Hyphal Death during Colony Development in *Streptomyces antibioticus*: Morphological Evidence for the Existence of a Process of Cell Deletion in a Multicellular Prokaryote

Elisa M. Miguélez, Carlos Hardisson, and Manuel B. Manzanal

Laboratorio de Microbiología, Facultad de Medicina, Universidad de Oviedo, 33006 Oviedo, Spain

Abstract. During the life cycle of the streptomycetes, large numbers of hyphae die; the surviving ones undergo cellular differentiation and appear as chains of spores in the mature colony. Here we report that the hyphae of *Streptomyces antibioticus* die through an orderly process of internal cell dismantling that permits the doomed hyphae to be eliminated with minimum disruption of the colony architecture. Morphological and biochemical approaches revealed progressive disorganization of the nucleoid substructure, followed by degradation of DNA and cytoplasmic constituents with transient maintenance of plasma membrane integrity. Then the hyphae collapsed and appeared empty of cellular contents but retained an apparently intact cell wall. In addition, hyphal death occurred at specific regions and times during colony development. Analysis of DNA degradation carried out by gel electrophoresis and studies on the presence of dying hyphae within the mycelium carried out by electron microscopy revealed two rounds of hyphal death: in the substrate mycelium during emergence of the aerial hyphae, and in the aerial mycelium during formation of the spores. This suggests that hyphal death in *S. antibioticus* is somehow included in the developmental program of the organism.

Key words: *Streptomyces antibioticus* • programmed cell death • colony development • hyphal death • multicellular development

Cell death has long been known to be a fundamental feature of animal development, but only very recently has it become a fashionable subject of general biological interest. This is because programmed cell death (PCD, an active physiological process of cell deletion) has been recognized as playing a relevant role in the turnover of self-renewing tissues, morphogenesis, embryonic development (Ellis et al., 1991; Cohen et al., 1992; Wyllie, 1992; Vaux et al., 1994; Sanders and Wride, 1995; Raff, 1996; Jacobson et al., 1997), and more recently in neurodegenerative diseases and cancer progression (Williams, 1991; Sen and D’Incalci, 1992; Meisen, 1994; Barr and Tomei, 1994; Kerr et al., 1994; Cohen et al., 1996; Kusiak et al., 1996). Most studies on cell death have been carried out in animal cells. By contrast, little or no attention has been given to the process of cell death in prokaryotes, although they have been widely used as models for the study of many other basic cellular processes.

Members of the genus *Streptomyces* (mycelial gram-positive soil bacteria) possess two characteristics which make them attractive microorganisms for the study of cell death at the prokaryote level: first, unlike *Escherichia coli* (the most studied and best known single-cell prokaryote), which continuously divides by binary fission into two functionally and structurally identical daughter cells (which, therefore, are potentially immortal), the streptomycetes grow through the formation of long, multinucleoid hyphae that, with time, undergo senescence and die; second, they execute a complex developmental cycle that represents one of the probably several evolutionary attempts at multicellularity (Champness, 1988; Shapiro, 1988; Chater, 1989a; Chater and Losick, 1997). In fact, colonies of streptomycetes are now viewed as multicellular organisms containing morphologically and biochemically differentiated populations of hyphae organized into separate somatic and germ cell lineages (Chater, 1993; Champness and Chater, 1994; Bruton et al., 1995), the development of which is governed by an intricate system of intercellular communication (Horinouchi and Beppu, 1992; Kaiser and Losick, 1993; Willey et al., 1993). The colony growth cycle of the streptomycetes is initiated when a spore germinates, giving rise to one or more long multinucleoid filaments. These filaments elongate and branch repeatedly, originat-
ing a vegetative mycelium (substrate mycelium) that develops over, and into, the culture medium. As the colony ages, specialized branches emerge from the substrate mycelium and grow away from the surface of the colony, originating the reproductive aerial mycelium. Then, the aerial hyphae sequestate into chains of uninucleoid compartments, which finally metamorphose into thick-walled spores (McVittie, 1974; Härdisson and M anzanal, 1976; H odgson, 1992; C hater, 1998). A long this cycle, large numbers of hyphae (including the original substrate hyphae and any portion of the aerial mycelium which does not differentiate into spores) degenerate and die. Since this phenomenon was reported for the first time (Wildermuth, 1970), autolysis has been the most frequently used in the literature to describe the process of hyphal death which accompanies colony development in streptomycetes (Kalakoutsikii and A gre, 1976; E nsigh, 1978; L ochi and S hariples, 1984; M éndez et al., 1985; B rana et al., 1986; C hater, 1989a,b; H odgson, 1992; K elemen et al., 1995).

We present studies demonstrating that autolysis (a form of cell death that follows degradation of the bacterial cell wall by the uncontrolled, lytic action of murein hydrolases) is not the predominant mechanism for hyphal death in Streptomyces antibioticus. Instead, the hyphae undergo progressive disorganization of internal cell constituents (including extensive genome digestion), preceding loss of plasma membrane integrity. Cell wall degradation, if it occurs, is just a very late event. In addition, analysis of DNA degradation carried out by gel electrophoresis revealed two rounds of hyphal death during colony development: during emergence of the aerial hyphae and during formation of the spores. We conclude that during the life cycle of S. antibioticus the mycelium does not undergo a random process of autolysis, but a highly regulated process of PCD.

Materials and Methods

Strains and Media

S. antibioticus ATCC 11891 was used in this work. The microorganism was grown as lawns on glucose/asparagine/yeast extract (GAE) medium (containing 1% glucose, 0.1% asparagine, 0.05% yeast extract, 0.05 K H P O , 0.05% MgSO , 0.001% FeSO , 0.7% KHPO , 100 mM MOPS buffer [pH 7.0], and 2% agar). Plates were inoculated by spreading confluent 0.2 ml of a suspension (10 spores/ml; Hardisson et al., 1978), followed by incubation at 28°C. The developmental stage of the lawns was monitored by visually observing the changes in coloration of the surface of the cultures (waxy-yellow appearance for cultures with only substrate mycelium, powder-white appearance during aerial mycelium development, and powder-gray appearance during sporulation; Wildermuth, 1970). For biochemical studies the microorganism was cultured on sterile cellophane membranes which had been overlaid previously onto the solidified culture medium. This cultivation procedure facilitates the harvesting and handling of large mycelial masses while allowing the organism to express all stages of its growth cycle (M éndez et al., 1985; Brana et al., 1986; M iguez et al., 1994).

Microscopy

At different times of incubation, samples of the cultures (exhibiting uniformity of development) were obtained and processed for microscopy as follows. Blocks of agar containing mycelium were cut out from the culture medium and dissected into small pieces (2–4 mm in width and 8–10 mm in length). The pieces were fixed overnight at room temperature in 1% wt/vol osmium tetroxide in 0.1 M veronal acetate buffer (pH 6.0), and postfixed with 0.5% wt/vol uranyl acetate in 0.1 M veronal acetate buffer (pH 6.0) for 2 h. After this, pieces were dehydrated through graded acetone solutions over a 2-h period at room temperature, embedded in Epox resin and polymerized at 60°C for 36 h. Before polymerization, pieces were properly positioned to facilitate vertical sectioning of the whole mycelium. For electron microscopic observations, ultrathin sections of silver-gray interference color (thickness, 60–90 nm) were obtained with an LKB Ultratome III equipped with a diamond knife and mounted on Formvar-coated copper grids. To improve contrast, ultrathin sections were poststained in the dark for 10 min on uranyl acetate drops (2% wt/vol aqueous uranyl acetate), followed by counterstaining with lead citrate (pH 12, 1.5 min). Ultrathin sections were examined in a Philips EM 300 electron microscope at an operating voltage of 60 kV, and photographed with a Siemens electron microscopy film (AGFA; developed for 4 min in Kodak D 19). For light microscopic observations, thin sections (~1 μm thick) were mounted on slides, stained in Toluidine blue (0.1% wt/vol Toluidine blue in 0.1% aqueous sodium borate) for 1 min, and examined in a Nikon light microscope.

For high-resolution scanning electron microscopy, agar blocks containing mycelium were fixed with osmium tetroxide (1% wt/vol in 0.1 M veronal acetate buffer, pH 6.0) for 2 h, passed through increasing concentrations of acetone, and dried to critical point with a Balzers CPD-300 apparatus. The dried samples were mounted on aluminum stubs, coated with gold by vacuum evaporation (SCD-004 sputter coat; Balzers) and examined with a Jeol J SM-6100 scanning electron microscope.

Image Analysis

High-contrast photographic negatives were digitized using a Hewlett Packard 4C slide scanner. The digitized images were imported into Scion Image (Beta 2 version for Windows 95; Scion Corp.) for digital analysis. Plots of pixel intensity from zones of interest were obtained by positioning rectangular selections (452 pixels width, 600 pixels height) over such zones, or by tracing linear selections (5 pixels width) perpendicularly to the wall, passing from resin to the middle of the cells. Plots were generated by measuring pixel intensities (intensity range of each pixel 256 gray levels [8 bits]) along the linear selections (Profile Plot function) or in the rectangular selections (Surface Plot function). By using the LUT tool, pseudocolor images were generated in which the different components of the cell displayed an arbitrarily different color. All plots were obtained from hyphae considered to be cut diametrically (i.e., showing a clear-cut cell wall profile consisting of two electron-dense zones separated by a less electron-dense zone). Final images were composed and prepared for printing by using A dobility Photoshop software (4.0 version; A dobility Systems). All the images were printed using a Epson SC-800 printer.

Analytical Procedures

At various times of incubation, the mycelium from three plates (8.5 cm diam) was gently scraped from the cellophane with a plastic spatula, pooled, and suspended in 10 mM potassium phosphate buffer (pH 7.0). The samples were adjusted at full, or half, and a 1 μl sample was spotted on a MS E Soninprep 150 sonicator. Samples of disrupted mycelial suspensions (1.5 ml, by triplicate) were taken, 1.5 ml of 0.5 N perchloric acid was added, and the samples were maintained for 30 min at 0°C in an ice bath. A filter centrifugation, the pellets were extracted three times with 0.5 N perchloric acid at 70°C. Supernatants were pooled and assayed for RNA by the orcinol method (Schneider, 1957). Pellets were dissolved in 1.0 N NaOH and assayed for protein (Lowry et al., 1951). For dry cell weight determinations, samples (1.5 ml by triplicate) of disrupted mycelial suspensions were collected in preweighted glass vials and dried at 100°C to constant weight.

DNA Extraction and Agarose Gel Electrophoresis

Standard methods were used for DNA extraction (Hopwood et al., 1985). In brief, samples of mycelium collected from cellophane membranes were suspended in a lysing solution consisting of 5 mM Tris-HCl (pH 8.0), 25% sucrose, and lysozyme (1 mg/ml). After 1 h at 37°C, the solution was successively treated with Pronase (1 mg/ml) and 1% SDS. After phenol/chloroform extraction, the DNA solution was incubated for 1 h at 37°C in the presence of RNAse (40 μg/ml) and then precipitated by adding an equal volume of 100% ethanol. The precipitated DNA was washed with 70% ethanol and suspended in TE buffer. The final DNA solution was checked spectrophotometrically for purity and concentration. Equal amounts of DNA from each sample were electrophoresed in 1% agarose gels in TBE buffer at 60–80 mA for 1–2 h. The gels were stained with ethidium bromide and photographed under UV light.
Results

Morphological Analysis of Hyphal Death

The morphological characteristics of hyphal death in *S. antibioticus* were analyzed by transmission and scanning electron microscopy. With this purpose, *S. antibioticus* was cultured at 28°C on GAE medium. Between 36 and 96 h of incubation, samples of the cultures were harvested and processed for electron microscopy as described in Materials and Methods. As observed previously (see below), during this period of incubation the cultures were particularly rich in hyphae undergoing cellular degeneration. These hyphae could be distinguished by the presence of large electron-transparent areas in the cytoplasm and by the aberrant shapes they displayed (see Figs. 1 and 3).

From inspection of numerous hyphae at different stages of cellular degeneration, a logical progression of cytological changes for hyphal death could be deduced (Fig. 1). The first recognizable cytological change characteristic of hyphal death was a disorganization of the nucleoid which loses its condensed, axially disposed form and expands into a large, open network of DNA. During this process, the fibrillar substructure of the nucleoid becomes progressively less electron-dense and more disorganized. As the degree of disorganization of the nucleoid increases, the cytoplasm becomes less electron-dense. Such a clearing of the cytoplasm probably results from degradation of ribosomes and other macromolecular constituents, as is suggested by the marked reduction in the RNA and protein contents which the colony undergoes during this period of development (see Fig. 9). Finally, what remains of the nucleoid is a large electron-translucent area that shows no signs of fibrillar substructure and fills most of the intracellular space of the hypha. Only small electron-dense aggregates of unknown chemical nature were present in that region of the cytoplasm.

While all this is occurring, the plasma membrane shows no signs of ultrastructural disorganization and remains as a continuous, well-stained structure close to the inner face of the wall (Fig. 1, d’ and e’). Only after the cytoplasm has completely cleared and the fibrillar components of the nucleoid have almost disappeared does the plasma membrane retract from the wall and dissociate into a number of vesicles (Fig. 1 f). Moreover, an undamaged, triple-layered cell wall was persistently seen at all stages of cellular degeneration. All of these ultrastructural changes were particularly evident in the plots of pixel intensity obtained by digital analysis of the images. As Fig. 2 shows, all the plots generated from hyphae undergoing cellular degeneration revealed progressive disorganization of the nucleoid and loss of cytoplasmic electron density with maintenance of plasma membrane and cell wall integrity.

In the upper zone of the substrate mycelium and throughout the aerial mycelium, dying hyphae undergo a series of drastic changes in their morphology. As Fig. 3 shows, accompanying degeneration of the nucleoid, the hyphae collapse and undergo distortion of the hyphal shape (Fig. 3, a–c). The end result is an aberrant hyphal structure, empty
of cellular contents but still retaining a continuous, apparently intact cell wall (Fig. 3d). These morphological changes were also examined by using scanning electron microscopy. As Fig. 4a shows, normal hyphae typically appear as long, cylindrical cells the surfaces of which are totally smooth. In degenerating hyphae, however, the wall intrudes and forms small depressions at many sites along the hypha (Fig. 4, b and c). Then, the wall collapses and the hyphae progressively shrink, acquiring an irregular, tubular-deflated appearance (Fig. 4, d–g). These aberrant, but otherwise characteristic, hyphal shapes probably arise as a consequence of the loss of water and cellular contents in degenerating hyphae or, more likely, of their displacement towards the growing parts of the colony (see below).
Figure 3. Transmission electron micrographs of degenerating hyphae of \textit{S. antibioticus} showing changes in hyphal morphology. (a and b) Longitudinal sections of hyphae showing zones where the wall appears collapsed (arrowheads). (c) Longitudinal section of a hypha showing a cross-sectioned, deeply collapsed zone of the cell wall (arrowheads). (d) Collapsed hypha displaying aberrant shapes. Numbers indicate the age of the cultures. Bars: (a-c) 0.5 μm; (d) 1 μm.

**Time Course of Hyphal Death during Colony Development**

A series of light and electron microscopic studies was performed to examine the time course of hyphal death during colony development in \textit{S. antibioticus}. Accordingly, the microorganism was grown in GAE medium. At different times during growth, samples of the cultures were harvested, fixed, and processed for microscopy (Materials and Methods). Large semithin sections were used to examine the general organization of the colony in the light microscope. Ultrathin sections obtained from areas of interest were used for the ultrastructural characterization of the process in the electron microscope. The results obtained are shown in Figs. 5–8.

During the first 24 h of incubation (Fig. 5, left), the substrate mycelium grew on the surface of the culture medium by forming a thin compact layer of hyphae. Within it, however, the hyphae penetrated to depths of 40 μm. Throughout these colonies, both above and within the culture medium, the hyphae were indistinguishable ultrastructurally: they appeared very electron-dense and showed no apparent symptoms of cellular degeneration (Fig. 5, c and c’). The general organization of the colonies was basically similar after 36 h of incubation (Fig. 5, right). However, the presence of hyphae surrounded by a thin sheath growing upwards from the substrate mycelium revealed that the aerial mycelium begins form at this stage of development. Ultrathin sections corresponding to these colonies revealed few hyphae with symptoms of degeneration, and these appeared at or near the boundary with the culture medium (Fig. 5 b).

After 48 h of incubation, the colonies contained a well-developed aerial mycelium (Fig. 6, left). In semithin sections, the aerial mycelium appeared as a loose network of hyphae that develop upwards into the air (Fig. 6 a). Ultrathin sections through representative zones of these colonies, illustrating the vertical distribution of hyphae undergoing cell degeneration, are shown in Fig. 6, b–d. No hyphae with symptoms of cellular degeneration were encountered in the aerial mycelium at this stage of development, where a majority of the hyphae showed a dense, heavily stained cytoplasm (Fig. 6 b). Below this zone and near the boundary with the culture medium (not only above, but also within it), the substrate mycelium appeared as an intricate network of hyphae in different stages of cellular degeneration, ranging from hyphae with electron-transparent areas in their cytoplasm to hyphae in which the cytoplasmic components had completely disappeared (Fig. 6, c and d). At the bottom of the colony (~130 μm below the surface of the culture medium), a minority of the hyphae displayed a quite different morphological form of cell death, which was identifiable in 5–10% of the hyphae present in that region (Fig. 6, right). These hyphae did not collapse nor did they undergo progressive disorganization of the nucleoid and cytoplasm with maintenance of cell wall integrity. Instead, there was an early rupture of the wall and plasma membrane followed by rapid release of cellular contents into the surrounding medium, as suggested by the almost total absence of hyphae exhibiting stages of cellular degeneration later than those shown in Fig. 6, b and c. Only very few hyphae showing the initial stages of cell wall degradation could be examined and all these hyphae showed lightly stained cytoplasm with nucleoids of various sizes and shapes irregularly distributed through it.

After 72 h of incubation, the colonies displayed abundant sporulation in the aerial mycelium (Fig. 7). Ultrathin sections of these colonies did not reveal changes in the substrate mycelium, which appeared composed almost entirely of empty dead hyphae. In the aerial mycelium, however, a fraction of the hyphal population metamorphoses into chains of spores, while the remainder (nonsporulating hyphae) degenerates and dies. Dead hyphae appeared throughout the aerial mycelium, but they were maximally abundant towards the boundary with the substrate mycelium (data not shown).

The oldest colonies examined, 5 d old, consisted entirely of dead hyphae and mature spores (Fig. 8). All that remained in the older parts of these colonies was an intricate, pseudo-skeletal structure formed by the walls of the dead hyphae. Many such hyphae displayed aberrant morphologies, but near to the boundary with the culture medium a majority of dead hyphae retained their original tubular shape. This is probably because in such a zone of the colony the mycelium was less affected by the emergence of the aerial hyphae. Since the aerial mycelium completely depends upon translocation of water and nutrients from the substrate mycelium (a part of which is provided by dy-
ing hyphae themselves; Méndez et al., 1985), its development will draw fluids from the nearest subjacent hyphae which, as a consequence of this, will undergo collapse and distortion of shape.

Hyphal Death and Associated Biochemical Changes

*S. antibioticus* was cultured on sterile cellophane membranes overlaid on solid GAE medium. At different times during growth, samples of mycelium were carefully removed from the cellophane membranes and used either to examine the state of DNA by agarose gel electrophoresis or to estimate changes in the macromolecular content of the hyphae.

Changes in DNA Integrity. As Fig. 9 a shows, DNA extracted from samples collected during the first 24 h of incubation (lanes 2 and 3) did not show signs of degradation. However, after 36 h of incubation and continuing through later time points (Fig. 9 a, lanes 4–9), DNA degradation was apparent as a smear beneath a band of high molecular weight DNA. As can be seen, DNA degradation coincided in time with the presence in the colony of hyphae undergoing nuclear degeneration (see Figs. 5–7). On the other hand, there were two time points at which DNA degradation reached a maximum (Fig. 9 b, 48 and 72 h of incubation). These corresponded to the same time points at which the colonies were found to form aerial hyphae and to differentiate into spores, respectively (see Figs. 6 and 7).

Changes in Dry Cell Weight and Macromolecular Content. As Fig. 9 c shows, over the period 12–32 h after inoculation the cultures entered a phase of rapid growth during which all the growth parameters increased steadily. This was followed by a long period of slow biomass accumulation (which began with the emergence of the aerial mycelium and extended throughout its development), during which the total contents of RNA and protein in the mycelium varied significantly: there was a marked decrease during emergence of the aerial hyphae (between 32 and 44 h of incubation), followed by a slight increase between 44 and 64 h of incubation and by a moderate but persistent decrease during the period of spore formation.

Discussion

As in most, if not all, bacterial systems exhibiting multicellularity (Mendelson et al., 1997; Shapiro, 1997; Shimkets and Dworkin, 1997), colony development in streptomycetes is maintained by a tight balance between cell proliferation and cell death processes. Much of what we currently know about hyphal death in streptomycetes comes from an early study on colony development in *Streptomyces coelicolor*, carried out by Widermuth almost 30 years ago (Widermuth, 1970). In that work, it was reported that the mycelium undergoes extensive breakdown during the life cycle of the colony and that this process is accompanied by the presence of vacuole-like spaces in the cytoplasm (later identified as deposits of some fatty storage material; Olukoshi and Packter, 1994; Plaskitt and Chater, 1995), condensation of the nucleoid, and the rupture of the plasma membrane. However, this study was limited in two
relevant aspects: these cytological alterations were not precisely placed within the time framework of the overall death process; and there was no information on early cytological changes and, therefore, no clear picture of the sequence of morphological events leading to hyphal death could be deduced from these observations. Since that time, hyphal death has been reported repeatedly to occur as a normal part of colony development in streptomycetes (Kalakoutskii and Agre, 1976; Ensign, 1978; Locci and Sharples, 1984; Chater, 1989a, 1993; Hodgson, 1992), but no attempts have been made to elucidate the mechanisms underlying this process. Moreover, despite decades of extensive morphological studies on colony development in streptomycetes, it is remarkable that we still know so little concerning the ultrastructural aspects of hyphal death.

Therefore, the main objectives of this study were to characterize the sequence of ultrastructural changes leading to hyphal death in S. antibioticus, to identify the zones of the colony where they occur, and to determine the developmental time at which they appear. Our electron microscopy study has revealed some previously unrecognized aspects of hyphal death in streptomycetes. We have discerned that nucleoid degradation is a relatively early event in the hyphal death process and that this degradation clearly precedes the rupture of the plasma membrane. Moreover, we also observed that nucleoid degradation is accompanied by progressive digestion of cytoplasmic contents and distortion of the hyphal shape, and that all this occurs with maintenance of cell wall integrity and without disturbing the general organization of the colony.

In addition, we have observed that the hyphae die in specific zones and at specific times during the colony life cycle. Studies on the presence and distribution of degenerating hyphae within the mycelium carried out by electron microscopy and analysis of DNA degradation carried out by gel electrophoresis revealed two rounds of hyphal
death during colony development in S. antibioticus. The first round coincided with development of the aerial mycelium. It caused massive death in the substrate mycelium but had no apparent effect on the emergent aerial hyphae. The second round was not triggered until sporulation had been initiated and was more selective, since it only affected the basal, nonsporulating parts of the aerial hyphae. These observations are also interesting because they suggest that hyphal death is somehow included into the developmental program of the colony.

All together, the results obtained in this study demonstrate that the hyphae of S. antibioticus do not die via autolysis (see Introduction). Instead, they undergo an orderly process of internal cell dismantling (including extensive genome digestion), followed by shrinkage and distortion of the hyphal shape that resembles PCD in animal development. However, there are some aspects of the cytology and functions of hyphal death in S. antibioticus that distinguish it from PCD in higher organisms. First, dying hyphae do not display features such as reduction in nuclear size, condensation of chromatin, and internucleosomal cleavage of DNA (which gives a characteristic ladder when analyzed by electrophoresis), as is characteristically seen in eukaryotic cells undergoing PCD via apoptosis. This is probably a consequence of the quite different ultrastructural organization of the prokaryotic nucleoid, which lacks a nuclear membrane and contains an extensively folded DNA molecule not arranged into nucleosomes (Kellenberger and Arnold-Schulz-Gahmen, 1992; Robinow and Kellenberger, 1994). Second, dead hyphae do not completely disappear, but form part of the colony structure where they still could potentially perform two, nonmutually exclusive roles: they could provide a mechanical support for aerial hyphae to develop far from the surface of the culture medium, and they could serve as a conducting system for passage of water and solutes within the colony.
colony (a process of considerable importance for the aerial hyphae which develop in the absence of a surrounding liquid medium). Moreover, as dying hyphae provide nutrient support for development of the aerial mycelium in S. antibioticus (Méndez et al., 1985), maintenance of cell wall integrity would allow the cytoplasmic contents of these hyphae to be degraded and reused for growth without disturbing the general architecture of the colony.

In conclusion, our study has provided morphological evidence for the existence of a process of PCD in a prokaryotic organism. This adds new support to the hypothesis that the basic structure of the cell death processes has been preserved and extended throughout evolution (Vaux et al., 1994; Vaux and Strasser, 1996; Hochman, 1997). Establishing the mechanisms and signals that regulate such a process will be a major challenge for the future. In this respect, it is important to note that several bacterial plasmids carrying genes capable of killing their host have been reported recently to be responsible for the death of specific subpopulations of bacterial cells (Bugge and Gerdes, 1995; Naito et al., 1995; Yarmolinsky, 1995; Chaloupka and Vinter, 1996; Franch and Gerdes, 1996; Holčík and Lyer, 1997). Interestingly, two such plasmid-encoded killer systems seem to be present in Streptomyces spp. (Holčík and Lyer, 1997, and references therein). Finally, the colony growth cycle of the streptomycetes provides a useful prokaryotic system for the study of the mechanism and role of cell death in development. Such studies may provide insights into the role of cell death in more complex eukaryotic systems and may also provide insights into the evolution of this important phenomenon.
We thank A. Quintana for his help with the scanning microscopy and A.M. Nistal for helping with digital imaging.

This work was supported by grants BIO94-1025 and PM97-0197 from the CICYT, Spain.

Received for publication 4 January 1999 and in revised form 25 March 1999.

References


