Inactivation of Bacterial Spores by Hydrostatic Pressure

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SUMMARY

Spores of various species of the genera Bacillus and Clostridium were inactivated by hydrostatic pressures up to 8000 atmospheres. Inactivation was a function of holding time at pressure rather than of the compression and decompression stages. Inactivation generally proceeded more rapidly at high than at low temperatures; below about 50° there was a well defined optimum pressure for inactivation, but above about 50° an increase in pressure up to 8000 atmospheres caused progressively more inactivation.

Inactivation was decreased at extremes of pH value and by high ionic strength solutions. A proportion of the spores pressurized under certain conditions became heat-sensitive. These observations, and the chemical, phase-contrast, and electron-microscopic changes seen in pressurized spores, suggested that pressure caused inactivation of spores by first initiating germination and then inactivating the germinated forms.

INTRODUCTION

Hydrostatic pressure above 1000 atmospheres (atm.) causes a rapid inactivation of many vegetative bacteria (Hite, 1899; Hite, Giddings & Weakley, 1914; Heden, 1964) whereas spores are more resistant and may survive pressures above 12,000 atm. (Larson, Hartzell & Diehl, 1918; Basset & Macheboeuf, 1932). Recently we noticed, however, that under certain conditions inactivation of spores occurred more rapidly at lower hydrostatic pressures than at higher pressures. This observation was surprising, and was difficult to reconcile with the expected effect of a physical treatment on a biological system: an increase in the intensity of the treatment ought to result in an increase in the degree of inactivation of the bacteria (cf. heat, ultraviolet and ionizing radiation, electric shock, ballistic disintegration; see Roberts & Hitchins, 1969). This anomalous effect of pressure led us to study and define the effects of environmental variables on pressure inactivation of spores more closely, with a view to explaining, first, how spores are inactivated by pressure and, secondly, why lower pressures can cause greater inactivation than higher pressures.

METHODS

Organisms. Bacteria used were Bacillus coagulans NCTC 3991 and B. cereus NCTC 8035 (National Collection of Type Cultures, Colindale), B. cereus strains T and px, B. polymyxa M1, B. subtilis var. niger (syn. globigii), B. subtilis A (rough strain; Edwards, Busta & Speck, 1965) and Clostridium sporogenes PA 3679.

Production of spores. Spores of the aerobes were grown on potato yeast-extract

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Glucose agar (Gould & Ordal, 1968) at 30° (Bacillus cereus strains) or 37° (B. coagulans, B. polymyxa and B. globigii) or on the Fortified Nutrient Agar of Edwards et al. (1965) at 45° (B. subtilis A). Spores of Clostridium sporogenes were grown at 37° in the liquid Reinforced Clostridial Medium (Oxoid) of Gibbs & Hirsch (1956). When sporulation and lysis of sporangia were complete the spores were washed six times with cold distilled water (using a centrifuge) and stored at 4° in water at a concentration of approximately 20 mg. dry wt spores/ml. Suspensions of ‘superdormant’ spores of B. cereus PX and B. polymyxa (Gould, Jones & Wrighton, 1968) were prepared as follows: Spores were activated by heating at 70° for 30 min., then incubated at a concentration of about 10⁸ spores/ml. for 1 hr in yeast glucose broth at 37°. The suspension was then heated at 70° for 30 min. to inactivate germinated spores whilst leaving viable the superdormant forms. The spores were finally recovered, washed and stored as above. Mixed spores from soil were prepared by extracting 10 kg. portions of soil with 1 l. portions of cold distilled water, filtering through glass wool to remove coarse debris and concentrating the spores by repeated differential centrifugation. The final suspension was heated at 70° for 30 min. to inactivate non-spore forms and stored in water, as above.

Application of pressure. Suspensions (2 ml.) were enclosed in sachets made from 1 in. broad polythene Layflat Tubing (Transatlantic Plastics Ltd., 45 Victoria Road, Surbiton, Surrey), using an impulse heat sealer (A. H. Bland Ltd., Winchelsea Road, Harlesden, London N.W.10). The final seal was made through the suspension in order to exclude air bubbles. The sachets were immersed in a container of water or oil, to which the pressure was transmitted by hydraulic oil.

The high pressure intensifiers were designed following conventional techniques (Manning, 1963), and pressures up to 8000 atm. could be reached. Pressure was measured by a manganin gauge, and temperature close to the sachets by a mineral-insulated metal-sheathed thermocouple. Temperature was controlled by circulating heated glycol from a thermostated bath through a jacket on the outside of the pressure cylinder. Pressure was normally raised in less than half a minute and lowered in less than 5 sec., unless otherwise stated. Adiabatic heating of the samples amounted to approximately 3° per 1000 atm.; the temperature transient decayed in 10 min.

Radiation. Suspensions (1 ml.) were sealed into glass ampoules and subjected at ambient temperature to γ-radiation from a Co 60 source at a dose rate of about 0·6 mrad./hr.

Estimation of viability. Spores surviving the various treatments were enumerated by poured plate viable counts using nutrient agar, and incubation temperatures as listed above. Heat-sensitive forms were inactivated by heating samples at 70° for 30 min. before plating as indicated in the Results section.

Chemical estimations. Calcium was estimated by flame photometry on samples of bacterial pellets dry-ashed (500°, 16 hr) and extracted with 0·1 N-HCl. Dipicolinic acid in bacterial pellets was assayed by the method of Janssen, Lund & Anderson (1958). For estimation of hexosamine, bacterial pellets were first hydrolysed (6 N-HCl, 5 hr, 100°) in sealed evacuated ampoules. Hexosamine was then determined by the Boas (1953) modification of the Elson and Morgan reaction using sodium carbonate blanks to lower non-specific colour due to sugars and amines, as suggested by Immers & Vasseur (1950); glucosamine hydrochloride was the standard (Hamilton & Stubbs, 1967).
Inactivation of spores by pressure

Electron microscopy. Bacterial suspensions were centrifuged and the pellets re-suspended, for fixation, in potassium permanganate (2%, w/v) at 22° for 90 min. (spores) or 60 min. (germinated spores) (Mollenhauer, 1959). The 'fixed' pellets were washed at least six times by centrifugation prior to embedding in Epon 812, wet-sectioned and stained with uranyl acetate (1%, w/v) for 10 min. and examined in a JEM 6S electron microscope.

RESULTS

Lethal effect of hydrostatic pressure

When suspensions of spores of different species were subjected to different pressures for a constant time at 20° the pattern of survivors (Table 1) showed that pressure had inactivated the spores, but the greatest inactivation was not caused by the highest pressure. Intermediate pressures, i.e. between about 1000 and 3000 atm. (depending on the organism), were most lethal.

Table 1. Inactivation of spores of different species by hydrostatic pressure

<table>
<thead>
<tr>
<th>Organism</th>
<th>Survivors (%)* following pressurization† at 20° for 1 hr at (atm.)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1000</td>
</tr>
<tr>
<td>Bacillus cereus NCTC 8035</td>
<td>18</td>
</tr>
<tr>
<td>B. cereus PX</td>
<td>98</td>
</tr>
<tr>
<td>B. subtilis var. niger (syn. globigii)</td>
<td>100</td>
</tr>
<tr>
<td>B. polymyxa</td>
<td>100</td>
</tr>
</tbody>
</table>

* Survivors were enumerated by poured plate viable counts.
† Spores (c. 10⁶/ml.) were suspended in sodium phosphate buffer (0.1 M, pH 8.0) during pressurization.
‡ Not tested.

Table 2. Comparison of continuous and discontinuous application of pressure on viability of spores

<table>
<thead>
<tr>
<th>Organism</th>
<th>Application of pressure†</th>
<th>Survivors (%)* following pressurization</th>
<th>Unheated spores</th>
<th>Heated spores‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacillus subtilis var. niger (syn. globigii)</td>
<td>Continuous</td>
<td>3.5</td>
<td>1.6</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Discontinuous</td>
<td>7.7</td>
<td>4.6</td>
<td></td>
</tr>
<tr>
<td>B. cereus PX</td>
<td>Continuous</td>
<td>1.6</td>
<td>0.30</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Discontinuous</td>
<td>0.45</td>
<td>0.30</td>
<td></td>
</tr>
</tbody>
</table>

* Spores (c. 10⁶/ml.) were suspended in sodium phosphate buffer (0.1 M, pH 8.0). Survivors were enumerated by poured plate viable counts.
† Pressure (2000 atm., 20°) was applied either continuously for 1 hr, or discontinuously as a series of twelve 5 min. exposures.
‡ Spores were heated for 30 min. at 70° following pressurization.

Spores in suspensions subjected to pressure for 1 hr (as in Table 1) were inactivated to about the same extent whether held under pressure continuously or discontinuously, i.e. compressed and decompressed twelve times, each treatment being for a period of 5 min. (Table 2). This, and other experiments, showed that the extent of the inactivation...
depended on the duration of pressure treatment and not on the compression or decompression steps.

Spore concentration did not affect the extent of the inactivation, e.g. the survivor levels of spores of *Bacillus coagulans* pressurized in sodium phosphate (0.1 M, pH 8.0) at 2000 atm., 45° for 1 hr, then heated at 70° for 1 1/2 hr were 0.13, 0.23, 0.21, 0.23 and 0.24 respectively for initial viable spore concentrations of 2.6 x 10^9, 2.7 x 10^8, 2.5 x 10^7, 2.5 x 10^6 and 2.6 x 10^5 per ml.

*Effect of temperature on the inactivation of spores by pressure*

Inactivation of spores by pressure was strongly influenced by temperature. Figure 1a summarizes the results of experiments with spores of *Bacillus coagulans*. Similar but less exhaustive experiments were performed with spores of *B. cereus* strains T, PX

![Fig. 1. Inactivation of spores of Bacillus coagulans at different pressures and temperatures. (a) Spores (c. 10^9/ml.) were heated at 70° for 30 min., then pressurized at the indicated temperatures and pressures for 30 min. The suspending medium was sodium phosphate buffer (0.1 M, pH 8.0). Survivors were enumerated by viable counts using poured plates. Temperatures of pressurization were: 25° (○); 35° (●); 45° (□); 55° (■); 65° (▽); 75° (△). (b) As Fig. 1a, but spore suspensions were additionally heated at 70° for 30 min. following pressurization in order to inactivate forms which had been heat-sensitized but not inactivated by pressure. (Arrows indicate that survivors were not detected and were therefore less than indicated by the symbol.)

and NCTC8035, *B. subtilis* var. niger (syn. globigii), *B. polymyxa* M1 and *Clostridium sporogenes* PA3679. The same qualitative pattern was seen with all these organisms. Four well-defined regions of response to pressure could be recognized (Fig. 1a): (1) a region of increased inactivation with increased pressure up to about 1000 atm.; (2) a tendency to plateau around 2000 to 3000 atm.; (3) a region of further increased
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Inactivation; (4) a region above about 4000 atm. where the inactivation decreased at temperatures below 55º but increased at higher temperatures.

When samples of spores from the experiment reported in Fig. 1a were heated at 70º for 30 min. following pressurization below 65º, the inactivation was increased (Fig. 1b); the increase was particularly evident in the lower temperature and pressure regions (compare Fig. 1a, b). Since the temperature of 70º (for 30 min.) was well below that which inactivates untreated spores, pressure, in addition to inactivating spores, increased the sensitivity of a proportion of the survivors to heat.

The storage temperature following pressure treatment was unimportant, except as indicated in Fig. 1a, b. For instance, the survivor levels of spores of Bacillus coagulans following pressurization at 1000 atm. (at 70º for 30 min.) and incubation for 1 hr at 0, 25, 37, 45, 55 and 65º were respectively 1·2, 1·2, 1·1, 1·1, 1·1 and 0·9%.

Kinetics of inactivation of spores by pressure

Figure 2 shows the rates of inactivation of spores of three species by pressure at 70º. No single response was apparent: semilog inactivation curves were typically concave, but with an initial rapid drop in numbers of survivors for Bacillus coagulans at the higher pressures (Fig. 2a); the rate of inactivation of the more pressure-resistant spores of B. subtilis A was slow but approximately exponential (Fig. 2b); spores of Clostridium sporogenes held at 3000 atm. were about 99·99% inactivated within a few minutes, but the surviving 0·01% of the population resisted further inactivation for at least 2 hr (Fig. 2c).

Appearance of spores

The optical density of spore suspension fell when the spores were inactivated, or merely sensitized to heat by pressure. On examination by phase-contrast microscopy,
spores which had been inactivated or sensitized to heat by pressure always had changed from bright to dark, as happens when spores germinate. Examination with the electron microscope revealed structural changes which were typical of the changes seen in germinating spores (Plate 1).

Chemical changes

The chemical changes that occur on germination were sought and found to occur when spores were subjected to pressure. Pressure caused spores to excrete calcium, dipicolinic acid and hexosamine-containing material (Table 3).

<table>
<thead>
<tr>
<th>Organism</th>
<th>Pressure* (atm.)</th>
<th>Time (hr)</th>
<th>Dipicolinic acid (%)</th>
<th>Calcium (%)</th>
<th>Hexosamine (%)</th>
<th>Phase-dark spores (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacillus cereus PX</td>
<td>2000</td>
<td>1</td>
<td>91.7</td>
<td>92</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>B. polymyxa M1</td>
<td>2000</td>
<td>1</td>
<td>86</td>
<td>69.5</td>
<td>48</td>
<td>--</td>
</tr>
<tr>
<td>B. subtilis var. niger (syn. globii)</td>
<td>1000</td>
<td>1/2</td>
<td>72.8</td>
<td>--</td>
<td>--</td>
<td>81</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1</td>
<td>74.2</td>
<td>--</td>
<td>--</td>
<td>83</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td>85</td>
<td>--</td>
<td>--</td>
<td>88</td>
</tr>
<tr>
<td></td>
<td>6000</td>
<td>1/2</td>
<td>87.3</td>
<td>--</td>
<td>--</td>
<td>97</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1</td>
<td>98.2</td>
<td>--</td>
<td>--</td>
<td>97</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td>91.7</td>
<td>--</td>
<td>--</td>
<td>96</td>
</tr>
</tbody>
</table>

* Temperature during pressurization was 20°C. Spores (c. 10⁷/ml.) were suspended in water.

Effect of heat and of γ-radiation prior to pressurization

When spores of some species were sublethally heated prior to pressurization, their sensitivity to pressure at the lower temperatures and pressures increased. However, spores pressurized at higher temperatures (e.g. 65°C) could be rendered more resistant to pressure by prior heating above about 70°C (Fig. 3). (The low survivor level at 90°C in Fig. 3 was due to inactivation by the high temperature alone.)

Exposure to γ-radiation sufficient to inactivate a proportion of the spores in a suspension caused an increase in the pressure sensitivity of the survivors (Table 4).

Inactivation of spores by pressure at different pH values

Figure 4 shows the effect on survivor levels of pressurizing spores in various buffers at different pH values. (Although pressure will change the pH value of buffer solutions, the changes in these experiments were estimated not to exceed about 0.4 pH units; Distèche, 1959.) The nature of the buffer seemed unimportant, and spores were inactivated by pressure over a wide pH range. However, the inactivation was greatest near neutrality and was definitely lowered at extreme pH values.

Decrease in pressure inactivation of spores by ions

Spores were pressurized in various solutions of known water activity (a_w). Figure 5 summarizes four experiments which suggested that non-ionic solutes (sucrose, glycerol) at low a_w had little effect on the inactivation of spores by pressure, whereas ionic
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solutes (NaCl and, more effectively, CaCl₂) decreased the inactivation. The spores were therefore more likely protected from the inactivating effects of pressure by high concentrations of ions rather than by low a_w per se. Below a_w about 0.96, additional salt had proportionally less effect than above a_w 0.96.

Spores of different species

When suspensions of spores of different species were subjected to pressure, the same phenomena were observed as recorded above, i.e. some spores were inactivated

![Graph](image)

**Fig. 3.** Decreased pressure-sensitivity of preheated spores of *Bacillus coagulans*. Spores (c. 10⁸/ml. in sodium phosphate buffer; 0.1 M, pH 8.0) were heated for 30 min. at the temperatures indicated prior to pressurization at 1000 atm. for 30 min. at 65°C.

**Fig. 4.** Inactivation of spores of *Bacillus coagulans* by pressure at different pH values. Spores (c. 10⁸/ml.) were heated at 70°C for 30 min., then pressurized at 1000 atm. for 30 min. at 65°C. The symbols refer to different buffers (0.1 M) as follows: sodium citrate/phosphate (○); sodium phosphate (●); tris (hydroxymethyl) aminomethane/HCl (□); sodium bicarbonate/carbonate (■); sodium carbonate/hydroxide (△).

**Table 4. Increased pressure sensitivity of γ-irradiated spores of Bacillus coagulans**

<table>
<thead>
<tr>
<th>Pressure and temperature</th>
<th>Control (no irradiation)</th>
<th>0.2 mrad</th>
<th>0.5 mrad</th>
</tr>
</thead>
<tbody>
<tr>
<td>1500 Atm., 45°C</td>
<td>5.5 × 10⁻²</td>
<td>8.0 × 10⁻³</td>
<td>3.5 × 10⁻⁴</td>
</tr>
<tr>
<td>1000 Atm., 65°C</td>
<td>1.9 × 10⁻²</td>
<td>3.5 × 10⁻³</td>
<td>1.0 × 10⁻⁴</td>
</tr>
</tbody>
</table>

* Survivors were enumerated by poured plate viable counts. Radiation alone inactivated spores to the following extent: 0.2 mrad, 20% survivors; 0.5 mrad, 2.2% survivors.

† Spores (c. 10⁸/ml.) were irradiated and pressurized in sodium phosphate buffer (0.1 M, pH 8.0), and were heated at 70°C for 30 min. prior to counting.
and some became heat-sensitive in the manner shown in Fig. 1a, b, and individual spores became phase-dark as if germinated. However, the extent of inactivation varied greatly with different organisms (Table 5). Those spores which are most easily germinated by ‘physiological’ germinants at 1 atm. (e.g. Bacillus polymyxa, B. cereus) were in general more pressure-sensitive than those spores which are most dormant at 1 atm. (e.g. B. subtilis, and mixed spores from soil; Table 5). ‘Superdormant’ spores of B. polymyxa and B. cereus (i.e. those spores in a population which remained ungerminated in a medium which caused germination of the majority) were likewise more resistant than the less dormant ones to the lethal effect of pressure (Table 5).

Fig. 5. Effect of ionic and non-ionic solutes on inactivation of Bacillus coagulans spores by pressure. Spores (c. 10⁸/ml.) were heated at 70° for 30 min., then resuspended in media of differing water activity (a_w) and pressurized at 1000 atm. for 30 min. at 65°. Compounds used to adjust a_w were: sucrose (○); glycerol (□); sodium chloride (●); calcium chloride (■). In addition, the suspensions contained buffer (0.1 M, pH 8.0; not taken into account in a_w estimation) which was tris (hydroxymethyl) amino methane/HCl when calcium chloride was present, or otherwise sodium phosphate.

Table 5. Inactivation of ‘superdormant’ spores by pressure

<table>
<thead>
<tr>
<th>Organism</th>
<th>Survivors (%) following pressurization† for 1 hr at</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacillus cereus PX</td>
<td>0.037     —       —       —       —</td>
</tr>
<tr>
<td>B. cereus PX ‘super-dormant’ fraction</td>
<td>0.75      —       —       —       —</td>
</tr>
<tr>
<td>B. polymyxa M1</td>
<td>0.094     —       —       —       —</td>
</tr>
<tr>
<td>B. polymyxa M1 ‘superdormant’ fraction</td>
<td>1.5       —       —       —       —</td>
</tr>
<tr>
<td>B. cereus NCTC 8035</td>
<td>0.08      —       —       —       —</td>
</tr>
<tr>
<td>B. subtilis var. nigri (syn. globigii)</td>
<td>1.7       0.09    0.0015  0.0003  —</td>
</tr>
<tr>
<td>B. coagulans NCTC 3991</td>
<td>11.5      0.1     0.0075  0.0037  —</td>
</tr>
<tr>
<td>Mixed soil spores</td>
<td>—         —       —       —       —</td>
</tr>
<tr>
<td>B. subtilis A</td>
<td>100       64     3.3      4.6     —</td>
</tr>
</tbody>
</table>

* Survivors were enumerated by poured plate viable counts.
† Spores were suspended in sodium phosphate buffer (0.1 M, pH 8.0) during pressurization.
DISCUSSION

The lethality of hydrostatic pressures in excess of about 1000 atm. for vegetative forms of bacteria is well documented. The most effective pressure depends on the organism, its habitat (e.g. terrestrial, or marine barophile) and stage of growth (see Introduction, and Zobell & Johnson, 1949). Inactivation of vegetative bacteria, by pressure, also depended upon environmental factors; for instance, being most rapid at extreme pH values and lowered by the addition of protective substances such as salts or sugars (Timson & Short, 1965).

The mechanism of inactivation of vegetative bacteria by pressure was suggested by Heden (1964) and Rutberg (1964) to involve damage to the processes of DNA replication, transcription or translation (Landau, 1967). Timson & Short (1965) reasoned that damage may arise via electrostriction and the increase in ionization of weak electrolytes and proteins which is known to result from increase in pressure. Indeed, many studies with pure proteins have shown that high hydrostatic pressures initiate conformational changes which may result in denaturation (Johnson, Eyring & Pollissar, 1954; Kalckar, 1962). Thus the increase in inactivation of vegetative bacteria that occurs with increase in pressure may result from denaturation of proteins.

In contrast to vegetative bacteria, spores resisted pressures well above those which cause the denaturation of free proteins. Furthermore, the pattern of inactivation of spores has been found in this study to differ from that of vegetative forms in a number of fundamental ways. For instance, inactivation of spores by pressure was most rapid near neutral pH values rather than in acid or alkali. In some instances, the denaturation of proteins by pressure, when it involves a decrease in protein molecular volume, can be counteracted by increasing the temperature, which generally causes an increase in protein molecular volume (Eyring, Johnson & Gensler, 1946; Johnson et al. 1954). For example, Morita & Haight (1962) reported that malic dehydrogenase was active at 101° under 1300 atm. pressure. Analogous protection by pressure of vegetative bacteria (Johnson & Lewin, 1946) and spores (Johnson & Zobell, 1949) from inactivation by heat, and vice versa, has been reported. Results in the present paper, however, show that the temperature coefficient for inactivation of spores by pressure is very high, and that, far from counteracting the effect of pressure, heat that was insufficient alone to inactivate spores increased the inactivation caused by pressure.

The discovery of optimal values of pH, temperature and pressure for inactivation of spores by pressure argues against the inactivation having resulted mainly from effects on nuclear material, translation, denaturation of proteins or increased solvation (like the lethal mechanism postulated for vegetative forms by Heden, 1964; Timson & Short, 1965; and Landau, 1967, or for spores by Clouston & Wills, 1969). These mechanisms should generally cause progressively more inactivation as the pressure is raised. The possibility that hydrostatic pressure physically distorted spores sufficiently to cause inactivation was also unlikely since again one would expect high pressures always to cause greater inactivation than low pressures, and one would not expect the high temperature coefficient that was observed. Furthermore, hydrostatic pressure is not vectorial and therefore would not cause distortion of the sort that can disrupt and inactivate spores; i.e. squeezing under a microscope slide cover glass (Lewis, Snell & Burr, 1960), shaking with glass beads (Curran & Evans, 1942) or scraping with a wire loop (Knaysi & Curran, 1961). The regions of pressure and temperature
investigated were such that transitions through different ice phases under pressure (Bridgman, 1937), which can disrupt vegetative bacteria (Edebo & Heden, 1960), did not occur.

The observations that pressure caused heat sensitization of spores, initiated leakage of calcium, dipicolinic acid and hexosamine-containing material, and resulted in fall in optical density of suspensions and phase-darkening of individual spores, together with the electron-microscopical evidence, showed that pressure initiated changes in spore structure and function identical to those observed on germination (Hamilton & Stubbs, 1967). Further evidence is provided by the pressure inactivation being greatest near neutral pH values. The effects of heat (Curran & Evans, 1945) and γ-radiation (Gould & Ordal, 1968) as 'activating' treatments for germination could explain their potentiation of the lethality of pressure for spores. Germination (at 1 atm.) is inhibited by ionic and non-ionic solutes in a similar manner to the inhibition of pressure inactivation of spores (Fig. 5). Spores which are most difficult to germinate completely with 'physiological' germinants at 1 atm. (e.g. soil spores; 'superdormant' spores, Table 5; Gould et al. 1968) were also the most resistant to inactivation by pressure. Taken altogether, the evidence leads to the conclusion drawn by Clouston & Wills (1969) that pressure initiates germination (Gould & Sale, 1970).

Presumably the shapes of the inactivation-pressure curves reflect the sensitivity to pressure of the germination as well as the sensitivity of the germinated spores to pressure inactivation. At lower temperatures the lowest pressures caused germination and heat sensitization but were insufficient to cause appreciable inactivation of the resulting germinated forms. Medium pressures caused considerable germination and were high enough to inactivate a large proportion of the resulting germinated population. The overall effect was therefore to inactivate many but not all of the spores. The highest pressures caused less germination and again only a proportion of the germinated spore population was inactivated, so the overall effect was that the highest pressures caused less inactivation than medium pressures. At higher temperatures the germination was increased and most of the germinated spores were inactivated; always more spores were germinated and became heat-sensitive than were inactivated. Above 65° the germination and inactivation curves coincided, suggesting that the pressure-germinated spores were directly inactivated by heat.

Although pressure induces germination, evidently not all the germinated spores are inactivated by pressure. It is also evident that at near room temperature, spores, whether germinated or not, have a remarkable resistance to extreme pressure.

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References
Bridgman, P. W. (1937). The phase diagram of water to 45,000 kg./cm.². Chemical Physics 5, 964.
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A. Ungerminated spore of *Bacillus cereus* PX, showing wide cortex zone (CX).

B. Spore of *Bacillus cereus* PX germinated by incubation at 37° for 10 min. in buffer (sodium phosphate, 0.1 M, pH 8.0) plus the germinants inosine (100 μM) and L-alanine (1 mM).

C. Spore of *Bacillus cereus* PX germinated by pressurization at 2000 atm. (20°, 1 hr) in nutrient-free buffer (sodium phosphate, 0.1 M, pH 8.0).