
The Sporulation of *Bacillus sphaericus* stimulated by Association with other Bacteria: an Effect of Carbon Dioxide

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SUMMARY: Sporulation of *Bacillus sphaericus*, NCTC 7582, in a complex medium was strongly stimulated by growth in mixed culture with a sporing strain of *B. cereus* and with sporogenous and asporogenous strains of *B. subtilis*, NCTC 85. There was a similar, but less pronounced, effect with a variety of Gram-positive and Gram-negative organisms. The degree of sporulation of *B. sphaericus* in these mixed cultures depended on (a) an external stimulus from the medium and (b) the previous cultural history of the *B. sphaericus* inoculum upon which depended its ability to respond to the external stimulus. Sporulation of *B. sphaericus* in pure culture was strongly stimulated by bicarbonate and ketoglutarate, suggesting that increased carbon dioxide production was the stimulating factor in mixed cultures.

The stimulatory effect of one organism upon the growth of another has been observed in bacteria and in fungi (e.g. Grassberger, 1897; Bacon, Burrows & Yates, 1951; Hawker, 1947). In fungi there are examples of stimulation of both growth and sporulation in mixed cultures (Asthana & Hawker, 1936; Hawker, 1939). This report describes a similar phenomenon in the genus *Bacillus* and suggests a mechanism for the effect.

ORGANISMS AND METHODS

A culture of *Bacillus sphaericus*, NCTC 7582, grown for 24 hr. at 37° on casein hydrolysate yeast extract (CCY) agar (Gladstone & Fildes, 1940) was suspended in 50 ml. glucose serum broth (Fry & Greaves, 1951) and freeze-dried in 5 ml. batches. Starting with one of these freeze-dried batches, a series of sporulation tests was made, during which time the organism was maintained by subculturing every 48 hr. on CCY agar, and on potato-extract agar (Robinow, 1951) enriched with 1/10 vol. of CCY medium (Pot. CCY).

The strongly sporing strain of *Bacillus cereus* isolated from air and the sporogenous and asporogenous strains of *Bacillus subtilis*, NCTC 85, were maintained on CCY agar. The other organisms used were subcultured at least twice in Pot. CCY medium before the mixed culture tests.

For the sporulation tests, which were set up in duplicate, 10 ml. batches of liquid Pot. CCY medium in 50 ml. conical flasks were inoculated with 0.5 ml. of a just visibly turbid suspension of organisms and shaken at 37° for periods up to 36 hr.

RESULTS

There was rapid growth and remarkably vigorous sporulation of *Bacillus cereus* and *B. subtilis* in the test medium, which was selected for this reason. The first signs of sporulation appeared at 14 hr., and at 17 hr. almost every
Sporulation of B. sphaericus

A cell contained a well-developed spore. B. sphaericus taken from the first CCY agar subcultures showed approximately 1–5% sporulation after 36 hr. At 18 hr, 20–30% of the cells appeared to have reached a half-way stage of sporulation, but disintegrated before the spore had properly developed. After about twenty subcultures on CCY agar, although growth in the test medium was not diminished, the sporing ability of the organism became almost negligible, until finally no sign of sporulation could be detected. During this time the effect of inoculating test flasks with both B. cereus and B. sphaericus was also studied. The first B. sphaericus subcultures, with some tendency to sporulate when grown alone, showed almost complete sporulation after 18 hr. growth with B. cereus (Pl. 1, figs. 1, 2). Later subcultures, which showed no sign of sporulation when grown alone, were stimulated to produce pre-spores, i.e. refractile non-staining areas in the presence of B. cereus, and then appeared to disintegrate. After further subcultures on CCY agar, B. sphaericus lost its ability to respond to the B. cereus stimulus.

When an original freeze-dried stock culture of Bacillus sphaericus was maintained by subculture on Pot. CCY agar, there was a marked increase of sporing ability in pure culture. Similarly, cultures which had become asporogenous after repeated subculture on CCY agar, sometimes regained their ability to sporulate after approximately twenty subcultures on Pot. CCY agar. Thus, of four such asporogenous cultures, two regained their original sporing ability completely. The other two remained asporogenous but produced cells containing large granules staining black by the Albert (1921) method and purplish pink with methylene blue and toluidine blue. Some of the granules appeared to be free. Examined in wet preparation, the granules were less refractile than developing spores from which they could readily be distinguished. The phosphorus:nitrogen ratio in washed cells from a 'granular' culture was 0·37 compared with 0·26 in a culture not containing granules. The granules therefore probably contained volutin, i.e. polymetaphosphate (Wiaime, 1947).

Sporogenous cultures of Bacillus sphaericus grown for 16 hr. in Pot. CCY medium were centrifuged and the cells resuspended in the medium from 9 and 14 hr. cultures of B. cereus. After 5 hr. shaking there was only poor sporulation, but large purple-pink granules appeared in most cells (Pl. 1, figs. 4, 5). The appearance of these resuspended cultures was similar to that of the asporogenous granular cultures just described (Pl. 1, fig. 3). After 24 hr. shaking, the resuspended culture consisted almost entirely of non-sporulating cells free from large granules. Similar results were obtained with 10 hr. cultures of B. sphaericus.

A slight stimulation of Bacillus sphaericus sporulation was observed when the organism was grown in a cellophan tube immersed in a shaken growing culture of B. cereus, but this result was not always repeatable.

The effect of growing Bacillus sphaericus in mixed culture with other organisms was then studied. Both the sporogenous and asporogenous strains of B. subtilis stimulated rapid and almost complete sporulation of the feebly sporing B. sphaericus, and increased the 'granulation' of the asporogenous strain. Staphylococcus aureus (Oxford strain), Corynebacterium xerosis
NCTC 7904, and laboratory strains of *Chromobacterium prodigiosum* (*Serratia marcescens*) and a *Leuconostoc* species had a similar but less pronounced effect. *Escherichia coli*, NCTC 1100, and a laboratory strain of *Pseudomonas aeruginosa* (*pyocyanea*) grew well in mixed culture with *Bacillus sphaericus* but had no apparent effect on its sporulation.

*Bacillus sphaericus* cells resuspended in culture fluid from a stimulating organism showed very poor sporulation (see above). It appeared therefore that the stimulus in mixed culture was produced during associated growth of the stimulating organism, and was probably due to some non-specific effect such as (i) a more rapid exhaustion of nutrients in mixed culture, or (ii) a reduction of dissolved oxygen or an increase of carbon dioxide concentration in the medium to some critical level. The effect of adding 0.04 M-sodium bicarbonate to the medium as a means of increasing carbon dioxide concentration was next tested. Control flasks were adjusted to the same pH as those to which bicarbonate was added, i.e. pH 8.2. At 18 hr. there was very poor sporulation in the controls, but almost complete sporulation in the presence of bicarbonate (Pl. 1, figs. 6, 7). Final pH values were 9.0 and 9.2 respectively. There was equally strong stimulation of sporulation in the presence of 0.04 M-ketoglutarate (Pl. 1, figs. 8, 9) with initial and final pH values of 7.6 and 8.7 both in the controls and ketoglutarate media. Although the other possibilities mentioned above were not explored, it seemed highly probable that stimulation of *B. sphaericus* sporulation in mixed culture was due to increased carbon dioxide concentration in the medium.

**DISCUSSION**

Under the conditions described it appeared that the degree of sporulation of *Bacillus sphaericus* depended on (a) a stimulus from the medium which could be intensified by growth in mixed culture with a variety of bacteria, and (b) an intracellular factor which controlled the ability of the cell to respond to the outside stimulus. Although a high CO₂ concentration in the medium may not have been of prime importance for the initiation of sporulation, its presence appeared to be essential in order to make the process rapid and complete. This may hold for sporing organisms in general, since it has been repeatedly observed that rapid and complete sporulation occurs after a period of vigorous growth, i.e. high CO₂ production. It is specially interesting in this connexion that Cantino (1952) showed that bicarbonate stimulated the production of thick-walled resistant sporangia in the mould *Blastocladiella*. The function of CO₂ as a sporulation stimulant in the genus *Bacillus* may be connected with its possible stimulatory effect on dipicolinic acid synthesis (Powell, 1953). It seems likely, especially since conversion of lysine to pipelic acid has been demonstrated (Rothstein & Miller, 1958; Lowy, 1958) that dipicolinic acid is derived from αε-diaminopimelic acid by deamination followed by ring closure and dehydrogenation. It is possible that CO₂, by inhibiting the decarboxylation of αε-diaminopimelic acid (an effect already observed by Hoare & Work, 1954, personal communication), might allow the
deamination reaction and subsequent build-up of dipicolinic acid to proceed more rapidly.

In this investigation we obtained an asporogenous culture of Bacillus sphaericus by repeated subculture of the organism on a relatively rich medium (CCY). The sporulating power of this culture was in some cases restored by repeated transfer to a less nutritious medium (Pot. CCY). The sensitivity, i.e. the degree of response of a vegetative cell of B. sphaericus to the external stimulus, may depend on its ability to divert one or more metabolic pathways in ‘abnormal’ directions, e.g. for the synthesis of dipicolinic acid, and it is possible that this ability might be enhanced or lowered by adaptive processes on different media, or by variation (Brunstetter & Magoon, 1932; Schmidt, 1950). We found that there was often an increased production of volutin by the asporogenous strain of B. sphaericus under conditions which stimulated sporulation of the sporogenous strain. This was not due to the development of acid conditions in the culture (cf. Duguid, Smith & Wilkinson, 1954). It may represent an incomplete response to the external stimulus, i.e. a derangement of metabolism in cells whose metabolic path was blocked as a result of conditions in the medium and could not be diverted in the direction of sporulation. There was also volutin production in sporogenous cells, especially after resuspension in culture medium from a stimulating organism. This may again represent an incomplete response in cells which were capable of sporulation but were receiving an insufficiently sustained stimulus from the medium to drive the sporulation process to completion.

We wish to thank Mrs W. Fryer for technical assistance and Mr R. E. Strange for nitrogen and phosphorus analyses. Acknowledgement is made to the Chief Scientist, Ministry of Supply, for permission to publish this paper.

REFERENCES


EXPLANATION OF PLATE

All films heat-fixed and stained with methylene blue. Magnification ×1500.

Fig. 1. 18 hr. sporogenous B. sphaericus in pure culture.
Fig. 2. 18 hr. sporogenous B. sphaericus in mixed culture with B. cereus.
Fig. 3. 18 hr. asporogenous B. sphaericus subcultured repeatedly on Pot. CCY agar, showing granule production in test medium at pH 8·1.
Fig. 4. 21 hr. sporogenous B. sphaericus.
Fig. 5. 16 hr. sporogenous B. sphaericus resuspended for 5 hr. in B. cereus medium.
Fig. 6. 18 hr. sporogenous B. sphaericus at pH 9·0.
Fig. 7. 18 hr. sporogenous B. sphaericus sporulating with 0·04M-bicarbonate at pH 9·2.
Fig. 8. 18 hr. sporogenous B. sphaericus at pH 8·7.
Fig. 9. 18 hr. sporogenous B. sphaericus sporulating with 0·04M-ketoglutarate at pH 8·7.

(Received 8 December 1954)

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