Demonstration of Succinic Dehydrogenase in the Mesosomes of the Mycelial Phase of Paracoccidioides brasiliensis

By SYLVA LEIVA AND L. M. CARBONELL
Centre for Microbiology, Instituto Venezolano de Investigaciones Científicas, Apartado 1827, Caracas, Venezuela

(Accepted for publication 10 March 1970)

SUMMARY

In the hypha of Paracoccidioides brasiliensis the succinic dehydrogenase activity is localized in the mesosomes, plasma membrane and mitochondria. Short permanganate fixation after incubation enhances the positive cytochemical reaction. The higher activity is localized in mesosomes and it is observed as high electron-dense zones and concentric ring-like patterns. The membrane shows a spotty reaction and the mitochondrion shows a positive zone in the intracristae space.

INTRODUCTION

Mesosomes or intracytoplasmic membrane systems have been demonstrated in several fungi (Edwards & Edwards, 1960; Moore & McAlear, 1961; Carbonell & Rodriguez, 1968). Light microscope demonstrations of reductases have been done in Saccharomyces cerevisiae (Reiss, 1967a), Neurospora crassa (Reiss, 1967b, 1968) and Paracoccidioides brasiliensis (Carbonell & Kanetsuna, 1966). Different types of oxido-reductase have been demonstrated cytochemically and biochemically in the mesosomes of Gram-positive bacteria (Vanderwinkel & Murray, 1962; van Iterson & Leene, 1964; Sedar & Burde, 1965; Ferrandez, Chaix & Ryter, 1966; Cesari, Rieber & Imaeda, 1969; Reaveley & Rogers, 1969). Morphologically, the mesosomes of fungi and bacteria are very much alike and the similarities could be confirmed if they were shown to have similar cytochemical characteristics. The purpose of the present work is to demonstrate succinic dehydrogenase activity in the mesosomes of Paracoccidioides brasiliensis.

METHODS

The mycelial phase of Paracoccidioides brasiliensis (IVIC-pb9) was inoculated into a GGY medium (glycine 1%, glucose 2%, yeast extract 0.2%) in Erlenmeyer flasks which were placed on a reciprocal shaker at 100 oscillations/min., stroke amplitude 5 cm., at 20 to 22°.

Four-day-old samples were harvested by low speed centrifugation and incubated for 30 to 50 min. in the following mixture: 3.5 ml 0.8 M-sodium succinate, tetranitro blue tetrazolium (TNBT) 1 mg./ml., which had been previously dissolved in N,N-dimethyformamide (Sedar, Rosa & Tsou, 1962; Sedar & Burde, 1965). For controls, sodium malonate (0.5 g.) was added to the incubated mixture as a competitive inhibitor; the material was incubated without substrate; and the enzyme was inactivated...
with glutaraldehyde, or by treating at 65° to 80° for 10 min. The material was then fixed.

As previously shown (Carbonell & Kanetsuna, 1966), the intrinsic factors which reduced the tetrazolium salt are eliminated by treating the samples with cold acetone for 10 min. before incubation, and washing in a mixture of 0.2 M-phosphate buffer (pH 7.4) and Ringer solution (1:9, v/v). Following incubation, the samples were harvested by low speed centrifugation. Acetate-veronal buffer of Michaelis, pH 6.1 (Kellenberg, Ryter & Séchaud, 1958), was used for preparing the fixative, for washing and for the agar solution.

After incubation, different fixation procedures were used: (a) 1% osmium tetroxide at 20 to 22° for 12 h.; (b) 5% glutaraldehyde at 4° for 12 h.; (c) fixation as in (b), and postfixed as in (a), for 3 h.; and (d) permanganate fixation for 10 min. or less at 4° following the technique of Kellenberg et al. (1958). To facilitate manipulation, the fixed material was embedded in 2% agar. It was then dehydrated with ethanol and embedded in Maraglas (Freeman & Spurlock, 1962). Sections were cut with diamond knives and examined with a Hitachi H1 electron microscope. For morphological details, samples were also fixed in glutaraldehyde and postfixed in osmium tetroxide, embedded in Maraglas and stained with lead nitrate and uranyl acetate.

RESULTS

The fine structure of the untreated mycelia of Paracoccidioides brasiliensis has been reported (Carbonell & Rodriguez, 1968). The following brief description will deal only with the features relevant to the present topic. The structure of the plasma membrane varied according to the sectioning angle. When attached to the cell wall it appeared as a wide electron-dense layer, or as a three-layered structure (an outer wide electron-dense layer, a low-density middle space and a very narrow inner electron-dense layer). The invaginations of the plasma membrane are interpreted as the beginning of mesosomes. This membranous system, formed by the apposition of the infolded plasma membrane, underwent additional invaginations, forming multivesicular or lamellar structures that are interpreted as tubular infoldings of the plasma membrane seen in different sectioning angles (Pl. 1, fig. 1; Pl. 3, fig. 7; Pl. 4, fig. 11). In addition to the plasmalemma and mesosomes, a cell wall with septum, septal bodies (Woronin bodies), nuclei, mitochondria, glycogen and vacuoles were identified (Pl. 1, fig. 1; Pl. 3, fig. 6, 7).

In media containing succinate, TBNT and buffer, TBNT showed reduction within a few minutes. When malonate was present and when enzymes were inactivated, the reduction of the TBNT was minimal.

The best fixation procedures used were glutaraldehyde+osmium and KMnO₄ fixations. Fixation with glutaraldehyde before incubation almost completely destroyed the enzymic activity (Seligman et al. 1967) and there was no increase in morphological details.

Deposits of TBNT-formazan (TNF) were found in the plasma membrane (Pl. 1, fig. 2; Pl. 2, fig. 5; Pl. 3, fig. 6) and mesosomes (Pl. 1, fig. 2, 3; Pl. 2, fig. 4; Pl. 3, fig. 8). In the plasma membrane they appeared scattered as electron-dense lines. The positive reaction of the plasma membrane was present near the mesosomes and in the membrane that embraced it (Pl. 1, fig. 2; Pl. 2, fig. 5). In the mesosomes, TNF appeared
as a homogeneous confluent electron-dense deposit (Pl. 2, fig. 4; Pl. 4, fig. 11) or as a concentric ring-like pattern (Pl. 2, fig. 4; Pl. 3, fig. 8). In some mesosomes one or other of the two patterns predominated. When the concentric ring-like pattern appeared it seemed that the reaction was not completely positive in the whole structure (Pl. 1, fig. 2; Pl. 2, fig. 4). Some mesosome structures showed heterogeneity in their reaction, as demonstrated by a clear positive zone near to a completely negative area (Pl. 4, fig. 11). In rare instances a deposit of needle-like microcrystals lying outside the cell wall was seen. Their significance cannot yet be explained. The reaction of the mitochondria in the material studied was localized in the intracristae space and in the outer membrane (Pl. 4, fig. 12). The small number of cristae in the mitochondria is noteworthy.

With malonate and the incubation mixture without substrate, or with inactivation of the enzyme with heat or fixation, some electron-density persisted in the mesosome (Pl. 4, fig. 9, 10).

**DISCUSSION**

The results show that the succinic dehydrogenase (SDH) activity of *Paracoccidioides brasilienis* was mainly found in the mesosomes and less in the plasma membrane and mitochondria. The fine structure and the localization of the positively reacting zones near the septum and the cell wall identify these as mesosomes (Carbonell & Rodriguez, 1968). The positive reaction observed in the mesosome of a hypha of *P. brasilienis* was due to SDH activity, since it appeared only when the incubation mixture had succinate as substrate and was not observed when the enzyme was inactivated or inhibited, or when the incubation mixture was used without substrate, as shown in Pl. 4, fig. 9, 10. Incubation with succinate followed by shorter permanganate fixation than the one commonly used (Kellenberg et al. 1958; Mollenhauer, 1959) proved to be the best procedure to demonstrate SDH activity. It was better than glutaraldehyde+osmium fixation because the electron-dense zones were more clearly delimited and the electron density was higher.

We presume that the high electron density is due to three factors: the electron density given by the TNF (Seligman et al. 1967), by the potassium permanganate when used alone (Bradbury & Mek, 1960; Hopwood, 1969) and by the oxidation of formazan by potassium permanganate. The most important factor is the enhancement of the reaction when permanganate is used. Due to the fact that mesosomes are made by complex infoldings of the plasma membrane, and the SDH is associated with the infoldings, the electron densities vary with the concentration of SDH in a given area.

The heterogeneity of mitochondria with respect to SDH activity has been shown by Seligman et al. (1967) and Ogawa & Barnett (1964). The first authors saw an all-or-none reactivity of the mitochondrion, while Ogawa & Barnett (1964) observed heterogeneity of the reaction inside the same mitochondrion. Dense zones and negative areas were seen in our material, indicating heterogeneity of reaction.

It is interesting to note that the hypha of *Paracoccidioides brasilienis* has mitochondria and mesosomes (Carbonell & Rodriguez, 1968) and that SDH activity is found in both membranous systems, but much more in the mesosomes than in the mitochondria. Carbonell & Rodriguez (1968) demonstrated a close ultrastructural relationship between the mesosome and the formation of the septum which is part of the cell
Since SDH activity was found in mesosomes, it is possible that respiratory activity present in the mesosome has something to do with the building of the cell wall.

REFERENCES


Plate 2

S. LEIVA AND I. M. CARBONELL
Succinic dehydrogenase in *P. brasiliensis*

EXPLANATION OF PLATES

In the following figures, *G* signifies glycogen; *N*, nucleus; *M*, mitochondria; *V*, vacuole; *PM*, plasma membrane; *CW*, cell wall; *Me*, mesosomes; *S*, septum; *WB*, Woronin bodies.

PLATE 1

Fig. 1. Mycelial phase of *Paracoccidioides brasiliensis*. Observe the nucleus, mitochondria, vacuoles plasma membrane, cell wall and mesosomes which are formed by a whorl of membranes and tubulo-vesicular structures. Glutaraldehyde + osmium tetroxide fixation with lead nitrate + uranyl acetate stain. × 24,000.

Fig. 2. Observe deposits of TNF indicating succinic dehydrogenase activity in the mesosome and a portion of plasma membrane. Incubation mixture, KMnO₄ fixation without stain. × 76,000.

Fig. 3. Mesosomes at the tip of a hypha. Observe the heterogeneous reactivity of the mesosome, the electron-dense zones in contrast with others of less electron density. Note the low electron density of the cell wall. Incubation mixture, KMnO₄ fixation without stain. × 68,000.

PLATE 2

Fig. 4. A large mesosome with many positively and negatively reacting areas. Note (arrows) that in some of the concentric ring-like structures the membrane has different electron densities. Incubation mixture, glutaraldehyde + osmium fixation, without stain. × 40,000.

Fig. 5. This shows a well-defined and positive reaction at the mesosome, and at the plasma membrane which delimited it. The cell wall is completely negative. Incubation mixture, KMnO₄ fixation without stain. × 100,000.

PLATE 3

Fig. 6. Septal mesosomes. Portions of the plasma membrane show an electron density similar to that observed in the mesosomes. Nucleus, mitochondria and three Woronin bodies are also seen. Incubation mixture, KMnO₄ fixation without stain. × 60,000.

Fig. 7. Septal mesosome in which a clearly delimited membrane is seen. Glutaraldehyde + osmium fixation with stain. × 45,000.

Fig. 8. In this mesosome note high electron density of each whorl. Incubation mixture, KMnO₄ fixation without stain. × 144,000.

PLATE 4

Fig. 9. Observe the very low electron density in the mesosome. Incubation mixture without substrate; KMnO₄ fixation without stain. × 90,000.

Fig. 10. Observe a very low electron density in the mesosome. Incubation mixture without substrate. Glutaraldehyde + osmium fixation without stain. × 60,000.

Fig. 11. Heterogeneity of the reaction in the mesosome structure. Compare well delimited structure with negative reaction (*Me₂*) with other (*Me₂*) which shows a positive reaction. Incubation mixture, KMnO₄ fixation without stain. × 70,000.

Fig. 12. A mitochondrion with positive reaction located in the outer membrane and in the intracristae space. Incubation mixture, KMnO₄ fixation without stain. × 80,000.