Specific and non-specific resistance to aminoglycosides in *Escherichia coli*

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**SUMMARY** The turbidimetric responses of a strain of *Escherichia coli* K12 to gentamicin and tobramycin were investigated. Both agents showed antibacterial activity below the conventionally measured minimum inhibitory concentration, but exposure to such subinhibitory concentrations of either agent generated a bacterial population which was able to grow in previously inhibitory concentrations at a rate equivalent to that of the parent culture. The increase in resistance was non-specific in that both aminoglycosides were equally affected, and was unstable on multiple passage in drug-free broth.

The response to tobramycin was unaffected by the presence of an R factor conferring gentamicin resistance, but exposure of the R factor bearing strain to gentamicin caused a concomitant increase in the resistance to tobramycin, apparently by a non-specific adaptive mechanism similar to that observed with the parent strain. It is suggested that prior treatment of a gentamicin-resistant organism with gentamicin (as may occur during blind therapy) may adversely affect the subsequent response to other aminoglycosides.

Gentamicin is now widely used as the treatment of choice for many serious Gram-negative infections. Tobramycin has a similar spectrum of activity to gentamicin (Benveniste and Davies, 1973; Andrews et al., 1973) but is unaffected by some of the bacterial enzymes which inactivate gentamicin. The difference between an adequate therapeutic dosage and toxic dosage of these drugs is small, especially in patients with renal failure (Noone et al., 1974; Mawer et al., 1974). It is therefore desirable that the most efficient way of using them be established.

To gain a better understanding of the interaction between drug and bacterium we have investigated the response of a strain of *Escherichia coli* K12 to various concentrations of gentamicin and tobramycin by continuous turbidimetric monitoring and have attempted to assess the effect of possession of an R factor conferring resistance to gentamicin alone on the responses to gentamicin and tobramycin.

**Material and methods**

*Escherichia coli* K12 strain J53 (F−, pro−, met−); Clowes and Hayes, 1968) and a derivative of this strain, J53 (RS28) carrying the R factor RS28, were kindly provided by Dr Naomi Datta.

The R factor RS28 was originally derived from a strain of *Serratia marcescens* and belongs to the compatibility group M, which has an unusually wide host range (Hedges et al., 1975). RS28 confers resistance to ampicillin, streptomycin, tetracycline, chloramphenicol, and gentamicin.

Growth medium was the 'complete' medium of Greenwood and O'Grady (1973) modified by replacing the phosphate buffer by tris (hydroxymethyl) aminomethane (TRIS) in order to avoid interference with aminoglycoside activity by phosphates. The medium (CMT broth) has the following constituents per litre: glucose 5 g, casitone pancreatic digest (Difco) 10 g, yeast extract (Difco) 5 g, NaCl 5 g, TRIS buffer 6 g adjusted to pH 7.2 with HCl.

Gentamicin was obtained from British Schering Ltd, and tobramycin from Eli Lilly & Co Ltd, both as pure powders. Stock solutions were prepared in sterile distilled water from which appropriate working dilutions were made.

Minimum inhibitory concentrations (MICs) of gentamicin and tobramycin for the test strains were estimated by a conventional tube dilution method using CMT broth.
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Continuous turbidimetric measurements were obtained using the 12-channel opacity monitoring system described by Mackintosh et al. (1973). Six tubes containing 19 ml of CMT broth and 1 ml of an appropriate dilution of antibiotic or sterile distilled water (drug-free control) were seeded from an overnight broth culture to achieve an inoculum of ca 10⁶ organisms per ml. At the end of each experiment subcultures onto solid media were made to check that growth in antibiotic was not due to contamination. Adaptation was studied by seeding a fresh series of antibiotic-containing tubes with bacteria growing in the presence of peri-MIC concentrations of antibiotic in a previous experiment.

The stability of induced resistance was tested by serial daily subculture of adapted strains in antibiotic-free CMT broth, the turbidimetric response to graded antibiotic concentrations being investigated after 1, 7, 12, 18, and 24 subcultures.

Results

The MICs of gentamicin and tobramycin for E. coli K12 J53 were 4 and 2 μg per ml and for J53 (RS28) were 32 and 4 μg per ml, respectively.

Response of E. coli K12 J53

Continuous turbidimetric records of E. coli K12 J53 grown in the presence of gentamicin and tobramycin are shown in Fig. 1a and b.

In the presence of 0·25 μg gentamicin per ml or 0·5 μg tobramycin per ml the growth curve closely followed that of the control. In the presence of 0·5 μg gentamicin per ml (MIC/8) or 1 μg tobramycin per ml (MIC/2) growth was suppressed for a time, although once growth occurred, the slope of the curve was the same as for the drug-free control. Thus the minimum antibacterial concentration (MAC), the lowest concentration of antibiotic achieving an observable antibacterial effect in this system, was 0·5 μg gentamicin per ml and 1 μg tobramycin per ml.

As the concentration of antibiotic was further increased, the time taken for growth to attain a level of 50% of maximum opacity (T₅₀) increased correspondingly until a concentration (corresponding to the MIC) was reached at which growth was suppressed for the 22-hour period of observation. Subcultures onto solid media showed that delayed growth was not due to contamination, although the colonies of the previously suppressed cultures were much more variable in size and shape than those of the original strain.

Adaptation to Increased Resistance

The culture which grew in 2 μg gentamicin per ml (T₅₀ = 21 h, Fig. 1a) was used as the inoculum for a fresh series of gentamicin-containing CMT broths. In broth containing 2 μg gentamicin per ml (that is, the concentration to which the culture had previously been exposed) this culture grew as rapidly as it and its parent J53 did in antibiotic-free medium (T₅₀ = 3·8 h) (Fig. 2). Gentamicin, 4 μg per ml (the MIC of the original strain), delayed the growth of this culture (T₅₀ = 6 h) and higher concentrations (≥ 8 μg/ml) suppressed growth for the 22-hour period of observation. Thus by a single passage in gentamicin the MAC increased to equal the original MIC, an eight-fold increase. By this means a culture was derived which grew in 4 μg gentamicin per ml but at a reduced growth rate (T₅₀ = 7·3 h).

Further adaptation could be slowly achieved by serial passage in increasing concentrations of antibiotic. The ability to grow in increasing concentra-

Fig. 1 Continuous opacity records of E. coli K12 J53 in the presence of various concentrations (μg per ml) of gentamicin. The inoculum was derived from a culture growing in 4 μg gentamicin per ml in a previous experiment.
tions of gentamicin was accompanied by increasingly irregular colonial morphology on solid media. Virtually identical results were obtained using tobramycin.

CROSS RESISTANCE OF ADAPTED STRAINS
The strain of *E. coli* K12 J53 adapted to grow in 4 μg gentamicin per ml was also tested against tobramycin. An identical response was observed, the MAC now equalling the original MIC for tobramycin.

STABILITY OF ADAPTIVE RESISTANCE
The strain of *E. coli* K12 J53 adapted to grow in 4 μg gentamicin per ml was serially passaged through antibiotic-free CMT broth. An immediate reversion to greater susceptibility to gentamicin was not observed. However, a slow progression towards greater susceptibility was observed, but the process of reversion was not quite complete even after 24 passages (Figs. 1a, 2, and 3).

EFFECT OF R FACTOR RS 28

Turbidimetric response
The rate of growth of J53 (RS28) in antibiotic-free broth was the same as that of its parent, J53. In the presence of 2 μg gentamicin per ml J53 (RS28) grew uninterruptedly, but with each increase in concentration above the MAC, 4 μg gentamicin per ml, there was an increase in the time taken for the cultures to reach 50% opacity (Tso) (Fig. 4). This pattern of response to different concentrations of gentamicin was similar to that shown by the parent J53 to eight-fold lower concentrations of gentamicin. The MIC of gentamicin for J53 (RS28) was greater than 32 μg per ml. However, the possession of the R factor appeared to have little effect upon the sensitivity to tobramycin; the MAC was 0·5 μg per ml, and the MIC remained unchanged.

**Adaptation to increased resistance**
Cultures of J53 (RS28) growing after initial suppression of growth showed a similar pattern of adaptation to that observed with the parent strain J53. The MAC of a culture initially suppressed by 32 μg gentamicin per ml, but growing in this concentration during the overnight incubation period, increased to 64 μg gentamicin per ml, and the MIC to >128 μg gentamicin per ml. Furthermore, the increased resistance to gentamicin was reflected in a concomitant increase in resistance to tobramycin, the MAC and MIC being raised to 4 and 8 μg tobramycin per ml, respectively.

Discussion

Although the measurement of MIC and MBC values is generally regarded as the definitive test of the in vitro activity of antimicrobial agents, many factors may influence the results obtained. For aminoglycosides the result may be affected by various components of the medium, its pH, and the method of performing the test. What is not generally recognised is that the conventionally determined tube dilution MIC value represents a measurement taken at an arbitrary point in time during a continuing series of events. Continuous turbidimetric monitoring of cultures exposed to aminoglycosides as they would be in a conventional tube dilution test clearly shows this. An eight-fold difference in MIC was observed between a reading taken after six hours' incubation, when the control tube was fully grown, and one taken after 24 hours' incubation (Fig. 1a). Which of these results is the correct one in clinical terms is a matter of speculation, but as the MIC end-point appears to be controlled by the ability of the culture to adapt to otherwise inhibitory concentrations of the drug, the MAC is at least likely to be more reproducible and less affected by the density of the bacterial inoculum.

Bacteria growing at concentrations of antibiotic between the MAC as defined above and MIC as measured by a conventional tube dilution technique form a very small proportion of the original bacterial
population (Waterworth, 1972), but under the selective pressure of antibiotic levels approaching the MIC they quickly become the majority population. In the present study *E. coli* K12 J53 was easily trained to grow in concentrations of gentamicin and tobramycin close to those to which bacteria may be exposed *in vivo* for a large part of conventional therapeutic regimes. Although peak serum levels may exceed the level of adaptation which is easily achieved, such levels are attained only transiently. It is generally believed that strains adapted to resistance in this way are unimportant clinically owing to loss of virulence. Weinstein *et al.* (1971) demonstrated loss of virulence for mice in strains of *Pseudomonas aeruginosa* which had been adapted to quite high gentamicin levels. However, at this level of adaptation these strains exhibited (a phenomenon that we confirm) grossly abnormal growth characteristics. Continuous turbidimetric monitoring of *E. coli* K12 J53 showed that no measurable change in the rate of growth, as measured by the ability to achieve a 50% opacity level (T50), occurred at low levels of adaptation with this organism and that slightly increased resistance was not a severe handicap since after 24 subcultures in antibiotic-free broth a population adapted to a moderate increase in resistance had not completely returned to its original fully sensitive state. An important feature of this type of resistance is that it appears to extend to other aminoglycosides (Houang and Greenwood, 1977). It is possible that those ‘coliforms’ frequently isolated from superficial lesions which have a decreased sensitivity to all aminoglycosides are partially adapted strains.

The effects of an R factor (RS28) conferring resistance to gentamicin but not to tobramycin by the production of an inactivating enzyme are different from those produced by adaptation (Figs 4 and 2). In this case delayed growth in concentrations of gentamicin that are initially inhibitory may simply reflect the time taken for the gentamicin to be enzymically inactivated. However, re-exposure of the emergent bacterial population to gentamicin revealed that adaptation was contributing to the overall effect. Furthermore, the sensitivity to tobramycin was also substantially decreased by previous exposure to gentamicin, although possession of the R factor did not itself affect the intrinsic susceptibility of the organism to tobramycin.

The demonstration that exposure of a gentamicin-resistant organism to gentamicin can also decrease the susceptibility of the organism to another aminoglycoside suggests that blind therapy with one aminoglycoside may compromise the success of subsequent treatment with another, even if the infecting organism proves to be resistant to the first drug by an enzymic mechanism which does not affect the second. If these *in vitro* considerations are substantiated clinically, it may be that blind therapy with aminoglycosides might have to be used more circumspectly than it is at present, at least in those centres where gentamicin resistance is a problem.

This work formed part of a project undertaken by one of us (SLM) during a course of study leading to the degree of MSc in MedicalMicrobiology, University of London. We are indebted to Dr N. Datta for kindly providing the cultures of *E. coli* K12 J53 used in these experiments.

References


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