Oxidation of medium-chain acyl-CoA esters by extracts of Aspergillus niger: enzymology and characterization of intermediates by HPLC

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The activities of β-oxidation enzymes were measured in extracts of glucose- and triolein-grown cells of Aspergillus niger. Growth on triolein stimulated increased enzyme activity, especially for acyl-CoA dehydrogenase. No acyl-CoA oxidase activity was detected. HPLC analysis after incubation of triolein-grown cell extracts with decanoyl-CoA showed that β-oxidation was limited to one cycle. Octanoyl-CoA accumulated as the decanoyl-CoA was oxidized. β-Oxidation enzymes in isolated mitochondrial fractions were also studied. The results are discussed in the context of methyl ketone production by fungi.

Keywords: β-oxidation, Aspergillus niger, methyl ketones, medium-chain fatty acids

INTRODUCTION

The development (Causey & Bartlett, 1986; Bartlett et al., 1988) and application (Watmough et al., 1989) of HPLC techniques for the separation and identification of intermediates in the β-oxidation of fatty acids has led to improved knowledge of the details of this process in mammalian mitochondria (Watmough et al., 1989), mammalian peroxisomes (Bartlett et al., 1990) and bacteria (Broadway et al., 1992). Detailed information has been obtained on the intermediates temporarily accumulating, rate-limiting step(s), metabolite channeling and degree of chain shortening.

It is widely held, although not universally (Yagi et al., 1991), that methyl ketones, which are important flavour compounds in blue cheeses, are formed by fungi because of incomplete β-oxidation (possibly peroxisomal) of medium-chain fatty acids (Lawrence & Hawke, 1968). The oxidation is supposed to stop with the formation of 3-ketoacyl-CoA esters. Decacylation is then followed by decarboxylation, either spontaneously or under the influence of a decarboxylase. The process has been studied by many workers over the past 70 years but the above hypothesis has never been proved. Questions such as which compartment of the cell is involved or why the oxidation aborts in the first cycle of oxidation have not been answered.

Kunau and co-workers (Kionka & Kunau, 1985; Thieringer & Kunau, 1991; Kunau et al., 1988) have provided much detailed enzymological data on peroxisomal β-oxidation in fungi, particularly Neurospora crassa. It seemed to us there was an opportunity to apply this type of enzymological work and HPLC techniques (Bartlett et al., 1988) for identifying the intermediates formed during methyl ketone production in fungi. This work is described here.

METHODS

Growth of the micro-organism. Aspergillus niger N423 (cpsAt nicAI) was grown routinely in Pontecorvo complex medium (Pontecorvo et al., 1953) with the modification that KH₂PO₄ concentration was increased 10-fold. Growth was achieved by inoculating 200 ml medium (in 1 litre conical flasks), containing a suitable carbon source and urea (10 mM), with 1 × 10⁸ spores. For isolation of mitochondria, growth was achieved by inoculating 200 ml yeast extract/peptone/glucose (YEPA, 1 litre conical flasks, containing a suitable carbon source, with 1 × 10⁸ conidia. YEPA contained (per litre): proteose peptone, 20 g; yeast extract, 10 g; glucose, 20 g; Pontecorvo vitamin solution, 10 ml. Cultures were incubated on an orbital shaker (145 r.p.m.) at 30°C. Solid cultivation and maintenance were performed on potato/dextrose agar (39 g l⁻¹).

Spores were obtained after growth on potato/dextrose agar at 30°C for 3–4 d or until sporulation. Spores were harvested by gentle suspension in a sterile solution of Tween 80 (0.8%, w/v) and subsequent filtration through glass-wool. Spore suspensions were stored at −20°C in portions (up to 1.5 ml) containing 1 × 10⁶ spores. The stock was renewed every 2–3 months.

Abbreviations: ABTS, 2,2′-azino-bis(3-ethylbenzthiazoline-6-sulphonic acid); DCPIP, 2,6-dichlorophenolindophenol; DTNB, 5,5′-dithiobis(2-nitrobenzoic acid); DTP, 4,4′-dithiodipyridine; PMS, phenazine methosulphate; triolein, 1,2,3-trioctadecanoylglycerol; YEPD, yeast extract/peptone/glucose.
Transference of mycelia. As the growth on certain carbon sources such as fatty acid methyl esters and triglycerides (especially the ones containing medium-chain fatty acids) was poor, the behaviour of A. niger after incubation with these substrates was studied by transference: after the first incubation, the mycelia were harvested by filtration through a pre-sterilized filter (Miracloth, Calbiochem), washed thoroughly with sterile distilled water and transferred aseptically to a second medium containing fatty acid, methyl ester, triglycerol, etc., before incubation for the required period.

Harvesting of mycelia and preparation of cell-free extracts. Mycelia were harvested at room temperature by vacuum filtration through Whatman no.1 filter paper and washed thoroughly with distilled water. The resultant cell cake was weighed (wet weight) and used for preparation of cell-free extracts, or stored at -20°C. Cell-free extracts were obtained by passing the cell-suspension (1 g cell cake per 15 ml 50 mM HEPES pH 7.5 containing 1 mM DTT) three times through a pre-cooled French pressure cell (35 MPa). (For some experiments studying methyl ketone production cell walls were weakened with 1% Caylase C and 0.25% cellulase at pH 9.5 and 37°C before passage through the French pressure cell.) The fungal homogenate was centrifuged at 7700 g for 10 min at 4°C to remove whole cells and cell debris. The supernatant was taken as cell-free extract and was either used immediately or stored in small aliquots (1.5 ml) in liquid nitrogen until required.

Enzyme assays. All enzyme assays were carried out at 30°C in a Unicam 8625 UV/VIS spectrophotometer.

Acyl-CoA dehydrogenase (EC 1.3.99.3). The reaction mixture included the following components (final volume 1 ml): 0.15 M HEPES pH 8.0, 0.6 ml; 1 mM DCPIP, 0.1 ml; 1 mM PMS, 0.05 ml; 2 mM acyl-CoA, 0.05 ml; water and enzyme preparation, 0.2 ml. The reaction was started by acyl-CoA addition. An absorbance coefficient of 21 500 M^-1 cm^-1 was used for DCPIP at pH 8.0. The assay follows the decrease in absorbance at 600 nm due to DCPIP reduction.

Acyl-CoA oxidase (EC 1.3.3.6). The reaction mixture included (final volume 1 ml): 100 mM HEPES pH 7.5, 0.8 ml; 25 mM ABTS, 0.05 ml; 1 mg ml^-1 peroxidase, 0.05 ml; 2 mM acyl-CoA, 0.05 ml; enzyme preparation, 0.05 ml. The reaction was started by acyl-CoA addition. The absorbance coefficient for the ABTS radical cation is 18 400 M^-1 cm^-1. The increase in absorbance at 405 nm due to ABTS radical cation was used.

Acetyl-CoA estimation. Acetyl-CoA was assayed using citrate synthase and oxaloacetate. CoASH released on condensation of acetyl-CoA and oxaloacetate was measured by reaction with DTNB. An aliquot of reaction mixture (200 µl) was added to trichloroacetic acid (60 µl, 15%, v/v) and precipitated protein was removed by centrifugation (5 min). Then 200 µl of the supernatant was added to 0.8 M Tris/HCl buffer pH 8.0 (0.8 ml) containing oxaloacetate (approx. 2 mg solid) and DTNB (250 nmol) and the absorbance at 412 nm (A412) was read against a blank assay (without substrate). Citrate synthase (pigeon breast muscle, 1 U) was added to the mixture and the increased absorbance at 412 nm (A412) measured. The amount of acetyl-CoA was given by the difference A412−A412 due to the release of CoASH from acetyl-CoA and subsequent reaction with DTNB. An absorbance coefficient of 13 600 M^-1 cm^-1 for DTNB was used.

Carnitine acyltransferase (EC 2.3.1.7). This assay is based on that of Ramsay & Tubbs (1975) for carnitine acetyltransferase using l-carnitine. The reaction mixture contained (final volume 1 ml): 250 mM sodium phosphate buffer pH 7.8, 0.1 ml; 0.3 mM acetyl-CoA, 0.3 ml; 1 mM DTP, 0.1 ml; 1 M KCl, 0.1 ml; water and enzyme preparation, 0.35 ml; 40 mM L-carnitine, 0.05 ml. The reaction was started by l-carnitine addition and an absorption coefficient of 19 800 M^-1 cm^-1 was used to calculate rates. The increase in absorbance at 342 nm, due to reduction of dithiobispyridine (DTP), was followed.

Citrate synthase (EC 4.1.3.7). The method follows the increase in absorbance at 412 nm due to mercaptide formation, resulting from the reaction between DTNB and CoASH (Parvin, 1969). The reaction mixture contained (final volume 1 ml): 0.5 M Tris/HCl pH 8.0, 0.2 ml; 1 mM oxaloacetate (freshly prepared in 100 mM Tris/HCl pH 8.0), 0.1 ml; 1 mM acetyl-CoA, 0.1 ml; 5 mM DTNB (prepared in 100 mM 13.75 mM Triton X-100, 0.02 ml; 12.5 mM CoASH, 0.02 ml; 45 mM ATP, 0.05 ml; 0.3 M MgCl2, 0.02 ml; cell-free extract and water, 0.145 ml; 12.5 mM potassium oleate (substrate), 0.02 ml. An aliquot of the previous assay (60 µl) was withdrawn and added to the following mixture: 50 mM sodium phosphate buffer pH 6.5, 2.6 ml; 25 mM ABTS, 0.15 ml; 1 mg peroxidase ml^-1, 0.15 ml; 1 mM H2O2, 0.05 ml. The decrease in absorbance at 405 nm due to free CoASH present in the assay was measured and subtracted from a blank in which the substrate was omitted. An absorption coefficient of 18 400 M^-1 cm^-1 for ABTS radical cation was used.

β-Oxidation indirect assay. In this assay, acyl-CoA esters are generated in situ by the acyl-CoA synthetases already present in the extract. The assay contained (final volume 1 ml): 0.5 M HEPES buffer pH 8.0, 0.5 ml; 40 mM NaN3, 0.1 ml; 40 mM MgCl2, 0.1 ml; 17 mM NAD+, 0.1 ml; 6 mM CoASH, 0.1 ml; cell-free extract and fatty acid (0.68 µmol, added in 7 µl DMSO). The reaction was initiated by substrate addition and activity was measured by following the increase in absorbance at 340 nm due to NAD+ reduction (ε340 = 6220 M^-1 cm^-1).

β-Oxidation direct assay. The assay contained (final volume 1 ml): 150 mM HEPES buffer pH 8.0, 0.6 ml; 17 mM NAD+, 0.1 ml; 6 mM CoASH, 0.05 ml; cell-free extract and acyl-CoA ester (100 nmol introduced in the assay as an aqueous solution). The reaction was started by acyl-CoA addition and activity was measured by following NAD+ reduction at 340 nm. All incubation mixtures contained between 0.1 and 0.15 mg of protein. NADH formation was routinely checked by addition of yeast alcohol dehydrogenase (30 µg) and acetaldehyde (0.5 µmol) to the assay mixture. The resulting decrease in absorbance at 340 nm (NADH oxidation) corresponded to the NADH formed in the assay.

Enzymes. All enzyme assays were carried out at 30°C in a Unicam 8625 UV/VIS spectrophotometer.

Acyl-CoA dehydrogenase (EC 1.3.99.3). The reaction mixture included the following components (final volume 1 ml): 0.15 M HEPES pH 7.5, 0.6 ml; 1 mM DCPIP, 0.1 ml; 1 mM PMS, 0.05 ml; 2 mM acyl-CoA, 0.05 ml; water and enzyme preparation, 0.2 ml. The reaction was started by acyl-CoA addition. An absorbance coefficient of 21 500 M^-1 cm^-1 was used for DCPIP at pH 8.0. The assay follows the decrease in absorbance at 600 nm due to DCPIP reduction.

Acyl-CoA oxidase (EC 1.3.3.6). The reaction mixture included (final volume 1 ml): 100 mM HEPES pH 7.5, 0.8 ml; 25 mM ABTS, 0.05 ml; 1 mg ml^-1 peroxidase, 0.05 ml; 2 mM acyl-CoA, 0.05 ml; enzyme preparation, 0.05 ml. The reaction was started by acyl-CoA addition. The absorbance coefficient for the ABTS radical cation is 18 400 M^-1 cm^-1. The increase in absorbance at 405 nm due to ABTS reduction was followed (Werner et al., 1970).

Enoyl-CoA hydratase (crotonase; EC 4.2.1.17), L-3-hydroxyacyl-CoA dehydrogenase (EC 1.1.1.35) and 3-ketoacyl-CoA thiolase (EC 3.1.2.20). These enzymes were assayed according to Binstock & Schulz (1981).

Acyl-CoA hydrolase (EC 3.1.2.11). This assay is based on that of Ramsay & Tubbs (1975) for carnitine acetyltransferase. The assay mixture contained: 50 mM HEPES pH 7.5, 0.5 ml; water, 0.3 ml; 1.2 mM DTP, 0.1 ml; 2.1 mM acyl-CoA, 0.05 ml; enzyme preparation, 0.05 ml. The reaction was started by acyl-CoA addition. An absorption coefficient of 19 800 M^-1 cm^-1 for DTP was used to calculate rates. The increase in absorbance at 324 nm due to reduction of DTP was followed.

Acyl-CoA synthetase (EC 6.2.1.3). The assay was undertaken in two steps: first the formation of the acyl-CoA ester by acyl-CoA synthetase, and second the detection of the amount of CoASH consumed, by reaction with ABTS. The first step of the assay was based on the assay for acyl-CoA synthetase of Ichihara & Shibasaki (1991) and contained (in a final volume of 0.5 ml): 0.43 M MOPS/NaOH buffer pH 7.6, 0.175 ml;
β-Oxidation in Aspergillus niger

Tris/HCl pH 8.0, 0.05 ml, and water. The reaction was started with enzyme addition. The molar absorption coefficient for DTNB was 13600 M⁻¹ cm⁻¹.

Protein content. Protein was estimated by the method of Bradford (1976), using BSA as a standard.

Synthesis of acyl-CoA esters

Enoyl-CoA esters. Enoyl-CoA esters were synthesized from the saturated acyl-CoA esters by acyl-CoA oxidase. The mixture contained (final volume 1 ml): 0.2 M sodium phosphate buffer pH 6.5, 0.75 ml; 5 U acyl-CoA oxidase ml⁻¹, 0.05 ml; 2 mM acyl-CoA ester, 0.05 ml; the reaction was started by acyl-CoA addition.

3-Hydroxyacyl-CoA esters. 3-Hydroxyacyl-CoA esters were synthesized from the saturated acyl-CoA esters by acyl-CoA oxidase and enoyl-CoA hydratase. The reaction was undertaken in two steps: first the formation of the enoyl-CoA, and second the hydration of the enoyl-CoA double bond followed at 263 nm. The reaction mixture contained (final volume 1 ml): 0.2 M sodium phosphate buffer pH 6.5, 0.75 ml; 5 U acyl-CoA oxidase ml⁻¹, 0.05 ml; 5 U enoyl-CoA hydratase ml⁻¹, 0.005 ml; 2 mM acyl-CoA ester, 0.05 ml. The reaction was started by acyl-CoA addition.

β-Ketoacyl-CoA esters. β-Ketoacyl-CoA esters were formed from the corresponding saturated acyl-CoA esters by the combined action of acyl-CoA oxidase, enoyl-CoA hydratase and L-3-hydroxyacyl-CoA dehydrogenase. This synthesis is based on the assay of thiolase described by Seubert et al. (1968). The assay follows the increase in absorbance at 303 nm due to the formation of the enol form of the β-ketoacyl-CoA ester. The assay mixture contained (final volume 1 ml): 0.13 mM Tris/HCl buffer pH 8.9, 0.75 ml; 0.2 M MgSO₄, 0.025 ml; 10 mg ml⁻¹ BSA, 0.05 ml; 40 mM sodium pyruvate, 0.0025 ml; 20 mM NAD⁺, 0.025 ml; 2 mM acyl-CoA esters, 0.05 ml; 970 U lactate dehydrogenase ml⁻¹, 0.005 ml; 5 U acyl-CoA oxidase ml⁻¹, 0.045 ml; 5 U enoyl-CoA hydratase ml⁻¹, 0.005 ml (freshly diluted from a 50 U ml⁻¹ solution); 10 U L-3-hydroxyacyl-CoA dehydrogenase ml⁻¹, 0.045 ml. The reaction was started by the addition of the acyl-CoA ester.

Extraction and analysis of acyl-CoA esters by HPLC. Acyl-CoA esters were extracted from β-oxidation reaction mixtures as described by Causey & Bartlett (1986) with minor modifications. An aliquot (0.5 ml) taken from a β-oxidation assay was quenched by the addition of glacial acetic acid (100 μl). An internal standard (acyl-CoA ester, 50 nmol) was added and the mixture extracted three times with a total of 15 ml diethyl ether to remove fatty acids. Saturated (NH₄)₂SO₄ (50 μl) was added to the aqueous phase and a mixture of methanol/chloroform (2:1, v/v, 3 ml) added slowly. The mixture was kept for 20 min at room temperature. The salt/protein precipitate was then removed by centrifugation (1000 g, 5 min) and the supernatant separated. The resultant pellet was extracted with a further 3 ml methanol/chloroform (2:1, v/v) and the supernatants combined. The solvent was removed by evaporation under a gentle stream of N₂ at 55 °C. The residue was resuspended in 200 μl KH₂PO₄ (50 mM, pH 5.3) and 20 μl analysed by HPLC.

HPLC was carried out with a system equipped with a Kontron 420 HPLC pump and a Kontron 425 HPLC gradient former using a Lichrosorb 10RP18 column (250 mm x 4.6 mm) at room temperature. Column eluates were monitored by using a Kontron 432 HPLC detector at 260 nm. Peak areas were determined by a Kontron I-459 integrator. Acyl-CoA esters were resolved by the following gradient of acetonitrile in 50 mM KH₂PO₄ (pH 5.3): isocratic 5% (v/v) for 5 min, linear to 40% (v/v) for 10 min, linear to 50% (v/v) for 5 min. The flow rate was 1 ml min⁻¹ throughout and all solvents were continuously de-aerated with helium.

Extraction and determination of methyl ketones, secondary alcohols and fatty acids. After harvesting the mycelia, the pH of the spent medium was adjusted to 1–2 with H₂SO₄ and internal standards were added (methyl ketone and fatty acid). The medium was then extracted twice with chloroform (2 x half the volume of medium) and the chloroform layers collected and combined. The residual water was removed with anhydrous MgSO₄ and the solvent reduced down to a convenient volume by rotary evaporation.

Methyl ketones, secondary alcohols and fatty acids were analysed by using a PYE Unicam PU 4500 chromatograph. equipped with an integrator and a flame ionization detector, fed with hydrogen (44 ml min⁻¹) and air (600 ml min⁻¹). The apparatus was fitted with a 10% (w/v) DEGS (diethylene glycol succinate) packed column (130 cm x 4 mm), using nitrogen as a carrier gas (40 ml min⁻¹). The temperature of the column was maintained at 130 °C, and that of the injector and detector at 160 °C. Compounds were quantified by automatic integration.

Electron microscopy of mycelia. The method used for the preparation of cells for electron microscopy was that of Graves et al. (1976), slightly modified: after embedding, 4% (w/v) glutaraldehyde in 0.1 M cacodylate buffer pH 7.5 was used and fixation was in 1% (w/v) OsO₄.

Isolation of mitochondria. After harvesting and washing, the mycelia (6 g wet weight) were ground gently with washed sand (2 g per g mycelia) in 50 mM sodium phosphate buffer pH 7.0 containing 5 mM EDTA, 0.6 M sucrose and 0.5% (w/v) BSA (6 ml) isolation buffer (Mannela, 1982), in a pre-cooled pestle and mortar (3–4 min). The resultant paste was suspended in an additional 34 ml of the same buffer solution and PMSF (0.5 mM final concentration) added to the mixture. The homogenate was centrifuged at 1000 g for 10 min, the pellet was discarded, and the supernatant was filtered through two layers of muslin and centrifuged at 16000 g for 10 min. The resultant pellet was taken as the crude mitochondrial fraction; this was gently taken up in isolation buffer, washed once and resuspended. For further purification, the crude mitochondrial pellet was suspended in isolation buffer containing 25% (v/v) Percoll and centrifuged at 48000 g for 1 h. After centrifugation, the gradient was displaced with a 2 M sucrose solution. Fractions (1 ml) were collected from the top of the gradient.

Respiration measurements on mitochondria. The respiratory activity of the mitochondrial preparations was measured by the use of a Clark-type O₂ electrode (Rank Bros). The experiments were conducted at 30 °C and assays contained (in a final volume of 3 ml): 10 mM sodium succinate, 0.2 mM ADP, mitochondrial preparation and Lambowitz buffer (8 mM NaH₂PO₄, 8 mM Tris, 5 mM MgCl₂, 0.7 mM EDTA, 0.3 M sucrose, pH 7.2) (Lambowitz et al., 1972). Mitochondrial suspensions were disrupted on ice by sonication (3 x 20 s), prior to enzyme assays.

Chemicals. Saturated acyl-CoA esters (except acetyl-CoA), fatty acids, fatty acid methyl esters, medium-chain triacylglycerols, triolein (practical grade), acyl-CoA oxidase (from Candida sp.), acyl-CoA synthetase (from E. coli), crotonase, 3-hydroxyacyl-CoA dehydrogenase (type V), peroxidase (type I, from horseradish), citrate synthase (from pigeon breast muscle) and Percoll were purchased from Sigma. Alcohol dehydrogenase (EC 1.1.1.1, from yeast) and cellulase were
from Botrenger Mannheim. Caylase C₉ was from Cayla (Toulouse, France). DMSO was obtained from Fisons Scientific Apparatus. Authentic methylalkyl ketones and their respective secondary alcohols were obtained from Aldrich. Yeast extract, proteose peptone and potato/dextrose agar were from Oxoid. Acetyl-CoA was synthesized according to Simon & Shemin (1953).

RESULTS

Selection of micro-organism

Aspergillus niger was selected for this work. It showed rapid growth on glucose and on triolein and proved a good producer of methyl ketones, especially 2-heptanone and 2-nonanone. These findings confirm those of Yagi et al. (1990).

Methyl ketone production by cells and cell extracts

Fig. 1 shows the time course for the formation of decanoic acid and 2-nonanone on presenting triolein-grown A. niger with 0-5% (w/v) methyl decanoate. There appeared to be a slight lag in 2-nonanone production, but since the appearance of this product was likely to be consequent upon the liberation of decanoic acid from methyl decanoate by cellular lipase(s) this was not surprising. There was also the possibility that residual triolein or oleic acid associated with the cells interfered in the processing of decanoic acid. It was always very difficult to remove triolein from cells prior to transfer to a second medium. In an experiment using 1,2,3-tridecanoylglycerol as the substrate for 2-nonanone production, similar results were obtained, but now about 35 h was required to generate the levels of product seen at 4 h in Fig. 1. Presumably, the lipase(s) involved were much less active with 1,2,3-tridecanoylglycerol.

Incubation of glucose-grown A. niger with 0-5% (w/v) 1,2,3-tridecanoylglycerol for 16 h resulted in complete loss of cell viability and only about 30% of the initial cell mass could be recovered. The same effect was seen with 4 mM decanoic acid, suggesting that the effect with 1,2,3-tridecanoylglycerol was due to the release and accumulation of decanoic acid. Electron microscope examination of the cell residue showed no sign of mitochondria, peroxisomes, etc., and the cell walls were often ruptured. As the incubations produced some 2-nonanone initially, much as did the triolein-grown cells, it was tempting to suggest that methyl ketone production was somehow linked to those degradative effects. However, in experiments with cells pre-grown on triolein then incubated with 0-5% (w/v) methyl decanoate or 1,2,3-tridecanoylglycerol over 16 h the cells remained viable, showed no signs of damage in the electron microscope and, as shown in Fig. 1, produced 2-nonanone. It is not known why triolein-grown cells were more robust towards decanoic acid. Perhaps it was because these cells had a much enhanced β-oxidation activity (see below).

Incubations of extracts from triolein-grown cells with octanoic or decanoic acids (1 mM) and ATP (1-10 mM) and MgCl₂ (1-10 mM), or of the cell extracts with decanoyl-CoA (0-1 mM), yielded no measurable quantities of 2-heptanone or 2-nonanone over a 12 h period. Extracts of cells which, prior to cell breakage, had been producing 2-heptanone from octanoic acid, also failed to form 2-heptanone. After 24 h, some 2-heptanone was produced from octanoic acid, but in these cases, it was always possible to demonstrate the presence of mycelia. When cycloheximide (1 mg ml⁻¹) was included in the 24 h incubations, no 2-heptanone could be found.

Our results do not support the findings of Yagi et al. (1991), who reported the formation of 2-nonanone from the oxidation of decanoic acid by a cell-free system from Penicillium decumbens. We did find, as did Yagi et al. (1991), that cell extracts could be apparently separated into a heat-labile pellet (3000 g) and heat-stable supernatant fraction (3000 g) which when combined together generated 2-heptanone from octanoic acid. Also, we found that methyl ketone formation was stimulated (six- to sevenfold) by 1-3 mM Ca²⁺ (CaCl₂). However, our incubations required 24 h and not 4 h to generate 2-heptanone, and mycelia could always be detected.

Activities of β-oxidation enzymes

The activities of the enzymes of β-oxidation found in glucose- and triolein-grown A. niger are shown in Table 1. Some of the enzymes, but particularly acyl-CoA dehydrogenase, were induced by triolein. With triolein, the activities of the enzymes were very similar to those seen for Neurospora crassa (Kionka & Kunau, 1985) and Candida tropicalis (Dommes et al., 1983) after contact with oleic acid. This was a little surprising in view of the differences in substrate and assay conditions used, but it may be that these enzymes are rather insensitive to these changes. As with N. crassa (Kionka & Kunau, 1985) there was no trace of acyl-CoA oxidase.

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**Fig. 1.** Production of 2-nonanone by A. niger triolein-grown mycelia on incubation with methyl decanoate. Mycelia were first grown on triolein (0-5%, w/v) for 40 h and then transferred to a second incubation medium (Pontercorvo complex medium, 200 ml) containing 0-5% (w/v) methyl decanoate. ▲, 2-Nonanone; ●, 2-nonanol; ○, decanoic acid. Data are the means of duplicate experiments.
activity in *A. niger*; this contrasts with *C. tropicalis*, where a 700-fold induction of this enzyme activity [330 nmol min⁻¹ (mg protein)⁻¹] occurred on transfer to oleic acid (Dommes et al., 1983).

Activities for the $\beta$-oxidation enzymes in glucose-grown *A. niger* (Table 1) differed quite markedly from those of *N. crassa* (Kionka & Kunau, 1985) and *C. tropicalis* (Dommes et al., 1983). For acyl-CoA dehydrogenase, we found barely detectable levels of activity (see below) and these were much lower (50-fold) than found for the other organisms. For enoyl-CoA hydratase and 3-ketoacyl-CoA thiolase, our measured activities were by contrast much higher (10- to 100-fold).

Assays of acyl-CoA dehydrogenase as performed here used PMS as electron carrier and DCPIP as electron acceptor and they were very sensitive to the presence of any free thiol (e.g. CoASH) because it rapidly reduced the electron acceptors. Accordingly, with extracts containing an active acyl-CoA thioesterase there was an ‘apparent’ acyl-CoA dehydrogenase activity. Extracts of glucose-grown *A. niger* showed an acyl-CoA thioesterase activity of 2 nmol min⁻¹ (mg protein)⁻¹. When the effect of this on acyl-CoA dehydrogenase assays was corrected for, the resulting activities were very low (Table 1). Most (> 90%) of the observed activity was due to the acyl-CoA thioesterase. This no doubt meant that our estimate for acyl-CoA dehydrogenase in glucose-grown cells was inaccurate, although clearly much more informative than an uncorrected value. Kionka & Kunau (1985) and Dommes et al. (1983) used a different electron-acceptor system, but they did not mention acyl-CoA thioesterase or any necessary correction in discussing their data. Accordingly, it is not clear that our results for acyl-CoA dehydrogenase in glucose-grown cells can be compared. This problem does not arise with the results for triolein-grown cells of *A. niger*. In these extracts, the activity of the acyl-CoA thioesterase was about the same as for glucose-grown cells while the acyl-CoA dehydrogenase was 400-fold more active.

### $\beta$-Oxidation assays

Attempts to follow the $\beta$-oxidation of decanoyl-CoA by extracts of glucose-grown cells were unsuccessful because no significant formation of NADH could be demonstrated. This was not surprising in view of the very low activity for acyl-CoA dehydrogenase recorded in Table 1. 'Indirect' assays using decanoic acid, ATP, MgCl₂, CoASH and NAD⁺ and extracts of triolein-grown cells were also unsuccessful. The detection of activity here was made more difficult by the very low activity of acyl-CoA synthetase [25 nmol min⁻¹ (mg protein)⁻¹ with decanoic acid as substrate] in these extracts.

Assays of $\beta$-oxidation with extracts of triolein-grown cells incubated with decanoyl-CoA gave rapid and continuous production of NADH on addition of substrate. The reactions were completed within 30–60 min, by which time approximately 1 mol NADH was produced per mol added substrate. Similar results were obtained with octanoyl-CoA. Thus, the activity was easily detected, although the overall process was limited in extent. If, by contrast, the reaction mixture which contained about 0.01 U acyl-CoA dehydrogenase was supplemented with 0.1 U commercial acyl-CoA oxidase the initial rate of NADH production was increased 10- to 20-fold and after 20 min about 3 mol NADH were produced per mol added substrate.

The rates of decanoyl-CoA oxidation in $\beta$-oxidation and in acyl-CoA dehydrogenase assays were compared over a range (25–300 µM) of substrate concentrations at 30 °C, pH 8.0. Results were reproducible to within ± 10% and the Lineweaver–Burk plots were linear. Values of $V_{\text{max}}$ = 38 nmol min⁻¹ (mg protein)⁻¹, $K_m$ = 130 µM, and $V_{\text{max}}$ = 15 nmol min⁻¹ (mg protein)⁻¹, $K_m$ = 30 µM were obtained in $\beta$-oxidation and in acyl-CoA dehydrogenase assays, respectively. The fact that the two assays did not agree was not surprising because, even if $\beta$-oxidation was strictly limited by acyl-CoA dehydrogenase, the electron acceptor system was quite different for the two types of assay. The nature of the ‘natural’ electron acceptor for acyl-CoA dehydrogenase in $\beta$-oxidation assays was uncertain, especially so if the enzyme was of microbody origin.

### HPLC analysis of products of $\beta$-oxidation

Incubations of tetradecanoyl-CoA with extracts of triolein-grown *A. niger* over a 30 min period and analysis showed that the formation of dodecanoyl- and decanoyl-CoA could be identified as the reaction progressed. No other intermediates were found. Mixing of decanoyl-CoA with extracts of triolein-grown cells revealed a much more limited process. Only one CoA ester product could be identified over a 120 min period (Fig. 2) and this product accumulated as the decanoyl-
M. F. BALTAZAR, F. M. DICKINSON and C. RATLEDGE

**Table 2. Enzyme activities in fractions (S_{16000} and P_{16000}) of A. niger cell homogenate**

Myceila were grown on glucose (2%, YEPD) for 16 h. A crude mitochondrial pellet was prepared as described in Methods, and was suspended in isolation buffer. Homogenate fractions were disrupted (by sonication) for enzyme assays. S_{16000}, 16000 g supernatant (150 mg protein); P_{16000}, 16000 g pellet (20 mg protein).

<table>
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<tr>
<th>Enzyme</th>
<th>S_{16000} (%)</th>
<th>P_{16000} (%)</th>
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<tr>
<td>Citrate synthase</td>
<td>37.5 (49)</td>
<td>39 (51)</td>
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<tr>
<td>Enoyl-CoA hydratase</td>
<td>8 (50)</td>
<td>8 (50)</td>
</tr>
<tr>
<td>3-Hydroxyacyl-CoA dehydrogenase</td>
<td>0.1 (12.5)</td>
<td>0.7 (87.5)</td>
</tr>
<tr>
<td>3-Ketoacyl-CoA thiolase</td>
<td>5 (92.6)</td>
<td>0.4 (7.4)</td>
</tr>
<tr>
<td>Carnitine acetyltransferase</td>
<td>5-2 (55)</td>
<td>4.3 (45)</td>
</tr>
</tbody>
</table>

CoA decreased (Fig. 3). The experiment was repeated and the results were confirmed. The retention time of this compound identified it as octanoyl-CoA, which correlated well with the finding of 0.8 mol acetyl-CoA released per mol decanoyl-CoA consumed. With a 1 h incubation mixture in the absence of NAD^+, no octanoyl-CoA or acetyl-CoA was formed, but instead, 3-hydroxydecanoyl-CoA and 2-decenoyl-CoA in the proportion of 10:1. Now 3-hydroxyacyl-CoA dehydrogenase was inoperative and there was accumulation of intermediates prior to the blocked step. Similar findings have been found by other workers with other systems (Broadway et al., 1992; Stanley & Tubbs, 1974, 1975; Watmough et al., 1989). With decanoyl-CoA, NAD^+ and 0.1 U acyl-CoA oxidase included, a 20 min incubation produced dramatic change. The 10-fold increase in the decanoyl-CoA processing capacity gave rise in 5 min to >95% loss of the initial 100 nmol decanoyl-CoA substrate and formation of 40 nmol hexanoyl-CoA, 18 nmol 3-hydroxyoctanoyl-CoA or hexenoyl-CoA and 30 nmol octanoyl-CoA. After 20 min there appeared to be 95 nmol butanoyl-CoA and most of the other intermediates had disappeared. This experiment confirmed that the limited β-oxidation activity of triolein-grown cell extracts was primarily due to the low activity of acyl-CoA dehydrogenase. It also showed that when the rate-limiting step for the overall process was changed, then intermediates other than the saturated CoA esters accumulated, at least temporarily.

**β-Oxidation enzymes in mitochondria**

Mitochondria were obtained from non-induced, glucose-grown cells by sand grinding and centrifugation. The mitochondria exhibited respiratory ratios (P:O) of 1:5-2:0 on addition of 0.3 mM ADP to assays measuring O_2 uptake consequent upon sodium succinate (10 mM) oxidation. Density-gradient centrifugation of a mitochondrial fraction on a Percoll gradient showed one sharp band of citrate synthase (mitochondrial marker) in the middle of the gradient and little activity in peripheral fractions. The distribution of β-oxidation enzymes, together with citrate synthase and carnitine acetyltransferase, is shown in Table 2 for supernatant and mitochondrial fractions of crude cell extract. About
50% of the original mitochondria were apparently preserved intact. The distribution of the \( \beta \)-oxidation enzymes was rather variable and reflected the contributions of the mitochondrial and microbody fractions. However, the low activity of 3-ketoacyl-CoA thiolase in the mitochondria seemed noteworthy. The experiment was repeated and the results were essentially the same. Kionka & Kunau (1985) found only enoyl-CoA hydratase in mitochondria from sucrose-grown N. crassa.

Isolation of mitochondria was also attempted from triolein-grown cells (0-25%, w/v), but the results were less satisfactory. The cells were more difficult to extract and only 20% of the initial mitochondria remained intact, as judged by citrate synthase and carnitine acetyltransferase activities. The respiratory ratios (P:O) were 12–14 and density-gradient centrifugation of a mitochondrial fraction showed citrate synthase distributed throughout the gradient. Despite these deficiencies the distribution of the \( \beta \)-oxidation enzymes appeared much the same as indicated by the results of Table 2.

**DISCUSSION**

The main objective of the present work was to test the hypothesis (Lawrence & Hawke, 1968) that methyl ketone production from medium-chain-length fatty acids is primarily associated with \( \beta \)-oxidation. This view is widely held, but there is little direct experimental evidence to support it. An alternative scheme based on work with extracts of *Penicillium decumbens* was proposed (Yagi et al., 1991) involving formation of the secondary alcohol in the cell wall layer followed by oxidation to methyl ketone. We have reproduced observations of the type used to support the alternative mechanism, but we have been unable to reproduce the central requirement of cell-free extracts capable of producing methyl ketones from fatty acids.

Our studies showed that \( \beta \)-oxidation enzymes of *A. niger* were induced by growth on triolein with strong induction (400-fold) of acyl-CoA dehydrogenase. Electron microscopy also showed that large numbers of peroxisomes appeared over this period. However, the inability of extracts to produce 2-heptanone from octanoic acid showed that the specificity of \( \beta \)-oxidation enzymes cannot alone explain methyl ketone formation. Some additional factors are needed.

\( \beta \)-Oxidation experiments and HPLC analyses (Figs 2 and 3) showed that the capacity for \( \beta \)-oxidation of medium-chain-length fatty acids by extracts of induced cells was limited to one turn of the spiral. This limitation appeared to be entirely due to the rate-limiting activity of specificity of acyl-CoA dehydrogenase, because added acyl-CoA oxidase led to more extensive degradation and the appearance of a range of chain-shortened CoA-esters. The rate-limiting behaviour of acyl-CoA dehydrogenase is often encountered with \( \beta \)-oxidation systems as, for example, in *Corynebacterium* 7E1C (Broadway et al., 1992) and mammalian liver mitochondria (Wattmough et al., 1989). Peroxisomal \( \beta \)-oxidation in rat liver was shown to be incomplete although the degree of chain shortening varied with the experimental conditions (Osmundsen et al., 1979; Lazarow, 1982; Bartlett et al., 1990). The present work is unusual in describing a \( \beta \)-oxidation system predisposed to give only one turn of the cycle with medium-chain-length fatty acids. The finding seems to offer at least a partial explanation of why, for example, whole cells produced only 2-nonanone from decanoic acid and no 2-heptanone or 2-pentanone.

The results on \( \beta \)-oxidation do not provide the basis for a comprehensive explanation of methyl ketone formation from medium-chain-length fatty acids. Such an explanation could not be expected, in view of the inability of the extracts to synthesize these compounds. However, it is clear that octanoyl-CoA would be bound to accumulate from decanoate oxidation and if it was also the case that CoA supplies were limited, then a pool of 3-ketodecanoyl-CoA would form by equilibration through 3-ketoacyl-CoA thiolase with octanoyl-CoA and acetyl-CoA. Under these circumstances, deacylation of some 3-ketodecanoyl-CoA and decarboxylation of 3-ketodecanoate to 2-nonanone can be envisaged. Lawrence & Hawke (1968) suggested that methyl ketone formation is due to lack of free CoA and that deacylation of 3-ketoacyl-CoA allows CoASH to be recycled quickly to other parts of the cell. Also, Hatton & Kinderlehrer (1991) provided evidence that the rate of methyl ketone formation is increased when the cells are stressed by some form of nutritional stringency.

The results with isolated mitochondria raised the possibility that these organelles could be the source of methyl ketone synthesis because of low 3-ketoacyl-CoA thiolase activity and possibly also low concentrations of CoA. Some support for the idea also came from the finding that glucose-grown *A. niger* proved effective at producing 2-heptanone from a low concentration of octanoic acid (1 mM). Such cells contained very few microbodies initially although the number increased quickly in a 4 h incubation. Against this proposal, however, was the finding that the mitochondria from either glucose- or triolein-grown cells showed no detectable activity for acyl-CoA dehydrogenase or carnitine acetyltransferase. The latter was not a serious problem because Sherratt & Sparwary (1994) stated that medium-chain fatty acids can enter mitochondria. However, the lack of acyl-CoA dehydrogenase suggests that 3-ketoacyl-CoA esters could not accumulate in mitochondria unless there was an alternative route for supplying enoyl-CoA or 3-hydroxyacyl-CoA esters. The finding of high mitochondrial enoyl-CoA hydratase activity in both glucose- and triolein-grown cells (Table 2; see also Kionka & Kunau, 1985, with *N. crassa*) suggests that the substrates for the enzyme might be available under some circumstances.

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277
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