The mechanism of protection of infant mice from intestinal colonisation with *Campylobacter jejuni*

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**Summary.** BALB/c mice, vaccinated intraperitoneally with a heat-killed (62°C) suspension of *Campylobacter jejuni* before mating, completely protect c. 90% of their own infants from intestinal colonisation. This protection has now been investigated further in fostering experiments. Fostering by vaccinated dams within the first 24 h of life prevented intestinal colonisation in 50% of infants from non-vaccinated dams, and reduced colonisation in a further 25%. Infants from vaccinated dams, even if allowed to receive their own mothers' colostrum and milk, became susceptible to challenge when subsequently fostered by non-vaccinated dams. Immunity in experimentally infected infant mice depended upon the consumption of immune milk at and after the time of challenge. High concentrations of IgG antibodies specific for *C. jejuni* were found in the serum and mammary secretion of vaccinated dams, but there was very little specific IgA antibody.

**Introduction**

*Campylobacter jejuni* infection is a common cause of diarrhoea that is usually self-limiting and of varying severity in children and adults (Butzler and Skirrow, 1979). It can cause death in the elderly and the debilitated and severe chronic infection in immunodeficient patients (Ahnen and Brown, 1982). The pathogenesis of campylobacter-induced disease is not clear (Klipstein *et al.*, 1985; Taylor *et al.*, 1985). Little is known about protective mechanisms, but the immune responses, mainly antiflagellar, have been described (Kaldor *et al.*, 1983; Wenman *et al.*, 1985).

An ideal animal model of the disease is not available, but after oral infection the alimentary tract of infant mice becomes colonised without giving rise to illness (Field *et al.*, 1981). Protection against this colonisation can be produced by vaccinating the females before mating (Dolby and Newell, 1986). Like other intestinal bacterial pathogens, *C. jejuni* adheres to mucosal surfaces (Newell and Pearson, 1984; Fauchère *et al.*, 1985) and it seems likely that protection in the infant mouse is brought about by interference with adherence and hence with colonisation. Flagella, although aiding colonisation (Newell *et al.*, 1985), apparently play little role in stimulating protection against colonisation in the infant mouse, because a non-flagellate variant effectively immunised against the flagellate parent strain (Dolby and Newell, 1986).

The purpose of this study was to examine, by means of fostering experiments, the mechanism by which the infant offspring of immunised female mice are protected from intestinal colonisation with *C. jejuni*.

**Materials and methods**

**Vaccination**

The non-flagellate variant SF-2 (NCTC 11827; National Collection of Type Cultures, Colindale, London NW9 5HT) of *C. jejuni* strain 81116 (NCTC 11828) was used for the vaccine as previously described (Dolby and Newell, 1986). Briefly, the organisms were harvested from blood agar in phosphate-buffered saline (PBS), pH 7.2. The suspension was adjusted to 50 International Opacity Units and then heat-killed in a water bath at 62°C for 45 min. Female BALB/c mice were vaccinated intraperitoneally four times at weekly intervals with 0.2 ml of the vaccine and then mated. This regimen was previously found to give about 80% protection (Dolby and Newell, 1986).

**Fostering**

Conception was detected by the presence of vaginal plugs, and caesarian section was performed on some of
the pregnant females 18 days later. Caesarian-delivered infants from vaccinated dams were placed immediately with non-vaccinated foster dams that had themselves just given birth. Infants born to non-vaccinated dams were placed with vaccinated dams within the first 24 h of life, and vice versa. In a separate set of experiments, some naturally born infants were fostered at the age of 4–5 days. A few naturally born infants of vaccinated and non-vaccinated dams were left with their own mothers as controls. For the sake of convenience the dams were usually allowed to suckle either their own offspring or fostered offspring, but not a mixture of both.

Challenge

The wild flagellate strain of \( C. \) jejuni strain 81116, (NCTC 11828) was grown for 24 h on blood agar at 37°C in micro-aerophilic conditions. A suspension of the growth was made in Brucella Broth (Difco), containing Campylobacter Supplement SR 84 (Oxoid) and adjusted to 10 International Opacity Units. An equal volume of sterile skimmed milk was added and each infant mouse was fed orally from a plastic-tipped needle and syringe 4–5 days after birth with 0·025 ml containing approximately 10^7 viable organisms (Dolby and Newell, 1986).

Degree of colonisation

Infant mice were killed at intervals of up to one week after challenge. A segment (c. 5 mg) of the colon was removed and viable organisms were estimated as described previously (Dolby and Newell, 1986). These estimations were made on 2–3 mice/litter at 2–3 and 5–6 days after challenge. The two estimations were invariably similar and provided a confirmation of results within each litter. In the present study, protection of individual infants rather than of families was assessed, by applying the scheme used by Dolby and Newell (1986) for complete protection and expressing the results as percentages. The \( \chi^2 \) test was used for statistical evaluation. A \( p \) value of < 0·05 was considered significant.

Collection of serum and milk

Blood was collected from vaccinated and non-vaccinated dams 6–7 weeks after the final dose of vaccine (i.e., 3–4 weeks after birth of the young and 2–3 weeks after challenge). The dams were anaesthetised and bled out from the heart. Serum was separated, heated at 56°C for 30 min and stored at 4°C.

Milk was collected from the stomachs of infant mice, killed about 7 days after birth (2–3 days after challenge), and stored at −70°C. It was thawed, homogenised in an equal volume of PBS and centrifuged at 12 000 g, and the supernate was collected for testing.

Antibody determination by the enzyme-linked immunosorbert assay (ELISA)

The concentration and Ig class of specific antibody in samples of serum and milk were measured. Strain 81116 cells, disrupted ultrasonically, were used to coat the wells (Newell, 1986), but the conjugate consisted of alkaline phosphatase and goat anti-mouse IgG or IgA (Sigma, Fancy Road, Poole, Dorset BH17 7NH). The substrate consisted of a tablet (5 mg) of the disodium salt of \( p \)-nitrophenyl phosphate (Sigma) dissolved in 5 ml of 10% diethanolamine buffer, pH 9·8 (Don Whitley Scientific Ltd, Shipley, West Yorkshire BD17 5JS). The optical density was read on a micro ELISA reader at 490 nm.

Results

In this series of experiments, the protection conferred by vaccinated dams on their own infants against intestinal colonisation by \( C. \) jejuni was 91% (table I).

Table I. \( C. \) jejuni infection in infant mice fostered within 24 h of birth

<table>
<thead>
<tr>
<th>Infant mouse group</th>
<th>Number of litters</th>
<th>Number of infants</th>
<th>Percentage protected</th>
</tr>
</thead>
<tbody>
<tr>
<td>I/U left with own dam</td>
<td>11</td>
<td>37</td>
<td>0*</td>
</tr>
<tr>
<td>I/V left with own dam</td>
<td>10</td>
<td>23</td>
<td>91*</td>
</tr>
<tr>
<td>I/V fostered by UD</td>
<td>22</td>
<td>49</td>
<td>0</td>
</tr>
<tr>
<td>I/U fostered by UD</td>
<td>18</td>
<td>48</td>
<td>50†</td>
</tr>
</tbody>
</table>

I/U = infants from unvaccinated dam (UD). I/V = infants from vaccinated dam (VD).

*Confirms the findings of Dolby and Newell (1986).
†A further 25% showed a 50–75% reduction in colonisation.

Failure of placental antibody or colostral antibody ingested during the first 24 h of life to protect against oral challenge

Initially, caesarian section was performed to ensure that mammary secretion from vaccinated dams was not taken by their own infants before they were transferred to unvaccinated foster mothers. The colonisation was compared with that of naturally born infants of vaccinated dams fostered within 24 h of birth by non-vaccinated dams. There was no protection in either group (fig. 1), indicating that neither placental antibody nor colostral taken within the first 24 h of life by infants from vaccinated dams influenced the effects of a challenge given 4–5 days later. In the light of these findings the next experiments were performed with naturally born infants, transferred to foster dams within 24 h of birth.

Protection of infant mice fostered within 24 h of birth

The colonisation of naturally born offspring of vaccinated dams transferred within the first 24 h of
PROTECTION AGAINST *C. JEJUNI* IN INFANT MICE

Fig. 1. The colonisation at 2–3 days after challenge of infants of vaccinated dams fostered on to non-vaccinated dams. (a) Caesarian-delivered infants, fostered immediately. (b) Naturally delivered infants fostered within 24 h. Each point represents one member of a 2–6 member litter. Viable counts are expressed as cfu/mg of mouse colon.

Fig. 2. The colonisation at 6 days after challenge of infants of vaccinated and non-vaccinated dams fostered within 24 h of delivery on to non-vaccinated and vaccinated dams respectively. (a) Vaccinated foster dams. (b) Non-vaccinated foster dams. Each point represents one member of a 2–6 member litter. Viable counts are expressed as cfu/mg of mouse colon.
life to non-vaccinated dams, and vice versa, was next investigated. The degree of colonisation in 10 litters is given in fig. 2. Of 24 infants born to five non-vaccinated dams and transferred to vaccinated foster dams, 13 showed no sign of colonisation and the remaining 11 infants were colonised with no more than $10^{5.5}$ cfu/mg of gut (fig. 2a). In contrast, almost all of the 23 infants born to five vaccinated dams and transferred to non-vaccinated foster dams were colonised with $>10^{4.5}$ cfu/mg of gut, some with as many as $10^6$ (fig. 2b).

Vaccinated foster dams were, therefore, able to reduce significantly, by means of their mammary secretion, the ability of *C. jejuni* to colonise the infant mouse colon. However, they could protect only those infants that were with them at the time of challenge. Their own infants, transferred to non-vaccinated foster mothers within 24 h of birth, were not protected by the immune colostrum they had consumed. The summed results of all the experiments (61 litters) are given in table I. The proportion of infant mice completely protected by immune mammary secretion was greater among those left undisturbed with their own mothers (91%) than among those transferred from unvaccinated mothers to vaccinated foster mothers within 24 h of birth (50%). This difference ($p<0.001$) was reduced by the inclusion of partly protected mice (table I).

**The protective effect of immune mammary secretion taken up to the time of challenge**

The offspring of vaccinated females were left with their own mothers for the first 4 days of life until c. 1 h before challenge to allow any absorption of maternal immunoglobulins from the mammary secretion via the infant gut wall to take place. Immediately after challenge each litter was transferred to a non-vaccinated foster dam. Table II shows that immune mammary secretion taken up to the time of challenge now protected 10% of infant mice from colonisation; on the other hand, colostrum taken by control infants transferred to unimmunised dams within 24 h of birth gave no protection—a confirmation of all our previous results (table I and fig. 1). Table II also shows that 56% of infants from non-vaccinated mothers transferred at challenge to vaccinated dams were protected compared with 71% transferred within 24 h of birth.

These observations seem to suggest that protection was better the longer the infants were left with vaccinated dams before challenge. With the numbers of mice used, however, these small differences between the two groups of figures (0% and 10%; 56% and 71%) fall short of being statistically significant.

**Table II. *C. jejuni* infection in infant mice fostered 4 days after birth**

<table>
<thead>
<tr>
<th>Infant mouse group</th>
<th>Fostered at age (days)</th>
<th>Number of litters</th>
<th>Number of infants</th>
<th>Percentage protected</th>
</tr>
</thead>
<tbody>
<tr>
<td>I/V fostered by UD</td>
<td>1*</td>
<td>4</td>
<td>18</td>
<td>0</td>
</tr>
<tr>
<td>I/U fostered by UD</td>
<td>1*</td>
<td>4</td>
<td>17</td>
<td>71</td>
</tr>
<tr>
<td>I/V fostered by UD</td>
<td>4</td>
<td>6</td>
<td>30</td>
<td>10†</td>
</tr>
<tr>
<td>I/U fostered by UD</td>
<td>4</td>
<td>6</td>
<td>25</td>
<td>56§</td>
</tr>
</tbody>
</table>

*These mice are included for comparison, and to confirm earlier results (table I and fig. 1).
† An extra 7% were partially protected.
§ An extra 16% were partially protected.

Abbreviations as in table I.

**Antibody assays (ELISA)**

High titres of campylobacter-specific IgG antibody were demonstrated in the serum (162 000–486 000) and milk (2700–72 900) of vaccinated dams. Generally, a high concentration of IgG antibody in the serum of a mouse was reflected in its milk, though at a slightly lower concentration. There was, however, very little IgA in either milk or serum. Non-vaccinated dams had no demonstrable campylobacter-specific IgG or IgA.

**Discussion**

In mice, maternal immunity is acquired via the placenta and colostrum (Fahey and Barth, 1965), but the trans-placental proportion is small (Brambell, 1966). Antibodies, mainly as IgG in colostrum and milk, may act locally in the gut or be absorbed (Halsey et al., 1982; Appleby and Catty, 1983). The relative importance of circulating and intestinal antibody in *C. jejuni* diarrhoea in man has not been established. In mice we could investigate only the prevention of colonisation and not the prevention of diarrhoea, from which mice rarely suffer.

Our experiments with infant mice show that immune mammary secretion protects strongly against intestinal colonisation by *C. jejuni*, in natural and fostered offspring, provided that it is taken at and after the time of challenge. Transplacental immunity plays no part.

From our experiments, it is difficult to attribute
firmly any protective role to circulating IgG antibodies absorbed from milk in the gut. There was a suggestion, however, of a slight protective effect. This was similar to that achieved by injecting hyperimmune antiserum repeatedly into infant mice (Abimiku et al., 1986). This suggestion was evident in two observations. (1) The proportion of infants protected when left undisturbed with their vaccinated dams was larger (91%) than that of infants fostered on to vaccinated dams (50–75%); the difference in complete protection between the two groups was statistically significant (p < 0.001). However, different feeding patterns in fostered and natural offspring might have produced quantitative differences in antibody transmission. (2) There was a slight protective effect (10–17%) in infants left with their own vaccinated dams for 4–5 days before being challenged and fostered on to non-vaccinated dams but none in infants fostered 24 h after birth. In this instance the difference between the two groups was not statistically significant and the slight apparent protective effect might have been attributable either to circulating antibodies absorbed from the milk or to the residual milk antibody in the stomachs of infants fostered at the time of challenge on to non-vaccinated dams. The part played by mammary secretion from vaccinated dams in the prevention of colonisation by C. jejuni resembles that in rotavirus infection of neonatal mice (Sheridan et al., 1984).

We found high concentrations of IgG specific for C. jejuni in the milk and serum of vaccinated dams but little specific IgA antibody. Mice differ from most mammals in having a low concentration of IgA antibody in their milk during the first 4 days of lactation, but an increase thereafter has been reported (Halsey et al., 1982). Most of the milk in these experiments was collected 2–3 days after a challenge given 4–5 days after birth, and IgA antibodies should have been demonstrable if present. Assays were, however, made on unseparated immunoglobulins; this may have led to a blocking of the IgA assay by IgG antibody. On the other hand, the intraperitoneal vaccination of the dams probably stimulated a humoral IgG antibody response (Lange and Holmgren, 1978), which may well have influenced the Ig class in the milk (Parrot, 1979).

We thank Dr Diane Newell, Centre for Applied Microbiology and Research, Porton Down, for the strains of C. jejuni and for help with the ELISA; Drs Susan Stephens and S. P. Borriello of the Clinical Research Centre for helpful discussions; Miss Moira Hegan for supervising the caesarian sections and infant transfer; and Mr P. Hayes, Mr C. Windler and other members of the Animal House staff.

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