Growth of *Arthrobacter globiformis* in soil observed by fluorescent antibody and ELISA techniques

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The growth and survival of *Arthrobacter globiformis* (NCIMB 10683) in a wheatfield and a sand-dune soil were investigated by introducing rod-shaped cells on the surface of microscope slides into the soil. Slides were recovered and stained with fluorescent antibodies. In the wheatfield soil, new microcolonies of considerable size were observed on or around both soil particles and fungal hyphae throughout the incubation period of 48 d. In both sterile and non-sterile wheatfield soil, the majority of the rods reverted to cocci after 2 weeks incubation. In the sand-dune soil, cells did not grow and only a few were found after 48 d. Growth of *A. globiformis* in the wheatfield soil was also investigated using a direct soil inoculation technique. Soil samples were recovered after different incubation periods and cells extracted using a centrifugation method. The number of cells in extracts was estimated using ELISA and direct count techniques. In the latter technique, extracted cells were double-stained with fluorescent antibodies and ethidium bromide. Growth took place in both sterile and non-sterile wheatfield soil. The introduced rods reverted to cocci after a few days, as on the slides. However, cells introduced directly into soil grew faster than those introduced on slides.

**Keywords:** soil microbiology, *Arthrobacter globiformis*, cell enumeration, fluorescent antibodies

INTRODUCTION

Many workers have studied growth and activity of soil bacteria, including *Arthrobacter* spp., by introducing laboratory-grown bacterial cultures into sterile soil and determining viable cell numbers by conventional plating techniques after incubation of the soil (Robinson *et al.*, 1965; Salonius *et al.*, 1970; Lowe & Gray, 1973a, b). The information obtained using this approach is misleading because of the differences between sterilized and natural soils as media for microbial growth. Sterilization, in addition to eliminating the viable population, alters the physical, chemical and nutritional properties of the soil. Furthermore, the types and amounts of nutrients available for bacterial growth in soil depend partly on the presence of other micro-organisms, their products and dead remains.

Thompson *et al.* (1990, 1992), used plate culture procedures to investigate the survival of two ecologically distinct bacteria (*Flavobacterium* and *Arthrobacter*) introduced into sterile and non-sterilized unplanted and rhizosphere soils both in laboratory soil microcosms and in the field. However, they found difficulties in discriminating the organism under study from the rest of the soil microflora. It was also impossible, by conventional plate count techniques, to appreciate the spatial inter-relationships between micro-organisms within the soil, or to examine the morphology (rods, cocci or mycelia) of the arthrobacters in soil.

The organism under study can be distinguished from the rest of the soil microflora by using fluorescent antibodies. Siala & Gray (1974) investigated both the occurrence of native cells and the growth and germination of vegetative and spore forms of *Bacillus subtilis* in sterile and non-sterile soil from the A, and C horizons of a sand dune soil in this way.

The use of immunological techniques for quantitative autecological studies of micro-organisms in soil has received little attention. Development and evaluation of a procedure for the enumeration of a specific population in natural habitats using an immunofluorescence cell

**Abbreviations:** FITC, fluorescein isothiocyanate; RITC, rhodamine isothiocyanate.
counting technique were reported by Bohlool & Schmidt (1973) and Schmidt (1974). More recently, enzyme-linked immunosorbent assay (ELISA) has been developed for quantitative studies (Engvall & Perlmann, 1971). The ELISA technique has been used successfully for the identification of Rhizobium strains (Kishinesey & Gurfel, 1980; Olsen et al., 1983; Fuhrmann & Wollum, 1985) and reports indicated that the rhizobial numbers are estimated reliably by this assay in soil (Martensson & Gustafsson, 1985; Nambiar & Anjaiath, 1985). However, Lochner et al. (1988) reported some limitations of this assay for routine determination of legume inoculant quality.

The preparation of relatively specific antisera (Mansoor, 1992) to a number of Arthrobacter reference strains and the staining of both rod and coccoid forms of these strains with the same fluorescence intensity made it possible to distinguish the strain introduced into soil, either on the surface of sterile slides or added directly to the soil, from the rest of soil microflora. Thus, it became possible to study the location, growth and morphology of Arthrobacter globiformis (NCIMB 10683) in the presence of a complete soil microflora, using both immunofluorescence and ELISA techniques. This is important, as A. globiformis is one of the commonest isolates from soil and its function is poorly understood. It was also possible to test predictions on the behaviour of this organism made from studies on its physiology in continuous culture.

METHODS

Soils. Soil samples were collected from the top 6 cm of a wheatfield (Brick House Farm, Peldon, Essex, UK; OS Map 168, TL 997154) and from an embryo sand dune (near Chapel of St Peter ad Vincula, Bradwell, Essex; OS Map 168, TM 031089). The wheatfield soil was a silty clay loam (Windsor series) with a pH of 6.5 which had been planted only with wheat for 30 successive years and had been regularly supplied with NPK fertilizers. The sand-dune soil (pH 8.3) came from an area of wind-blown sand and crushed shells just above the spring high-tide level and was colonized by Agropyron pungens and Carex arenaria.

The samples were sieved to remove large roots and twigs. The moisture content of each soil was determined and approximately 40 g of each soil were placed in each of several Petri dishes. For the preparation of sterile soil, deionized water was added to double the moisture content. The soil was autoclaved for 30 min at 121°C on two separate occasions, with 24 h incubation at room temperature in between. After the second autoclaving, plates of soil were cooled to room temperature, by which time the water content of each soil had fallen to about 10% level. The soil samples were maintained at this level throughout the experiment by addition of sterile water following further decreases in weight.

Preparation of the inoculum. Arthrobacter globiformis NCIMB 10683 was grown on nutrient agar (Oxoid) in medical flat bottles at 25°C for 18 h. The cells were harvested by centrifugation, washed four times in sterile deionized water and resuspended in 25 ml deionized water in 100 ml Erlenmeyer flasks. After dispersion of aggregates by stirring, the suspension was diluted with sterile deionized water to give an appropriate concentration of cells. The suspension was checked microscopically to confirm that cells were rod-shaped.

Contact slides. Clean, grease-free microscope slides were sterilized and inoculated by spreading 0.01 ml of a suspension of rod-shaped cells over one side of the slide with a flame-sterilized needle. The slides were allowed to dry in sterile Petri dishes and then placed on the surface of soil samples with the inoculum in contact with the soil. This resulted in an inoculum of about 10^7-10^8 cells mm^-2. Inoculated slides were incubated in sterile and non-sterile wheatfield soil, and in sterile and non-sterile sand-dune soil at 15°C. Controls of non-inoculated slides and slides inoculated with heat fixed cells were also used.

Recovery, staining and observation of slides. Triplicate slides were recovered from each treatment after varying periods of incubation. The slides were heat fixed, rinsed in PBS for 5 min and air dried. Gelatin-FITC conjugate (Bohlool & Schmidt, 1968) was allowed to flood the slide surface and dried as a film at 60°C. The A. globiformis-specific antiserum (0.1 ml) at a dilution of 1:20 (antiserum:PBS) was added to the slide. The slides were incubated in a humid chamber for 1 h at 25°C and washed for 15 min before air drying. FITC-conjugated anti-rabbit serum was used at a dilution of 1:20 to stain the preparations, which were incubated and washed as before. The stained bacteria were covered with a mixture of a mountant (Mowiol) and an antifade agent (p-phenylenediamine) (1:9) and a cover glass. Preparations were observed with a ×100 oil-immersion lens, using non-fluorescent immersion oil, and viewed with a Leitz Dialux 20 microscope, fitted with a Ploemopak 2.4 (Leitz Instruments) under short-wavelength blue light. Fifty fields of view were observed on each slide, the fluorescent cells counted and the dimensions of 10 randomly selected cells in each field of view measured. The mean number of fluorescent cells and their biovolume per field of view were determined. Photographs were taken with a WILD MPS 55 photo automatic system (Leitz Instruments) on Kodak Ektachrome film (ASA 400).

Soil inoculation. Forty gram quantities of sterile and non-sterile wheatfield soil in Petri dishes were inoculated with appropriate concentrations of cell suspension prepared as before but in a small volume of water and mixed thoroughly with a sterile sparula to disperse the cells throughout the soil. The soils were incubated at 15°C.

Cell extraction. Triplicate Petri dishes were recovered from each treatment after different periods of incubation. A soil extraction procedure for bacteria based on that of Holben et al. (1988) and modified at the University of Aberdeen, UK (J. I. Prosser, personal communication) was used. The contents of each Petri dish (40 g soil) were mixed with 150 ml 0.1% (w/v) sodium deoxycholate (pH 7.5) and 10 g acid-washed polyvinylpyrrolidone (Evans et al., 1972) and homogenized in a 500 ml capacity Atomix blender for 3 min intervals (with 1 min cooling on ice between each blending cycle). The homogenate was placed on ice to settle and transferred to a 250 ml centrifuge bottle. The soil debris was pelleted by centrifugation at 1500 g for 10 min at 4°C. The supernatant (SN1) was transferred to a fresh 250 ml centrifuge bottle and kept on ice for further treatment. The soil pellet was subjected to two more rounds of homogenization and centrifugation. The three supernatants obtained (SN1, SN2 and SN3) were combined and centrifuged at 10000 g for 30 min at 4°C. The bacterial pellet was washed twice in 200 ml 0.1% (w/v) sodium pyrophosphate and twice in 2% (w/v) sodium hexametaphosphate (pH 8.5) by centrifugation at 10000 g for 30 min at 4°C. The bacterial pellet was suspended in 100 ml TE buffer (0.33 M Tris/HCl, pH 8.0 + 1 mM EDTA) and centrifuged as above. The supernatant was decanted and the final bacterial pellet weighed and suspended in 10 ml Tris/sucrose/EDTA buffer (pH 8.0) in a 25 ml capacity Erlenmeyer flask.

Enumeration of cells in extracts from soil. The extracted cell suspension was agitated to disperse the bacterial cells and serial
dilutions (10⁻¹–10⁻⁷) were prepared. Portions (0.1 ml) of appropriate dilutions were spread over the surface of sterile peptone yeast extract agar (Goodfellow et al., 1968) in Petri dishes. Three plates were prepared at each dilution and all were incubated at 25 °C for 10 d. The mean numbers of bacteria developing were calculated and the number of viable bacteria per g oven-dry soil determined.

Appropriate dilutions of extracted cell suspensions (0.01 ml) were spread over exactly 1 cm² area of new, unused microscope slides. Three slides were prepared at each dilution for each treatment. The preparations were dried at 45 °C and stained using fluorescent antisera as above. The stained preparations were dried at room temperature and counter-stained with 0.1 ml ethidium bromide solution (0.2 mg ml⁻¹) for 3 min. The slides were rinsed in deionized distilled water and dried. The preparations were mounted under cover slips and viewed with a Leitz Dialux 20 microscope, with either short-wavelength blue or green light. The number of green-fluorescing and red-fluorescing cells occurring in 50 fields of view inside an oblong (area: 0.0004 cm²) marked on the eyepiece graticule on each slide were counted and the dimensions of 10 randomly selected fluorescent cells in each field of view measured. The number and biovolume of the cells per g oven-dry soil was determined.

ELISA. Fifty-microlitre portions of appropriate dilutions of extracted cell suspensions prepared as above were placed in wells of a 96-well microtitre plate. An indirect ELISA procedure was used to estimate cell concentrations (Mansoor, 1992).

RESULTS

Determination of a standard curve for ELISA tests on soil samples

Plate counts and indirect-ELISA values obtained with serially diluted suspensions of 15 sterile wheatfield soil samples inoculated with A. globiformis were used to determine the correlation between A₄₅₀ values and plate counts and to construct a standard curve. Log cell numbers were plotted against three dilutions of each sample (Fig. 1). A highly significant (P = 0.001) linear relationship was obtained (r = 0.98) and analyses of variance indicated that the straight line was statistically a good fit when soil samples contained more than 10⁷ cells (g soil)⁻¹.

The reliability of the curve for estimating cell numbers from indirect-ELISA values was tested with 13 sterile soil samples inoculated with different cell concentrations of A. globiformis. A comparison of indirect-ELISA estimates with plate counts of A. globiformis is shown in Fig. 2. Statistical analyses (t-test) showed that there were no significant differences (P = 0.05) between the two counting methods when plate counts were in the range 10⁶–10⁸ cells (g soil)⁻¹. The results also showed (Fig. 2) that it was not possible to detect A. globiformis cells using the indirect ELISA test when plate counts were below 10⁶ cells (g soil)⁻¹. The standard curve was used in a comparison of the fate of rods of A. globiformis in sterile and non-sterile wheatfield soil.

Growth of bacteria in non-sterile wheatfield soil

Fate of bacteria on contact slides. Triplicate slides were recovered immediately after placing the slides in contact with the soil and at 6 d intervals for 48 d. Biovolumes of the fluorescing cells were calculated (Fig. 3). The number and biovolume of cells dropped during the first 6 d of incubation. Examination of stained slides recovered at this stage indicated that some of the fluorescing cells were washed off the slides during the staining procedure (Fig. 4a). There were also green halos around sites of cell growth, suggesting that the cells were secreting antigenic materials and that cells had been lost during preparation. An increase in the number and biovolume of the fluorescing cells was observed after 12 d and continued to the 30th day, after which the number and biovolume of the cells rapidly declined. Stained slides recovered at and after 18 d showed microcolonies of considerable size formed on or around soil particles (Fig. 4b). Examination
of slides recovered at and after 24 d showed the development of fungal hyphae and actinomycete mycelium; they also showed microcolonies of considerable size on or near fungal hyphae (Fig. 4c). Fewer microcolonies were associated with actinomycete mycelium. After 12 d, at least 80% of the rods had reverted to cocci (Fig. 5).

**Fate of bacteria added directly to soil.** Triplicate soil samples were recovered immediately after adding bacteria and at weekly intervals for seven weeks. Bacteria were extracted from soil using the centrifugation extraction method and the biovolume of *A. globiformis* cells per g soil estimated using direct cell counts and ELISA techniques. Agreement between fluorescent antibody and ELISA estimates was good. A twofold decrease in the biovolume of cells was observed during the first week of incubation, followed by a gradual 10-fold increase that continued for 2 weeks, after which the biovolume of cells declined steadily to the end of the experiment (Fig. 6). Examination of stained slides of extracted cells recovered during the first week of incubation showed that the majority of cells were already coccoid.

**Growth of bacteria in non-sterile sand-dune soil**

The growth of *A. globiformis* in the non-sterile sand-dune soil was only investigated using the contact slide technique. A marked decrease in the biovolume of fluorescing cells was observed during the first 6 d of incubation and continued to the end of the experiment (Fig. 3). No growth was observed during the whole experiment. Stained slides recovered at and after 6 d of incubation showed that the majority of the fluorescing cells were dying or lysing cocci (Fig. 5). Fungal hyphae or actinomycete mycelium were very rare on the slides.

**Growth of bacteria in sterile wheatfield soil**

**Fate of bacteria on contact slides.** *A. globiformis* cells introduced on slides and placed in contact with the sterile wheatfield soil decreased in biovolume (as in the non-
Growth of Arthrobacter in soil

Fig. 5. Changes with time in the morphology of *A. globiformis* NCIMB 10683 cells on contact slides in sterile (●) and non-sterile (▲) wheatfield soil and sterile (●) and non-sterile (▼) dune sand.

Fig. 6. Growth curves of *A. globiformis* NCIMB 10683 inoculated into non-sterile wheatfield soil. Based on estimates from ELISA (●) and fluorescent antibody methods (■).

Fig. 7. Growth curves of *A. globiformis* NCIMB 10683 on contact slides in sterile wheatfield soil (●) and dune sand (■) detected with fluorescent antibodies.

Fig. 8. Growth curves of *A. globiformis* NCIMB 10683 inoculated into sterile wheatfield soil. Based on estimates from ELISA (●) and fluorescent antibody methods (■).

Fig. 9. *A. globiformis* NCIMB 10683 cells after 7 d growth in sterile wheatfield soil, showing the development of a myceloid cell as well as many cocci. Bar, 5 μm.

sterile soil) during the first 6 d of incubation (Fig. 7). However, an approximately 100-fold increase in the biovolume of the fluorescing cells was observed during the next 2.5 weeks, after which it dropped steadily. As in the non-sterile soil, examination of stained slides recovered from the sterile soil at and after 18 d of incubation showed micro-colonies of considerable size formed on or around soil particles; the majority of the fluorescing cells were cocci (Fig. 5).

Fate of bacteria added to soil. In sterile soil, the introduced cells behaved differently from those introduced into non-sterile soil. A significant 200-fold increase in the biovolume of the cells was observed during the first week of incubation, followed by a slight but significant decline during the second week (Fig. 8). Biovolumes measured between weeks 2 and 5 were not significantly different but a further significant decrease had taken place by week 7. The majority of cells from samples recovered during the first week of incubation were cocci, but a few were myceloid (Fig. 9). Agreement between the two
methods used to estimate cell numbers was good, but fluorescent antibody counts were slightly higher than ELISA estimates.

**Growth of bacteria in sterile sand-dune soil**

Growth of *A. globiformis* cells introduced into sterile sand-dune soil was only investigated using the contact slide technique. Cells placed in contact with sterile sand-dune soil showed similar behaviour to the cells in non-sterile sand-dune soil. A marked decrease in the biovolume of the fluorescing cells was observed during the first 6 d and continued to the end of the experiment (Fig. 7). Stained slides recovered from this soil showed similar results to those from non-sterile soil.

**Generation times of cells in soil**

Table 1 shows the generation times of *A. globiformis* in sterile and non-sterile wheatfield soil, estimated during exponential phase, using soil inoculation and contact slide techniques.

<table>
<thead>
<tr>
<th>Technique</th>
<th>Counting method</th>
<th>Mean generation time (h)</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>Sterile soil</td>
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<tr>
<td>Soil inoculation</td>
<td>Fluorescent antibody count</td>
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<tr>
<td></td>
<td>Indirect ELISA</td>
<td>28</td>
</tr>
<tr>
<td>Contact slide</td>
<td>Fluorescent antibody count</td>
<td>124</td>
</tr>
</tbody>
</table>

The mean generation time for *A. globiformis* in the non-sterile wheatfield soil was substantially higher than in the sterile soil (Table 1), a result similar to that obtained for *Rhizobium japonicum* by Schmidt (1974). In unsterile soil, an approximately twofold decrease in biovolume occurred during the first week of incubation, followed by a gradual 10-fold increase during the next 2 weeks. By contrast, in the sterile soil, a 200-fold increase in biovolume was observed during the first week of incubation. Survival of inocula in non-sterile habitats is known to be adversely affected by predation and competition (Habte & Alexander, 1977; Casida, 1980).

The maximum growth rate of *A. globiformis* NCIMB 10683 in continuous culture has been shown to be about 0.37 h⁻¹, equivalent to a generation time of 2.7 h (Luscombe & Gray, 1974). Luscombe & Gray (1971, 1974) showed that the transformation of rods into cocci depended upon the growth rate of the culture and that rods were characteristic of rapidly growing cultures. On the basis of such data, a change from coccus to rod in the natural environments was predicted if the generation time was greater than 13.8 h. The results obtained in the present study are consistent with this prediction. Direct observation of cells extracted from soil after one week of incubation showed that most of the rod-shaped cells in the inoculum had changed to cocci or short rods: they never reverted to rod forms, even though growth was occurring. This shows that cocci are not dormant forms but are the commonest growth form in soil for this organism.

**DISCUSSION**

Estimations of the biovolume of a particular species can be made both by counting cell numbers using fluorescent antibodies and by ELISA methods. However, ELISA was found to be insensitive when detecting cells in sterile soils and could not be used when populations fell below $10^6$ (g soil)⁻¹. When cell numbers exceeded this level, ELISA and fluorescent antibody results from soil samples containing a mixture of organisms correlated well. However, when there were differences they were not consistent: sometimes the ELISA estimates were higher and sometimes lower.

Ecological conclusions about bacteria should not be drawn from experiments on populations in pre-sterilized soil. The mean generation time for *A. globiformis* in the non-sterile wheatfield soil was substantially higher than in the sterile soil (Table 1), a result similar to that obtained for *Rhizobium japonicum* by Schmidt (1974). In unsterile soil, an approximately twofold decrease in biovolume occurred during the first week of incubation, followed by a gradual 10-fold increase during the next 2 weeks. By contrast, in the sterile soil, a 200-fold increase in biovolume was observed during the first week of incubation. Survival of inocula in non-sterile habitats is known to be adversely affected by predation and competition (Habte & Alexander, 1977; Casida, 1980).

The results obtained in the present study thus extend those obtained by Thompson *et al.* (1990, 1992) for *Arthrobacter* strain A109. In non-sterile soils, although marked decreases in the biovolume of *A. globiformis* cells were observed in the first week, division and formation of micro-colonies of considerable size on or around soil debris did occur in the next 2–3 weeks. The formation of micro-colonies was also observed around or near fungal hyphae and occasionally near actinomycete mycelium. However, the growth of *A. globiformis* cells was not conditional upon the growth and development of fungal hyphae, since their growth started well before fungal growth and observations on sterile soils showed similar large colonies of arthrobacters. This contrasts with the behaviour of *Bacillus subtilis* on contact slides in acidic
sand-dune soil, when fungal growth was a precondition for bacterial growth (Siala & Gray, 1974).

It has generally been assumed that contact slides change the environment of bacteria being studied in the soil and stimulate activity since they favour the condensation of water and provide a fresh and different physical surface for colonization. However, the use of both direct inoculation and inoculated contact slides in this study showed that contact slides actually slowed the rate of growth of bacteria, especially when they were in contact with otherwise sterile soil. Bacteria added directly to the soil will be in greater contact with nutrients, and oxygen diffusion to the cells may be restricted at slide surfaces: losses of cells from slides during preparation could also lead to inaccuracies.

Contact slides placed in sterile wheatfield soils induced the growth of a small number of myceloid cells. The formation of myceloid cells in a number of Arthrobacter strains in culture has been ascribed to increased osmotic stress (Deutch & Perera, 1992): others have related it to vitamin deficiencies. It is not clear what is responsible for it in soil but it was relatively rare under the conditions of this experiment.

A. globiformis NCIMB 10683 was isolated from a sand-dune soil planted with pine trees (Lowe & Gray, 1972), and Lowe & Gray (1973a) showed that it could grow in sterile alkaline dune soil from the C horizon. However, it failed to establish itself and died out in surface non-sterile alkaline dune sand in the present study. This may reflect the salt content of the sand used here which came from very close to the high-tide line.

REFERENCES


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