PROBIOTICS

Maintaining remission of ulcerative colitis with the probiotic Escherichia coli Nissle 1917 is as effective as with standard mesalazine

W Kruis, P Frič, J Pokrotnieks, M Lukáš, B Fixa, M Kaščák, M A Kamm, J Weismueller, C Beglinger, M Stolte, C Wolff, J Schulze

Background and aim: Evidence exists for the pathogenic role of the enteric flora in inflammatory bowel disease. Probiotics contain living microorganisms which exert health effects on the host. We compared the efficacy in maintaining remission of the probiotic preparation Escherichia coli Nissle 1917 and established therapy with mesalazine in patients with ulcerative colitis.

Patients and methods: In total, 327 patients were recruited and assigned to a double blind, double dummy trial to receive either the probiotic drug 200 mg once daily (n = 162) or mesalazine 500 mg three times daily (n = 165). The study lasted for 12 months and patients were assessed by clinical and endoscopic activity indices (Rachmilewitz) as well as by histology. The primary aim of the study was to confirm equivalent efficacy of the two drugs in the prevention of relapses.

Results: The per protocol analysis revealed relapses in 40/110 (36.4%) patients in the E coli Nissle 1917 group and 38/112 (33.9%) in the mesalazine group (significant equivalence p = 0.003). Subgroup analyses showed no differences between the treatment groups in terms of duration and localisation of disease or pretrial treatment. Safety profile and tolerability were very good for both groups and were not different.

Conclusions: The probiotic drug E coli Nissle 1917 shows efficacy and safety in maintaining remission equivalent to the gold standard mesalazine in patients with ulcerative colitis. The effectiveness of probiotic treatment further underlines the pathogenetic significance of the enteric flora.

UC, ulcerative colitis; IBD, inflammatory bowel disease; CAI, clinical activity index; EI, endoscopic index; ITT, intention to treat; CAI, clinical activity index; GCP, good clinical practice.

Materials and methods

The study was conducted according to the Helsinki Declaration (revised version of Hong Kong) and adhered to good clinical practice (GCP) guidelines. The study was approved by the Ethikkommission der Ärztekammer Nordrhein, Germany, as well as by the local ethics committees of the participating centres. All patients received material in their own language and gave written informed consent. Patients were included in the study if aged 18–70 years and diagnosed with UC in remission (clinical activity index (CAI) ≤ 4, endoscopic index (EI) < 4, and no signs of acute inflammation on histological examination). In addition, inclusion criteria comprised at least two acute attacks of UC prior to the study and a duration of the current remission of no longer than 12 months. Exclusion criteria were: active UC; proctitis with up to 10 cm proximal spread; Crohn’s disease; infectious colitis; severe accompanying illnesses or major colonic surgery; use of antibiotics, sulphonamides, steroids, or other therapies for UC at entry into the trial; administration of EcN within the previous six months before trial entry; as well as known intolerance to salicylates.

Study medication

The investigational drug was a bacterial preparation for oral use containing non-pathogenic Escherichia coli of strain Nissle 1917 (serotype O6:K5:H1). Capsules were enteric coated to protect the microorganisms from gastric juice and contained 2.5–25×10⁷ viable bacteria (MutafloR 100 mg; Andeypharma GmbH, Herdecke, Germany). The control preparation was mesalazine, consisting of enteric coated 5-aminosalicylic acid.
acid (Salofalk500 mg; Dr Falk Pharma GmbH, Freiburg, Germany). The test group received one capsule of Metaflor 100 mg once daily and one tablet of placebo three times daily from day 1 to day 4, and two capsules of Metaflor 100 mg once daily and one tablet of placebo three times daily from day 5 to the end of the study. The control group received one capsule of placebo once daily and one tablet of Salofalk 500 mg three times daily from day 1 to day 4, and two capsules of placebo once daily and one tablet of Salofalk 500 mg three times daily from day 5 to the end of the study.

No concomitant medication for UC was allowed throughout the study.

Study design
This was a randomised, double blind, double dummy trial comparing the relapse preventing effects and safety of a bacterial preparation containing viable EcN and mesalazine for 12 months in patients with UC in remission. The study was conducted in 60 hospitals and private settings in 10 European countries (see list of participating investigators in the appendix).

Randomisation was carried out in a double blind manner in blocks of four patients using 1:1 allocation to the two treatment groups. Only complete blocks of random numbers were used for each centre. If patients were eligible for study entry, they were assigned to random numbers (= patient numbers) in ascending order within each centre according to the chronological order of their randomisation and were given the corresponding study medication.

Evaluation
Clinic visits were required at the start and end of the study as well as after 1, 2, 3, 6, and 9 months of treatment.

The primary objective of the study was to compare the number of patients experiencing a relapse of UC during the 12 month observation period between the two treatment groups. Patients were classified as suffering a relapse when at least one dose of the study medication, and a per protocol population.

Table 1  Demographic data and prestudy clinical characteristics

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>ITT population</th>
<th>PP population</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>EcN (n = 162)</td>
<td>Mesalazine (n = 165)</td>
</tr>
<tr>
<td>Demographic data</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Males</td>
<td>56.8%</td>
<td>52.7%</td>
</tr>
<tr>
<td>Age (y) (median (range))</td>
<td>43 (19–69)</td>
<td>41 (19–82)</td>
</tr>
<tr>
<td>Localisation of UC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sub-/total</td>
<td>26 (16.1%)</td>
<td>35 (21.3%)</td>
</tr>
<tr>
<td>Distant</td>
<td>102 (63.0%)</td>
<td>88 (53.4%)</td>
</tr>
<tr>
<td>Duration of UC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;5 y</td>
<td>71 (43.8%)</td>
<td>84 (50.9%)</td>
</tr>
<tr>
<td>&gt;5 y</td>
<td>91 (56.2%)</td>
<td>81 (49.1%)</td>
</tr>
<tr>
<td>Treatment before study (single/combined therapy)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oral salicylates (mg)*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;1,500</td>
<td>60 (37.4%)</td>
<td>68 (41.2%)</td>
</tr>
<tr>
<td>1,500-3,000</td>
<td>46 (28.4%)</td>
<td>47 (28.5%)</td>
</tr>
<tr>
<td>&gt;3,000</td>
<td>7 (4.3%)</td>
<td>7 (4.2%)</td>
</tr>
<tr>
<td>Clinical activity index</td>
<td>0.9 (1.2)</td>
<td>0.9 (1.2)</td>
</tr>
<tr>
<td>Endoscopic index</td>
<td>1.9 (1.6)</td>
<td>1.8 (1.5)</td>
</tr>
<tr>
<td>Histology (rectum)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>No signs of active disease</td>
<td>77.8%</td>
<td>79.4%</td>
</tr>
<tr>
<td>Quality of life score</td>
<td>24 (4.1)</td>
<td>25.2 (3.7)</td>
</tr>
<tr>
<td>Smoker</td>
<td>10 (6.2%)</td>
<td>11 (6.7%)</td>
</tr>
</tbody>
</table>

UC, ulcerative colitis; EcN, Escherichia coli Nissle 1917; ITT, intention to treat population; PP, per protocol population.

*Partly combined with steroids.

No significant differences between treatment groups.
population (PP). According to generally accepted standards for equivalence and non-inferiority trials,\(^*\) primary analysis of the main objective (difference in relapse rates) was based on the PP population. Assuming 25% protocol violators, a total number of 160 patients in each treatment group was therefore planned.

Baseline comparability and statistical analysis of secondary objectives was assessed using Fisher’s exact test (two sided; \( \alpha = 0.05 \)). In addition, Kaplan–Meier curves were plotted. If no CAI or other parameter was documented at the individual study end, the “last observation carried forward” method was applied. Results are given as mean (SD). Statistical tests were executed using SPSS software package version 10.0 under the Microsoft Windows NT operating system. For exploratory comparisons (tables 2, 3), the Student’s \( t \) test was used.

RESULTS

Patient characteristics

In total, 327 patients were enrolled and randomised to either the EcN preparation (n = 162) or mesalazine (n = 165). The two patient groups were matched with regard to demographic, clinical, and pretreatment characteristics (table 1). The time gap between the end of the last relapse before the study and entry into the study was not longer than four weeks in 11.1% of patients receiving EcN and in 9.1% receiving mesalazine, and not longer than three months in 25.9% and 25.5% of EcN and mesalazine patients, respectively. All 327 randomised patients received at least one dose of the study medication and thus were included in the ITT and safety analysis set.

Before unblinding the study, a steering committee assessed protocol violations in 105/327 (32.1%) patients. Major protocol deviations comprised violation of inclusion criteria (CAI \( \leq 4 \), EI \( \leq 4 \), and no signs of acute inflammation on histological examination) (32 patients in both groups), premature discontinuation of the study without relapse (see below), and unknown or not unequivocally assessed end

| Table 2 Reasons for premature discontinuation of the study |
|-----------------|-----------------|-----------------|
| Reason for premature discontinuation* | EcN (n = 162) | Mesalazine (n = 165) |
| Deterioration of disease (relapse not included) | 1 (0.6%) | 1 (0.6%) |
| Newly emerged exclusion criterion during study | 4 (2.5%) | 2 (1.2%) |
| Patient’s request | 13 (8.0%) | 9 (5.5%) |
| Adverse events | 3 (1.9%) | 6 (3.6%) |
| Insufficient patient compliance | 6 (3.7%) | 6 (3.6%) |
| Insufficient patient cooperation (diary) | 6 (3.7%) | 3 (1.8%) |
| Patient did not appear anymore | 2 (1.2%) | 5 (3.0%) |
| Other reasons | 1 (0.6%) | 2 (1.2%) |
| Patients with premature discontinuation | 19 (11.7%) | 20 (12.1%) |

*Multiple reasons possible.

EcN, Escherichia coli Nissle 1917.
Relapse (primary objective)

PP analysis revealed relapse in 40/110 (36.4%) patients in the EcN group and in 38/112 (33.9%) patients in the mesalazine group (fig 2), resulting in significant equivalence between the two groups (p = 0.003). The corresponding one sided upper 95% confidence limit for the difference in treatment was two groups (p = 0.003). The number of patients in the study at the scheduled visits is shown in fig 1. Premature discontinuation of the study for reasons other than relapse of disease occurred in 39/327 (11.9%) patients (in 19/162 (11.7%) patients in the EcN group and in 20/165 (12.1%) patients in the mesalazine group) (table 2). Newly emerged exclusion criteria during the study were start of concomitant medication in four patients on EcN. One patient on mesalazine became afraid of participating in the study. No deaths but 17 serious adverse events were reported in 13/327 (4%) patients (EcN 7, mesalazine 6). Each serious adverse event occurred only once.

Figure 3 depicts the probability of remaining in remission by Kaplan-Meier curves. Median time to relapse in the EcN group could not be calculated due to the large number of late censorings. In the mesalazine group it was 386 days.

ITT analysis confirmed these results, showing a relapse rate of 45.1% in the EcN group and 37.0% in the mesalazine group (significant equivalence p = 0.013). The upper limit of the 95% confidence interval for the difference in treatment was 16.9%.

Subgroup analyses (secondary objectives)

All subgroup analyses were performed in the ITT population. CAI increased in all patients by 1.8 (3.4) points over the study period, showing a slightly larger increase in the EcN group (2.4 (3.7)) than in the mesalazine group (1.2 (3.0)). No differences were observed in EI or histology between the start and end of the study (fig 4). Table 3 lists relapse rates with regard to duration, localisation, and pretreatment. There were no significant differences between the treatment groups for any of these characteristics. Quality of life scores on admission were 24.5 (3.9) in the EcN group and 24.4 (4.0) in the mesalazine group. Respective values after 12 months were 24.3 (5.2) and 25.1 (3.9). No significant changes occurred during the 12 month observation period.

Safety and tolerance

As rated by the patients, overall tolerance was very good or good in the EcN group in 80.0% and in the mesalazine group in 86.0%. According to the physician’s assessment, the respective values were 85.1% and 90.3%.

Discontinuation of the study medication due to adverse events (relapse included) occurred in 22 (6.7%) patients (11 (6.8%) in the EcN group and 11 (6.9%) in the mesalazine group). Most frequent reasons were gastrointestinal disorders such as bloody stools, nausea, diarrhoea, mucous secretion (EcN 4.3%, mesalazine 4.2%), and abdominal pain (EcN 0.6%, mesalazine 2.4%).

Generally, no unexpected drug reactions occurred during the study. No deaths but 17 serious adverse events were reported in 13/327 (4%) patients (EcN 7, mesalazine 6). Each serious adverse event occurred only once.

Adverse events were reported in 68/162 (42.0%) patients treated with EcN and in 85/165 (35.2%) patients treated with mesalazine. Many adverse events reflect symptoms common for active UC such as bloody stools (4.6%), diarrhoea (5.8%),...
and abdominal pain (8.5%). The most frequent non-intestinal adverse events were viral infections (EcN 4.9%, mesalazine 4.2%), nausea (3.1%, 3.0%) and headache (1.9%, 0.6%). Laboratory tests showed no significant alterations.

**DISCUSSION**

Most controlled trials are designed to test differences in efficacy. In contrast, our trial was aimed at proving equivalence. Indeed, we demonstrated that the probiotic EcN provides significantly equivalent efficacy in preventing relapses of UC and is not inferior to the established gold standard mesalazine. This result was not only confirmed by statistical analysis of the PP population, which is preferred in equivalence studies, but also by ITT analysis.

Therapeutic efficacy is usually demonstrated by superiority in a placebo controlled trial. In serious disease however when effective therapy exists that has already been tested by comparison with placebo, additional placebo controlled trials may be considered unethical. A meta-analysis reviewed 16 studies of maintenance therapy involving 2341 patients with UC. In four of these 16 trials, preparations containing 5-ASA were compared with placebo; in the remaining 12 studies sulphasalazine was compared. 5-ASA was observed to be significantly more effective than placebo in all dosage subgroups (<1 g/day, 1–1.9 g/day, ≥2 g/day). A dose dependent trend was not observed. Indeed, some studies comparing at least two doses were performed showing mainly negative or conflicting results: Pentasa 3 g/day was not superior to 1.5 g/day; balsalazide 4 g/day was better than 2 g/day; balsalazide 6 g/day was better than 3 g/day in one study but in another trial was similarly effective; and two studies with olsalazine reached different conclusions. Thus superior efficacy of doses higher than 1.5 g/day has not been established. It can be stated that mesalazine 1.5 g/day presently reflects the standard in the prevention of UC relapses and thus it qualifies as a control in an equivalence trial.

Previous studies on EcN were criticised for several reasons—for example, short observation period or heterogeneity of patients and outcome parameters. The present trial considered this critique and followed actual standards. The observation period was 12 months, only patients with UC in remission were included, and the clinical outcome was assessed by well established endoscopic and histological activity indices resulting in a low relapse rate for the mesalazine group comparable with previous publications. A total of 327 patients were included to achieve a statistical power sufficient to test for equivalence in a one sided set.

Most likely, IBD is caused by an unrestrained inflammatory response to as yet undefined agents. Although precise identification of the antigenic stimuli has not been determined, the intestinal microflora represents a likely culprit. To manipulate the resident gut bacteria therefore seems to offer a rational approach to maintaining remission in IBD. One way of doing this, which has gained credence over recent years, is by using probiotics.

Mechanisms which may account for probiotic activity include production of antimicrobial agents, inhibition of adhesion of pathogens, and influence on mucosal barrier function. It was reported that inhibition of nuclear factor κB could be mediated by probiotic microorganisms. The properties of EcN are well characterised and its genome has been extensively analysed. It carries non-pathogenic adhesion molecules. A specific lipopolysaccharide renders it immunogenic without showing any immunotoxic properties. Immunomodulating activity was demonstrated for specific immune responses as well as for induction of non-specific natural immunity in preterm infants. EcN develops antagonistic activity against enterobacteria such as *Salmonella enteritidis*, *Shigella dysenteriae*, *Yersinia enterocolitica*, and *Vibrio cholerae*. It prevents invasion of *Salmonella typhimurium* into intestinal cells, inhibits adhesion and invasion of adherent invasive *E coli*, and reduces concentrations of mucosa associated colonic microflora constituents in UC.

EcN is safe. Molecular genetics as well as functional analyses have revealed that EcN does not produce any virulence factors or carry any genes for pathogenicity traits. It does not bear genes for antibiotic resistance, transferable genes or plasmids, and does not take up foreign pathogenic DNA. No formation of enterotoxins, cytotoxins, or haemolysins has been observed and there is no serum resistance. Clinical studies have demonstrated a favourable safety profile for EcN compared with placebo, mesalazine, and lactulose. Our study confirms this excellent safety and tolerance record.

There are other controlled studies with different probiotics. Relapse prevention with *Lactobacillus* GG tested negatively for maintenance therapy in surgically induced remission of Crohn’s disease but a small study showed positive results when *Saccharomyces boulardii* was added to mesalazine.

Inflammation of the ileal pouch constructed after proctocolectomy and ileoanal anastomosis in patients with UC is of particular interest because bacterial growth seems to be of pivotal pathophysiological significance. Cases successfully treated with EcN have been reported. A formulation comprising eight different probiotic bacteria demonstrated convincing therapeutic effects in primary prevention and chronic pouchitis. In an uncontrolled study, this preparation was able to colonise the gut and maintain remission in patients with UC.

**Table 3** Relapse rates according to clinical characteristics (intention to treat population)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>EcN</th>
<th>Mesalazine</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>%</td>
<td>n</td>
</tr>
<tr>
<td>Duration of UC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;5 y</td>
<td>46.5</td>
<td>33/71</td>
</tr>
<tr>
<td>&gt;5 y</td>
<td>44.0</td>
<td>40/91</td>
</tr>
<tr>
<td>Localisation of UC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sub-total</td>
<td>46.2</td>
<td>12/26</td>
</tr>
<tr>
<td>Left sided</td>
<td>63.0</td>
<td>17/27</td>
</tr>
<tr>
<td>Distal</td>
<td>41.2</td>
<td>42/102</td>
</tr>
<tr>
<td>Pretreatment</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oral sulphasalazine*</td>
<td>45.1</td>
<td>51/113</td>
</tr>
</tbody>
</table>

*Partly combined with steroids.

No significant differences between treatment groups.
In conclusion, the use of probiotics in IBD is in accordance with its pathogenesis. They may prevent induction of inflammatory reactions. EcN shows therapeutic efficacy and safety in maintaining remission in UC. It can be considered as an alternative to mesalazine.

REFERENCES


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