A Spectrophotometric Technique for Recording Uptake of an Organomercurial by Mycelial Fungi

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(Received 11 January 1978)

A spectrophotometric technique is described by which uptake of 2-chloromercuri-4-nitrophenol (CMNP), an organomercurial compound toxic to fungi, may be continuously monitored for 30 min periods. The method is demonstrated using mercury-resistant and mercury-susceptible isolates of *Penicillium expansum* and *Cladosporium cladosporioides*. With *P. expansum* no consistent differences in uptake were recorded. Mercury-resistant isolates of *C. cladosporioides* consistently absorbed less CMNP into their hyphae than did mercury-susceptible isolates.

INTRODUCTION

Development of resistance to organomercurial fungicides in vivo has been demonstrated in a number of fungal genera including *Aspergillus* (Gibson, 1953), *Chrysosporium* (Williams & Pugh, 1975), *Leptosphaeria* (Harrower, 1976), *Pyrenophora* (Noble et al., 1966), *Cladosporium*, *Syncephalastrum* (Greenaway, 1972a), *Ulocladium* (Greenaway, 1973b) and *Penicillium* (Kiessling, 1961; Greenaway, Cripps & Ward, 1974).

Resistance to these compounds may be due to exclusion of the mercurial ions from the hyphae, to active excretion of mercurial ions which have entered the hyphae, or to detoxification of the compound within the hyphae. We therefore wished to measure the uptake of mercurials by mercury-resistant and mercury-susceptible fungi to see whether differential uptake occurred and have examined a colorimetric technique by which such uptake might be monitored.

Measurement of the decrease in absorbance (A) of a compound can provide a convenient method of monitoring uptake from aqueous solutions. This method is unfortunately difficult to apply to most organomercurial compounds which are used as fungicides, e.g. phenyl-Hg-X, ethyl-Hg-X, because these compounds absorb light maximally at low wavelengths (<340 nm) and fungi often release into the solution materials which also absorb at these wavelengths. However, the organomercurial 2-chloromercuri-4-nitrophenol (CMNP), which is not a commercial fungicide, has an absorption peak at 405 nm. This compound was found to be toxic to fungi in vitro and was used for studies comparing uptake by resistant and susceptible isolates of *C. cladosporioides* and *P. expansum*.

METHODS

Isolates, media and growth. The fungal isolates used were: *Penicillium expansum* Link ex S. F. Gray emend. Thom, mercury-susceptible s139 and mercury-resistant r143; *Cladosporium cladosporioides* (Fres.) de Vries, mercury-susceptible s37, s38, s39 and mercury-resistant r251, r252, r253. The isolates were maintained on minimal-malt agar (Greenaway, 1973a) at 20 °C and spores obtained from colonies grown under these conditions were used for inoculation of liquid media (minimal-malt without agar but containing 25 p.p.m. streptomycin sulphate).
Media were autoclaved at 121 °C for 15 min. In tests for resistance, CMNP in 0.1 M-KOH was added to autoclaved minimal-malt medium cooled to 50 °C.

Mycelium for uptake experiments was grown in 500 ml of minimal-malt liquid medium in 21 conical flasks. Medium was inoculated with a spore suspension, sufficient to produce about 100 mycelial pellets per flask, and incubated in a Gallenkamp cooled orbital incubator at 20 °C and 160 rev. min⁻¹ for 6 d (C. cladosporioides) or 5 d (P. expansum). This produced mycelial pellets of about 5 mm diam. Some isolates (e.g. C. cladosporioides r251 and r253) grew more rapidly in liquid medium than others of the same species and in these cases fewer mycelial pellets were used in uptake experiments because they were bigger.

During uptake experiments, mycelial pellets were maintained in a solution containing 0.1 M-glucose (to provide a respiratory substrate for the fungus) and Na₂HPO₄/NaH₂PO₄ (40 mM, pH 6.8) which was essential to stabilize the pH, since absorbance of CMNP at 405 nm is pH dependent.

Apparatus for spectrophotometric assessment of uptake. The mycelial pellets suspended in buffered glucose solution containing CMNP were incubated in a reservoir. The uptake was continuously monitored by circulating the suspending liquid from the reservoir through a Hellma semi-micro flow-through cell (no. 134) mounted in a Pye Unicam SP600 UV spectrophotometer (set at 405 nm) and back to the reservoir through narrow Teflon tubing (Fig. 1a). Small sections of silicone rubber tubing were used for joints. The solution was circulated by a peri-pump (Orme Scientific) fitted with a 60 rev. min⁻¹ motor and a 2.5 mm silicone rubber tube giving a flow of 12 ml min⁻¹. The total volume of liquid was 50 ml of which 3.0 ml was outside the reservoir at any one time.

The reservoir was fitted with a 15 mm diam. sinter disc, porosity 0, mounted behind a glass plate (15 mm diam.) perforated with numerous 1 mm holes. The perforated plate prevented the mycelium from contacting the sinter disc, which otherwise became rapidly blocked. The outlet from the reservoir had a small trap above it (Fig. 1b) to collect air bubbles which might otherwise have accumulated in the spectrophotometer cell. Air was bubbled into the buffered glucose solution through a sinter disc at 20 ml min⁻¹ and the mycelial pellets were kept in motion by a slowly revolving Teflon-coated magnetic stirring bar. The reservoir stood on a plastic Petri-dish lid to insulate it from the slight heating effect of the stirrer.

The logarithmic signal of absorption produced by the spectrophotometer was converted by a Pye Unicam SP45 linear–log converter to a linear signal, which was recorded continuously on a Smiths Servoscribe potentiometric recorder.

Experimental procedure for assessment of uptake. Buffered glucose solution was circulated through the system to check that it was functioning efficiently. The solution was then replaced by 50 ml of buffered glucose solution containing 60 μM-CMNP and this solution was circulated for 5 min to ensure a stable recorder baseline. No adsorption of CMNP by the circulation system was recorded during a 30 min test period.

Mycelial pellets were strained from the growth medium, rinsed in buffered glucose solution for 30 min in the orbital incubator and drained in a nylon bag. A 10 ml volume (about 50 to 80 mycelial pellets) was added to the mercurial solution and stirred gently for 30 min, and the A₄₃₅ was continuously recorded. The pH of the mercurial solution did not change during this period.

At the end of the uptake period we investigated the distribution of CMNP that was removed from the circulating solution by the mycelial pellets. The mycelial pellets were removed from the solution, rinsed for 5 min with gentle stirring and aeration in 30 ml of buffered glucose solution (to remove any CMNP loosely
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Fig. 2. Removal of 2-chloromercuri-4-nitrophenol from solution by mercury-resistant and mercury-susceptible isolates. (a) Cladosporium cladosporioides s37 (susceptible), dry wt of mycelium 130 mg; R252 (resistant), dry wt of mycelium 122 mg. (b) Penicillium expansum s139 (susceptible), dry wt of mycelium 69.2 mg; R143 (resistant), dry wt of mycelium 102 mg.

associated with the hyphae) and then for 5 min in 30 ml of 5 mM-cysteine solution, freshly made and adjusted to pH 8.0 with 0.5 M-KOH (to remove CMNP more tightly bound to the hyphae).

The amounts of CMNP removed from the mycelial pellets by rinse solutions were assessed by measurement of \( A_{405} \) after adjusting the solutions to 0.1 M with respect to KOH, because the \( A_{405} \) of CMNP is at its highest in 0.1 M-KOH (McMurray & Trentham, 1969), and also because the CMNP-cysteine complex is apparently destroyed in 0.1 M-KOH, thus making the two measurements directly comparable.

Measurement of interhyphal spaces in the mycelial pellet. The volume of the spaces between the hyphae of a mycelial pellet may affect uptake and rinsing patterns. The interhyphal spaces were therefore measured using blue dextran (Greenaway & Ward, 1978).

RESULTS

Effect of 2-chloromercuri-4-nitrophenol on mycelial growth

The resistant \( P. \text{ expansum} \) isolate grew at 65% of control growth in the presence of the CMNP at 80 \( \mu \text{M} \) and at 35% of control growth at 160 \( \mu \text{M} \). The susceptible isolate did not grow at concentrations above 80 \( \mu \text{M} \). The resistant \( C. \text{ cladosporioides} \) isolate R252 grew at 45% of the control growth in the presence of CMNP at 80 \( \mu \text{M} \) and at 10% of the control growth at 160 \( \mu \text{M} \). The other resistant isolates (R251 and R253) behaved similarly. The susceptible \( C. \text{ cladosporioides} \) isolates (s37, s38 and s39) grew hardly at all at 80 \( \mu \text{M} \).

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Susceptible isolates of \( C. \text{ cladosporioides} \) took up CMNP more rapidly than did resistant isolates, particularly during the first 5 min of uptake. Figure 2(a) shows results for susceptible (s37) and resistant (R252) isolates. On the basis of a set of six replicates, the total ‘uptake’ (in 30 min) by \( C. \text{ cladosporioides} \) R252 was 1.78 ± 0.30 \( \mu \text{g} \) (mg dry wt mycelium)\(^{-1}\) and the uptake by s37 was 2.88 ± 0.29 \( \mu \text{g} \) (mg dry wt mycelium)\(^{-1}\); these values are significantly different at the 5% level. For other isolates of \( C. \text{ cladosporioides} \), the average uptake (three replicates of each) was 1.54 and 0.95 \( \mu \text{g} \) (mg dry wt mycelium)\(^{-1}\) for R251 and R253, respectively, and 2.09 and 2.41 \( \mu \text{g} \) (mg dry wt mycelium)\(^{-1}\) for s38 and s39, respectively. Thus, on average, the susceptible isolates of \( C. \text{ cladosporioides} \) took up 74% more CMNP than did resistant isolates. Uptake of CMNP by the resistant and susceptible isolates of \( P. \text{ expansum} \) is shown in Fig. 2(b). On the basis of a set of 10 replicates, the uptake of CMNP by the resistant isolate of \( P. \text{ expansum} \), R143, was 3.56 ± 0.84 \( \mu \text{g} \) (mg dry wt mycelium)\(^{-1}\) and the uptake by the susceptible isolate, s139, was 2.67 ± 0.67 \( \mu \text{g} \) (mg dry wt mycelium)\(^{-1}\); these values are not significantly different.

The total ‘uptake’ included the CMNP that had passed into the space between the hyphae.
A dilution effect which perhaps shows up as the initial rapid 'uptake' in Fig. 2), that which became associated with wall constituents and that which was taken up into the cytoplasm. By rinsing the mycelial pellets that had taken up CMNP, first with buffered glucose and then with cysteine solutions, the distribution of the CMNP within the different compartments could be assessed. The amounts of CMNP released from both the resistant and susceptible isolates of *P. expansum* by glucose and cysteine rinses were similar, and similar amounts remained after the rinses. With the *C. cladosporioides* isolates, however, a larger percentage of the CMNP associated with the mycelial pellets was removed by the glucose rinse from the resistant isolates (~251, 11%; ~252, 15%; ~253, 18%) than from the susceptible isolates (~257, 11%; ~258, 2%; ~259, 4%). Cysteine rinses removed similar percentages of the remaining CMNP from all isolates. After the rinses, the CMNP associated with resistant *C. cladosporioides* isolates was on average less than half that associated with susceptible isolates.

The blue dextran-permeable space of mycelial pellets of resistant isolates of *C. cladosporioides* was smaller (approx. 1.9 ml in 10 ml pellet volume) than that in susceptible isolates (approx. 4.1 ml in 10 ml pellet volume). The blue dextran-permeable space could not be accurately measured in *P. expansum* (see Greenaway & Ward, 1978) but was close to 5 ml in 10 ml pellet volume. There appeared to be no obvious difference between resistant and susceptible isolates in this respect.

**DISCUSSION**

The apparatus described here for circulation of CMNP solution through a spectrophotometer is similar to that described by Haverkate, Brevoord & Verloop (1972). Our reservoir is of a simpler construction than that of Haverkate and co-workers, but our filter system would need modification for uptake studies using fungal spores or bacteria. Continuous recording of uptake for 30 min was achieved with ease and provided a direct recording of the decrease in concentration of CMNP in the solution to which mycelial pellets had been added.

The method we describe has the advantage that it permits any changes in the kinetics of uptake during the experimental period to be clearly seen. From Fig. 2 we may speculate that the initial more rapid 'uptake' represents the initial equilibration between the suspending solution and the measured blue dextran-permeable space. The extent of the initial uptake appears to be related to this space.

The $A_{405}$ of a solution of CMNP containing mycelium could also be monitored by periodically removing samples of the liquid, filtering through a sinter to remove detached hyphae and reading the $A_{405}$, samples then being returned to the solution containing mycelium. Such a procedure yields similar results to those presented here, but does not give a continuous record. A previous technique using dithizone to assess uptake of phenyl-mercury by *Pyrenophora avenae* Ito & Kuribay. (Greenaway, 1972b) is much less convenient than that described here.

As there was no consistent difference in amounts of CMNP associated with mycelium of resistant and susceptible *P. expansum* isolates, either before or after rinsing, resistance is presumed to depend either on metabolic detoxification or on the binding of mercurials to inessential metabolites within the hyphae which may prevent their access to the site of action within the cell. In contrast, less CMNP was usually associated with mycelial pellets of resistant isolates of *C. cladosporioides* than with those of susceptible isolates. The difference in uptake by these isolates appears to be related to differences in the measured blue dextran-permeable volume, both being smaller in resistant than in susceptible isolates. Although our experiments do not of course relate to natural conditions, they do suggest that a difference in the growth habit may result in the uptake of less organomercurial fungicide by resistant isolates of *C. cladosporioides*.

We have examined the walls of both resistant and susceptible *C. cladosporioides* using gas-liquid chromatography and infrared techniques but have detected no differences in
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their structure. Minor changes in structure would, however, not have been detected by these techniques.

This work was supported by a grant from the Agricultural Research Council.

REFERENCES


