Postprandial dysmetabolism and cardiovascular disease in type 2 diabetes

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The worldwide prevalence of type 2 diabetes mellitus has reached epidemic proportions. The so-called traditional risk factors cannot fully explain the excessive cardiovascular disease risk of type 2 diabetic patients. Numerous studies indicate that postprandial metabolic derangements, most notably hyperglycaemia and hypertriglyceridaemia, which are exaggerated and prolonged in type 2 diabetes, are important cardiovascular disease risk factors since they induce oxidative stress and endothelial dysfunctions. This review discusses the current evidence showing that postprandial dysmetabolism may indeed constitute an important cardiovascular disease risk factor as well as the mechanisms underlying this association. Finally, some possible therapeutic options and recommendations for future research are discussed.
In this review we will briefly discuss the present evidence for postprandial dysmetabolism as a potential cardiovascular disease risk factor, with special emphasis on postprandial glucose and lipid dysmetabolism. To this purpose, the association of postprandial dysmetabolism with the presence of (indicators of) atherosclerotic vascular disease and the possible underlying mechanisms will be reviewed.

**POSTPRANDIAL HYPERGLYCAEMIA**

Various mechanisms keep the plasma glucose levels in healthy subjects between strict limits, even after a carbohydrate load. The insulin response after a meal regulates the glucose uptake from the blood into the peripheral tissues and inhibits gluconeogenesis and glycogenolysis in the liver. Dysfunction of β-cells (for example, loss of the normal insulin secretion pattern) and insulin resistance contribute to glucose intolerance and both can be found very early in the disease process, finally leading to type 2 diabetes.4

In the United Kingdom Prospective Diabetes Study, lowering of glycated haemoglobin mainly reduced long term microvascular complications of diabetes, whereas effects on macrovascular disease were less convincing.18 In apparent contrast, several epidemiological studies have shown an association between two hour glucose concentrations after a 75 g glucose load (2hPG) and the occurrence of cardiovascular disease in the general population.19-20 A meta-analysis of 20 studies with more than 95 000 people demonstrated a continuous relationship between postload glucose levels and cardiovascular disease risk extending into the non-diabetic range.20 The Diabetes Epidemiology: Collaborative analysis Of Diagnostic criteria in Europe (DECODE) study demonstrated that the 2hPG concentrations, even in subjects with normal fasting glucose, were associated with mortality, independent of fasting plasma glucose concentrations.20

Chronic hyperglycaemia has been associated with impaired endothelial function.21 Recent studies in healthy and type 2 diabetic subjects indicate that acute hyperglycaemia causes endothelial dysfunction as measured by FMD.22 Also, circulating ICAM-1 plasma levels significantly increased in both diabetic and normal subjects after an oral glucose tolerance test, suggesting endothelial cell activation.23

Ceriello and co-workers have shown that postprandial hyperglycaemia is accompanied by several alterations of the coagulation system.24-25 An oral glucose load in both healthy and type 2 diabetes patients caused a shortening of the half life of fibrinogen and an increase in plasma fibrinopeptide A and the fragments of prothrombin and antithrombin III. In addition, acute, short term hyperglycaemia resulted in a transient hyper-reactivity of platelets to high shear stress, combined with a significant rise of plasma vWF in patients with type 2 diabetes.26 Taken together, these findings suggest that hyperglycaemia may induce a hypercoagulable state.

In healthy subjects and those with impaired glucose tolerance, consecutive pulses of intravenous glucose increased circulating cytokine concentrations (interleukin-6 and tumour necrosis factor-α) to a greater extent than during similar blood glucose levels which were kept stable during a hyperglycaemic clamp. This effect was more pronounced in subjects with impaired glucose tolerance.27 The same investigators showed changes in interleukin-6 (but not tumour necrosis factor-α) plasma concentrations in type 2 diabetic patients after a carbohydrate meal.28 Thus, blood glucose excursions may induce a proinflammatory response.

Several in vitro studies demonstrated cytotoxic effects of high glucose levels in various cell types.29-30 Of interest is the demonstration by Riso and colleagues that intermittent high glucose levels induced more apoptosis than constant corresponding glucose levels in human umbilical vein endothelial cells.30

Four main molecular mechanisms underlying the hyperglycaemia-induced vascular damage have recently been reviewed,31 all of which are the result of intracellular hyperglycaemia. These include increased polyol pathway influx; increased advanced glycation end-product formation; activation of protein kinase C isoforms; and increased hexosamine pathway flux. These seemingly different mechanisms are the result of a single process—that is, overproduction of superoxide by the mitochondrial electron transport chain. This hyperglycaemia-induced oxidative stress ultimately results in modification of intracellular proteins resulting in an altered function, DNA damage, activation of the transcription factor nuclear factor-κB, causing abnormal changes in gene expression, decreased production of nitric oxide, and increased expression of cytokines and inflammasome factors and procoagulant and proinflammatory molecules.31

Taken together, postload or postprandial glucose levels are associated with enhanced risk of cardiovascular disease. However, most epidemiological studies addressing the contribution of postload glucose levels to cardiovascular disease risk, especially the early ones, did not take into account the earlier mentioned classical risk factors, such as dyslipidaemia. In studies investigating the relationship between postload glucose and cardiovascular disease risk, adjustment for blood pressure, lipids and smoking, resulted in considerable attenuation of this association.24-31 These data indicate that postload glucose may not be an independent cardiovascular disease risk factor but rather a risk marker, suggestive of underlying other metabolic disturbances, such as insulin resistance and dyslipidaemia, that may have an even greater impact on cardiovascular disease risk.

In conclusion, evidence of postprandial hyperglycaemia as an independent risk factor is not convincing. Therefore, the observed association between postload glucose excursions with cardiovascular disease is at least partly explained by the presence of insulin resistance and related cardiovascular disease risk factors.

**POSTPRANDIAL HYPERTRIGLYCERAEMIA**

In the western diet, more than 40% of the energy intake is derived from fats. Figure 1 demonstrates a schematic representation of the metabolic pathways of dietary fats leading to triglyceride-rich lipoproteins (TRL) and endogenous TRL production.34 In the insulin-resistant state the production of very low density lipoprotein (VLDL) by the liver is inappropriately high. Together with a reduced lipoprotein lipase activity this results in high triglyceride concentrations, especially in the postprandial state. The large amount of TRLs and their prolonged residence time in the circulation may lead to increased exchange of the core lipid cholesteryl ester for triglycerides between TRL and LDL and HDL particles mediated by cholesteryl ester transfer protein. This process enriches LDL and HDL with triglyceride, and these particles are subsequently more readily hydrolysed by hepatic lipase resulting in smaller, denser LDL particles and lower concentrations of HDL. These abnormalities may explain the characteristic diabetic dyslipidaemia, which is now recognised to be very atherogenic.35

Already in 1959 an association between plasma concentrations and incident coronary heart disease was reported.35 However, the known inverse association between triglycerides and HDL cholesterol makes it difficult to show an independent association between plasma triglyceride and atherosclerotic vascular disease. A recently performed meta-analysis including data of 57 000 subjects from 17 studies demonstrated that fasting triglyceride concentrations were an independent risk factor for cardiovascular disease, also when adjusted for HDL cholesterol.36 A 1 mmol/l increase in
The remaining particles, the chylomicron remnants, are removed from the circulation by the liver through binding of their surface apoE to the low density lipoprotein (LDL) receptor or LDL receptor related protein. Very low density lipoprotein (VLDL) particles are triglyceride-rich apoB100 containing particles, synthesised by the liver. As with chylomicrons, VLDL triglycerides are hydrolysed by LPL. VLDL remnants or intermediate density lipoproteins (IDL) are taken up by liver receptors via apoE or converted to LDL. Chylomicrons, VLDL and their respective remnants (RLP, remnant lipoproteins) are termed triglyceride-rich lipoproteins (TRL). Under physiological conditions, insulin, which is raised in the postprandial state, suppresses lipolysis from adipose tissue and hepatic VLDL production, however, this insulin action is inappropriate in insulin resistance and type 2 diabetes, resulting in high TRL concentrations. The large amount of TRL and their prolonged residence time in the circulation increase the exchange of esterified cholesterol from high density lipoprotein (HDL) and LDL to TRL and of triglycerides to LDL and HDL particles, which is mediated by cholesterol-ester transfer protein (CETP). Triglyceride enrichment of LDL particles renders them better substrates for hepatic lipase (HL), which hydrolyses triglycerides from the core of LDL and turns them into smaller and denser particles. Small dense LDL are more atherogenic as they readily enter the subendothelial space and become oxidised. Triglyceride enriched HDL particles are smaller and are more rapidly catabolised, which may explain the observed low plasma HDL in insulin resistance and type 2 diabetes.

In general practice, serum lipid concentrations including triglycerides are measured in the morning after an overnight fast. However, the fasting value should be considered the nadir of the 24 hour triglyceride profile and could therefore be misleadingly low. In the past few years several clinical studies have suggested that high postprandial TRL may be related to coronary heart and/or carotid artery disease in non-diabetic and diabetic subjects. The Physician Health study, including 14 916 men aged 40–84 years, with a follow up of seven years, showed that the non-fasting triglyceride concentrations strongly predicted incident myocardial infarctions, with a relative risk of 1.40 (95% confidence interval 1.10 to 1.77) per 1.13 mmol/l increase. This study suggests that random or postprandial triglyceride concentrations are an important indicator of cardiovascular disease risk. Although fasting triglycerides are the most important determinant of postprandial triglycerides, it may be argued that in insulin resistant subjects with a delayed postprandial TRL clearance, non-fasting triglycerides should be used to approximate overall triglyceride exposure.

In male patients after a myocardial infarction, Karpe and co-workers found that the progression of coronary lesions over five years was related to the postprandial plasma levels of small chylomicron remnants (SF 20–60 apolipoprotein B48). Adjustment for the possible confounding effect of HDL and dense LDL apolipoprotein B concentrations did not substantially alter the strength of this association. In line with these findings are coronary angiography data described by Mero et al, which suggest that especially small chylomicron remnants are implicated in the progression of coronary artery disease.

In healthy and type 2 diabetic subjects, endothelial dysfunction, measured by ultrasound as FMD, has been associated with high triglyceride excursions. Correlations between postprandial TRL, impaired FMD and oxidative stress markers have been demonstrated, suggesting that free radical production may be an underlying mechanism. Supporting this hypothesis is the finding that in healthy subjects this effect could be attenuated by the antioxidant vitamin C.

In vitro studies demonstrated increased adhesion molecule expression in endothelial cells after incubating with chylomicrons and VLDL. Using rat arterial rings, Lundman et al showed impairment of endothelium-dependent relaxation following exposure to the triglyceride-containing fat emulsion Intralipid (Pharmacia-Upjohn, Uppsala, Sweden), however, exposure to VLDL did not affect vascular function.

Postprandial coagulation activation by TRL was demonstrated by several investigators, however the underlying mechanism(s) are not fully understood. An elegant study performed by Silveira et al suggests an important role for the intrinsic coagulation pathway, based on in vivo activation of factor XI by triglyceride. Other prothrombotic changes occurring with an oral fat load are increased plasmin activator inhibitor-1 (PAI-1) activity and PAI-1 antigen. Postprandial lipaemia enhanced platelet P-selectin expression without affecting other markers of platelet activation.
The effect of a high fat meal (50 g of fat) on cytokine concentrations, reflecting the inflammatory state, was studied in healthy and type 2 diabetic patients. In healthy subjects, significant correlations were found between postprandial triglyceride and tumour necrosis factor-α levels, whereas in diabetic patients also a positive correlation between postprandial plasma triglyceride and interleukin-6 concentrations was observed. Antioxidant supplementation lowered the rise of the cytokines, suggesting that the cytokine response to triglycerides was mediated by oxidative stress.

To summarise, historically, in diabetic patients, most emphasis was laid on hyperglycaemia, whereas recent evidence demonstrates the importance of dyslipidaemia, in particular hypertriglyceridaemia, as a cardiovascular disease risk factor. Although at present, epidemiological and long term intervention studies are largely lacking, in vivo data convincingly show an association between postprandial TRL and indicators of cardiovascular disease. Similar to postprandial hyperglycaemia, both in vivo and in vitro studies indicate that (postprandial) increases in triglycerides are proinflammatory, prothrombotic, and adversely affect several endothelial functions, by inducing oxidative stress (fig 2). Therefore, it is feasible that prolonged postprandial hypertriglyceridaemia leads to an atherogenic environment in vivo. However, as for postprandial hyperglycaemia, evidence for postprandial hypertriglyceridaemia as independent in cardiovascular disease is still scanty. More evidence, which can only be obtained from large prospective studies, is certainly required.

**THERAPEUTIC INTERVENTIONS**

Drugs that have been effective in reducing meal related glucose excursions are the α-glucosidase inhibitors, the short acting insulin analogues, and the meglitinides.50–52

Recently, the STOP-NIDDM study showed a lower incidence of hypertension and myocardial infarction after treatment with the α-glucosidase inhibitor acarbose.53 These findings should be interpreted with some caution as the study was not designed to assess the effect of acarbose on cardiovascular disease endpoints. The prospectively predefined endpoint was conversion to diabetes, however, in the final report 10 different cardiovascular disease endpoints were mentioned, including angina and peripheral vascular disease. In addition, it can not be excluded that the reduction in cardiovascular disease endpoints is the result of lowering triglycerides, since a triglyceride lowering effect of acarbose is described.54

Metformin is the only blood glucose-lowering drug that has been shown to lower diabetes related cardiovascular disease endpoints in obese type 2 diabetic patients. Based on the proposed working mechanism,55 an effect on meal related glucose excursions can not be expected, but metformin was shown to reduce postprandial chylomicron concentrations.56

Correction of postprandial hyperglycaemia, for example with insulin secretion enhancers, will not only affect glucose levels but probably also the postprandial lipid responses. However, recently Vakkilainen et al demonstrated that nateglinide and gliclazide increased postprandial insulin secretion and decreased postprandial glycaemia, but neither drug attenuated postprandial lipaemia in type 2 diabetic subjects with good glycaemic control.57

Intensive insulin treatment may improve diabetic dyslipidaemia to some extent, however, an entire correction of the atherogenic lipid profile, including postprandial hypertriglyceridaemia, may not be achieved.58

The lipid lowering drugs—that is, hydroxymethylglutaryl coenzyme A reductase inhibitors (statins), and peroxisome proliferator-activated receptor α-agonists (fibrates), are established as most efficient agents that reduce cardiovascular disease morbidity and mortality in various high risk populations.59–62 Statins inhibit cholesterol synthesis and up-regulate the hepatic LDL receptor, whereas fibrates increase LPL activity and limit hepatic VLDL secretion. Based on their respective working mechanisms, the most benefit on postprandial lipaemia may be expected from fibrates. Indeed, fibrate treatment reduced postprandial triglyceride levels by 30–50%.63 The proposed beneficial effect of fibrates on postprandial endothelial function measured by FMD, however, is disappointing.64–66 Statins tend to induce a modest lowering of both fasting and postprandial triglycerides.67–69 However, statin treatment showed a marked beneficial effect on postprandial induced oxidative stress and endothelial function.63 Independently of their cholesterol-lowering action, statins seem to have anti-inflammatory and vasculo-protective effects (“pleiotropic” effects).70

We conclude that the high cardiovascular disease morbidity and mortality associated with type 2 diabetes is at least partly due to a prolonged and exaggerated postprandial state in these patients. To date, however, controlled randomised intervention studies showing that postprandial both glucose and triglyceride lowering results in amelioration of clinically relevant endpoints are lacking.

These conclusions should in no way distract from the therapeutic aim to achieve target glycaed haemoglobin and lipid values in patients with type 2 diabetes.

**RECOMMENDATIONS FOR FUTURE RESEARCH AND TESTING POSTLOAD DYSMETABOLISM**

As discussed above, although the beneficial effect of therapy targeting postprandial dysmetabolism still needs to be established, studies assessing the true atherogenic exposure of the vascular system in high risk patients should abandon the classical glucose centred view and use physiological tests combining glucose and lipid loads. Most studies mentioned earlier demonstrated the effects of postprandial dysmetabolism on a single and rather artificial challenge, like a liquid 75 g glucose or liquid fat load. In daily life, most meals...
consumed are mixed and of solid consistence. Ceriello and co-workers showed a cumulative adverse effect of postprandial hypertriglyceridemia and hyperglycaemia on endothelial function. The effect of a single component challenge possibly underestimates the impact of real life postprandial dysmetabolism, cardiovascular disease, and diabetes.5

5. Evidence suggests that high postload or postprandial glucose levels constitute an independent risk factor for the development of cardiovascular disease.

6. In the insulin resistant state, high VLDL concentrations are the result of an increased flux of chylomicrons from the gut due to a reduced lipoprotein lipase activity.

7. The fasting triglyceride value should be considered the nadir of the 24 hour triglyceride profile and could therefore be misleadingly low.

8. Statins have been shown to reduce postprandial triglyceride levels more adequately than fibrates and have a beneficial effect on postprandial induced oxidative stress.


ACKNOWLEDGEMENTS

MET is supported by a grant from the Dutch Diabetes Foundation (grant no 2000.00.025).

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ANSWERS

1. False. The excessive global increase in the occurrence of type 2 diabetes may be explained by the vast spreading of western lifestyle patterns, including consumption of energy dense foods and less physical activity, in parts of the world with the highest population densities—that is, the developing countries.

2. False. Hyperglycaemia is the result of progressive pancreatic β-cell failure.

3. True. Absolute risk of death by cardiovascular disease was much higher for diabetic than non-diabetic men of every age stratum, ethnic background, and risk factor level—overall three times higher, with adjustment for age, race, income, serum cholesterol level, systolic blood pressure, and smoking.5

4. False. Endothelial function can only be estimated—indirectly—by provoking the endothelium to release nitric oxide resulting in vasodilatation that can be imaged and quantitated.

5. True. Postload glucose may not be an independent cardiovascular disease risk factor but rather a risk marker, suggestive of underlying metabolic disturbances that collectively impact on cardiovascular disease risk.

6. False. In the insulin resistant state inappropriate production of VLDL by the liver occurs, due to impaired suppression of VLDL triglyceride by insulin (hepatic insulin resistance) and an increased substrate (fatty acid) flux to the liver.

7. True. In clinical practice triglycerides are measured in the morning after an overnight fasting period. Triglyceride profiles of type 2 diabetic patients show that after breakfast their concentrations gradually increase after consecutive meals and the peak concentration is achieved between dinner and bedtime.

8. False. Based on their working mechanism, that is, increasing (postprandial) lipoprotein lipase activity, but also in practice, fibrates, rather than statins reduce postprandial triglycerides. Statins tend to induce a modest lowering of fasting and postprandial triglyceride levels.

9. True. A cumulative effect of postprandial hypertriglyceridaemia and hyperglycaemia on endothelial function has been demonstrated, possibly due to potentiation of oxidative stress mechanisms.
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Postgrad Med J 2005 81: 1-6
doi: 10.1136/pgmj.2004.020511

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