

# Lipid Profile Alterations in the Metabolic Syndrome the Latest Biochemical and Physiological Aspect

Dr. Ashfaq ul Hassan<sup>1</sup> and Dr. Shazia Nazir<sup>2</sup>

<sup>1</sup> SKIMS

Received: 5 April 2015 Accepted: 3 May 2015 Published: 15 May 2015

---

## Abstract

Cardiovascular and cerebrovascular complications form a major challenge in the management of the The Metabolic Syndrome. The risks of development are compounded by simultaneous occurrence of Type 2 diabetes mellitus (T2DM), dyslipid emias, hypertension and other medical conditions. The significant increase in morbidity, mortality along with premature death and disabilities resulting from these conditions in both developed and developing is a matter of great concern. The article makes a microanalysis of Biochemical alterations associated with dyslipidemias in cases of metabolic syndrome.

---

**Index terms**— apolipoproteins, esterification, syndrome, circumference, waist, glucocorticoids, lipase. increased waist circumference, hypertriglyceridermia, low high density lipoprotein (HDL)cholesterol, hypertension (130/85mmHg) and fasting glucose of 110 mg/dl or high.

Obesity, insulin resistance, physical inactivity, advanced age and hormonal imbalance have been suggested as the underlying risk factors for the development of metabolic syndrome.

The metabolic syndrome is not a new syndrome. It was first described in the 1920 by Kylin, a Swedish physician, as the association of high blood pressure (hypertension), high blood glucose. (hyperglycemia) and gout. Advances in the 1970s and early 1980s expanded our understanding of the link of these to coronary heart disease, even in the absence of diabetes, and linked the metabolic risk factors to atherosclerosis. 1 In 1988, G.M. Reaven grouped several metabolic disorders together as syndrome X and proposed that insulin resistance was the underlying event explaining dyslipidemia, high blood pressure, and diabetes, and this characterization was further examined by DeFronzo and Ferranini in 1991. These factors were then observed to be influenced by both genetic and environmental factors. 2 Many other names were proposed for this syndrome, including the plurimetabolic syndrome (1988) , the deadly quartet (1989), syndrome X plus (1991), metabolic syndrome X, the metabolic syndrome, the insulin resistance syndrome ( ??991 It is the transport of lipid from intestine to liver. Chylomicrons transport the dietary lipid from intestine to liver. In diet the major lipids are triglyceride and cholesterol. Cholesterol is absorbed as such in proximal small intestine and is esterified to cholesteryl ester (ChE). Triglycerides are hydrolysed by lipases to glycerol and fatty acids which are absorbed in intestine. Inside intestinal cells triglyceride is synthesized by fatty acids. Chylomicrons are synthesized in the small intestine that contain triglyceride, cholesteryl ester, cholesterol, phospholipids, and apoprotein B-48 (apo B-48) and apo-A. These chylomicrons are secreted in the intestinal lymph and reach the systemic circulation via thoracic duct. In the circulation apo E and apo C are transferred to chylomicrons by HDL, so now chylomicrons contain apo A, apo B-48, apo-E, & apo-C. In the circulation, triglycerides of chylomicrons are hydrolysed by lipoprotein lipase (LPL) present on endothelial cells of vessels of skeletal muscles, adipose tissue and heart, but not in liver. The released fatty acids are utilized locally by these tissues. Adipose tissue uses these fatty acids to store them as TGs and heart and skeletal muscle use them as source of energy. Thus, function of chylomicrons is to transport exogenous (dietary) triglyceride to adipose tissue (for storage), heart (for energy) and muscle (for energy). The chylomicron particle progressively shrinks in size by action of LPL and, cholesterol, phospholipids, apo-A and apo C are transferred to HDL, creating chylomicron remnants that contains more cholesterol, less triglycerides, apo-E & apo B-48.

#### 1 b) Transport of Hepatic lipds (Endogenous pathway)

VLDL is synthesized in liver that contains high triglyceride, ChE, cholesterol, phospholipid and Apo B-100. (VLDL particles resemble chylomicrons in composition except that VLDL contains Apo B-100 instead of Apo B-48). VLDL particles are secreted in the plasma and as with chylomicron, Apo E and Apo C are transferred from HDL to VLDL. Now VLDL contains Apo B-100, Apo E and Apo C. In plasma, triglycerides of VLDL are hydrolysed by same lipoprotein lipase and apo C is transferred to HDL and the remnants are called IDL. 40-60% of IDL is removed by liver via LDL receptor mediated endocytosis, this process requires Apo E which acts as ligand for LDL receptors. Remaining IDL is remodeled by hepatic lipase which hydrolyzes more triglyceride to form LDL that contains maximum cholesterol. 70% of LDL is removed by liver via LDL receptor and 30% is utilized by peripheral tissues as a source of cholesterol.

#### 2 c) The Alterations in Metabolic Syndrome

The genetic and environmental factors play a major role in metabolic and cardiovascular of obesity. The way genetic factors modifying the effects of obesity are largely unknown. Also there are the subjects who are defined as obese by various guidelines but do not have insulin resistance and conversely insulin resistance can be present in lean individuals. Abdominal obesity is related, to greater risk of cardiovascular events and T2DM. Initially waist: hip ratio was considered but the current clinical approach to Metabolic Syndrome uses sex specific waist circumference to define the body mass component contributing to increase in cardiovascular events, greater fasting insulin levels and increased insulin resistance. Waist-circumference is a general index of central fat mass and it reflects both abdominal subcutaneous adipose tissue (SAT) and abdominal visceral adipose tissue (VAT) and is major determinant of metabolic and cardiovascular complication of obesity 5. Visceral obesity may partly be a marker of a dysmetabolic state and partly a cause of the metabolic syndrome. Although waist circumference is a better marker of abdominal fat accumulation than the body mass index, an elevated waistline alone is not sufficient to diagnose visceral obesity and have proposed that an elevated fasting triglyceride concentration could represent, when waist circumference is increased, a simple clinical marker of excess visceral/ectopic fat. 6 For the diagnosis of abdominal obesity three simple measurements, namely height, weight and maximal abdominal girth should be made on all patients. The body mass index (BMI); 18.5-25 kg/m<sup>2</sup> is normal, 25-30 kg/m<sup>2</sup> is overweight or >30 kg/m<sup>2</sup> is obese. According to the ATP III criteria the waist circumference of >35 inches (88 cm) for women, >40 inches (102cm) for men] defines excess abdominal adiposity. The weight reduction diet is usually choice for the obese patient but the most common approach for overweight person who may have the Metabolic syndrome is a balanced calorie diet. Hypothyroidism is a major cause of secondary dyslipidemia, the cause of which resides in a decrease of cholesterol excretion and in a marked increase in apo B lipoproteins because of a decreased catabolism and turnover by a reduced number of LDL receptors on the liver cell surface. LDL can be removed from plasma by receptors, which are regulated at mRNA level by thyroid hormone. Thus increased levels of total and LDL cholesterol are a common finding in hypothyroidism and may represent an increased risk factor for coronary heart disease. 7 Abdominal obesity, due to intra-abdominal adiposity, drives the progression of multiple cardiometabolic risk factors independently of body mass index. This occurs both through altered secretion of adipocyte-derived biologically active substances (adipokines), including free fatty acids, adiponectin, interleukin-6, tumour necrosis factor alpha, and plasminogen activator inhibitor-I, and through exacerbation of insulin resistance and associated cardiometabolic risk factors. 8 The prevalence of abdominal obesity is increasing in western populations, due to a combination of low physical activity and high energy diets, and also in developing countries, where it is associated with the urbanization of populations. The measurement of waist circumference, together with an additional comorbidity, readily identifies the presence of increased cardiometabolic risk associated with abdominal obesity. For example, 80% men with waist circumference 90 cm and triglycerides (TG) 2 mmol/L were found to have an atherogenic triad of elevated apolipoprotein B, fasting hyperinsulinaemia, and small, dense LDL, which had been strongly associated with adverse cardiovascular outcomes in a previous observational study. 9 Accordingly, measurement of waist circumference should become a standard component of cardiovascular risk evaluation in routine clinical practice. As a simple initial screening approach to distinguish viscerally obese from subcutaneously obese patients, has previously proposed that the simultaneous presence of fasting hypertriglyceridemia and of an increased waist circumference (hypertriglyceridemic waist) could represent a simple clinical phenotype to identify patients with an excess of visceral adipose tissue, with ectopic fat and with the related features of the metabolic syndrome. 10 Lifestyle modification remains the initial intervention of choice for this population, with pharmacological modulation of risk factors where this is insufficiently effective. Looking ahead, the initial results of randomized trials with rimonabant, the first CBI (cannabinoid) receptor blocker, indicate the potential of corrective overactivation of the endogenous endocannabinoid system for simultaneous improvement of multiple cardiometabolic risk factors. 11

### 3 III. Increased Triglycerides

Dyslipidemia is widely established as an independent risk factor for CVD. Low HDL Cholesterol and hypertriglyceridemia have been found to be independent and significantly related to myocardial infarction and stroke in patient with Metabolic syndrome. The dyslipidemia in Metabolic syndrome patient may be caused by combination of overproduction of very low density lipoprotein (VLDL), apoB-100, decreased catabolism of apo B containing

---

106 particle and increased catabolism of HDL apo-AI particle. In T2DM, insulin resistance and obesity combine to  
107 cause hypertriglyceridemia and low HDL-C, due to over production of VLDL. Plasma LDL-C levels are normal  
108 in Metabolic syndrome though LDL particles are smaller and denser than normal which may be associated with  
109 increased cardiovascular risk. The effects of low HDL-C and high triglyceride levels., are related to greater  
110 risk of coronary disease due to variety Of environmental and genetic factors. For instance lower HDL-C levels  
111 are formed in cigarette smokers, obese person, sedentary individuals, and androgen or progesterone users. The  
112 genetic variants of lipoprotein lipase, hepatic lipase, cholesterol ester transfer protein and peroxisome proliferator-  
113 activated receptors have effects on HDL-C and TG levels in population which contributes to the development of  
114 Metabolic syndrome.

115 The primary defect is probably focused in the inability to incorporate, the FFA to triglycerides (TGs) by the  
116 adipose tissue (inadequate esterification) 12 . This result is reduced fatty acid trapping and consequent retention  
117 by the adipose tissue. The insulin resistance also causes reduced retention of FFA by adipocytes. Both these  
118 abnormalities lead to the increased flux of FFA back in the liver. Adipose tissue is a prominent source of cholesteryl  
119 ester transfer protein. 13 Cholesteryl ester transfer protein is an important determinant of lipoprotein composition  
120 because of its capacity to mediate the transfer of cholesteryl esters from cholesteryl ester rich lipoproteins to TG  
121 rich lipoproteins in exchange for TGs. 14 In obese subjects, cholesteryl ester transfer protein activity and mass  
122 are increased. 15 Increased flux of FFA from periphery to the liver in increasing resistance state stimulates  
123 hepatic TG synthesis which in turn promotes the assembly & secretion of TG containing VLDL” as well as  
124 apo B production in liver. Under normolipidemic conditions in humans, VLDL secretion is affected by TG and  
125 cholesterol availability and recent studies suggest an association between’ cholesterol synthesis and. production  
126 of smaller-VLDL particles (VLDL2). 16 While insulin suppresses the formation, of large VLDL particles, VLDL  
127 receptors does not, have any impact on the production of the smaller VLDL2 fraction n When insulin resistance  
128 occurs, the high insulin values make the liver resistant to the inhibitory effects of insulin on VLDL secretion  
129 increasing insulin resistance is proposed to be the precursor for two events. in the presence of insulin resistance  
130 the visceral adipocytes is more sensitive to the metabolic effect of the lipolytic hormone glucocorticoids and  
131 catecholamines. The hormonal lipolytic activity produces an increased release of FFA Volume XV Issue III  
132 Version IYear 2 015 ( D D D D ) F

133 into portal system which serves as hepatic substrate to assemble TGs & TG rich VLDL. Secondly increasing JR  
134 leads to increased production of apo B as a consequent to increased synthesis & secretion of TG containing VLDL  
135 cholesterol particles. The fall off clearance rates is likely to reflect the-rates of lipolysis and could be attributable  
136 to a change in lipoprotein lipase activity (decreased in insulin resistance state) and other factors such as the  
137 apoC-II content or the apo-CII/CIII ratio (modulators of lipoprotein lipase activity) in VLDL. 17 In the insulin  
138 resistance, LDL levels are usually within normal limit or only mildly raised, however LDL particle is often of  
139 abnormal composition (small dense LDL) Hypertriglyceridemia is responsible for abnormal composition.-It has  
140 been found that small dense LDL is not seen in hypothyroidism until plasma TG level exceeds 1.5 mmol/L. The  
141 elevation of TGs in hypothyroidism is caused by a reduced removal rate of TG from plasma due to a decrease in  
142 the activity of hepatic TG lipase. Under these condition large TG rich VLDL (VLDL1) molecules accumulates  
143 when VLDL is lyophilized by LPL a population of LDL particle with changed apo B conformation is produced.  
144 These particles fail to bind efficiently LDL receptor and so have a prolonged residence time in circulation. By the  
145 action of CETP, cholesterol esters are replaced by TG in LDL & HDL particle. TG rich LDL is a good substrate  
146 for hepatic lipase that finally generates small dense LDL which is associated with increased cardiovascular risk.  
147 Many studies have shown that small, dense LDL particles have proatherogenic properties such as: (a) reduced  
148 LDL receptor mediated clearance, (b) increased arterial wall retention,(c) increased susceptibility to oxidation.  
149 19

## 150 4 IV. Low hdl-Cholesterol

151 Low HDL-Cholesterol in patient with Metabolic syndrome is often considered as secondary to raised TG. Lipid  
152 exchange is taking place between LDL & HDL particles mediated by cholesterol ester transport protein (CETP).  
153 These TG rich but cholesterol depleted HDLC is more prone to be catabolized. They undergo hydrolysis of their  
154 TGs components and dissociation of their protein component apo-A (the main protein of HDL). 20 There is  
155 circumstantial evidence that low HDL-C contributes to increase risk of atherosclerosis in this Condition. There is  
156 also evidence that HDL-cholesterol regime-therapy may reduce the risk of atherosclerosis in Met yn. 21, ???, ???  
157 The dyslipidemia contributing to diagnosis of Metabolic syndrome is defined by estimation of lipid levels after  
158 12 hour fasting in usual healthy state. A low HDL-C level <40 mg/dl in men and <50 mg/dl in female and TG  
159 level >150 mg/dl in either sex are the components for the diagnostic of Metabolic syndrome. The complexity of  
160 HDL, in terms of both composition and metabolism is striking. It is clear, however, that low HDL-C levels are a  
161 very common characteristic of the dyslipidemia of IR and T2DM. In fact, it may be the most common finding;  
162 HDL-C.and apo A-I levels in plasma can be low even when plasma TG levels are normal. Why are HDL-C and  
163 apo A-I levels reduced in individuals with JR and T2DM.

164 As was the case for LDL, the answer lies mainly, but not completely, with CETP. The reverse cholesterol  
165 transport system moves cholesterol from tissues to nascent, cholesterol-poor apo A-I phospholipid discs, where  
166 the cholesterol is esterified. Progressive accumulation of CE. (Cholesterol esters) leads to the maturation of  
167 the HDL into a spherical lipoprotein that delivers the CE and some of the unesterified cholesterol to the liver.

168 Alternatively, HDL2 particles can participate in the CETP-mediated exchange of their CE for TG in VLDL,  
169 VLDL remnants, and chylomicron remnants. The impact of this exchange on reverse cholesterol transport is  
170 unclear. If the TG rich lipoproteins that receive the CE from HDL are removed by the liver, there may be no  
171 effect on reverse cholesterol transport, assuming that delivery of CE via uptake of apo lipoproteins results in  
172 delivery of the lipid to the biliary tract with the same efficiency as does delivery via SRB 1. Unfortunately,  
173 increased CETP-mediated exchange of HDL CE for TGs is associated with lipolysis of HDL TG by HL, and this  
174 produces a small, lipid-depleted HDL that resembles nascent particles. Apo A-I binding to this HDL particle is  
175 diminished, causing apo A-I to dissociate from the particle and be cleared more rapidly from the plasma. Thus,  
176 fractional catabolism of apo A-I is increased in patients with T2DM, reducing the number of HDL particles in the  
177 circulation. Additionally, if the CE-enriched apo B-lipoproteins are not removed by the liver, but rather find their  
178 way into the artery wall, then reverse cholesterol transport will have been "shortcircuited" and atherogenesis may  
179 be accelerated.

180 There are other metabolic pathways that can also generate low HDL in IR and Metabolic syndrome. IR is  
181 associated with increased HL (hormone lipase) activity, which can increase hydrolysis of HDL, TG and generate  
182 smaller HDL. The potential roles of defective ABCA1-mediated efflux of cellular-free cholesterol, defective LCAT  
183 activity, or increased selective delivery of HDL cholesterol ester to hepatocytes as causes of the low HDL levels  
184 present in IR are under investigation. However, the observation that low HDL-C and apo A-I are frequently  
185 present even when TG levels are relatively normal suggests that non-CETP mechanisms are clearly important in  
186 the pathogenesis of low HDL-C concentrations in IR.

## 187 5 V. Conclusion

188 Elevated triglyceride is the most prevalent (81%) component of metabolic syndrome in all thyroid patients of  
189 present study. Hence hypertriglyceridemia on routine. There is a significant correlation ( $p= 0.03$ ) between waist  
190 circumference and triglyceride levels. It is the visceral fat which is responsible for hypertriglyceridemia. Hence  
191 it can be concluded that those having high TG levels and high waist circumference, have visceral adiposity.  
192 Therefore screening for other parameters of metabolic syndrome may be warranted as visceral adipose tissue  
193 causes the most metabolic derangements. All the results indicate that lipid profile undergoes a change in  
194 metabolic syndrome and in clinical practice the fatal outcomes and complications are directly related to the  
195 severity of lipid abnormalities. Hence the change in Lipid Profile can be a prognostic factor for Metabolic  
196 Syndrome and the changes should be seriously taken care of and addressed at the earliest.

## 197 6 Volume XV Issue III Version I



Figure 1:

1

Hypertension	Raised arterial pressure(140/90 mmHg) or antihypertensive medication
Dyslipidemia	Raised plasma triglyceride(>1.7 mmol/L) or low HDL cholesterol(<0.9 mmol/L) in men and (<1.0mmol/L) in women
Central or general obesity	Waist to hip ratio >0.90 in men: >0.85 in women or BMI>30Kg/m <sup>2</sup>
Microalbuminuria	Urinary albumin excretion rate >20ug/min or albumin: creatinine ratio>30 mg/g

Figure 2: Table 1 :

2

Risk factors	Defining levels*
Abdominal obesity (waist circumference)	>102 cm(> 40 inch)
Men	>88cm (> 35 inches)
Women	
Triglycerides	>150 mg/dl
HDL Cholesterol	
Men	<40mg/dl
Women	<50mg/dl
Blood Pressure	>130/85 mm Hg
Fasting Glucose	>110 mg/dl
Metabolic syndrome: diagnosis is established when >3 of these risk factors are present	

II. Normal Physiology of Lipid Transport

[Note: a) Transport of dietary lipid (Exogenous pathway)]

Figure 3: Table 2 :

---

<sup>1</sup>© 2015 Global Journals Inc. (US)

<sup>2</sup>© 2015 Global Journals Inc. (US) Lipid Profile Alterations in the Metabolic Syndrome the Latest Biochemical and Physiological Aspect

- 
- 199 [Ross et al. ()] ‘Abdominal obesity, muscle composition, and insulin resistance in premenopausal women’. R Ross  
200 , J Freeman , R Hudson , I Janssen . *J Clin Endocrinol Metab* 2002. 87 p. .
- 201 [Shafirir ()] ‘Animal models of syndrome X. Current Topics in’. E Shafirir . *Diabetes Research* 1993. 12 p. .
- 202 [Ginsberg et al. ()] ‘Apolipoprotein B metabolism in subjects with deficiency of apolipoproteins C-III and AI;  
203 evidence that apolipoprotein CIII inhibits catabolism of triglyceride-rich lipoproteins by lipoprotein lipase in  
204 vivo’. H N Ginsberg , N-A Le , I J Goldberg . *J Clin Invest* 1986. 78 p. .
- 205 [Morton ()] ‘Cholesteryl ester transfer protein and its plasma regulator: lipid transfer inhibitor protein’. R E  
206 Morton . *Curr Opin Lipidol* 1999. 10 p. .
- 207 [Danese et al. ()] ‘Clinical review 115: effect of thyroxin therapy on serum lipoproteins in patients with mild  
208 thyroid failure: a quantitative review of the literature’. M D Danese , P W Ladenson , C L Meinert , N R  
209 Powe . *J Clin Endocrinol Metab* 2000. 85 p. .
- 210 [Drayna et al. ()] ‘cloning and sequencing of the human cholesteryl ester transfer protein cDNA’. D Drayna , A  
211 S Jamagin , J Mclean . *Nature* 1987. 327 p. 6324.
- 212 [Lemieux et al. ()] ‘Despre’s JP. Hypertriglyceridemic waist. A marker of the atherogenic metabolic triad  
213 (hyperinsulinemia, hyperapolipoprotein B, small, dense LDL) in men’. I Lemieux , A Pascot , C Couillard ,  
214 B Lamarche , A Tchernof , N Alme Ras , J Bergeron , D Gaudet , G Tremblay , D Prud’homme , A Nadeau  
215 . *Circulation* 2000. 102 p. .
- 216 [Despres et al. ()] ‘Effects of rimonabant on metabolic risk factors in overweight patients with dyslipidemia’. J  
217 P Despres , A Golay , L Sjostrom . *N Engl J Med* 2005. 353 p. .
- 218 [Executive Summary of the Thyroid Report of National Cholesterol Education Program (NCEP)] *Executive*  
219 *Summary of the Thyroid Report of National Cholesterol Education Program (NCEP)*,
- 220 [Expert Panel on detection, evaluation and treatment of high blood cholesterol in adults JAMA ()] ‘Expert  
221 Panel on detection, evaluation and treatment of high blood cholesterol in adults’. *JAMA* 2001. 285 p. .
- 222 [Lamarche and Tchernof ()] ‘Fasting insulin and apolipoprotein B levels and low-density lipoprotein particle size  
223 as risk factors for ischemic heart disease’. B Lamarche , A Tchernof , Mauriege . *JAMA* 1998. 279 p. .
- 224 [Barter and Rye (16996)] ‘High density lipoproteins and coronary heart disease’. P J Barter , K A Rye .  
225 *Atherosclerosis* 16996. 121 p. .
- 226 [Arai et al. ()] ‘Increased plasma cholesteryl ester transfer protein in obese subjects: a possible mechanism for  
227 the reduction of serum HDL cholesterol levels in obesity’. T Arai , S Yamashita , K Hirano . *Arterioscler*  
228 *Thromb* 1994. 14 p. .
- 229 [Lempiainen ()] ‘Insulin resistance syndrome predicts coronary heart disease events in elderly nondiabetic men’.  
230 P Lempiainen . *Circulation* 1999. 100 (2) p. .
- 231 [Malmstro et al. ()] ‘Metabolic basis of hypotriglyceridemic effects of insulin in normal men’. R Malmstro , C J  
232 Packard , T D Watson . *Arterioscler Thromb Vase Biol* 1997. 17 p. .
- 233 [Wajchenberg ()] ‘Subcutaneous and visceral adipose tissue: their relation to the metabolic syndrome’. B L  
234 Wajchenberg . *Endocr Rev* 2000. 21 p. .
- 235 [Chatt et al. ()] ‘Susceptibility of small, dense, low-density lipoproteins to oxidative modification in subjects with  
236 the atherogenic lipoprotein phenotype, pattern B’. A Chatt , R L Brazg , D L Tribble . *Am J Med* 1993. 94  
237 p. .
- 238 [Nieves et al. ()] ‘The atherogenic lipoprotein profile associated with obesity and insulin resistance is largely  
239 attributable to intraabdominal fat’. D J Nieves , M Cnop , B Retzlaff . *Diabetes* 2003. 52 p. .
- 240 [Nieves et al. ()] ‘The atherogenic lipoprotein profile associated with obesity and insulin resistance is largely  
241 attributable to intraabdominal fat’. D J Nieves , M Cnop , B Retzlaff , C E Walden , J D Brunzell , R H  
242 Knopp , S E Kahn . *Diabetes* 2003. 52 p. .
- 243 [Ginsberg and Haug ()] ‘The insulin resistance syndrome: impact on lipoprotein metabolism and atherothrom-  
244 bosis’. H N Ginsberg , L S Haug . *J Cardiovasc Risk* 2000. 7 p. .
- 245 [Wilson et al. ()] ‘The metabolic syndrome as a precursor of cardiovascular disease and type 2 diabetes’. Pwf  
246 Wilson , D ’ Agostino , R B Parise , H Sullivan , L Meigs , JB . *Circulation* 2005. 112 p. .
- 247 [Wilson and Grundy ()] ‘The metabolic syndrome: practical guide to origins and treatment: part I’. P W Wilson  
248 , S M Grundy . *Circulation* 2003. 108 p. .