Selection of R- and S-phase Colonies from Dimorphic *Alcaligenes odorans* var. *viridans*, and Their Isolation in Pure Culture

By B. BRZIN

Department of Microbiology, Medical Faculty, Ljubljana, Yugoslavia

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*Alcaligenes odorans* var. *viridans* was first described by Mitchell & Clarke (1965), who observed colonial pleomorphism of the bacteria. They stated that neither type of colony bred true on subculture. Many strains were later isolated and studied by Brzin (1971), who found that fringe formation was occasionally observed in young colonies even though, in general, only older colonies, or, more often, only parts of them, were surrounded by a fringe. This early fringe formation was suspected to be due to the beginning of the R-phase dissociation of colonies of *A. odorans* var. *viridans*. Therefore selection of R- as well as S-phase colonies and their isolation in pure culture was attempted simply by successive subculturing of the bacteria from typical parts of colonies. The two phases of *A. odorans* var. *viridans* were isolated in pure culture, tested for stability and their physiological and biochemical characteristics were compared.

METHODS

Strains of *Alcaligenes odorans* var. *viridans* were subcultured on 5 % bovine blood agar every 48 h. For the first 24 h they were incubated at 37 °C and later at room temperature (about 22 °C). The largest and most pronounced fringe was then picked at its extreme border, transferred to a fresh medium, and incubated for 24 h at 37 °C and subsequently at room temperature for another 24 h. The bacterial growth at the extreme border of the fringe around the new colonies was in this way serially subcultured, to give a pure culture of the R-type colony.

In a similar way, a pure culture of the S-type of *Alcaligenes odorans* var. *viridans* was obtained by successively subculturing the colonies which remained completely rounded and smooth even after prolonged cultivation at room temperature.

RESULTS AND DISCUSSION

In this simple way, the R- as well as the S-variant of *Alcaligenes odorans* var. *viridans* became more and more pronounced until, finally, upon continued selection of typical R- and S-type growths, pure cultures of R- and S-colonies were isolated (Fig. 1 a, b).

In an effort to help the differentiation and isolation of the two phases of *Alcaligenes odorans* var. *viridans*, various media and other growth conditions were used, but with no significant effect.

Owing to its punctiform, greening growth and its ability to grow in salt broth, the S-colony type of *Alcaligenes odorans* var. *viridans* was even more similar to *Enterococcus* than was the original dimorphic culture; whereas the R-type strongly resembled *Pseudomonas aeruginosa*. 
During the first month after the isolation of cultures of the R- and S-type, the cultures were still unstable. When the bacterial growth to be transferred to fresh medium was picked at random, i.e. not at the extreme border of the fringe or from a completely rounded, smooth colony, the new culture soon tended to revert to a mixed one. After continued subcultivation the R- as well as the S-type of growth became stable. Cultures were influenced neither by varying physical factors (incubation temperature, degree of dryness of the medium)
nor varying chemical factors (composition and pH of the medium, type and percentage of blood in the medium). The physiological and biochemical characteristics of the two colony types were found to be identical. In the antibiogramme test no difference was found between the reaction of either type to antibiotics or chemotherapeutics. Moreover, there was no difference between the two colony types in the characteristic features observed by Brzin (1971), namely their ability to grow in salt broth (6.5%), and the tendency of the bacteria to elongate in this medium.

Since with some bacteria the rough-smooth colony variation is a reflection of bacterial size (Grula, 1960a, b), the size of bacteria of each colony form was compared repeatedly and under various conditions. No significant difference in microscopic morphology was found between the R- and S-variant. In this case the rough-smooth colony variation did not seem to be a reflection of bacteria size or of the presence or absence of a capsule. The observed change in colonial properties of these bacteria was probably due to spontaneous mutation involving the surface character. No cause-and-effect relationship was found between the chemical composition, pH or state of dryness of the media, and the readiness of these bacteria either to dissociate or to be isolated in a pure and stable culture of R- or S-type.

A change in type of growth in fluid media is sometimes coupled with aberrant bacterial morphology when culture conditions are not optimal. For example, the flocculent growth of Arthrobacter induced by too low a vitamin B₁₂ supply was found to be associated with abnormal bacterial morphology (Chaplin & Lochhead, 1956). It is therefore possible that the absence of any significant difference in microscopic morphology between the R- and S-variants was the reason why no great difference between the mode of growth in fluid media of each variant was observed. The R-variant grew more diffusely than the S-variant in nutrient broth, but showed only a weak tendency to creep along the inner side of the vessel and only occasionally formed a pellicle after prolonged incubation at room temperature.

Both colony-types of Alcaligenes odorans var. viridans are deposited in the National Collection of Industrial Bacteria (NCIB), held at the Ministry of Agriculture Fisheries and Food, Torry Research Station, 135 Abbey Road, Aberdeen, AB9 8DG; Scotland.

REFERENCES


