An Updated Review on Hyperlipidemia

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Abstract: Hyperlipidemia is a disease portrayed by an extension, in any incident, one of the plasma lipids, including triglycerides, cholesterol, cholesterol esters just as plasma lipoproteins including outstandingly low thickness and low thickness lipoproteins close by diminished high thickness lipoproteins levels. This stature of plasma lipids is among the leading risk factors related to cardiovascular illnesses. The present review focuses on the sorts of hyperlipidemia, lipid processing, the pathophysiology of hyperlipidemia, treatment frameworks for raised lipid profiles. In the interim statins and fibrates remain the major antihyperlipidemic administrators for the treatment. The fortune spot of the plant domain has different plants to treat this ailment. The indigenous course of action of the drug gives bountiful data about plants available for the treatment of hyperlipidemia. The review also lightens the role of nutraceuticals in hyperlipidemia therapy. The search was performed in PubMed, PsycInfo, Web of Science, Scopus, and Medline databases. A quality assessment yielded a total of 60 papers to be considered for the review by using strict inclusion and exclusion criteria.

Keywords: Lipoproteins, hyperlipidemia, hypercholesterolemia, atherosclerosis, and statin.

INTRODUCTION

CHOLESTEROL

Cholesterol was a waxy fat-like substance that was found in all cells of the body and in some of the foods. Cholesterol was a necessary nutrient for many functions including:

- Repairing the cell membranes.
- Construction of vitamin D on the skin’s surface
- Producing hormones such as that estrogen, testosterone
- Probably helping cell links in a brain that is essential for learning and memory.

Regardless of these benefits when cholesterol levels rise the blood and that can have dangerous consequences depending on the type of cholesterol and its production stimulated by saturated fat [1].

LIPOPROTEIN

Lipids are insoluble in the blood (plasma), they must be transported to the cells by special carriers called lipoproteins. Lipoproteins are spherical particles of high molecular weight. Each lipoprotein particle contains a non-polar core and a hydrophilic surface. The hydrophilic surface makes the lipoprotein soluble in plasma and acts as an interface between the plasma and lipid core. The center comprises of hydrophobic lipids, triglycerides, and cholesterol esters, encompassed by a hydrophilic surface layer of phospholipids, unesterified cholesterol, and explicit proteins named apolipoproteins or apoproteins. The apolipoproteins provide structural integrity to the lipoproteins and determine the lipoprotein’s metabolic fate by serving as binding sites for receptors and activating enzymes involved in lipid metabolism [2].

LIPID METABOLISM

Almost all the dietary fats are ingested from the intestinal lumen into the intestinal lymph, what’s more, swollen into chylomicrons. These lipoproteins shift into the circulation system where they got hydrolyzed by endothelial
lipoprotein lipase which hydrolyzes the triglyceride into glycerol and nonesterified unsaturated fats. After which the chylomicron remainders are invested in the liver and bundled with cholesterol, cholesteryl esters, and ApoB100 to shape VLDL. After the arrival of VLDL into the circulation system, it will be changed over into IDL by the activity of lipoprotein lipase and hepatic lipase, where phospholipids and apolipoproteins moved back to HDL. Besides, after the hydrolysis by hepatic lipase, IDL will be altered over to LDL and disaster more apolipoproteins [2]. Cholesterol is come back to the liver by the turn around the cholesterol transport pathway utilizing HDLs which are initially orchestrated by the liver, what's more, discharged into the blood. In the blood, HDL cholesterol is esterified by LCAT to cholesteryl ester, besides, moved to VLDL and chylomicrons to return to the liver through the LDL receptor. Cholesteryl ester is moved to LDL particles by CETP and afterward exposed to LDL-receptors interceded endocytosis. At long last, cholesteryl esters are hydrolyzed to cholesterol and separated from the body as bile acid [3].

### CLASSIFICATION OF LIPOPROTEIN

The cholesterol along with some other types of fats cannot be dissolved in the blood. Moreover so as to be shipped to and from cells, they must be particularly conveyed by specific particles called lipoproteins, which comprise an external layer of protein with an internal center of cholesterol and triglycerides [4, 5]. In expansion, the lipoproteins have been discovered fundamental for cholesterol to move around the body. The lipids can be classified as TC, triglycerides, LDL, HDL, and very low density lipoprotein (VLDL) cholesterol.

#### Total Cholesterol

As per rules of the National Cholesterol Education Program (NCEP), TC fixation underneath 200 mg/dL has been considered as attractive, while, focuses more prominent than 240 mg/dL are near as hyperlipidemia. In any case, epidemiological proof recommends that the danger of heart occasions diminishes as TC levels fall roughly to 150 mg/dL. Also, TC ought to be under 180 mg/dL for kids [6, 7].

#### Triglyceride

Triglycerides are another kind of fat that is conveyed in the blood by VLDL. The abundance of calories, liquor, or sugar in the body gets changed over into triglycerides and put away in fat cells all through the body [8]. The triglyceride focus under 150 mg/dL is viewed as ordinary, though, centralizations of 200-499 mg/dL are considered as high. Besides, centralizations of 500 mg/dL or higher are viewed as hazardous for the turn of events and movement of different CVDs [9].

#### LDL cholesterol

LDL is normally known as the terrible cholesterol, which is created by the liver and conveys cholesterol and different lipids from the liver to various territories of the body like muscles, tissues, organs, and heart. The significant levels of LDL demonstrate considerably more cholesterol in the circulatory system than would normally be appropriate and consequently, increment the danger of heart disease [8, 10]. As indicated by NCEP rules, LDL cholesterol fixations underneath 100mg/dL are viewed as ideal, while focuses on the scope of 160-189 mg/dL are considered to the higher side. In any case, expanding proof backings that typical human LDL cholesterol fixation can be as low as 50 to 70 mg/dL [11]. Besides, it has been exhaustively observed that the danger of CVDs diminishes as LDL cholesterol fixation diminishes.

#### HDL cholesterol

HDL is ordinarily alluded to as the great cholesterol, which is delivered by the liver to convey cholesterol and different lipids from tissues back to the liver for degradation [10]. Significant levels of HDL cholesterol have been considered as a decent marker of a sound heart. The centralizations of 60 mg/dL or higher have been considered as ideal, though, HDL concentrations underneath 40 mg/dL are considered as a significant hazard factor for CVDs. Notwithstanding, HDL is regularly deciphered with regards to TC and LDL focuses, and subsequently might be viewed as less critical when LDL is low [8, 10].

#### VLDL Cholesterol

VLDL is like LDL cholesterol as it contains for the most part fat and very little protein. VLDL cholesterol is the lipoproteins that convey cholesterol from the liver to organs and tissues in the body [11]. They are shaped by a blend of cholesterol and triglycerides. Additionally, VLDLs are heavier than LDL and are likewise connected with atherosclerosis and heart disease [12].

### ENZYMES INVOLVED IN LIPOPROTEIN TRANSPORT & METABOLISM

There are several enzymes and transfer proteins that play an important role in lipoprotein metabolism. Some of the enzymes and their functions are mentioned in Table 1.
Table 1: Importance Of Enzymes In Lipid Transport & Metabolism [14, 15]

<table>
<thead>
<tr>
<th>Enzymes</th>
<th>Functions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lipoprotein lipase</td>
<td>Hydrolysis of TG rich particles some phospholipase activity activated by APO C-II</td>
</tr>
<tr>
<td>Hepatic lipase</td>
<td>Hydrolysis of Tri-, Di- and mono acylglycerols, Acyl-CoA thioester and phospholipids conversion of HDL2 to HDL1 activated by APO A-III</td>
</tr>
<tr>
<td>Pancreatic lipase</td>
<td>Hydrolysis of FAs at positions 1 and 3 of emulsified TGs in the intestine.</td>
</tr>
<tr>
<td>Lecithin cholesterol</td>
<td>Catalysis of lecithin with cholesterol to give lysolecithin and cholesteryl ester activated by APO A-I and APO c-I.</td>
</tr>
<tr>
<td>Acyltransferase LCAT</td>
<td>Esterification of cholesterol in the intestinal lumen</td>
</tr>
<tr>
<td>Pancreatic cholesterol esterase</td>
<td>Esterification of cholesterol within the cells</td>
</tr>
<tr>
<td>Acyl co A-cholesterol acyl transferase ACAT</td>
<td>Esterification of cholesterol within the cells</td>
</tr>
<tr>
<td>Cholesterol ester transfer protein CETP</td>
<td>Transfers esterified cholesterol from HDLs to VLDLs and LDLs.</td>
</tr>
<tr>
<td>HMG Co-A reductase</td>
<td>Rate limiting enzyme of cholesterol synthesis</td>
</tr>
</tbody>
</table>

HYPERLIPIDEMIA

Hyperlipidemia is a condition when abnormally high levels of lipids i.e. The fatty substance are found in the blood. This situation is also known as hypercholesterolemia [16]. The human body is complex machinery and for maintaining the homeostasis of various organ and organ systems. Any undesirable change will disturb the balance resulting in a diseased state. Lipids are fats in the circulatory system, usually separated into cholesterol and triglycerides. Cholesterol flows in the circulatory system and is engaged with the structure and capacity of cells. Triglycerides (TG) are best seen as a vitality that is either utilized promptly or put away in fat cells [17] tg is fabricated in the liver from the nourishments or by being consumed from the digestive system. Virchow in 19th century who identified cholesterol crystals in the atherosclerotic lesion and stated that endothelial cell injury initiates atherogenesis in a modification of this hypothesis, it was proposed that the endothelium normally influences the behavior of arterial smooth muscle cells by providing a barrier to the passage of plasma proteins and that the major effect of hemodynamic or other factors that injure endothelium is to reduce the effectiveness of the barrier [18]. Arteries are normally smooth and unobstructed on the inside, but in case of increased lipid level, a sticky substance called plaque is formed inside the walls of arteries. This leads to reduced blood flow, leading to stiffening and narrowing of the arteries. It has been proved that elevated plasma levels of cholesterol and LDL are responsible for atherosclerosis in man, and epidemiological data suggest that elevated plasma levels of HDL have a protective effect [19, 20].

CLASSIFICATION

TYPES OF HYPERLIPIDEMIA

Hyperlipidemia can be broadly divided into:
1. Primary hyperlipidemia is due to Single gene defect: It is familial and called monogenic or genetic. Polygenic gene defect, Multiple genetic defects, dietary, and physical activity are caused due to it.

Table 2: Disorder

<table>
<thead>
<tr>
<th>TYPE</th>
<th>DISORDER</th>
<th>CAUSE</th>
<th>OCCURRENCE</th>
<th>ELEVATED PLASMA LIPOPROTEIN</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Familial lipoprotein lipase</td>
<td>Genetic</td>
<td>Very rare</td>
<td>Chylomicrons</td>
</tr>
<tr>
<td>Ia</td>
<td>Familial hypercholesterolemia</td>
<td>Genetic</td>
<td>Less common</td>
<td>LDL</td>
</tr>
<tr>
<td>Ib</td>
<td>Polygenic hypercholesterolemia</td>
<td>Multifactorial</td>
<td>Commonest</td>
<td>LDL</td>
</tr>
<tr>
<td>II</td>
<td>Familial Dysbetalipoproteinemia</td>
<td>Genetic</td>
<td>Rare</td>
<td>IDL, Chylomicrons Remnants</td>
</tr>
<tr>
<td>III</td>
<td>Hypertriglyceridemia</td>
<td>Multifactorial</td>
<td>Common</td>
<td>VLDL</td>
</tr>
<tr>
<td>V</td>
<td>Familial combined hyperlipidemia</td>
<td>Genetic</td>
<td>Less common</td>
<td>VLDL, LDL</td>
</tr>
</tbody>
</table>

2. SECONDARY HYPERLIPIDEMIA

It is associated with diabetes, myxoedema, nephritic syndrome, chronic alcoholism, with the use of drugs like corticosteroids, oral contraceptives, Beta-blocker [22].
Type 1 Hypercholesterolemia

**REASONS**

**Hypothyroidism**
- Anorexia nervosa
- Acute intermittent porphyria
- Obstructive liver disease
- Nephrotic syndrome
- Drugs: Progestins, thiazide, diuretics, glucocorticoids, beta-blockers, isotretinoin, protease inhibitors, cyclosporine, mirtazapine, sirolimus.

**Hypertriglyceridemia**
- Obesity
- Pregnancy
- Lipodystrophy
- Acute hepatitis
- Diabetes mellitus
- Ileal bypass surgery
- Glycogen storage disease
- Systemic lupus erythematosus
- Monoclonal gammopathy: multiple myeloma
- Drugs: Alcohol, estrogens, isotretinoin, beta-blockers, glucocorticoids, bile-acid resins, thiazides, asparaginase,azole antifungals, mirtazapine, anabolic steroids, sirolimus, bexarotene.

Hypocholesterolemia
- Malnutrition
- Malabsorption
- Chronic liver disease
- Myeloproliferative diseases Chronic infectious diseases: AIDS, tuberculosis

Low HDL
- Malnutrition
- Obesity
- Drugs: anabolic steroids, probucol, isotretinoin, progestins.

**CAUSES OF HYPERLIPIDEMIA [23]**
1. Way of life propensities or treatable ailments. Way of life patrons incorporate stoutness, not working out, and smoking
2. Cushing’s Syndrome
3. Kidney illness
4. Pregnancy
5. An underactive thyroid organ
6. Ecological and hereditary elements
7. Liquor consumption
8. Monoclonal Gammopathy
9. Nephrotic Syndrome
10. Obstructive Jaundice
11. Decrease in thyroidism
12. Diabetes (type 2).
13. Anorexia Nervosa
14. Drugs • Thiazide Diuretics • Ciclosporin • Glucocorticoids • Beta Blockers • Retinoic Acid
15. High dietary straight forward sugars
16. Estrogen treatment

**PATHOPHYSIOLOGY OF HYPERLIPIDEMIA**
LDL cholesterol ordinarily courses in the body for 2.5 days, and in this manner ties to the LDL receptor on the liver cells, experiences endocytosis, and is processed. LDL is excluded, and the arrangement of cholesterol by the liver is smothered in the HMG-CoA reductase pathway [24]. FH, LDL receptor work is diminished or missing, and LDL flow for a normal length of 4.5 days, bringing about essentially expanded degree of LDL cholesterol in the blood with ordinary degrees of different lipoproteins [25]. In changes of ApoB, the decreased power of LDL elements to the receptor...
causes the stretched degree of LDL cholesterol. It isn’t known how the change causes LDL receptor dysfunction in changes of PCSK9 and ARH is a pathologic procedure described by the aggregation of lipids, cholesterol, and calcium and t]. In spite of the fact that atherosclerosis happens in a specific way in all individuals, FH patients may create quickened atherosclerosis because of the overabundance level of LDL. The level of atherosclerosis roughly depends on the number of LDL receptors despite everything communicated and the usefulness of these receptors. In various heterozygous types of FH, the receptor effort is just somewhat injured, and LDL levels will stay quite low. In the more genuine homozygous structures, the receptor isn’t communicated at all [26]. A few investigations of FH associates recommend that extra chance variables are for the most part having an effect on everything when an FH persistent create atherosclerosis [27]. Notwithstanding the exemplary hazard factors, for example, smoking, high blood weight, and diabetes, hereditary examinations have appeared that a typical variation from the norm in the prothrombin quality (G20210A) builds the danger of cardiovascular occasions in patients with FH [28]. A few examinations create that a considerable level of lipoprotein(a) was an extra risk cause for ischemic coronary illness [29]. The dangerous situation was moreover seen as higher in patients with a particular genotype of the angiotensin-changing over chemical (ACE) [30, 31].

**COMPLICATIONS**

**Inconveniences of hyperlipidemia**

**Atherosclerosis**

Hyperlipidemia is the most significant hazard factor for atherosclerosis, which is a significant reason for cardiovascular sickness. Atherosclerosis the advancement of sinewy plaques inside the dividers of enormous and medium arteries [32].

**Coronary Artery Disease (CAD)**

Atherosclerosis, the significant reason for coronary corridor illness, portrayed by the collection of lipid and the arrangement of sinewy plaques inside the mass of the conduits bringing about narrowing of the veins that gracefully blood to the myocardium, and results in restricting bloodstream and deficient measures of oxygen to address the issues of the heart. Raised lipid profile has been associated with the advancement of coronary atherosclerosis [33].

**Myocardial Infarction (MI)**

MI is a condition that happens when blood and oxygen supplies are in part or totally hindered from streaming in at least one cardiovascular courses, bringing about harm or passing of heart cells. The impediment might be because of burst atherosclerotic plaque. The investigations show that around one-fourth of overcomers of myocardial dead tissue was hyperlipidemic [33].

**Ischemic stroke**

Stroke is the fourth driving reason for death. Typically, strokes happen because of blockage of a course by blood coagulation or a bit of atherosclerotic plaque that loosens up in a little vessel inside the mind. Numerous clinical preliminaries uncovered that bringing down of low-thickness lipoprotein and all-out cholesterol by 15% altogether diminished the danger of the main stroke [34].

**CLINICAL MANIFESTATION**

The National Cholesterol Education Program suggests that a fasting lipoprotein profile (FLP) including absolute cholesterol, LDL, HDL, and triglycerides ought to be estimated in all grown-ups 20 years old or more established at any rate once like clockwork [35].

Estimation of plasma cholesterol (which is about 3% lower than serum judgments), triglyceride, and HDL levels following a 12-hour or longer quick is significant, in light of the fact that triglycerides might be raised in nonfasted people; total cholesterol is just humbly influenced by fasting [35].

Two conclusions, 1 to about two months separated, with the patient on a steady eating regimen and weight, and without intense sickness, are prescribed to limit changeability and to acquire a solid benchmark. In the event that the all-out cholesterol is more prominent than 200 mg/dL, a subsequent assurance is suggested, and if the qualities are in excess of 30 mg/dL separated, the normal of three qualities ought to be utilized (Barbara et al., 2005). After a lipid irregularity is affirmed, significant parts of the assessment are the history (counting age, sex, and, if female, menstrual and estrogen substitution status), physical assessment, and research center examinations.

A total history and physical assessment ought to survey (1) nearness or nonattendence of cardiovascular hazard factors or unequivocal cardiovascular sickness in the individual; (2) family ancestry of untimely cardiovascular malady or lipid issue; (3) nearness or nonappearance of auxiliary reasons for hyperlipidemia, including simultaneous meds; and (4) nearness or nonappearance of xanthomas, stomach agony, or history of pancreatitis, renal or liver ailment, fringe
vascular infection, stomach aortic aneurysm, or cerebral vascular ailment (carotid bruits, stroke, or transient ischemic assault).

Diabetes mellitus is currently viewed as a CHD chance proportional. That is, the nearness of diabetes in patients without known CHD is related to a similar degree of hazard as patients without diabetes yet having affirmed CHD [35].

In the event that the physical assessment and history are lacking to analyze a familial issue, at that point agarose-gel lipoprotein electrophoresis is helpful to figure out which class of lipoproteins is influenced. On the off chance that the triglyceride levels are underneath 400 mg/dL and neither sort III hyperlipidemia nor chylomicrons are identified by electrophoresis, at that point, one can compute VLDL and LDL fixations: VLDL = triglyceride/5; LDL = absolute [35].

Cholesterol - (VLDL + HDL). Beginning testing utilizes absolute cholesterol for case finding, yet ensuing administration choices ought to be founded on LDL.

Since all out cholesterol is made out of cholesterol got from LDL, VLDL, and HDL, assurance of HDL is valuable when all-out plasma cholesterol is raised. HDL might be raised by moderate liquor ingestion (less than two beverages for each day), physical exercise, smoking suspension, weight reduction, oral contraceptives, phenytoin, and terbutaline. HDL might be brought down by smoking, stoutness, an inactive way of life, and medications, for example, ß blockers [36].

Conclusion of lipoprotein lipase inadequacy depends on low or missing protein movement with typical human plasma or apolipoprotein C-II, a cofactor of the catalyst.

To have an okay of coronary illness, the lipid levels must be kept up as follows:
1. LDL under 130 mg/dL or < 70 on the off chance that you have set up the conclusion of diabetes
2. HDL more prominent than 40 mg/dL (men) or 50 mg/dL (ladies)
3. Complete cholesterol under 200 mg/dL and
4. Triglycerides under 200 mg/dL or 150 on the off chance that you have set up coronary illness or diabetes.

<table>
<thead>
<tr>
<th>S. NO</th>
<th>TEST NAME</th>
<th>NORMAL VALUES</th>
<th>INDICATORS</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Total Cholesterol</td>
<td>Total Cholesterol: &lt; 200 mg/dL (desirable) (&lt; 180</td>
<td>200-239 mg/dL = Borderline High (borderline</td>
</tr>
<tr>
<td></td>
<td></td>
<td>optimal)</td>
<td>mg/dL = Hypercholesterolemia)</td>
</tr>
<tr>
<td>2</td>
<td>Total Cholesterol</td>
<td>&lt; 180 mg/dL</td>
<td>&gt; 180 mg/dL may lead to premature</td>
</tr>
<tr>
<td></td>
<td>for children</td>
<td></td>
<td>Atherosclerosis</td>
</tr>
<tr>
<td>3</td>
<td>Triglyceride Levels</td>
<td>Less than 150 mg/dL</td>
<td>150-199 mg/dL is Borderline High 200-499 mg/dL</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>is High 500 mg/dL or above is very High</td>
</tr>
<tr>
<td>4</td>
<td>HDL Cholesterol</td>
<td>≥ 60 mg/dL is desirable</td>
<td>In general, HDL levels &lt; 40 mg/dL increases</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>risk for CHD. Women with levels &lt; 47 mg/dL</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>and men &lt; 37 mg/dL have increased risk.</td>
</tr>
<tr>
<td>5</td>
<td>LDL Cholesterol</td>
<td>&lt; 100 mg/dL (optimal) 100-129 mg/dL (near</td>
<td>130-159 mg/dL Borderline High 160-189 mg/dL</td>
</tr>
<tr>
<td></td>
<td></td>
<td>optimal/above optimal)</td>
<td>mg/dL High ≥190 mg/dL Very High</td>
</tr>
<tr>
<td>6</td>
<td>C-reactive Protein</td>
<td>CRP &lt; 1 mg/dl</td>
<td>CRP &gt; 1 mg/dl (&gt; 10 mg/dl often seen) suggests</td>
</tr>
<tr>
<td></td>
<td>(CRP)</td>
<td></td>
<td>inflammation</td>
</tr>
<tr>
<td>7</td>
<td>VLDL Cholesterol</td>
<td>The VLDL normal range is between 0–40 mg/dL and the</td>
<td>&gt; 40 suggests can increase the risk of</td>
</tr>
<tr>
<td></td>
<td></td>
<td>recommended optimum range is between 0–30 mg/dL</td>
<td>developing heart disease</td>
</tr>
</tbody>
</table>

PHARMACOLOGICAL TREATMENT
Many hypolipidemic synthetic drugs are available in the market for the treatment of hyperlipidemia and other associated disease [37]. The existing hypolipidemic drugs with there mechanism of action use and side effects are listed in Table 4.
Table 4: Drugs Treatment

<table>
<thead>
<tr>
<th>Categories</th>
<th>Drugs</th>
<th>MOA</th>
<th>Use</th>
<th>Side effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>HMG-CoA reductase inhibitors</td>
<td>Lovastatin</td>
<td>HMG-CoA reductase catalyzes the synthesis of mevalonic acid from HMG-CoA and is the rate-limiting step in cholesterol biosynthesis Leads to the up-regulation of LDL receptors in the liver.</td>
<td>Great for all hyperlipidemia involving increased levels of LDL or cholesterol Atherosclerosis; stroke prevention Primary prevention of CADV</td>
<td>Headache Nausea sleep disturbance elevations in hepatocellular enzymes and alkaline phosphatase. CI in hepatic dysfunction</td>
</tr>
<tr>
<td></td>
<td>Atorvastatin</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Rosuvastatin</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bile acid sequestrants</td>
<td>Cholestyramine</td>
<td>Anion-exchange resin – binds bile acids in intestinal lumen preventing enterohepatic circulation (this increases excretion of bile which is made from cholesterol), this causes an up-regulation of hepatic LDL receptors and increased production of cholesterol.</td>
<td>Uses hyperlipidemias involving isolated increases of LDL.</td>
<td>Constipation Flatulence Dyspepsia Hypertriglyceridemia Hyperchloremic acidosis</td>
</tr>
<tr>
<td></td>
<td>Colestipol</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Colesevelam</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fibrates</td>
<td>Gemfibrozil,</td>
<td>activation of nuclear transcription receptor to increase LPL synthesis (removes TGs from lipoproteins); enhanced removal of VLDL from plasma</td>
<td>DOC fro type III lipoproteinemia (familial dysbetalipoproteinemia) • Hypertriglyceridemias • useful for Pruritus in biliary obstruction</td>
<td>Skin rash, gastrointestinal (nausea, bloating, cramping) myalgia; lowers blood cyclosporine levels; potentially nephrotoxic in cyclosporine treated patients. Avoid in patients with CrCl &lt;30 mL/min. (Fenofibrate)</td>
</tr>
<tr>
<td></td>
<td>Fenofibrate,</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Micronized</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nicotinic acid (Niacin)</td>
<td>Niacin</td>
<td>It is a potent inhibitor of lipolysis in adipose tissues which decreases mobilization of FFAs (a major precursor of TGs) to the liver which in turn decreases VLDL (after few hours) Increases HDL levels.</td>
<td>Uses hyperlipidemias with very high VLDL and LDL. • Patients with very low HDL.</td>
<td>dry skin myositis Prostaglandin-mediated cutaneous flushing, warm sensation Headache Pruritus, Nausea, Vomiting, diarrhea hyperpigmentation Decreased glucose tolerance Hepatotoxicity (check AST, ALT levels) Rhabdomyolysis Hyperuricemia</td>
</tr>
<tr>
<td>cholesterol absorption inhibitors</td>
<td>Ezetimibe</td>
<td>• Localizes at the brush border, selectively inhibits intestinal absorption of cholesterol and related sterols (only blocks exogenous sterol intake)</td>
<td>Used in hypercholesterolemia a together with statins &amp; diet regulation.</td>
<td>Diarrhea Abdominal pain CI liver dysfunction</td>
</tr>
<tr>
<td>PCSK9 inhibitor</td>
<td>Alirocumab</td>
<td>Proprotein exchange subtilisin Kexin 9 (PCSK9) is a serine protease formed predominantly in the liver that goes ahead to the degradation of hepatocyte LDL receptors and improved LDL-C levels</td>
<td>• This group of lipid-lowering therapy appears hopeful in a variety of clinical situations.</td>
<td>the effect is injection site reaction</td>
</tr>
<tr>
<td></td>
<td>Evolocumab</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
NON PHARMACOLOGICAL TREATMENT

The targets of dietary treatment are to diminish the admission of complete fat, soaked unsaturated fats (i.e., immersed fat), and cholesterol logically and to accomplish an alluring body weight. This includes:

1. Concentrated immersed fat admission to 7 percent of everyday calories.
2. Lessen complete fat admission to 25 to 35 percent of everyday calories.
3. Constrained dietary cholesterol to fewer than 200 mg for each day.
4. Eating 20 to 30 g daily of dissolvable fiber, which is found in oats, peas, beans, and certain natural products; and
5. Expanded admission of plant stanols or sterols, substances found in nuts, vegetable oils, corn, and rice, to 2 to 3 g day by day. Different nourishments that can help control cholesterol incorporate cold-water fish, for example, mackerel, sardines, and salmon. These fish contain omega-3 unsaturated fats that may bring down triglycerides. Soybeans found in tofu and soy nuts and numerous meat substitutes contain an incredible cell reinforcement that can bring down LDL [38].

HERBS USED AS ANTIHYPERLIPIDEMIC ACTIVITY

Herbal medicines have been the chief source of primary health care worldwide. Since ancient times, plants have been cooking as a rich source of effective and safe medicines. Most of the world's population still relies on traditional medicines. A list of some medicinal plants which have antihyperlipidemic activity is mention in table [39-51].

Table 5: Herbs Used As Antihyperlipidemic Activity

<table>
<thead>
<tr>
<th>S. NO</th>
<th>PLANTS NAME</th>
<th>BIOLOGICAL SOURCE</th>
<th>PARTS USED</th>
<th>ROLE IN HYPERLIPIDEMIA</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Basil Ocimum basilicum</td>
<td>Lamiaceae</td>
<td>Leaf</td>
<td>Lowering LDL and total cholesterol, an increase of HDL</td>
</tr>
<tr>
<td>2</td>
<td>Blueberry Vaccinium corymbosum</td>
<td>Ericaceae</td>
<td>Fruits</td>
<td>Lowering triglyceride and LDL levels</td>
</tr>
<tr>
<td>3</td>
<td>Celery Apium celeri</td>
<td>Family Apiaceae</td>
<td>Seed</td>
<td>Decreasing serum triglyceride, total cholesterol, LDL-C, and hepatic triglyceride</td>
</tr>
<tr>
<td>4</td>
<td>Dandelion Taraxacum officinale</td>
<td>Compositae</td>
<td>Root leaves</td>
<td>Decreasing serum triglyceride, total cholesterol, LDL-C and increasing HDL-C</td>
</tr>
<tr>
<td>5</td>
<td>Dill Anethum graveolens</td>
<td>Apiaceae</td>
<td>Dried ripe fruit</td>
<td>Decreasing serum triglyceride</td>
</tr>
<tr>
<td>6</td>
<td>Evening primrose oil</td>
<td>Oenothera biennis</td>
<td>Root, herb.</td>
<td>Decreasing serum triglyceride, total cholesterol</td>
</tr>
<tr>
<td>7</td>
<td>Fenugreek Rigonella foenum</td>
<td>family Fabaceae.</td>
<td>Seeds and leaves</td>
<td>Decreasing serum triglyceride, total cholesterol, LDL-C</td>
</tr>
<tr>
<td>8</td>
<td>Ginger Zingiber officinale</td>
<td>Zingiberaceae,</td>
<td>Rhizomes</td>
<td>Decreasing serum LDL-C and increasing HDL-C</td>
</tr>
<tr>
<td>9</td>
<td>Ginseng Ginsing panax</td>
<td>Araliaceae</td>
<td>Dried root</td>
<td>Decreasing serum triglyceride, total cholesterol, LDL-C and increasing HDL-C</td>
</tr>
<tr>
<td>10</td>
<td>Grape Vites vinifera</td>
<td>Vitaceae</td>
<td>Fruits</td>
<td>Lowering oxidized LDL and elevate HDL-C level</td>
</tr>
<tr>
<td>11</td>
<td>Green tea Camellia sinensis</td>
<td>Myrtaceae</td>
<td>Leaves</td>
<td>Lowers the adiposity and affects the appearance of lipid metabolism genes</td>
</tr>
<tr>
<td>12</td>
<td>Nigella Nigella Sativa</td>
<td>Ranunculaceae</td>
<td>Seed</td>
<td>The decrease in triglyceride and increase in HDL-C</td>
</tr>
<tr>
<td>13</td>
<td>Psyllium Plantago Psyllium</td>
<td>Plantaginaceae</td>
<td>Ripe seeds</td>
<td>Decrease in LDL</td>
</tr>
<tr>
<td>14</td>
<td>Aswagandha Withania Somnifera</td>
<td>Solanaceae</td>
<td>Root</td>
<td>help improve heart health by reducing cholesterol and triglyceride levels</td>
</tr>
<tr>
<td>15</td>
<td>Indigo Berry Randia dumetorum</td>
<td>Rubiaceae</td>
<td>Fruit</td>
<td>a decrease in total cholesterol level in serum at 18 hours and 24 hour</td>
</tr>
<tr>
<td>16</td>
<td>Guggul Commiphora wigatti</td>
<td>Burseraceae</td>
<td>Gum resin</td>
<td>a decrease in total cholesterol level in serum at 18 hours and 24 hour</td>
</tr>
<tr>
<td>17</td>
<td>Berberis Berberis aristata</td>
<td>Berberaceae</td>
<td>Root resin</td>
<td>decrease in lipid profile in serum when compared to the dyslipidaemic HFD control</td>
</tr>
<tr>
<td>18</td>
<td>Black piper Piperum nigrum</td>
<td>Piperaceae</td>
<td>Fruit</td>
<td>It begins the digestion route by helping quick break down of better fat molecules into easily digestible simple molecules and prevents the growth of fat in the body.</td>
</tr>
<tr>
<td>19</td>
<td>snake jasmine Rhinacanthus nasatus</td>
<td>Acanthaceae</td>
<td>Whole plant</td>
<td>lipoprotein level decreased</td>
</tr>
<tr>
<td>20</td>
<td>Liquorice Glycyrrhiza Glabra</td>
<td>Fabaceae</td>
<td>Root</td>
<td>decrease in lipid profile in serum when compared to the dyslipidaemic HFD control</td>
</tr>
</tbody>
</table>
ROLE OF NUTRACEUTICALS IN HYPOLIPIDEMIC THERAPY

Nutraceutics is a region of pharmacology concerning food segments or dynamic fixings that might be utilized as helpful specialists. This incorporates countless mixes, for example, a functioning fixing, food supplements (i.e., supplements the ordinary eating routine), and useful nourishments (i.e., food sources advanced with parts with explicit helpful or defensive capacities), just as arrangements dependent on restorative herbs. Most mixes are vegetable begun, however, there are additional substances with creature source (e.g., fish oil). Ongoing examinations have demonstrated promising outcomes for these medications in different neurotic confusions, for example, diabetes, atherosclerosis, cardiovascular infections, malignancy, and neurological issue. There is an assortment of nutraceuticals with a potential lipid-bringing down impact, and in this manner helps in cardiovascular ailment anticipation and treatment [52].

Policosanols

These are a blend of regular long-chain aliphatic alcohols that got from a wide assortment of plants. It has been recommended that policosanols may hinder the movement of HMG-CoA reductase, however, this isn't completely affirmed. Various clinical examinations proposed a lipid-bringing down impact of policosanols in various kinds of decrease in LDL-cholesterol like that of statins (about 25%) and a 10% expansion of HDL-C [53].

Polyphenols

This is an exceptionally enormous group of substances accessible in the plant world. The fundamental element is the nearness of different phenolic bunches having a strong enemy of oxidant impact. It has additionally been hypothesized that polyphenols can restrain HMG-CoA reductase, just as ACAT2 and MTP, supporting a hypocholesterolemic effect [54].

Probiotics

Probiotics could bring down cholesterol ingestion through direct cell impacts or intervene by bile metabolism [55]. A few examinations on various patients have archived huge decreases in all-out cholesterol, up to 40% [56, 57]. Despite, accurate components of activity were not distinguished, More as of late, it has been proposed a job of the outflow of the quality of bile salt hydrolase movement in the lactobacilli strains to clarify the cholesterol-bringing down activity, regardless of whether this theory itself doesn't appear to be totally convincing [58].

Guggul

This is a tar extricated from the bark of Commiphora Mukul, a little prickly tree, otherwise called the tree of myrrh he dynamic segments, guggulsterone E and Z, have been exhibited to have an adversarial activity of FXR, an atomic receptor engaged with the bile digestion.

Plant Sterols

Plant sterols decay intestinal maintenance of cholesterol through The lessening of the substance of cholesterol inside the micelles and a subsequent lower degree of absorbable cholesterol [59]. Besides, a couple of examinations suggest that phytosterols can fight with cholesterol in the transporter of the intracellular incorporation (NPC1L1), and moreover increase the activity of transmembrane proteins at risk for the release of cholesterol (ABCA1) and plant sterols (ABCG5 and ABCG8) in the stomach related framework and liver with the net effect of extending the appearance of the two sterols into the intestinal lumen by enterocytes and in the bile conductors in the liver [60, 61]. The lower intestinal ingestion of cholesterol-incited by plant sterols reduces cholesterol pool of liver, which response by growing the affirmation of LDL receptors, finally realizing higher take-up of plasma LDL and as such in a net hypocholesterolemic sway.

Fermented Red Rice

The maturing of red rice by a parasite (Monascus purpureus). Produces a substance called monacolin K, which thwarts the amalgamation of cholesterol [62]. The monacolin K is in any case called lovastatin, atstatine open in the market far and wide. The red yeast rice in like manner conveys diverse monacolins that may update the restriction of HMG CoA reductase. Moreover, late data show that stood out from the old-style lovastatin, monacolin K removed from red yeast rice have even a higher bioavailability with a higher sufficiency at the comparable dosage [63].

Berberine

This substance, with an intolerant taste and excellent yellow concealing, is nearby in the bark, roots, and stems, including the underground (rhizome) of plants of the family Berberis, for instance, barberry (Berberis vulgaris L.). Of late, most thought has been on the metabolic properties of berberine. In 2004, Konge, have demonstrated that berberine diminished plasma cholesterol by 29%, triglycerides by 35%, and LDL cholesterol by 25%, however, it didn't alter HDL-cholesterol levels [64]. Berberine extends the number of LDL receptors on the hepatic cell surface, correspondingly to
statins. During statin treatment, the presentation of LDL-receptors on cell layers follows the reducing of the endogenous cholesterol association and the subsequent abatement of intracellular cholesterol pool, however, the action of berberine has all the earmarks of being associated with the ability to spoil a protein (PCSK9) at risk for the poor adulteration of LDL receptors in the liver. In the future, berberine may have synergistic contacts with statins [65, 66].

**CONCLUSION**

Hyperlipidemia has been influencing humankind for a very long time. In spite of the fact that numerous medications are accessible to treat Hyperlipidemia, the issue of upgraded cholesterol levels in the blood is as yet common and is being a reason for some coronary issues. As of late, certain synthetic compounds (drugs) of the common beginning have seen light in rewarding these cholesterol levels, lessening the danger of coronary failures. To diminish the hazard because of hyperlipidemia, the different ways are to follow a solid way of life and medications that can decrease the cholesterol and triglyceride levels in the blood.

**REFERENCES**


