The Stability of Antibiotics in Soils

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SUMMARY: Ten antibiotics have been included in this study; eight of them were metabolic products of fungi isolated from soils of the Bagshot Sand type. Their stability in Bagshot Sand soils and in a neutral garden loam has been investigated. Some were more stable than others, the rate of inactivation varied from soil to soil, but all exhibited a fair degree of stability in some of the soils. Four types of inactivation could be distinguished: (1) The natural pH of the soil was sometimes that at which the antibiotic was intrinsically unstable; this was noted specially with albidin, frequentin, gliotoxin, penicillin and viridin. (2) Inactivation caused by some form of biological activity, indicated by less rapid inactivation in heat-treated than in untreated soil, was observed with griseofulvin, mycophenolic acid and patulin. (3) Adsorption on the soil was noticeable only in the case of streptomycin, the only basic antibiotic studied; acid-washed sand was able to bind appreciable quantities of this antibiotic. (4) Some other form of inactivation, probably chemical in nature, was concerned in the inactivation of gladiolic acid, penicillin and streptomycin.

Discussion of the natural significance of antibiotics logically follows the discovery of their production by micro-organisms. Similarly, consideration of their possible role in soil ecology follows from the fact that many soil micro-organisms are capable of producing them. Prerequisites of a balanced judgement on the subject include information on: (a) the types of micro-organism commonly isolated from various types of soil; (b) the antibiotics produced by such soil micro-organisms; (c) the stability of such antibiotics in the soil from which their producers were isolated; (d) the mode of inactivation of these antibiotics where this is found to occur; (e) the possibilities of detecting antibiotic production and accumulation in natural soils; (f) competition between strains of micro-organisms which do and do not produce antibiotics; (g) the significance—if any—of antibiotics to the micro-organisms which produce them.

The present paper is concerned with the stability and mode of inactivation of antibiotics in soil. In order to relate this as closely as possible to conditions under which antibiotics might naturally be produced, the majority of the antibiotics chosen for study were produced by fungi isolated from the Bagshot Sand type of soil, in which production of antibiotics under natural conditions has been suspected (Brian, Hemming & McGowan, 1945; Rayner, 1945). Furthermore, among the soils selected for study were soils from different levels of a Bagshot Sand soil profile. Streptomycin has been included, although actinomycetes in general are scarce in these acid soils, to afford a basis of comparison with the somewhat similar studies of Siminoff & Gottlieb (1951) and Pramer & Starkey (1951).
MATERIALS

Soils. Four of the soils used were obtained from the profile of a pit dug in a typical Bagshot Sand podsol on Wareham Heath, Dorset. The fifth sample, included for comparison, was a neutral light garden soil from Welwyn, Hertfordshire. The origin and properties of these soils are shown in Table 1.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Horizon</th>
<th>Depth (in.)</th>
<th>Description</th>
<th>pH value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wareham I</td>
<td>A₀</td>
<td>0-2</td>
<td>Litter and peat</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td>A₁</td>
<td>2-6</td>
<td>Black sandy peat</td>
<td>4·2</td>
</tr>
<tr>
<td></td>
<td>A₂</td>
<td>6-10</td>
<td>Slightly leached grey</td>
<td>5·0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>sand</td>
<td>5·4</td>
</tr>
<tr>
<td>Wareham II</td>
<td>A₃</td>
<td>10-18</td>
<td>Leached sand</td>
<td>4·2</td>
</tr>
<tr>
<td>Wareham III</td>
<td>B₁₋₂</td>
<td>18-22</td>
<td>Podsol pan</td>
<td>4·3</td>
</tr>
<tr>
<td>Wareham IV</td>
<td>C</td>
<td>22-26</td>
<td>Brown sandy gravel—</td>
<td>4·9</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>some clay</td>
<td>5·0</td>
</tr>
<tr>
<td>Garden</td>
<td></td>
<td>0-3</td>
<td>Light neutral loam</td>
<td>7·2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>7·2</td>
</tr>
</tbody>
</table>

* Soils from these horizons were not used in the experiment.

After collection the soils were air-dried, coarsely crushed and stored in tins at room temperature. The sand used in some experiments was washed for 24 hr. in running water, then for 12 hr. in c. n-HCl, and finally for another 36 hr. in running water. After this treatment very little clay remained and it had virtually no buffering capacity. Assays made as later described on aqueous extracts of these soils, at the concentration used in the experiments, showed no antifungal or antibacterial activity. Thus, all activity measured in the experiment can only have been caused by the antibiotic added.

Antibiotics. The following antibiotics were studied: albidin (Curtis & Grove, 1947), frequentin (Curtis, Hemming & Smith, 1951), gladiolic acid (Brian, Curtis & Hemming, 1948), gliotoxin (Brian, 1946), griseofulvin (Brian, Curtis & Hemming, 1946), mycophenolic acid, patulin, viridin (Brian & McGowan, 1945), penicillin and streptomycin. All these, except the last two, were prepared in these laboratories from fungi isolated from soils of the Bagshot sand type. These organisms are listed in Table 2.

Frequentin, patulin, penicillin and streptomycin are soluble in cold water, so that solutions of the required concentration could easily be prepared. The remaining antibiotics being only sparingly soluble in water are most conveniently brought into solution in a small quantity of a water-miscible organic solvent followed by dilution with water. To complicate the interpretation of results by the introduction of organic solvents into the system was undesirable, as these might well alter the microbiological activity of soil, so this method was not used. Gladiolic acid and mycophenolic acid were obtained as aqueous solutions of their sodium salts by solution in sodium
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hydroxide of appropriate strength. Gliotoxin and viridin were unstable in simple aqueous solution and were prepared in a dilute McIlvaine buffer by refluxing in a citric acid solution of appropriate strength, and adding the phosphate of the buffer in appropriate quantities later. Albidin was dissolved in the cold in water slightly acidified with HCl. Griseofulvin was dissolved by refluxing in water. The concentration of griseofulvin, viridin and albidin solutions was determined, after preparation of the solutions, by biological assay.

Table 2. Origin, potency and method of assay of antibiotics

<table>
<thead>
<tr>
<th>Antibiotic</th>
<th>Produced from</th>
<th>Potency</th>
<th>Assay</th>
</tr>
</thead>
<tbody>
<tr>
<td>Albidin</td>
<td><em>Penicillium albidum</em> Sopp (BRL248)</td>
<td>0-6</td>
<td>BA</td>
</tr>
<tr>
<td>Frequentin</td>
<td><em>P. frequentans</em> Westl. (BRL737)</td>
<td>3-0</td>
<td>BA</td>
</tr>
<tr>
<td>Gladiolic acid</td>
<td><em>P. gladioli</em> McCull &amp; Thom (BRL59)</td>
<td>7-0</td>
<td>BA</td>
</tr>
<tr>
<td>Gliotoxin</td>
<td><em>P. terikoskii</em> Zal. (BRL536) <em>Trichoderma viride</em> Pers. ex Fr. (BRL211)</td>
<td>3-5</td>
<td>BA</td>
</tr>
<tr>
<td>Griseofulvin</td>
<td><em>P. nigricans</em> (Bainier) Thom (BRL250)</td>
<td>1-1</td>
<td>BA</td>
</tr>
<tr>
<td>Mycophenolic acid</td>
<td><em>P. stoloniferum</em> Thom (BRL438)</td>
<td>0-4</td>
<td>BA</td>
</tr>
<tr>
<td>Patulin</td>
<td><em>P. expansum</em> Link (BRL440)</td>
<td>—</td>
<td>SCP</td>
</tr>
<tr>
<td>Viridin</td>
<td><em>Trichoderma viride</em> Pers. ex Fr. (BRL213)</td>
<td>0-06</td>
<td>BA</td>
</tr>
<tr>
<td>Penicillin</td>
<td><em>P. chrysogenum</em> Thom</td>
<td>1690 units/mg.</td>
<td>SCP</td>
</tr>
<tr>
<td>Streptomycin</td>
<td><em>Streptomyces griseus</em> (Krainsky) Waksman &amp; Henrici</td>
<td>423 units/mg.</td>
<td>SCP</td>
</tr>
</tbody>
</table>

Notes. The numbers appearing in parentheses after names of organisms are the accession numbers in the culture collection of Butterwick Research Laboratories (BRL). Penicillin G was supplied by I.C. Pharmaceuticals Ltd. Streptomycin was supplied by Boots Pure Drug Co. Ltd. Potencies refer (except in the case of streptomycin and penicillin) to least concentrations (μg./ml.) preventing germination of *Botrytis allii* spores or (in case of griseofulvin) producing curling of *Botrytis allii* hyphae. Methods of assay: BA = *Botrytis allii* spore germination test; SCP = *Bacillus subtilis* cylinder plate assay.

METHODS

Experimental layout. The stability of each antibiotic in a McIlvaine buffer (0-01 M-citric acid + 0-02 M-Na,HPO₄) over a pH range of 3-7 was first determined. Next, the stability of similar solutions was determined in the presence of acid-washed sand. Finally, the stability of solutions of the antibiotic was determined in the presence of each of the five soils previously described, using both air-dried but otherwise untreated soil (subsequently referred to as fresh) and soil autoclaved at 15 lb./sq. in. for 25 min. (subsequently referred to as partially sterile). In most cases both water solutions and buffer solutions of the antibiotic were exposed to fresh soil, the buffer being of a pH value close to the natural pH value of the soil; buffer solutions only were exposed to partially sterile sand and soil. For reasons already explained experiments with viridin and gliotoxin could only be carried out in buffer. Since the buffers
used were very dilute, the pH value occasionally drifted slightly during the course of the experiment.

In the soil and sand experiments 250 ml. antibiotic solution was placed in a 'Glaxo' culture vessel and 25 g. (air-dry weight) of soil added. This gave a 2 mm. layer of soil or sand with a 10 mm. supernatant layer of solution. It is appreciated that such conditions might seem to be unduly anaerobic and that an excessive proportion of antibiotic solution was exposed to a given volume of soil. It was thought, however, that under such conditions processes of inactivation would be slowed down and more readily studied. Moreover, preliminary experiments in which this condition was compared with the more obviously aerobic conditions of a soil percolator (Jefferys & Smith, 1951), indicated that there was little difference between the two experimental conditions. The static technique, being far simpler to handle, was therefore adopted.

In all experiments two concentrations of antibiotic were used. The lower was the lowest that could be conveniently assayed and the higher was in all cases ten times greater than the lower.

**Methods of assay.** Samples for assay were aseptically taken and every effort was made to remove at the same time a part of the soil present, in order to maintain an approximately constant ratio between soil and solution. Assays were made within 1 hr. of setting up the experiment and thereafter at suitable intervals. All samples were filtered through No. 5 Whatman filter-paper before assay.

Patulin, penicillin and streptomycin were assayed by the cylinder plate technique (Foster & Woodruff, 1944). The test organism in each case was *Bacillus subtilis*, a strain received from I.C. Pharmaceuticals Ltd. being used for assay of patulin and penicillin, and a strain (NCTC 7241) obtained from Boot's Pure Drug Co. Ltd. for streptomycin. The assay agar contained peptone, Lab-Lemco meat extract and Marmite. Standard curves were constructed daily and the concentrations of the antibiotics in the unknown solutions estimated graphically from these curves.

The remaining antibiotics were assayed by a serial dilution spore germination test using conidia of *Botrytis allii*. Brian & Hemming (1945), describing the method, used the inhibition of germination as the end-point of the assay. In the present work, the end-point used was based on loss of the germ tubes, which occurs at higher dilutions than inhibition of spore germination. This results in a more sensitive assay. Titres are expressed in 'dilution units'. This term is the number of consecutive twofold dilutions, through which inhibition of germination or stunting of germ-tubes can be recognized. These units are proportional to the logarithm of the concentration of the active material.

The assay of griseofulvin is different in that the limiting concentrations needed to produce the various characteristic morphological changes in germ-tubes were recorded; results are presented similarly in dilution units.
RESULTS

These are expressed in Figs. 1–19, and commented on in the text below.

Albidin

Albidin is unstable throughout the range pH 3.3–6.9 in buffer solution (Fig. 1). In the more acid solution (pH 3.3–4.8), after an initial rapid fall in activity, a steady level of activity is reached. This can be interpreted as meaning that albidin breaks down to a second antifungally active material of greater stability at low pH. At pH 5.9 and above, the active degradation product is also unstable. Sand had little effect on the rate of inactivation of albidin. The soils varied in their effect (Fig. 2). Wareham II and IV had little effect. In Wareham I, and less noticeably in Wareham III, the rate of inactivation in fresh soil is greater than would be expected merely from the pH of the solution, though partially sterile soil had little effect. This can best be interpreted as due to microbial activity; such differential inactivation in fresh and partially sterile soil is termed 'biological inactivation' in the following pages. In the garden soil inactivation is extremely rapid, as would be expected from the high pH; indications of biological inactivation are slight.

Frequentin

Buffer solutions of frequentin (Fig. 3) are highly stable in the range pH 3.3–4.6. Activity is slowly lost at pH 5.7 and more rapidly at pH 6.9. Sand has no effect on the rate of inactivation at neutrality, but at lower pH it may somewhat increase the rate of inactivation (Fig. 3). The rate of inactivation in Wareham II and IV soils (Fig. 4) is similar to that in buffer or in buffer and sand. In Wareham I and III soils inactivation is more rapid, and in Wareham I the greater rate of inactivation in fresh soil as compared with the rate in partially sterile soil, suggests that here again biological inactivation is taking place. Inactivation is very rapid in garden soil; this is mainly attributable to the high pH, but there is some indication of biological inactivation in the fresh soil.

Gladiolic acid

Gladiolic acid solutions in buffer (Fig. 5) are fairly stable in the range pH 3.4–7.0 and addition of sand has no effect. Once again Wareham II and IV soils had no effect. Inactivation at a much greater rate takes place in Wareham I and III and garden soils (Fig. 6); the rate of inactivation is little greater in fresh than in partially sterile soil. This indicates that in these soils inactivation is mainly due to some cause other than microbial decomposition or inactivation attributable to pH.

Gliotoxin

Buffer solutions of gliotoxin (Fig. 7) are highly stable at pH 3.4 and 4.9. Above pH 4.9 slow inactivation takes place, being more rapid the higher the pH. Sand had little if any effect on the rate of inactivation. In the presence of all the soils such inactivation as was observed (Fig. 8) could be explained by the pH of the system.
Fig. 1. Fungistatic activity of solutions of albidin (initially c. 30 µg./ml.,

Fig. 2. Fungistatic activity of solutions of albidin (initially c. 30 µg./ml.,
Figure 3. Fungistatic activity of solutions of frequentin (initially c. 100 μg./ml., ■, and c. 10 μg./ml., □) stored at 25° in buffer in the presence and absence of sand.

Figure 4. Fungistatic activity of solutions of frequentin (initially c. 100 μg./ml., ■, and c. 10 μg./ml., □) stored at 25° in contact with variously treated soils.
Fig. 5. Fungistatic activity of solutions of gladiolic acid (initially c. 30 μg./ml., ■, and c. 8 μg./ml., □) stored at 25° in buffer in the presence and absence of sand.

Fig. 6. Fungistatic activity of solutions of gladiolic acid (initially c. 30 μg./ml., ■, and c. 8 μg./ml., □) stored at 25° in contact with variously treated soils.
Fig. 7. Fungistatic activity of solutions of gliotoxin (initially c. 200 µg./ml., ■, and c. 20 µg./ml., □) stored at 25° in buffer in the presence and absence of sand.

Fig. 8. Fungistatic activity of solutions of gliotoxin (initially c. 200 µg./ml., ■, and c. 20 µg./ml., □) stored at 25° in contact with variously treated soils.
Fig. 9. Fungistatic activity of solutions of griseofulvin (initially c. 30 μg./ml., ■, and c. 3 μg./ml., □) stored at 25° in contact with variously treated soils.

Fig. 10. Fungistatic activity of solutions of mycophenolic acid (initially c. 20 μg./ml., ■, and c. 2 μg./ml., □) stored at 25° in buffer in the presence and absence of sand.
Fig. 11. Enzymatic activity of solutions of mycoamylase (initially c. 2000 μg/ml, ■) stored at 25°C in buffer and in contact with variously treated soils.

Fig. 12. Estimated concentrations of solutions of mycoamylase (initially c. 2000 μg/ml, ■) stored at 25°C in buffer and in the presence and absence of sand.
Fig. 13. Estimated concentrations of solutions of patulin (initially c. 2000 µg./ml., ■, and c. 200 µg./ml., □) stored at 25° in con-

Fig. 14. Estimated concentrations of solutions of penicillin (initially c. 50 units/ml., ■, and c. 1 units/ml., □) stored at 25° in buffer in the presence and absence of
Fig. 15. Estimated concentrations of solutions of penicillin (initially c. 50 units/ml, □, and c. 5 units/ml, □) stored at 25° in contact with variously treated soils.

Fig. 16. Estimated concentrations of solutions of streptomycin (initially c. 400 units/ml, □, and c. 40 units/ml, □) stored at 25° in buffer in the presence and absence of sand.
Fig. 17. Estimated concentrations of solutions of streptomycin (initially c. 400 units/ml., ■, and c. 40 units/ml., □) stored at 25° in contact with variously treated soils.

Fig. 18. Fungistatic activity of solutions of viridin (initially c. 100 μg./ml., ■, and c. 10 μg./ml., □) stored at 25° in buffer in the presence and absence of sand.
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**Griseofulvin**

Solutions of griseofulvin were completely stable over the range pH 3.4–7.0 and addition of sand had no effect, and accordingly detailed data are not presented. Rapid biological inactivation occurred in the fresh garden soil (Fig. 9). It was stable in Wareham II and IV soils; it was also stable in Wareham III, though in the solutions in water an inexplicable rapid loss in activity took place towards the end of the experiment. In Wareham I soil the more concentrated solutions in buffer showed no indication of biological inactivation, and quite high titres were recorded after 40 days; however, the rather lower general level of activity as compared with that recorded in buffer solution with no soil present, contrasted with the rapid and complete disappearance of griseofulvin from the less concentrated solutions, suggests that some such process as a physical adsorption took place.

**Mycophenolic acid**

Mycophenolic acid solutions in buffer at pH 4.6–7.0 are only very slowly inactivated; at pH 4.0 and below inactivation is rather more rapid (Fig. 10). Sand had no effect. The antibiotic was very stable in Wareham II, III and IV soils (Fig. 11). In the Wareham I soil, whether fresh or partially sterile, activity gradually disappeared over 16 days and thereafter began to reappear. In fresh garden soil rapid biological inactivation took place.
Patulin

Patulin was almost completely stable in buffer solution in the pH range 3.3–6.8; at pH 6.8 slow inactivation took place (Fig. 12). Sand had no appreciable effect on the rate of inactivation. It was very stable in all soils of the Wareham series (Fig. 13). Inactivation was more rapid in partially sterile garden soil and was greatly accelerated in fresh garden soil, indicating a considerable biological element among the factors causing inactivation.

Penicillin

Penicillin is stable at neutrality but decreases in stability as the pH is lowered (Fig. 14). This accords with previously published data (Florey, Chain, Heatley, Jennings, Sanders, Abraham & Florey, 1949), though it was rather more stable at pH 5.8 and 6.9 in our experiments than would have been expected from the earlier data. Sand had no effect on the rate of inactivation. The garden soil, whether fresh or partially sterile, rapidly inactivated penicillin, in spite of a pH favourable to stability. Inactivation was rapid in Wareham I and III, as would have been expected from the low pH. In Wareham II and IV it was surprisingly stable with but the slightest indication of biological inactivation (Fig. 15).

Streptomycin

Over the pH range 3.5–6.9, no activity was lost from buffer solutions of streptomycin over the period of the experiment (Fig. 16). This agrees with results quoted by Waksman (1949). Addition of sand to the less concentrated streptomycin solutions (40 units/ml) caused an immediate and complete loss of activity. At the lower pH levels it caused an immediate reduction in the activity of the more concentrated solution. This result is best explained by some form of adsorption. Siminoff & Gottlieb (1951) have shown that streptomycin is bound by clay minerals; apparently carefully washed sand is also sufficiently acidic to bind the streptomycin anion. This adsorption appears to be most marked in acid solutions.

Streptomycin was also adsorbed on all the soils (Fig. 17), and this resulted in immediate and complete inactivation of the 40 units/ml solution. In Wareham II and IV, once the adsorptive capacity of the soil was saturated, no further loss of activity took place. In the remaining soils the rate of inactivation, after the initial adsorption, was influenced by the presence of buffer. In Wareham I, streptomycin in water solution was inactivated slowly in sterile soil, rapidly in fresh soil; inactivation did not occur in fresh soil in buffer solution. A similar result was obtained in Wareham III, though here water solutions in fresh and partially sterile soils were inactivated equally rapidly. Inactivation was very rapid in garden soil, whether fresh or partially sterile.

Viridin

In buffer at pH 3.3 viridin (Fig. 18) is moderately stable; above pH 5.0 it is very rapidly inactivated. Detailed study of the data will show that in all cases inactivation follows a course equivalent to a first-order reaction and...
that the rate is directly related to the pH. Sand increases the rate of inactivation throughout the pH range tested. The stability of viridin in soil (Fig. 19) is primarily dependent on the pH of the soil. Inactivation is rapid in the neutral garden soil. Wareham II and IV accelerate inactivation to the same extent as sand; Wareham III has a rather more pronounced accelerating effect. In none of these soils is there any sign of biological inactivation. In fresh Wareham I soil, on the other hand, inactivation is much more rapid than in the same soil partially sterilized, in which inactivation is at the same rate as in sand.

CONCLUSIONS

These results show that the inactivation of antibiotics in soil may be the result of one or more distinct processes. These include: inactivation resulting from intrinsic instability of the antibiotic at the pH of the soil; inactivation as the result of microbiological activity; adsorption on some soil constituent, of which the clay colloids and humus particles are probably of greatest importance; some other kind of inactivation, probably chemical in nature, at present imperfectly understood.

The effect of soil pH on the rate of inactivation was most noticeable with the antibiotics albidin, frequentin, gliotoxin, penicillin and viridin.

Biological inactivation, expressed by more rapid inactivation in fresh than in partially sterile soil, occurred only in garden soil, Wareham I and Wareham III. These are the soils with the highest organic matter content and the most abundant microflora. Biological inactivation was most notable with the antibiotics griseofulvin, mycophenolic acid and patulin, but it was a contributory factor to the inactivation of several others.

Adsorption was an important factor in the inactivation of streptomycin solutions; it is noteworthy that carefully washed sand was capable of binding significant amounts of this basic antibiotic. The amount adsorbed was related to the pH of the solution. Adsorption was a less important factor in the inactivation of the acidic or neutral fungal antibiotics.

The rapid inactivation of gladiolic acid, penicillin and streptomycin in partially sterile garden soil, and some other similar observations, under pH conditions favourable to stability, suggests strongly that some other form of inactivation may occur. It is conceivable that the antibiotic in such cases is reacting chemically with a soil constituent, or that some soil constituent acts as a catalyst.

Finally, it may be noted that those antibiotics produced by fungi isolated from the acid Bagshot Sand soils were more stable in the acid Wareham series of soils than in the neutral garden soil. While no direct evidence is presented here that any of these antibiotics are produced naturally in soil, at least it may be said that all show a sufficient degree of stability, in some of the soils at least, for them to have a significant biological effect if produced.

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REFERENCES


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