Cytolysis of *Bacillus subtilis* by *Fusarium oxysporum*

W. D. Grant,1* B. A. Prosser1 and St J. Wakefield2

1Cawthron Institute, Private Bag, Nelson, New Zealand
2Department of Pathology, Wellington Clinical School of Medicine, Wellington Hospital, Wellington, New Zealand

(Received 8 August 1990; revised 2 October 1990; accepted 23 October 1990)

Growth of *Fusarium oxysporum* on heat-killed *Bacillus subtilis* cells was accompanied by the loss of bacterial cytoplasmic contents, and this 'cytolysis' could be catalysed in heat-treated bacteria by the fungal culture fluids. In electron micrographs the bacterial walls appeared undamaged, and the absence of wall-lytic enzymes was confirmed by use of isolated bacterial walls as substrate. Appearance of cytolytic activity in cultures was paralleled by the production of proteolytic activity in the cultures. Proteolysis and cytolysis had similar pH optima at 8.8–9.0. Cultures grown on casein, but not glucose, produced high cytotytic activity. Rapid cytolysis occurred when heat-treated *B. subtilis* cells were incubated with trypsin, subtilisin or pronase E. Viable bacteria, however, were not attacked, either by concentrated culture fluids or by the commercial protease preparations.

**Introduction**

A wide range of soil fungi can attack heat-killed bacterial cells, utilizing these cells as a sole source of C, N and P (Fermor & Wood, 1981; Grant et al., 1986). Although certain fungi produce extracellular bacteriolytic enzymes which attack the cell walls of the bacterial substrate, many others, including some *Fusarium* species, apparently degrade the cytoplasm of the bacteria without affecting the wall structure (Grant et al., 1986). This process has been termed cytolysis. We have now demonstrated quantitatively the growth of *Fusarium oxysporum* on *Bacillus subtilis* cells, and have investigated the process of cytolysis in this system. We present evidence that extracellular fungal proteases play an important part in this process. Our results suggest that cytolysis is part of a non-specific utilization of proteinaceous organic matter by the fungus, rather than a specific degradation of bacterial cytoplasm.

**Methods**

*Organisms.* The strain of *Fusarium oxysporum* used for this study was isolated from garden soil. Its identity has been confirmed by Dr P. G. Broadhurst, Plant Diseases Division, DSIR, Auckland, New Zealand. *B. subtilis* strains 168 and FJ6 were gifts from, respectively, Professor J. Mandelstam, University of Oxford, UK, and Dr R. S. Buxton, National Institute for Medical Research, London, UK.

*Culture media.* The standard growth medium for *F. oxysporum* was adapted from that of Scheffer & Walker (1953) with 3% rather than 5% (w/v) glucose, the final composition being glucose 0.167 M; NH4NO3, 0.175 M; KH2PO4, 37 mM; MgSO4, 10 mM; FeCl3, 74 μM. The pH was 7.0. When bacteria were used as C, N and P source, the glucose, NH4NO3, and KH2PO4 were replaced with freeze-dried cells of *B. subtilis* 168 (3 mg ml−1) prepared as described below. Glucose and NH4NO3 were omitted from the medium when the fungus was grown on casein (BDH, 1%, w/v) as C and N source. *B. subtilis* was grown in either Nutrient Broth (Oxoid) or the defined glucose/glutamate sporulation medium, here termed S medium, of Donnellan et al. (1964) supplemented with L-tryptophan (0.1 mM) for strain 168 or L-methionine (1 mM) for strain FJ6. All media were sterilized at 121 °C for 15 min, with the exception of L-tryptophan, which was passed through a 0.2-μm filter.

*Preparation of B. subtilis cells as fungal growth substrate.* *B. subtilis* 168 was grown in 1 litre batches of Nutrient Broth (Oxoid) in 5 litre Erlenmeyer flasks on a rotary shaker (150 r.p.m.) at 25 °C. Growth was monitored by turbidity (OD600) readings (Pye-unican SP6-400 spectrophotometer), and cells were harvested by centrifugation (5000 g, 15 min) near the end of exponential growth. The pellets were washed once with distilled water, resuspended in distilled water and heated at 100 °C for 15 min. The heat-killed cells were washed again in distilled water then freeze-dried from suspension in distilled water.

*Growth of F. oxysporum on B. subtilis.* A starter culture (25 ml of the *B. subtilis* medium) was inoculated from a potato-dextrose agar slant of *F. oxysporum* and incubated statically at 25 °C for 3 d. The mycelium was then broken into small pieces by shaking with sterile glass beads in a wrist-action shaker (Griffin & George). Aliquots (0.5 ml) of the resulting suspension were transferred to 33 ml of *B. subtilis* medium in 250 ml Erlenmeyer flasks which were incubated statically at 25 °C. The flask contents were swirled daily by hand. At intervals, replicate flasks were harvested, as well as an uninoculated control flask, by centrifuging (5000 g, 30 min), washing the pellet once with distilled water, re-centrifuging, then freeze-drying to constant weight. The biomass of *F. oxysporum* grown on *B. subtilis* medium was estimated indirectly by extraction and measurement of the ergosterol present in the harvested material using the adaptation of the method of Seitz et al. (1979) described previously (Grant et al., 1990). Cultures of...
F. oxysporum grown in the standard glucose medium were used to confirm that a linear relationship existed between fungal dry weight and extractable ergosterol.

Growth and treatment of B. subtilis for cytology experiments. B. subtilis 168 was grown at 25 °C on a rotary shaker (150 r.p.m.), harvested by centrifugation (5000 g, 15 min) at mid-exponential phase (OD₆₀₀ 0.4-0.5), washed once with sterile S medium at 0 °C, then treated to inactivate autolytic enzymes. Cells were suspended in S medium, heated at 85 °C for 10 min, chilled in an ice bath, centrifuged at 5000 g, and resuspended in fresh medium. The treatment caused no detectable release of 280 nm-absorbing material, although >99% loss of viability occurred. Cells, whether heat-treated or not, were always used within 2 h of harvesting.

Cytolysis assay. Cytolysis was monitored by the decrease in OD₆₀₀ caused by the loss of bacterial cytoplasmic contents. Viable or heat-treated B. subtilis cells were suspended in sterile S medium at 25 °C so that on fourfold dilution the OD₆₀₀ was 0.7–0.8. Sterile 0.1 M-glycine/NaOH buffer pH 9.0 (2 ml) was added to 1 ml of the cell suspension, followed by 1 ml of F. oxysporum culture fluid, previously sterilized by passage through a 0.2 μm filter. Blanks were identical buffered cell suspensions incubated with 1 ml of sterile salts solution of the standard F. oxysporum medium. The initial rate of change of OD₆₀₀ was taken as a measurement of cytolysis rate. Commercial enzyme preparations were also tested for cytolytic activity using this procedure. Final concentrations of these enzymes and the appropriate buffers were: trypsin (Sigma, type III), 0.5 mg ml⁻¹ in 0.05 M-sodium phosphate buffer, pH 7.8; pronase E (Sigma type XIV), 0.25 mg ml⁻¹ in 0.01 M-sodium phosphate buffer, pH 7.5; subtilisin (Sigma, type VIII), 0.25 mg ml⁻¹ in 0.01 M-sodium phosphate buffer pH 7.5; lipase (Sigma, type VI), 0.2 mg ml⁻¹ in 0.01 M-sodium phosphate buffer pH 7.7; phospholipase C (Sigma, type IX), 0.2 mg ml⁻¹ in 0.01 M-Tris/HCl buffer pH 7.3.

Protease assay. The procedure was an adaptation of the method of Rinderknecht et al. (1968) in which 10 mg Hide Powder Azure (Sigma) was suspended in 2 ml of 0.1 M-glycine/NaOH buffer pH 9.0 in a 75 × 8 mm test tube, then 1 ml of F. oxysporum culture fluid was added. The mixture was incubated for 60 min at 25 °C, agitating gently every 5 min. The tube was cooled in an ice bath and the dye released was measured at 595 nm. The blank was 1 ml F. oxysporum standard culture medium incubated with a buffered sample of Hide Powder Azure under identical conditions.

Electron microscopy. Cells were fixed in half-strength Karnovsky’s fixative, post-fixed in osmium tetroxide, dehydrated in alcohol and embedded in Epon 812. Thin sections were stained in uranyl acetate and Reynold’s lead citrate and examined in a Siemens 102 electron microscope.

Other methods. Cell walls of B. subtilis 168 were prepared and wall degradation assayed as described previously (Grant et al., 1990). Reducing and amino groups were measured by the procedures of Park & Johnson (1949) and Ghuysen et al. (1966) respectively. The buffers used in establishing pH optima were 0.1 M-Tris/HCl (pH 7.3–9.0) and 0.1 M-glycine/NaOH (pH 9.0–11.0). Where required, culture fluids were concentrated by ultrafiltration on an Amicon YM-10 filter.

Results

F. oxysporum will grow at the expense of viable B. subtilis cells, but the process is difficult to investigate because the bacteria autolyse before appreciable fungal growth occurs. Growth of the fungus on heat-killed (100 °C) F. oxysporum will grow at the expense of viable B. subtilis cells, but the process is difficult to investigate because the bacteria autolyse before appreciable fungal growth occurs. Growth of the fungus on heat-killed (100 °C) B. subtilis cells as sole C, N and P source occurred over 9 d incubation at 25 °C, as shown by the increase in ergosterol content of the biomass from <0.05 (detection limit) to 3.59 ± 0.42 μg (ml culture⁻¹) (mean of four flasks, ± SD), and decrease in the total dry weight of cell material (fungus plus bacteria) from 2.90 ± 0.14 to
1.65 ± 0.05 mg (ml culture)⁻¹ (mean of five flasks each, ± SD). As the fungus grew, the bacteria appeared to lose their cytoplasmic contents until, when viewed under phase-contrast microscopy, they resembled empty cell walls. This phenomenon, which we have termed cytoly-sis (Grant et al., 1986), also occurred when cell-free supernatants of *F. oxysporum* cultures were incubated with heat-killed (100 °C) *B. subtilis* cells, or with freshly-grown bacteria which had been more mildly heat-treated (85 °C for 10 min) to inactivate autolytic enzymes without causing major structural damage. Supernatants of the fungus grown on the standard glucose medium
showed little or no cytolytic activity. The decrease in OD₆₀₀ which consistently occurred during cytolysis was used to quantify the process (Fig. 1a).

As the cytolytic activity of the _F. oxysporum_ supernatants was not dialysable, and was destroyed by heating, it seemed likely that the process was enzymic. An alkaline protease activity was detected in the culture supernatants, and it was then discovered that incubation of heat-treated _B. subtilis_ with the proteases trypsin, subtilisin or pronase E brought about very rapid cytolysis. Typical results of several such experiments are shown in Fig. 1(b). Incubation with lipase or phospholipase C had no apparent effect on the heat-killed bacteria (not shown). The time courses of appearance of the proteolytic enzyme and of cytolytic activity in duplicate cultures of _F. oxysporum_ grown on _B. subtilis_ were remarkably similar (Fig. 2). The pH profiles of the two activities were also similar, with identical optima at pH 8.8–9.0. _F. oxysporum_ cultures grown on casein, which also produced extracellular alkaline protease activity, were found to possess cytolytic activity similar to that obtained during growth on the bacteria. In contrast, cultures grown in the standard glucose medium produced only low levels of activity (Fig. 1a), and protease activity was not detectable by the Hide Powder Azure method in supernatants of these cultures.

Electron microscopy of _B. subtilis_ cells after treatment with cytolytic supernatants of _F. oxysporum_, or with pronase E, showed severe disruption of the bacterial cytoplasm (Fig. 3), but no detectable deterioration of the cell walls. Incubation of supernatants with suspensions of isolated _B. subtilis_ cell walls buffered at pH 2.8, 4.0, 7.0 and 9.0 for up to a week at 25 °C resulted in no decrease in turbidity compared with controls, and no changes in free amino or reducing groups in the walls. Wall degradation was not observed, even with 20-fold concentrates of culture supernatants.

In spite of their effectiveness on heat-killed bacteria, culture supernatants of _F. oxysporum_ were consistently unable to bring about cytolysis of viable cells of _B. subtilis_, either in culture or in buffered suspensions. To investigate this apparent resistance of viable _B. subtilis_ to cytolysis, tenfold concentrates of _F. oxysporum_ culture supernatant were incubated with freshly grown exponential-phase _B. subtilis_ cells. In contrast to the relatively rapid attack of such concentrates on heat-killed bacteria, no cytolysis of the viable cells occurred, although the autolytic degradation of the cells over the long incubation period made interpretation difficult. To avoid this problem, pronase E, which brings about rapid cytolysis (Fig. 1b) was used, but again there was no detectable cytolysis of the viable cells. Trypsin and subtilisin, similarly, had no effect on live bacteria. In other experiments to test cytolysis of viable cells, the bacteria were pretreated with chloramphenicol to inhibit autolysis (Rogers & Forsberg, 1971) or, alternatively, the autolysin-deficient mutant F16 (Fein & Rogers, 1976) was used as substrate. These approaches allowed the extension of the incubation periods to 26–36 h, but in neither case was cytolysis observed, with either the concentrated supernatants or the commercial proteases.

**Discussion**

When fungi are grown on a substrate of heat-killed bacteria, they produce a range of extracellular hydrolytic enzymes, including several proteases (Fermor & Wood, 1981; Fermor, 1983). Although certain fungi also produce bacterial wall-degrading hydrolases under these growth conditions (Grant et al., 1986), many do not, yet are still capable of utilizing the bacteria as their nutrient source. In these cases, the bacterial cytoplasm appears to be gradually digested away, leaving behind an apparently empty cell wall, a process which we have termed cytolysis (Grant et al., 1986).

We have now investigated cytolysis quantitatively during the growth of a strain of _F. oxysporum_ on a medium containing heat-killed cells of _B. subtilis_. The process appears to involve the action of at least one extracellular protease, produced by the fungus in response to bacterial cells or protein in the medium, and results in severe disruption of the bacterial cytoplasm. Although proteases which can attack bacterial wall peptidoglycan are known (e.g. Ensign & Wolfe, 1965; Tsai et al., 1965), the evidence suggests that cytolysis does not affect wall structure, and this is supported by the finding that trypsin, subtilisin and pronase E could also catalyse cytolysis. None of these proteases attack peptidoglycan, and indeed they have frequently been used during purification of this polymer (e.g. Brown et al., 1976; Schleifer, 1985).

Our studies show that cytolysis by _F. oxysporum_ seems to be restricted to attack on heat-treated bacteria only. All attempts to bring about cytolysis of viable bacteria were unsuccessful, even with concentrated fungal culture fluids or with the commercial proteases. These results suggest that some degree of damage, probably of the cytoplasmic membrane, was a prerequisite for cytolysis to occur: even the relatively mild conditions used to inactivate the _B. subtilis_ autolytic enzymes would have caused some membrane damage (Miller & Ordal, 1972), thus making the cell contents vulnerable to attack by external hydrolases, proteases in particular.

Although bacterial cytolysis by fungi has not been described previously, proteolytic 'emptying' of bacteria has been reported several times in the literature. Norris (1957) found a 'lytic principle' in the supernatant fluid of
Bacillus cereus M8 which lysed thiomersal-treated or heat-killed cells of seven Bacillus species, leaving intact cell walls 'almost devoid of contents'. Shirai & Aida (1980) showed that trypsin rapidly degraded heated cells of Bacillus colistinus, although in this case significant lysis occurred with viable cells. Their electron micrograph of trypsin-treated B. colistinus was remarkably similar to ours of B. subtilis in Fig. 3. Finally, Bush (1985) showed that purified proteases 'lysed' cells of several methanogenic bacteria, leaving the cell walls largely intact. Cultures or freshly harvested cells were again insensitive to this lysis. Bush (1985) was also unable to identify the site of damage responsible for lysis. A corollary from these reports and the present work is that true lysis of bacteria, in the sense of destruction of the cell structure caused by enzymic hydrolysis of the wall, cannot simply be assumed when an extract or culture filtrate brings about a turbidity loss in a bacterial suspension. Proteolytic 'emptying' of the cell has the same effect (Fig. 3). Assumptions of wall hydrolysis have been made in the past (e.g. Lynn, 1989). Such assumptions need to be verified by electron microscopy and by use of isolated cell walls as enzyme substrates.

Although it is possible that F. oxysporum and other fungi could produce proteases specifically for the degradation of bacteria in their immediate environment, it seems likely that such enzymes would be capable of attacking live bacteria. As this has not been observed, a more plausible explanation of cytolysis is that it represents a non-specific proteolysis induced by the presence of proteinaceous material. As shown in Fig. 1, cytolytic activity was low in the supernatant of glucose-grown F. oxysporum, and the cytolytic properties of supernatants after growth on casein support this interpretation. The capacity of several different proteases to catalyse cytolysis is further evidence for a non-specific process. Finally, one would expect an ability to attack proteinaceous material to be widespread in soil fungi: in a recent survey of soil fungi, we detected cytolysis in 73 out of 89 isolates (B. A. Prosser & W. D. Grant, unpublished observations). Since cytolysis occurs only with damaged bacteria, the fungi which carry out this process are acting as opportunistic 'saprophytes', in contrast to those species which can produce enzymes to degrade bacterial walls. These latter fungi can be regarded, in a sense, as 'predatory', since they are capable of attacking and utilizing viable bacteria as their nutrient source.

We thank Mr Alfred Harris, formerly of Wallaceville Agricultural Research Centre, Upper Hutt, New Zealand, for his help with preliminary electron microscopy work. This project was carried out as part of a research contract with the New Zealand Government, and supported by the New Zealand Lottery Grant Board.

References


