Tracing the fate of dietary fatty acids: metabolic studies of postprandial lipaemia in humans

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Abstract

Most postprandial studies have investigated the response of a single meal, yet the ingestion of sequential meals is more typical in a Western society. The aim of this review is to explain how natural and stable isotope tracers of fatty acids have been used to investigate the metabolism of dietary fat after single and multiple meals, with a focus on in vivo measurements of adipose tissue metabolism. When stable isotope tracers are combined with arteriovenous difference measurements, very specific measurements of metabolic flux across tissues can be made. We have found that adipose tissue is a net importer of dietary fat for 5 h following a single test meal and for most of the day during a typical three-meal eating pattern. When dietary fat is cleared from plasma, some fatty acids ‘spillover’ into the plasma and contribute up to 50% of postprandial plasma non-esterified fatty acid concentrations. Therefore, plasma non-esterified fatty acid concentrations after a meal reflect the balance between intracellular and extracellular lipolysis in adipose tissue. This balance is altered after the acute ingestion of fructose. The enzyme lipoprotein lipase is a key modulator of fatty acid flux in adipose tissue and its rate of action is severely diminished in obese men. In conclusion, in vivo studies of human metabolism can quantify the way that fatty acid trafficking modulates plasma lipid concentrations. The magnitude of fatty acid flux from adipose tissue has implications for ectopic fat deposition in tissues such as the liver and muscle.

The immediate fate of dietary fat is important to health. Indeed, atherogenesis has been described as a postprandial phenomenon (1), and postprandial lipaemia (the rise in plasma triacylglycerol (TAG) concentrations after a meal) is thought to be involved (2,3). Although the exact mechanisms that link postprandial lipaemia remain to be elucidated, small chylomicron remnants have been implicated in the progression of coronary artery disease (4). Postprandial lipaemia has also been shown to be associated with oxidative stress and inflammation as recently reviewed (5). Therefore it is important to understand factors that influence the duration and magnitude of postprandial lipaemia. A key tissue in the disposal of meal fatty acids is adipose tissue. The importance of adipose tissue in this respect is outlined in the so called adipose tissue expandability hypothesis (6). This hypothesis proposes that ‘a failure in the capacity for adipose tissue expansion, rather than obesity per se is the key factor linking positive energy balance and type 2 diabetes’. With increasing adiposity in some individuals, it is proposed that the capacity of adipose tissue to store further TAG is reduced, and lipids begin to accumulate in other tissues. The aim of this paper is to summarise the metabolic studies in humans performed by ourselves and others that have helped to understand the way in which adipose tissue metabolises and stores dietary fat.

The single meal model

A typical protocol for studying postprandial lipaemia is to study volunteers after an overnight fast followed by a single meal. Whilst fasting, the role of adipose tissue is to release fatty acids into the systemic plasma in order to supply tissues with a high requirement for fatty acids, such as skeletal muscle and the heart. Thus fasting plasma NEFA concentrations are high (Figure 1). After a test meal is given, adipose tissue metabolism is coordinated in order to deal with the nutrient load that is given. A typical metabolic response to a high fat meal in a healthy non-obese male is shown in Figure 1. As fat enters the bloodstream in the form of chylomicron-TAG, the concentration of plasma TAG increases. The increase in plasma TAG is also partly due to an
increase in the concentration of endogenous TAG in very low density lipoprotein (VLDL, synthesised in the liver) \(^{(7)}\) and plasma taken after a meal containing fat is often cloudy because these large lipoproteins scatter light. The concentration of plasma TAG starts to fall as it is cleared from the plasma (Figure 1). This is largely mediated by the action of adipose tissue lipoprotein lipase (LPL) \(^{(8)}\), situated at the capillary endothelium. LPL hydrolyses chylomicron-TAG, releasing fatty acids to be taken up by adipose tissue. This pathway is upregulated by chylomicron-TAG, which increases rapidly in response to the carbohydrate content of the meal (Figure 1). It has long been known that LPL is inhibited by apoCII but it has recently been shown that LPL is also inhibited physiologically by angiopoietin-like protein (Angptl)-4 \(^{(9)}\). The expression of this protein appears to be decreased in response to food, thus lifting the inhibition and allowing TAG hydrolysis to proceed at a higher rate. Studies from knock-out mice have shown that LPL also appears to require GPIHBP1, a cell-surface glycoprotein synthesised by the endothelium \(^{(10)}\).

In contrast to the increase in concentration of plasma TAG after a mixed meal, the concentration of plasma NEFA rapidly decreases, but often rebounds above postabsorptive values at the end of the postprandial period (Figure 1). The initial decrease is due to the action of plasma insulin on the suppression of intracellular lipases. Thus insulin is a key mediator of changes in adipose tissue fatty acid trafficking in the transition from fasting to fed states; any meal given with no carbohydrate (e.g. pure fat load) would fail to illicit the metabolic responses that depend on the increase in plasma insulin concentrations. However, adipose tissue is very sensitive to insulin. Therefore, even a small insulin excursion can reduce plasma insulin concentrations (Figure 2, after fructose ingestion).

**The two-meal model**

The single meal model is very useful in understanding the key pathways involved in the metabolism of dietary fatty acids, but in reality, most people on a Western diet would eat a second meal less than 5-8 h later. It became clear that postprandial TAG metabolism became complicated by a second, later meal, giving an unusual plasma TAG profile with more than one peak \(^{(11)}\). We further investigated plasma TAG concentrations in response to two meals by using naturally occurring fatty acids in foods as tracers of metabolism \(^{(12)}\). After an overnight fast, we gave healthy volunteers a high-fat breakfast that was enriched in linoleic acid (18:2 n-6). Five h later, the volunteers consumed a lunch meal that was enriched in oleic acid (18:1 n-9). After the second meal, there was a rapid peak in chylomicron TAG, and the fatty acid composition of this early peak was remarkably similar to that of the breakfast meal. This suggested that the breakfast fat had been residing in a storage pool before being released by a stimulus from lunch. In an elegant study, Robertson et al \(^{(13)}\) obtained biopsy material from volunteers who had consumed fat 5 h previously, and used electron microscopy to show that the fat was in fact residing in the enterocyte, and that it was released in response to a nutrient stimulus.

**Spillover fatty acids**

An interesting observation from the two-meal study described above was that the plasma NEFA profile was quite unusual. Whilst we normally expect plasma NEFA concentrations to decrease after a meal, an increase was observed in response to lunch. Moreover, the composition of plasma NEFA after lunch was similar to that of breakfast fat (and the ‘early chylomicron TAG’ peak). This gave support to the notion that when chylomicron TAG is hydrolysed by LPL, not all the
fatty acids are taken up; some spillover or ‘leak’ into the systemic plasma \(^{14-16}\). ‘Spillover’ fatty acids can be quantified using stable isotope tracers. The assumptions are that LPL does not discriminate between different fatty acids originating from the same meal \(^{17}\), and that spillover fatty acids derive solely from chylomicron-TAG ie that within the time course of the study, appearance in the plasma NEFA fraction of meal fatty acids that have arisen via recycling through VLDL-TAG is minimal \(^{18}\).

In the following example, the contribution of whole body ‘spillover fatty acids’ to the systemic plasma NEFA pool has been calculated. The data are from a study in which we investigated the effect of fructose v glucose on postprandial lipaemia. Two hundred and fifty mg of \[^2\text{H}_2\]palmitate was given as part of a liquid test meal and was traced into chylomicron-TAG. The appearance of \[^2\text{H}\]palmitate in the plasma NEFA fraction was therefore assumed to be due to spillover from LPL-mediated hydrolysis. From this data it was possible to derive the proportion of fatty acids in the total plasma NEFA pool that had arisen from spillover (Figure 2). It was found that the higher plasma NEFA concentration observed after the glucose test meal (compared with fructose) was entirely due to the higher concentration of fatty acids originating from the spillover route (Figure 2). In other words, spillover fatty acids account for the ‘rebound’ effect whereby plasma NEFA concentrations are higher at the end the study period than the beginning (Figure 1). The proportion of chylomicron fatty acids that spillover has been found to be highest (as much as 80 %) in the late postprandial period \(^{19}\). This may be due to the fact that fatty acids are less easily taken up into adipose tissue against a concentration gradient ie when adipose tissue lipolysis is higher \(^{20,19}\). This would be in accordance with observations that tissues without a high lipolytic flux such as forearm muscle would appear to have as much as 100 % efficiency in terms of the uptake of LPL-derived meal fatty acids \(^{19}\).

**The technique of arteriovenous difference**

Adipose tissue metabolism can be quantified in specific adipose tissue depots by the technique of arteriovenous difference. Blood is sampled simultaneously from an artery (or arterialized vein) and a small vein draining the adipose tissue. The difference in concentration of plasma or blood metabolites between the two sites represents metabolism during one pass through the tissue. We have used this technique to study adipose tissue metabolism of meal fatty acids, initially in human abdominal subcutaneous tissue \(^{21}\) and more recently in the femoral depot \(^{22}\). In order to quantify flux through the tissue, adipose tissue blood flow (ATBF) must be measured allowing the net transcapillary flux of fatty acids across the tissue can be calculated. This represents the net balance of different pathways of fatty acid trafficking through the tissue. In the postabsorptive state, there is a large concentration gradient between plasma NEFA entering the tissue (artery) and plasma NEFA leaving the tissue (adipose venous drainage). In a group of healthy men (BMI 23-34), the mean values were 591 ± 41 and 1208 ± 131 respectively \(^{23}\), and the mean net flux of NEFA from the tissue was 1080 nmol per 100 g tissue per min. Using an average adipose tissue fatty acid composition \(^{24}\), this equates to approx 250 µg adipose tissue TAG hydrolysed per 100 g tissue per min. For a person in energy balance with a 20 kg total fat mass this represents approx 24 g TAG to be replaced (0.12 % of adipose tissue TAG).

**Adipose tissue uptake of meal fatty acids**

Using the technique of arteriovenous difference and measurements of mass balance, adipose tissue has been found to be a net importer of fatty acids for 5 h following a mixed meal in a number of
studies as reviewed (25). The uptake of meal fatty acids can be studied more specifically using either radioactive (26) or stable isotope tracer methodology. With the addition of stable isotope tracers to our protocol of arteriovenous difference described above, we were able to develop a very powerful model to study adipose tissue metabolism in vivo. In this model, \([^{13}\text{C}]\text{palmitic acid}\) is given orally, in order to represent exogenous fatty acids, and \([^{2}\text{H}_2]\text{palmitate}\) is given as a continuous intravenous infusion, in order to label endogenous pools (plasma NEFA and VLDL-TAG), see Figure 3. Using a specific antibody to apoB100, combined with immunoaffinity chromatography, we are able to separate VLDL from chylomicrons (27). This represents a very important advancement in the study of postprandial fatty acids as there is a large overlap in the size and density of these two lipoprotein classes.

In the study of healthy men mentioned above (23), we traced 100 mg \([^{13}\text{C}]\text{palmitic acid}\) into the chylomicron fraction where it was measurable at 60 min and peaked at 240 min. We calculated TAG extraction in adipose tissue as (arteriovenous difference of \([^{13}\text{C}]\text{palmitic acid}\) in plasma TAG) x ATBF, and this peaked at 120 min. The time course of fractional TAG extraction is very interesting. At 60 min after the test meal, it is as high as 30 % in some individuals but falls to about 10 % at 180 min. Presumably, this is because it is only during the early time points that newly released chylomicrons are available for hydrolysis. These newly-released chylomicrons would be comparatively large, and therefore a good substrate for LPL (28). In addition, at later time points, plasma TAG-[\(^{13}\text{C}\)]palmitic acid is no longer confined to chylomicrons, and may represent newly synthesised VLDL-TAG. We also calculated the fractional extraction of plasma \([^{2}\text{H}_2]\text{palmitate-TAG}\), representing the hydrolysis of VLDL-TAG. This was less than the fractional extraction of plasma \([^{13}\text{C}]\text{palmitate-TAG}\), indicating preferential hydrolysis of chylomicron TAG rather than VLDL-TAG, in agreement with in vitro findings that LPL binds to chylomicrons with a greater affinity than VLDL (28). Moreover, we demonstrated preferential uptake of fatty acids derived from chylomicron-TAG compared with fatty acids taken up directly from the plasma NEFA pool. The latter pathway has only recently been recognised as a significant route of fatty acid uptake into adipose tissue (29).

**Adipose tissue dietary fatty acid metabolism in response to a high-carbohydrate diet**

It is well known that a high carbohydrate diet, particularly one that is high in extrinsic sugars, leads to an increase in fasting plasma TAG concentrations (30). We explored the capacity of subcutaneous adipose tissue to ‘cope’ with clearing meal fatty acids in the context of hypertriacylglycerolaemia promoted by the ingestion of a high carbohydrate, low fat diet for three days (31). This rather extreme diet (75 % energy from carbohydrate) lead to a doubling in fasting plasma TAG from 1.0 to 2.0 mmol/L compared with a diet that was only 45 % energy from carbohydrate. In response to a standard test meal, the iAUC for postprandial lipaemia was similar after both diets. However, after the high carbohydrate diet, postprandial plasma TAG concentrations did not return to baseline by the end of the study (360 min). Fasting and postprandial plasma NEFA concentrations were identical after the two dietary regimens. We traced the meal fat with \([^{13}\text{C}]\text{palmitic acid}\) and found that plasma \([^{13}\text{C}]\text{palmitate-TAG}\) extraction and TAG clearance in adipose tissue were unaffected by the background diet. Thus it would seem that despite an increase in the fasting concentration of plasma TAG after the high-fat diet, the ability of adipose tissue to dispose of dietary fat was unaffected. The metabolism of other tissues was however affected; TAG clearance (also measured by arteriovenous difference) was significantly lower across the forearm after the high carbohydrate diet versus the high fat diet.
Also meal fatty acids tended to be repartitioned away from oxidation, towards esterification in the liver and muscle in response to short-term adaptation to the high carbohydrate diet.

24 hour studies of human adipose tissue metabolism

To follow-on from the arteriovenous difference and two-meal models described above, we investigated postprandial fatty acid metabolism over a 24h period, in order to study the response to three typical Western-style mixed meals (18). We used a continuous intravenous infusion of [2H2]palmitate as before, to label systemic fatty acids and used three unique uniformly-labelled fatty acid tracers to trace the meal fatty acids. To label the breakfast meal, we used [U13C]linoleic acid, and to label lunch and dinner, we used [U13C]oleic acid and [U13C]palmitic acid respectively. The uniformly labeled notation indicates that all the carbon atoms are stable isotopes and an advantage of this analytically is that there is minimal background (when using GC-MS) compared with fewer carbons being labeled. Additionally, the high density of labeling offers greater potential for tracing metabolic pathways (32). The [U13C]fatty acid tracers appeared in the plasma TG pool after each respective meal but although the concentration started to fall after about 5h, there always remained some tracer present. Thus, 24 h after the ingestion of [U13C]linoleic acid, there was measurable [U13C]linoleic acid in the plasma TG pool. This was assumed to represent chylomicron remnants and tracer that had become incorporated into VLDL-TAG via hepatic recycling. Likewise, [U13C]linoleic acid in the plasma NEFA pool, representing spillover fatty acids was still present at 24 h. Using a radioactive tracer of meal fatty acids, it has been estimated that 0.9 % of meal fatty acids remain in the circulation after 24 h (33). In our study, we were able to calculate the total transcapillary flux of fatty acids across subcutaneous abdominal adipose tissue in healthy non-obese men. This calculation represents the total fatty acids, from all pools, crossing in/out of adipose tissue from the plasma. Over the 24 h period, there was net uptake of fatty acids immediately after the first meal, and this continued until approximately 17 h after breakfast i.e during the whole of the daytime adipose tissue takes up and stores fatty acids. Although the plasma TAG taken up after a meal was mainly from chylomicrons, a proportion was taken up from VLDL-TAG. The quantitative significance of this pathway increased with each meal and made up one-third of the total of the transcapillary flux after the 3rd meal. Plasma NEFA were also taken up directly, but this remained small. We estimated the amount of fat from the meal that was taken up into adipose tissue at a whole body level. This increased from 15 % of the meal fat after the first meal, to 48 % after last meal. The calculation is only approximate since it depends on an assumption that regional variation of adipose tissue metabolism is not significant. However, the increase in stored fat with time is still a valid observation. This observation was dependent on an increase in LPL rate of action during the 24 h period, but even more so by an increase in fatty acid re-esterification (i.e. stepwise formation of TAG via the action of the enzymes monoacylglycerol transferase and diacylglycerol transferase). These lean subjects had a 2-fold increased LPL rate of action over 24 h but a 3-fold increase in net fatty acid uptake.

Meal fatty acid handling in insulin resistance/obesity

Fasting plasma TAG concentrations are typically higher in obesity, and this is associated with enhanced hepatic secretion of VLDL (34). This is associated with higher postprandial TAG concentrations due to increased VLDL and chylomicron remnants (35). This is also associated with a prolonged lipaemia, representing delayed clearance of dietary fat. The higher overnight fasting
and postprandial concentration of VLDL-TAG in obesity are intimately related to plasma insulin concentrations (35). We recently found that the higher postprandial TAG concentrations in insulin resistant, compared with insulin sensitive men (matched for BMI) was accounted for differences in the contribution of splanchnic sources (from de novo lipogenesis, visceral adipose tissue lipolysis or hepatic TAG) rather than from dietary sources (36). We found that meal fat clearance, as measured by clearance of [U\(^{13}\)C]palmitate in chylomicron TAG was lower in the insulin resistant men.

The burden of the higher fasting plasma TAG that often accompanies obesity can be expected to impact upon postprandial adipose tissue metabolism. We used the 24 h model described above to compare the abdominal subcutaneous adipose tissue metabolism of healthy lean and obese men (37). Despite higher fasting and postprandial plasma TAG concentrations, plasma NEFA concentrations were very similar throughout the whole of the 24 h period. How plasma NEFA homeostasis in the postprandial period is maintained in the face of an expanded adipose tissue mass and disturbed TAG metabolism is an interesting question. Using data derived from the intravenous infusion of \([^{2}H_2]\) palmitate to calculate RaNEFA, we were able to determine that whole body lipolysis at all time points was considerably down-regulated when expressed per unit of fat mass. At the tissue level, adipose tissue NEFA release was markedly lower in the obese men. Perhaps even more dramatically, the hydrolysis of meal fatty acids was severely diminished in the obese men; LPL rate of action was very low and failed to be upregulated during the course of the day (Figure 4). Total fatty acid trafficking was considerably lower in the obese group. The adipose tissue quiescence seemed to be achieved via two main mechanisms. Firstly, gene expression of key enzymes relating to fatty acid trafficking such as LPL, DGAT1, DGAT2 and HSL were lower in a similar group of obese men (37). Secondly, subcutaneous adipose tissue blood flow (ATBF) was markedly lower and very unresponsive for the whole day in the obese compare with lean group. This is in agreement with findings after a single meal; ATBF is highly associated with insulin sensitivity (38). ATBF is integral to adipose tissue function, and an impaired blood flow will lead to a reduced supply of TAG substrate and reduced removal of products of metabolism within the adipocyte or at the capillary endothelium. The difference in adipose tissue handling of meal fatty acids meant that overall, even accounting for differences in adipose tissue mass, the obese group stored a lower proportion of fatty acids from the three meals in adipose tissue. Notably, they did not increase the efficiently of storage over the day, as was found in the lean group. The implications for this are important, an excess of meal fatty acids would lead to increased exposure to other tissues and could lead to the possibility of ectopic fat deposition.

Metabolism of specific fatty acids

Some of the measurements described above have been made with the assumption that palmitic acid behaves as a typical fatty acid. Differences in the postprandial metabolism of specific fatty acids can be studied in one of two ways. Either different fatty acids are given within the same test meal, and compared, or the response of different meals (containing different fatty acids) is compared.

We have investigated the adipose tissue metabolism of different fatty acid species consumed in a single test meal using the technique of arteriovenous difference, using commercially produced structured triacylglycerols or natural fats. Perhaps surprisingly, we found that adipose tissue did not discriminate between the fatty acids that we were testing for either type of test meal. Using structured TAGs containing oleic acid plus either stearic or palmitic acid, we
found no difference in LPL-mediated hydrolysis of the different fatty acids, or in their subsequent uptake by the tissue. The structural position of the individual fatty acids within the TAG molecule was also unimportant. We investigated the metabolism of different fatty acids given in the same test meal (14:0, 16:0, 16:1 n-7, 18:0, 18:1 n-9, 18:2 n-6, 20:5 n-3, 22:6 n-3). Although the molar proportion of the fatty acids in the meal was not maintained in chylomicrons, net uptake into adipose tissue was entirely proportional to their presence in chylomicrons. Nonetheless, overall, this meant that the storage of meal fatty acids into adipose tissue was in the order n-3 polyunsaturated < saturated < n-6 polyunsaturated < monounsaturated; oleic acid was stored in the greatest amount. Note that these studies did not use stable isotope tracers; therefore represent net uptake. We have recently compared the metabolism of stable isotope tracers of 16:0 ([U-13C]palmitate), 18:1 n-9 ([U-13C]oleate) and 18:2 n-6 ([U-13C]linoleate; all tracers were given simultaneously in the same test meal. We found that their incorporation into chylomicron-TAG was similar. However there was a tendency for the incorporation of 18:1 n-9 to be highest. This pattern was maintained as the fatty acids became incorporated into plasma NEFA and VLDL-TAG. This was explained by the low partitioning of [U-13C]oleate into plasma cholesteryl ester (CE) and phospholipid (PL) fractions, and also erythrocyte PL. Meal fatty acids are taken up into adipose tissue mainly via chylomicron-TAG, VLDL-TAG and plasma NEFA. Therefore the isotopic studies have shown that preferential uptake of meal oleate, compared with other meal fatty acids, could occur due to greater availability in these fractions.

Another aspect of comparing the metabolism of different fatty acids is to compare their postprandial metabolism when given separately. In this case, the fatty acids could behave differently due to different physico-chemical properties of the chylomicrons formed from the different test meals. This is difficult to study in humans because of the rapidity by which chylomicrons are cleared in the blood. However, animal studies in which the lymph has been accessed have shown that different dietary fatty acids form different sized chylomicrons. For example linoleic acid forms larger chylomicrons than palmitic acid, and there is some evidence to suggest that this is the case in humans too, as reviewed in (39).

**Regional differences in fatty acid metabolism**

Not only is the amount of adipose tissue important to health, body fat distribution is also important. The role of visceral fat (upper body fat, associated with a large waist) in relation to the metabolic syndrome has recently been reviewed (40) but the relationship is not clear (41,42). Paradoxically, greater accumulation of fat on the hips is beneficial in terms of risk of myocardial infarction (43).

Because of the importance of body fat distribution, studies have attempted to look at mechanisms behind differential accumulation of fat in different depots. Regional meal fat storage has been studied by giving a 3H or 14C fatty acid tracer with a meal and then measuring specific activity in adipose tissue depots 24 h later. An advantage of radioactive tracers in this respect is that they are very sensitive; an important consideration since the tracer is dispersed into a large pool.

Regional differences in the uptake of meal fatty acids into adipose tissue depots has been comprehensively investigated by Jensen et al using a [3H]triolein tracer given with a liquid test meal (33). Fatty acid oxidation was determined by incorporation of the tracer into the plasma water pool and it was estimated that approximately 50% of the tracer was oxidized over the 24 h.
Intrabdominal fat took up the tracer with the greatest avidity (Table 1) but the upper body depot, being the largest depot, accounted for over half of the uptake of fatty acids not oxidized. After a high-fat, high-energy meal, women store more dietary fatty acids in leg fat than men (44). Moreover, meal fatty acid storage (mg meal fat/g adipose tissue lipid) was greater in women with more leg fat than those with less leg fat. Also, after a high-fat meal, the depots with more fat seemed to take up meal fat less avidly than those with less fat.

Mcquaid et al have recently developed a technique to apply the technique of arteriovenous difference to femoral adipose tissue by cannulation of the saphenous vein (22). In accordance with the work of Jensen and colleagues (Table 1), it was found that net uptake of meal fatty acids was lower in femoral than subcutaneous abdominal tissue. ATBF was also lower in the femoral depot. An interesting finding was that whilst chylomicron-TAG was the preferred source for postprandial fat deposition in the subcutaneous abdominal depot, the femoral depot discriminated less against VLDL-TAG. It was therefore hypothesized that femoral adipose tissue may accumulate dietary fatty acids that have been recycled as VLDL and non-esterified fatty acids.

Conclusions

Healthy adipose tissue adapts rapidly to the ingestion of a mixed meal. Intracellular lipolysis is immediately suppressed, reducing the flux of fatty acids leaving the tissue. An increase in ATBF facilitates the entry of meal fat into the tissue, thus providing substrate for LPL which is activated by the action of insulin. The fractional extraction of chylomicrons is very efficient, as much as 30% in the early postprandial period. It is possibly higher than this at earlier time points. Not all fatty acids released by the action of LPL are taken up by the tissue; some are released into the plasma, particularly in the late postprandial period where as much as 50% of the plasma NEFA pool is composed of meal fatty acids. Whilst these dietary fatty acids may be taken up by non-adipose tissues, there is the potential to recycle back to adipose tissue and be taken up directly from the plasma NEFA pool. Adipose tissue is a net importer of dietary fat for 5 h following a single test meal and for most of the day during a typical three-meal eating pattern. The action of LPL seems to increase sequentially after meal intake, but uptake of meal fatty acids into adipose tissue increases to a greater extent, suggesting upregulation of pathways of esterification. Obesity and insulin resistance are associated with higher fasting and postprandial plasma TAG concentrations and reduced efficiency of adipose tissue meal fat storage. This offers a possible explanation to explain ectopic fat deposition associated with obesity. Hypertriglycerolaemia due to a high carbohydrate diet in non-obese individuals does not affect the ability of adipose tissue to clear dietary fat. There is a marked difference in the way that different adipose tissue depots handle dietary fat and this may be related to the metabolic phenotypes associated with different body fat distribution patterns.
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References

36. Hodson L, Bickerton AS, McQuaid SE, et al. (2007) The contribution of splanchnic fat to VLDL-triglyceride is greater in insulin resistant than insulin sensitive men and women: studies in the postprandial state. Diabetes
Figure legends

Figure 1. Typical postprandial concentrations of plasma metabolites in a healthy male after a mixed meal. Data are from Bickerton et al (23).

Figure 2. Origins of plasma NEFA after ingestion of glucose or fructose, 0.75 g sugar/kg body wt plus 500 mg [2H2]palmitate. Data taken from Chong et al (45). Data are shown as mean and SEM, n=12. Systemic plasma NEFA concentrations are shown as filled circles (P<0.05 comparing effect of sugars (45). Fatty acids estimated to have arisen from adipose tissue intracellular lipolysis are shown as open circles (P=ns comparing effect of sugars). The difference is the estimated concentration of plasma NEFA derived from dietary-TAG ‘spillover’ (see text), calculated as follows: Spill over fatty acids (µmol/L) = pNEFA (µmol/L)/ (% palmitate in chylomicron-TAG) * 100, where pNEFA is the concentration of fatty acids in the plasma NEFA pool derived from chylomicron spillover. pNEFA = concentration of [2H2]palmitate in plasma/chylomicron TTR. TTR = the tracer tracee ratio for [2H2]palmitate in plasma/chylomicron.

Figure 3. Pathways of postprandial adipose tissue fatty acid trafficking studied by isotopic labelling of plasma lipid pools. Endogenous pathways are labelled using an intravenous infusion of [2H2]palmitate (potassium salt) complexed with human albumin. This equilibrates with systemic plasma non-esterified fatty acids (NEFA) and is taken up by the liver where it is esterified to triacylglycerol (TAG) and exported in very low-density lipoprotein (VLDL). Exogenous pathways are labelled by the ingestion of [13C]palmitic acid with a test meal. This is incorporated into chylomicron TAG but soon appears in the plasma NEFA pool via the ‘spillover route (a) after hydrolysis by the enzyme lipoprotein lipase (LPL). [13C]Palmitic acid also becomes incorporated into VLDL in the liver via plasma NEFA uptake, and through chylomicron remnant uptake. In adipose tissue, such recycled VLDL-TAG fatty acids are not thought to contribute quantitatively to the spillover route. Fatty acids can be taken up directly from the plasma NEFA pool (b).

Figure 4 Meal fatty acid uptake into adipose tissue. Adipose tissue lipoprotein lipase (LPL) rate of action (assumed to be equivalent to TAG extraction) and uptake of LPL derived fatty acids (TAG extraction minus spillover fatty acids) in 9 lean (A) and 10 abdominally obese men (B), over a 24 h period, during which three mixed meals were consumed, at t=0, 5 and 10 h (dotted lines). Data are calculated from fatty acid stable isotope tracers, from data in the study of McQuaid et al (37).