Poliovirus antibody titres, relative affinity, and neutralising capacity in maternal milk

S Zaman, B Carlsson, A Morikawa, S Jeansson, I Narayanan, K Thiringer, F Jalil, L Å Hanson

Abstract
Varying titres of secretory IgA antibodies to poliovirus type 1 were found previously in the milk of unvaccinated, lactating Pakistani mothers during two different years, reflecting the antigenic exposure on mucosal membranes. To study further the changes in the extent and the form of antigenic exposure reflected in the human milk, human milk samples from Pakistani, Indian, Japanese, and Swedish mothers were collected. The quality and the neutralising capacity of the antibodies was also studied. Secretory IgA, IgG, and IgM antibodies to poliovirus type 1 were determined using enzyme linked immunosorbent assay (ELISA) and relative affinity was measured in ELISA by elution with potassium thiocyanate. Microneutralisation tests were also performed.

The higher secretory IgA antibody titres to poliovirus type 1 in the unvaccinated, naturally exposed Pakistani and Indian mothers' milk, compared with the Swedish and Japanese mothers, presumably reflect the epidemiological situation in these countries. Neutralising capacity and the relative antibody affinity seemed to be higher both in the Pakistani mothers and the group without natural exposure but only given inactivated poliovirus vaccine, that is the Swedish mothers, than the group meeting only live vaccine strains, that is the Japanese mothers.

In our previous study we found varying levels of secretory IgA antibodies to poliovirus type 1 in the milk of lactating, unvaccinated mothers living in the poor urban slum areas of Lahore, Pakistan.1 Secretory IgA antibodies in the milk of these mothers apparently reflect antigenic exposure on mucosal membranes, especially in the gut, and hence may follow the changes in the extent and form of antigenic exposure.1-9

The neutralising antibodies in the milk to poliovirus have usually been assumed to be primarily secretory IgA antibodies as suggested by earlier observations with isolated and even degraded secretory IgA.10 The poliovirus antibody titres in colostrum from mothers in Lahore, determined with the enzyme linked immunoabsorbent assay (ELISA), were high as was the neutralising capacity.1 This may be of special interest as in many developing countries live, oral poliovirus vaccine (OPV) is now given already at or soon after birth (Expanded Programme on Childhood Immunisation, Pakistan, interprovincial immunisation cover-age evaluation and disease surveillance 1987, unpublished)11 and interference between virus and milk antibodies could occur if the infant is breast fed.12 13

Moreover, several studies have reported protection provided by breast milk against various enteric infections,14-18 reflecting not only the function of the amount of specific secretory IgA antibodies present but most likely also relating to the quality of the antibodies, for example avidity.19 This may have future implications in optimally using the vaccines in childhood immunisations in the developing countries.

This study was conducted in order to determine the presence, titres, and relative affinity of secretory IgA antibodies to poliovirus type 1 as well as neutralising capacity in the Colostral and mature milk samples from Pakistan, India, Japan, and Sweden, where the extent and nature of exposure of the mothers to poliovirus varies.

Subjects and methods
STUDY POPULATIONS
This study was performed with breast milk samples from four groups of mothers. (1) Unvaccinated Pakistani mothers who were living in an area still endemic for poliomyelitis, but presently with almost 80% infants fully vaccinated from birth onwards using OPV.11 20 (2) Unvaccinated mothers from India—where poliomyelitis is still endemic and 45% of infants are presently vaccinated using OPV.21 (3) OPV vaccinated mothers in Japan—where poliomyelitis does not exist and only OPV is used for vaccinations.11 (4) Swedish mothers vaccinated exclusively with inactivated poliovirus vaccine (IPV)—where poliomyelitis as well as circulation of poliovirus in the community has been prevented.21

Nine samples of colostrum from Pakistani mothers living in the poor urban slum area of Lahore were obtained (mean day of collection was three days after delivery). Ten, eight, and 14 samples of Colostrum each were obtained from the Indian, Japanese, and Swedish mothers respectively (mean day of collection was three days after delivery). The Indian mothers came from a poor section of New Delhi.

Mature milk samples were also collected from 13 Pakistani, 14 Japanese, and 15 Swedish lactating mothers between 13 to 360 days of lactation. The milk samples were obtained directly in plastic tubes after gentle expression of the breasts. The samples were kept at −20°C until transported on dry ice. All milk samples were analysed at the same time period.
Antibody determinations

The samples of milk and colostrum were thawed and centrifuged twice to eliminate fat and cells. ELISA was performed as already described.\(^2\)\(^2\)\(^3\)\(^4\) Poliovirus type 1 antigen produced by the Rijksinstituut voor Volksgezondheid en Milieuhygiene, Bilthoven, the Netherlands was used for coating microtitre plates. Rabbit antisera to \(\alpha, \gamma, \) and \(\mu\) chains (Dakopatts AS) conjugated to alkaline phosphatase, and sheep antiserum against secretory component (Seward Labs) were used for reading the samples tested in four 10-fold dilutions. Titres were expressed as the interpolated reciprocal reading giving an absorption of 0.2 above background at 405 nm.

Antibody avidities were determined in ELISA,\(^2\)\(^4\)\(^5\) where potassium thiocyanate (diluted in phosphate buffered saline containing 0.05% Tween 20) was used in eight different concentrations ranging from 0.25 M to 4 M to elute the antibodies bound to the antigen on the solid phase. All samples were first diluted to similar antibody activity (a reading of 0.6±0.8 optical density for uneluted samples). Relative affinity index was calculated as the molarity of potassium thiocyanate where 50% of the antibodies were eluted.

Neutralisation titres were determined in a modified technique using microtitre plates with poliovirus type 1 virus. Starting dilution was 1:10.\(^6\)\(^7\) \(^8\)

Statistical analysis

Log\(_{10}\) transformations were done on the reciprocal ELISA titres. Neutralisation titres were given as geometric mean titres. These means were compared using Wilcoxon's rank sum test. The statistical significance was chosen at the 0.05 level with a two tailed test.\(^28\)

Results

The reciprocal ELISA titres of IgA, IgG, and IgM in samples of colostrum against poliovirus type 1 are shown in table 1. The mean IgA titres were the highest in the Pakistani and Japanese samples and were significantly higher than the Swedish and Japanese colostral samples (p<0.01). IgG antibodies were rarely found. All the Pakistani mothers had high IgM titres. Six out of 10 Indian and only 2/14 Swedish and 1/8 Japanese mothers had detectable IgM antibodies (table 1).

The mature milk samples from the Pakistani mothers showed the highest mean IgA titres to poliovirus type 1 compared with the Swedish or the Japanese milk samples (p<0.05). IgG antibodies were detected in a few Swedish and Japanese samples. IgM antibodies were rarely found in the Swedish or Japanese milk samples, and they were not determined in Pakistani milk samples (table 2).

The relative affinity index calculated for mature milk antibodies from the Swedish mothers was significantly higher than in the Swedish colostral samples (p<0.001). Swedish mature milk samples also had a higher affinity index than the Japanese mature milk (p<0.005).

No other significant differences were seen between or within the groups (table 3).

Geometric means of neutralisation titres to poliovirus type 1 were the highest in the Pakistani samples of colostrum compared with the Indian (p<0.05), Swedish (p<0.05), or Japanese samples, which had very low neutralisation titres. The mean neutralisation titres were similar in the Pakistani and the Swedish mature milk samples and were very low in all the samples from the Japanese mothers (table 4).

Table 3 Mean (range) avidity index for IgA antibodies in milk expressed as molarity of potassium thiocyanate giving 50% antibody activity of uneluted sample

<table>
<thead>
<tr>
<th>Country</th>
<th>Colostrum</th>
<th>Mature milk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pakistan</td>
<td>0.85 (0.25-1.53)</td>
<td>1.04 (0.45-1.60)</td>
</tr>
<tr>
<td>Japan</td>
<td>0.82 (0.1-1.38)</td>
<td>0.77 (0.06-1.38)</td>
</tr>
<tr>
<td>Sweden</td>
<td>0.64 (0.1-1.4)</td>
<td>1.59 (0.8-2.50)</td>
</tr>
</tbody>
</table>

*Swedish mature milk significantly higher than Swedish colostrum (p<0.001) and Japanese mature milk (p<0.005).

Table 4 Neutralisation titres expressed as geometric mean titres representing the neutralising capacity of the samples of colostrum and mature milk in mothers from different countries

<table>
<thead>
<tr>
<th>Country</th>
<th>Neutralisation titres*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Colostrum</td>
<td>Mature milk</td>
</tr>
<tr>
<td>Pakistan</td>
<td>60**</td>
</tr>
<tr>
<td>Japan</td>
<td>&lt;30</td>
</tr>
<tr>
<td>Sweden</td>
<td>30</td>
</tr>
</tbody>
</table>

*Neutralisation titres expressed as geometric mean titres. **Showing significant differences between India, Pakistani, and Swedish colostrum (p<0.05).

Geometric mean titres (SD) 3.67 (0.64) and 3.67 (0.43) for Pakistani and Swedish mature milk samples (p<0.05). ND=not done.

\(^*\)Titres as assessed by ELISA expressed as Log\(_{10}\). Comparison of Indian mothers in secretory IgA antibodies with Pakistani mothers (>0.05) and with Japanese and Swedish mothers (p<0.01) using Wilcoxon's rank sum test.\(^*\)Number of samples positive among the total number of samples.

\(^*\)Geometric mean titres (SD) 3.67 (0.64) and 3.67 (0.43) for Pakistani and Swedish mature milk samples (p<0.05). ND=not done.
Discussion

Human milk contains a variety of antibodies to intestinal microbes to which the mother is exposed. The milk antibodies against salmonella and shigella reflected well the higher extent of exposure of the Guatemalan compared with Swedish mothers. A different antibody response to cholera vaccination and to live and killed poliovirus vaccinations was seen in Pakistani and Swedish mothers as another illustration of the connection between intestinal exposure and milk antibody response.

The significantly higher secretory IgA antibody titres in the mature Pakistani milk samples compared with Japanese and Swedish milk samples may suggest that exposure is maintained throughout the lactation period in the Pakistani mothers. The lower values in the Japanese and Swedish mothers presumably reflect the result of a long term immunological memory because their exposure to the poliovirus in the form of vaccine had taken place years earlier.

The present data agree with previous suggestions that milk antibodies may reflect the epidemiological situation in a community. In India and Pakistan wild as well as vaccine poliovirus strains are still in circulation, and nationwide vaccinations of children against poliomyelitis are being done using OPV. As a result, the Indian and Pakistani mothers had higher secretory IgA antibody titres against poliovirus type 1 in their milk than the Japanese and Swedish mothers. The fact that the Pakistani mothers had significantly higher mean secretory IgA antibody titres (p<0.01) compared with the Japanese and Swedish mothers presumably illustrates the high endermicity of poliomyelitis in Pakistan. The Japanese mothers are presently exposed only to vaccine strains as no cases of acute poliomyelitis have been reported since 1981 and OPV has been the vaccine used since 1955. In contrast, the only permitted vaccine in Sweden has been IPV since 1955, eliminating wild as well as vaccine virus strains during the late 1970s. The secretory IgA antibody titres to poliovirus type 1 may still appear in this situation of repeated IPV intake of IPV can induce a secretory IgA response.

The secretory IgA antibodies occurring after repeated IPV also seemed to be of higher avidity than those after both natural exposure and OPV. The avidities of the antibodies in the Swedish mothers even increased during lactation. It seems that there can be quite variable patterns in the milk antibody avidities as we have been able to demonstrate both decreases and increases as lactation goes on. Milk from Costa Rican as well as Swedish mothers showed decreasing avidities against Escherichia coli O antigens, comparing colostrum and mature milk. Earlier, we were not able to see such avidity decreases in antibodies to E coli O antigens to Pakistani mothers. It was also remarkable that the Swedish mothers had higher neutralising titres than the Japanese mothers. It is not known whether this can be related to the higher avidity of the Swedish milk antibodies.

The higher antibody avidity in the secretions indicates a more mature immune response produced after continuing antigen exposure. Thus IPV can probably give rise to memory cells, which then can migrate to the lactating mammary gland many years after the vaccination. Nutritional status, repeated infections, and a possible genetic mechanism may be responsible in determining the antibody avidity. However different routes of antigenic exposures seem to be a more prominent factor in our study.

The low neutralisation titres in the Japanese colostrum samples indicate that natural exposure and repeated IPV vaccinations can induce a better neutralising antibody response in secretions than vaccinations with live vaccines. Also, a higher avidity index can be obtained by repeated vaccinations with IPV.

This study was supported by grants from the Swedish Agency for Research Cooperation with Developing Countries; the Ellen, Walter and Lennart Hesselman Foundation for Scientific Research; and the Faculty of Medicine, University of Göteborg and the Swedish Medical Research Council (No.215). We wish to thank Eeva Niinimäki, Ingela Karlsson, and Ann-Marie Mäntyniemi for skilled technical assistance and Mrs Surraiaya for help in the field work.

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Arch Dis Child 1993 68: 198-201
doi: 10.1136/adc.68.2.198

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