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Citation: MEANWELL, R.J.L. and SHAMA, G., 2006. Chitin in a dual role as substrate for Streptomyces griseus and as adsorbent for streptomycin produced during fermentation. Enzyme and microbial technology, 38(5), pp. 657-664

Additional Information:

- This is a journal article. It was published in the journal, Enzyme and microbial technology [© Elsevier] and the definitive version is available at: doi:10.1016/j.enzmictec.2005.07.017

Metadata Record: https://dspace.lboro.ac.uk/2134/3482

Publisher: © Elsevier

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Chitin in a dual role as substrate for *Streptomyces griseus* and as adsorbent for streptomycin produced during fermentation.

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Abstract

Streptomycin production from chitin by *Streptomyces griseus* was compared using two different types of bioreactor. The first was equipped with a combined U-shaped paddle and Rushton impeller. Also employed was a bioreactor of novel design in which the chitin was contained in a wire mesh basket that was totally submerged in a liquid salts medium. During operation the chitin was gently fluidised by air admitted into the basket. Fermentation was continued in both bioreactors until maximum antibiotic titres were achieved whereupon operation was interrupted to allow the streptomycin adsorbed to the chitin substrate to be extracted into pH 3.0 buffer before continuing fermentation of the same batch of chitin a second time. At a chitin concentration of 10 % (w/v) the highest streptomycin yields (e. 5.5 mg/L) were obtained using the stirred bioreactor, however growth occurred more rapidly in the vertical basket bioreactor.

*Keywords*: Chitin; *Streptomyces*; Streptomycin; Stirred Bioreactors; Vertical Basket Bioreactor

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1 Introduction

Chitin is an abundant biopolymer that occurs in the shells of crustaceans, the carapaces of insects and in the cell walls of fungi. Although many of these sources of chitin are too dispersed to be exploited, commercial fishing of crustaceans, and their subsequent processing, generates large quantities of chitin-rich waste [1]. This waste can be purified by either chemical or biological methods [2] to yield chitin that can be used in a number of applications within the food [3] medical and veterinary [4,5] or environmental protection industries [6].

One common factor in many of the more traditional applications to which chitin and the closely related partially deacetylated polymer, chitosan, have been put is based on the powerful chelating properties of these polymers. Both chitin and chitosan have been used to remove a diverse range of compounds from waste streams including heavy metals [7] and dyes [8,9]. Chitin has also been used in the microbial production of chitinases as fungal control agents [10,11] and as a source of polyene antibiotics produced by a number of species of Monascus [12]. More recently, it was shown that a strain of Pseudomonas fluorescens cultured on chitin produced thermally stable antifungal compounds active against a number of phytopathogens [13].

Although the ability to utilise chitin is fairly widespread amongst both marine and terrestrial micro-organisms [14], nearly all species of the genus Streptomyces are chitinolytic [15]. The streptomycetes, are economically important because they produce a number of high value secondary metabolites including many antibiotics.
Streptomycin, the first aminoglycoside antibiotic discovered, is produced by *Streptomyces griseus*, and remains a useful weapon in the fight against the resurgence of tuberculosis [17].

Streptomyces fermentations are characterised by high broth viscosities owing to the filamentous nature of the organism, and the maintenance of adequate mixing and aeration in conventional stirred bioreactors is achieved by vigorous agitation. Operation under these conditions leads to high shear rates and can result in reduced yields, difficulties with subsequent downstream processing and even product degradation [18,19]. The incorporation of chitin, a solid, in culture media in conventional stirred bioreactors would result in additional stresses being imposed on the streptomycetes filaments due to the abrasive action of the chitin particles. Consequently, in investigating the cultivation of *S. griseus* on chitin for the production of streptomycin, we sought to employ bioreactors designed to minimise physical damage to the organism. We employed two different types of bioreactor; the first was a stirred bioreactor equipped with an impeller of special design. This comprised a combination of U-shaped paddle and Rushton impeller that was designed to be operated at low speed whilst maintaining the chitin in suspension. The second type of bioreactor investigated here was one of novel design and incorporated a static vertical basket in which the chitin was contained whilst being immersed in a salts medium. Air was passed directly into the basket so as to both provide oxygen to the culture of *S. griseus*, and to gently fluidise the chitin flakes.

In addition to serving as substrate, the chitin also acted as adsorbent for the streptomycin produced during fermentation. Because the chitin is solid and easily separated from the aqueous broth, it was possible to interrupt the fermentation so that
adsorbed streptomycin could be recovered. Following this, the fermentation was allowed to proceed again before conducting a further cycle of streptomycin recovery.

2 Materials and methods

2.1 Micro-organism & Cultivation

*Streptomyces griseus* (NCIMB 8136) was purchased in lyophilised form (NCIMB, Aberdeen, UK). Spores of the organism were maintained on sterile soil and were cultured on yeast malt extract agar (NCIMB Medium No. 29) at 28° C when required for bioreactor experiments. The liquid medium used for such experiments comprised flakes of chemically purified chitin (Sigma Aldrich plc, Poole, Dorset, UK) at concentrations specified in the text and the following salts per litre of distilled water; K₂HPO₄, 0.76 g; KH₂PO₄, 0.3 g; MgSO₄, 0.5 g; FeSO₄, 0.01 g; ZnSO₄, 0.0018 g; MnCl₂, 0.0016 g. The pH of the medium was adjusted to 6.5 and was autoclaved at 121° C for 20 minutes. Inoculum for bioreactors comprised 200 ml of culture grown in the salts medium supplemented with 1 % (w/v) chitin for 96 hours in an incubator-shaker at 100 rpm and 28 °C.

2.2 Streptomycin Determination

The concentration of streptomycin was determined using a standard bioassay procedure [20] and was based on the inhibition of the growth of *Bacillus subtilis* (NCIMB 8054). The only variation introduced into the procedure was in the method
of producing that the spores of *B. subtilis* [21]. Culture broths were first filtered through 0.2 μm Whatman cellulose nitrate membrane filters (Fisher Scientific, Loughborough, Leics.) and subsequently using 2 kDa membranes (Dow Danmark A/S, Nakskov, Denmark). Aliquots of filtered culture broths (150 μL) were added to wells cut into agar plates which were incubated for 48 hours at 30° C. Zones of inhibition were measured using Vernier callipers. All determinations were performed at least in duplicate. A standard calibration curve was obtained using aqueous solutions of streptomycin (Sigma Aldrich plc).

2.3 CO₂ Determination

Gas samples were taken from the air exit line of the bioreactor using a gas-tight syringe and directly injected into a gas chromatograph (Model 104, Pye Unicam, Cambridge, UK) equipped with a thermal conductivity detector operated at 180° C. The glass chromatography column was 6 mm diameter and 175 cm long and was packed with Molecular Sieve 5A (Phase Separations, Deeside, Clwyd, UK).

2.4 Contamination Monitoring

Samples of fermentation broth were routinely plated onto Tryptone Soy Agar (Oxoid, Unipath Ltd., Basingstoke, Hants) and Malt Extract Agar (Oxoid) and incubated at 30 and 25° C respectively and examined after 2 and 4 days for the presence of contamination.
2.5 Bioreactors

2.5.1 Stirred Bioreactor

A standard stirred 2-litre glass bioreactor vessel (LH Engineering Co. Ltd., Stoke Poges, Bucks, UK) was used with a working volume of 1.5 litres. The air flowrate to the sparger was 100ml/min. Agitation was provided by an impeller that comprised a U-shaped paddle and a Rushton impeller of 5 cm diameter (Figure 1). The U-shaped paddle was 23 cm wide and 18 cm high and fabricated from stainless steel strips 1cm wide and was operated at 55 rpm. After filling with medium, the bioreactor was autoclaved at 121º C for 40 minutes.

2.5.2 Vertical Basket Bioreactor

This comprised a cylindrical wire frame of height 105 cm and diameter 65 cm (Figure 2) fabricated around a hollow central support member and held in place by circular end plates. Wire mesh (0.2 mm aperture) was wound round the frame to create a chamber into which the chitin could be contained. The base plate was fabricated from sheet steel fitted with a wire mesh plate 1cm above it. Air was admitted into the basket via the hollow central support. The air emerged from four 0.5 mm diameter holes at the base of the support and below the wire mesh and flowed upwards in the form of fine bubbles through the basket. The liquid working volume was 1.0 litre. An air flowrate of 100 ml/min was found to be sufficient to bring about gentle fluidisation of the chitin flakes contained in the chamber. In addition, two aerators were installed outside of the basket to both further aerate the liquid medium and to provide agitation;
the flowrate to each of these aerators was 150 ml/min. The total liquid volume was 1 litre. The mass of chitin flakes added to the basket was 35 g. After filling with medium, the bioreactor was autoclaved at 121º C for 40 minutes.

2.6 Recovery of Adsorbed Streptomycin from Chitin

2.6.1 Stirred Bioreactor

At the termination of fermentation experiments the chitin was allowed to sediment and the culture broth drawn off. An equal volume of pH 3.0 buffer (made by mixing 597 ml citric acid (0.1 M) with 153 ml dibasic sodium phosphate (0.2 M)) and diluting to 1500ml buffer was added. Extraction was allowed to proceed at ambient temperature for 24 hours in situ in the bioreactor at ambient temperature at an impeller speed of 55 rpm. The treated chitin was then added to the salts medium in the bioreactor and re-autoclaved as described above. The bioreactor was re-inoculated once the temperature of the medium had reached 28º C.

2.6.2 Vertical Basket Bioreactor

At the termination of fermentation experiments the aqueous portion of the medium was pumped out of the bioreactor and replaced with an equal volume of sterile pH 3.0 buffer solution. Extraction was then allowed to proceed for 24 hours under conditions of fluidisation. After extraction of the streptomycin the buffer was pumped out of the
bioreactor and replaced with an equal volume of salts medium. Fermentation was allowed to proceed immediately without necessitating a second inoculation.

3 Results

3.1 Stirred Bioreactor

Figure 3 shows CO₂ profiles with time for two fermentations performed in succession on the same batch of chitin in the stirred bioreactor with in each case an initial chitin concentration of 7.5 % w/v. In the first fermentation the CO₂ peak occurs 48 hours earlier than in the subsequent one but in both cases the peak CO₂ concentrations are within 5 % of one another. For reasons given in the Discussion, antibiotic titres in Figure 4 and in all subsequent figures are reported as ‘apparent streptomycin concentration’. The streptomycin titres (Figure 4) display a broadly similar pattern to one another, however, that for the first fermentation shows a slightly more pronounced peak occurring 100 hours after the CO₂ maximum. The final streptomycin concentration reached at the end of the first fermentation at 2.5 mg/L is marginally greater than that achieved in the second fermentation (2.3 mg/L). The pH change over the entire time course of this and all subsequent fermentations never amounted to more than 0.4 units and suggests that the chitin provided some buffering capacity.

The use of the specially designed impeller enabled the chitin concentration to be increased to 10 % (w/v) without any settling out of the chitin or its accumulation in any of the ‘dead zones’ of the bioreactor. As for the fermentations described above,
the same batch of chitin was subjected to two successive fermentations. CO₂ profiles obtained under these conditions are shown in Figure 5. As for the previous fermentations, the profiles are very similar to one another and reach maximum concentrations at times comparable to those of Figure 4. Significantly, the CO₂ peak also occurs earlier in the second fermentation. Streptomycin titres are compared in Figure 6. The curve for the first fermentation displays a peak of 2.2 mg/L at approximately 380 hours, whereas that for the second fermentation peaks at 340 hours before declining and rising again to a final value of 2.0 mg/L. In both cases antibiotic titres were lower than those achieved at the lower chitin concentration of 7.5 %.

3.2 Vertical Basket Bioreactor

The procedure for conducting two successive fermentations with the same batch of chitin were similar to those for the stirred bioreactor but differed in that the chitin was not re-autoclaved after the first fermentation. Figure 7 shows the CO₂ concentration against time profiles for 10 % (w/v) chitin fermentation using the vertical basket bioreactor. These are similar to one another, both reaching maxima at approximately 175 hours, significantly earlier than was the case for the stirred bioreactor. The maximum streptomycin titres were also similar to one another (Figure 8). The maximum titre reached in the first fermentation was 2.8 mg/L at 300 hours and 3.0 mg/L at approximately 250 hours. Antibiotic titres were both greater than those obtained in the stirred bioreactor and were achieved in shorter times.

Table 1 shows streptomycin yields in the liquid medium, those recovered from the solid chitin and total yields. The former were obtained by multiplying streptomycin
titres at the end of the fermentation by final liquid volume remaining in the bioreactor – the working volume of the stirred bioreactor was 1.5 litres and that of the vertical basket bioreactor was 1.0 litre. The streptomycin yields recovered from solid chitin were obtained by multiplying streptomycin titres in the extraction buffer at 24 hours by the volume of buffer employed. The stirred bioreactor clearly resulted in the production of greater amounts of streptomycin both in the liquid medium and adsorbed on the chitin. Although total yields are comparable at both chitin concentrations (7.5 and 10.0 %), the results of Table 1 suggest that the contribution made to total yield by the ‘solid fraction’ was greater at the higher chitin concentration. Both ‘solid’ and ‘liquid’ streptomycin yields are lower in the vertical basket bioreactor.

4 Discussion

Preliminary studies (not reported here) conducted using shake flasks had revealed the presence of anti-microbial compounds in the early phases of growth of S. griseus on chitin. These were compounds that inhibited the growth of the bioassay bacterium (B. subtilis) but were evidently not secondary metabolites as they were produced during the idiophase. GC and HPLC assays were employed in the early phases of fermentation in order to eliminate definitively the presence of streptomycin and whilst none was detected, the tests were too insensitive for all but the highest concentrations of streptomycin encountered in this work [22]. Further tests revealed that these anti-
microbial compounds could be removed by filtering the culture broths through a 2 kDa membrane and therefore all samples taken during the course of fermentations for streptomycin determination by bioassay were filtered thus prior to testing by bioassay. However, Fourier Transform Infra Red (FTIR) analysis conducted on solid chitin remaining at the end of fermentation revealed the unambiguous presence of streptomycin [22]. As the quantity of solid chitin required for FTIR analyses was too large to permit the technique to be used routinely throughout the time course of fermentations, reliance had to be placed on the bioassay as the most sensitive method for detecting streptomycin. Under such conditions it was felt more appropriate to refer to ‘apparent streptomycin’ titres.

The presence of solid chitin flakes in the fermentation medium to which the growing streptomycete filaments became attached, prevented the direct determination of biomass concentration. Consequently, reliance had to be placed on CO₂ concentration in the exit gas stream from the bioreactor as an indicator of growth. Unpublished work conducted with the strain of *S. griseus* used here in a medium devoid of chitin had shown that a close correlation existed between the concentration of biomass and the concentration of CO₂ in the gas outlet from the bioreactor [22]. This correlation was important and confirmed the validity of using exit gas CO₂ concentration as an indirect indication of microbial growth. CO₂ evolution is commonly used as an indicator of growth and has been preferred to other indirect methods such as assays for cell wall components [23]. Work conducted to characterise the effects of agitation on cell lysis in *Streptomyces clavuligerus* also revealed concomitant increases in the concentrations of biomass and CO₂ [18].
Peak CO\textsubscript{2} evolution was achieved more rapidly in the vertical basket bioreactor than in the stirred bioreactor. This suggests that biomass growth was more rapid in the former (Figures 5 & 7). This was undoubtedly a reflection of the different shear regimes established in the two bioreactors, as mechanical stresses are known to influence not only the morphology of streptomycetes but to have the ability to induce cell lysis [18,24]. Evidently, the shears inside the basket were less intense than those in the stirred bioreactor.

The maximum yield of streptomycin achieved using the vertical basket bioreactor (2.8 mg/L) compare favourably with that achieved (3.0 mg/L) in a chitin-free medium containing yeast extract and malt extract [22]. Whilst growth was more rapid in the vertical basket bioreactor, total streptomycin titres were greater in the stirred bioreactor (Table 1). Convincing evidence has been accumulated to show that stresses of different types can enhance the production of secondary metabolites. The imposition of heat, salt or ethanol stresses stimulated antibiotic production by *Pseudomonas fluorescens* [25]. Similarly, greater yields of the antibiotic microcin were obtained when *E. coli* was cultured in a shake flask rather than in a rotating wall bioreactor intended to simulate conditions of microgravity [26].

The advantage offered by the vertical basket bioreactor configuration was that the ‘draw and fill’ process could be conducted with relative ease and without having to autoclave the chitin residue in between and without encountering subsequent contamination.
Taken together these findings suggest that the performance of the vertical basket bioreactor might be improved still further. Conditions similar to those described here could be maintained to achieve rapid biomass growth during idiophase. Once optimal biomass concentration had been reached, the hydrodynamic environment inside the basket could be altered so as to increase the level of mechanical stress imposed on S. griseus in order to maximise streptomycin yield. The most direct way of achieving this would be to increase the air flowrate to the basket which would have the effect of increasing the fluidisation intensity. Moreover, future studies of this type would need to be conducted using industrial strains of S. griseus which produce very much higher yields of streptomycin [27] in order to establish whether a process based on the vertical basket bioreactor could compete with the conventional method used to produce streptomycin.

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Figure 1 Impeller Arrangement for Stirred Bioreactor
Figure 3 CO₂ Profiles during successive fermentation of chitin (7.5 %) in a stirred bioreactor

Figure 4 Antibiotic Profiles during successive fermentation of chitin (7.5 %) in a stirred bioreactor

Figure 5 CO₂ Profiles during successive fermentation of chitin (10 %) in a stirred bioreactor
Figure 6 Antibiotic Profiles during successive fermentation of chitin (10 %) in a stirred bioreactor

Figure 7 CO₂ Profiles during successive fermentation of chitin (10 %) in a Vertical Basket bioreactor

Figure 8 Antibiotic Profiles during successive fermentation of chitin (10 %) in a Vertical Basket bioreactor
<table>
<thead>
<tr>
<th>Bioreactor</th>
<th>Stirred 1st Ferm</th>
<th>Stirred 2nd Ferm</th>
<th>Vertical Basket 1st Ferm</th>
<th>Vertical Basket 2nd Ferm</th>
</tr>
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<tbody>
<tr>
<td>Chitin Conc. (mg/L)</td>
<td>7.5</td>
<td>7.5</td>
<td>10.0</td>
<td>10.0</td>
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<tr>
<td>Streptomycin Yield (Liquid, mg)</td>
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<td>4.49</td>
<td>3.36</td>
<td>3.09</td>
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<tr>
<td>Streptomycin Yield (Solid, mg)</td>
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<td>1.37</td>
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<td>2.46</td>
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<tr>
<td>Total Streptomycin Yield (mg)</td>
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<td>5.86</td>
<td>5.61</td>
<td>5.55</td>
</tr>
</tbody>
</table>

Table 1. Streptomycin Yields Obtained in Stirred and Vertical Basket Bioreactors