The effect of salinity and compatible solutes on the biosynthesis of cyclopropane fatty acids in Pseudomonas halosaccharolytica

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The moderately halophilic eubacterium Pseudomonas halosaccharolytica has been grown at salinities over the range 5-25% (w/v), equivalent to 0.7-3.5 M-NaCl, and the fatty acid composition determined in the late-exponential and stationary phases of batch culture. There was an increase in the proportion of cyclopropane fatty acids (CFA) as the cultures went into stationary phase at all salinities; the overall proportion of CFA was higher in the media containing more salt. The biosynthesis of CFA in P. halosaccharolytica was determined using radiolabelled S-adenosylmethionine as the precursor incubated in cell-free extracts prepared by breaking bacteria with a French press. Compared with the activity obtained in 100 mM-phosphate buffer, the activity of CFA synthetase was inhibited by the addition of NaCl or KCl, but stimulated up to 12-fold by added glycinebetaine, with maximum activity at 3 M. Although the specific activity of CFA synthetase in lysates from cultures grown in 0.7 or 2.1 M-NaCl were similar in the presence of 3 M-glycinebetaine, the enzyme activity in low-salinity cultures was better adapted to function in 1 M-glycinebetaine. Shift-up experiments, in which CFA synthetase activity was assayed in cell-free extracts prepared at different times after increasing culture salinity from 0.7 to 2.1 M-NaCl, showed that the activity of the enzyme was immediately responsive to compatible solute concentration changes and indicated that enzyme induction would not be required to achieve the salt-dependent alterations in membrane lipid CFA composition in vivo. A range of other compatible organic solutes stimulated CFA synthetase activity to a much lesser extent (1-8-fold) compared with glycinebetaine. It is suggested that a compatible solute, which is normally accumulated during osmo(halo)adaptation by an organism in order to contribute towards osmotic balance, does not behave passively towards intracellular proteins but can also stimulate enzyme activity.

Introduction

Halophilic eubacteria require > 3% (w/v) NaCl and can grow in up to 25% (w/v) NaCl (Kushner, 1978; Larsen, 1986). They are important in the spoilage of preserved foods and in saline environments, and so the increasing consumption of ‘fast’ foods and problems of soil salinization of large tracts of artificially-irrigated land worldwide make studies of these bacteria timely. In particular, it is necessary to understand the mechanisms which these bacteria utilize in order to enable them to withstand the osmotic and ionic stresses imposed by high salinity.

The two main ways in which eubacteria (and other micro-organisms) respond to raised external salinity is by the accumulation of intracellular compatible solutes and by modification of membrane composition and function (Russell, 1989a, 1992; Csonka, 1989). In aerobic eubacterial halophiles the compatible solutes most commonly accumulated as major components are the amino acids glycinebetaine, ectoine and glutamate; smaller amounts of the sugar trehalose or other polyols and amino acids may be present (Imhoff, 1986; Regev et al., 1990; Wohlfarth et al., 1990). The compatible solute composition usually depends on the type of culture medium used and in many species, including Pseudomonas halosaccharolytica (Severin et al., 1992), glycinebetaine predominates when complex broth media are employed. There is good evidence that the membrane is involved in the sensing and response to external salinity (Russell & Kogut, 1985; Stock et al., 1989; Csonka & Hanson, 1991), and usually the lipid composition is altered during haloadaptation. Two types of lipid change are observed: in the first, the phospholipid

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Abbreviation: CFA, cyclopropane fatty acid(s).
composition is altered so as to preserve the integrity of the bilayer phase; in the second, there is an alteration in the fatty acid composition of the lipids (Russell, 1989a, 1992). The most frequently reported change in fatty acid composition of Gram-negative eubacteria, including *P. halosacharolytica* (Monteoliva-Sanchez & Ramos-Cormenzana, 1987), is a salt-dependent increase in the proportion of cyclopropane fatty acids which may also increase as batch cultures enter the stationary phase of growth.

Cyclopropane fatty acids (CFA) are formed by the addition of a methylene group across the double bond of a monounsaturated fatty acid (Jarrell et al., 1983), and the thermal properties of CFA-containing lipids (Cronan et al., 1979) or membranes (Jarrell et al., 1983) are subtly distinct. NMR studies support the hypothesis that CFA increase the organizational stability of membranes, whilst providing a level of fluidity comparable with that of monounsaturated fatty acids (Macdonald et al., 1985). Since the CFA synthetase enzyme in different bacteria is either an integral membrane protein or it associates with the membrane and is regulated by activation or induction (Harwood & Russell, 1984), it could well be involved in a self-balancing, salt-triggered system for modifying membrane properties when external NaCl and therefore internal compatible solute concentrations are increased. Therefore, we have investigated the effects of growth phase and the addition of NaCl and organic compatible solutes on CFA synthetase activity and the membrane lipid fatty acid composition in the moderately halophilic eubacterium *P. halosacharolytica*. This was selected for study because it is known to accumulate cyclopropane fatty acids (Ohno et al., 1979) and it is capable of growing over a wide range of salinities.

**Methods**

**Bacterial strains.** *Pseudomonas halosacharolytica* (CCM 2851) was used as the experimental organism. The quotation marks indicate that the name was not included in the Approved Lists of Bacterial Names or its supplements (Skerman et al., 1980; Moore et al., 1985).

**Cultural conditions.** For experimental purposes, cultures were grown in a liquid medium containing 0.5% proteose peptone (Difco), 0.5% yeast extract (Difco) and 0.2% glucose. This medium was supplemented with appropriate dilutions of a stock solution of a balanced mixture of sea salts (Subov, 1931) containing (g l⁻¹) NaCl (243), MgCl₂, 6H₂O (21), MgSO₄, 7H₂O (28.8), CaCl₂, 2H₂O (1.08), KCl (6), NaHCO₃ (0.18) and NaBr (0.078) to give media with final concentrations of 0.7-3.5 M-NaCl.

Cultures were grown in 100 ml of medium contained in 250 ml Erlenmeyer flasks shaken at 100 r.p.m. in a Gallenkamp gyrorotatory incubator at 30 °C. Growth was monitored by measurement of optical density at 500 nm using a spectrophotometer (Novospec, LKB). Stock cultures were maintained on the same liquid medium containing 10% total salts, solidified with agar (2%, w/v), and sub-cultured at weekly intervals; freshly inoculated slopes or plates were grown for 2 d at 30 °C and stored at room temperature.

**Salinity shift-up.** Exponentially growing cultures were shifted-up from 0.7 to 2.1 M-NaCl using the method of Russell et al. (1985). Briefly, this involved minimum centrifugation to collect bacteria and gentle resuspension in prewarmed growth medium. Two types of control were performed in which cultures were grown and resuspended in 0.7 M-NaCl (pre-shift control) or 2.1 M-NaCl (post-shift control). Following shift-up, cultures resumed exponential growth immediately at the same rate as post-shift controls.

**Preparation of cell-free extracts.** Liquid batch cultures were grown to the required growth phase and the bacteria collected by centrifugation at approx. 7000 gₘ, for 10 min.

The bacterial pellets were resuspended in a volume of 100 mM-potassium phosphate buffer, pH 7.6, which was equivalent to 1/10th of the original culture volume. Compatible solutes were predissolved at the appropriate concentration in this buffer solution as required. The bacteria were broken by two passages through a French pressure cell (Amisco) at 44 MPa pressure. Cellular breakage was estimated by phase-contrast light microscopy to be > 99%. It was not necessary to remove the residual intact bacteria because they are impermeable to the radioactive precursor used to assay CFA synthetase activity (vide infra). The lysates were used the same day for determination of CFA synthetase activity.

**Assay of cyclopropane fatty acid synthetase.** Attempts were made to assay CFA synthetase activity in intact bacteria by incubating growing cultures or washed cell suspensions with radiolabelled sodium acetate. However, although the major fatty acids, 16:0, 16:1 and 18:1, were not shown. Therefore, it was decided to use cell-free lysates. CFA synthetase activity was measured by incubating lysates with S-adenosyl-l-[methyl-¹⁴C]methionine (Amersham; final concn 3.7 kBq ml⁻¹, specific radioactivity 2.22 GBq mol⁻¹) with shaking at 30 °C. The reaction was stopped by mixing with an equal volume of KOH (40%, w/v). This mixture was heated at 70 °C for 1 h which releases the fatty acids by saponification. After cooling, an equal volume of water was added, and the solution was acidified to pH < 2 using approx. 6 M-HCl. The fatty acids were extracted with three aliquots of light petroleum (b.p. 40–60 °C) and the combined extracts placed in plastic scintillation vials. The solvent was evaporated and 10 ml of scintillant (FisoFlour '1', Fisons) added and the radioactivity determined using a LKB Rackbeta liquid-scintillation counter programmed with an external-standard channels-ratio facility for the estimation of counting efficiency. CFA synthetase activity was expressed as 'd.p.m. ¹⁴C-radioactivity incorporated (mg lysate protein)⁻¹'. Although there were differences in the activity of individual lysates prepared from separate cultures, the variation in replicate determinations for any one lysate was < 5% and the trends observed (e.g. in the effects of compatible solutes) were consistently reproducible. Protein concentration was measured using the modification of the Lowry method described by Markwell et al. (1978).

**Lipid extraction and analysis.** For the determination of bacterial fatty acid composition, cellular lipids were extracted with chloroform/methanol mixtures using the method of Bligh and Dyer as described by Kates (1986). The chloroform extracts containing total lipid were evaporated to dryness under nitrogen gas and redissolved in fresh chloroform. Aliquots of this extract were used for the preparation of fatty acid methyl esters by transmethylation using conc. H₂SO₄ (2.5%,
v/v) in dry MeOH (Russell & Volkman, 1980). The fatty acid composition was determined by gas-liquid chromatography using a Perkin-Elmer F33 gas chromatograph equipped with glass columns packed with 10% SP2330 on 100–120 Supelcoport (Supelco) as the stationary phase. Separations were performed isothermally at 170 °C.

Radio-GLC was performed using a Pye-Unicam GCD gas chromatograph equipped with a flame ionization detector and connected to a Raga Raytest detector system (LabLogic, Sheffield, UK) utilizing a gas-flow proportional counter. The injector and detector temperatures were 200 °C, and the oven temperature was 170 °C. The carrier gas was 5% (v/v) CO₂/argon. Fatty acid methyl esters were separated isothermally at 170 °C on a glass column (1.5 m x 5 mm i.d.) packed with 5% (w/w) SP-2100 on 100/120 Supelcoport (Supelco).

**Results**

**Fatty acid composition**

Cultures of *Pseudomonas halosaccharolytica* were grown in liquid medium containing 5–25% total salts, equivalent to 0.7–3.5 M-NaCl, and harvested at the end of the exponential phase and 48 h after entering the stationary phase (Fig. 1) for the determination of fatty acid compositions which are given in Table 1. There was little effect of an increase in salinity from 0.7 to 1.4 M-NaCl on the growth rate of cultures; at higher concentrations there was a decrease in growth rate and, particularly at the highest salinities a marked increase in the lag period before exponential growth commenced (Fig. 1). Further increases in salt concentration gave larger increases in the lag period and slower exponential growth rates.

There was an increase in the proportion of CFA as the cultures went from the exponential into the stationary phase (Table 1). The addition of salt to the growth medium also resulted in an increase in CFA content, whether cultures were analysed at the end of exponential or in the stationary phase, with the greatest increase in CFA occurring at or above 2 M-NaCl (Table 1). The increases in total CFA content were matched by corresponding decreases in total unsaturated fatty acid contents.

**Cyclopropane fatty acid synthetase activity**

When lysates prepared in 100 mM-potassium phosphate buffer from cultures grown in 0.7 M-NaCl were incubated with S-adenosyl[methyl-14C]-methionine, there was linear incorporation of the radiolabel into fatty acids for up to 3 h, at a rate of approx. 600 d.p.m. h⁻¹ (mg lysate protein)⁻¹ (Fig. 2). Changing the buffer composition, including increasing its ionic strength, did not increase this rate of incorporation (data not shown), which was too low for radio-GLC analysis to be used for determining the identity of the radiolabelled fatty acid products. However, when glycinebetaine was added (by breaking bacteria in buffers containing increasing concentrations of the compatible solute) there was a large increase in the incorporation of radioactivity of up to 12-fold in 3 M-glycinebetaine (Fig. 2). Further increases in compatible solute concentration up to 6 M-glycinebetaine did not further stimulate incorporation (Fig. 3).

A number of other organic compatible solutes were tested in lysates of cultures grown in 0.7 M-NaCl medium, but they did not stimulate CFA synthetase to the same extent as did glycinebetaine (Table 2). Sucrose and trehalose gave 1.8- and 1.5-fold increases, respectively, but glycerol, proline and sodium glutamate gave activities that were only marginally greater than those of control lysates prepared in phosphate buffer alone. When lysates were prepared in buffer containing 1 M-NaCl, there was 75% inhibition of CFA synthetase activity, and with 3 M-NaCl or -KCl the inhibitions were 89 and 93%, respectively (Table 2).

Radio-GLC analysis of the fatty acid products of S-adenosyl[14C]methionine incorporation by lysates showed that 81–85% of the radioactivity was present in the two CFA, 17:0(cyc) and 19:0(cyc), with the remainder being found in 18:0 and an unknown fatty acid eluting after 19:0(cyc) (Table 3). The precursor was incorporated into the same products whether lysates were prepared in phosphate buffer containing 1 or 3 m-glycinebetaine, from cultures grown in 0.7 or 2.1 M-NaCl medium (Table 3).

The specific activities of CFA synthetase in lysates prepared from cultures grown in 0.7 and 2.1 M-NaCl were compared in the presence of 1 and 3 M-

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**Fig. 1.** Effect of salinity on growth of *P. halosaccharolytica*. Batch cultures were grown at 30 °C in a rich medium (see Methods for composition) containing 5–25% (w/v) of a balanced mixture of sea salts, equivalent to 0.7 (%), 1.4 (●), 2.1 (▲), 2.8 (▼) and 3.5 (□) M-NaCl. Culture growth was monitored by optical density measurements at 500 nm.
Table 1. The effect of culture age and salinity on the fatty acid composition (wt %) of *P. halosaccharolytica*

Cultures were grown in broth medium containing 0·7–3·5 m-NaCl as indicated (equivalent to 5–25%, w/v, total salts) and harvested at the end of the exponential phase and 24 h after entering stationary phase for lipid extraction and preparation of fatty acid methyl esters by GLC analysis as described in Methods. Abbreviations: tr., < 0·1%; exp, exponential phase of growth; stat, stationary phase of growth.

<table>
<thead>
<tr>
<th>Growth phase...</th>
<th>Fatty acid composition (wt %) at equivalent concn of NaCl (m) in culture medium</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0·7</td>
</tr>
<tr>
<td>Fatty acid</td>
<td></td>
</tr>
<tr>
<td>16:0</td>
<td>43·1</td>
</tr>
<tr>
<td>16:1</td>
<td>6·8</td>
</tr>
<tr>
<td>17:0cyc</td>
<td>0·2</td>
</tr>
<tr>
<td>18:0</td>
<td>0·7</td>
</tr>
<tr>
<td>18:1</td>
<td>44·5</td>
</tr>
<tr>
<td>19:0cyc</td>
<td>4·7</td>
</tr>
</tbody>
</table>

Fig. 2. Effect of glycinebetaine on the incorporation of S-adenosyl-L-[methyl-14C]methionine in CFA by cell-free lysates of *P. halosaccharolytica*. Cultures were grown in 5% (w/v) total sea salts, equivalent to 0·7 m-NaCl, to the end of exponential phase. Harvested bacteria were resuspended in 100 mM-potassium phosphate buffer, pH 7·6, alone (○) or containing 0·5 (●), 1·0 (▲), 1·5 (▼), 2·0 (□) and 3·0 (■) m-glycinebetaine, and cell-free lysates prepared by breaking the bacteria with a French press. CFA synthetase activity was determined by the incorporation of S-adenosyl-L-[methyl-14C]methionine (see Methods) and is expressed as 'd.p.m. incorporated (mg lysate protein)-1'.

Fig. 3. Effect of glycinebetaine on the activity of CFA synthetase in cell-free lysates of *P. halosaccharolytica*. Cell-free lysates were prepared in 100 mM-potassium phosphate buffer, pH 7·6, alone or containing 0·5, 1·0, 1·5, 2·0, 3·0, 4·0, 5·0 and 6·0 m-glycinebetaine. The initial rate of CFA synthetase activity was determined and is expressed as 'd.p.m. incorporated h-1 (mg lysate protein)-1'.

glycinebetaine (Fig. 4). Glycinebetaine stimulated the activity of both enzyme preparations, and although the activities in 3 m-glycinebetaine were very similar, in 1 m-glycinebetaine the lysates from low-salinity cultures gave higher activities compared with those of the high-salinity cultures (Fig. 4).

Effect of salinity shift-up on cyclopropane fatty acid synthetase activity

Cultures growing in 0·7 m-NaCl medium were shifted-up to 2·1 m-NaCl and the activity of CFA synthetase determined in lysates prepared in 1 and 3 m-
Table 2. The effect of some salts and organic compatible solutes on CFA synthetase activity in lysates of P. halosaccharolytica

Cultures were grown in broth medium containing 0.7 M-NaCl (equivalent to 5% total salts) and harvested in late exponential phase for the preparation of lysates by breaking bacteria in 0.1 M-phosphate buffer alone ('none') or containing the solute indicated. The activity of CFA synthetase is expressed as ‘d.p.m. incorporated h⁻¹ (mg lysate protein)⁻¹’.

<table>
<thead>
<tr>
<th>Addition</th>
<th>CFA synthetase activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>712</td>
</tr>
<tr>
<td>1 M-NaCl</td>
<td>180</td>
</tr>
<tr>
<td>3 M-NaCl</td>
<td>75</td>
</tr>
<tr>
<td>3 M-KCl</td>
<td>47</td>
</tr>
<tr>
<td>1 M-Glycinebetaine</td>
<td>3041</td>
</tr>
<tr>
<td>3 M-Glycinebetaine</td>
<td>7488</td>
</tr>
<tr>
<td>3 M-Trehalose</td>
<td>1040</td>
</tr>
<tr>
<td>3 M-Sucrose</td>
<td>1302</td>
</tr>
<tr>
<td>3 M-Glycerol</td>
<td>897</td>
</tr>
<tr>
<td>3 M-Proline</td>
<td>851</td>
</tr>
<tr>
<td>3 M-Sodium glutamate</td>
<td>751</td>
</tr>
</tbody>
</table>

Table 3. Effect of culture medium salinity and glycinebetaine concentration on the percentage distribution of radioactivity from S-adenosyl[¹⁴C]methionine in fatty acids synthesized by lysates of P. halosaccharolytica

Cultures were grown in broth medium at the salinities indicated and cell-free lysates prepared by breaking bacteria in buffer containing 1 or 3 M-glycinebetaine as shown. Lysates were incubated for 2 h with S-adenosyl[¹⁴C]methionine and the lipid fatty acids analysed by radio-GLC as described in Methods. There were no significant differences between the fatty acid mass compositions of lysates and the intact bacteria from which they were derived (the latter are given in Table 1).

<table>
<thead>
<tr>
<th>Fatty acid</th>
<th>Glycinebetaine concn (m)</th>
<th>0.7</th>
<th>2:1</th>
</tr>
</thead>
<tbody>
<tr>
<td>16:1</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>16:0</td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
</tr>
<tr>
<td>17:0:ycyc</td>
<td>12.1</td>
<td>8.9</td>
<td>10.6</td>
</tr>
<tr>
<td>18:1</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>18:0</td>
<td>5.1</td>
<td>6.2</td>
<td>6.6</td>
</tr>
<tr>
<td>19:0:ycyc</td>
<td>69.2</td>
<td>73.8</td>
<td>70.4</td>
</tr>
<tr>
<td>Unknown</td>
<td>13.6</td>
<td>11.2</td>
<td>12.4</td>
</tr>
</tbody>
</table>

Abbreviations: ND, not detected; tr., trace, < 0.1%.

glycinebetaine 2 and 6 h after the shift (Table 4). Controls were performed in which cultures growing in 0.7 or 2:1 M-NaCl underwent exactly the same ‘shift-up’ procedures except that the bacteria were resuspended in fresh growth medium having the same salinity as the corresponding pre-shift culture (Table 4). The CFA synthetase activities of lysates prepared in 3 M-glycinebetaine measured 2 and 6 h after a shift-up were 73 and 48%, respectively, compared with the pre-shift...
salinity controls; in contrast, the activities of these 'shifted lysates' were 149 and 91%, respectively, compared with the post-shift salinity controls. In both the shifted and control lysates there was a marked stimulation of CFA synthetase activity by glycinebetaine, ranging from 4- to 8-fold.

Discussion

_Pseudomonas halosaccharolytica_ was chosen for the present study, since it was known to accumulate CFA in its membrane lipids in response to an elevation in culture medium salinity (Ohno _et al._, 1979). The present data demonstrate the interplay between culture age and salinity on CFA content, which was important to establish under our cultural conditions before studying the effects of compatible solutes on CFA synthetase activity since it was necessary to ensure that cultures were harvested at clearly defined stages of the growth cycle in batch culture. Salt concentrations of 0·7 and 2·1 M were chosen as a suitably wide range (3-fold difference) within which salinity had only a small effect on the exponential growth rate (Fig. 1), since growth rate changes could also influence fatty acid composition. Thus we were able to study the effects of salt and compatible solutes on CFA synthetase activity _in vitro_ in lysates prepared from cultures which had been grown at the same rate but in different salinities and with different fatty acid compositions.

Since CFA accumulate in a salt-dependent manner, it had been anticipated that we should find that the specific activity of CFA synthetase would be greater in lysates prepared from cultures grown in 2·1 M-NaCl compared with those grown in 0·7 M-NaCl. When account is taken of the (presumed) total intracellular concentration of compatible solutes, this was indeed the finding, there being 2–3-fold more activity at the higher salinity. However, most of this difference appears to be due to the stimulation by glycinebetaine, although the enzymes from low- and high-salinity cultures did react differently to NaCl in that much lower activities were obtained at low salinity with the lysates from 2·1 M-NaCl compared with 0·7 M-NaCl-grown cultures (Fig. 3). Clearly the enzymes react differently and there is a marked effect of the compatible solute, but it is difficult to draw more precise conclusions because it was not possible to mimic more closely the intracellular compatible solute composition. The cells would have contained ectoine as well as glycinebetaine when grown in the broth medium (Severin _et al._, 1992). However, we did not have ectoine available to add to lysates, although they would have contained some of the endogenous compounds which remain during cell breakage. Therefore, both the composition and total concentration of compatible solutes in lysates are approximations.

In order to investigate further the relationship between CFA synthetase activity and culture medium salinity, shift-up experiments were performed. There was no lag period following a shift-up from 0·7 to 2·1 m-NaCl and, after the shift, cultures resumed exponential growth at essentially the same rate. It was decided to sample cultures 2 and 6 h after the shift-up, because this was known to cover the early and later stages of the adaptive process in respect of changes in fatty acid composition. However, over this time-scale we did not observe any adaptive changes in CFA synthetase activity. It appeared that within 2 h after the shift, the enzyme responded in a manner more like that of cultures grown in the post-shift salt concentration, and that the major determinant of enzyme activity was the compatible solute concentration. The time-scale of changes in the intracellular concentrations of glycinebetaine and ectoine in this bacterium is not known, but the data obtained with cell-free lysates indicate that enzyme activity would respond immediately to any changes and that enzyme induction would not be necessary to achieve the changes in fatty acid composition.

A major function of the organic compatible solutes which are accumulated inside micro-organisms in response to salinity is assumed to be the balancing of osmotic pressures outside and inside the cell, although exactly equivalent balancing has not been proven for any organism. Moreover, it is not clear whether they have any other function(s) inside cells. Their accumulation to high concentrations inside cells means that they must at the very least not interfere with the activity of intracellular protein and other macromolecular functions. There are several theories of how compatible solutes achieve their functions and, although none provide a conclusive explanation, it is clear that the maintenance of protein hydration becomes critical for halophiles growing in high salinities, assuming that they maintain osmotic equilibrium between the cytoplasm and the culture medium (Trüper & Galinski, 1990). A mechanism proposed for glycinebetaine is that it functions as a compatible solute by acting on the cluster structure of water and thereby indirectly influences the hydration of proteins. The present data show that glycinebetaine, one of the major compatible solutes which are accumulated by _P. halosaccharolytica_ cultured in complex growth media (Severin _et al._, 1992), has a marked positive effect on CFA synthetase activity. Therefore, whatever the molecular explanation for glycinebetaine action in this organism, it clearly is not a passive one; moreover, it is selective in the sense that other organic compatible solutes which are found in different halophilic micro-
organisms stimulate CFA synthetase activity of *P. halosaccharolytica* much less or only marginally. Compatible solutes such as glycinebetaine (Pollard & Wyn Jones, 1979) and ectoine (Lippert & Galinski, 1992) are known to be capable of stabilizing enzyme activities, but it is not known if this plays a specific role during osmo(halo)-adaptation. If the naturally accumulated compatible solute stimulates other enzymes, besides CFA synthetase, which are involved in osmo(halo)-adaptation to the same extent or differentially, this would be a major factor in the regulation of intracellular metabolism during this process and would necessitate a revision of ideas about the functional significance of the interactions of compatible solutes with cellular proteins.

In marked contrast with the organic compatible solutes, NaCl and KCl strongly inhibited CFA synthetase, despite the fact that bacteria were grown in the presence of \( \geq 0.7 \text{m-NaCl} \). Salt is known to inhibit the activities of a number of intracellular enzymes in moderately halophilic eubacteria and optimum salt concentrations in the range \( 0.1-0.6 \text{m-NaCl} \) have been reported (Kamekura, 1986). The only comparable published data on fatty acid synthetases is that of Pugh et al. (1971) who observed a cell-free preparation of the fatty acid synthetase from *Escherichia coli* was 90% inhibited by 1 m-NaCl. In addition, we have observed similar levels of inhibition by salt of the fatty acid synthetase in *Vibrio costicola* (M. Kogut and N. J. Russell, unpublished data). This inhibition is probably a result of the high ionic strength, rather than a specific effect of Na+, because we observed a comparable inhibition by KCl. It is not surprising that the fatty acid synthetase from *V. costicola* is sensitive to high concentrations of NaCl or KCl because we have shown in inhibition by KCl. It is not surprising that the fatty acid synthetase, which are involved in osmo(halo)-adaptation to the same extent or differentially, this would be a major factor in the regulation of intracellular metabolism during this process and would necessitate a revision of ideas about the functional significance of the interactions of compatible solutes with cellular proteins.


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