Oxygen Uptake and Mitochondrial Enzyme Activities in the Mitotic Cycle of *Physarum polycephalum*

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The rate of oxygen uptake by single macroplasmodia of *Physarum polycephalum* increased in two steps during each synchronous mitotic cycle. Plateaux in the respiratory pattern, of 0.22 cycles duration, occurred in mid-interphase and in the period up to and including mitosis. A fall in the rate of respiration was frequently associated with mitosis itself. This pattern of respiration continued for more than 10 mitotic cycles after inoculation and was unaffected by omission of an initial routine period of starvation. A similar stepped pattern of respiration was observed in synchronously germinating spherules during the period between outgrowth and the first mitosis. The specific activities of succinate dehydrogenase, fumarase and malate dehydrogenase remained relatively constant during the mitotic cycle, while fluctuations in cytochrome oxidase activity paralleled those in specific respiratory activity. Possible mechanisms for controlling the pattern of respiration are discussed with reference to published data on protoplasmic streaming and ATP concentrations during the mitotic cycle.

**INTRODUCTION**

Energy metabolism in the cell cycle is frequently monitored by measuring the development of respiratory activity in synchronous cultures. These experiments have, however, often yielded conflicting results. For example, according to different reports the rate of O₂ uptake by one species of yeast, *Saccharomyces cerevisiae*, increases continuously (Cottrell & Avers, 1970), discontinuously in steps (Scopes & Williamson, 1964; Greksak & Hanicova, 1973) or in a series of peaks (Nosoh & Takamiya, 1962; Kuenzi & Fiechter, 1969; Wiemken *et al.*, 1970; Dharmalingam & Jayaraman, 1973). This inconsistency suggests that the pattern of respiratory activity may be affected by the history of the cultures, and particularly by the different procedures used to synchronize the cells. Distortion of the cell cycle has been shown to occur even when one of the least disruptive synchronizing procedures – selecting cells of uniform age from a sucrose gradient – is employed (Mitchison, 1977). It was therefore of interest to study respiration in the mitotic cycle of *Physarum polycephalum*, an acellular slime mould in which nuclear division is naturally synchronous.

In addition to respiratory activity we have monitored the accumulation of a number of mitochondrial enzymes in *P. polycephalum*. Two models for the control of mitochondrial biogenesis in the cell cycle predict synchronous growth and division of the mitochondrial population within a single cytoplasm (Lloyd *et al.*, 1971; Barath & Küntzel, 1972). Although the available evidence suggests that replication of the organelle occurs asynchronously in *P. polycephalum* (Braun & Evans, 1969; Guttes *et al.*, 1969), it remains possible that the
synthesis of individual mitochondrial components, especially those coded for by nuclear DNA (Schatz & Mason, 1974), is associated with particular stages of the mitotic cycle.

METHODS

Culture conditions. Physarum polycephalum (strain M, b) was maintained at 24 °C as a microplasmodial culture growing in semi-defined liquid medium on a reciprocal shaker, as described by Daniel & Baldwin (1964). Synchronous macroplasmodial cultures were prepared by fusion of a microplasmodial suspension and were grown in Petri dishes at 26 °C on filter paper or membrane filter supplied with semi-defined medium (Daniel & Baldwin, 1964; Sachsenmaier, 1964).

Spherules were obtained and prepared for synchronous germination according to J. Mohberg (personal communication). A microplasmodial culture, inoculated at low density, was grown for 7 to 10 d into stationary phase, when spherules developed and formed a ‘collar’ on the glass above the medium. The medium, containing most of the slime, was decanted and the spherules were washed off with sterile distilled water. The spherules were sedimented at 500 g for 2 min and washed three more times with sterile distilled water. The final pellet was suspended in 0.2 ml water and 0.1 to 0.2 ml was inoculated as a thin disc on dry filter paper which rested on glass beads in a Petri dish. Growth medium, diluted with 2 vol. water, was pipetted underneath the filter paper and the spherules were incubated in darkness at 26 °C. Synchronous germination normally followed within 9 h.

Measurement of respiration rate. The rate of O₂ uptake was measured manometrically at 26 °C in the dark. To monitor respiratory activity in a whole surface culture for up to 12 h, a microplasmodium of diameter 1.5 cm, growing on a 2.1 cm filter paper disc, was placed in a Warburg flask containing 8 to 9 ml fresh growth medium so that the filter paper rested on the surface of the medium and was supported by the center well. Carbon dioxide was absorbed by 0.4 ml 10% (w/v) KOH placed in the side-arm with a strip of filter paper. Mechanical shaking was not used and O₂ uptake was followed at intervals. At least five manometer readings were taken during a 20 to 40 min period to determine the rate of O₂ uptake at each time. Flask and manometer taps were left open between each set of readings. The time of mitosis (telophase) was estimated by phase-contrast microscopic examination of 95% (v/v) aqueous ethanol-fixed smears (Guttes et al., 1961) from sister macroplasmodia inoculated simultaneously and growing at 26 °C in an incubator. When it was possible to estimate mitotic times directly from the culture at the end of the experiment, these were found to be unaffected by conditions in the Warburg flask.

Preparation of homogenates for enzyme assay. A section of a surface culture, with the central inoculum spot omitted, was washed with ice-cold 50 mm-Tris/HCl buffer pH 7.5, frozen on solid CO₂ in 1 to 3 ml of this buffer containing 25% (w/v) glycerol and stored at −20 °C overnight. Immediately after thawing, the plasmodium was treated ultrasonically for 3 to 5 s (MSE 150 W Ultrasonic Disintegrator, 3 mm diam, Ti probe, 6.5 μm amplitude, frequency approx. 20 kHz) in a glass tube cooled by an ice/water mixture. An additional 45 s ultrasonic treatment was used to release latent cytochrome oxidase activity. The crude homogenate was used for enzyme assay.

Enzyme assays. The assay for fumarase (EC 4.2.1.2) was based on that of Racker (1950). The reaction mixture contained, in a final volume of 3 ml: 150 μmol L-malate, 100 μmol potassium phosphate buffer pH 7.4 and 20 to 100 μl enzyme extract. The rate of fumarate production was followed at 250 nm and 30 °C and increased linearly with the amount of enzyme for rates of ΔAA₅₅₀ up to at least 5.5 x 10⁻² min⁻¹.

Succinate dehydrogenase (EC 1.3.99.1) activity was measured according to Breidenbach et al. (1967). The assay mixture contained, in a final volume of 3 ml: 120 μmol sodium succinate, 0.65 μg phenazine methosulphate, 0.18 mg 2,6-dichlorophenolindophenol, 30 μmol KCN, 340 μmol Tris/HCl buffer pH 7.5 and 20 to 100 μl enzyme extract. The enzyme activity could be increased as much as threefold by incubating the extract at succinate concentrations above 50 mM, so each sample was routinely ‘substrate activated’ (Singer et al., 1973) for 15 min immediately before adding phenazine methosulphate to start the reaction. The rate of dye reduction at 30 °C was measured at 600 nm against that in control reaction mixtures lacking succinate. The rate increased linearly with the amount of enzyme extract for rates of −ΔAA₅₅₀ up to at least 4.4 x 10⁻² min⁻¹.

Malate dehydrogenase (EC 1.1.1.37) activity was assayed by following the reduction of NAD with L-malate as substrate, according to the method of Teague & Henney (1973).

Cytochrome oxidase (EC 1.9.3.1) activity was estimated by following the reduction of cytochrome c at 550 nm and 30 °C (Smith, 1961). The assay mixture contained, in a final volume of 3 ml: 150 μmol potassium phosphate buffer pH 7.3, 120 nmol cytochrome c and 20 to 100 μl enzyme extract. Before use the cytochrome c was reduced with dithionite and the excess dithionite was oxidized by aeration. High concentrations of cytochrome c inhibit cytochrome oxidase activity and the concentration used was the highest at which Michaelis–Menten kinetics were followed. Consequently, when the rate of −ΔAA₅₅₀ was less than 1.5 x 10⁻³...
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min\(^{-1}\) (and the rate of substrate consumption therefore below 2 \(\%\) min\(^{-1}\)), the reaction approximated at first to zero order and the initial linear rate was measured.

Enzyme activities are expressed as \(\mu\)mol substrate converted min\(^{-1}\) (mg protein\(^{-1}\)).

Protein estimation. The protein in two samples of each homogenate was precipitated with an equal volume of 8 \(\%\) (w/v) trichloroacetic acid and washed twice in 4 \(\%\) (w/v) trichloroacetic acid. The pellets were solubilized in 0-4 M-NaOH at 45 °C for 60 min and assayed in duplicate by the method of Lowry using bovine serum albumin as standard.

RESULTS

Respiration during the mitotic cycle

The respiration rate of a single macroplasmodium of *Physarum polycephalum* increased discontinuously in two steps during the synchronous mitotic cycle (Fig. 1). Two curves are shown, one for a culture which had been subjected to an initial period of starvation at the time of inoculation, the other for a culture inoculated simultaneously but supplied immediately with growth medium. Depriving the culture of medium for a short period is a routine step for obtaining thin, rapidly growing plasmodia, but it is not essential for the preparation of a synchronous culture. The third and fourth mitoses in the culture fed immediately after inoculation preceded those in the second culture by approximately the length of the starvation period. No difference, however, was observed between the two plasmodia in the degree of synchrony or in the pattern of respiration. In each case interruptions to the increase in respiration rate occurred twice per mitotic cycle: once at about the time of mitosis and once in mid-interphase.

To determine whether these fluctuations in the pattern of respiration decay with time after inoculation, or if they are sustained even after long periods of synchronous growth, we cultured small macroplasmodia on filter paper for 4 d (approximately 10 to 12 synchronous mitotic cycles). The pattern of O\(_2\) uptake during the mitotic cycles of two such cultures is shown in Fig. 2. In both cultures a plateau in the rate of respiration coincided approximately with the estimated time of mitosis; in one (upper curve) a mid-interphase plateau is also clearly apparent.

The extent of the increase in respiration rate between mitoses in Fig. 2 (1-97 and 1-82 for the upper and lower curves, respectively) suggests that growth in these cultures was more balanced than in those shown in Fig. 1, where the rate in each increased only 1-75-fold between the third and fourth mitoses. This may indicate that more than 4 mitotic cycles were required before the macroplasmodial cultures recovered from the effects of the inoculation procedures. Despite this, however, the pattern of respiration was virtually unchanged over 6 to 8 mitotic cycles (cf. Figs 1 and 2). Thus it is unlikely that the same perturbations which led to temporarily unbalanced growth could account for the discontinuities in the respiratory pattern.

The results from a number of separate experiments in which respiration rate was followed in single macroplasmodia are summarized in Fig. 3. The mean times for the mid-points of the two plateaux in the respiratory pattern, expressed as a proportion of the mitotic cycle (with standard deviations), were 0-40 ± 0-095 (6) and 0-94 ± 0-044 (10); the mean values for the duration of the plateaux were 0-22 ± 0-011 (6) and 0-22 ± 0-079 (10) cycles, respectively. A decrease in the rate of O\(_2\) uptake (varying between 4 and 17 \(\%\)) was frequently detected towards the end of the second plateau, at or close to the time of mitosis (e.g. Figs 1 and 2), but no similar decrease was ever observed in mid-interphase.

Respiration during spherule germination

Spherules are the vegetative resting stage of the *P. polycephalum* plasmodium. If they are washed free of slime and allowed to germinate on filter paper in close proximity to one another, as described in Methods, they do so with a high degree of synchrony. Following germination they quickly fuse to form a single macroplasmodium which undergoes a normal
synchronous mitosis approximately 9 h later. This system not only permitted us to study respiration during the transition from the quiescent state to the rapidly growing plasmodium, but also enabled us to compare the pattern of respiration during the first mitosis with that in a normal synchronous culture.

The spherules were already respiring within 45 min of replacing stale medium with diluted growth medium, and the rate of $O_2$ uptake increased continuously throughout the 8 h germination period (Fig. 4). After outgrowth had begun, however, the increase in respiration rate changed to a discontinuous pattern remarkably similar to that seen during the normal mitotic cycle (Figs 1 and 2). This pattern is evident in both cultures monitored after germination. One of these had been transferred to full-strength growth medium immediately after outgrowth began, while the other was transferred to fresh diluted medium at this time.
Fig. 4. Oxygen uptake during spherule germination. Cultures of synchronously germinating spherules were prepared as described in Methods. Immediately after washing and inoculating the spherules on to filter paper (at the time indicated by an arrow) one culture was transferred to a Warburg flask and O₂ uptake was monitored for the period up to and including the initial stages of outgrowth (●); after outgrowth had begun, two further cultures were transferred to Warburg flasks containing either fresh one-third strength growth medium (○) or fresh undiluted growth medium (◆) and O₂ uptake was measured until after the first synchronous mitosis (M).

In the latter culture an interphase plateau lasting 120 min was observed, compared to only 60 min in the former culture where the overall rate of increase was greater and the synchrony of germination less good. The gradual decrease in the rate of O₂ uptake towards the end of the experiment in the culture growing on diluted growth medium was probably due to exhaustion of the medium. In the other culture, where the full-strength medium supported continued growth, a further plateau was recorded at about the time of the first mitosis. Thus the pattern of respiration was almost identical to that in the normal plasmodium.

Mitochondrial enzyme activities during the mitotic cycle

No marked oscillations were detected in the specific activities of succinate dehydrogenase, fumarase or malate dehydrogenase (Fig. 5). (The significance of a small decrease in the specific activities of each of these three enzymes at or around the time of each mitosis is not certain.) In contrast, two peaks in cytochrome oxidase activity were seen at 0.3 and 0.9 of the cycle (Fig. 5). The same oscillatory pattern of cytochrome oxidase activity was observed in a number of separate experiments. In one such experiment (Fig. 6) samples were assayed for both cytochrome oxidase and respiratory activity and the rate of O₂ uptake (expressed here on a total protein basis) also fluctuated in a biphasic manner, with maxima coinciding with those in cytochrome oxidase activity at 0.2 and 0.7 of the cycle. This result was obtained by taking samples at intervals from large macroplasmodia (5 to 7 cm diam.) and monitoring them briefly for respiration rate, rather than by growing small macroplasmodia (1.5 cm diam.) continuously in the Warburg flask as for Figs 1 and 2. The close agreement between the results obtained by the different experimental procedures can be seen when the data are expressed on a plasmodial basis (Fig. 6). Conversion of the data to this form was made on the assumption that the protein content of these cultures increased exponentially, although a similar result would have been obtained if, for example, the pattern of protein accumulation reported by Birch & Turnock (1977) had been used instead as the basis for our calculations.
DISCUSSION

The persistence of the stepped pattern of respiration into at least the tenth mitotic cycle after inoculation (Fig. 2), and the demonstration that the same pattern could be seen under several conditions of culture (Figs 1, 4 and 6), indicates that it is a normal feature of synchronous macroplasmodial growth in *P. polycephalum*.

Robbins & Morrill (1969) have suggested that respiratory activity in the cell cycle reflects the changing demands of macromolecular synthesis, and this hypothesis is supported by the frequent correlation, in a variety of cell types, between a fall in the rate of O₂ uptake and the cessation of RNA synthesis at the time of mitosis (reviewed by Mitchison, 1971). A biphasic pattern of RNA and protein synthesis in *P. polycephalum*, which closely resembles the pattern of specific respiration rate, has been reported on the basis of pulse-labelling studies with radioactive uridine (Mittermayer *et al.*, 1964) and lysine (Mittermayer *et al.*, 1966). A criticism of such studies, however, is that the rate of labelled precursor incorporation is influenced by factors other than the rate of macromolecular synthesis, notably by changes in the size of the endogenous precursor pool. It may therefore be significant that Fink (1975) has reported fluctuations in the UTP pool in synchronously growing *P. polycephalum* which might account for the pattern of uridine incorporation observed by Mittermayer *et al.* (1964). Evidence obtained by other techniques indicates that, apart from a brief stoppage during mitosis itself, the rate of synthesis of RNA (Nygaard *et al.*, 1960; Hall & Turnock, 1976) and of protein (Mittermayer *et al.*, 1966; Brewer, 1972; Birch & Turnock, 1977) increases continuously between one mitosis and the next. An alternative explanation may therefore have to be found for the pattern of respiration in *P. polycephalum*.

One possibility is that oscillations in a complex feedback control system are responsible for the respiratory pattern (Poole *et al.*, 1973). Low frequency oscillations in the respiration...
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Fig. 6. Cytochrome oxidase activity and O₂ uptake in the synchronous mitotic cycle. Oxygen uptake and cytochrome oxidase activity were measured at intervals in samples from a large (5 to 7 cm diam.) macroplasmodium. A 90 to 120° segment, from which the inner inoculum spot had been removed, was carefully teased off the filter paper and submerged in 2.0 ml fresh growth medium in a Warburg flask. The centre well contained 0.35 ml 10% (w/v) KOH and a filter paper strip to absorb CO₂. Oxygen uptake was followed manometrically for 20 to 25 min with shaking at 100 strokes min⁻¹. The rate of O₂ uptake by any sample did not exceed 1.8 μl min⁻¹. The samples used in manometric measurements were also used, after homogenization, for the assay of cytochrome oxidase activity; this is expressed as μmol cytochrome c oxidized min⁻¹ (mg protein)⁻¹. The rate of oxygen uptake is expressed both per mg protein (●) and, to aid comparison with Figs 1 and 2, on a plasmodial basis (---) in arbitrary units. The latter curve was obtained by recalculating the manometric data on the assumption that the protein content of the plasmodium increased twofold in an exponential manner between successive mitoses. The periods of mitosis (MII and MIII) were accurately determined by examination of smears taken from both the macroplasmodial cultures and from the samples themselves before and after the brief period of manometric measurements.

Changes in the rate of protoplasmic streaming might account for both the depletion of the ATP pool and the frequently observed decrease in the rate of respiration close to the time of mitosis (Figs 1, 2, 4 and 6). This would be the case if the shuttle streaming which is such a distinctive feature of this stage of the *P. polycephalum* life cycle has a role in facilitating gaseous exchange in the unusually large syncytium. The rate of O₂ uptake would then be limited by diffusion during the 20 min period before telophase when streaming ceases (Sachsenmaier *et al.*, 1973). From Fig. 6, where the experimental procedure allowed accurate timing of changes in respiratory activity relative to mitosis, it is evident that the respiration rate drops by up to 20% during the same 20 min period, and recovers at the start of the next mitotic cycle, just as streaming is reinitiated (Sachsenmaier *et al.*, 1973). The sudden decrease in ATP concentration at mitosis is consistent with this idea, since it indicates that controls other than those which operate to maintain energy balance are responsible for the drop in the rate of respiration.

The specific activities of fumarase, succinate dehydrogenase and malate dehydrogenase remained relatively constant during the mitotic cycle (Fig. 5), indicating that their synthesis...
roughly parallels that of total protein. In this way they resemble the majority of those enzymes monitored in *P. polycephalum* which are not associated with nucleic acid metabolism (Sachsenmaier & Ives, 1965; Hüttermann et al., 1970; Wolf et al., 1973). In the case of cytochrome oxidase, however, an oscillating pattern similar to that of respiratory activity was observed (Figs 5 and 6). This does not necessarily imply discontinuous synthesis of enzyme protein, since it has been shown that a stepped pattern of increase in cytochrome oxidase activity occurs in the cell cycle of *S. cerevisiae* despite a continuous accumulation of its structural counterpart, the cytochrome *aa₃* complex (Cottrell et al., 1975).

While periodicity in the accumulation of respiratory enzymes, including cytochrome oxidase, has been reported in a number of synchronous micro-organisms, the pattern has not previously been found to coincide with that of O₂ uptake (Cottrell & Avers, 1970; Forde & John, 1973; Poole & Lloyd, 1974). It is therefore unlikely that fluctuations in the specific activity of the terminal oxidase can account for the respiratory oscillations observed here. In agreement with this it can be calculated that the minimum *in vitro* activity of cytochrome oxidase was sufficient to support a rate of O₂ uptake of 0.64 µl min⁻¹ (mg protein)⁻¹, almost 20% faster than the maximum respiratory activity observed in the same experiment (Fig. 6). Assuming that the coincidence between the patterns of increase in cytochrome oxidase activity and respiration rate is not simply fortuitous, it is therefore likely that each is modulated in the mitotic cycle by the same, as yet unidentified, controls.

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