Nuclease activities and cell death processes associated with the development of surface cultures of *Streptomyces antibioticus* ETH 7451

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The presence and significance of developmentally regulated nucleases in *Streptomyces antibioticus* ETH 7451 has been studied in relation to the lytic processes occurring during differentiation. The cell-death processes have been followed in surface cultures by a propidium iodide viability assay. This has allowed the visualization of dead (membrane-damaged, red fluorescent) and live (membrane-intact, green fluorescent) mycelium during development, and has facilitated the analysis of the role of nucleases in these processes. A parallel activity-gel analysis showed the appearance of 20–22 kDa, 34 kDa and 44 kDa nucleases, the latter appearing only when aerial mycelium is formed. The appearance of these nucleases shows a remarkable correlation with the death process of the mycelium during differentiation and with chromosomal DNA degradation. The 20–22 kDa enzymes are possibly related to the lytic phenomena taking place in the vegetative substrate mycelium before the emergence of the reproductive aerial mycelium, whereas the function of the 44 kDa nuclease seems to be related to the sporulation step. The 20–22 kDa nucleases require Ca$^{2+}$ for activity and are inhibited by Zn$^{2+}$. The nucleases are loosely bound to the cell wall from where they can be liberated by simple washing. Conceivably, these enzymes work together and co-ordinate to achieve an efficient hydrolysis of DNA from dying cells. The results show that the biochemical reactions related with the lytic DNA degradation during the programmed cell death are notably conserved in *Streptomyces*. Some of the features of the process and the biochemical characteristics of the enzymes involved are analogous to those taking place during the DNA fragmentation processes in eukaryotic apoptotic cells.

**Keywords:** programmed cell death, substrate mycelium, chromosomal DNA degradation, viability staining, differentiation

INTRODUCTION

The cell cycle best characterized of all the actinomycetes is that of *Streptomyces*, a genus with outstanding biotechnological importance: about two thirds of the antibiotics and other secondary metabolites of interest in medicine and agriculture are produced by this genus (Baltz, 1998). In *Streptomyces* the exponential growth phase coincides with the vegetative phase, which is characterized by the presence of hyphae with few septa between the cells. Next an aerial mycelium phase occurs, which is made up of thicker hydrophobic hyphae perpendicular to the surface. These suffer a massive septation and segregation of the genetic material, forming compartments which will finally give rise to spores. The intracellular effectors connecting the environmental signals with the regulatory genes are not well known, although it is conceivable that changes in particular metabolites are a determinant for the onset of the differentiation (Chater, 1998). The aerial growth of *Streptomyces* follows a period of decrease in the synthesis of macromolecules (Granozzi et al., 1990) and in fact, the formation of aerial mycelium coincides with a massive death in the substrate mycelium, which contributes to the nutrient support of the aerial hyphae.

Abbreviation: PI, propidium iodide.
The hyphae of *S. antibioticus* undergo an orderly process of internal cell dismantling, including extensive genome digestion, that resembles the programmed cell death in animal development (Miguelez *et al.*, 1999). Our group has been interested in clarifying the mechanisms that intervene in the reuse of the above-mentioned DNA constituents. With this purpose in mind we previously analysed the presence of nucleolytic activities that could play a role in such processing in *S. antibioticus* ATCC 11891. Initially, we detected and purified a nutritionally regulated 29 kDa periplasmic nuclease that nick double-stranded DNA at dG/dC-rich sequences, leaving 35–250 bp end products with 3' hydroxyl and 5' phosphate termini (Cal *et al.*, 1995). We then detected two main extracellular nucleases of 18 and 34 kDa, which were, as in the former, nutritionally regulated (i.e. they did not appear in rich media, which repress differentiation). Their biochemical characteristics made them suitable for the degradation and recycling of the DNA building blocks, as for example their lack of specificity on cutting DNA sequences and the formation of 5'-phosphate mononucleotides as predominant end products (Nicieza *et al.*, 1999). These enzymes show a dependence on Ca²⁺ for their activity and co-operate efficiently with the periplasmic nuclease to completely hydrolyse the DNA. In the present work we carry out such an analysis during the differentiation of *S. antibioticus* ETH 7451, a strain which is also currently being investigated by our group due to its remarkable capability to sporulate in submerged conditions (Novella *et al.*, 1992). This will facilitate the physiological analysis of the death process. We started our investigations in this strain by following the process of the mycelium death in cultures growing on agar, in order to subsequently compare this with the cell death processes taking place in submerged cultivation. To achieve this, we applied a propidium iodide viability stain technique, which allowed us to show the damage to the cytoplasmic membrane. In this way we have shown the evolution of the cell death processes in relation to the appearance of the nucleases and chromosomal DNA degradation. The main role in the substrate DNA hydrolysis is probably performed by 20–22 kDa nucleases, which seem to be the equivalent of the previously described 18 kDa nuclease from *S. antibioticus* ATCC 119891 (Nicieza *et al.*, 1999). We also report a 44 kDa nuclease, not described previously, which appears when the aerial mycelium is formed. In conditions in which the nuclease activities are inhibited, such as in the presence of Zn²⁺, the chromosomal DNA appears less degraded. The nuclease are loosely bound to the cell wall from where they can be liberated by simple washing. The results show that the function of these enzymes is likely conserved within *Streptomyces* and lend further support to the possible role of the specific activities in development.

**METHODS**

**Organism and culture conditions.** *S. antibioticus* ETH 7451 was grown in Petri dishes with GYM medium (glucose, yeast extract, malt extract; Novella *et al.*, 1992). Plates were directly inoculated with a dense spore suspension to obtain confluent growth. Where indicated, plates were covered with sterile cellophane disks of the same diameter, before inoculation. In the inhibition experiments, an aqueous solution of 0.5 and 1 mM ZnCl₂ (1 ml) or water (control) was added directly to the plates after 3–6 h incubation.

**Viability assay.** A permeability assay similar to that described previously for submerged *Streptomyces* cultures was used (Fernández & Sánchez, 2001). This involves the staining of damaged (leaky) cells with a polar (cell-impermeant) stain (propidium iodide, PI) in order to detect the dead cell population of *S. antibioticus*. GAE plates prepared with Noble agar (Difco) and inoculated as described above were used to obtain blocks of the GAE agar with a scalpel. These were further trimmed to cubes of about 7 mm in size and introduced into a microtome (11 mm hole diameter) previously cooled to 4°C, with the growth surface side-oriented. Sections of about 0.3–0.4 mm were obtained with a razor blade, deposited in the same orientation on a clean slide and covered with a SYTO 9 plus PI (LIVE/DEAD BacLight Bacterial Viability Kit; Molecular Probes, L-13152) stain mix prepared as recommended by the manufacturer (1:1, v/v). The SYTO 9 green fluorescent nucleic acid stain labels all the cells, that is, those with intact membranes and those with damaged ones. The PI red fluorescent nucleic acid stain only enters the damaged *Streptomyces* cells (Fernandez & Sanchez, 2001), causing a reduction in the SYTO 9 stain because of its higher nucleic acid affinity. After staining for at least 10 min in the dark, the sample was observed under a Bio-Rad MRC600 laser confocal microscope, at 488 and 568 nm excitation and 530 (green) or 630 nm (red) emission. Both images were mixed with the Confocal Assistant version 4.02 program (Todd Clark Breije, 1994–1996, freeware program distributed by Bio-Rad Laboratories) in order to make up the final dual-colour image.

**Nuclease activity gel and protein analysis.** Nuclease activities were analysed in samples obtained at the three developmental stages: substrate mycelium, aerial mycelium and sporulation. Samples were obtained in three different ways: in one of them, the bacterium was grown directly on the agar surface and the agar was extruded by forcing it through the hole of a plastic syringe (Nicieza *et al.*, 1999). The resulting suspension was centrifuged at 4°C, for 30 min at 17000 g and the supernatant was used as the source of the enzyme. In other samples the mycelium was grown on the surface of cellophane disks, and was scraped out with a plain spatula, resuspended in 20 mM Tris/HCl pH 8.0, 1 mM EDTA, 7 mM 2-mercaptoethanol buffer and ruptured in an MSE Soniprep 150, in 6 cycles of 10 s, on ice. After centrifuging at 10000 r.p.m. (Eppendorf 5415C microcentrifuge) for 30 min at 4°C, the supernatant was used as the source of the activity. Finally, in other experiments the mycelium collected from the cellophane disks as mentioned above was shaken for about 10 min in a Vortex before analysing the activity. To avoid differences in the results (due to the different sizes of the collected samples), we processed...
identified agar volumes in the extruded samples. For the sonicated and shaken samples, the weight/buffer volume relationships or the amount of protein were identical. The bands corresponding to the different nuclease activities were visualized in each sample by measuring the ‘in situ’ DNA hydrolysis after separating the proteins in denaturing SDS-polyacrylamide gels containing denatured DNA (heated at 100 °C for 10 min and then chilled on ice) (Rosenthal & Lack, 1977). After the electrophoresis the proteins were renatured and the activity detected in 20 mM Tris-HCl pH 8.0, 7 mM 2-mercaptoethanol, 10 mM MgCl₂, 5 mM CaCl₂, 10% DMSO buffer, as reported elsewhere (Nicieza et al., 1999). Gels were incubated for 90 min to 3 h at 37 °C. Micrococcal nuclease and bovine pancreatic DNase I (Amersham Pharmacia) were used as controls. Proteins were analysed by SDS-PAGE in 12% polyacrylamide gels (Laemmli, 1970) after mixing the above samples with sample buffer (M. Fernandez & J. Sanchez, unpublished results) and heating at 100 °C for 5 min before loading. Proteins were stained with silver (Cal et al., 1995). Molecular masses were estimated by their mobility with reference to marker proteins (Low Range, Bio-Rad).

**DNA fragmentation assay.** DNA degradation in the mycelium of *S. antibioticus* ETH 7451 was analysed in samples collected from cellophane discs after a careful extraction and subsequent agarase gel electrophoresis. The mycelium was suspended in Tris-HCl pH 8.0, 1 mM EDTA (TE buffer), pH 8.0 plus 2 mg lysozyme ml⁻¹ and incubated at 30 °C for 1 h. Then EDTA (100 mM final concn) and proteinase K (50 µg ml⁻¹ final concn; Roche Molecular Biochemicals) were added and the sample incubated at 30 °C for 5 min. SDS (1%, final) was added and the samples incubated further for 2 h at 37 °C. An equal volume of phenol/chloroform (1:1) was added and the solution incubated for 5 min at 37 °C. The DNA was analysed by 8% agarose gel electrophoresis in 0.5 vol. 2-propanol, washed by centrifugation with chloroform (1:1) and the aqueous phase was collected. The process was repeated until a completely clean aqueous solution was obtained. DNA was precipitated with 0.1 vol. 3 M sodium acetate and 1 vol. 2-propanol, washed by centrifugation with 70% ethanol and resuspended in TE. RNase A (100 µg ml⁻¹) and RNase T₁ (1000 U ml⁻¹) (Sigma) were added and the samples incubated at 37 °C for 2 h. The phenol/chloroform extraction, precipitation and washing were repeated as above. The DNA was analysed by 0.8% agarose gel electrophoresis and stained with SYBR gold nucleic acid gel stain (Molecular Probes) which presents more than tenfold sensitivity over the previously used ethidium bromide (Nicieza et al., 1999). The gels were visualized on a 300 nm UV transilluminator and photographed with a Polaroid 667 black-and-white print film and a photographic filter (S-7569; Molecular Probes). The nature of the fluorescent band was confirmed by digestion with DNase I (0.1 µg ml⁻¹) for 2 h at 37 °C.

**RESULTS**

**Viability staining of the Streptomyces colonies**

*S. antibioticus* ETH 7451 appears transparent and slightly orange-dark on the plates of GYM medium. The phase of substrate (vegetative) mycelium is unusually short (about 9 h) and is followed by the appearance of a thick opaque white layer (formed by aerial mycelium) projecting from the substrate mycelium. After 24 h the surface is completely covered by a grey layer of a dense mass of spores. Assays of viability were carried out at the different developmental phases. The SYTO 9 plus PI stain gives an outstanding view of the vegetative hyphae embedded in the agar (Fig. 1). At the phase of vegetative (substrate) mycelium (6 h) the hyphae appear green-fluorescent, indicating that the membrane is intact (Fig. 1a). When the culture develops (about 6–9 h), red-fluorescent membrane-damaged hyphae are observed within the substrate mycelium embedded in the agar, together with a red layer of substrate mycelium in the upper zone of the agar surface (Fig. 1b). By 9 h, some zones of the plate cultures appear completely red both on the surface and within the agar (not shown). In other areas a green layer of mycelium can be observed within the internal red layer of mycelium (Fig. 1c). The thickness of the upper red area can vary from sample to sample in the same culture, sometimes appearing thinner and the green-fluorescent layer thicker (not shown). Aerial mycelium starts to develop after 12 h, on the upper layer of red hyphae (Fig. 1d), showing a deep agar zone with a membrane-compromised substrate mycelium and an upper green-fluorescent hyphae layer which should correspond to an incipient aerial mycelium. As the aerial mycelium development continues, two different regions can be observed: a green layer in the lower part of the sample (nearer to the agar surface) and a red network in the bulk of the air-projecting mycelia with live spore chains interwoven within the red layer (Fig. 1e). In more mature zones, all aerial mycelium appear red-fluorescent; only the chains of newly formed spores are viable (Fig. 1f).
Appearance and characteristics of the nucleases and chromosomal DNA fragmentation in the course of development

Nuclease activities were analysed in samples of plate cultures of *S. antibioticus* ETH 7451 at the three stages of development: substrate mycelium, aerial mycelium and sporulation. Each of these were processed in three different ways as specified in Methods. In the first, the agar from plates in which the bacterium was growing directly on the surface was extruded by forcing it through the hole of a plastic syringe; while the mycelium remained intact, this treatment disrupted the structure of the gel and the proteins could be recovered in the supernatant after centrifugation. This method permits the analysis of the nucleases excreted by the mycelium into the surrounding agar. In another series of samples, the mycelium grown on the surface of cellophane disks was scraped with a plain spatula and ruptured to analyse the intracellular nuclease activity. Finally, in a third series of experiments, the mycelium grown on the cellophane disks was collected in a buffer (see Methods) and shaken. All the samples thus obtained were analysed for nuclease activity in DNA-polyacrylamide gels. By these combined analyses a complete view of the nucleases and their distribution was obtained (Fig. 2).

Two main bands of 34 kDa and ~22 kDa were detected at 9 h of cultivation (no activity was detected at 6 h; data not shown), but their intensity was higher in the washed mycelium sample (Fig. 2a). In addition, another ~20 kDa (and a minor 18 kDa band) band appears in the washed mycelium. The 20 and 22 kDa bands could be different forms of the same nuclease, as both show a strict requirement of Ca$^{2+}$ for their activity (Fig. 2c). The 34 kDa nuclease does not have such a strict requirement as the 20–22 kDa nucleases (Fig. 2c). In the sonicated mycelium, the two 20 and 22 kDa bands are again visible (the higher intensity in this case corresponds to the 20 kDa band) in addition to one thin 54 kDa band and
the scarcely visible 18 kDa band. The chromosomal DNA appears already partially degraded at this time (Fig. 2e). When the aerial mycelium developed completely (15 h samples), the 20–22 kDa bands could not be distinguished, but the 34 kDa band plus a 44 kDa band were clearly seen (a fine 20 kDa band can be also seen in the sonicated mycelium; Fig. 2a). The 44 kDa nuclease, as with the 34 kDa enzyme, does not require Ca²⁺ for the activity (Fig. 2c). At this point, a very extensive degradation of chromosomal DNA is observed; only a fraction of the DNA appears intact in the upper part of the gel, the rest being fragments of approximately 100–250 kb (Fig. 2e). Similar results were obtained in the sporulation phase (24 h), in which aerial mycelium continued to form, but the intensity of the 44 kDa nuclease is somewhat higher. The aspect of the chromosomal DNA on the gel is similar to the 15 h sample (Fig. 2e), although the fraction which migrates more slowly on the gel is higher. The 34 kDa nuclease was present in all the developmental steps. The fact that the above nuclease can be washed away from the cells by moderate shaking of the intact mycelium and also detected in the extruded agar samples indicates that they are externally located and loosely bound to the cell wall. This is supported by the comparatively high nuclease activity present in the washed-mycelium samples (Fig. 2a).

As previously reported for S. antibioticus ATCC 11891 (Nicieza et al., 1999), the development of S. antibioticus ETH 7451 is impaired in the presence of 0.5–1 mM Zn²⁺. This means that whereas in the control cultures at 15 h the aerial mycelium is completely formed, the cultures with Zn²⁺ are retarded in the phase of substrate mycelium. After 24 h, the control cultures have sporulated; the plates with Zn²⁺ are in the aerial mycelium phase, with no sign of septation within the hyphae (not shown). When the nucleasees were analysed in the extruded agar samples from both media at 15 h, the control cultures showed the presence of the 34 kDa and 45 kDa bands (Fig. 2c); however, in the cultures with 0.5 and 1 mM Zn²⁺ the 20–22 kDa band (the 20 kDa band was not seen with 1 mM Zn²⁺) and the 34 kDa band were visible. The gel activity analysis was repeated with the extruded agar sample from the 0.5 mM Zn²⁺ plates (Fig. 2e) but in this case 0.5 mM and 1 mM Zn²⁺ were added to the incubation buffer. The results (Fig. 2d) show that the activity of the three 20–22 and 34 kDa nucleases is inhibited by Zn²⁺, although the 20–22 kDa nucleases seem more sensitive than the 34 kDa enzyme; the 18 kDa band present in the control is not affected. It can be concluded that it is the activity, and not the synthesis of the enzymes, which is inhibited by Zn²⁺. This inhibition is likely to be responsible for the fact that in the 24 h cultures the chromosomal DNA appears less degraded in the presence of 0.5 and 1 mM Zn²⁺, when the control cultures are sporulated and the DNA is substantially degraded (Fig. 2e; see above).

An SDS-PAGE analysis of the proteins of the washed substrate (vegetative) mycelium from S. antibioticus ETH 7451 showed a remarkably high proportion of proteins loosely bound to it (Fig. 2b). This suggests that the surface, and most likely the cell wall from the vegetative mycelium is a functionally very active structure which maintains a notable reservoir of proteins, some or most of them probably related with scavenger and/or degradative functions. These cell-wall located proteins are drastically reduced in number in the aerial mycelium and, in fact, only a few predominant proteins (33 kDa and 38 kDa) are observed at 15 h in the washed S. antibioticus ETH 7451 mycelium (Fig. 2b). These two proteins plus another of about 86 kDa, and minor bands of 42 kDa and 36 kDa are all that can be seen in the sporulation phase gels (Fig. 2b). This will notably facilitate the purification and biochemical characterization of the nucleases and the subsequent cloning of the genes.

**DISCUSSION**

**Analysis of the viability of substrate and aerial mycelium of S. antibioticus**

The viability test has been recently applied by our group to *Streptomyces* submerged cultures (Fernandez & Sanchez, 2001). In this study it has been used for the first time in cultures growing on the surface. The technique has shown its outstanding feasibility for evaluating the physiological state of the mycelium of *Streptomyces* growing on the agar surface. This type of analysis will constitute a valuable complement of other biochemical or morphological studies. The ultrastructural studies of the colony, such as those performed previously in *S. antibioticus* ATCC 11891 (Miguelez et al., 1999), report the internal details of the particular hyphae (as for example the progression of nucleoid degradation or the changes in the cytoplasm) whereas the viability method contributes by providing the real view of the general physiological state of the colony. For example, it has been claimed that nucleoid degradation is an early event in the hyphal death process and that this degradation precedes the rupture of the plasma membrane (Miguelez et al., 1999). However, this is only based on the ultrastructural aspect of the nucleoid within specific hyphae (i.e. a disorganization of the electron-dense nucleoid and a continuous well-stained membrane structure). Whereas the internal disorganization of some hyphae could coexist with a relative integrity of the plasma membrane, what seems clear is that at the stage at which the nucleolytic processes conceivably take place (shortly before and during aerial formation and sporulation steps) both the presence of the nucleases and the presence of massively membrane-damaged hyphae coexist (Fig. 1b–f; Fig. 2a). As mentioned in the Results, no nuclease activity was detected at 6 h cultivation. During this time the membrane, as shown by the viability staining (Fig. 1a) remains intact. Apart from this, the above-mentioned ultrastructural analysis of the cell death process carried out in surface cultures (Miguelez et al., 1999) shows remarkable analogies with our viability results in *S. antibioticus* ETH 7451.
described by the above authors in S. antibioticus ATCC 11891, the first round coinciding with the formation of aerial mycelium (which should correspond to the death phases shown in Fig. 1b–d). This massive death of the substrate mycelium does not hinder the emergence of the aerial hyphae, as occurs in our study (see Fig. 1d). The second round takes place when sporulation starts and affects the nonsporulating parts of the aerial hyphae (Fig. 1e, f). At the end, the S. antibioticus substrate mycelium appears empty and all nonsporulating aerial hyphae degenerate and die (Miguelez et al., 1999), this phenomenon being maximally abundant towards the boundary with the substrate mycelium (as also shown in the stained sample in Fig. 1f). The utility of the analysis by fluorescence confocal microscopy described in our work extends far from the phenomenon analysed here, as it opens the possibility to investigate the influence of the medium and nutritional conditions on the viability state of this biotechnologically relevant group of bacteria.

Nuclease production, chromosomal DNA degradation and differentiation are co-ordinated events

The impairment of the aerial mycelium formation by Zn$^{2+}$ could be due to additional effects of this cation on other enzymes, for example the serine proteases. A trypsin-like enzyme has been described in S. antibioticus ATCC 11891 (Nicieza et al., 1999) and a similar protease has been also detected in S. antibioticus ETH 7451 (M. Fernandez, unpublished results). Its role has been related to proteolytic processing of a hypothetical inactive precursor of the 18 kDa and/or the 34 kDa nucleases (Nicieza et al., 1999; J. Huergo & J. Sanchez, unpublished results). The 20–22 kDa nucleases seem to be equivalent to the 18 kDa nuclease detected previously in the surface-sporulating strain S. antibioticus ATCC 11891. In our previous work (Nicieza et al., 1999) we reported that the 18 kDa nuclease peaks in the aerial mycelium phase (about 36 h). This apparent difference with respect to the results obtained in the ETH 7451 strain can be readily explained by the difference in the growth kinetics between the two strains. The development of S. antibioticus ATCC 11891 is slower than the ETH 7451 strain and thus, the phase of substrate (vegetative) mycelium in the ATCC 11891 strain lasts about 24 h. It is conceivable that both vegetative and aerial mycelium coexist at 48 h, whereas in the ETH 7451 strain, which develops significantly more rapidly, only vegetative or aerial mycelium would predominate at each phase (as suggested by the photographs in Fig. 1e, f). This could explain the higher intensity of the 18 kDa nuclease band in the activity gels from the aerial mycelium phase of S. antibioticus ATCC 11891. The similarity between the 18 kDa and the 20–22 kDa nucleases is further supported by the fact that the ETH 7451 enzyme shows, as occurred with the ATCC 11891 18 kDa nuclease (Nicieza et al., 1999), a strict requirement for Ca$^{2+}$. The activity of the 44 kDa and 34 kDa nucleases, by contrast, is not noticeably impaired in the absence of that cation [this is also true for the S. antibioticus ATCC 11891 34 kDa nuclease, which was previously claimed to need Ca$^{2+}$ (Nicieza et al., 1999; J. Huergo & J. Sánchez, unpublished results)]. Moreover, the 20–22 kDa nucleases are synthesized only in the vegetative (substrate) mycelium (plates incubated for 9 h) and could play, as postulated in the above strain for the 18 kDa nuclease (Nicieza et al., 1999), a preferential role in the DNA lytic processes which conceivably take place in the substrate mycelium just before and during the course of emergence of the aerial mycelium. This hydrolysis is likely facilitated by the disorganization of the cytoplasmic membrane, as mentioned above, and shown by the viability stain (Fig. 1c, d). The role of these nucleases and possibly the 34 kDa enzyme, in the substrate mycelium DNA degradation, is further supported by the delaying of the chromosomal DNA hydrolysis in the presence of Zn$^{2+}$ (Fig. 2e). The 44 kDa nuclease appears specifically at the aerial mycelium phase; when the appearance of the aerial mycelium is impaired by Zn$^{2+}$, the nuclease is not visible (but the 20–22 kDa enzymes are present; Fig. 2c), thus supporting the relationships between the synthesis of the enzyme at the differentiation phase and its potential role in the lytic processes which take place in the aerial mycelium before spore formation. A nuclease of similar molecular mass has been recently detected in the previously studied S. antibioticus ATCC 11891 strain (M. Fernández & J. Sánchez, unpublished results). This enzyme, as shown here, appears specifically when the aerial mycelium is forming (M. Fernández & J. Sánchez, unpublished results). The 34 kDa nuclease could be related to both lytic processes, probably contributing to the concerted activity of the mentioned nucleases to a rapid and extremely efficient degradation of the DNA, as has been shown previously with the 18 kDa and 34 kDa enzymes from S. antibioticus ATCC 11891 (Nicieza et al., 1999).

Chromosomal DNA appears partially degraded when aerial mycelium starts to form in S. antibioticus ATCC 11891 growing on plates (Nicieza et al., 1999). In the sporulation phase the DNA degradation is much more intense (Nicieza et al., 1999). These kinetics are also observed in S. antibioticus ETH 7451 (Fig. 2e). When the aerial mycelium is completely formed (15 h) the DNA is extensively degraded, with the exception of a small slowly migrating fraction visible as a defined band on the agarose gel. This likely represents the DNA of the few spores present at this time, which are partially sensitive to lysozyme (Novella et al., 1992). This band, although more intense, is also seen in the 24 h sporulated cultures. The remaining DNA visible on the gel is completely degraded to 100–250 kb fragments; the smaller oligonucleotides and mononucleotides conceivably formed in the hydrolysis (Nicieza et al., 1999) will not be recovered by the 2-propanol/sodium acetate step and thus are not visible on the gel. As already pointed out, S. antibioticus ETH 7451 has been previously used in our group to analyse differentiation, as it has an exceptional capability to sporulate synchronously after a nutritional down-shift under submerged conditions (Novella et al., 1992). The bacteria harbour a NaeI-
Nucleases and cell-death processes in *Streptomyces*

**Streptomyces as a developmental model**

These and our previous data (Nicieza *et al.*, 1999) point to the existence of a series of co-ordinated and notably conserved biochemical events, related to chromosomal DNA degradation during the lytic processes that accompany development and differentiation in *Streptomyces*. In the hypothetical sequence for cell death induction, a change in the nutritional environment, growth rate or both, will transmit a still uncertain signal to the *Streptomyces* cells which in turn could activate a cytoplasmic effector (perhaps with the co-operation of 


The formation of the nucleases which intervene in the DNA degradation process is conceivably under a strict control, part of which could be the induction of the above-mentioned proteolytic processing of an inactive high molecular mass precursor which would give rise to the active 18 and 34 kDa nuclease forms (Nicieza *et al.*, 1999; J. Huergo & J. Sánchez, unpublished results). A second round of DNA hydrolysis could take place in the aerial mycelium, when spores are forming. In this, the 44 kDa enzyme plus the 34 kDa nuclease could play the role. We have investigated the existence of nucleases with the characteristics and location of those described here, in several other species of *Streptomyces*, such as *Streptomyces coelicolor*, *Streptomyces albus*, *Streptomyces acromogenes* or *Streptomyces lividans*; our initial results show the presence, in all of them, of 34 kDa and 18–23 kDa cell-wall located nucleases (no activity could be detected within the cytoplasm) with similar biochemical characteristics to those described here and in *S. antibioticus* ATCC 11891 (J. Huergo & J. Sánchez, unpublished results).

It has been suggested that micro-organisms which present a complex developmental life (such as *Streptomyces*, *Bacillus*, *Anabaena*, *Caulobacter*, *Rhizobium* or the myxobacteria) would have programmed cell death mechanisms which could be considered as the phylogenetic precursors of the eukaryotic programmed cell death (Yarmolinsky, 1995; Hochman, 1997). This has been shown for *S. antibioticus*, in which an orderly process of internal cell dismantling, rather than an uncontrolled autolytic process, takes place (Miguez *et al.*, 1999). However, as shown in this study and in our work, a remarkable difference between this mechanism and the programmed cell death of higher organisms is that dead hyphae from *Streptomyces* do not completely disappear, but remain to form part of the colony structure, thus allowing the passage of the recycled nutrients. On the other hand, several eukaryotic protein kinases and apoptotic proteins have homologues in *Streptomyces* (Zhang, 1996; Aravind *et al.*, 1999) and some of the apoptotic nucleases, as also occurs with the 18 kDa and the 20–23 kDa *Streptomyces* enzymes, have Mg\(^{2+}\) and Ca\(^{2+}\) requirements and are inhibited by high NaCl or KCl concentrations, Zn\(^{2+}\) or aurin tricarboxylic acid (Peitsch *et al.*, 1994; Montague & Cidlowski, 1996; Hale *et al.*, 1996; Hughes & Cidlowski, 1999; Widlak & Garrard, 2001). It seems that the study of the biochemical and genetic basis of the programmed cell death in this bacterium could contribute to a better understanding of the role and evolution of this important process in eukaryotic cells.

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