosclerosis and CAD, that is, the “beyond cholesterol” concept. Evidence started pointing toward the fact that it is in particular the sdLDL which is responsible for its atherogenic property.

LDL and Its Subtypes

LDL is the main cholesterol carrying lipoprotein. It is broadly defined as lipoprotein fraction with density 1.019–1.063 g/mL. Structurally, it consists of a core of cholesterol esters surrounded by a coat of phospholipids and protein (Apo B100 mainly). LDL delivers cholesterol to the cells, where it is utilized for synthesis of membranes, steroid hormones, etc. Krauss and Blanche used density gradient ultracentrifugation and separated LDL particles of normal subjects according to their size and density. Austin et al (1990) conducted a study on 301 subjects. They separated LDL using gradient gel electrophoresis and identified two distinct patterns, Pattern A with a predominance of large buoyant LDL particles (density 1.019–1.044 g/mL), and Pattern B consisting of mainly small dense particles (density 1.044–1.063 g/mL) (Fig. 1).

Pattern B (sdLDL) has been found to be associated with number of diseases including CHD, obesity, metabolic syndrome, acute ischemic stroke, and Type 2 diabetes mellitus (DM). High sdLDL itself is a component of “atherogenic phenotype,” consisting of high TGs, low HDL, and high sdLDL. This is now recognized as a distinct dyslipidemia linked to various genetic loci. Accordingly, sdLDL concentration was accepted as a risk factor for cardiovascular events by National Cholesterol Education Program (NCEP III).

Atherogenicity of sdLDL

The next question which arises is “Why is sdLDL atherogenic?” Small dense LDL differs from large buoyant LDL in certain properties. First, sdLDL has lesser affinity for the LDL receptor than other LDL subtypes. This leads to reduced LDL uptake and as a result, prolonged existence in the circulation. Second, sdLDL has greater affinity for arterial wall proteoglycans. The result is increased retention in the subendothelial space and hence greater opportunity for promoting atherogenic changes. These properties are a result of conformational changes in the Apo B part of sdLDL. These changes alter the properties of sdLDL; the result being decreased receptor-mediated uptake and increased proteoglycan affinity. Desialylation of sdLDL also contributes to this property.

Oxidized LDL is well known for its atherogenicity. LDL oxidation takes place in the circulation; it leads to the generation of specific epitopes which are recognized by receptors and induce immune response and inflammation. The altered lipid composition and lesser content of antioxidative vitamins reduce the resistance of sdLDL to oxidative stress and hence increase its atherogenic potential.

![Fig. 1 Low-density lipoprotein and its subtypes. LDL, low-density lipoprotein.](image-url)
Numerous studies have proven the atherogenic potential of sdLDL. Nishikura et al in a prospective cohort study of 7 years concluded that sdLDL is a promising biomarker to predict future cardiovascular events in the secondary prevention of stable CAD. In the Quebec Cardiovascular Study which was a prospective cohort study of 13 years, it was concluded that sdLDL is a strong independent predictor of ischemic heart disease (IHD). The Framingham Offspring Study which was a prospective cohort study on 1,680 women and 1,508 men concluded that sdLDL is a very strong predictor of IHD in women, whereas in men it is the sdLDL/LDL ratio which is a stronger predictor.

What Determines the Distribution between Large Buoyant and Small Dense LDL?

SdLDL is a subtype of LDL, so the basic synthetic pathway is the same. Triglyceride (TG)-rich very-low-density lipoprotein (VLDL) is synthesized in the liver and released into the circulation. TGs are acted upon by lipoprotein lipase and hepatic lipase producing intermediate-density lipoprotein and LDL. Simultaneously, LDL is acted upon by cholesterol ester transfer protein (CEPT) which removes cholesterol esters from LDL in exchange for TGs which it brings in from other lipoproteins. The distribution of LDL among large buoyant and small dense subtypes depends on various factors. The trait of sdLDL predominance or the atherogenic lipoprotein phenotype (high TGs, low HDL, and high sdLDL) is a distinct dyslipidemia linked to various genetic loci with a heritability of 35 to 45%. The prevalence of this trait is higher in older men and postmenopausal women. Other factors include diet, obesity, exercise, drugs, abdominal adiposity, and hormonal status (►Fig. 2).

However, the major factor determining LDL subtype distribution is the level of TGs. Studies have shown that pattern B subclass distribution is not seen till TG levels exceed 1.5 mmol/L or 120 mg/dL. Fifty percent of the variation in LDL size is determined by TG levels. When TG levels are high, it leads to increased synthesis of TG-rich VLDL1. Lipolysis of VLDL1 produces LDL particles with prolonged residence time of around 5 days in the circulation compared with LDL derived from smaller VLDL precursors which remain in the circulation for around 2 days. This smaller LDL has sufficient time to be remodeled by CETP; LDL loses cholesterol esters and gains TGs. If this happens to a certain degree, next exposure of TG-rich LDL to hepatic lipase (HL) promotes a particle shift to smaller denser range. Hence the levels of activity of CETP and HL also influence LDL subclass distribution (►Fig. 3).

sdLDL as a Drug Target

The importance of sdLDL as a predictive biomarker in CAD is well established and its atherogenic potential understood. Numerous studies are now being performed using different approaches of modifying the lipid profile. Instead of LDL, targeting sdLDL alone or along with high TGs and low HDL as a part of the atherogenic triad, can probably give a better outcome. Different approaches have been tested for reducing sdLDL with variable results.

Fibrates

Since high TGs are a direct influencing factor on sdLDL levels, TG reduction was one of the earliest approaches tried out.

**Fig. 2** Factors influencing SdLDL levels. CETP, cholesterol ester transfer protein; SdLDL, small dense low-density lipoprotein; TG, triglycerides; VLDL, very-low-density lipoprotein.
Fibrates are peroxisome proliferator activated receptor α agonists. They increase muscle and hepatic fatty acid oxidation and reduce lipogenesis. In addition, they increase lipoprotein lipase (LPL) activity and reduce VLDL levels through decreased synthesis and increased clearance. The effect is reduced TG levels, modest elevation of HDL, and shift of sdLDL to large buoyant LDL (lbLDL) while total LDL remains the same. Though apparently fibrates appear to be a good treatment modality, disappointing results were obtained in clinical trials in regard to CVD prevention. Drugs that have been tested include fenofibrate, bezafibrate, and gemfibrozil. Meta-analysis of the fibrate outcome trials has revealed that significant positive outcome is obtained in hypertriglyceridemic patients, who are but not in those who are normotriglyceridemic. At present, fibrates are being used as an add on to statins in high-risk patients with baseline high TG levels.

**Niacin**

Niacin reduces TG levels by decreasing TG synthesis and accelerating Apo B degradation leading to decreased secretion of VLDL, LDL, and sdLDL. High-dose niacin increases HDL. Immediate release of niacin causes flushing and worsens glycemic control in Type 2 DM patients. Slow release forms are hepatotoxic and less efficient in raising HDL levels. Extended release forms have favorable effect on the lipid profile without being hepatotoxic or worsening glycemic control. However, no additional benefit of adding niacin to statins has been demonstrated in any clinical trial for improving CVD outcome.

**CETP Inhibitors**

CETP mediates transfer of TGs from TG-rich lipoproteins to HDL and LDL in exchange for cholesterol esters. This is the main pathway of reverse cholesterol transport and hence, cholesterol breakdown and excretion through the liver. Use of CETP inhibitors, for example, torcetrapib increases HDL levels but also impairs reverse cholesterol transport, hence curbing its benefit. Until now good results have not been obtained using CETP inhibitors. Newer more potent drugs are being developed. Anacetrapib is one such drug which increases HDL, reduces LDL and Apo B 100, and increases the LDL TG/cholesterol ratio, LDL size, and Apo B 100 clearance.

**Statins**

Statins inhibit the enzyme HMG CoA reductase; hence they control the rate limiting step of cholesterol synthesis. They are highly effective in reducing LDL; they also reduce TGs and raise HDL. Rosuvastatin in high doses is highly effective in reducing sdLDL levels. Fukushima et al used atorvastatin to reduce LDL in patients with acute coronary syndrome and metabolic syndrome and found that the sdLDL reduction was 5.5 times greater than ACS patients without metabolic syndrome. Choi et al, however, reported that though statins reduce the absolute levels of both sdLDL and lbLDL, they are not effective in reducing the proportion of sdLDL to total LDL.

**Sodium-Glucose Co-transporter-2 Inhibitors**

Sodium-glucose co-transporter-2 (SGLT2) inhibitors are a group of drugs used in the treatment of Type 2 DM. They reduce blood glucose by inhibiting glucose re-absorption in the proximal tubules of the kidney. Clinical trials have revealed that SGLT-2 inhibitors reduce TGs and increase not only HDL but LDL as well. Hayashi et al in a study on the effect of the SGLT2 inhibitor dapagliflozin on the lipid profile of
Type 2 diabetics reported that dapagliflozin reduced the levels of sdLDL by 20% and increased LDL levels by 18% keeping the total LDL levels the same. They postulated that the reduction in sdLDL levels was brought about by reducing TG levels and increasing insulin sensitivity.35

**Future Prospects**

LDL reduction and subtype redistribution is an area of constant research. Intervventional methodologies are being developed which will target the primary pathophysiological aspects of the atherogenic dyslipidemia complex and will lead to HDL increase and LDL subclass modification as a secondary effect. Together this will lead to a better outcome for those patients at risk of CAD. Some of the ongoing research areas are discussed below.

LPL activity is one of the factors influencing sdLDL levels. In patients with homozygous loss of function mutations leading to severe hypertriglyceridemia, gene therapy has a good outcome.36 Strategies other than gene therapy are also being explored. APOC3 is an inhibitor of LPL. Inhibition of APOC3 using antisense therapy may reduce sdLDL levels.37 ANGPTL are another group of proteins which inhibit LPL. Antibody-mediated blockade of ANGPTL3 has given good outcome in animal models.38

Other methods being tried out include inhibition of certain enzymes of lipid metabolism. Monoaclglycerol acyl transferase and diacylglycerol acyl transferase are enzymes involved in TG synthesis.39 Acetyl CoA carboxylase catalyzes the rate limiting step in the fatty acid synthesis. Drug-mediated inhibition of these enzymes may decrease lipogenesis and increase lipid oxidation. Clinical trials are going on for these drugs.40

ETC-1002 (bempedoic acid) is another drug under investigation. It inhibits adenosine triphosphate (ATP) citrate lyase and activates AMP-activated protein kinase, thus reducing fatty acid synthesis and increasing its oxidation. A major benefit of this drug is that it reduces LDL levels regardless of the TG levels.41,42 However, further clinical trials are required to assess its efficacy and safety as a combination drug.27

Whatever the treatment strategy being developed in the future, the importance of lifestyle modification in lipid lowering should not be ignored. Diet, exercise, and maintenance of healthy body weight have a positive effect on atherogenic dyslipidemia. Improved-insulin sensitivity and hormonal balance which accompanies lifestyle changes are thought to bring about this effect.43-45

**Conclusion**

Small dense LDL is the atherogenic component of LDL. It is a predictive biomarker for CAD. Lipid lowering drugs are being routinely used as a treatment component in high-risk patients for primary and secondary prevention of CAD. But the outcome is not always favorable, in spite of significant LDL reduction. Targeting the atherogenic dyslipidemia complex stressing on sdLDL reduction may be the solution. Various drugs have been shown to be effective in reducing sdLDL. However, further research is necessary to draw clear guidelines for sdLDL reduction therapy in CAD treatment and prevention.

**Conflict of Interest**

None declared.

**References**


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