THE PRODUCTION OF DIARRHOEA IN BABY RABBITS BY THE ORAL ADMINISTRATION OF CELL-FREE PREPARATIONS OF ENTEROPATHOGENIC *ESCHERICHIA COLI* AND *VIBRIO CHOLERAE*: THE EFFECT OF ANTISERA

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**Plate XVII**

Since the initial investigations of Dutta, Panse and Kulkarni (1959), baby rabbits have been used fairly extensively in studying the diarrhoea-producing effect of orally administered cell-free products of *Vibrio cholerae*. Essentially the same technique, but with piglets, has been employed by Kohler (1968) and Smith and Gyles (1970a) in studying porcine enteropathogenic strains of *Escherichia coli*. Exploiting the fact that the enterotoxin produced by these porcine strains is controlled by a transmissible plasmid designated Ent (Smith and Halls, 1968), Smith and Gyles (1970a) were able to compare materials from strains that differed only according to whether they were Ent+ or Ent−. The value of such a well-controlled system became apparent when it was noted that under some conditions material from an Ent− strain could cause diarrhoea. Recently Smith and Linggood (1971) transmitted an Ent plasmid from a human enteropathogenic strain of *E. coli* to *E. coli* K12F−. Because the enterotoxin produced by this strain, like that produced by most other human enteropathogenic strains, is not active on ligated pig intestine, but is active on ligated rabbit intestine, it was decided to determine its effect when administered orally to baby rabbits. Enterotoxic preparations of a strain of K12F− harbouring an Ent plasmid belonging to a porcine strain, of a selection of wild human and porcine enteropathogenic strains and of *V. cholerae* were included for comparison. Antiserum-neutralisation tests were also performed, particularly as Gyles and Barnum (1969) had demonstrated, in ligated intestine studies, an immunological relationship between the enterotoxins of *V. cholerae* and of porcine enteropathogenic strains of *E. coli*.

**Materials and Methods**

*Description of strains of *Escherichia coli*.* The human enteropathogenic strains were given the prefix H and the porcine ones the prefix P. The strain designated K12F−(H19)Ent+ was the K12F− strain into which Smith and Linggood had introduced an Ent plasmid from the human enteropathogen H19 (antigenic structure O26:K60:H19). The K12F−(P307)Ent+ strain had been prepared in a similar manner by Smith and Gyles (1970a), its Ent plasmid having originated in the porcine enteropathogen P307 (antigenic structure O8:K87,88ab).

*Preparation of LT and ST forms of *E. coli* enterotoxin.* Since the enterotoxin of some
enteropathogenic strains exist in two forms, ST and LT, in culture media, both forms were studied in the present work, but greater emphasis was placed on the LT form because it is more active than ST on ligated rabbit intestine (Smith and Gyles, 1970b). They were prepared according to the methods of Smith and Gyles (1970a).

_E. coli_ antisera. These were prepared by the technique of Smith and Gyles (1970a).

### Table I

*The production of diarrhoea in baby rabbits by the oral administration of LT-type enterotoxin of strains of Escherichia coli*

<table>
<thead>
<tr>
<th>Strain from which LT was prepared</th>
<th>Antigenic structure</th>
<th>Dose (ml)</th>
<th>Number of rabbits that developed diarrhoea/number challenged</th>
</tr>
</thead>
<tbody>
<tr>
<td>K12F~(H19)Ent~</td>
<td>?</td>
<td>5</td>
<td>0/10</td>
</tr>
<tr>
<td>K12F~(H19)Ent+</td>
<td>?</td>
<td>5</td>
<td>0/10</td>
</tr>
<tr>
<td>K12F~(P307)Ent+</td>
<td>?</td>
<td>20*</td>
<td>6/8</td>
</tr>
<tr>
<td>K12F~(P307)Ent*</td>
<td>?</td>
<td>5</td>
<td>10/10</td>
</tr>
<tr>
<td>K12F~</td>
<td>?</td>
<td>2.5</td>
<td>2/2</td>
</tr>
<tr>
<td>K12F~</td>
<td>?</td>
<td>5</td>
<td>0/7</td>
</tr>
<tr>
<td>K12F~</td>
<td>?</td>
<td>20*</td>
<td>0/7</td>
</tr>
</tbody>
</table>

**Human enteropathogens**

- H4: O55:K59:H7
- H6: O26:K60:H11
- H7: O111:K58:H2
- H8: O119:K69:H6
- H19: O26:K60:H11
- H26: O125:K70:H21

**Porcine enteropathogens**

- P5: O141:K85ab,88ab
- P14: O147:K89,88ac
- P115: O149:K91,88ac
- P120: O138:K81,88ac

* Concentrated to 5 ml, by dialysis against polyvinyl-pyrrolidone, before administration.

K12F~(P307)Ent+ is a K12F~ strain to which the Ent plasmid has been transmitted from the porcine enteropathogen P307 (antigenic structure 08:K87,88ab); K12F~(H19)Ent+ is a similar strain except that its Ent plasmid originated in the human enteropathogen H19.

_Vibrio cholerae_ enterotoxin and antiserum. These were kindly supplied by the Wellcome Research Foundation, the enterotoxin consisting essentially of a freeze-dried cell-free broth culture of _V. cholerae_. For use, this was dissolved in distilled water to give a final concentration of 50 mg per ml. The antiserum had been made by injecting the enterotoxic preparation into rabbits.

The administration of enterotoxin to baby rabbits. Rabbits, 6–9 days old and weighing approximately 100 g, were removed from their mothers in turn and given, by means of a polythene stomach tube attached to a syringe, the preparation under test. Administration occupied only 1–1 min., after which each rabbit was returned to its mother. The rabbits were then examined frequently for evidence of diarrhoea. In antiserum-neutralisation tests, the enterotoxin-antiserum mixtures were held at 37°C for 1 hr before being given to the rabbits. The volume of _E. coli_ and _V. cholerae_ preparations used in these tests was 5 and 1 ml respectively.

Rabbit ligated intestine preparations. The technique of Smith and Gyles (1970b) was employed.
RESULTS

The results of giving baby rabbits orally LT preparations of *E. coli* K12F- (H19)Ent+, K12F- (P307)Ent+ (K12F- strains to which had been transmitted the Ent plasmids respectively from the human enteropathogenic strain, H19, and from the porcine enteropathogenic strain, P307), and of the original K12F- strain, are illustrated in table I. Results are also given for several wild enteropathogenic strains of human and porcine origin. The human strains had been specially selected in that they had previously yielded LT preparations that consistently dilated ligated rabbit intestine; the porcine strains were all strongly positive in this respect (Smith and Gyles, 1970b).

None of the ten rabbits given 5 ml doses of LT prepared from K12F- (H19)Ent+ or from the seven human enteropathogenic strains, including H19 itself, showed any sign of ill-health as a result. Six of eight given 20 ml of the LT of K12F- (H19)Ent+ that had been concentrated to 5 ml by dialysis against polyvinyl-pyrrolidone developed diarrhoea 10–18 hr after administration, the principal abnormality noted when they were killed being gross distension of the large intestine with a pale yellow watery fluid (fig. 1).

All of the ten rabbits given 5 ml of LT of K12F- (P307)Ent+ developed severe diarrhoea within 4 to 9 hr, nine of them dying within 8 1/2 to 20 hr. Apart from the diarrhoea, the most important clinical sign in these rabbits was intense dehydration. At autopsy, many of them had large amounts of watery

<table>
<thead>
<tr>
<th>Strain against which antiserum was prepared</th>
<th>Dose of antiserum (ml)</th>
<th>Number of rabbits that developed diarrhoea/number challenged, when the antiserum was pre-incubated with the enterotoxin of</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td><em>V. cholerae</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0/1</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>0/1</td>
</tr>
<tr>
<td></td>
<td>0.5</td>
<td>0/1</td>
</tr>
<tr>
<td></td>
<td>0.25</td>
<td>0/1</td>
</tr>
<tr>
<td></td>
<td>0.125</td>
<td>0/2</td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>V. cholerae</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td>1/3</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>1/1</td>
</tr>
<tr>
<td></td>
<td>0.5</td>
<td>1/1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1/1</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>...</td>
</tr>
</tbody>
</table>

* None = normal pig serum.

The dose of *V. cholerae* and K12F- (P307)Ent+ enterotoxins employed was 1 and 5 ml respectively.
fluid in the small intestine as well as in the large intestine (fig. 2). Similar findings were obtained in the rabbits given LT prepared from the wild porcine enteropathogenic strains. No sign of ill-health was noted in baby rabbits given ST, instead of LT, preparations of K12F−(P307)Ent+ or K12F−(H19)Ent+ in 15-ml amounts.

All four baby rabbits given 1 ml of the enterotoxic preparation of \textit{V. cholerae} developed severe diarrhoea 6 to 12 hr later, the clinical and post-mortem signs closely resembling those observed in the rabbits given the LT preparations of porcine enteropathogenic \textit{E. coli}. None of three rabbits given 0.5 ml of the \textit{V. cholerae} preparation developed diarrhoea.

The dilating effect of LT preparations of K12F−(H19)Ent+ and K12F−(P307)Ent+ in ligated segments of intestine was compared in three adult rabbits. Approximately four times the amount of the K12F−(H19)Ent+ LT preparation was required to produce the same degree of dilatation as that produced by the K12F−(P307)Ent+ preparation.

The results of studies on the diarrhoea-inhibiting effect of different antisera on the \textit{V. cholerae} and K12F−(P307)Ent+ enterotoxins are summarised in table 11. Although both enterotoxins were neutralised by either the \textit{V. cholerae} or the K12F−(P307)Ent+ antiserum, more heterologous than homologous antiserum was required in each case to achieve this neutralisation. The K12F−(H19)Ent+ antiserum and normal pig serum had no observable neutralising effect on these two enterotoxic preparations.

\textbf{DISCUSSION}

Because they were made in exactly the same manner, the only important difference between the LT preparations of K12F−(H19)Ent+ and K12F−(P307)Ent+ lies apparently in the fact that they represent the expression of two different Ent plasmids, one derived from a human enteropathogenic strain of \textit{Escherichia coli} and one from a porcine one. The results indicate that the "porcine" plasmid codes for a more powerful enterotoxin for rabbits, shown by the production of diarrhoea in baby rabbits and by the dilatation of ligated intestinal segments of adult rabbits. Furthermore, the tests with the wild human and porcine strains suggest that this difference is a general one.

The antiserum-neutralisation experiments lend some support to the observation of Gyles and Barnum (1969) in ligated intestine tests that the enterotoxins of \textit{V. cholerae} and porcine enteropathogenic \textit{E. coli} are immunologically related. The quantitative aspects of our experiments, however, suggest that they are not identical.

\textbf{SUMMARY}

Strains of \textit{Escherichia coli} K12F− to which Ent plasmids from a human and a porcine enteropathogenic strain of \textit{E. coli} had been transmitted yielded cell-free preparations that produced diarrhoea when administered orally to baby rabbits. The preparations obtained from the strain containing the "porcine" plasmid was more active in this respect than were those of the strain containing the "human" plasmid. This difference was also observed when preparations
Fig. 1.--The baby rabbit on the right had been given, 24 hr previously, 20 ml of an LT preparation of E. coli K12F–(H19)Ent+ concentrated by dialysis against polyvinyl-pyrrolidone to 5 ml before administration. The one on the left had been given a similar preparation of K12F– at the same time. About $\times 3$.

Fig. 2.--This baby rabbit had been given 5 ml of an LT preparation of E. coli K12F–(P307)Ent+ 20 hr previously. The small intestine in addition to the large intestine contained large amounts of watery fluid. About $\times 2.5$. 
of wild human and porcine enteropathogenic strains were compared. The diarrhoea produced in the baby rabbits resembled that produced by *Vibrio cholerae* enterotoxin.

The diarrhoea-producing activity of cell-free preparations of the K12 strain that contained the "porcine" Ent plasmid could be neutralised by antiserum against either this strain or *V. cholerae*; so could cell-free preparations of *V. cholerae*, but more heterologous than homologous antiserum was required in each case to achieve neutralisation.

I am grateful to Miss Carole Smith for her capable technical help and to Dr P. D. Walker of the Wellcome Foundation for the *V. cholerae* enterotoxin and antiserum.

REFERENCES


