Formation and resuscitation of ‘non-culturable’ cells of *Rhodococcus rhodochrous* and *Mycobacterium tuberculosis* in prolonged stationary phase

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After growth of *Rhodococcus rhodochrous* in Sauton’s medium, and further incubation for about 60 h in stationary phase, there was a transient (up to 5 log) decrease in the c.f.u. count, whereas the total count remained similar to its initial value. At the point of minimal viability, the most probable number (MPN) count was 10 times greater than the c.f.u. count. This difference was further magnified by 3–4 logs (giving values close to the total count) by incorporating supernatant taken from growing cultures. A small protein similar to Rpf (resuscitation-promoting factor of *Micrococcus luteus*) appeared to be responsible for some of the activity in the culture supernatant. The formation of ‘non-culturable’ cells of the ‘Academia’ strain of *Mycobacterium tuberculosis* was similarly observed following growth in Sauton’s medium containing Tween 80 in sealed culture vessels, and further incubation for an extended stationary phase. This resulted in the formation, 4–5 months post-inoculation, of a homogeneous population of ostensibly ‘non-culturable’ cells (zero c.f.u.). Remarkably, the MPN count for these cultures was $10^5$ organisms ml$^{-1}$, and this value was further increased by one log using supernatant from an actively growing culture. Populations of ‘non-culturable’ cells of *Mycobacterium tuberculosis* were also obtained by the filtration of ‘clumpy’ cultures, which were grown in the absence of Tween 80. These small cells could only be grown in liquid medium (MPN) and their viability was enhanced by the addition of culture supernatant or Rpf. The ‘non-culturable’ cells that accumulated during prolonged stationary phase in both the *R. rhodochrous* and the *Mycobacterium tuberculosis* cultures were small ovoid and coccoid forms with an intact permeability barrier, but with undetectable respiratory activity. The authors consider these non-culturable bacteria to be dormant. The observed activity of culture supernatants and Rpf with ‘non-culturable’ bacterial suspensions invites the speculation that one, or more, of the cognate *Mycobacterium tuberculosis* Rpf-like molecule(s) could be involved in mechanisms of latency and reactivation of tuberculosis *in vivo*.

**Keywords:** dormancy, Nocardiaceae, VBNC, Rpf, *Micrococcus luteus*

**INTRODUCTION**

It is generally believed that pathogenic, slow-growing mycobacteria, such as *Mycobacterium tuberculosis* or *Mycobacterium leprae*, can persist for long periods of time *in vivo* after initial infection, by passing into a dormant or non-replicating persistent state (Gangadharam, 1995; Parrish *et al*., 1998; Wayne & Sohaskey, 2001). Such dormant *Mycobacterium tuberculosis* cells can persist for many years in the host, causing latent disease.

**Abbreviations:** ADC, albumen, glucose and NaCl; CTC, 5-cyano-2,3-ditolyl tetrazolium chloride; MPN, most probable number; Rpf, resuscitation-promoting factor; SN, supernatant; VBNC, viable but non-culturable.
NAD pool under anaerobiosis, an transition to anaerobic persistence. For example, the activity of \textit{Mycobacterium bovis} is convincingly established. The most widely accepted model for the transition of \textit{Mycobacterium tuberculosis} to an apparently dormant state is Wayne’s model, in which cells are subjected to gradual oxygen depletion. This results in the formation of an anaerobic, drug-resistant, non-replicating state (Wayne, 1994; Wayne & Hayes, 1996). Under these conditions, the putatively dormant bacteria shut down protein synthesis, but this restarts after the reintroduction of oxygen (Hu \textit{et al.}, 1998). A similar phenomenon has been found for \textit{Mycobacterium smegmatis} (Dick \textit{et al.}, 1998) and \textit{Mycobacterium bovis} (Lim \textit{et al.}, 1999). \textit{Mycobacterium tuberculosis}, \textit{Mycobacterium smegmatis} and \textit{Mycobacterium bovis} revealed similar changes during the transition to anaerobic persistence. For example, the activity of some dehydrogenases was induced to maintain the NAD pool under anaerobiosis, an \(\alpha\)-crystallin-like protein was expressed, and cell-wall thickening occurred in \textit{Mycobacterium tuberculosis} and \textit{Mycobacterium bovis} BCG (Cunningham & Spreadbury, 1998; Hutter & Dick, 1998; Lim \textit{et al.}, 1999). Another approach to obtain dormant \textit{Mycobacterium tuberculosis} in vivo is the Cornell mouse model. Mice infected by \textit{Mycobacterium tuberculosis} are treated with antibiotics, after which the number of culturable cells extractable from organs falls to zero (Wayne, 1994). Notably, mycobacterial DNA can still be detected in such apparently ‘sterile’ tissues and quantitative PCR has revealed the presence of amounts of DNA equivalent to about 10\(^5\) organisms (g tissue\(^{-1}\)) (de Wit \textit{et al.}, 1995). Subsequently, culturable \textit{Mycobacterium tuberculosis} cells reappear, either spontaneously or in response to immune suppression (Flynn & Chan, 2001; McCune \textit{et al.}, 1966). The essential difference between the two models is the maintenance of the high culturability of \textit{Mycobacterium tuberculosis} cells in Wayne’s model and the presence of a non-culturable state in the Cornell model. However, the absence of experiments in which the presumably dormant bacteria in the Cornell model can be resuscitated and enumerated \textit{in vitro} creates some difficulties in data interpretation. On the one hand, PCR may detect a substantial population of non-culturable (dormant) cells that subsequently resuscitate \textit{in vivo}. On the other hand, it may detect a large proportion of dead cells, in which case the re-emergence of colony-forming units would simply reflect the re-growth of a very small number of viable cells left after treatment with antibiotics (Kell \textit{et al.}, 1998). These two possibilities have not yet been distinguished experimentally.

\textit{Micrococcus luteus} is a member of the \textit{Actinomycetales} and is therefore related to \textit{Mycobacterium tuberculosis}. This organism can persist in a dormant state, following growth to stationary phase in batch culture and starvation for several months in the spent growth medium (Kaprelyants \textit{et al.}, 1993; Kaprelyants & Kell, 1993a). Dormant \textit{Micrococcus luteus} cells have lost the ability to grow on agar plates; however, the addition of supernatant (SN) from growing \textit{Micrococcus luteus} cultures to dormant cells has been shown to restore the ability to divide freely, thereby resuscitating the cells to normal, colony-forming bacteria (Kaprelyants & Kell, 1996; Kaprelyants \textit{et al.}, 1994). \textit{Micrococcus luteus} cells secrete a resuscitation-promoting factor (Rpf), which promotes the resuscitation of dormant cells (Kaprelyants \textit{et al.}, 1996). Rpf has subsequently been shown to be a small protein that is active at picomolar concentrations (Kaprelyants \textit{et al.}, 1999; Mukamolova \textit{et al.}, 1998). Genes encoding Rpf-like proteins are widely distributed throughout the \textit{Actinomycetales}; database searches have revealed that similar genes are present in mycobacteria, corynebacteria and streptomyces (Kell & Young, 2000). If, as seems likely, the Rpf-like proteins of these organisms have similar functional roles to that of \textit{Micrococcus luteus} Rpf, we may be able to predict the existence of a ‘non-culturable’, dormant state in other members of the \textit{Actinomycetales}.

The aim of the present study was to find conditions for the \textit{in vitro} transition of two organisms belonging to the \textit{Corynebacterineae} (fast-growing \textit{R. rhodochrous} and slow-growing \textit{Mycobacterium tuberculosis}) to a ‘non-culturable’ (possibly dormant) state. We also tested the hypothesis that secreted bacterial proteins may stimulate resuscitation of these two organisms.

**METHODS**

\textbf{Organisms and media.} \textit{Rhodococcus rhodochrous} NCIMB 13805 was grown aerobically at 37 °C in shake flasks (125 ml medium in 750 ml flasks, agitation at 200 r.p.m.) in nutrient broth E (Lab M) or in modified Sauton’s medium. The latter contained (per litre): MgSO\(_4\)\(\cdot\)7H\(_2\)O, 0.5 g; l-asparagine, 4 g; glycerol, 60 ml; iron ammonium citrate, 0.05 g; sodium citrate, 2 g; ZnSO\(_4\)\(\cdot\)7H\(_2\)O, 0.1 ml of a 1% solution in H\(_2\)O; K\(_2\)HPO\(_4\), 775 g; NaH\(_2\)PO\(_4\), 2 H\(_2\)O, 4.25 g. When the culture had reached stationary phase, agitation was continued at 37 °C for up to 2 weeks. The avirulent Academia strain of \textit{Mycobacterium tuberculosis} (Ogloblina & Ravich-Birger, 1958) was obtained from the Phthysopulmonology Center (Moscow). Cells were grown in loosely capped (plastic screw-caps) 14 ml test tubes containing 2 ml of unmodified Sauton’s medium supplemented with albumen, glucose and NaCl (ADC), without shaking at 37 °C (Connell, 1994). Cultures were held in stationary phase at 37 °C for up to 8 months. To obtain a population of single cells, cultures were passed through two Schott glass filters (50 µm and 20 µm, respectively) followed by a series of five nitrocellulose filters (4 µm, 2 µm, 1-5 µm, 0.8 µm and 0.45 µm, respectively; Whatman). To produce less-aggregated and oxygen-starved populations, \textit{Mycobacterium tuberculosis} cells were also grown in 200 ml of the same medium in the presence of 0.05% Tween 80, with shaking (150–200 r.p.m.) in 750 ml flasks fitted with rubber seals. Needles were employed for sterile sampling without oxygen input.

\textbf{Scanning electron microscopy.} Cells were fixed for 30 min in 3% glutaraldehyde, followed by treatment with 0.1 M sodium cacodylate buffer (pH 7.2). Samples were air-dried, sputter-coated with 20 nm Au particles and examined using a GSM-840A (Japan) scanning electron microscope.
Respiratory activity, membrane energization and the permeability barrier. The respiratory activity of the bacterial cells was determined by using the fluorescent redox dye CTC (5-cyano-2,3-ditolyl tetrazolium chloride; Polysciences), as described by Kaprelyants & Kell (1993b). Briefly, the bacteria were incubated in the presence of 4 mM CTC (freshly prepared in 10 mM sodium phosphate buffer, pH 7.0) for 30 min at 37°C. Fluorescence was monitored with a Nikon fluorescence microscope with excitation at 530–550 nm. Cells were stained with propidium iodide (4 µM in phosphate buffer), to assess the state of the membrane permeability barrier. Rhodamine 123 was used to monitor membrane energization, as previously described (Kaprelyants & Kell, 1992). Rhodamine 123 accumulation was sensitive to the uncoupler carboxylcyanide-3-chlorophenyl-hydrazone (CCCP). Cells were studied under the fluorescence microscope (excitation at 510–560 nm and emission at 590 nm for propidium iodide; excitation at 450–490 nm and emission at 520 nm for rhodamine 123).

Estimation of the redox potential. The changes in the oxidation/reduction potential of the growth medium during R. rhodochrous growth were measured in undiluted samples (3 ml) taken from the bacterial culture. Measurement of the redox potential was done with a platinum electrode (and an AgCl reference electrode; normal potential at 20°C was 201 ± 2 mV) by using an I-130.2M ionometer (Soyusanalit-pribor) under aerobic conditions at 20°C, with intensive stirring until a constant value was reached.

Assessment of cell viability. Bacterial suspensions were serially diluted in growth medium and duplicate 100 µl samples were plated in triplicate onto agar-solidified nutrient broth E (for R. rhodochrous) or onto agar-solidified, ADC-supplemented Sauton’s medium (for Mycobacterium tuberculosis). Plates were incubated at 37°C. After 5 days of growth for R. rhodochrous and after 2 months of growth for Mycobacterium tuberculosis, c.f.u. values were enumerated. The detection limit was 5 x 10^5 c.f.u. ml⁻¹. The same serially diluted samples of Mycobacterium tuberculosis were also employed for resuscitation and most probable number (MPN) assays. Using a fresh pipette tip each time, 100 µl samples were taken from each dilution. Ten of these samples were added to 10 replicate 10 ml Pyrex screw-cap tubes, each containing 2 ml Sauton’s medium supplemented with ADC. Rpf was added to five of the tubes to give a final concentration of 100 pM. Five additional samples were added to five replicate 10 ml Pyrex screw-cap tubes containing 2 ml filter-sterilized SN taken from cultures of Mycobacterium tuberculosis in the late-exponential phase of growth (30–40 days post-inoculation). The 15 tubes for each dilution were incubated at 37°C without shaking for 2 months.

Resuscitation and MPN assays for R. rhodochrous were performed in 48-well plastic plates (‘Corning’) containing either 0.5 ml modified Sauton’s medium (see above) or 0.5 ml filter-sterilized SN taken from R. rhodochrous cultures. Some wells contained 125 pM Rpf. All wells were supplemented with 0.5% yeast extract (LabM). Appropriate serial dilutions of R. rhodochrous cells (50 µl) were added to each well. Plates were incubated at 37°C, with agitation at 150 r.p.m. for 5 days. Tubes or wells with visible bacterial growth were enumerated. The detection limit was 5 x 10^6 c.f.u. ml⁻¹. The same serially diluted samples of Mycobacterium rhodochrous were subjected to centrifugation (12000 g, 20 min). The SNs were sterilized by passage through a 0.22 µm filter (Whatman) before use. Proteinase treatment of the SNs was performed at 37°C by adding trypsin (200 µg ml⁻¹) or proteinase K immobilized on acrylic beads (0.04 U ml⁻¹; Sigma), followed by the addition of trypsin inhibitor (400 µg ml⁻¹) to samples containing trypsin. The time of incubation was 30 min for trypsin-treated SNs and 3 h for proteinase-K-treated SNs.

Bacterial counts. The total number of Mycobacterium tuberculosis cells was determined microscopically using a Helber’s chamber. A minimum of 100 bacteria was counted and the SD of the total counts was not in excess of 20%.

Preparation of recombinant Micrococcus luteus Rpf. The Rpf protein of Micrococcus luteus [histidine-tagged recombinant form] was obtained as described by Mukamolova et al. (1998). Mono Q ion-exchange purification was omitted in some experiments. The purified protein was stored in 10 mM Tris/HCl (pH 7-4) containing 50% (v/v) glycerol at −20°C for up to 2 weeks, and the protein concentration was determined spectrophotometrically. Before use, all preparations were screened for growth-promoting activity using a small inoculum of Micrococcus luteus, as described by Mukamolova et al. (1998). Some preparations had poor activity; only those with substantial activity were employed for these experiments.

Preparation of antibody columns. Rabbits were immunized three times at 3-week intervals by subcutaneous injection with 1 ml of a 50% (v/v) mixture of Rpf (1 mg ml⁻¹ in water) and incomplete Freund’s adjuvant (Sigma). Serum was collected 10 days after the last immunization, and the immunoglobulin fraction was purified by a standard protocol using polyethylene glycol. Similar methods were used by Micropharm (Newcastle Emlyn, Carmarthenshire, UK) to raise and purify antibodies to a truncated version of Rpf (residues A₁₀–E₁₁₅) in sheep. Immunoglobulins were conjugated to CNBr-activated Sepharose 4B (Sigma) (Osterman, 1985).

RESULTS

Formation of ‘non-culturable’ cells of R. rhodochrous

R. rhodochrous cells were initially grown in the rich medium, nutrient broth E, for different times and then employed as an inoculum (10⁶ cells ml⁻¹) for growth in modified Sauton’s minimal medium (the elevated concentrations of phosphates in this medium stabilize the culture pH during stationary phase). Stationary phase was established within about 24 h, and further incubation resulted in a gradual decrease of the c.f.u. value, starting about 30 h post-inoculation (Fig. 1a). A minimum value was reached between 80 and 100 h post-inoculation, after which time the c.f.u. value gradually recovered until it finally exceeded the initial value. After the initial growth phase, the total count (estimated microscopically; data not shown) and the optical density remained almost constant, except that the optical density fell slightly, coincident with the period of non-culturability (Fig. 1a). The precise time at which culturability (c.f.u. counts) reached its minimum was variable from one experiment to another, but it correlated with a transient increase in the redox potential of the culture (Fig. 1b). The extent of the transient loss of culturability depended on the age of the inoculum. The
Fig. 1. R. rhodochrous cells grown in batch culture in modified Sauton’s medium show a transient loss of culturability. A culture in nutrient broth E (inoculated 18 h previously) was employed as inoculum (10^6 cells ml\(^{-1}\)) in (a) and (b) and the number of colony-forming units were monitored over time. The OD\(_{600}\) (a) and Eh (b) were also monitored. In (c), the effect of inoculum age on the maximum loss of culturability is shown. This experiment was repeated 15 times; the results of a typical experiment are shown. For further details see Methods. The SD values for c.f.u. determinations were generally within the limits almost covered by the symbols in the Figure; for example, they were 38%, 3% and 11% for the points at 24 h, 48 h and 96 h, respectively, in part (a). The SD values on measurements of Eh and OD\(_{600}\) in stationary phase were about 3% and 5%. (a) ●, OD\(_{600}\) value; ○, c.f.u. value. (b) ●, Eh value; ▽, c.f.u. value.

largest effect was observed with an inoculum taken from a culture that had itself been inoculated some 18–19 h previously (Fig. 1c). Optimum conditions for observing the loss of culturability (i.e. the biggest effect) were as indicated in Methods. However, loss of culturability was observed over a range of agitation speeds, between 100 and 250 r.p.m., and at culture volumes corresponding to between 12% and 25% of the volume of the culture flask.

Microscopic examination of the cells during the period of non-culturability revealed that the majority were ovoid or coccoid forms; this morphology contrasted with the characteristic elongated curved rod shape of exponentially growing cells (Fig. 2a). Only about 10% of the cells in the population were metabolically active [according to fluorescence microscopy with Rhodamine 123 as an indicator of membrane energization (Kaprelyants & Kell, 1992) and CTC as an indicator of respiratory activity (Kaprelyants & Kell, 1993b)]. Between 80% and 90% of the cells failed to stain with propidium iodide, indicating that their membrane permeability barrier was intact (Davey et al., 1999).
Resuscitation of ‘non-culturable’ cells of *R. rhodochrous*

Resuscitation of cells taken during the period of non-culturability (Fig. 1a) was performed in liquid medium, using procedures previously developed for resuscitating non-culturable cells of *Micrococcus luteus* (Kaprelyants et al., 1994). Fig. 3(a) shows that dilution of the culture into fresh Sauton’s medium containing 0.5% yeast extract followed by incubation for up to 40 h did not give any evidence of resuscitation. Addition of catalase (3 μg ml⁻¹) or pyruvate (0.5 mM) to the medium did not promote resuscitation either in liquid or on solid medium. Resuscitation was observed when the cells were incubated in yeast-extract-supplemented (0.5%), filter-sterilized SN taken from a late-exponential phase culture (20 h post-inoculation) of bacteria grown in Sauton’s medium (Fig. 3a). The resuscitation activity of the SN depended on its concentration; a maximal effect was observed when the proportion of SN to cells was 50% (v/v) and above (Fig. 3b). For a numerical estimation of the resuscitation of ‘non-culturable’ cells, MPN values were determined. This procedure (Kaprelyants et al., 1994) permits true resuscitation to be distinguished from the re-growth of a small number of initially viable cells, since it occurs at high dilutions where initially viable cells are absent. Incubation of the ‘non-culturable’ cells in Sauton’s medium resulted in a 10-fold increase in viability measured by the MPN method when compared with c.f.u. counts (Fig. 4a). Different variations in medium composition did not result in any further increase of the MPN value. Values were similar in Sauton’s medium and in a rich medium (nutrient broth E) (Fig. 4a), and the addition of ADC supplement or yeast extract at concentrations greater than 0.5% (v/v) was inhibitory (data not shown).
Table 1. Influence of different treatments on resuscitation activity of SN taken from exponentially growing cultures of *R. rhodochrous*

<table>
<thead>
<tr>
<th>SN treatment/conditions</th>
<th>‘Resuscitation index’</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>2500 ± 150</td>
</tr>
<tr>
<td>Control</td>
<td>2500 ± 150</td>
</tr>
<tr>
<td>Heat</td>
<td>270 ± 30</td>
</tr>
<tr>
<td>Autoclaving</td>
<td>18 ± 0.4</td>
</tr>
<tr>
<td>Boiling 5 min</td>
<td>110 ± 10</td>
</tr>
<tr>
<td>Boiling 10 min</td>
<td>158 ± 6.1</td>
</tr>
<tr>
<td>Boiling 20 min</td>
<td>20 ± 0.4</td>
</tr>
<tr>
<td>Storage</td>
<td>4 °C, 2 days</td>
</tr>
<tr>
<td></td>
<td>270 ± 30</td>
</tr>
<tr>
<td></td>
<td>4 °C, 6 days</td>
</tr>
<tr>
<td></td>
<td>8.5 ± 4.3</td>
</tr>
<tr>
<td></td>
<td>Ambient, 1 day</td>
</tr>
<tr>
<td></td>
<td>40 ± 6.6</td>
</tr>
<tr>
<td>Trypsin</td>
<td>37 °C, 30 min</td>
</tr>
<tr>
<td></td>
<td>150 ± 50</td>
</tr>
<tr>
<td>Proteinase K</td>
<td>Ambient, 1 h</td>
</tr>
<tr>
<td></td>
<td>10.8 ± 1.1</td>
</tr>
<tr>
<td>Dialysis</td>
<td>4 °C, overnight</td>
</tr>
<tr>
<td></td>
<td>310 ± 50</td>
</tr>
<tr>
<td>Ultra-filtration (&lt;10kDa)</td>
<td>1.8 ± 0.2</td>
</tr>
<tr>
<td>Passage through anti-Rpf antibody column</td>
<td>400, 55</td>
</tr>
<tr>
<td>Anti-Rpf</td>
<td>100, 11</td>
</tr>
<tr>
<td>Anti-truncated Rpf</td>
<td></td>
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</tbody>
</table>

However, a dramatic increase in MPN values was observed when ‘non-culturable’ cells were incubated in SN taken from bacterial cultures (Fig. 4a). The age of the SN was critically important. SN taken from cultures 18–20 h and 118–132 h post-inoculation was active, whereas that taken from cultures 45–70 h post-inoculation was inactive (Fig. 4b). Moreover, the SN taken between 45 and 70 h contained some inhibitory material(s) that resulted in a decrease of the MPN value, compared with the value obtained in the absence of SN (Fig. 4b). The ‘inhibitory’ SN also suppressed the growth of exponential cells of *R. rhodochrous* (data not shown). The speed of agitation of the 48-well plates used to measure MPN value was also critical for maximum resuscitation. The optimal speed was 150 r.p.m.; above or below this speed, the MPN value was significantly decreased.

Some properties of the resuscitation-promoting activity present in the culture SN are shown in Table 1. Autoclaving or boiling the SN for 20 min resulted in a loss of activity. Activity was lost during storage at 4 °C over several days. The resuscitation-promoting activity was sensitive to proteolytic enzymes, removed by ultra-filtration (molecular mass >10 kDa) and partially retained by dialysis (molecular mass >10 kDa). The passage of the SN through affinity columns containing immobilized anti-Rpf (or anti-truncated-Rpf) antibodies resulted in a substantial loss of activity, suggesting that part of the observed resuscitation activity was due to the presence of Rpf-like proteins. This conclusion was reinforced by the finding that picomolar concentrations of recombinant Rpf also stimulated resuscitation in Sauton’s medium supplemented with 0.5% yeast extract, although the effect was less pronounced than that of culture SN (Fig. 4a).

Formation of ‘non-culturable’ cells of *Mycobacterium tuberculosis*

*Mycobacterium tuberculosis* (Academia strain) was grown in Sauton’s medium supplemented with ADC, without agitation in capped test tubes. After growth for 1 month (early-stationary phase), samples were taken periodically (one tube per time point) and the total count and c.f.u. count were estimated. Since the cells were grown in medium lacking detergent, large aggregates (and sometimes also a surface pellicle) were produced. To facilitate the manipulation of individual cells, cultures were passed through a series of filters of decreasing pore size (20–0.45 µm). Moreover, we expected that cells with a non-culturable phenotype would have a reduced size, as has previously been observed with *Micrococcus luteus* (Mukamolova et al., 1995), *Mycobacterium tuberculosis* isolated from murine macrophages (Biketov et al., 2000) and *R. rhodochrous* (see above). The culturability of the cells (estimated as the ratio c.f.u. count/total count) decreased with decreasing filter pore size. The 1-5 µm filtrate was unable to form colonies on agar (Fig. 5), despite the fact that microscopy revealed the presence of large numbers of cells (total count 4 × 10⁷ cells ml⁻¹).

Non-culturable cells in the 1-5 µm filtrate were a mixture of ovoid cells, with a length of about 1-4 µm and small individual cocci, with a diameter of 0.5–0.7 µm (Fig. 2b). Both types of cells had an intact membrane barrier (according to propidium-iodide staining) and did not reveal any respiratory activity (not stained by CTC). As seen with *R. rhodochrous*, the degree of non-culturability of the cells in the 1-5 µm filtrate was time dependent (Fig. 5b). Culturability was minimal about 4 months post-inoculation, after which time the c.f.u. count gradually increased and approached the value of the total count after 8 months of cultivation. The total count of bacteria in this fraction did not change significantly during the entire period of the experiment (Fig. 5b). Oxygen availability was certainly restricted in the tube cultures used for this experiment, because incubation of *Mycobacterium tuberculosis* in stationary phase in oxygenated shake flasks resulted in the main-
Viability of *Mycobacterium tuberculosis* in relation to cell size and culture age. (a) A culture of *Mycobacterium tuberculosis* (4 months post-inoculation) was filtered consecutively through 50, 4, 2, 1.5 and 0.45 μm filters and the c.f.u. count/total count for each of the filtrates was determined. In part (b), actively growing cells of *Mycobacterium tuberculosis* were inoculated (10⁵ cells ml⁻¹) into Sauton’s medium supplemented with ADC, and replicate cultures were incubated at 37 °C without agitation for 8 months in capped test tubes. Periodically, samples were withdrawn and filtered through a 2 μm filter. The c.f.u. (●; SD between 10% and 20%) and total (▲; SD about 17%) counts of the filtrates were determined. The results of a typical experiment are shown.

![Graph](image1.png)

**Fig. 5.** Viability of *Mycobacterium tuberculosis* in relation to cell size and culture age. (a) A culture of *Mycobacterium tuberculosis* (4 months post-inoculation) was filtered consecutively through 50, 4, 2, 1.5 and 0.45 μm filters and the c.f.u. count/total count for each of the filtrates was determined. In part (b), actively growing cells of *Mycobacterium tuberculosis* were inoculated (10⁵ cells ml⁻¹) into Sauton’s medium supplemented with ADC, and replicate cultures were incubated at 37 °C without agitation for 8 months in capped test tubes. Periodically, samples were withdrawn and filtered through a 2 μm filter. The c.f.u. (●; SD between 10% and 20%) and total (▲; SD about 17%) counts of the filtrates were determined. The results of a typical experiment are shown.

To avoid problems associated with heterogeneity, we also determined whether similar non-culturable cells could be formed in the more homogeneous cultures that are produced when *Mycobacterium tuberculosis* is grown in the presence of a suitable detergent. To this end, cultures were grown in flasks containing ADC-supplemented Sauton’s medium containing 0.05% Tween 80 (a) or no Tween (b) and were grown for 4–5 months, as indicated in Methods. (a) Samples of comparatively homogeneous cultures were withdrawn periodically and the c.f.u. values were determined without prior filtration (SD between 10% and 20%). This experiment was repeated twice with similar results. In part (b), the bacterial population at 4 months (post-inoculation) was passed through a 2 μm filter and the total and c.f.u. counts of the filtrate were determined. MPN assays (columns with bold outline) were performed in ADC-supplemented Sauton’s medium with and without recombinant Rpf (125 pm) and also in SN taken from an actively growing culture of *Mycobacterium tuberculosis* (3–4 weeks post-inoculation). The results of 20 independent experiments are summarized, ± SD.

![Graph](image2.png)

**Fig. 6.** Production and resuscitation of ‘non-culturable’ cells of *Mycobacterium tuberculosis* in Sauton’s medium. Growing cells of *Mycobacterium tuberculosis* were inoculated (10⁵ cells ml⁻¹) in ADC-supplemented Sauton’s medium containing 0.05% Tween 80 (a) or no Tween (b) and were grown for 4–5 months, as indicated in Methods. (a) Samples of comparatively homogeneous cultures were withdrawn periodically and the c.f.u. values were determined without prior filtration (SD between 10% and 20%). This experiment was repeated twice with similar results. In part (b), the bacterial population at 4 months (post-inoculation) was passed through a 2 μm filter and the total and c.f.u. counts of the filtrate were determined. MPN assays (columns with bold outline) were performed in ADC-supplemented Sauton’s medium with and without recombinant Rpf (125 pm) and also in SN taken from an actively growing culture of *Mycobacterium tuberculosis* (3–4 weeks post-inoculation). The results of 20 independent experiments are summarized, ± SD.

period (Fig. 6a). However, in contrast to cells grown in static culture without Tween 80, this state of non-culturability was maintained for the remainder of the experimental period (4–8 months post-inoculation).

**Resuscitation of ‘non-culturable’ cells of *Mycobacterium tuberculosis***

Resuscitation of the ovoid and coccoid cells in the 1.5 μm filtrate of 4-month-old cultures was performed in capped test tubes containing liquid Sauton’s medium supplemented with ADC, using an MPN assay. In contrast to *Micrococcus luteus* or *R. rhodochrous*, estimation of viability in liquid medium (MPN) led to a substantial (5 log) increase in the number of viable cells
similar methods to obtain non-culturable cells of *Kaprelyants & Kell, 1993a). In this study, we used mained in a dormant state for several months stationary phase cultures of dormant, inactive state. Previously we found that either maintain some detectable activity or persist in a questionable. Cells with non-culturable phenotypes may and medical significance must be considered highly questionable. Thomas M. O. Shleeva and others

**Table 2. Resuscitation of Mycobacterium tuberculosis**

grown with agitation and in the presence of Tween 80

<table>
<thead>
<tr>
<th>Count</th>
<th>Culture age (months)</th>
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<tbody>
<tr>
<td></td>
<td>5</td>
</tr>
<tr>
<td>Total</td>
<td>7.2 × 10⁸</td>
</tr>
<tr>
<td>c.f.u.</td>
<td>&lt;5*</td>
</tr>
<tr>
<td>MPN</td>
<td>1.1 × 10³</td>
</tr>
<tr>
<td>MPN(_{ss})</td>
<td>1.0 × 10⁶</td>
</tr>
</tbody>
</table>

*Limit of detection.

(Fig. 6b). The presence of picomolar concentrations of recombinant Rpf occasioned an additional 1–2 log increase in cell viability. SN taken from a culture of *Mycobacterium tuberculosis* cells growing exponentially in ADC-supplemented Sauton’s medium produced an equivalent increase in the viable count. In some experiments, such as that shown in Fig. 6(b), viability estimated using the SN was equivalent to the total bacterial count. We were unable to obtain resuscitation of cells that passed through the 0.45 μm filters under any of the above conditions (data not shown).

*Mycobacterium tuberculosis* cells grown in the presence of Tween 80 were also able to resuscitate in liquid medium. Table 2 shows that 5-month-old and 6-month-old cells, which had completely lost the ability to form colonies on solid medium, were resuscitated in liquid medium. An additional 1–2 log increase in resuscitation was obtained in the presence of SN. These results are very similar to those observed with cells obtained by filtration of heterogeneous cultures.

**DISCUSSION**

The phenomenon of non-culturability of bacterial cells is a matter of intensive debate [for reviews see Barer (1997), Barer & Harwood (1999), Barer et al. (1998) and Kell et al. (1998)]. There are many publications claiming the existence of ‘viable but non-culturable’ (VBN(C) bacteria, including several pathogenic species of bacteria. However, only in relatively few cases has the transition of such bacteria to a normal, viable (culturable) state been shown. The great majority of the experiments published to date fail to provide an unequivocal demonstration of the resuscitation of non-culturability of bacteria (Kell et al., 1998). These publications also suggest that the viability of the bacterial cell population may be limited by the presence of non-culturability of bacteria (Kaprelyants & Kell, 1998). Unless such ‘VBN(C) cells can be resuscitated, their microbiological and medical significance must be considered highly questionable. Cells with non-culturability of phenotypes may either maintain some detectable activity or persist in a dormant, inactive state. Previously we found that stationary phase cultures of *Micrococcus luteus* contain a large proportion of ‘non-culturable’ cells that remained in a dormant state for several months (Kaprelyants & Kell, 1993a). In this study, we used similar methods to obtain non-culturability of cells of *R. rhodochrous* and *Mycobacterium tuberculosis*, two members of the Corynebacterineae related to *Micrococcus luteus*.

As we found for *Micrococcus luteus*, rather strict culture conditions must be satisfied to observe the transition of the majority of the cell population to a non-culturable state (Mukamolova et al., 1998). For *R. rhodochrous*, inoculum age (Fig. 1b), medium composition, speed of flask agitation and culture volume in relation to the flask capacity significantly influenced the formation of non-culturability bacteria. However, in contrast to *Micrococcus luteus*, non-culturability was a transient phenomenon for *R. rhodochrous* cells. For each culture, there was a window of several hours during which about 99.9% of the bacteria in the culture had lost culturability. Some previously published results also show a transient decrease in the c.f.u. count when a fast-growing organism, *Mycobacterium smegmatis*, is held in stationary phase (Dick et al., 1998; Smulders et al., 1999). The authors did not draw this specific conclusion from their data and considered this as a ‘death’ phase. Recently, Keer et al. (2000) have found that several mutants of *Mycobacterium smegmatis*, defective in stationary phase survival under aerobic or anaerobic conditions, revealed a transient decrease in viability followed by an increase in culturability. The authors suggest at least two possible explanations for this effect: (a) re-growth (cryptic growth) of viable cells, and (b) formation of ‘non-culturable’ (possibly dormant) bacteria in stationary phase and their subsequent resuscitation (Keer et al., 2000, 2001). Since the authors did not measure culturability by MPN counts, direct evidence was neither sought nor obtained for resuscitation of non-culturability cells.

In our study, we applied MPN assays to measure viability. This method permits numerical estimation of the number of resuscitatable cells in liquid medium at high dilutions, effectively circumventing the problem of re-growth of residual viable cells during resuscitation (Kaprelyants et al., 1994; Kell et al., 1998). This assay revealed the presence of ‘non-culturable’ cells of *R. rhodochrous* during a rather short time interval in stationary phase. The transient character of the loss of cell viability during stationary phase suggests that there are sequential processes occurring. First, there is a transition to a ‘non-culturable’ state. This then seems to be followed by the (cryptic?) growth of residual viable bacteria, possibly accompanied by the growth of resuscitated bacteria after 86–96 h of incubation. The individual contributions of (a) residual viable cells and (b) resuscitated cells to the observed growth could vary, depending on the particular set of culture conditions (inoculum age, aeration conditions, growth medium, etc.). Non-culturability cells may be formed under many different conditions, but we have found conditions where resuscitatable cells make up the majority of the bacterial population (and are therefore experimentally accessible). Several questions remain. For example, why do *R. rhodochrous* cultures start to lose their culturability? Why do the viable bacteria fail to grow during the period of transition to the ‘non-culturable state’,
why do they start to grow subsequently? During the transition to ‘non-culturability’, we found evidence for the presence of an inhibitory component(s) in the SN, which is no longer present once the c.f.u. counts start to increase again. Further studies are needed to elucidate the possible role of accumulation/destruction of this inhibitor in culture behaviour.

The extent of resuscitation often exceeded the proportion of active cells in the population (about 10% in all experiments, according to Rhodamine-123 and CTC staining) and in some experiments approached 100%. This fact indicates that inactive cells, with a morphology characteristic of dormancy (small coccoidal cells with enhanced phase-contrast; Mukamolova et al., 1995), can resuscitate. Therefore, we suggest that incubation of *R. rhodochrous* cells under the conditions described resulted in the formation of a population of dormant bacteria, with a characteristic ‘non-culturable’ phenotype.

*Mycobacterium tuberculosis* held in stationary phase in capped tubes also revealed the transient formation of ‘non-culturable’ cells (on a different time scale). However, because of the marked non-homogeneity of *Mycobacterium tuberculosis* cultures grown without detergent, filtration was required to isolate the fraction of non-culturable cells. The filtered fraction containing non-culturable bacteria may not include all such bacteria in the population; aggregates, unable to pass through the filter, may also contain such bacteria. The increase in the c.f.u. count after 4 months of incubation in stationary phase (Fig. 5b) could be due to the growth of initially viable cells, to the resuscitation of non-culturable cells, or to a mixture of both of these factors in the highly heterogeneous population. The substantially homogeneous populations grown in the presence of Tween 80 were unable to form colonies for a long period of time (Fig. 6a). We suspect that it may also be possible to establish stable populations of ‘non-culturable’ cells for *R. rhodochrous* in stationary phase, but the appropriate conditions for this were not identified here.

There is a substantial difference between the ‘non-culturable’ cells of the two species studied, in respect of their ability to resuscitate. For *R. rhodochrous*, SN was needed for resuscitation (Fig. 4a), whereas significant resuscitation (up to 5 logs) of *Mycobacterium tuberculosis* cells occurred in liquid Sauton’s medium without any additions (Fig. 6b). SN or Rpf further increased resuscitation of *Mycobacterium tuberculosis* cells by up to 2 logs (Fig. 6b). These data clearly demonstrate the fact that the term ‘non-culturability’ has an operational meaning only (Barer, 1997; Kell et al., 1998); *Mycobacterium tuberculosis* cells that were unable to form colonies on plates were able to grow in liquid medium, as was reported by Biketov et al. (2000).

As previously seen with *Micrococcus luteus* (Kaprelants et al., 1994), SN taken from exponentially growing cells of both species studied possessed a ‘resuscitation-promoting’ activity. In *R. rhodochrous* SN, activity was associated with a small protein, which probably belongs to the Rpf family as SN passed through an affinity column containing anti-Rpf antibodies (M.O. Shleeva, unpublished results). Moreover, Southern hybridization of BamHI-, PstI-, PvuII- or Xbol-digested *R. rhodochrous* DNA using a rpf-specific probe revealed the presence of four hybridizing bands (data not shown). Recombinant *Micrococcus luteus* Rpf stimulated resuscitation of *R. rhodochrous*, but it was less active than SN.

A similar conclusion could be made with respect to the active compound in *Mycobacterium tuberculosis* SN, as Rpf also showed stimulatory activity (although it was less active than SN; Fig. 6b). Moreover, the Rpf-like proteins of *Mycobacterium tuberculosis* do indeed have growth factor activity and are able to stimulate the growth of non-culturable cells of *Mycobacterium bovis* BCG held in prolonged stationary phase (G.V. Mukamolova, unpublished results). In addition, ‘non-culturable’ cells have also been found after persistence of *Mycobacterium tuberculosis* in murine macrophages *ex vivo*. The viability of these cells was restored by incubation in the presence of Rpf in liquid medium (Biketov et al., 2000).

Zhang and co-workers have recently reported that non-culturable cells accumulate during prolonged incubation of *Mycobacterium tuberculosis* H37Ra cells in stationary phase in the presence of Tween 80 (Sun & Zhang, 1999; Zhang et al., 2001). In contrast to our study, their cell populations contained a significant number of colony-forming units. Slight differences in culture conditions may account for this (e.g. we maintained bacteria in capped tubes without oxygen input). They showed that SN from growing *Mycobacterium tuberculosis* cultures contained phospholipids, which increased the number of colony-forming units in aged cultures when added to plates and permitted growth from low inocula in liquid medium. Another active compound isolated from the SN was an 8 kDa protein, Rv1174c. Chemically synthesized peptides corresponding to three different segments of Rv1174c were added at micromolar concentrations (i.e. much higher than those at which Rpf is active) to liquid medium inoculated with old cells. After 5–10 days incubation, the number of recoverable colony-forming units was enhanced. We suggest that the phospholipids and micromolar concentrations of peptides employed by the authors may have promoted repair of injured cells (this may be especially true for the dried cells with zero c.f.u. counts used in this study) (Ray & Speck, 1973). Since the authors did not measure viability by MPN, re-growth of a residual portion of viable cells cannot be excluded in some experiments and the extent of the proposed resuscitation activity cannot be quantified.

The transition of mycobacterial cells to ‘dormancy’ in the Wayne model *in vitro* is based on the formation of non-replicating bacteria during adaptation to anaerobic conditions.
conditions (Wayne, 1994; Wayne & Hayes, 1996; Wayne & Sohaskey, 2001). Protein synthesis in this state is generally reduced (Hu et al., 1998), although certain genes, such as acr, are up-regulated (Desjardins et al., 2001; Hu et al., 1999; Yuan et al., 1996). The model has been established in several different species, viz. Mycobacterium tuberculosis, Mycobacterium smegmatis and Mycobacterium bovis BCG (Dick et al., 1998; Lim et al., 1999; Wayne, 1994). However, in contrast to our study, mycobacterial cells in Wayne’s model retain high viability and develop sensitivity to metronidazole as they become anaerobic (Wayne & Sramek, 1994), indicating that they remain metabolically active. Therefore, they may not be considered truly dormant, but rather in a state of (O2) starvation–survival (Kaprelyants et al., 1993). We suggest that the difference in culturability may reflect the different times of persistence of Mycobacterium tuberculosis cells in a non-growing state (days in Wayne’s model and months in our study). Restoration of culturability requires resuscitation in liquid medium, which may either occur spontaneously or require the provision of compounds (e.g. growth factors) present in the SN of growing cells. The proportion of cells able to resuscitate spontaneously and those requiring growth factors may vary from one experiment to another, and resuscitation may depend on the particular conditions employed for the establishment of ‘non-culturability’.

Prolonged incubation of Mycobacterium tuberculosis cells in stationary phase may reflect more accurately the situation in vivo. In this connection we may stress that the essence of the Cornell model of dormant Mycobacterium tuberculosis is the creation of a period of ‘sterility’ in vivo, induced by the administration of antibiotics after infection, when Mycobacterium tuberculosis cannot be detected by c.f.u. counts (Wayne, 1994) (in contrast to PCR; de Wit et al., 1995). As shown in Fig. 7, Wayne’s model may possibly represent a step towards the establishment of true dormancy in Mycobacterium tuberculosis, which is characterized by the emergence of ‘non-culturable’ cells that need to be resuscitated before they can resume active growth. The necessity for anaerobic conditions to produce ‘non-culturable’ cells of Mycobacterium tuberculosis in our experiments may reflect a mechanistic link between Wayne’s model and the model employed here. More work is required to establish whether the proposed sequence of events does indeed take place during the transition to, and reactivation from, the latent state in vivo. The observed activity of SN or Rpf with ‘non-culturable’ bacterial populations suggests that one or more of the cognate Mycobacterium tuberculosis proteins could be involved in mechanisms of latency and reactivation of tuberculosis in vivo.

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