45Obesity, Insulin Resistance and Free Fatty Acids

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Abstract

Purpose of Review—to describe the role of FFA as a cause for insulin resistance in obese people.

Recent Findings—elevated plasma FFA levels can account for a large part of insulin resistance in obese patients with type 2 diabetes. Insulin resistance is clinically important because it is closely associated with several diseases including T2DM, hypertension, dyslipidemia and abnormalities in blood coagulation and fibrinolysis. These disorders are all independent risk factors for cardiovascular disease (heart attacks, strokes and peripheral arterial disease). The mechanism by which FFA can cause insulin resistance, although not completely known, include generation of lipid metabolites (diacylglycerol), proinflammatory cytokines (TNF-α, IL1β, IL6, MCP1) and cellular stress including oxidative and endoplasmic reticulum stress.

Summary—increased plasma FFA levels are an important cause of obesity associated insulin resistance and cardiovascular disease. Therapeutic application of this knowledge is hampered by the lack of readily accessible methods to measure FFA and by the lack of medications to lower plasma FFA levels.

Keywords
free fatty acids; insulin resistance; obesity; cardiovascular disease

INTRODUCTION

Insulin resistance, defined as inhibition of insulin stimulation of several metabolic pathways including glucose transport, glycogen synthesis and anti-lipolysis, is of considerable clinical relevance because it is pathophysiologically linked to several serious medical problems including type 2 diabetes (T2DM), hypertension, atherogenic dyslipidemia, abnormalities of blood coagulation and fibrinolysis and non-alcoholic fatty liver disease (1) (Figure 1). Some of these disorders have been collectively labeled the “metabolic or insulin resistance syndrome” (2). Importantly, they are all independent risk factors for cardiovascular diseases (CVD), which may explain why obesity is associated with a 2-4 times increased risk for heart attacks, strokes and peripheral vascular disease (1). Insulin resistance can have many causes (3), but by far the most common cause, particularly in developed countries, is obesity. In the US, obesity has reached epidemic proportions where more than 2/3 of all adults are either overweight or obese (4). Exactly why and how obesity causes insulin
resistance is not completely understood. The most likely possibilities, however, are excessive nutrient intake and/or an expanded adipose tissue.

It is now recognized that adipose tissue is not only a storage site for excessive calories in the form of fat but is also a metabolically active tissue that synthesizes and secretes a large number of biologically active substances, such as proinflammatory cytokines, acute phase reactants, angiotensin II, leptin, resistin, adiponectin, PAI-1 and others (5). Some of these compounds, particularly, when administered in large amounts, can produce insulin resistance. Nevertheless, to be considered a physiological link between obesity and insulin resistance, an adipose tissue derived factor should meet the 3 criteria shown in Table 1.

So far, only FFAs meet the 3 criteria in human subjects. Therefore, this review focuses primarily on the effects of FFA on insulin action. This does not rule out that some of the other adipose tissue derived factors may eventually be recognized as physiologic causes for obesity associated insulin resistance.

**FFA levels are increased in obese people**

Plasma FFA levels are elevated in most obese subjects (6) because 1) the enlarged and stressed adipose tissue releases more FFA and 2) FFA clearance may be reduced (7). Moreover, once elevated, FFA will inhibit insulin’s antilipolytic action which will further increase FFA release into the circulation (8).

**Raising FFA levels causes acute insulin resistance**

Acute elevations of plasma FFA levels, for instance by infusion of heparinized triglyceride emulsions, decreases whole body insulin stimulated glucose uptake. This FFA induced insulin resistance develops after a delay of approximately 2 hours and disappears approximately 4 hours after normalization of plasma FFA, is dose dependent and affects men and women equally, regardless of whether or not they are diabetic (9-11). Because insulin stimulated glucose uptake under these euglycemic-hyperinsulinemic clamp conditions occurs mostly (greater than 80%) in skeletal muscle (12), it follows that FFA causes acute insulin resistance in skeletal muscle. Acute elevations of plasma FFA to levels frequently seen in obese individuals, also inhibit insulin suppression of hepatic glucose production. This acute FFA induced hepatic insulin resistance is primarily due to inhibition of insulin mediated suppression of glycogenolysis with little or no acute effect on gluconeogenesis (13). On the other hand, longer lasting FFA elevations are likely to also increase gluconeogenesis.

In endothelial cells, IV infusion of insulin has been shown to increase nitric acid production resulting in increased peripheral vascular blood flow (14). Physiologically elevated FFA levels produce insulin resistance in endothelial cells by inhibiting this insulin induced increase in nitric oxide and blood flow (15).

**Acute lowering of plasma FFA reduces chronic insulin resistance**

That the chronically elevated FFA levels seen in most obese individuals were responsible for a large part of their insulin resistance was demonstrated by acutely normalizing the chronically elevated plasma FFA levels in obese, diabetic and non-diabetic subjects. This resulted in normalization of insulin sensitivity in obese non-diabetic individuals and improved insulin sensitivity from ~ 25% to ~ 50% of normal in obese patients with T2DM (16). Similar results have been reported in subjects genetically predisposed to develop T2DM (17).
Moreover, a substantial part of the insulin resistance lowering activity of thiazolidinediones (TZDs) can be attributed to their lowering of plasma FFA levels (18), which they do by increasing FFA oxidation (19).

Mechanisms of FFA mediated insulin resistance

Despite much work, the mechanisms by which obesity and FFA cause insulin resistance are still not completely understood. Almost 50 years ago, Randle et al. proposed that FFA inhibited insulin stimulated glucose uptake in rat heart and diaphragm muscle by increasing fat and decreasing carbohydrate oxidation (20). Later studies in human skeletal muscle revealed, however, that the FFA induced increased fat and decreased carbohydrate oxidation was not a likely cause for the ensuing insulin resistance. Rather, FFAs were found to inhibit insulin action at the level of insulin stimulated glucose transport and/or phosphorylation by inhibiting insulin signaling (11,21). Recently, several mechanisms have been proposed to explain how obesity and/or FFA can interfere with insulin signaling.

The lipid metabolite hypothesis

This hypothesis is based on the finding that an increase in plasma FFA concentration results in intramyocellular and intrahepatic accumulation of triglycerides as well as several metabolites of the FFA reesterification pathway including longchain Acyl-CoAs and diacylglycerol (DAG) (22). DAG is a potent activator of conventional and novel protein kinase C (PKC) isoforms (23). In human skeletal muscle, elevation of plasma FFA levels has been shown to increase DAG and to activate PKC β2 and δ (24). PKC, a serine/threonine kinase, has been shown in rodents to cause insulin resistance by decreasing tyrosine phosphorylation of the insulin receptor substrate (IRS) 1/2 (25). Complicating the issue is that there are more than 20 serine/threonine consensus sites in IRS1 that can be phosphorylated. Not surprisingly, therefore, other serine/threonine kinases have been postulated to be involved in obesity/FFA induced inhibition of insulin signaling and action (see below).

The reason for the increase in DAG following lipid infusions is not entirely clear. A reasonable assumption is that the fatty acid reesterification pathway can be overwhelmed in high fatty acid flux situations. Supporting this notion is the demonstration that overexpression of DGAT (the enzyme which catalyzes the formation of triacylglycerol from DAG) in skeletal muscle of mice, protected against high fat diet induced insulin resistance, whereas DGAT deficiency increased FFA induced insulin resistance in isolated mouse muscles (26). Moreover, one single 90 minute session of moderate intensity exercise completely reversed FFA induced insulin resistance in healthy volunteers one day later (27). This was accompanied by increases in skeletal muscle triglyceride, GPAT, DGAT and IκB-α proteins and decreases in skeletal muscle DAG, glycogen and phosphorylated JNK levels (27). Nevertheless, so far, the DAG/PKC hypothesis in human tissue is based on correlative data and requires more direct evidence.

The inflammation hypothesis

It is now recognized that obesity is an inflammatory state associated with increased levels of proinflammatory cytokines and chemokines in the circulation and in tissues (28). Recent studies have shed some light on possible causes. For instance, FFA have been shown in vivo and in vitro to activate the canonical proinflammatory NF-κB pathway. In vivo, acutely increasing plasma FFA levels activated NF-κB in human skeletal muscle (24) and resulted in increased expression of several proinflammatory cytokines including TNF-α, IL1β and IL6 in rat liver as well as an increase in circulating MCP1 levels (29). The rise in plasma MCP1 is particularly interesting because MCP1 is well established to regulate macrophage recruitment to sites of inflammation (30). The rise in plasma MCP levels, therefore, may be
involved in the recently reported monocyte recruitment into adipose tissue of obese animals where they differentiate into macrophages and produce proinflammatory cytokines (31,32). In vitro, linoleic acid (C18:2) activated NF-κB and increased IL6 and TNF-α in cultured human adipocytes and stomal vascular cells (33). The concept that emerges from these observations is that elevated plasma FFA levels, either as result of obesity or high fat feeding, can produce a state of insulin resistance as well as low grade inflammation. The early events leading from a rise of circulating FFA to activation of NF-κB are not clear but include several possibilities. First, Gao et al. have recently shown that the FFA mediated activation of IKK (a kinase involved in the activation of NF-κB) in fat cells was PKC dependent (34). Thus, DAG mediated PKC activation may be an upstream effector of NF-κB activation connecting the lipid metabolite with the inflammation hypothesis. Second, there is evidence to suggest that FFA mediated activation of IKK and NF-κB may be, at least in part, mediated by the Toll like receptor 4 (TLR-4) (35). The TLR-4 pathway, which is essential for the development of innate immunity to pathogens, triggers production of inflammatory cytokines (36). Thus it appears, that the sensing of excess nutrients such as FFA and the sensing of infectious pathogens may use the same signaling pathway.

Possible mechanisms through which FFA mediated NF-κB activation and cytokine production can result in insulin resistance includes activation of JNK by cytokines (37,38) and induction of suppressor of cytokine signaling (SOCS) which can interfere with binding of IRS1/2 to the insulin receptor (39).

However, whether FFA induced inflammation actually causes insulin resistance in vivo is not certain. On one hand, there is little doubt that the very high concentration of proinflammatory cytokines observed during severe bacterial infection or burns or increasing TNF-α blood levels 10-fold by infusion of rh TNF-α, can result in insulin resistance (40). Also, Salicylate, a non-specific inhibitor of IKK, given in high doses (4-8 g/day), increased insulin sensitivity, supporting a role of NF-κB activation in the pathogenesis of insulin resistance (41,42). On the other hand, the evidence that the cytokine levels typically observed in response to lipid infusions or in obese people contribute to the insulin resistance, is less convincing (reviewed in ref. 43). For instance, TNF-α and IL-6 are considered important inhibitors of insulin action. Yet, administration of TNF-α neutralizing antibodies to obese diabetic patients did not reduce their insulin resistance (44,45), nor did infusion of IL-6 into normal volunteers produce insulin resistance (40) and IL-6 deficient mice are insulin resistant (46,47). This, however, does not exclude the possibility that even modest elevations of several of these cytokines together, may produce insulin resistance.

The oxidative and endoplasmic reticulum (ER) stress hypotheses

In response to converting excessive amounts of FFAs and other nutrients into fat, the adipocyte can develop signs of oxidative and ER stress. In healthy volunteers, a lipid infusion induced a rise in plasma FFA levels and produced an increase in plasma free radical concentrations (indicating oxidative stress) (48). In cultured adipocytes, Furukawa et al. have shown that FFAs activated NADPH oxidase and induced reactive oxygen species (ROS) production and that the ROS induced oxidative stress resulted in dysregulated production of proinflammatory cytokines from white adipose tissue (49). These proinflammatory cytokines may then be able to produce insulin resistance (see above).

In vitro, FFA have been shown to produce ER stress in cultured adipocytes (50), liver cells (51) and pancreatic β-cells (52,53). In vivo, heparinized lipid infusions increased plasma FFA and produced ER stress in rat liver (Boden et al., unpublished). ER stress can produce insulin resistance via activation of JNK (54,55), whereas relieving ER stress by overexpression of ORP150, a molecular chaperone that protects cells from ER stress, reduces insulin resistance in diabetic (dbdb) mice (56). Thus, there is evidence supporting
the notion that FFA induced oxidative and/or ER stress may contribute to the FFA mediated insulin resistance.

CONCLUSIONS

The pivotal role of elevated plasma FFA levels as a cause for insulin resistance and as a pathogenetic factor in the development of T2DM has been established. In addition, it has recently been recognized that FFAs simultaneously cause insulin resistance and activate the proinflammatory NF-κB pathway resulting in secretion of many proinflammatory and proatherogenic cytokines and chemokines. Thus, elevated plasma FFA levels in obese subjects can produce a low grade inflammatory state which may contribute to the accelerated atherosclerosis and non-alcoholic fatty liver disease, conditions whose prevalence is several-fold increased in obesity.

Challenges for the future include the prevention or correction of obesity and elevated plasma FFA levels mainly through decreased caloric intake and increased caloric expenditure, development of easy, fast and reliable methods to measure FFA in small blood samples and the development of efficient pharmacological approaches to normalize increased plasma FFA levels.

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REFERENCES


First demonstration that adipose tissue lipolysis and plasma FFA levels coincide with changes in the number of adipose tissue macrophages. These results suggest that plasma FFA levels and local lipid fluxes regulate adipose tissue macrophage recruitment.


- Acutely increasing plasma FFA levels produces insulin resistance
- Lowering of chronically elevated plasma FFA levels improves insulin resistance
- Elevated plasma FFA levels can account for up to 50% of the insulin resistance in obese patients with type 2 diabetes
- Acutely increasing plasma FFA levels is proinflammatory
Figure 1.
Relationship between positive energy balance, an expanded adipose tissue (obesity), insulin resistance and cardiovascular disease. A chronic positive energy balance results in an expanded adipose tissue, i.e., obesity and is associated with insulin resistance and the development of serious disorders including type 2 diabetes, hypertension, dyslipidemia and disorders of coagulation and fibrinolysis which are all independent risk factors for the development of cardiovascular disease.
### Table 1

Criteria for a physiologic link between obesity and insulin resistance.

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