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Thermic effect of food and sympathetic nervous system activity in humans

L Tappy

Institute of Physiology, University of Lausanne, Lausanne, Switzerland

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Summary — The intake of nutrients is known to increase energy expenditure. Measured thermic effects of nutrient are 0–3% for fat, 5–10% for carbohydrates and 20–30% for proteins. Stimulation of adenosine triphosphate (ATP) hydrolysis during intestinal absorption, initial metabolic steps and nutrient storage are responsible for this food thermic effect. The autonomic nervous system modulates the thermic effect of nutrients. Parasympathetic muscarinic antagonists reduce the thermic effect of orally administered nutrients, most likely by delaying gastric emptying and decreasing the amount of nutrient storage. Antagonists of the beta-adrenoreceptors decrease the thermic effect of glucose. The part of glucose induced thermogenesis which is eliminated by beta-adrenergic antagonists has been called 'facultative thermogenesis' and takes place, at least in part, in skeletal muscle. Insulin-induced stimulation of muscle sympathetic nerve activity may be involved in this facultative thermogenesis. The thermic effect of food is reduced in obese, insulin-resistant patients. The effect of thermogenesis represents about 50–150 kcal/day in such patients, and can explain only a minor part of their excess body weight. Defective thermogenesis may, however, contribute to weight gain, or impair weight loss in such patients.

energy expenditure / sympathetic nervous system / parasympathetic nervous system / obesity


dépense énergétique / système nerveux sympathique / système nerveux parasympathique / obésité
THERMIC EFFECT OF FOOD

It has long been recognized that oral or parenteral administration of nutrients in healthy individuals is associated with a stimulation of energy expenditure. This increase over baseline of energy expenditure associated with feeding is referred to as the thermic effect of food (TEF). In humans, the measurement of energy expenditure is most frequently performed using indirect calorimetry (Jéquier and Felber, 1987). With this technique, the thermic effect of a nutrient or of a mixture of nutrients can be calculated in two different ways: either a bolus of the nutrient is administered orally or intravenously, and the energy expenditure is monitored until the nutrient has been absorbed and disposed of completely and energy expenditure has returned to baseline values. In these conditions, the thermic effect of the nutrient (expressed as a percentage of its energy content) can be expressed as the incremental area above baseline of energy expenditure divided by the energy content of the nutrient administered; or the nutrient can be administered continuously (either parenterally or enterally) until steady states in both energy expenditure and the plasma concentrations of the nutrient administered and of its metabolites are attained. In these conditions, the thermic effect of the nutrient can be expressed as the difference between energy expenditure observed after its administration and basal energy expenditure, divided by the rate of nutrient energy administered (Acheson, 1993).

Measured TEF are 5 to 10% of the caloric content of carbohydrates administered, 0 to 3% of that of lipids and 20 to 30% of that of proteins or amino acids (Flatt, 1978; Ravussin et al, 1983; Thiébaud et al, 1983a,b; Tappy and Jéquier, 1993). In healthy subjects over a 24 h period, TEF represents about 10% of the total amount of energy expended (Ravussin et al, 1986).

MECHANISMS RESPONSIBLE FOR THE THERMIC EFFECT OF FOOD

In rodents, energy expenditure can be dramatically increased by exposure to cold or carbohydrate overfeeding. In these conditions, the animals oxidize large amounts of fat and glucose in brown adipose cells (Himms-Hagen, 1995). These cells form a specialized thermoregulatory organ, in which oxidative phosphorylation can be uncoupled and nutrients oxidized to release heat without synthesis of adenosine triphosphate (ATP). This uncoupling of oxidative phosphorylation in brown adipose tissue of rodents can be triggered by the activation of the sympathetic nervous system. Although brown adipose tissue is present in normal newborn humans and in human adults under special circumstances (such as pheochromocytoma), there is no conclusive evidence that the brown adipose tissue plays an important role in the control of TEF in normal human adults (Astrup, 1986; Brundin and Wahren, 1991). Rather, it appears that TEF in adult humans is related to the stimulation of energy-requiring processes during the postprandial period. The intestinal absorption of nutrients, the initial steps of their metabolism and the storage of the absorbed, but not immediately oxidized nutrients, all require ATP hydrolysis, and therefore consume a certain amount of energy. As illustrated in figure 1, the net amount of energy provided by a nutrient which can be used by the organism to produce mechanical or chemical work is therefore equal to the total amount of ATP molecules synthesized upon oxidation of the nutrient minus the amount of ATP which has been hydrolyzed in the initial steps of the metabolism of the nutrient. Consequently, the ratio of ATP used in initial metabolic steps to ATP made upon complete oxidation of the nutrient can provide an index of the thermic effect of the nutrient. Furthermore, the amount of ATP
The thermic effect of a nutrient is related to the number of adenosine triphosphate (ATP) used in the initial steps of its metabolism. ADP: adenosine diphosphate; b: Total ATP synthesis = total energy expenditure; a: ATP used for activation or storage of A to A⁺; b–a: net ATP available for energy-requiring processes; thermic effect of A is proportional to a/b.

Fig 1. The thermic effect of a nutrient is related to the number of adenosine triphosphate (ATP) used in the initial steps of its metabolism. ADP: adenosine diphosphate; b: Total ATP synthesis = total energy expenditure; a: ATP used for activation or storage of A to A⁺; b–a: net ATP available for energy-requiring processes; thermic effect of A is proportional to a/b.

Fig 2. Both glucose and fructose yield 38 adenosine triphosphate (ATP) synthesized from 38 adenosine diphosphate (ADP) upon complete oxidation. The number of ATP used prior to oxidation varies, however, according to the sugar considered and to the metabolic pathway followed prior to its oxidation, resulting in variable thermic effects.
metabolism. Compared to the thermic effect of 75 g oral glucose (6.5%), the thermic effect of fructose is substantially higher (10.2%) (Tappy et al, 1986). This is explained by the fact that fructose metabolism requires the hydrolysis of one additional ATP for the phosphorylation of glyceraldehyde, which accounts for the increased thermic effect of this sugar (Tappy and Jéquier, 1993) (fig 2).

When glucose is converted into glycogen, 2.5 mol ATP are also consumed, 0.5 mol for glucose absorption, 1 mol for glucose-6 phosphate synthesis and 1 mol for regeneration of uridine diphosphate (UDP) to uridine triphosphate (UTP) after uridyl diphospho-glucose synthesis. The thermic effect of glycogen synthesis from exogenous glucose is therefore about 2.5/38 = 6.6%. If glucose is released thereafter from muscle glycogen to be oxidized within the muscle, another mole of ATP will be required for the synthesis of fructose-1,6 diphosphate and the overall thermic effect will be 3.5/38 = 9.2%. If glycogenolysis occurs in the liver and glucose is released into the systemic circulation, 2 additional moles of glucose will be hydrolyzed for the synthesis of fructose-1,6 diphosphate, bringing the thermic effect to about 11%.

It can be seen then, that the amount of ATP used in metabolism can vary dramatically according to the metabolic pathway used. Changes in the rate of utilization of the endogenous substrate and the activation of energy requiring processes, such as glycogen synthesis, are believed to be responsible for the TEF.

ROLE OF THE AUTONOMIC NERVOUS SYSTEM IN TEF

A large number of studies have demonstrated that TEF can be modulated by pharmacological agents acting on the autonomic nervous system. Inhibition of the parasympathetic muscarinic receptors decreases the thermic effect of oral (Nacht et al, 1987), but not intravenous (Deriaz et al, 1989) glucose. This effect has been ascribed to a decreased glycogen synthesis due to a delay in gastric emptying and slower carbohydrate absorption (Schneeberger et al, 1991).

The beta-adrenergic receptors blockade decreases the thermic effect induced by infusions of glucose + insulin (Acheson et al, 1983). This part of the TEF of intravenous glucose which can be suppressed by beta-adrenergic antagonists has been called ‘facultative thermogenesis’ (Acheson et al, 1984). Activation by the sympathetic nervous system of energy-requiring processes, such as the Cori cycle (Kusaka and Ui, 1977) and the lipolysis-reesterification of free fatty acids (Wolfe et al, 1987; Breitenstein et al, 1990), is likely to be involved in this facultative portion of TEF. However, it was shown that a 2 week treatment with beta-adrenergic antagonists did not alter the overall energy expenditure of healthy humans (Acheson et al, 1988). This suggested that the facultative thermogenesis of intravenous glucose may be related to sustained increases in plasma insulin concentrations.

The role of insulin per se in glucose-induced thermogenesis has been investigated in several studies. Interestingly, insulin infusion has been reported to stimulate the sympathetic nervous system as demonstrated by an increase in plasma catecholamine concentrations (Rowe et al, 1981). More recently, it has been observed in healthy humans through in vivo microneurography of the lateral peroneal nerve that insulin specifically stimulates the sympathetic nerve fibers targeted to skeletal muscle (Andersen et al, 1991). This stimulation of muscle sympathetic nerve activity by insulin may be responsible for part of the facultative TEF, which has been suggested to take part predominantly in muscle (Astrup
et al, 1985). The relationship between the insulin-induced activation of the sympathetic nervous system in skeletal muscle and facultative thermogenesis remains far from being clearly demonstrated. Fructose administered either orally (Tappy et al, 1986) or intravenously (Schwarz et al, 1992) markedly stimulates energy expenditure. Moreover, a substantial portion of this fructose thermic effect can be inhibited by beta-adrenergic antagonists (Tappy et al, 1986; Schwarz et al, 1992). Fructose administered intravenously, however, does not elicit an increase in sympathetic muscle activity (Vollenweider et al, 1993), suggesting that other mechanisms are involved in the facultative thermogenesis of this sugar. In addition, the activation of muscle sympathetic activity by a lower body negative pressure (Brown et al, 1966; Scherrer et al, 1988) fails to stimulate basal energy expenditure (Tappy et al, 1995). This suggests that sympathetic muscle activity per se is not thermogenic. Interactions between actions of insulin and the sympathetic nervous system in muscle may be postulated to modify facultative, glucose-induced thermogenesis.

TEF IN OBESITY

TEF has been extensively studied in the context of the pathogenesis of obesity. Several studies initially reported that the thermic effect of glucose is decreased in obese patients (reviewed in Tappy et al, 1991). Furthermore, it was observed that glucose-induced thermogenesis is further depressed in post-obese patients (Schutz et al, 1984b). These results suggest that a defect in TEF may contribute to a positive energy balance and weight gain in pre-obese individuals. Other studies, however, markedly moderated these early conclusions. First, it was observed that a decreased TEF was not present in every obese patient. In fact, the studies that reported a normal TEF in obese patients are almost as common as those in which a decreased TEF was observed (D'Alessio et al, 1988). Second, it was noted that the decreased TEF of obese patients was observed after glucose feeding but not after fructose (Simonson et al, 1988) or amino acid administration (Tappy et al, 1993). Furthermore, the decrease in glucose-induced thermogenesis, when present in obese patients, was related to insulin resistance. During euglycemic, hyperinsulinemic clamps, overinfusion of insulin in obese subjects resulted in an overall glucose disposal similar to that observed in lean subjects and normalized glucose-induced thermogenesis (Ravussin et al, 1985). Third, it was realized that the decreased TEF in obese patients represented only about 50–150 kcal/day (Schutz et al, 1984a). Individuals who gain weight increase both their fat mass and lean body mass. As a consequence of this increased body mass and fat-free mass, it can be estimated that 24 h energy expenditure increases by 20–25 kcal for each kg body weight gained due to an increase in both basal energy expenditure and the energy expended in physical activity (Ravussin et al, 1986; Owen, 1988). As a consequence, the decreased TEF of obese patients can account for a gain of 2–8 kg, which is far less than the excess body weight of many obese individuals. This suggests that a defective TEF cannot be held as the major cause of weight gain in obese patients. It may nevertheless contribute to weight gain or the relapse of obesity after weight loss in some obese subjects.

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