CONTINUOUS-CULTURE STUDIES OF INTERACTIONS AMONG HUMAN SKIN-COMMENSAL BACTERIA

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The study of interactions among skin commensals has been prompted by the recent finding that more than one-fifth of normal persons possess antibiotic-producing bacteria on their skin (Selwyn and Ellis, 1972). The antagonistic effects of a strain of *Staphylococcus epidermidis* against a variety of bacteria have been investigated both semi-quantitatively on solid media and quantitatively in liquid batch cultures (Marsh and Selwyn, 1977). However, batch cultures permit only short-term growth studies, and the organisms in such a system undergo unrealistically wide changes in population density in comparison with those observed in most in-vivo situations. Studies have therefore now been performed in a chemostat, which provides stable long-term growth conditions.

MATERIALS AND METHODS

*Bacterial strains.* These were strain S6+, an antibiotic-producing strain of *S. epidermidis*, biotype 4 (Baird-Parker, 1974); strain S6-, an antibiotic non-producing *S. epidermidis* of biotype 4; and strain M7, a *Micrococcus* of biotype 7 (Baird-Paker, 1965). All three strains had been isolated from the forehead of one of us (S. S.). Previous studies on solid and in liquid media showed that strain M7 was antagonised by strain S6+ but not by strain S6-; the latter was itself unaffected by strain S6+ (Marsh and Selwyn, 1977).

*Media.* Peptone water was used as growth medium for the continuous cultivation of strains S6+, S6- and M7 (both pure and mixed) at two concentrations: 0.1% (w/v) for studies at low population levels and 0.5% (w/v) for high population levels.

Selective media were developed for strains S6+, S6- and M7 in accordance with the MICs of various antibiotics for them. These media enabled accurate viable counts of each strain to be made from mixed cultures. The three selective media used were: nutrient agar with bacitracin 1.25 μg per ml and fusidic acid 0.196 μg per ml for strain S6+; nutrient agar with bacitracin 1.25 μg per ml and benzylpenicillin 0.10 μg per ml for strain S6-; and nutrient agar with furazolidone 7.5 μg per ml and kanamycin 0.156 μg per ml for strain M7.

*Chemostat.* The design of the simple continuous culture apparatus was based on that of a "Porton-type" chemostat described by Evans, Herbert and Tempest (1970). It had two reservoirs for medium that could be used alternately through a two-way tap, thus allowing the effects of both 0.1% and 0.5% peptone water to be observed during the same experiment. The volume of the culture vessel was 100 ml, and a water jacket maintained the temperature of the culture at 33°C. A variable-speed flow-inducer, MRHE 22 (Watson-Marlow Ltd) was used to provide a constant dilution rate of 0.3 per h in all experiments.

*Viable counts.* Culture fluid for viable counts was removed twice daily from the culture vessel by a sampling device in a port on top of the culture vessel (Marsh, 1975). Colony counts per ml were made on the appropriate selective media by the method of Miles, Misra and Trwin (1938), and growth curves were constructed. Differences in counts greater than 0.4 log10 were significant at the 5% level.

RESULTS

Pure cultures of all three strains in 0.5% peptone water reached steady-state conditions at $10^7 \pm 0.1$ for strain M7, $10^7 \pm 0.1$ for strain S6- and $10^7 \pm 0.3$ for S6+. In 0.1% peptone water the three strains stabilised at a population level of $10^6 \pm 0.1$ in pure culture.

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TABLE

Colony counts of antibiotic-producing (strain S6+) and non-producing (strain S6−) staphylococci and indicator micrococcus (strain M7) when grown in pairs or as pure cultures in 0.5% peptone water

<table>
<thead>
<tr>
<th>Time (days)</th>
<th>Mean log_{10} colony counts per ml in pairs (and pure cultures)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Pair 1</td>
</tr>
<tr>
<td></td>
<td>Strain S6−</td>
</tr>
<tr>
<td>0</td>
<td>7.5(*)</td>
</tr>
<tr>
<td>1</td>
<td>7.0 (7.8)</td>
</tr>
<tr>
<td>2</td>
<td>7.3 (7.7)</td>
</tr>
<tr>
<td>3</td>
<td>7.3 (7.8)</td>
</tr>
<tr>
<td>4</td>
<td>7.2 (7.9)</td>
</tr>
<tr>
<td>5</td>
<td>7.7 (7.7)</td>
</tr>
<tr>
<td>6</td>
<td>7.5 ...</td>
</tr>
<tr>
<td>7</td>
<td>...</td>
</tr>
<tr>
<td>8</td>
<td>...</td>
</tr>
<tr>
<td>9</td>
<td>...</td>
</tr>
</tbody>
</table>

* Started as batch cultures and therefore not comparable with paired cultures until day 1.
... = Not done.

The results of growing two strains together in the chemostat in 0.5% peptone water confirm our previous finding on solid media and in liquid batch cultures that the growth of strain S6− had no effect on that of strain M7. Under similar conditions, strain S6+ inhibited the growth of strain M7 by a factor of >3 log_{10} (the table). Both in pure and mixed culture the results were not influenced by the inoculum size.

When the three strains were grown together in 0.1% peptone water, the viable counts were similar to those obtained in pure cultures and no antagonism was detected. However, in 0.5% peptone water, when strain S6+ was inoculated after 2 days, the viable count of strain M7 fell over 200-fold after some delay, while that of strain S6− stayed relatively constant (fig. 1).

The results when all three strains were grown together in mixed culture and the growth medium was changed during the course of the experiment are shown in fig. 2. In 0.1% peptone water the three strains maintained steady-state conditions with no demonstrable antagonism. When, after 3 days, the inflow was changed to 0.5% peptone water, strains S6+ and S6− responded immediately to the richer medium, and the viable counts of both increased to levels similar to those obtained in pure cultures. However, strain M7 failed to respond to the increased concentration of nutrients; after a small fall, the viable count stayed at a level slightly lower than that in 0.1% peptone. Nevertheless, strain M7 was able to respond to this change in the medium when grown in mixed culture without strain S6+, although it reached a lower steady-state population than did strain S6− (fig. 3). When, however, strain S6+ was introduced after 5 days, the viable count of strain M7 fell by a factor of 1 log_{10} while that of strain S6− remained constant.

**DISCUSSION**

Antagonism of strain M7 was seen only in 0.5% peptone water, in which viable counts of strain S6+ reached at least 10^7; evidently the smaller populations of strain S6+ in 0.1% peptone water did not produce inhibitory concentrations of antibiotic. This explanation is supported by assays of the crude inhibitory agent in samples of media examined during
FIG. 1.—Growth in mixed chemostat cultures of strains S6− and M7, with subsequent addition of strain S6+, in 0.5% peptone water.

FIG. 2.—Growth in mixed chemostat cultures of strains S6+, S6− and M7 in 0.1% peptone water, and subsequently in 0.5% peptone water.

FIG. 3.—Growth in mixed chemostat cultures of strains S6− and M7 first in 0.1% peptone water, then in 0.5% peptone water, and finally after the addition of strain S6+.
our previous batch-culture studies (Marsh and Selwyn, 1977), as well as by the effects of changing the growth medium during the course of an experiment (fig. 2). Antagonism took between 12 h and 3 days to become manifest in continuous culture. This is considerably longer than we had observed in batch culture, in which "washing-out" of antibiotic does not occur (Marsh and Selwyn, 1977). In all the chemostat experiments, however, the viable count of strain S6\(^-\) remained fairly constant under stable conditions, showing that the inhibition of strain M7 was not due to fluctuations in the operation of the culture system.

The micrococci are obligate aerobes and so are sensitive to inadequate aeration of the culture medium. The results obtained in batch cultures (Marsh and Selwyn, 1977), and those given in the table show, however, that oxygen limitation was not responsible for the inhibition of strain M7. When a mixture of strains S6\(^-\) and M7 was grown in 0.5% peptone, both strains reached viable counts approaching those obtained under the same conditions in pure culture. In contrast, when strains S6\(^+\) and M7 were grown together under identical conditions, strain M7 was inhibited.

In its original habitat on normal human skin, strain S6\(^+\) was unable to suppress strain M7, and both grew in a relatively steady state with strain S6\(^-\) (Selwyn, 1975). However, when skin hydration was greatly increased, strain S6\(^+\) readily exerted an inhibitory effect against strain M7 and other sensitive organisms (Selwyn, Marsh and Sethna, 1976). Under these conditions, skin is comparable to a suitable agar medium, supporting good bacterial growth and production of antibiotic substances without allowing excessive dilution of the latter. This concept is well illustrated when comparing mixed-culture results on solid and in liquid media. Previous work (Marsh and Selwyn, 1977) showed that the bactericidal action of the strain S6\(^+\) inhibitory agent could be readily demonstrated on solid media, but only the extremely sensitive group-D diphtheroid was completely inhibited in liquid batch cultures. Continuous-culture techniques offer many obvious advantages over batch cultures, but suffer from the unavoidable disadvantage, common to all liquid systems, of diffusion and dilution of metabolic products. In agar, high local concentrations can readily reach bactericidal levels, whereas in broth the titres produced by relatively fewer cells dispersed in a comparatively large volume of fluid appear usually to attain only bacteriostatic levels.

An improved system is clearly required for the long-term quantitative study of bacterial interrelationships on solid surfaces. This type of system should combine the quantitative steady-state conditions of traditional continuous-culture techniques with a solid surface for the growth of the desired organisms and the retention of microbial products. This would not only be more suitable for investigating antagonisms of the type discussed and more relevant to skin ecology, but would also find applications in the study of other microbial interactions. Such a system is under development and the preliminary results have been reported (Milyani, 1976).

**SUMMARY**

Chemostat studies were made of an antibiotic-producing *Staphylococcus epidermidis* strain (S6\(^+\)), a similar, but antibiotic non-producing *S. epidermidis* strain (S6\(^-\)), and a sensitive indicator strain of *Micrococcus* sp. (M7). Pure and mixed continuous cultures were investigated at low population levels (in 0.1% peptone water) and at higher levels (in 0.5% peptone water). Strain S6\(^+\) antagonised the growth of strain M7 when its colony count was maintained above 10\(^7.0\) per ml, while strain S6\(^-\) remained unaffected.

**REFERENCES**
