Characterization of an autostimulatory substance produced by *Escherichia coli*

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**INTRODUCTION**

Bacterial populations are capable of cooperation and coordination of their activities to an extent which reminds one of the behaviour of multicellular organisms (Shapiro, 1998). Recently, the role of intercellular communication in the regulation of these social phenomena has received widespread attention, and a variety of diffusible chemical signals involved in communication has been discovered. The phenotypes regulated by intercellular communication include luminescence, gene transfer, sporulation, swarming, fruiting body formation and the production of siderophores, exoenzymes and other virulence factors, as well as the synthesis of antibiotics and other secondary metabolites. The chemical nature of the signals affording these types of communication depends on the organism and phenotype studied; in some cases they are highly species-specific, and are thus considered 'pheromones'. Bacterial pheromones described so far include peptides, nucleotides, acylated homoserine lactones, amino acids, fatty acids, cyclic dipeptides, a quinolone and a ketone (Fuqua & Greenberg, 1998; Fuqua et al., 1996; Holden et al., 1999; Hull et al., 1998; Kaiser & Losick, 1993; Kell et al., 1995; Pesci et al., 1999; Salmond et al., 1995; Swift et al., 1996). In certain Gram-positive organisms, intercellular growth stimulation mediated by proteinaceous cytokines has also recently been demonstrated (Kaprelyants et al., 1999; Kell & Young, 2000; Mukamolova et al., 1998a).

In *Escherichia coli*, no signalling molecule with a defined role has been chemically identified so far to our knowledge, although several papers have reported the action of such substances. Induction of certain *E. coli* genes by the addition of supernatant has been described (Baca-DeLancey et al., 1999; Li et al., 1997), and growth-inhibiting effects of supernatants have been reported in *E. coli* and other organisms (Barrow et al., 1996; Gray et al., 1996; Srinivasan et al., 1998). So far, however, the substances involved in these processes have not been identified. While homoserine lactone derivatives were shown to have growth-inhibiting effects by several authors (Garcia-Lara et al., 1996; Sitnikov et al., 1996; Surette & Bassler, 1998; Surette et al., 1999; Withers & Nordström, 1998), the production of these molecules by *E. coli* has not been demonstrated. On the other hand, published accounts of cases in which *E. coli*
supernatants stimulate growth, survival and activity of the producing strain appear to be rare. The only case known to us involves the autoinducer-mediated stimulation of the growth of a variety of Gram-negative organisms in serum-containing media by exogenously supplied noradrenaline (norepinephrine), as described by Lyte, Williams and colleagues (e.g. Freestone et al., 2000; Lyte et al., 1996). This, however, might not be considered a pure case of autostimulation because of the involvement of the eukaryotic hormone.

It is of course a common observation that low inocula lead to extremely long lag times unless the cells are supplied with ‘spent medium’ (also called ‘conditioned medium’ or ‘culture filtrate’; Lodge & Hinshelwood, 1943). Investigations of the physiological or chemical basis of this kind of effect are, however, rarely found in the literature. Ajl & Werkman (1948) reported stimulation of E. coli by CO\(_2\) which potentially accumulates in spent medium (cf. Dixon & Kell, 1989; Neidhardt et al., 1974). We here describe an autostimulatory effect of E. coli supernatants which is independent of the requirement for CO\(_2\) and which to our knowledge has not been characterized before.

### METHODS

**Bacterial strains and maintenance.** Escherichia coli ZK126 [W3110 ΔlacU169 tria2 (Connell et al., 1987), received from R. Kolter, Harvard Medical School] was used throughout this study as the model strain. By using other E. coli strains [ATCC 25922, and ZK1000 (ZK126 rpsO::kan, R. Kolter)], it was ascertained that the observed effects are not strain-specific. All strains were maintained on LB agar plates.

**Chemicals.** All chemicals used were analytical grade and obtained from Sigma, except for cAMP, which was obtained from Boehringer, and N-hexanoylhomoserine lactone (HHL) and butyrylhomoserine lactone (BHL), which were a gift from Dr Michael Givskov (Technical University of Denmark, Lyngby, Denmark). HPLC grade solvents were purchased from Fisher Scientific.

**Growth conditions and media.** All cultures were inoculated from 2-d-old single colonies on LB plates into MM medium [MOPS-buffered minimal medium (Neidhardt et al., 1974), see below]. The cultures were grown at 37°C with orbital shaking for 20 h, diluted 1:200 in fresh MM medium, grown for another 20 h, and finally diluted once again 1:200 in fresh MM medium. This method ensures that the culture-over material from the LB-grown colonies (which might contain a large proportion of 3-4 litres of initial inoculate is assumed for cultures of 2 ml). At the same time this way both the homogeneity and the reproducibility of the physiology of cells in the final culture were ascertained.

Medium MM was mixed freshly on the day of experiments from the following three stock solutions: 44 ml 1.5 mM KH\(_2\)PO\(_4\), 5 ml sterile filtered ‘10× MOPS’ (see below, stored frozen at -20°C), and 0.75 ml sterile filtered 0.67 M NaHCO\(_3\) (final concentration: 10 mM). Appropriate carbon sources were added at the concentrations indicated: glucose or succinate were added from sterile filtered stock solutions of 10% (w/v) (final concentrations: 0.001–0.05%, w/v), and glycerol was added from autoclaved stock solutions.

10× MOPS is a 10× stock of buffer substances and salts containing 400 ml 1 M MOPS (pH 7.4), 40 ml 1 M Tricine (pH 7.8), 10 ml 10 mM FeSO\(_4\), 50 ml 1.9 M NH\(_4\)Cl, 10 ml 0.276 M K\(_2\)SO\(_4\), 5 ml 0.5 mM CaCl\(_2\), 5 ml 0.528 M MgCl\(_2\), 100 ml 5 M NaCl, 10 ml ‘MNS’ (Micro-Nutrient-Solution, see below) and 370 ml H\(_2\)O to give 1000 ml. ‘MNS’ contained 10 µM ZnSO\(_4\), 80 µM MnCl\(_2\), 10 µM CuSO\(_4\), 30 µM CoCl\(_2\), 0.4 mM H\(_3\)BO\(_3\) and 3 µM (NH\(_4\))\(_2\)(MoO\(_4\))\(_2\). Final concentrations of MOPS, NaHCO\(_3\) and all inorganic constituents of MM medium are identical to those published by Neidhardt et al. (1974).

**Vitamins.** Vitamins were tested at the following concentrations (l\(^{-1}\)): 50 µg riboflavin, nicotinamide and b-panthothenic acid; 20 µg para-aminobenzoic acid, folinic acid and d-biotin; 15 µg pyridoxine; 10 µg pyridoxal. HCl and DL-3,8-thiazic acid; 5 µg pyridoxamine; 2 µg cyanocobalamin; and 0.03 µl tocopherol. This combination of vitamins is considered complete (Cote & Gherna, 1994) and is referred to as ‘12 vitamins’ in Results. In addition, thiamin was tested at concentrations of 2, 4, 10, 25, 50 and 200 µg l\(^{-1}\).

**Preparation of supernatants and testing of effects on growth.** Supernatants from exponentially growing or stationary-phase cells were prepared in the following fashion. The cells were grown in MM medium containing 0.04% (w/v) glucose unless stated otherwise with shaking at 37°C, harvested by centrifugation (20°C, 12000 g, 10 min), and the supernatants were sterile filtered twice (0.2 µm; Sartorius). The sterile filters were always washed with at least 40 ml HPLC grade water immediately prior to using them to filter supernatants or any other solutions to reduce the possibility of substances being released from the filter cartridges into the filtrates. The filtered supernatants were frozen at -20°C until they were processed. The activity of supernatants of cells grown in vessels of different materials and sizes was tested, but no difference in activity was found between cultures grown in plastic screw cap tubes (50 ml total volume, holding 5 ml cultures) or in different sizes of glass Erlenmeyer flasks (up to 1 l culture in flasks of 3 l total volume). For production of supernatants of cells that were limited due to depletion of carbon sources were kept at 37°C with shaking after growth in MM medium with 0.04% (w/v) glucose, and harvested at the times indicated. For supernatants of cells that were carbon-limited after resuspension in medium lacking carbon source, the cells were grown in MM medium containing 0.8% (w/v) glucose, harvested in exponential phase at 3×10\(^{10}\) c.f.u. ml\(^{-1}\), and resuspended in MM medium without glucose.

To test the recovery of cells in different growth phases, populations of bacteria were grown in MM medium containing 0.04% (w/v) glucose (or 0.05%, w/v, glycerol or succinate) under the same conditions, and left at 37°C with shaking. At the time of experiment, the populations were diluted in fresh MM medium to a final cell density of 100–500 c.f.u. ml\(^{-1}\), volumes of 200 µl each were dispensed into prewarmed 100-well plates, and growth was recorded as turbidity using an automatic growth analyser (BioScreen, Labsystems). The growth analyser was set to a temperature of 37°C and shaking to periods of 20 s of maximum intensity min\(^{-1}\). Optical density (OD\(_{360}\)) was monitored every minute with the wavelength at the ‘wide band’ setting. Apparent lag phases and growth rates were estimated by curve fitting of the OD\(_{360}\) data, using a three-phase fitting programme: the first phase (lag phase) was fitted to y = y\(_0\); the second (growth) phase to y = y\(_0\) exp{(t–lag)}; the third (stationary) phase was fitted to y = y\(_{max}\). The parameters are y\(_0\) = initial OD, t = time in hours, lag = lag time in hours and g = growth rate in divisions h\(^{-1}\). True lag times were determined from c.f.u. data.
data by plating out dilutions of samples on LB agar in intervals of 30–60 min.

**Characterization of active compounds.** Acid and alkali sensitivity were determined by adding HCl or NaOH, respectively, and keeping the treated samples at pH 4.2 or 12.8, respectively, for 45 min at 25 °C, after which the pH was readjusted to pH 7 with NaOH or HCl. Proteinase sensitivity was tested by incubating samples with 2 and 10 mg Proteinase K–acyrlic beads (Sigma) ml⁻¹ for 30 and 60 min at 37 °C.

Dialysis was performed against distilled water for 20 h at 25 °C using either standard dialysis tubing or SpectraPor 3 membrane (3.5 K cut-off). Ion-exchange affinity was tested by employing DE52 and CM52 resins (Whatman) at 4 g (ml supernatant)⁻¹ for 20 min at pH 4, 7 or 10 at 25 °C. Residues of the resins were removed by centrifugation (4000 g, 10 min) and filtration through 0.22 μm cellulose acetate membrane filters (Sartorius).

**Preparation of extracts.** Salts were removed from sterile filtered supernatants by consecutive ion-exchange treatment with DE52 and CM52 at pH 7 (see above) and the treated supernatants were evaporated to dryness at 60 °C using a rotary evaporator. Extracts were resuspended in 1/60 volume of distilled water, and frozen at −20 °C. The extracts obtained by this procedure are referred to as ‘supernatant concentrates’ in this study.

Ethyl acetate extractions of supernatants were performed at pH 4, 7 and 9.6, the solvent was evaporated, and the extracts were resuspended in HPLC grade water. Ethanol extracts were performed after saturation of supernatants with K₂CO₃ (0.9 g ml⁻¹), employing three times equal volumes of HPLC grade ethanol. The ethanol was evaporated off, the extracts were resuspended in HPLC grade water, and the extraction was repeated twice. This procedure allows effective removal of salts and concentration of up to 300-fold.

**Analysis of supernatant extracts by HPLC and LC-MS.** Supernatant concentrates and extracts were separated by HPLC (Waters 2690) employing a variety of columns and solvents. The columns used included reverse-phase columns such as Spherelone C18 (Phenomenex), Symmetry C18 (Waters) and Supersil LC18S (all 5 μm, 3.5 × 150 mm), a normal phase column (NOVAPAK Silica; Waters), a phenyl column (NOVAPAK Phenyl) and a size-exclusion column (Ultrahydrogel 120; Waters). Solvent systems were mixtures of HPLC grade water with either 0.5 mM acetic acid or 3 mM ammonium acetate, and with varying concentration of methanol or acetonitrile. Separation was performed at 30 °C, and elution was monitored using a diode-array detector. Fractions were taken every 0.5–1 min, evaporated, resuspended in HPLC grade water, and tested for activity. Electrospray MS (LCT instrument; Micromass) was performed either on active fractions directly (by using a syringe pump at 5 μl min⁻¹) or by linking the HPLC directly to the LC-MS interface.

### RESULTS

**Effects of the addition of stationary phase supernatants to cultures of *E. coli***

Batch cultures of *E. coli* strains were grown in MM medium containing 0.04% (w/v) glucose at 37 °C with shaking. Cultures reached a final cell density of between 7 × 10⁸ and 1.5 × 10⁹ c.f.u. ml⁻¹, and were kept at the same temperature with further shaking. Cells of *E. coli* ATCC 25922 were kept in stationary phase for 2 d, at which time the cell density was 1.3 × 10⁸ c.f.u. ml⁻¹. When these cells were diluted in fresh growth medium to a final cell density of 284 c.f.u. ml⁻¹, their recovery was strongly stimulated by addition of supernatant of 2-d-old stationary phase cells of the same strain. The apparent lag times (as estimated from BioScreen OD590 data) of unamended stationary phase cells were 20–3 h (all data means of five experiments; SD = 1.51), while the same cultures amended with 30% (v/v) filtered supernatant displayed lag times of only 13.2 h (SD = 0.44). Cells of the same strain in the exponential phase of growth (at 4.05 × 10⁹ c.f.u. ml⁻¹, diluted to a final cell density of 560 c.f.u. ml⁻¹) were also stimulated by the addition of stationary phase cell supernatant. Untreated growing cell populations showed mean lag times of 10.9 h (SD = 0.22), while populations treated with 30% (v/v) of the same supernatant as employed above were observed to display an apparent lag of 8.5 h (SD = 0.21). In summary, stimulation of recovery by supernatant addition was observed reproducibly with growing or starving cells, but stimulation of stationary phase cells was much more pronounced than that of growing cells of the same strain in the same medium.

A detailed analysis of supernatant effects on stationary phase cells is shown in Fig. 1. Here cells of *E. coli* ZK126 that had been carbon-starved for 53 d were diluted in fresh, fully supplemented MM medium (containing 0.04%, w/v, glucose) amended with different amounts of supernatants of a culture of the same strain that had been starved for 2 d under identical conditions (see Methods; +, control; ▼, addition of 30% untreated supernatant; ○, 50% supernatant; ▲, 80% supernatant). Datasets shown are representative sets of five parallel sets each.

![Fig. 1. Addition of untreated supernatant during recovery of starved cells of *E. coli* ZK126. Cells starved in MM medium for 53 d were diluted (to a final density of 144 c.f.u. ml⁻¹) in fully supplemented MM medium (containing 0.04%, w/v, glucose) amended with different amounts of supernatants of a culture of the same strain that had been starved for 2 d under identical conditions (see Methods; +, control; ▼, addition of 30% untreated supernatant; ○, 50% supernatant; ▲, 80% supernatant). Datasets shown are representative sets of five parallel sets each.](Image 362x596 to 540x733)
Influence of density and age of cell populations on activity of supernatants

As expected, the density of supernatant-producing cultures (dependent on the concentration of glucose added at the onset of the experiment) had some influence on the effects of the supernatants. The supernatants obtained from a high-density culture of ZK126 (MM medium with 0.4%, w/v, glucose; 4 × 10^9 c.f.u. ml^-1) had a stronger stimulatory effect than that of a culture with fivefold lower cell density (0.04%, w/v, glucose; 8 × 10^8 c.f.u. ml^-1). In one experiment with recovering stationary phase cells of ZK126, the untreated control cells showed mean apparent lag times of 17.2 h. Here, the addition of 30% (v/v) high-density supernatants reduced lag times to 11.1 h, and addition of the same proportion of low-density supernatant led to lag times of 13.8 h (all data means of five experiments each; similar results were obtained with ATCC 25922).

The effect of the age of cell populations on the stimulatory activity of their supernatants was tested in a series of experiments. In one set of experiments, populations of exponentially growing cells of E. coli ZK126 were harvested at 3.75 × 10^8 c.f.u. ml^-1, and 2-d-old stationary cells of the same strain were harvested at 8.5 × 10^9 c.f.u. ml^-1, and the supernatants obtained were sterile filtered. In parallel experiments, the two supernatant preparations were added at the same concentrations to 2-d-old stationary phase cells of the same strain (cell density 9.1 × 10^9 c.f.u. ml^-1) diluted in fresh MM medium to a final density of 120 c.f.u. ml^-1. Both supernatants significantly shortened the lag times of the recovering cells, but the supernatant of stationary phase cells clearly had a stronger effect than that of growing cells. Exponential phase supernatant shortened the apparent lag times from 17-2 h (untreated cells, SD = 0.47, n = 5 in all experiments) to 14.9 h (SD = 0.39) when added at 15% (v/v), and to 13.4 h (SD = 0.25) when added at 30% (v/v). Stationary phase supernatants, on the other hand, reduced the apparent lag times of the same population of recovering cells to 12.8 h (SD = 0.26) when added at 15% (v/v), and to 11.5 h (SD = 0.21) when added at 30% (v/v). Supernatants of cell populations kept in stationary phase for 3 or more days showed similar levels of activity as those kept for 2 d (data not shown). Thus for simplicity a standard time of 2 d of stationary phase (exactly 40 h) was adopted prior to harvesting for supernatant production.

Effect of supernatant addition on true lag times

True lag times were determined by plating on LB agar at intervals of 30–60 min. After 2 d in stationary phase, the mean true lag time displayed by populations of E. coli ZK126 was 2.2 h (mean of triplicates). After 8 d in stationary phase, populations of equal density were observed to have a mean lag phase of 4.5 h, after 48 d 5.5 h, and after 138 d 5.6 h (means of triplicates). Interestingly, the c.f.u. counts were observed to drop transiently during the lag phase of recovering cells which had been kept in stationary phase for 48 or more days.
The decreases were between 20 and 85% of the initial c.f.u. counts, with the lowest counts observed between 3 and 5 h after dilution in fresh medium. When 30% (v/v) supernatant was added to 2-d-old stationary cells, true lag times were reduced to 1–3 h (data not shown). When 138-d-old cells were diluted in fresh medium (final cell density of 275 c.f.u. ml⁻¹) amended with supernatant extract (corresponding to addition of 30%, v/v, supernatant), the true lag times were found to be between 1.8 and 2.5 h (data not shown). When supernatants or supernatant extracts were present during recovery, a decrease of c.f.u. counts such as described above for untreated suspensions was never observed. Growth rates were also increased by a factor of 1.5 in the presence of supernatant components. Corrected for the decrease in generation time, this reduction in lag time by supernatant addition amounts to 58–62% of the lag time at 25 °C.

Assessment of the number of recovering cells

The number of recovering cells in the presence and absence of supernatant was tested using an MPN (most probable number) technique: serial dilutions of stationary phase cells of ATCC 25922 or ZK126 were inoculated into fresh media with or without 30% (v/v) addition of the supernatant of a 2-d stationary phase culture of ATCC 25922. Ten parallels were employed for each treatment, and growth was monitored over 10 d using the BioScreen growth analyser (with OD₅₉₀ recorded every hour). No difference, however, was observed between MPN estimates of the recovery of treated and untreated samples in either of the strains: the percentages of replicate cultures that had grown were identical in the presence or absence of supernatant (data not shown). Thus we conclude that the components present in supernatants do not increase the number of recovering cells, but merely accelerate their recovery and subsequent growth.

Characterization and extraction of the active compound(s)

It was found that the growth-stimulating compound(s) produced by *E. coli* ATCC 25922 and ZK126 was preserved after 30 min boiling (Fig. 2, data shown for ATCC 25922 only). Further, the activity was found to be acid- and alkali-stable: exposure to pH 4.2 or 12.8 (at 25 °C) for 45 min each did not reduce the activity. Also, treatment with proteinase K (see Methods) had no effect on the growth-stimulating component (data not shown). Hence, the active component is heat-, acid-, alkali- and proteinase-resistant. On the other hand, heating to 250 °C for 17 h destroyed the activity completely (data not shown). Dialysis quantitatively removed the active constituents from supernatants: dialysed supernatant had no stimulatory activity whatsoever – in fact, significant inhibitory effects were observed after addition of dialysed supernatants (Fig. 2). Identical results were obtained for supernatants and cells of *E. coli* ATCC 25922 and ZK126.

Ion-exchange resins DE52 and CM52 were not found to retain significant amounts of active material at pH 4, 7 or 10. In fact, inhibitory material was removed by consecutive treatment with both ion-exchange resins (at neutral pH and 25 °C), allowing growth to higher final cell densities as compared to growth in the presence of untreated supernatant (Fig. 3). Activity was also main-
tained during rotary evaporation of supernatants or extracts (Fig. 3), which allowed concentration of samples up to 80-fold.

Extraction with ethyl acetate was not successful: no activity was detected in the organic phase after extraction at any pH tested (see Methods), while more than 95% of the activity remained in the aqueous phase even after several consecutive extractions with the solvent (data not shown).

After saturation with salt, the active component(s) could be recovered by extraction with ethanol (as outlined in Methods). Quantitative recovery could be demonstrated after three rounds of extraction (Fig. 4). Thus ethanol extraction of carbonate-saturated supernatants was shown to be a convenient and elegant way of concentrating the active compound(s).

HPLC was used as described in Methods to separate substances present in supernatant concentrates and ethanol extracts. More than 20 substances could be detected using the diode-array detector: uracil, cytosine, xanthine and thymine were tentatively identified by LC-MS to be present in the extracts. Stimulatory activity co-migrated with the uracil peak, irrespective of the type of column employed. Uracil, however, was shown not to be active (see below), and no other reproducible peak correlating with activity could be detected by LC-MS. A range of conditions (flow rates, solvents, voltages, etc.) was tested, but none of these attempts proved successful for identifying the active substance chemically.

Testing vitamins and other growth factors

In order to test the possibility that the supernatant components affording the growth stimulation are vitamins or other growth factors, a complete set of vitamins, a set of purines and pyrimidines, and a selection of amino acids were tested in their effect on recovery and growth.

A complete set of 12 vitamins, as detailed in Methods, stimulated growth to a certain extent, but not as much as brought about by supernatant addition. This was observed for cells that had been in stationary phase for 2 or 45 d for both strains tested (ZK126 and ATCC 25922; data shown for ZK126 in Fig. 5). Addition of thiamin at a range of concentrations (see Methods), in addition to the set of 12 vitamins, had no further effect on lag times or growth rates, and thiamin addition alone had no effect on lag times (data not shown).

As expected, addition of yeast extract or Casamino acids (both at 0.03–0.0003%, w/v) allowed recovery to proceed much faster than in the untreated controls. Supplementation with yeast extract or Casamino acids at concentrations of 0.003% (w/v) or more led to faster recovery than in the basal media amended with 30% (v/v) supernatants (data not shown). None of the single amino acids tested (methionine, threonine, leucine, isoleucine and glutamate), however, could be shown to be responsible for growth stimulation. None of the following substances stimulated recovery of stationary phase E. coli: sodium acetate, fumarate, l-homoserine lactone, l-homoserine, dL-homocysteine, butyrylhomoserine lactone (BHL) and N-hexanoylhomoserine lactone (HHL), and glutathione (disodium salt). Amino acids were tested at concentrations between 1 µM and 1 mM, sodium acetate was tested at concentrations ranging from 0.0001 to 1% (w/v), fumarate at between 0.01 and 0.5% (w/v), l-homoserine lactone was tested at between 0.2 and 500 µM, l-homoserine and dL-homocysteine at between 0.2 and 10 µM, BHL and HHL from 0.1 to 100 µM, and glutathione from 0.01 to 0.5% (w/v) (ZK126 tested only, data not shown). Addition of
reduced and oxidized forms of glutathione (added from freshly prepared stock solutions) did not reduce lag times of stationary phase cells of *E. coli* ZK126 at concentrations between 0.01 and 0.5% (w/v), but rather had dose-dependent inhibitory effects. Also, addition of noradrenaline (Arterenol, bitartrate salt; Sigma A9512) at a concentration of 50 µM (as described by Lyte *et al.*, 1996) did not lead to a reduction, but to an increase of the lag times observed in our assays with ZK126 (data not shown).

Due to the presence of purines and pyrimidines in supernatant extracts and the stability of active compounds (see above), purines and pyrimidines and their heat-stable derivatives were screened employing HPLC and BioScreen assays. The reverse-phase HPLC retention times of the following chemicals were much longer than those of the active fractions, and were thus not tested further: 1-methyl-adenine, 2-O-methyl-adenine, 3-O-methyl-adenine, 8N-methyl-adenine, adenosine, adenosine-monophosphate, adenosine-5’-diphosphate, adenosine-2’,5’-diphosphate, P1, P2-di(adenosine-5’)-pyrophosphate, P1, P2-di(adenosine-5’)-tetraphosphate, 5-adenosinyl-methionine, cytosine, iso-cytosine, guanosine, cGMP, thymine, 1-methyl-uracil, 3-methyl-uracil, 6-methyl-uracil, 5-ethyl-uracil, uracil-4-acetic acid and the cyclic dipeptides (diketopiperazines) cyclo-(Phe-Ser), cyclo-(His-Pro) and cyclo-(His-Phe). Of all the chemicals tested, only uracil and 5-hydroxy-methyl-uracil displayed HPLC retention times similar to those of the active substance(s). Uracil and 5-hydroxy-methyl-uracil, however, were without effect on recovery of *E. coli* ZK126 or ATCC 25922 when applied at concentrations between 0.01 and 0.5% (w/v, glucose) were diluted (1:100) and extracts, strain ZK126 was grown in MM medium not amended with iron (with 0.04% NaCl and 0.015%, w/v, glucose). The final cell densities of the cultures were 2.02 x 10^8 and 2.34 x 10^8 c.f.u. ml^{-1}, respectively. These cultures were then split into aliquots, one-half of which were amended with supernatants, supernatant extracts, or active fractions from HPLC at concentrations corresponding to those which had previously been shown to yield stimulation. Other media ingredients such as fresh nutrients were not added in this set of experiments, in order to test whether supernatant extracts themselves could supply carbon or nitrogen sources. During the following incubation at 37°C, OD_{590} and c.f.u. were monitored, and the experiment was repeated at least five times with each treatment. Upon addition of supernatant extracts or active fractions, the OD_{590} was observed to increase slightly immediately in comparison to the control. In carbon-limited cultures, the OD_{590} in the amended samples was 0.168 as compared to 0.164 (means of 12 measurements; standard deviations 0.017 and 0.016, respectively), and in nitrogen-limited cultures 0.184 as opposed to 0.177 (means of 11 experiments; standard deviations 0.022 in both cases). When the OD_{590} measurements were extended over 4 h, no further increases were observed. The c.f.u. counts in the treated cultures, however, were not higher than in the untreated cultures in any experiment at any time (samples were taken 40 and 170 min after addition) – actually, carbon-limited cultures displayed significantly lowered c.f.u. counts after supernatant extracts had been added, while nitrogen-limited cultures were unaffected (P < 0.05). In short, no growth of carbon- or nitrogen-limited cultures could be observed by addition of supernatant or supernatant extracts at concentrations which elicit the stimulation of growth described above. The slightly elevated OD_{590} data in treated samples can be explained by light absorption of the extracts themselves or by slight modification of the refractive index of the cells which is largely responsible for the light scattering in measurements of this type (Davey & Kell, 1996).

A series of experiments was performed in order to test the hypothesis that siderophore production might be responsible for the stimulatory activity of supernatants. To this end, the effect of supernatants of iron-limited cells on recovery was tested. Stationary phase cells (ZK126 grown overnight in fully supplemented MM medium with 0.04%, w/v, glucose) were diluted (1:100) in MM medium not amended with iron (with 0.04% NaCl and 0.015%, w/v, glucose), and growth was allowed to proceed into stationary phase as in the standard experiments. The addition of the supernatants of these (iron-starved) cells, however, had a smaller effect on lag times of recovering cells (strains ATCC 25922 and ZK126 tested). Thus siderophore production can be excluded as an explanation for the autostimulatory effect.

**Testing the presence of nutrients or siderophores in supernatants**

To test for the presence of nutrients in the supernatants and extracts, strain ZK126 was grown in MM medium containing 0.04% (w/v) glucose for 16 h and then diluted 1:500 in MM medium with either limiting nitrogen source (0.49 mM NH₄Cl and 0.5%, w/v, glucose) or limiting carbon source (9.5 mM NH₄Cl and 0.015%, w/v, glucose). The final cell densities of the cultures were 2.02 x 10^8 and 2.34 x 10^8 c.f.u. ml⁻¹, respectively. These cultures were then split into aliquots, one-half of which were amended with supernatants, supernatant extracts, or active fractions from HPLC at concentrations corresponding to those which had previously been shown to yield stimulation. Other media ingredients such as fresh nutrients were not added in this set of experiments, in order to test whether supernatant extracts themselves could supply carbon or nitrogen sources. During the following incubation at 37°C, OD_{590} and c.f.u. were monitored, and the experiment was repeated at least five times with each treatment. Upon addition of supernatant extracts or active fractions, the OD_{590} was observed to increase slightly immediately in comparison to the control. In carbon-limited cultures, the OD_{590} in the amended samples was 0.168 as compared to 0.164 (means of 12 measurements; standard deviations 0.017 and 0.016, respectively), and in nitrogen-limited cultures 0.184 as opposed to 0.177 (means of 11 experiments; standard deviations 0.022 in both cases). When the OD_{590} measurements were extended over 4 h, no further increases were observed. The c.f.u. counts in the treated cultures, however, were not higher than in the untreated cultures in any experiment at any time (samples were taken 40 and 170 min after addition) – actually, carbon-limited cultures displayed significantly lowered c.f.u. counts after supernatant extracts had been added, while nitrogen-limited cultures were unaffected (P < 0.05). In short, no growth of carbon- or nitrogen-limited cultures could be observed by addition of supernatant or supernatant extracts at concentrations which elicit the stimulation of growth described above. The slightly elevated OD_{590} data in treated samples can be explained by light absorption of the extracts themselves or by slight modification of the refractive index of the cells which is largely responsible for the light scattering in measurements of this type (Davey & Kell, 1996).

A series of experiments was performed in order to test the hypothesis that siderophore production might be responsible for the stimulatory activity of supernatants. To this end, the effect of supernatants of iron-limited cells on recovery was tested. Stationary phase cells (ZK126 grown overnight in fully supplemented MM medium with 0.04%, w/v, glucose) were diluted (1:100) in MM medium not amended with iron (with 0.04% NaCl and 0.015%, w/v, glucose), and growth was allowed to proceed into stationary phase as in the standard experiments. The addition of the supernatants of these (iron-starved) cells, however, had a smaller effect on lag times of recovering *E. coli* ZK126 than supernatants from fully supplemented cells of the strain. Also, neither excess addition of FeSO₄ (20 or 50 µM) nor addition of 2,3-dihydroxybenzoate (2–100 µM) had any effect on lag times of recovering cells (strains ATCC 25922 and ZK126 tested). Thus siderophore production can be excluded as an explanation for the autostimulatory effect.

**DISCUSSION**

In eukaryotes, cellular growth, behaviour and survival are known to be highly dependent on extracellular signals excreted by surrounding cells (Christensen *et al.*, 1998; Raff, 1992). Recently, such social phenomena are becoming more and more evident in a variety of aspects of prokaryote biology. Productivity and virulence of bacteria are now in many cases recognized to be subject to social control (Fuqua & Greenberg, 1998; Kaiser & Losick, 1993; Salmond *et al.*, 1995; Swift *et al.*, 1994). Growth and survival of prokaryotes, however, are still considered to be largely independent of the presence of sister cells. In this sense, the concept has persisted that microbial cells are independent units interacting mainly through means of competition and antibiotics. There are
many observations, though, which indicate that this concept is only partly true, and that indeed bacterial cells under certain conditions require the presence of sister cells (or products excreted by them) for growth and survival (Barer & Harwood, 1999; Kaprelyants & Kell, 1996; Kaprelyants et al., 1999; Kell et al., 1995). Publications on such observations are, however, relatively rare, and only in very few cases an explanation of these effects could be offered. For example, the lag times of cells aged for less than 8 h (‘early lag’) of Klebsiella aerogenes could be abolished by addition of filtrates from growing cells of the organism (Lodge & Hinshelwood, 1943), but the identity of the active chemical(s) has not been elucidated. The lag times displayed by aged cells (older than 9 h, and thus comparable to the aged cells employed in our study) could not be reduced by such addition (Lodge & Hinshelwood, 1943). In one well-characterized case, Micrococcus luteus produces a peptide which acts as both a cytokine and pheromone: stationary phase cells of M. luteus produce a protein which dormant cells of the organism require for growth (Davey & Kell, 1996; Kaprelyants & Kell, 1993; Kaprelyants et al., 1994; Mukamolova et al., 1998a, b). Similar observations were made for aged cultures of Mycobacterium tuberculosis and other Mycobacterium species (Mukamolova et al., 1998a), and related molecules appear to be widespread among the actinomycetes (Kell & Young, 2000).

In this paper, we report on the effects of the addition of spent medium on the regrowth of starved E. coli cells. Several phenomena could be observed: most strikingly, lag phases were strongly reduced by substances present in supernatants of stationary phase cells. Addition of raw supernatants at high concentrations, on the other hand, leads to inhibition of growth by reduction of growth rate and yield. A significant fraction of inhibitory material could be removed by ion-exchange treatment. Thus, the inhibitory effects are not merely a consequence of the lowered substrate concentration due to addition of supernatants, but a consequence of inhibitory products. The heat-labile inhibitory compound(s) might consist of peptides or proteins, and might constitute an inhibitory ‘signal’ excreted by the cells in stationary phase. These inhibitory compounds were, however, not characterized further, as the main aim of this study was to shed light on the stimulation of growth by excreted products.

**Stimulation of recovery of E. coli by supernatant addition**

Stationary phase cells of three strains of E. coli were shown to produce stimulatory substances, independent of the carbon source, the limiting nutrient or the age of cells. Susceptibility of cells to stimulation was also observed generally for all types of populations of the three strains tested, although stationary phase cells were stimulated more strongly than exponentially growing cells. Thus the production of and susceptibility to the stimulatory substance(s) are not restricted to specific strains or conditions. Preliminary screening of other bacterial species indicates that stationary phase cells of other *Escherichia* species (*Escherichia vulneris*, *Escherichia fergusonii*) and related species such as *Salmonella typhimurium*, *Enterobacter cloacae*, *Klebsiella pneumoniae* and *Citrobacter freundii* are stimulated significantly by supernatants obtained from stationary phase cells of *E. coli* ZK126 (data not shown). As the observed effect is not confined to the producer strain, any substance involved in the stimulation described here is not a true ‘pheromone’ as it is not species-specific.

In contrast to the events during resuscitation of *Micrococcus luteus*, growth of stationary phase cells of *E. coli* does not strictly require factors which are found in spent medium of the organism, because the number of recovering cells was independent of the presence of such factors. Hence the stimulation of growth of *E. coli* cells by supernatant addition is restricted to the reduction of lag times and increase in growth rates. Theoretically, such stimulation of growth could be brought about by amino acids, vitamins or similar nutritious material which could be leaking out of lysing cells. Our experiments with additions of supernatants or extracts to starving cells, however, did not reveal any potential role of this material as nutrient, because cells could not be observed to grow in the absence of added glucose. Also, it appears unlikely that nutritious substances would be present at significant concentrations in supernatants of stationary phase cultures grown in minimal medium, as the starving cells would be expected to be effective scavengers of these substances. Further, the chromatographic properties of the active principle were not similar to the above kinds of substance.

In our experiments, addition of supernatants or extracts of supernatants leads to significant shortening of lag phases of *E. coli*, particularly when the cells producing the substances and the cells tested were in prolonged stationary phase. This was observed both when apparent lag times (measured by optical density) or true lag times (monitored by c.f.u.) were determined during recovery. The discrepancies between apparent lag times and true lag times are simply a result of the sensitivity of the BioScreen growth analyser. We have determined that the lower limit of detection of the growth analyser for *E. coli* ZK126 and ATCC 25922 is $3 \times 10^7$ c.f.u. ml$^{-1}$, which corresponds to an OD$_{590}$ reading of 0.08 (blank readings with uninoculated MM medium gave readings between 0.065 and 0.07). Consequently, the first phases of recovery and growth of the diluted cultures (which have initial densities of 100–500 c.f.u. ml$^{-1}$) cannot be observed with the BioScreen. On the other hand, c.f.u. counts (which in our hands are sensitive down to 10 c.f.u. ml$^{-1}$) can monitor the development of cell numbers throughout the lag phases in our experiments, well before the limit of detection of the BioScreen is reached. The BioScreen, however, is a reliable source of growth data such as apparent lag times and can be employed to screen many substances or fractions quickly, which is an impossible task when relying on c.f.u. determinations.
The fact that the stimulatory effects of supernatants could not be observed when cells were grown in complex media is possibly due to a combination of the fact that the stimulatory compound is present in yeast extract and/or that lag times generally are minimal in the presence of a complete set of amino acids and vitamins. This is demonstrated by the fact that addition of yeast extract or Casamino acids at extremely low concentrations (0.0003%, w/v) shortened lag times of E. coli ZK126 significantly.

While untreated cell suspensions of freshly diluted stationary phase cultures displayed a slight decrease in c.f.u. during the lag phase, identical suspensions amended with supernatant components do not display this decrease. In theory, this effect could be indicative of the fact that a part of the population is transiently injured, and that this injury is avoided or alleviated in the presence of supernatant components. The fact, however, that the number of recovering cells in our experimental system is not influenced by the presence of supernatant components does not support this possibility. We conclude that substances present in the supernatants trigger or facilitate cell division during recovery, especially after prolonged stationary phase, rather than protect the cells from any deleterious effect of refeeding or dilution in fresh media. It is possible that a certain proportion of cells in all freshly diluted suspensions is (at least transiently) adversely affected by the process of handling and dilution. The discrepancy in the development of viable cells observed in the presence and absence of supernatant components might then merely be due to the fact that in the treated suspensions growth commences immediately, and thus compensates and obscures the decrease in c.f.u. in these samples. In summary, we hypothesize that E. coli produces a (or several) signal compound(s) or growth regulator(s) which stimulates (or stimulate) recovery from the stationary phase.

Characterization of stimulatory substances in E. coli supernatants

In order to elucidate the identity of the stimulatory compound, a range of chemicals was tested in their effect on lag times and growth rates. The chemicals were selected based on the possibility that they might be found in supernatants or that they might stimulate growth of E. coli. In the literature, several substances have been reported to be secretion products of the organism in the stationary phase. Amongst them are reduced glutathione (Loewen, 1979; Owens & Hartman, 1986) and uracil, xanthine and hypoxanthine (Rinas et al., 1995). The presence of uracil and xanthine in stationary phase supernatants could be confirmed by electrospray MS. The autostimulatory activity, however, could not be duplicated by addition of any of these substances, alone or in combination, nor by any other chemicals tested. In particular, it was ascertained that a complete set of vitamins is not sufficient to mimic the supernatant effect on recovery. It is, however, possible that the autostimulatory substance constitutes a rare form of vitamin (or a combination of growth factors) which has not been included in the literature because it is not essential for growth.

So far, the active compound present in supernatants of E. coli has not been identified. To our knowledge, we have excluded all the substances that have been reported to be present in supernatants or which might stimulate growth significantly. Also, we have excluded a range of possible signal substances such as peptides or other ionic or zwitterionic substances (such as amino acids) by demonstrating activity in extracts after heat and protease treatment, or after ion-exchange treatment, respectively. Due to the lack of extraction in ethyl acetate, we believe that we have also excluded N-acylated homoserine lactones.

Unfortunately, no way of selectively extracting the active material could be found. Consequently it has not been possible to isolate large amounts of the active compound at high purity. The possibility remains that a mixture of substances is responsible for the growth stimulation observed. This is, however, unlikely because activity could be detected in a single fraction after several consecutive rounds of fractionation on different HPLC columns. The active material was found to co-migrate with only two identifiable substances, namely uracil and 5-hydroxy-methyl-uracil; in fact, high levels of uracil could be detected in all supernatant extracts and in all active fractions. As stated above, however, no significant stimulation was observed after addition of uracil or 5-hydroxy-methyl-uracil. A range of other uracil derivatives and their isomers displayed HPLC retention times differing strongly from those of active fractions. No other substance could be identified in the active fractions, even when employing highly sensitive electrospray MS (LC-MS). Thus the chemical identity of the substance produced by E. coli, and stimulating stationary phase cells of itself and other bacteria, still remains to be elucidated. Given the large number of genes of unknown function in the E. coli genome (Blattner et al., 1997; Hinton, 1997), we may suppose that many more metabolites of this type will be identified.

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