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**SUMMARY**

DNA breakdown was detected 3 to 4 min. after addition of colicin E2 to sensitive cells; inhibition of cell division followed 5 to 10 min. later, but inhibition of DNA synthesis was observed only after several more minutes. Adsorption of E2, which takes place even at $4^\circ$, led to the formation of a specific surface complex (I). Complex I did not promote DNA breakdown. We suggest that the transition from this complex to a surface complex (II) which promoted DNA breakdown depended upon several factors which include temperature, concentration of E2, specific membrane proteins and, under certain conditions, high concentrations of extracellular KH$_2$PO$_4$. The formation of complex II did not depend on concomitant DNA or protein synthesis. The continued promotion of DNA breakdown by complex II and its associated nuclease was blocked by inhibition of energy metabolism. In addition, the removal of E2 from the cell surface by trypsin treatment during the early stages of the process greatly decreased the rate of DNA breakdown. E2-induced DNA breakdown, which appears to commence from a limited number of chromosomal sites, proceeded normally in UVr−, RecB−, RecC−, Hsr−, Hss−, PolA− and in several tsDNA replication mutants.

**INTRODUCTION**

Colicins appear to act by inducing physicochemical changes in the membrane, which promote specific intracellular changes in the sensitive cell. Evidence has been presented that: (1) Colicin E2 adsorbs to, but does not penetrate, the surface layer of sensitive bacteria (Maeda & Nomura, 1966); (2) E2 does not induce detectable changes in the permeability of the membrane (Nomura, 1964); (3) bacteria may be rescued from the lethal effects of E2 for several minutes after adsorption, by digestion of the extracellular colicin with trypsin (Reynolds & Reeves, 1963; Maeda & Nomura, 1966); (4) mutants which still adsorb E2 but remain insensitive to its presence appear to have altered membranes (Holland et al., 1970; Samson & Holland, 1970).

To explain the mode of action of colicin, Nomura (1964) proposed that specific transmission systems must exist in the membrane which connect the extracellular colicin to its intracellular target. Such a transmission system or pathway should therefore have specific and identifiable intermediates between the initial colicin-surface complex and the final modified cell component. Since a primary effect of E2 action is rapid degradation of DNA (Nomura, 1963; Holland, 1968), we assumed for the
purposes of this investigation that a final product in the E2 pathway is a specific DNA–nuclease complex which promotes rapid degradation of DNA. We further assumed that the release of soluble breakdown products of DNA provides a reasonable measure of formation of this complex.

In this study, attempts have been made to establish more clearly the presence of intermediate steps in colicin E2 action and to determine the nature of the processes which affect their formation. The mechanism of DNA breakdown itself appears to proceed via a specific mechanism unrelated to degradative systems involved in known repair or recombination processes.

METHODS

**Strains.** *Salmonella typhimurium* LT2 strain 906, a colicinogenic strain carrying the E2 (P9) factor, was the source of colicin E2 (Holland, 1968). *Escherichia coli* K12 laboratory strain 206 (Hfr thy− his− λ−) was used as the colicin-sensitive strain. The bacteria were normally grown in Difco nutrient broth supplemented with appropriate amounts of thymine.

**Colicin E2.** Colicin was used as a crude sterile lysate prepared from a mitomycin C induced culture of strain 906 as described previously (Hill & Holland, 1967). This material contained no antibacterial activity in addition to E2 (Hill & Holland, 1967) and behaved in all respects like highly purified preparations of E2 (Holland, 1968). For the assay of colicin, serial dilutions of the E2 preparations were spotted on nutrient broth agar plates previously seeded with strain 206. After 14 h. at 37° the highest dilution giving the minimal inhibition of growth was determined and the reciprocal of this dilution taken as the colicin titre. Preparations usually contained 2 × 10^5 arbitrary units of colicin E2/ml. and were stable for several months at 4°. Concentrations of 0.2 to 0.4 units of E2/ml./10^8 bacteria normally killed 50% at 37°.

**Bacterial counts.** For total bacterial counts, 0.05 ml. culture samples were diluted into non-pyrogenic N-saline (Polyfusor, Boots Pure Drug Co. Ltd, Nottingham) and counted in duplicate in a model F Coulter counter at densities around 2 × 10^4 bacteria/ml.

**Determination of DNA breakdown in treated cultures.** Strain 206 was aerated in nutrient broth + 2.5 μg. thymine/ml., until it reached 5 × 10^7 bacteria/ml. The culture was diluted tenfold into the same medium containing 40 μCi methyl-[3H]thymine (21.8 Ci/mmole)/ml.; incubation was continued for four to five generations to about 2 × 10^8 bacteria/ml. The culture was washed twice in nutrient broth + 25 or 100 μg. cold thymine/ml. and finally suspended in this medium at about 2 × 10^8 bacteria/ml. The labelled culture was then incubated at 37° for 10 min., colicin E2 was added and samples (0.2 ml.) were removed at intervals, mixed with 0.2 ml. ice-cold 10% trichloroacetic acid. After standing 60 min. on ice, the samples were analysed for cold and hot acid-soluble radioactivity as described by Howard-Flanders & Theriot (1966). Radioactive samples were counted with a Packard Tri-Carb scintillation counter.

**Measurement of DNA synthesis.** Incorporation of [3H]thymine into acid-precipitable material was used as a measure of DNA synthesis in treated cultures as described previously (Holland, 1968).

**Materials.** Mitomycin C, trypsin (type III, twice crystallized) and trypsin inhibitor (Soybean, type 1-S, twice crystallized) were obtained from the Sigma Chemical Company. Nalidixic acid was a gift from Bayer Products, Surbiton-upon-Thames, Surrey.
Induction of DNA breakdown by colicin E2

RESULTS

Kinetics of DNA breakdown and inhibition of cell division induced by colicin E2

Previous studies indicated that colicin E2 causes inhibition of cell division in addition to induction of rapid DNA breakdown (Holland, 1968). To determine which was the primary effect, the kinetics of the appearance of these two consequences of colicin action were measured in cultures of Escherichia coli strain 206 prelabelled with [3H]thymine and exposed to different concentrations of colicin E2. In other experiments the effect of E2 on division and DNA synthesis were measured simultaneously. Fig. 1 presents a composite figure of some typical results. The colicin concentration used in Fig. 1 (0.5 units/10^7 bacteria/ml.) was normally sufficient to kill 99% of the bacteria; DNA breakdown was first detected at 20 min. whilst the increase in bacterial number continued at the normal rate until minute 30. Thereafter total biomass (turbidity) increased at the normal rate for at least 25 min. and the bacteria became filamentous. DNA synthesis also continued at the normal rate for at least
50 min. after adding E2. Similar experiments were carried out with several different concentrations of E2 and, although the time of onset of DNA breakdown varied (see also Fig. 2), the result was always the same: DNA degradation began 5 to 10 min. before inhibition of division was observed. Thus inhibition of bacterial division did not appear to be a primary effect of colicin E2 action nor did it result from inhibition of DNA synthesis. The effect of E2 upon the division machinery therefore remains obscure.

Factors affecting initiation of DNA breakdown by E2

Effect of E2 concentration on DNA breakdown in growing bacteria. A striking effect of colicin E2 action observed in these studies was that the time of commencement of DNA breakdown in exponentially growing cultures depended on the amount of E2 added. A similar effect was observed by Nomura (1964). Labelled cultures of

![Graph showing effect of colicin E2 on growth and division of exponentially growing cultures of Escherichia coli strain 206 in nutrient broth treated with E2 at time zero.](image-url)

Fig. 3. Effect of colicin E2 on growth and division of exponentially growing cultures of *Escherichia coli* strain 206 in nutrient broth treated with E2 at time zero. Figures against the curves are the colicin concentrations added/10^7 bacteria; (a) extinction (E) rate changes, (b) total bacterial counts.

*Escherichia coli* strain 206 growing exponentially in nutrient broth were treated with various concentrations of E2 and the kinetics of DNA breakdown measured. Fig. 2 shows that, over a 100-fold range of E2 concentration, DNA breakdown was detected as early as 4 min. or as late as 50 min. (equivalent to 1.5 generations). Similar results were obtained even when adsorption of E2 was done at 4°C and the bacteria washed to remove excess E2 before raising the temperature to 37°C. Measurement of the survival
kinetics of bacteria treated with various E2 concentrations also showed that the delayed induction of DNA breakdown by small amounts of E2 was not due to slow adsorption of the colicin. When the effect of E2 upon bacterial division was measured at different colicin concentrations, similar results to those on the induction of DNA breakdown were obtained. Addition of E2 was followed by normal increases in bacterial number until, abruptly, the rate of increase changed at a time determined in some way by the amount of E2 present (Fig. 3).

Table 1. Effect of colicin E2 multiplicity on induction of DNA breakdown

Exponential broth cultures of *Escherichia coli* strain 206 (3 × 10^7 bacteria/ml.) treated at time zero with colicin E2 at 37° as in Fig. 2. Rates of DNA breakdown calculated from the slopes of the curves shown in Fig. 2.

<table>
<thead>
<tr>
<th>E2 concentration</th>
<th>DNA breakdown</th>
<th>Rate % sol. [^H]thymine released/min.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Units/10^7 bacteria</td>
<td>Estimated no. molecules/bacterium</td>
<td>Bacterial survival (after 60 min.)</td>
</tr>
<tr>
<td>0.025</td>
<td>25</td>
<td>53 %</td>
</tr>
<tr>
<td>0.05</td>
<td>50</td>
<td>23 %</td>
</tr>
<tr>
<td>0.1</td>
<td>100</td>
<td>16 %</td>
</tr>
<tr>
<td>0.25</td>
<td>250</td>
<td>0.4 %</td>
</tr>
<tr>
<td>0.5</td>
<td>500</td>
<td>0.1 %</td>
</tr>
<tr>
<td>5</td>
<td>5,000</td>
<td>0.01 %</td>
</tr>
<tr>
<td>50</td>
<td>50,000</td>
<td>0.001 %</td>
</tr>
</tbody>
</table>

This effect of concentration of colicin E2 on the pattern of DNA breakdown in sensitive bacteria was investigated in many experiments, and the relationship between E2 concentration, the timing of DNA breakdown and the subsequent rate of this process is examined in Table 1, which also shows the bactericidal effect of E2 and presents each colicin concentration as the number of E2 molecules/cell. This can be estimated since the molecular weight of E2 is known (Herschman & Helinski, 1967) and the specific activity of purified E2 (1000 units/μg. protein) was determined previously (Holland, 1968). On the basis of these calculations, the maximum number of E2 molecules adsorbed/organism was 2000, which is in good agreement with the figure of 2000 to 3000 E2 receptors calculated by Maeda & Nomura (1966). The results in Table 1 show that when sufficient E2 was added to saturate all receptors there was a minimum delay of 4 min. before breakdown was detected. At E2 concentrations sufficient to kill only 47% of the bacteria, less than 5% of the receptors should be occupied, and in this case breakdown was only detected at 40 min. These results indicate that, with exponentially growing bacteria, very few of the initial receptor colicin complexes promoted a lethal interaction between colicin and membrane. At high E2 concentrations, at least one lethal hit was quickly attained but at subsaturation concentrations growth and division appeared to be needed before a lethal hit, as expressed by the induction of DNA breakdown, took place. The data in Table 1 also show that, in contrast to its effect upon the timing of breakdown, increasing the concentration of E2 did not greatly influence the rate of DNA breakdown. Although E2, when added in excess, stimulated higher rates of breakdown, at subsaturation concentrations the rate of degradation was virtually independent of the E2 concentration, particularly when the fraction of bacteria killed by the colicin is taken into account.
These results do not rule out the possibility that DNA breakdown was initiated by the co-operative effect, upon the membrane, of many colicin molecules, i.e. a multihit mechanism. Since, however, there is evidence that the killing action of colicin is a single-hit process (Jacob, Siminovitch & Wollman, 1952; Nomura, 1963; Shannon & Hedges, 1967), we favour the alternative interpretation that DNA degradation is initiated by an all or none process, and that in growing cultures the main effect of increasing the E2 concentration was to increase the probability of an early single hit which then initiated DNA breakdown (or inhibition of division) at the maximum rate.

![Figure 4](image-url)

**Effect of inhibition of DNA and protein synthesis upon E2-induced DNA breakdown.** Energy is required for the conversion of the initial colicin E2–cell complex into a state from which viable bacteria can no longer be recovered by trypsin action (Reynolds & Reeves, 1969). Nomura & Maeda (1965) have shown that no DNA breakdown takes place in bacteria treated with E2 in the presence of 2,4-dinitrophenol. In addition, the results described in the previous section indicate that, particularly at low concentrations of E2, some growth was required to initiate DNA breakdown. The question whether
protein synthesis was needed for the formation of, for example, a specific colicin-surface complex or for the synthesis de novo of a specific DNase was therefore studied. Cultures prelabelled with [3H]thymine were exposed to different concentrations of colicin E2 with chloramphenicol (200 µg./ml.). As shown in Fig. 4, neither the length of the prebreakdown lag nor the subsequent rate of breakdown were affected. In a few experiments chloramphenicol did produce some decrease in rate of breakdown, particularly with low E2 concentrations, but this effect was not reproducible and may have been due to disturbance in the growth of the bacterial surface induced by chloramphenicol (see Rothfield & Pearlman-Kathencz, 1969). Amino acid starvation before and during treatment with colicin E2 was also without effect upon the DNA breakdown pattern. These results clearly showed that the promotion of DNA breakdown by E2 did not require the synthesis de novo of a DNase. In addition, these results showed that the lag period before the initiation of DNA breakdown by low E2 concentrations was apparently not associated with the synthesis of any cellular protein component.

The effect of inhibition of DNA synthesis upon colicin E2 action is also shown in Fig. 4. Labelled cultures of *Escherichia coli* strain 206 were exposed to E2 after treatment in various ways with nalidixic acid (NAL). NAL did not inhibit DNA breakdown induced by high or low E2 concentrations. Furthermore, experiments discussed below showed that E2-induced DNA breakdown proceeded normally at 42° in mutants defective in ability to replicate DNA at high temperature. Promotion of DNA breakdown by E2 could therefore be initiated at any point in the DNA replication cycle and degradation did not depend upon continued synthesis. Fig. 4d also shows that pretreatment with NAL under conditions which promoted more chromosomal replication forks per bacterium (see Pritchard, Barth & Collins, 1969) did not stimulate the rate of DNA breakdown on subsequent addition of colicin E2. This result suggests that DNA breakdown in E2-treated bacteria is not specifically triggered from the replication point on the chromosome.

*Induction of DNA breakdown by E2 in non-growing bacteria.* The results described above indicated that, particularly at low E2 concentrations, initiation of DNA breakdown depended on growth or accumulation of some cellular component. This interpretation predicted that, in non-growing bacteria (e.g. in buffer suspensions), high E2 concentrations should initiate breakdown more or less normally, but small amounts of E2 should initiate breakdown only poorly. Contrary to expectation, as shown in Fig. 5, extensive breakdown of DNA was observed with both high and low E2 concentrations. In the latter case the delay in initiation of DNA breakdown observed in growing bacteria was not found, and soluble [3H]thymine was detected in all cultures within a few minutes of adding E2. DNA breakdown in buffered suspensions quickly became linear in contrast to the exponential rates found with growing bacteria. Nevertheless, initial rates were usually at least half those for broth suspensions treated with comparable E2 concentrations. Again, as found with growing bacteria, the rate of DNA breakdown in buffer suspension was not very dependent upon the colicin concentration: over at least a 100-fold range of E2 concentration the rate of DNA breakdown increased at most sixfold.

The long delays in initiation of DNA breakdown observed in broth cultures were thus due to some effect unique to growing bacteria and were not essential for E2 action. Possible explanations for the effect of E2 multiplicity upon the timing of initiation of DNA breakdown will be discussed below.
Effect of phosphate on initiation of DNA breakdown in buffer suspensions. Bacteria in tris + HCl buffer showed very little DNA breakdown in the presence of colicin E2 although adsorption and killing by E2 were normal. Addition of KH₂PO₄ to treated bacteria in tris + HCl buffer stimulated DNA breakdown, and rates comparable to those in phosphate buffer were obtained with 10⁻⁸ M-phosphate. As shown in Fig. 6, KH₂AsO₄ was quite effective in stimulating breakdown in tris buffer, in contrast to NaCl, KCl or lower concentrations of KH₂PO₄. The stimulation produced by added phosphate was usually six- to eightfold although in a few experiments DNA breakdown in tris buffer alone was more extensive, and the effective stimulation by added phosphate was only two- to threefold.

In an attempt to determine which particular step in the E2 pathway was affected, the kinetics of inactivation of bacteria treated with E2 in tris buffer were examined and found to be similar whether phosphate was present or not. Phosphate was not
Induction of DNA breakdown by colicin E2

therefore required for irreversible binding of colicin E2. Similarly, phosphate was not required for the degradative process itself since removal of phosphate from the suspending medium after the onset of breakdown did not affect further degradation (Table 2). Several experiments, however, indicated that exogenous phosphate was required for formation of the specific colicin-envelope complex which rapidly initiated phosphate-independent DNA breakdown. Table 2 also shows: (1) suspension of bacteria in phosphate buffer did not stimulate DNA breakdown when exogenous phosphate was removed before addition of E2; (2) phosphate starvation of bacteria for at least 30 min. before addition of E2 did not affect subsequent phosphate stimulation of DNA breakdown; (3) after preadsorption of E2, bacteria showed increased DNA breakdown immediately upon addition of phosphate; treatment of organisms with E2 in the presence of phosphate at 4° did not stimulate subsequent DNA breakdown when phosphate was removed before incubation at 37°.

Table 2. Effect of phosphate on initiation of DNA breakdown by colicin E2

<table>
<thead>
<tr>
<th>Adsorption temperature</th>
<th>Presence of phosphate</th>
<th>DNA breakdown* % total DNA released</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expt. 1</td>
<td>37°</td>
<td>Throughout</td>
</tr>
<tr>
<td></td>
<td></td>
<td>During adsorption and for further 10 min.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>After adsorption only</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ten min. before addition of E2 only</td>
</tr>
<tr>
<td>Expt. 2</td>
<td>4°</td>
<td>Throughout</td>
</tr>
<tr>
<td></td>
<td></td>
<td>After adsorption only</td>
</tr>
<tr>
<td></td>
<td></td>
<td>During adsorption only</td>
</tr>
</tbody>
</table>

* The amount of breakdown shown is that obtained after 90 min. postadsorption incubation at 37°. When phosphate was present throughout, the amount of breakdown obtained was 25 to 40 % over several experiments; without phosphate, between 3 and 8 % of DNA was usually broken down in 90 min. at 37°.

In summary, the above results indicate that, in bacteria suspended in tris buffer, the initial colicin–cell complex was only poorly active in initiating DNA breakdown. In the presence of exogenous phosphate, a temperature-dependent process then converted the colicin–surface complex into an active form which promoted DNA breakdown.

Mechanism of DNA breakdown in bacteria treated with colicin E2

The degradative process requires energy. Since earlier studies showed that energy was required to promote colicin E2 action, the possibility was examined that the DNA breakdown process itself might be energy-dependent. As shown in Fig. 7, 2,4-dinitrophenol (DNP) immediately and dramatically decreased the rate of breakdown by treated bacteria. The effect was reversible and upon removal of DNP, breakdown immediately resumed at the original rate (Fig. 7b). Fig. 7a shows that DNP was equally effective in suppressing breakdown when added at late times. The breakdown
process itself is therefore inhibited by DNP. Energy metabolism in E2-treated bacteria was also inhibited at various times by cyanide \((2 \times 10^{-8} \text{M})\) or colicin K (an uncoupler of oxidative phosphorylation; see Levinthal & Levinthal cited by Luria, 1964) and in each case the rate of breakdown was greatly decreased, confirming that energy is required to maintain the colicin–membrane–DNA complex in a state propitious for breakdown, or alternatively that the E2-specific nuclease itself is ATP-dependent.

**Fig. 7.** Effect of inhibition of energy metabolism upon colicin E2-induced DNA breakdown. (a) Exponential cultures \((2 \times 10^7 \text{ bacteria/ml.})\) of *Escherichia coli* strain 206 in nutrient broth at 37° were treated with colicin E2 \((2 \text{ units/10}^7 \text{ bacteria})\) at time zero, \(\bullet-\bullet\). At the intervals indicated by the curves \(\circ-\circ\), 2,4-dinitrophenol (DNP) \((\text{final concentration } 2 \times 10^{-4} \text{M})\) was added. Release of acid-soluble \(^{3}H\)thymine was then measured as described in Methods. (b) Strain 206 in nutrient broth treated with E2 \((2 \text{ units/10}^7 \text{ bacteria/ml.})\). Separate fractions were treated with DNP at 20 min. and at 40 min.; duration of DNP treatment is indicated by broken lines. At the times indicated by arrows, DNP was removed by centrifugation and incubation of bacteria in nutrient broth resumed. The control culture \((\bigcirc)\) was treated with DNP only.

**Effect of trypsin on E2-induced DNA breakdown.** Nomura & Nakamura (1962) previously showed that macromolecule synthesis, completely suppressed by colicin K, was quickly resumed after addition of trypsin. The membrane alteration induced by colicin K is therefore only maintained in its presence and reverts to normal when it is removed. Since a primary consequence of colicin E2 action is the promotion of DNA breakdown, we determined whether nuclease activity also depended completely upon the continued presence of E2. In the presence of small amounts of E2 \((\text{Fig. 8a})\), addition of trypsin to *Escherichia coli* strain 206 greatly inhibited further breakdown even when added at 30 min.; during the early lag period it completely
Induction of DNA breakdown by colicin E2

suppressed DNA breakdown. This latter effect provides further confirmation that the formation of an active surface-E2 complex, which promotes DNA breakdown, is delayed at low E2 concentrations. At saturating concentrations of E2, trypsin was most effective when added up to 10 min. after initiation of breakdown and, as shown in Fig. 8b, was progressively less effective when added at later times. Phase-contrast

![Graphs and images showing the effect of trypsin on colicin E2-induced DNA breakdown.](image)

Fig. 8. Effect of trypsin on colicin E2-induced DNA breakdown. Cultures of Escherichia coli strain 206 were labelled in the usual way, washed and suspended in nutrient broth, pH 7.8 (a, b), or in tris buffer, pH 7.8 (c, d), containing 10^{-8} M-KH_{2}PO_{4}, 10^{-8} M-MgSO_{4} and 10^{-4} M-CaCl_{2}. Solid lines represent the primary cultures treated with colicin E2, broken lines denote samples removed from primary culture and treated with trypsin (1 mg./ml.). (a) Nutrient broth culture treated with colicin E2 (0.2 units/10^7 bacteria) at zero time and trypsin treatment done at the times indicated. (b) Nutrient broth cultures treated with E2 (2 units/10^7 bacteria). (c) Bacteria suspended in tris phosphate buffer and treated with E2 (8 units/10^7 bacteria). (d) Preadsorption of E2 (10 units/10^7 bacteria) for 45 min. at 4^\circ, cultures then centrifuged and deposit suspended in tris phosphate buffer before trypsin treatment.
microscopy of such cultures showed that trypsin was also most effective in preventing filament formation when added early.

Three possible explanations of these features of trypsin inhibition are:

1. initiation of DNA breakdown is not synchronized in all of the population, and trypsin only affects those bacteria in which E2-induction of DNA breakdown is not yet established;
2. colicin E2 becomes, with time, increasingly inaccessible to the trypsin molecule;
3. the specificity of the degradative process is progressively lost as breakdown becomes extensive.

In an attempt to test the first possibility, the effect of trypsin on E2 action in non-growing bacteria was examined in the hope that any asynchrony arising from some aspect of growth would be minimal. The effect of trypsin upon DNA breakdown was, however, very similar to that obtained with growing cultures (Fig. 8c). Early addition of trypsin completely abolished subsequent breakdown, and the trypsin effect was progressively lost as breakdown proceeded. Nevertheless, even when breakdown was well under way, addition of trypsin still produced a significant, although sometimes delayed, inhibitory effect. Similar results were obtained when adsorption of E2 was completed at 4°C and non-adsorbed E2 removed by centrifugation before shifting the treated bacteria to 37°C, conditions under which colicin is irreversibly adsorbed although at a slower rate. It therefore seems unlikely that trypsin affects only those bacteria in which DNA degradation has not yet been initiated.

Several relevant control experiments were also made: (a) under the cultural conditions normally used, trypsin treatment had no effect upon mitomycin C-induced DNA breakdown; (b) with E2-treated bacteria, subsequent removal of trypsin did not lead
Induction of DNA breakdown by colicin E2

to a resumption of the normal rate of DNA breakdown, and when equimolar amounts
of trypsin and trypsin inhibitor were added breakdown was not inhibited; (c) incubation
of bacteria with trypsin followed by removal of the enzyme and addition of colicin E2
had no inhibitory effect upon subsequent DNA degradation.

The observed effects of trypsin upon colicin E2 action appear therefore to be the
result of the proteolytic digestion of the surface-bound colicin and not to any non-
specific effects. On the basis of all these results it seems reasonable to conclude that,
at least at early times, removal of E2 by trypsin treatment can have a direct inhibitory
effect upon the E2-specific nuclease.

Effect of E2 on Escherichia coli K12 mutants defective in nuclease functions. A number
of strains of Escherichia coli mutants possibly lacking nuclease functions (see Table 3)
have been studied. All of them were sensitive to colicin E2 and showed normal E2-
induced DNA breakdown at 37° (or at 42° in the case of temperature-sensitive DNA-
replication mutants) including strains which lack an ATP-dependent nuclease (Buttin

Table 3. Escherichia coli mutants tested for colicin E2-sensitivity and
E2-induced DNA breakdown

The tsDNA mutants, E279 and E613, which show immediate cessation of DNA synthesis
on shift to 42° were kindly provided by Dr J. Wechsler. The tsDNA mutant 16 was kindly
provided by Mr H. G. Nandadasa, this strain and all the other ts mutants tested show
delayed inhibition of DNA synthesis on shift to high temperature.

<table>
<thead>
<tr>
<th>E. coli mutant strain</th>
<th>Genotype or phenotype</th>
<th>Linked to chromosomal locus</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>JG 64</td>
<td>polA−</td>
<td>metE</td>
<td>Gross &amp; Gross (1969)</td>
</tr>
<tr>
<td>4K</td>
<td>his−</td>
<td>serB</td>
<td>Glover (1970)</td>
</tr>
<tr>
<td>5K</td>
<td>ksr−</td>
<td>serB</td>
<td>Hubacek &amp; Glover (1970)</td>
</tr>
<tr>
<td>KMBL 90</td>
<td>urrB</td>
<td>gal</td>
<td>Van de Putte, Van Sluis, Van Dillewijn &amp; Rösch (1965)</td>
</tr>
<tr>
<td>JC 4457</td>
<td>recB</td>
<td>argA</td>
<td>Clark (1967)</td>
</tr>
<tr>
<td>KMBL 243</td>
<td>recC</td>
<td>argA</td>
<td>Van de Putte, Zwenk &amp; Rösch (1966)</td>
</tr>
<tr>
<td>E279</td>
<td>tsDNA</td>
<td></td>
<td>From Dr J. Wechsler</td>
</tr>
<tr>
<td>E613</td>
<td>tsDNA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>FA 21</td>
<td>tsDNA</td>
<td>str</td>
<td>Fangman &amp; Novick (1968)</td>
</tr>
<tr>
<td>16</td>
<td>tsDNA</td>
<td>ilv</td>
<td>From Mr H. S. Nandadasa</td>
</tr>
<tr>
<td>T 83</td>
<td>tsDNA</td>
<td>ilv</td>
<td></td>
</tr>
<tr>
<td>T 42</td>
<td>tsDNA</td>
<td>malB</td>
<td>Hirota, Ryter &amp; Jacob (1968)</td>
</tr>
<tr>
<td>T 46</td>
<td>tsDNA</td>
<td>ilv</td>
<td></td>
</tr>
</tbody>
</table>

& Wright, 1968) and restriction-deficient strains, which probably lack an ATP-
dependent endonuclease (Meselson & Yuan, 1968). As found previously, colicin E2
also promoted normal breakdown in a Uvr− mutant (Holland, 1967) although
mitomycin C-induced DNA breakdown was greatly decreased compared to wild-type
strains (Fig. 9). Uvr− mutants appear to lack the capacity to excise thymine dimers
(Howard-Flanders & Theriot, 1966) and to repair mitomycin C-induced damage
(Boyce & Howard-Flanders, 1964), and since such mutants show normal sensitivity to
colicin E2 it appears most unlikely that E2-directed DNA breakdown involves any
of the normal repair nucleases. Several mutants that were temperature-sensitive for
DNA replication were also examined, including mutants possibly defective in either
replication or the initiation of replication (Table 3), but all showed normal E2-induced
DNA breakdown at high temperature.
DISCUSSION

Factors affecting initiation of DNA breakdown by colicin E2

On the basis of our studies and those of other workers, we envisage the following scheme for the pathway of colicin E2 action:

bacteria + E2 → Surface complex I → Surface complex II (promotes DNA breakdown).

The initial adsorption step is a two-stage process (Reynolds & Reeves, 1969) leading to an irreversible state, complex I. This appears to require magnesium (Reynolds & Reeves, 1969) but not energy and is completely reversible by trypsin treatment. The specificity for complex I is shared by colicin E2 and E3 (Maeda & Nomura, 1966), although the two colicins have quite different intracellular effects. The chemical nature of the primary receptor is not known but it may well be analogous to a phage receptor. Some intermediate stages, presumably involving interaction between E2 and specific membrane proteins, may then be involved before the pathway culminates in the formation of complex II, a conformational state of the membrane which promotes rapid degradation of DNA and inhibition of cell division.

The formation of complex II is a complicated process, the efficiency and timing of which varies in relation to the colicin concentration and to the physiological conditions of the organism. Nevertheless, the initiation of DNA breakdown always appears to be an all-or-none process, with increasing concentrations of colicin E2 hastening initiation of breakdown rather than its ultimate rate. Although this multiplicity effect of E2 is not understood, it is clear that synthesis of neither protein nor DNA is required for the development of the final surface complex which triggers DNA breakdown. Differential growth or extension of various surface layers in the presence of small concentrations of E2 may, however, be important in determining the timing of complex II formation, but it cannot be an absolute requirement since in non-growing bacteria low doses of E2 still trigger DNA breakdown without appreciable delay.

It is still not clear whether complex II formation is energy-dependent. Promotion of E2 action is temperature-dependent and is inhibited by 2,4-dinitrophenol, but the finding that the DNA breakdown process itself requires energy makes it difficult to establish whether any of the intermediate steps are also energy-requiring.

Genetic analyses have previously shown that mutants refractory to colicin E2 may be blocked at several postadsorption steps (Hill & Holland, 1967; Nomura & Witten, 1967; Nagel de Zwaig & Luria, 1967) probably by alteration of specific proteins (Nomura & Witten, 1967; Holland, 1968; Nagel de Zwaig & Luria, 1969). Recent biochemical studies have shown that refractory mutants may lack or contain altered membrane proteins (Samson & Holland, 1970; C. Schnaitman, personal communication), suggesting that membrane or surface proteins may participate in complex II formation. Colicin E2 may therefore not interact directly with a membrane-bound nuclease but may act indirectly through the intermediacy of specific membrane proteins, possibly as suggested by Changeux & Thiery (1967) via conformational changes of repeating membrane protomers, leading to a final specific protein or membrane configuration which then promotes nuclease action. The finding that resting bacteria in tris buffer require added phosphate for efficient initiation of DNA breakdown but not for adsorption of colicin, provides further evidence for the presence of intermediate steps in E2 action.
Induction of DNA breakdown by colicin E2

We have suggested above that the final formation of complex II simply involves a series of physical changes in the cell membrane in response to the fixation of colicin. Several lines of evidence suggest that the ultimate biochemical changes induced by different colicins, including E2, derive from the altered properties of membrane-bound proteins or enzymes in situ and not from the activities of, for example, enzymes displaced from the membrane and acting through the cytoplasm. Thus the effects of colicin K upon energy metabolism and macromolecular synthesis are completely reversed upon subsequent removal of the colicin from the cell surface by trypsin treatment (Nomura & Nakamura, 1962). Moreover, although modification of 30S ribosomal subunits is induced by colicin E3 fixation to sensitive bacteria, extracts from such bacteria are incapable of modifying normal ribosomes in vitro (Konisky & Nomura, 1967). DNA breakdown induced by colicin E2 can, at least in the early stages, be halted or decreased by digestion of the adsorbed colicin with trypsin. Thus it appears that the membrane conformation induced by colicin E2 may return, as in the case of colicins E1 and K, to its original state when colicin is removed. The failure of trypsin to inhibit DNA breakdown effectively when added at later times is unexplained; asynchronous initiation of DNA breakdown within the population appears to be excluded as a major factor. More likely, the specificity of the degradative process is soon lost or the colicin becomes inaccessible to trypsin as DNA breakdown proceeds and division is halted.

Specificity of E2-induced DNA breakdown

With the formation of complex II the altered membrane induces rapid degradation of DNA and, as a consequence or as an independent effect, the inhibition of division. The degradative process appears to be highly specific to the Escherichia coli chromosome; in T4-infected bacteria colicin E2 does not promote phage DNA breakdown (Nomura, 1963), and invading λ DNA is not degraded in E2-treated bacteria and appears to replicate normally (unpublished results). Also, since DNA replication may continue in E2-treated bacteria without loss of newly synthesized strands until at least 10% of the chromosome is degraded (Fig. 1), it appears probable that breakdown is initiated at a few strictly localized sites, possibly at the replication fork or at the chromosomal origin, both of which have been shown to be membrane-bound (Sueoka & Quinn, 1968). The kinetics of breakdown of pulse-labelled DNA in colicin E2-treated bacteria (unpublished results), however, indicate that breakdown does not begin at the replication fork, unless recently copied parental strands only are affected. The possibility is now being examined that breakdown is initiated at the chromosomal origin.

Enzymes concerned in thymine dimer excision and in host specificity mechanisms are not activated by colicin E2, nor is the ATP-dependent nuclease which is determined by the RecB and RecC genes (Buttin & Wright, 1968; Barbour & Clark, 1970). We found no decrease in colicin E2-directed DNA breakdown in a variety of Escherichia coli mutants which are temperature-sensitive for DNA replication. Obinata & Mizuno (1970) have shown that at least in the early stages of E2-induced DNA breakdown, a specific endonuclease (but not endonuclease I) is involved. Further screening among DNA repair and replication mutants may well yield strains which lack one or more of the E2-specific nucleases.
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ADDITION

After preparation of this paper, Ringrose (Biochimica et biophysica acta, 1970, vol. 213, pp. 320–334) reported that DNA breakdown induced by E2 proceeds initially via single-strand breaks and that these can be repaired and their further formation halted if the bacteria are treated at an early stage with trypsin. This report further supports the hypothesis proposed here that membrane changes induced by E2 are reversible and that these changes directly affect the activity of a membrane-bound nuclease.

REFERENCES


Induction of DNA breakdown by colicin E2


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