Trimethoprim-sulphamethoxazole in the treatment of persistent diarrhoea: a double blind placebo controlled clinical trial

N H Alam, P K Bardhan, R Haider, D Mahalanabis

Abstract
The efficacy of an absorbable antimicrobial agent trimethoprim-sulphamethoxazole (TMP-SMX) in the management of children with persistent diarrhoea was evaluated in a double blind, randomised, and placebo controlled trial. Of the 55 patients studied, 28 received TMP-SMX, and 27 received placebo. A trend in stool weight reduction was observed from the third day after the drug was started, and the reduction was statistically significant on day 6 and day 7. However, the difference in total stool output (g/kg) up to day 7 was not significantly different between the two groups. The proportion of children whose diarrhoea resolved by day 7 (therapeutic success) was significantly more in the treatment group compared with the placebo group (23 v 15). Additionally, mean duration of diarrhoea in the group that received TMP-SMX was less compared with the placebo group (6-0 v 8-3 days); this difference, however, was not significant. Hospital infection (probably nosocomial infection) was significantly less in the TMP-SMX treated group (1 v 10). The results of our study indicate that TMP-SMX has a clinical benefit in respect of reducing the stool output, and higher recovery rate within seven days of treatment. In addition, it prevented possible hospital acquired infection.

(Keywords: trimethoprim-sulphamethoxazole, persistent diarrhoea, randomised controlled trials.

Persistent diarrhoea is one of the common clinical problems seen in developing countries. One of the mechanisms suggested in the pathogenesis of persistent diarrhoea is small bowel bacterial overgrowth. Treatment of persistent diarrhoea is still based on dietary management, fluid treatment, and general supportive measures. Clinical experiences of the use of antibacterials and anti diarrhoeal agents are limited. A combination of oral gentamicin, metronidazole, and cholestyramine was found to be effective in terminating diarrhoeal illness in most treated patients in an early report. The study, however, lacked a control group. Another study evaluated the efficacy of oral gentamicin, metronidazole, and cholestyramine in persistent diarrhoea in a controlled trial; gentamicin resulted in a significant decrease in stool weight. Tropical sprue, a syndrome of chronic diarrhoea associated with malabsorption in adults and older children is widely prevalent in this subcontinent and often responds to a course of tetracycline. Trimethoprim-sulphamethoxazole (TMP-SMX) is a widely used antimicrobial having very good activity against many Gram negative bacteria.

The present study was carried out to evaluate the efficacy of TMP-SMX in the treatment of children suffering from persistent diarrhoea.

Methods
Patient selection
The study was conducted at the Clinical Research and Service Centre of the International Centre for Diarrhoeal Disease Research, Bangladesh (ICDDR),B. The protocol was approved by the research review and ethics review committees of the centre. Patients eligible for the study were children of 5 months to 15 months of age with a history of diarrhoea acute in onset and persisting for more than 14 days but less than six weeks, and who had: (1) at least four liquid stools over 24 hours before admission, (2) stool output more than 50 g/kg body weight in 24 hours during the observation period, (3) no systemic infection requiring antibacterial treatment (for example, pneumonia, meningitis, and septicaemia, etc), (4) no severe malnutrition (for example, marasmus or kwashiorkor), (5) no history of antibacterial drugs at least seven days before admission, (6) no sulphonamide or co-trimoxazole hypersensitivity, (7) initial stool culture negative for Vibrio cholerae, salmonella, shigella, and microscopic stool examination negative for Entamoeba histolytica/Giardia lamblia during the observation period. Consent was obtained from the parent(s) or legal guardian before the patient entered into the study.

Patients were hospitalised in the clinical research unit of the centre throughout the study period. Initial evaluation included a standard clinical history, physical examination including anthropometry, and stool output measurements for 48 hours on hospital standard diet (milk-rice cereal, oil mixture). During this initial 48 hours (observation period) some laboratory investigations were also carried out. These included microscopic stool examination for pus cells, parasites (including G lamblia, E histolytica, and cryptosporidium), stool culture for shigella, salmonella, V cholerae, Campylobacter jejuni, and diarrhoeagenic Escherichia coli, and enzyme
Table 1  Baseline clinical characteristics of patients; values are mean (SD) or number

<table>
<thead>
<tr>
<th></th>
<th>TMP-SMX group (n=24)</th>
<th>Placebo group (n=27)</th>
</tr>
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<tbody>
<tr>
<td>Age (months)</td>
<td>9-0 (2-9)</td>
<td>8-5 (2-3)</td>
</tr>
<tr>
<td>Male/female</td>
<td>20/9</td>
<td>19/8</td>
</tr>
<tr>
<td>Body weight on admission (kg)</td>
<td>6-6 (1-0)</td>
<td>6-4 (0-9)</td>
</tr>
<tr>
<td>Duration of diarrhea before admission (days)</td>
<td>21-6 (7-2)</td>
<td>21-2 (7-6)</td>
</tr>
<tr>
<td>No of stools 24 hours before admission</td>
<td>13-6 (8-5)</td>
<td>10-6 (4-6)</td>
</tr>
<tr>
<td>Stool output in first 24 hours* (g/kg)</td>
<td>123-0 (42-1)</td>
<td>126-4 (48-2)</td>
</tr>
<tr>
<td>No of vomiting 24 hours before admission</td>
<td>2-9 (4-9)</td>
<td>2-2 (2-6)</td>
</tr>
<tr>
<td>No breast feeding on admission</td>
<td>27</td>
<td>27</td>
</tr>
<tr>
<td>Stool pathogens</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rotavirus</td>
<td>6</td>
<td>8</td>
</tr>
<tr>
<td>Enterotoxigenic E coli</td>
<td>5</td>
<td>7</td>
</tr>
<tr>
<td>Enteroadherent E coli</td>
<td>4</td>
<td>7</td>
</tr>
</tbody>
</table>

*Observation period.

linked immunosorbent assay (ELISA) test for rotavirus. E coli colonies were saved, and enterotoxigenic E coli and enteroadherent E coli were identified by specific gene probes. Blood was also tested for total and differential counts and for packed cell volume.

TREATMENT ALLOCATION
After enrolment the patients were randomly assigned to one of the two treatment groups; one group received TMP-SMX, and the other received placebo. The bottles of both treatment groups looked identical and were arranged and numbered sequentially to correspond to patients' serial numbers and the randomisation code. TMP-SMX and placebo were supplied and randomised by the World Health Organisation, Geneva. The randomisation code was kept sealed until data were fully computerised and provided to the centre's administration. For analysis the code in the form of group A and B, without disclosing their identity, was provided to the investigators. On submission of the data analysis tables the group identity was provided to the investigators for preparing the final report.

CASE MANAGEMENT

Fluid treatment
The fluid balance in patients was maintained using either intravenous fluids containing sodium 133 mmol/l, chloride 98 mmol/l, potassium 13 mmol/l, and acetate equivalent bicarbonate of 48 mmol/l, or oral rehydration solution. The patients with high purging rate (stool output >100 g/kg/day) were given only intravenous fluid for maintaining hydration.

Diet
Initial diet after admission into the study (that is on commencement of medicine intake) was based on rice, glucose, egg albumin, vegetable oil and salts mixture, which is routinely used for persistent diarrhea patients at ICDDR,B.8 This mixture has an energy density of 209 kJ (50 kcal)/100 g. Breast milk was allowed ad libitum. The patients, in addition, received multivitamin drops 10 drops twice daily plus zinc acetate 5 mg three times daily orally as per hospital practice.

Drug treatment
In children receiving TMP-SMX, the dose was adjusted to give 10 mg of trimethoprim/kg of body weight per day divided into two 12 hourly doses for seven days. The medicine contained a mixture of sulphamethoxazole and trimethoprim in a conventional ratio of 5:1. The patients were treated in the hospital with close monitoring until the diarrhea had stopped and remained under observation in the hospital for at least two diarrhoea free days. Body weight, intake of intravenous/oral rehydration fluid, water and diet, and output of stool, vomit, and urine were measured every eight hours, and clinical evaluation reports were recorded every morning and afternoon. Cessation of diarrhoea was defined as the appearance of soft or formed stool and no diarrhoea (liquid stool) for 48 hours. Liquid stool can be poured from one container to another, soft stool takes the shape of the container, and the formed stool retains its shape. Therapeutic success was defined as the cessation of diarrhoea within seven days of starting the TMP-SMX or placebo treatment. Hospital infection was identified as isolation of diarrhoeagenic pathogen on subsequent stool culture after the drug was started and continuation of diarrhoeal symptom, and/or signs of other infection (for example, fever, cough, toxicity, crackles on auscultation of chest, etc) that required antibacterial treatment.

DATA ANALYSIS
All data were computerised using StatPack Gold software package (Walnick Associates) in a microcomputer. Data analyses were done using a data analysis package (SPSS/PC+ Inc). Continuous variables were compared between the groups with Student's t test; x2 test was used for dichotomous variables. A non-parametric test (Mann-Whitney U-Wilcoxon rank sum W) was also used to compare the skewed data. Survival analysis was also done for the duration of diarrhoea and compared between the groups with the log rank test.

Results
Of the 55 patients studied, 28 received TMP-SMX and 27 received placebo. Baseline clinical characteristics of patients were comparable between the two groups (table 1). A trend in reduction of stool weight was observed from the third day after the drug was started,
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Table 2 Comparison of total stool output up to day 7, energy intake, duration of diarrhoea, outcome (success/failure), and hospital infection between two groups; values are mean (SD) or number

<table>
<thead>
<tr>
<th></th>
<th>TMP-SMX group (n=28)</th>
<th>Placebo group (n=27)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total 7 day stool output (g/kg)</td>
<td>577.0 (321.5)</td>
<td>756.4 (425.1)</td>
</tr>
<tr>
<td>Energy intake (kJ [kcal]/kg/day) from hospital diet</td>
<td>181.6 (60) [43.4-14.3]</td>
<td>188.6 (61) [44.9-14.6]</td>
</tr>
<tr>
<td>Duration of diarrhoea (days)</td>
<td>6.0 (4.5)</td>
<td>6.3 (5.9)</td>
</tr>
<tr>
<td>Outcome (success/failure)</td>
<td>23/25*</td>
<td>15/22</td>
</tr>
<tr>
<td>Hospital infection</td>
<td>1</td>
<td>10**</td>
</tr>
</tbody>
</table>

χ² test: *p=0.03; **p=0.003.

and the stool outputs (g/kg) were significantly less (fig 1) on day 6 and 7 (p=0.02 for each comparison). The difference in total stool output (g/kg) up to day 7 was, however, not statistically significant (table 2). Intake of oral rehydration solution, intravenous fluid infusion, water intake, and output of vomit and urine were similar in both groups (data not presented). The energy intake of patients of both groups was similar (table 2). The proportion of patients whose diarrhoea stopped within seven days after the drug was started (therapeutic success) was significantly higher (table 2) in the TMP-SMX treated group (p=0.03). The mean duration of diarrhoea was less in the TMP-SMX treated group; however, the difference was not significant. The survival analysis for duration of diarrhoea also showed a similar trend toward reduced duration of diarrhoea in the TMP-SMX group (fig 2) compared with the controls (p=0.055, log rank test). Hospital infections (probably nosocomial infection) were significantly less in the TMP-SMX treated group (p=0.003). The only patient in TMP-SMX treated group developing hospital infection had pneumonia and was treated with ampicillin and gentamicin injections. Among the patients who developed hospital acquired infection in the placebo group three patients did not resolve diarrhoea after seven days; subsequent stool culture yielded C jejuni and they were treated with oral erythromycin. Another patient developed clinical sepsis with ileus and was treated with ampicillin, gentamicin, and metronidazole injections. Six patients had clinical and radiological features of pneumonia: two of them were treated with penicillin injection, three were given ampicillin injection, and one received oral erythromycin. No drug related untoward effect was observed in the study patients.

Discussion

The results of the present study suggested that TMP-SMX in the treatment of severe persistent diarrhoea has a clinical benefit in terms of reducing the stool weight, duration of diarrhoea, and providing a higher rate of recovery from diarrhoea within seven days of treatment. In addition, it prevented hospital acquired infections. The mechanism of this apparent clinical benefit can only be speculated. TMP-SMX may have been effective in treating and preventing small bowel colonisation. A potential benefit of using antimicrobial agents in the treatment of persistent diarrhoea was suggested by two lines of evidence from study reports on persistent diarrhoea: (a) association of persistent episodes with small bowel bacterial overgrowth, principally with Gram negative enteric bacteria1-3 and (b) the frequent isolation in such episodes of various types of enterobacterioid E coli.9-11 A few reports have shown advantages in terminating diarrhoeal episodes with oral gentamicin,4 an unabsorbable broad spectrum antimicrobial, although one community based study failed to show this clinical benefit.12 The advantages of an absorbable oral antimicrobial is that besides its activity against gut infection, it might control other infections (for example, respiratory tract infection, urinary tract infection, etc). TMP-SMX is a widely used antimicrobial active against many Gram positive and Gram negative bacteria and might have been involved in preventing hospital infections in our patients. This could be an alternative or additional mechanism through which the patients receiving TMP-SMX have shown a clinical benefit.

The results of our study indicate that TMP-SMX was effective in reducing the stool output, duration of diarrhoea, and there was a higher recovery rate within seven days of treatment. In addition, it prevented possible hospital acquired infections. Further studies, both in hospitals and communities, are needed to support the recommendation of TMP-SMX in the treatment of persistent diarrhoea in children.

Figure 2 Survival plot for duration of diarrhoea after treatment with TMP-SMX or placebo in patients with persistent diarrhoea.

This study/publication was funded by the United States Agency for International Development under grant No DPE-0934-D-1-1009-00 with the International Centre for Diarrhoeal Disease Research, Bangladesh (ICDDR,B). The ICDDR,B is supported by the aid agencies of the Governments of Australia, Bangladesh, Belgium, Canada, China, Denmark, Germany, Japan, the Netherlands, Norway, Republic of Korea, Saudi Arabia, Sweden, Switzerland, the United Kingdom, and the United States; international organisations including the Arab Gulf Fund, Asian Development Bank, International Atomic Energy Centre, the United Nations Children’s Fund (UNICEF), the United Nations Development Programme (UNDP), the United Nations Population Fund (UNFPA), and the World Health Organisation; private foundations including the Ford Foundation, Population Council, Rockefeller Foundation and the Sasakawa Foundation; and private organisations including American Express Bank, Beyer AG, CARE, Helen Keller International, the Johns Hopkins University, Swiss Red Cross, and the University of California Press. We would like to thank the World Health Organisation, Geneva for supplying us with the drug and the placebo. Thanks are due to Mr M A Rahman Farway for secretarial assistance.
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Arch Dis Child 1995 72: 483-486
doi: 10.1136/adc.72.6.483

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