Brown adipose tissue activity in pyrexial cases of cot death

M E J LEAN, G JENNINGS
From the Dunn Nutrition Laboratory, Medical Research Council and University of Cambridge, Cambridge

SUMMARY Brown adipose tissue was investigated in two cases of cot death in which core temperatures were above 40°C on arrival at the mortuary. Evidence was obtained from mitochondrial (8-³H) guanosine diphosphate (GDP) binding and oxygen uptake of active thermogenesis with uncoupled mitochondrial respiration which was recoupled by GDP. Thermogenic capacity of brown adipose tissue, estimated by radioimmunoassay of the specific mitochondrial membrane "uncoupling protein" responsible for heat production, was similar to that measured in other infants or in experimental animals acclimated to moderately warm conditions (23°C).

Brown adipose tissue thermogenesis, occurring inappropriately in a warm, well insulated infant, could be a cause of some cases of cot death.

Some cot deaths, or sudden infant death syndrome (SIDS), have recently been considered to be related to problems with thermoregulation, resulting in overheating. Overinsulation with clothes or bedding may be an important factor, especially if the room temperature is kept warm, but excessive endogenous heat production could also play a part. Suggestions as to the source of excess heat production have not been made, but Naeye and coworkers reported increased amounts of brown adipose tissue in cot death infants.

The possibility that the activity of brown adipose tissue could, in some instances, be involved in the mechanisms leading up to cot death by generating heat inappropriately is difficult to explore. We have, however, had the opportunity of investigating the properties of brown adipose tissue in two pyrexial cases of cot death from which it was possible to obtain fresh tissue. The results suggest that brown adipose tissue is thermogenically active in such cases. Although its activity may not be different from that of normal infants, thermogenesis in this situation is inappropriate and could indeed contribute to death.

Material and methods

The two infants were both male and aged 5 months: they had died at home in their cots between 7–8 am. Body weights were 7 and 8 kg, respectively. These were both typical cases of cot death: previous developmental history and health were unremarkable and no underlying cause of death was found. The rectal temperature in each case, however, was above 40°C on arrival at hospital, and postmortem examinations were carried out within two hours of death.

Adipose tissue was removed from sites of axillary, cervical, interscapular and perirenal brown adipose tissue, and cytochrome oxidase activity (E.C.1.9.3.1) was measured as an index of mitochondrial mass by a spectrophotometric method. For electron microscopical examination, tissue was fixed in osmium tetroxide and embedded in Spurr's resin and sections were stained with uranyl acetate and lead citrate. Mitochondrial preparations were prepared within two hours, and thermogenic activity was assessed by a (8-³H) guanosine diphosphate (GDP) binding assay, according to previously described methods. Mitochondrial uncoupling protein content was measured as an index of thermogenic capacity using a solid phase radioimmunoassay with a lower limit of detection of 0.8 µg/mg total mitochondrial protein, using a specific rabbit anti-human uncoupling protein antiserum. In case 2 direct measurements of mitochondrial respiration were made using an oxygen electrode (YSI Co, Ohio, USA) at 25°C with 10 mM α-glycerophosphate as substrate in a medium (pH 7.1) containing 100 mM sucrose, 10 mM glycyglycine, 5 µM rotenone and bovine serum albumin 2.5 mg/ml. Initial respiration rate was obtained and GDP added (1 mM final concentration) to obtain the fully coupled respiration rate. Finally, 5 µM of the uncoupling agent FCCP (Sigma Chemical Co, Poole, Dorset) was added to estimate fully uncoupled oxygen uptake.
Biochemical measurements made on fresh brown adipose tissue from cases 1 and 2 are shown in the table: histological appearances were typical of active brown adipose tissue, with high vascularity, rounded nuclei, and multivesicular cytoplasm packed with mitochondria, and without evidence of necrosis (fig 1).

Uncoupling protein content was significantly correlated with cytochrome oxidase \( (r = 0.97, n = 6, p < 0.001) \) and GDP binding activity \( (r = 0.85, p < 0.02) \). All were highest in axillary and cervical sites. Scatchard analyses (fig 2) at GDP concentrations of 0.5–25 \( \mu \text{M} \) were interpreted as indicating single site mitochondrial GDP binding characteristics with dis
brown fat in cot deaths

Active thermogenesis, such as oxygen utilisation however, possibly indicating hypoxia, tissue. Other in reported primates oxidase protein appears much more that roughly equivalent greatest in acclimated to warm tissue.6

Acute stimulation. Its association constant (Kₜ) of 6·3 μM (axillary) or 5·1 μM (perirenal). Basal oxygen uptake of mitochondria from case 1 was similar to that after the addition of the uncoupling agent FCCP, indicating substantially uncoupled respiration in the basal state. Addition of GDP produced a clear reduction in oxygen uptake, indicating recoupling of respiration.

Discussion

The similarity of brown adipose tissue in human infants to the “hibernating gland” of small animals was recognised by Hatai in 1902.7 It is present in widespread intra-abdominal and paravascular sites in man, usually distinguishable from subcutaneous white adipose tissue by its darker colour and firmer consistency, features which derive from lower lipid content, higher cytochrome content, and greater vascularity. Its thermogenic function seems to be the same in all species studied, through uncoupled respiration regulated by catecholamines, dependent on the unique mitochondrial uncoupling protein. In a previous study we showed activation of brown adipose tissue thermogenesis and increased mitochondrial uncoupling protein content in adipose tissue from adults exposed to high circulating catecholamines from phaeochromocytoma.8

Uncoupling protein is generally present in detectable amounts in mitochondria from infant brown adipose tissue.6 The concentration was similar in 10 cot death infants (perirenal 6·6 ± 1·6 μg/mg, axillary 11·7 ± 2·2 μg/mg protein) to the remainder of a group of 17 infants who died suddenly from a variety of causes (table), which does not suggest any chronic stimulation. Acute sympathetic stimulation of brown adipose tissue may still contribute to some cot deaths, however. Increased synthesis of uncoupling protein in response to cooling or to noradrenaline occurs only over a period of two days or more, although the mRNA that codes for synthesis of the uncoupling protein appears much more rapidly.9 Cytochrome oxidase and GDP binding activities in both babies reported here showed characteristics similar to those in other species and suggested a thermogenic activity roughly equivalent to the brown adipose tissue of mice acclimated to warm laboratory conditions, being greatest in the axillary and cervical sites, which in primates comprise the largest single block of brown adipose tissue.

Naeye interpreted an increased amount of brown adipose tissue in cot deaths as a result of tissue hypoxia, possibly indicating pulmonary disease.3 Hypoxia, however, usually leads to a suppression of oxygen utilisation for thermogenesis10 as this could only continue at the expense of other vital organs. Active thermogenesis, such as we have shown near the time of death in two cases of cot death, involves increased oxygen uptake and would clearly be inappropriate in the setting of hypoxia. This consideration questions the recent suggestion of Stephenson and Variend11 that necrosis of brown adipose tissue in cot death results from continued metabolic activity in hypoxic tissue. The viral infective cause suggested by Grist and Urquhart seems more plausible,12 given the long known attractiveness of brown adipose tissue as a site of primary viral multiplication.13

The oxygen uptake of maximally stimulated brown adipose tissue in 23°C acclimated mice is about 1·5 ml O₂/minute, equivalent to a heat production of 8–10 calories/minute/g tissue.14 If Hull’s figure15 of 30 g of brown adipose tissue in a neonate is accepted, then the heat output under maximal acute stimulation, but without any long term trophic stimulation, would be about 250 calories/minute. Thermogenesis of this order would increase the basal metabolic rate of a neonate by about 200%—more than sufficient to account for the observed metabolic response to cold temperature or noradrenaline infusion, and enough to raise core temperature by 5–6°C/hour. The lower thermogenesis of brown adipose tissue of adult rats, acclimated to a temperature of 33°C and stimulated by noradrenaline measured at 0·4 ml O₂/minute/g by Foster and Frydman,16 would still produce an almost 100% increase in the metabolism of an infant with 30 g brown adipose tissue. If this occurred inappropriately and suddenly in a well clothed and insulated infant, the core temperature might be raised to dangerous levels and induce apnoea. Our biochemical findings indicate that this might have occurred in our two cases of pyrexial cot death. This conclusion can be made, irrespective of a separate question, which cannot be answered at present: is brown adipose tissue equally active in infants who have died of other causes?

We thank Professor A Gresham, department of morbid anatomy, School of Clinical Medicine, University of Cambridge, for his encouragement and help in providing rapid access to samples.

References


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Requests for reprints to: Dr M E J Lean, Diabetic Clinic, Woolmanhill Hospital, Aberdeen Royal Infirmary, Aberdeen AB9 1GS, Scotland.