HOST RESPONSE TO INFECTION

Induction of systemic and mucosal immune response in mice immunised with porins of Salmonella typhi

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Porins, purified from Salmonella typhi strain 0901 provided 90% protection to BALB/c mice against a lethal dose (300 × LD50) of S. typhi Ty2 when given intraperitoneally. To measure the porin-induced cellular immune responses, macrophages and lymphocytes were isolated from spleen and lamina propria (LP) of porin immunised-challenge mice and of infected and control mice; T-cell phenotypes, lymphocyte proliferation and cytokine production were studied. The secretory IgA (sIgA) antibody level in the intestinal fluid was also measured to study mucosal immune response. After immunisation, the splenic lymphocytes exhibited a significant increase in total T-cell count and CD4+/CD8+ ratio, while the LP lymphocytes (LPL) exhibited an increase in CD4+/CD8+ ratio only. They also exhibited a significant increase in porin-specific proliferative response and cytokine levels (IL-1, IL-2, IFN-γ and IL-4). After immunisation, slgA antibody was also found to be increased. These results suggest that porins given intraperitoneally induce cellular and humoral immune responses both at systemic and mucosal levels.

Introduction

Typhoid fever is an important public health problem in many developing countries [1, 2]. It has been estimated that the worldwide incidence of typhoid fever exceeds 12.5 million cases/year with a 1% mortality rate [3]. Typhoid fever is caused by Salmonella typhi, which replicates within the cells of the reticulo-endothelial system. The immune response to S. typhi infection is complex and involves cellular, secretory and serum components. In the mouse model, cellular components are involved in resistance to salmonella infection [4–6]. Similarly, it is believed that the cell-mediated immune (CMI) response to S. typhi in man plays a key role in defence against this infection.

The emergence of antibiotic multi-resistant Salmonella strains [7] and the high incidence of adverse reactions associated with traditional whole-cell killed vaccine have led to the development of a new generation of typhoid vaccines. Several attenuated S. typhi strains and subunit vaccines such as capsular polysaccharide (Vi) have been evaluated [8–12]. Presently, attention is focused on the role of porins in the induction of specific immunity. It has been observed that porins induce both humoral and cellular immune responses [13–16]. For CMI response, an earlier study has shown elicitation of delayed-type hypersensitivity and lymphocyte proliferation with porins [15].

The present study aimed to investigate the ability of porins of S. typhi to modulate specific immune status both at systemic and mucosal levels in the host and the protection provided thereby. It also aimed to evaluate the porin-specific induction of lymphocyte transformation, CD4+/CD8+ ratio, cytokine production and secretory IgA (sIgA) in mice immunised with porins.

Materials and methods

Bacterial strains

S. typhi strain 0901 obtained from the Central Research Institute, Kasauli, HP, India, was used to prepare porins. S. typhi Ty2 (standard pathogenic strain) obtained from the same institute was used to challenge the mice.

Animals

Inbred BALB/c mice weighing 16–20 g were used for the experiments and also for isolation of thymocytes for interleukin-1 (IL-1) assay. All the mice were given sterile water and feed ad libitum throughout the experimentation.
Purification of porins

Porins were purified from S. typhi strain 0901 by a modification of the method of Nikaido et al. [17]. Briefly, S. typhi cultured in L-broth containing D-glucose 0.5% at 37°C for 18 h was lysed with a French pressure cell. These lysed cells were treated for 30 min with sodium dodecyl sulphate (SDS) 2%. After centrifugation at 100,000 g for 30 min, the pellet was treated for 2 h with NaCl buffer (50 mM Tris-HCl, pH 7.7, 0.4 mM NaCl, SDS 1%, 5 mM EDTA, mercapto-ethanol 0.05%, 3 mM NaN3). After centrifugation at 100,000 g for 30 min the supernate containing the crude porin was treated with trypsin 0.5 mg/ml at 37°C for 2 h and then passed through a Sephacryl S-200 column (1.6 × 86 cm; Pharmacia) pre-equilibrated with NaCl buffer. The fractions containing the porins were pooled and dialysed against 0.5 mM Tris-HCl (pH 7.4) for 8 days. For the characterisation of porins, SDS-PAGE was performed as described previously [18]. The concentration of lipopolysaccharide (LPS) as a contaminant of porins was determined by Limulus amoebocyte lysate (LAL) assay [19].

Protection study

Protection against challenge was assessed as described previously [20]. Briefly, inbred BALB/c mice were divided into five groups of 10. The mice in four groups were actively immunised with different concentrations of porins (2.5, 5.0, 10 or 20 μg) given intraperitoneally (i.p.) on days 0 and 14. The control group was given normal saline. At 7 days after the last immunisation, the mice were challenged with 300 × LD50 of S. typhi Ty2. The numbers of deaths and survivors were recorded up to 15 days after challenge.

Immunisation of mice

For the study of other immunological parameters, four groups comprising 50 inbred BALB/c mice each were set up as detailed below. (1) Control group: 0.2 ml of normal saline was injected i.p. into each mouse on days 0 and 14. (2) Infected group: mice were challenged i.p. with 103 organisms of S. typhi in 0.5 ml of hog mucin. (3) Immunised group: mice in this group were immunised i.p. on days 0 and 14 with the minimum dose of porins (10 μg) that provided maximum protection to mice against 300 × LD50 of S. typhi. (4) Immunised-challenge group: this group was first immunised as above and 7 days after the last immunisation injection, they were challenged with 300 × LD50 of S. typhi Ty2 in 0.5 ml of hog mucin 5%. The mice were killed 10 and 30 days after immunisation/challenge and various parameters were studied.

Isolation of splenic cells

Cells from the spleen were separated as described by Trizio and others [21]. Briefly, the spleen was removed aseptically and teased to separate the cells. The dispersed cells were filtered through a glass wool column and washed once in RPMI medium by centrifugation at 400 g for 10 min. The red blood cells were lysed with chilled NH4Cl 0.9% solution and after centrifugation cells were suspended in tissue culture medium (RPMI-1640, Sigma, 2 mM L-glutamine, 20 mM HEPES, streptomycin 100 U/ml, 5 mM β-mercaptoethanol (TCM) and fetal calf serum (FCS) 10% and incubated for 90 min in Petri plates. Both adherent (macrophages) and non-adherent (lymphocytes) cells were collected.

Gut fluid collection

The entire small intestine from the duodenum to the ileoocaecal junction was removed and washed with 0.5 ml of chilled phosphate-buffered saline (PBS). This fluid was centrifuged at 4000 g at 4°C for 5 min. The supernate was collected, protease inhibitors were added and the material was stored at −70°C until required.

Isolation of lamina propria (LP) cells

The cells were separated with some modifications of a protocol described previously [22]. The entire small intestine from duodenum to the ileoocaecal junction was removed and washed with cold sterile normal saline. The Peyer's patches, blood vessels, fat and mesenteric tissues were completely removed. The intestine was first opened longitudinally and then cut laterally into small pieces 0.5 cm in length and washed extensively in Hanks's Balanced Salts Solution (HBSS). The pieces were treated three times for 30 min with HBSS containing 5 mM EDTA at 37°C and transferred to 25 ml of TCM containing FCS 5%, DNAse 0.1 mg/ml and collagenase (type II) 0.15 mg/ml. After incubation for 45 min at 37°C with slow stirring, the supernate was collected and the procedure was repeated once more. After the second washing the pieces were pumped through a syringe to effect total disruption. The dead cells and tissue debris were removed by passing the tissue suspension through a sterile glass wool column. The cell suspension was washed with TCM without FCS twice and incubated in a glass Petri plate at 37°C in air with CO2 5% for 1.5 h. The adherent cells were separated as macrophages and the non-adherent cell suspension was subjected to discontinuous gradient centrifugation to enrich for lymphocytes. The gradient was prepared by layering from the bottom 75% (2 ml) and 40% (2 ml) concentrations of Percoll. Approximately (3–4) × 107 cells suspended in TCM with FCS 5% were loaded on the top of the Percoll 40% layer. After centrifugation at 600 g for 20 min at 20°C the layer formed at the interface to the 75% and 40% Percoll solutions was collected. This contained lymphocytes with > 90% viability. The yield was approximately (7–8) × 106 cells/mouse.

Immunophenotyping of T lymphocytes

T cells isolated from spleen and LP were immunophenotyped as described previously [23]. Samples of
2 × 10⁶ lymphocytes isolated from spleen or LP were incubated separately with 10 μl of anti-CD3, anti-CD4 or anti-CD8 monoclonal antibodies (MAbs) labelled with either fluorescein isothiocyanate (FITC) or phycoerythrin (PE) for 30 min at room temperature in the dark. The cells were then washed with 2 ml of PBS and fixed in 0.5 ml of fixative (paraformaldehyde 0.05%). The cells were identified on CELL QUEST software of FACScan (Becton Dickinson).

**Lymphocyte transformation**

Splenic lymphocytes were suspended in TCM and 2 × 10⁶ cells/ml were cultured with porins 20 μg or concanavalin A (ConA) 5 μg/ml for 72 h at 37°C in air with CO₂ 5%; 6 h before the completion of the 72-h period, tritiated thymidine (³H) was added. After 72 h, the cells were harvested and radioactivity was measured on a β-counter (LKB, Sweden). The results were expressed as cpm.

**Cytokine assay**

**IL-1 production and measurement.** This was done according to the method of Blydan et al. [24]. Briefly, 2 × 10⁶ splenic or LP macrophages/ml were cultured with 20 μg of porins or 10 μg of LPS at 37°C in air with CO₂ 5% for 24 h. After centrifugation, the supernate was collected and used for IL-1 assay. Volumes of 100 μl of lymphocytes (2 × 10⁷ cells/ml) from 4–6-week-old BALB/c mice were cultured with 100 μl of serially two-fold diluted culture supernates and 20 μl of PHA (5 μg) in 96-well culture plates (Nunc). Six h before the completion of the 72-h incubation period, 0.5 μCi of tritiated thymidine was added. After 72 h, the cells were harvested and radioactivity counts were measured on a β-counter. The results were expressed as stimulation index derived from:

³H-thymidine incorporated by stimulated thymocytes cpm
³H-thymidine incorporated by unstimulated thymocytes cpm

**IL-2, IFN-γ and IL-4 production and measurement.** This was done according to the method of Hudson and Hay [25]. Briefly, splenic or LP lymphocytes (LPL) 2 × 10⁶/ml were cultured with 20 μg of porins at 37°C in air with CO₂ 5% for 48 h. The test included a standard challenge with ConA 10 μg. After 48 h, the culture was centrifuged and the supernate was collected for cytokine assays.

IL-2 and IL-4 assays were done with the HT2 cell line. For IL-2 assay, HT2 cells were incubated with anti-IL-2 MAbs and for IL-4 assay, the cells were incubated with anti-IL-2 and -IL-2R MAbs (a cocktail of TIB 222, HB 8794 and CRC-1698). Then 100 μl of 1 × 10⁶ HT2 cells/ml were added into 100 μl of supernate of lymphocyte culture taken in duplicate wells of a culture plate (Costar). For the negative control, instead of supernate, 100 μl of TCM was added; 0.5 μCi of tritiated thymidine was added to each well 6 h before the completion of a 24-h incubation period at 37°C in air with CO₂ 5%. The cells were then harvested and the radioactive counts were measured with a β-counter. The results were expressed as pg/ml. The IFN-γ assay was done similarly on the WEHI-279 cell line.

**Measurement of secretory IgA response in intestinal fluid**

Anti-porin slgA in intestinal fluid was measured by ELISA as described previously [26]. Briefly, 100 μl of intestinal fluid diluted 1 in 256 in bovine serum albumin (BSA)-phosphate-buffered saline (PBS) were added to duplicate wells of a 9-well microtitration plate (Nunc) coated with 100 μl of porins (5 μg/ml) and blocked with BSA 0.05% w/v in PBS. The plate was incubated at 37°C for 2 h. After washing with PBS-Tween 20, anti-mouse IgA horseradish peroxidase conjugate was added and the plate was incubated at 37°C for 2 h. The plate was then washed and o-phenylenediamine was added as substrate. The reaction was stopped after 30 min with 1 N H₂SO₄ and the plate was read in an ELISA reader at 492 nm.

**Statistics**

Data were analysed for statistical significance by an unpaired Student's t test and results were considered to be significant at p < 0.05.

**Results**

Porins from S. typhi were purified by gel filtration chromatography and the peak obtained after 68 ml contained pure porins. These were in trimer form and under reducing conditions they showed three bands of mol wt 34, 35 and 36 kDa as described previously [27]. The results of LAL assay showed LPS contamination to be 0.02% in purified porins.

**Protection study**

When mice immunised with different concentrations of porins were challenged with 300 × LD₅₀ of S. typhi Ty2 in hog mucin 5%, maximum protection was observed with 20 μg of porins injected i.p in two equal doses. Subsequently, this dose of porins was used as the immunisation dose in BALB/c mice.

**Phenotypic characterisation of T lymphocytes**

T lymphocytes isolated on days 5, 10 20 and 30 from spleen and LP of control, infected, porin immunised and porin immunised-challenge groups were immunophenotyped. The total T cell counts and CD4+/CD8+ ratio of splenic T cells are
depicted in Fig. 1. The total T-cell count in the immunised and immunised-challenge groups was significantly increased as compared with the control group \( (p < 0.001) \) throughout the study period except on day 5 in the immunised group. In the infected group, the total T-cell count was increased significantly on day 10 and significantly decreased on day 30. An increase in CD4+/CD8+ ratio was observed in both immunised and immunised-challenge groups as compared with the control group and this increase remained significant on days 5, 10 and 20 of the study period in the infected group. The increase in CD4+/CD8+ ratio on days 5 and 10 in the immunised group and on days 5, 10 and 20 in the immunised-challenge group was also significant as compared with the infected group.

As shown in Fig. 2a there was no significant change in total T-cell count from the LP among the study groups throughout the study period. The CD4+/CD8+ ratio of T lymphocytes (Fig. 2b) isolated from LP of immunised and immunised-challenge groups was significantly increased as compared with the control group on days 5 and 10. In the infected group this ratio was significantly increased as compared with the control group on day 5 only.

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**Fig. 1.** Phenotypic characterisation of splenic T cells. (a) Total T cells; (b) ratio of CD4+/CD8+ cells. The values are mean and SD of four separate experiments. □ Control group; □ infected group; □ porin immunised group; ■ porin immunised-challenge group. *\( p < 0.01 \) as compared with control group; **\( p < 0.05 \) as compared with control group; +\( p < 0.01 \) as compared with infected group; + +\( p < 0.05 \) as compared with infected group; #\( p < 0.01 \) as compared with immunised group.

**Fig. 2.** Phenotypic characterisation of lamina propria T cells. (a) Total T cells; (b) ratio of CD4+/CD8+ cells. The values are mean and SD of four separate experiments. □ Control group; □ infected group; □ porin immunised group; ■ porin immunised-challenge group. **\( p < 0.05 \) as compared with control group.
**Lymphocyte proliferation**

As shown in Table 1, there was a significant (p < 0.01) increase in porin-induced proliferative response of lymphocytes isolated from the spleens of the immunised group as compared with the control group.

**Cytokine production**

Porins induced IL-1, IL-2, IFN-γ and IL-4 production by splenic and LP cells isolated on days 5, 10, 20 and 30 from the control, infected, immunised and immunised-challenge groups.

**IL-1 production.** Fig. 3a shows the results of porin-induced IL-1 production by splenic macrophages. In the infected, immunised and immunised-challenge groups, IL-1 production on days 5 and 10 was significantly increased (p < 0.01) as compared with the control group. This increase in the immunised and immunised-challenge groups was also significant (p < 0.01) when compared with the infected group. As shown in Fig. 3b there was no significant increase in IL-1 production by macrophages isolated from the LP of infected, immunised and immunised-challenge groups as compared with that of the control group throughout the study period, except on day 5.

**IL-2 production.** The values for IL-2 production are shown in Fig. 4a. IL-2 production on days 5, 10 and 20 was significantly increased in the infected group (p < 0.01) as compared with the control group. There was a significant increase in IL-2 production in the immunised and immunised-challenge groups as compared with the control group throughout the study period. In these groups the increase on days 5, 10 and 20 was also significant (p < 0.01) as compared with the infected group. With ConA, IL-2 production by splenic lymphocytes isolated on day 10 from control, infected, immunised and immunised-challenge groups was 3025 SD 253.2, 950.6 SD 83, 1237 SD 125 and 1213 SD 112.4 pg/ml, respectively.

As shown in Fig. 4b, there was a significant increase in IL-2 production in the porin-stimulated culture of LPL isolated from immunised and immunised-challenge groups on days 5, 10 and 20 as compared with the control group. The increase in IL-2 production in

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**Table 1. Lymphocyte stimulation with porins**

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Control group</th>
<th>Porin immunised group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unstimulated</td>
<td>1286 (±207)</td>
<td>1353 (±200)</td>
</tr>
<tr>
<td>Stimulated with porins (20 µg/ml)</td>
<td>4341 (±628)</td>
<td>*58 255 (8340)</td>
</tr>
<tr>
<td>Non-specific stimulation (ConA 5 µg/ml)</td>
<td>95 720 (±15 393)</td>
<td>68 739 (±12 008)</td>
</tr>
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Results are expressed as mean (±SD) of four separate experiments. *p < 0.01 as compared with control group.

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**Fig. 3.** IL-1 production in cultures of (a) splenic macrophages and (b) lamina propria macrophages stimulated with porins. The values are mean and SD of six and four separate experiments with splenic and lamina propria macrophages respectively. ○ Control group; □ infected group; ▼ porin immunised group; ▧ porin immunised-challenge group. *p < 0.01 as compared with control group; **p < 0.05 as compared with control group; +p < 0.01 as compared with infected group; #p < 0.01 as compared with immunised group.
the porin-stimulated culture of LPL isolated from the infected group was significant as compared with the control group on day 5. With ConA, IL-2 production in cultures of LPL isolated on day 10 from control, infected, immunised and immunised-challenge groups was 2270 SD 239.8, 1134 SD 133.3, 1074 SD 87.45 and 1082 SD 107.5 pg/ml, respectively.

*IFN-γ production.* As for IL-2 production, IFN-γ was significantly increased in the infected, immunised and immunised-challenge groups as compared with the control group throughout the study (Fig. 5a). This increase in the immunised-challenge group was also significant as compared with the infected group throughout the study period. Similarly, the increase in IFN-γ production on days 20 and 30 in the immunised group was also significant as compared with the infected group. Lymphocytes isolated on day 10 from control, infected, immunised and immunised-challenge groups, upon stimulation with ConA (10 μg/ml)
released 1404 SD 170, 781 SD 90, 1227 SD 158 and 1085 SD 152 pg of IFN-γ/ml, respectively.

The IFN-γ production by LPL isolated from immunised and immunised-challenge groups was also increased significantly on days 5, 10 and 20 as compared with the control group (Fig. 5b). Lymphocytes isolated on day 10 from LP of control, infected, immunised and immunised-challenge groups on incubation with ConA (10 μg/ml) released 1531 SD 168, 1174 SD 143, 1511 SD 172 and 1509 SD 188 pg of IFN-γ/ml, respectively.

IL-4 production. Fig. 6a shows IL-4 production elicited by porins in cultures of splenic lymphocytes. Although IL-4 production in the infected, immunised and immunised-challenge groups was significantly increased as compared with the control group on days 5 and 10 of the study period, this was lower in comparison with porin-induced IL-2 and IFN-γ production (Figs. 4 and 5). With ConA, IL-4 production in cultures of splenic lymphocytes isolated on the day 10 from control, infected, immunised and immunised-challenge groups was 160 SD 23.9, 142.7 SD 17, 128.7 SD 19.52 and 164 SD 22.5 pg of IL-4/ml, respectively.

The investigation of IL-4 production by LPL showed that there was no significant increase in IL-4 production by LPL isolated from infected, immunised and immunised-challenge groups as compared with the control group throughout the study period, except on day 5 in immunised and immunised-challenge groups (Fig. 6b). With ConA, IL-4 production in cultures of LPL isolated from control, infected, immunised and immunised-challenge groups was 79.2 SD 9.6, 71.5 SD 12.3, 83.3 SD 13.5 and 104.7 SD 19.0 pg/ml, respectively.

Secretory IgA response in intestinal fluid

Fig. 7 shows the slgA antibody response in intestinal fluid. It was observed that there was a significant increase in slgA antibody in the immunised and immunised-challenge groups as compared with the control group on days 5 and 10. By days 20 and 30, the slgA antibody level had decreased to the control level. In the infected group there was no significant increase in slgA level. Intestinal fluid collected from mice orally infected with \(1 \times 10^6\) cells of \(S.\) typhimurium was used as positive control.

Discussion

This study investigated the specific systemic and mucosal cellular immune responses to porins of \(S.\) typhi along with the mucosal humoral immune response. Phenotypic change, proliferation and cytokine production were studied with splenic and LP cells isolated from infected, porin immunised and immunised-challenge groups along with the control group. SlgA antibody response was also measured to study the mucosal immune response. The porins purified from \(S.\) typhi strain 0901 contained little LPS contamination (0.02%). Earlier studies showed that this amount of LPS contamination does not influence the immune response [13]. The porins provided 90% protection
against 300 × LD50 of *S. typhi* Ty2, in BALB/c mice, confirming earlier results [28].

Immunophenotyping indicated an increased total T-cell count and CD4+/CD8+ ratio in spleens of porin-immunised and porin-immunised-challenge groups. Earlier it was observed that the mice immunised with porins of *S. typhimurium* showed induction of IL-2 production and DTH response because of the infiltration of helper-effector cells to the site where they generate CMI responses against porins [16]. The increase in CD4+/CD8+ ratio in porin immunised and porin immunised-challenge groups may fit in with this hypothesis. The present study showed that mice immunised with porins produced good lymphocyte proliferation responses to porins. The lack of significant difference in thymidine incorporation in the absence of antigen in control and porin immunised mice shows that immunisation with porins does not result in the appearance of non-specifically activated lymphocytes. The increase in CD4+/CD8+ ratio and porin-specific lymphocyte proliferation shows that porins induce a CMI response.

Previously it has been observed that the pro-inflammatory effects of porins, functioning as chemo-attractants, cause the release of histamine from rat peritoneal cells and induce rat paw oedema [29]. IL-1 production in the present study was significantly increased in the porin immunised and porin immunised-challenge groups as compared with the control group. A significantly high chemiluminescence response to porins has been observed in the monocytes of typhoid patients (unpublished data). This finding, as well as the production of IL-1 seen in the present study, may mean that monocytes/macrophages in blood and spleen are activated by porins. IL-1 has several pro-inflammatory effects, which include activation of T cells and polymorphonuclear leucocytes neutrophil and induction of monokine production (including IL-6 or IL-8).

Studies in mice and man have shown that, on the basis of reciprocal cytokine secretion patterns, T-helper (Th) lymphocytes can be divided into sub-populations termed Th1 (characterised by the production of IL-2 and IFN-γ), Th2 (characterised by the production of IL-4, IL-5, IL-6, IL-10 and IL-13) and Th0 (these secrete cytokines that do not fall into the Th1 and Th2 patterns) [30–32]. In general Th1 cells mediate CMI, provide B-cell help and activate macrophages to kill intracellular pathogens, whereas Th2 cells predominantly provide help to B cells for antibody production. IFN-γ involved in resistance to intracellular bacteria plays an important role in the defence against salmonella infection, probably by augmenting macrophage bactericidal activity [33].

The present study found that production of IL-2, IFN-γ and IL-4 was increased in infected, porin immunised and porin immunised-challenge mice, and that production of IL-2 and IFN-γ was greater than that of IL-4. The results suggest that immunisation with porins elicits both Th1- and Th2-type responses with predominant Th1-type T-cell responses, characterised by induction of lymphocyte proliferation and predominance of IL-2 and IFN-γ production. This type of Th response would be expected to play a critical role in resistance to *S. typhi* infection at the systemic level, by contributing to the elimination of *S. typhi* in localised macrophages of the reticuloendothelial system and in other cells. The significant increase in Th1-type responses in immunised-challenge groups shows that the Th1-type responses play an important role in protection against challenge with virulent strains of *S. typhi*.

The study also showed a significant increase in anti-porin antibody production in the serum of porin immunised and porin immunised-challenge groups (data not shown).

As *S. typhi* enters the host through the gut and as studies with several bacterial pathogens have shown that local antibodies secreted on to a mucosal surface can prevent colonisation and subsequent disease [34–
36], it may be desirable to induce a local immune response against *S. typhi*. The present study investigated whether immunisation of mice with porins induced a mucosal immune response. The sIgA antibody response, immunophenotyping of T cells and cytokine production by LP cells were examined. The significant increase in sIgA antibody level, CD4+/CD8+ ratio and cytokine production (IL-1, IL-2, IFN-γ and IL-4) in porin immunised and porin immunised-challenge groups showed that porins, when given i.p., also induced mucosal immune response. Previous studies with cholera toxin have indicated that a site-restricted sIgA response and cytokine production (IL-1, IL-2, IFN-γ and IL-4) in porin immunised and porin challenge groups showed that porins, as good immunogens against typhoid fever. In a separate experiment it was observed that when porin immunised animals were challenged with *S. typhi*, the organisms could be cultured from the spleen and peritoneal cavity for up to 7 days, but not afterwards (data not shown) so the immunity was not sterile. Therefore, porins can protect mice against *S. typhi*, possibly through both cellular and humoral immune responses induced systemically and locally. If this is also true in the human situation, then porins may act as good immunogens against typhoid fever.

### References


