THE EFFECTS OF BIOCOMPATIBLE COMPARED WITH STANDARD PERITONEAL DIALYSIS SOLUTIONS ON PERITONITIS MICROBIOLOGY, TREATMENT, AND OUTCOMES: THE BALANZ TRIAL

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♦ Background: A multicenter, multi-country randomized controlled trial (the BALANZ study) recently reported that peritonitis rates significantly improved with the use of neutral-pH peritoneal dialysis (PD) solutions low in glucose degradation products (“biocompatible”) compared with standard solutions. The present paper reports a secondary outcome analysis of the BALANZ trial with respect to peritonitis microbiology, treatment, and outcomes.

♦ Methods: Adult incident PD patients with residual renal function were randomized to receive either biocompatible or conventional (control) PD solutions for 2 years.

♦ Results: The safety population analysis for peritonitis included 91 patients in each group. The unadjusted geometric mean peritonitis rates in those groups were 0.30 [95% confidence interval (CI): 0.22 to 0.41] episodes per patient–year for the biocompatible group and 0.49 (95% CI: 0.39 to 0.62) episodes per patient–year for the control group [incidence rate ratio (IRR): 0.61; 95% CI: 0.41 to 0.90; p = 0.01]. When specific causative organisms were examined, the rates of culture-negative, gram-positive, gram-negative, and polymicrobial peritonitis episodes were not significantly different between the biocompatible and control groups, although the biocompatible group did experience a significantly lower rate of non-pseudomonal gram-negative peritonitis (IRR: 0.41; 95% CI: 0.18 to 0.92; p = 0.03). Initial empiric antibiotic regimens were comparable between the groups. Biocompatible fluid use did not significantly reduce the risk of peritonitis-associated hospitalization (adjusted odds ratio: 0.80; 95% CI: 0.48 to 1.34), but did result in a shorter median duration of peritonitis-associated hospitalization (6 days vs 11 days, p = 0.05). Peritonitis severity was more likely to be rated as mild in the biocompatible group (37% vs 10%, p = 0.001). Overall peritonitis-associated technique failures and peritonitis-related deaths were comparable in the two groups.
Conclusions: Biocompatible PD fluid use was associated with a broad reduction in gram-positive, gram-negative, and culture-negative peritonitis that reached statistical significance for non-pseudomonal gram-negative organisms. Peritonitis hospitalization duration was shorter, and peritonitis severity was more commonly rated as mild in patients receiving biocompatible PD fluids, although other peritonitis outcomes were comparable between the groups.


KEY WORDS: Biocompatibility; glucose degradation products; peritonitis; outcomes; randomized controlled trial; technique survival; end-stage renal disease.

Peritonitis is a serious complication of peritoneal dialysis (PD), resulting in considerable morbidity and mortality, depending on the underlying organism (1–10). Overall rates of peritonitis have declined since the early 1990s (11–13) because of advances in connectology and Staphylococcus decolonization protocols (12,14–16).

Recent evidence suggests that the type of dialysis fluid used for PD may also have an impact on peritonitis rates through local effects on the peritoneal membrane and host defenses against infection. Conventional PD fluids are considered “unphysiologic,” based on their acidic pH (5.0–5.8), high lactate concentration (30–40 mmol/L), high osmolality (320–520 mOsm/kg), high glucose concentration (31–236 mmol/L), and contamination by glucose degradation products (GDPs) generated during the heat sterilization process (17). Such solutions reduce the viability and growth of peritoneal mesothelial cells and fibroblasts in vitro, alter the turnover of structural collagen, and modify the homeostatic balance of cytokines and growth factors (18–20). The viability and function of peritoneal phagocytic cells such as leukocytes and macrophages are also impaired by standard PD fluids (17,18,21–23). Most of these adverse effects of dialysate on the peritoneal membrane appear to be accounted for by acidic pH and high GDP concentrations, because in vivo studies showed that they were largely abrogated by the use of neutral-buffered low-GDP (“biocompatible”) fluids (17,19,24–26).

The recently published balANZ randomized controlled trial (27) found that, in incident PD patients, clinically important and statistically significant reductions in peritonitis were associated with administration of a neutral-pH, lactate-buffered, low-GDP fluid (Balance: Fresenius Medical Care, Bad Homburg, Germany) compared with a conventional lactate-buffered PD solution (Stay-Safe: Fresenius Medical Care). Specifically, patients receiving biocompatible fluid had a lower probability of experiencing peritonitis (30% vs 50%, \( p = 0.006 \)), a significantly longer time to a first peritonitis episode (\( p = 0.01 \)), and a significantly lower overall peritonitis rate (0.30 episodes vs 0.49 episodes per patient-year, \( p = 0.01 \)), which persisted after adjustment for age, sex, body mass index, diabetes, cardiovascular disease, baseline renal function, and peritoneal transport status.

To further evaluate the impact of biocompatible fluid on peritonitis microbiology and outcomes, the present study aimed to determine whether the beneficial effects on peritonitis of neutral-pH, low-GDP (“biocompatible”) PD fluid compared with conventional dialysate in the balANZ trial were specific for particular microorganisms or led to improved peritonitis outcomes (peritonitis-associated hospitalization, peritonitis severity, peritonitis relapse, peritonitis-related technique failure, and peritonitis-related death).

METHODS

The study design and methodology have previously been described (28), as have the results of the main primary and secondary analyses specified in the statistical analysis plan (27). The trial was registered with the Australian New Zealand Clinical Trials Registry (ACTRN1260600044527), and the study protocol was approved by ethics committees at all participating centers. All patients provided written informed consent before trial participation.

Incident adult PD patients who at enrollment had both a measured residual glomerular filtration rate of 5 mL/min/1.73 m² or more, and a measured urine volume of 400 mL/day or more, were included in the study. Pregnant or breastfeeding patients, patients expected to die within 12 months, patients participating in trials targeting residual renal function in PD, and patients with a significant cancer history in the preceding 5 years, with acute infection at enrollment, with contraindications to PD, with any physical or mental disorder that appreciably hampered study protocol compliance, or with a known or suspected allergy to the trial product or related products were excluded.

Using central computer-generated randomization over a Web-based link, participants were randomly allocated to receive either Balance (neutral-pH, lactate-buffered, low-GDP solution) or Stay-Safe (conventional lactate-buffered PD solution), stratified by center and diabetic nephropathy. The PD connectology was identical for both systems, and patients were treated according to local unit practices. Table 1 shows the chemical composition...
of the biocompatible and control fluids used in the baINZ trial.

Peritonitis was defined as a dialysate white cell count greater than 100/μL, with more than 50% polymorphonuclear leukocytes in a patient with a compatible clinical picture (abdominal pain, fever, or cloudy dialysate) (29). The organisms present in each episode of peritonitis during the study, excluding relapses, were classified as gram-positive (coagulate-negative staphylococci, S. aureus, Streptococcus, or other), gram-negative (Pseudomonas or non-pseudomonal), fungal, culture-negative, and polymicrobial (2 or more organisms isolated on effluent culture). When a combination of classifications existed for a peritonitis episode, the episode was included in both classifications for the purposes of the Poisson analysis.

Peritonitis-associated hospitalization was defined as hospitalization primarily for treatment of peritonitis. Peritonitis severity was rated subjectively as mild, moderate, or severe on serious adverse event forms by local clinicians at the time of each peritonitis episode. Peritonitis relapse was defined according to the International Society for Peritoneal Dialysis guidelines as peritonitis with the same organism or a sterile episode occurring within 4 weeks of an earlier episode (29–31). In keeping with International Society for Peritoneal Dialysis recommendations (30,32), a peritonitis episode complicated by relapse was counted as a single episode. Peritonitis-related death was recorded if the patient’s death was directly attributable to peritonitis in the clinical opinion of the treating nephrologist. Each patient was followed for up to 24 months, and the results were analyzed on an intention-to-treat basis.

### Study participants and clinicians were not blinded to PD fluid treatment.

### STATISTICAL ANALYSIS

Results are expressed as frequencies and percentages, means ± standard deviations, or medians with range, depending on data distribution. Group comparisons were performed using the chi-square test, unpaired t-test, or Mann–Whitney test as appropriate. Organism-specific peritonitis rates were compared using Poisson regression, with no adjustment for multiple episodes or multiple organisms per subject. Kaplan–Meier survival analyses and multivariate Cox proportional hazards model analyses were used for time-to-event analyses. The Poisson and Cox analyses were adjusted for center, diabetic nephropathy status, baseline glomerular filtration rate, and PD modality (automated PD or continuous ambulatory PD). Data were analyzed by Statistical Revelations Pty Ltd. (http://www.statisticalrevelations.com.au/). Values of \( p < 0.05 \) were considered statistically significant.

### RESULTS

### PATIENT CHARACTERISTICS

The study randomized 185 patients to receive either biocompatible \( (n = 92) \) or control fluid \( (n = 93) \). Of those patients, 91 in each group \( (182 \text{ overall}) \) were included in the safety analysis in the present study. As previously reported (27), the two groups were well matched for all baseline characteristics, including age, sex, cause of end-stage renal failure, presence of cardiovascular disease, body mass index, initial dialysis modality, prescribed medications, blood pressure, prescribed dialysate volumes and glucose exposure, residual renal function and urine volume, peritoneal ultrafiltration, peritoneal permeability, and laboratory parameters (serum albumin, calcium, hemoglobin). There were also no significant differences between the biocompatible and control groups with respect to the proportions of patients receiving peritonitis prophylaxis measures during the study, including exit-site mupirocin prophylaxis \( (32\% \text{ vs } 40\%, p = 0.35) \), nasal screening for S. aureus \( (69\% \text{ vs } 62\%, p = 0.35) \), prophylactic antibiotic administration at catheter insertion \( (93\% \text{ vs } 90\%, p = 0.59) \), and prophylactic antifungal therapy at the time of peritonitis antibiotic therapy \( (55\% \text{ vs } 52\%, p = 0.77) \). No patients in the study used exit-site gentamicin cream. Of the 16 units participating in the study, 14 routinely prescribed antibiotics at the time of PD catheter insertion; 6 routinely prescribed antibiotics at the time of procedures (for example, colonoscopy);
10 routinely screened PD patients for nasal *S. aureus*; 10 routinely performed topical mupirocin eradication of nasal staphylococcal carriage; 4 routinely prescribed exit-site mupirocin; and 7 routinely prescribed antifungal prophylaxis concomitantly with antibiotic therapy.

**PERITONITIS MICROBIOLOGY**

Based on the 38 episodes of peritonitis that occurred in 27 patients in the biocompatible group and the 67 episodes that occurred in 45 controls, the unadjusted geometric mean peritonitis rates in the biocompatible and control groups were 0.30 [95% confidence interval (CI): 0.22 – 0.41] and 0.49 (95% CI: 0.39 to 0.62) episodes per patient–year respectively, with an incidence rate ratio (IRR) of 0.61 (95% CI: 0.41 to 0.90; *p* = 0.01) in favor of the biocompatible group.

Table 2 shows the micro-organisms recovered during peritonitis episodes. Figure 1 shows a forest plot representing the IRR for the analysis of each group of organisms and all organisms isolated from peritonitis episodes. Compared with control patients, patients in the biocompatible group experienced generally lower rates of culture-negative, gram-positive, gram-negative, and polymicrobial peritonitis episodes, although the reduction was statistically significant only for non-pseudomonal gram-negative peritonitis (IRR: 0.41; 95% CI: 0.18 to 0.92; *p* = 0.03).

**INITIAL PERITONITIS TREATMENT**

Table 3 shows the initial antibiotic regimens used to treat peritonitis episodes in the intervention and control groups. The three most common initial empiric antibiotic regimens used in each group were (in descending order) cefepime (primarily cephazolin) and gentamicin, vancomycin and gentamicin, and first- and third-generation cephalosporins. There were no significant differences in the initial empiric antibiotic selections between the groups (*p* = 0.63).

**PERITONITIS-ASSOCIATED HOSPITALIZATION**

Excluding relapses, hospitalization was required for 23 peritonitis episodes (61%) in the Balance group and for 41 episodes (61%) in the control group (*p* = 0.89). Using negative binomial regression with adjustment for age, sex, body mass index, diabetes mellitus, cardiovascular disease, PD modality, baseline glomerular filtration rate, and peritoneal transport status, biocompatible PD fluid was not associated with a significant reduction in the rate of peritonitis-associated hospitalization (IRR: 0.80; 95% CI: 0.48 to 1.34). The only independent predictor of hospitalization was male sex (*p* = 0.03).

Exposure to biocompatible PD fluid was associated with a shorter duration of hospitalization for peritonitis that just failed to reach statistical significance (*p* = 0.05, Figure 2). The median durations of peritonitis-associated hospitalization were 6 days (95% CI: 4 to 9 days) in the Balance group and 11 days (95% CI: 5 to 15 days) in the

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**TABLE 2**

<table>
<thead>
<tr>
<th>Variable</th>
<th>Dialysate type</th>
<th>Bio compatible</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient–years at-risk (n)</td>
<td>127.7</td>
<td>136.7</td>
<td></td>
</tr>
<tr>
<td>Gram-positive</td>
<td>20 (0.16)</td>
<td>29 (0.21)</td>
<td></td>
</tr>
<tr>
<td><em>Staphylococcus epidermis</em></td>
<td>5 (0.04)</td>
<td>8 (0.06)</td>
<td></td>
</tr>
<tr>
<td>CNS</td>
<td>4 (0.03)</td>
<td>7 (0.05)</td>
<td></td>
</tr>
<tr>
<td><em>S. aureus</em></td>
<td>2 (0.02)</td>
<td>5 (0.04)</td>
<td></td>
</tr>
<tr>
<td><em>Corynebacterium</em></td>
<td>3 (0.02)</td>
<td>2 (0.01)</td>
<td></td>
</tr>
<tr>
<td><em>Rhodococcus</em></td>
<td>1 (0.01)</td>
<td>0 (0)</td>
<td></td>
</tr>
<tr>
<td><em>Streptococcus pneumonia</em></td>
<td>3 (0.02)</td>
<td>6* (0.04)</td>
<td></td>
</tr>
<tr>
<td><em>Enterococcus</em></td>
<td>1 (0.01)</td>
<td>0 (0)</td>
<td></td>
</tr>
<tr>
<td><em>Nocardia</em></td>
<td>0 (0)</td>
<td>1 (0.01)</td>
<td></td>
</tr>
<tr>
<td>Diphtheroid species</td>
<td>1 (0.01)</td>
<td>0 (0)</td>
<td></td>
</tr>
<tr>
<td>Gram-negative</td>
<td>9 (0.07)</td>
<td>18 (0.13)</td>
<td></td>
</tr>
<tr>
<td><em>Escherichia coli</em></td>
<td>2 (0.02)</td>
<td>7 (0.05)</td>
<td></td>
</tr>
<tr>
<td><em>Klebsiella species</em></td>
<td>0 (0)</td>
<td>3 (0.02)</td>
<td></td>
</tr>
<tr>
<td><em>Acinetobacter species</em></td>
<td>3 (0.02)</td>
<td>3 (0.02)</td>
<td></td>
</tr>
<tr>
<td><em>Pseudomonas species</em></td>
<td>2 (0.02)</td>
<td>2 (0.01)</td>
<td></td>
</tr>
<tr>
<td><em>Enterobacter species</em></td>
<td>1 (0.01)</td>
<td>2 (0.01)</td>
<td></td>
</tr>
<tr>
<td>Other*</td>
<td>1 (0.01)</td>
<td>1 (0.01)</td>
<td></td>
</tr>
<tr>
<td>No growth</td>
<td>7 (0.05)</td>
<td>14 (0.10)</td>
<td></td>
</tr>
<tr>
<td>Multiple organisms</td>
<td>2 (0.02)</td>
<td>5 (0.04)</td>
<td></td>
</tr>
<tr>
<td><em>Mycobacterium TB</em></td>
<td>0 (0)</td>
<td>1 (0.01)</td>
<td></td>
</tr>
<tr>
<td><strong>TOTAL</strong></td>
<td>38 (0.30)</td>
<td>67 (0.49)</td>
<td></td>
</tr>
</tbody>
</table>

CNS = coagulase-negative *Staphylococcus*.  
* Results are presented as episodes per patient–year. The reported gram-positive, gram-negative, and mycobacterial peritonitis episodes do not include organisms isolated in polymicrobial peritonitis episodes.  
* One sample grew both *S. capitis* and *S. epidermidis*.  
* One sample grew both *Strep. salivarius* and *Strep. mitis*.  
* Includes *K. pneumoniae* and *K. oxytoca*.  
* Includes *A. baumannii* and *A. johnsonii*.  
* Includes *P. aeruginosa* and *P. oryzihabitans*.  
* Includes *Pantoea*.  
* Includes *Sphingomonas paucimobilis* and *Ochrobactrum anthropi*.  
* Includes *Stenotrophomonas*, *Citrobacter*, *Pseudomonas*, *Clostridium*, *Proteus*, *K. pneumoniae*, *Enterobacter*, *E. coli*, *Enterococcus faecalis*, Morganella morganii, and *Hafnia alvei*.
When peritonitis-associated hospitalization was examined according to causative organism, exposure to biocompatible fluid was associated with a significant reduction in time to hospital discharge for culture-negative peritonitis ($p = 0.04$) and a trend to a reduction in time to discharge for gram-negative peritonitis ($p = 0.07$). No differences were observed between the biocompatible and control groups for time to discharge for gram-positive ($p = 0.36$) or polymicrobial peritonitis ($p = 0.16$). On multivariate Cox proportional hazards model analysis, no significant interaction was seen between treatment assignment (biocompatible vs control) and causative organism ($p = 0.20$). Duration of peritonitis-associated hospitalization varied significantly between centers ($p = 0.002$) and was significantly longer in patients with diabetic nephropathy ($p = 0.005$). Hospitalization duration was not significantly associated with baseline glomerular filtration rate, PD modality, or causative organism.

**PERITONITIS SEVERITY**

Peritonitis severity was rated by attending clinicians in the biocompatible group as mild in 14 episodes (37%), moderate in 17 episodes (45%), and severe in 7 episodes (18%). In contrast, peritonitis episodes were significantly more likely to be rated moderate rather than mild in the control group ($p = 0.05$). When peritonitis-associated hospitalization was examined according to causative organism, exposure to biocompatible fluid was associated with a significant reduction in time to hospital discharge for culture-negative peritonitis ($p = 0.04$) and a trend to a reduction in time to discharge for gram-negative peritonitis ($p = 0.07$). No differences were observed between the biocompatible and control groups for time to discharge for gram-positive ($p = 0.36$) or polymicrobial peritonitis ($p = 0.16$). On multivariate Cox proportional hazards model analysis, no significant interaction was seen between treatment assignment (biocompatible vs control) and causative organism ($p = 0.20$). Duration of peritonitis-associated hospitalization varied significantly between centers ($p = 0.002$) and was significantly longer in patients with diabetic nephropathy ($p = 0.005$). Hospitalization duration was not significantly associated with baseline glomerular filtration rate, PD modality, or causative organism.

**TABLE 3**

Initial Antibiotic Regimens Used to Treat Peritonitis Episodes in Trial Participants

<table>
<thead>
<tr>
<th>Antibiotic combination used as initial regimen</th>
<th>Biocompatible ($n=38$)</th>
<th>Control ($n=67$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vancomycin + gentamicin [$%$]</td>
<td>11 (29)</td>
<td>14 (21)</td>
</tr>
<tr>
<td>Cephalosporin + gentamicin [$%$]</td>
<td>14 (37)</td>
<td>35 (52)</td>
</tr>
<tr>
<td>1st- and 3rd-generation cephalosporin [$%$]</td>
<td>7 (18)</td>
<td>8 (12)</td>
</tr>
<tr>
<td>Gentamicin alone [$%$]</td>
<td>1 (3)</td>
<td>4 (6)</td>
</tr>
<tr>
<td>Cefepime alone [$%$]</td>
<td>2 (5)</td>
<td>1 (1)</td>
</tr>
<tr>
<td>Vancomycin alone [$%$]</td>
<td>2 (5)</td>
<td>2 (3)</td>
</tr>
<tr>
<td>Vancomycin + 3rd-generation cephalosporin [$%$]</td>
<td>1 (3)</td>
<td>2 (3)</td>
</tr>
<tr>
<td>Vancomycin + piperacillin/tazobactam [$%$]</td>
<td>0 (0)</td>
<td>1 (1)</td>
</tr>
</tbody>
</table>

* The overall difference between the groups was not statistically significant ($p = 0.63$).
the control group [mild, 7 episodes (10%); moderate, 52 episodes (78%); severe, 8 episodes (12%); \(p = 0.001\)].

RELAPSED, RECURRENT, AND REPEAT PERITONITIS

In the biocompatible group, 5 peritonitis episodes (13%) were complicated by 7 relapses, which compares with 2 peritonitis episodes (3%) complicated by 2 relapses in the control group (\(p = 0.11\)). In the biocompatible group, 5 relapses (71%) necessitated hospitalization; in the control group, 1 relapse (50%) necessitated hospitalization.

Two episodes of recurrent peritonitis occurred in both the biocompatible and the control group (IRR: 1.07; 95% CI: 0.15 to 7.6; \(p = 0.95\)).

Repeat peritonitis occurred on 4 occasions in the biocompatible group and on 5 occasions in the control group (IRR: 0.86; 95% CI: 0.23 to 3.19; \(p = 0.82\)).

TECHNIQUE SURVIVAL AND PATIENT SURVIVAL

Table 4 shows the causes of withdrawal from PD among the \(b\)al\(A\)NZ trial participants. In both the biocompatible group and the control group, 5 patients had their catheters removed and were converted to hemodialysis as a result of peritonitis (\(p = 1.0\)). One death related to \(P\)seudomonas peritonitis occurred in a patient who was using biocompatible fluid.

DISCUSSION

This large multicenter, multi-country randomized controlled trial demonstrates that, compared with prescription of conventional dialysate, prescription of neutral-pH, lactate-buffered, PD solution low in glucose degradation products (Balance) in PD patients was associated with a marked reduction in overall peritonitis rates. When specific causative organisms were examined, the rates of culture-negative, gram-positive, gram-negative, and polymicrobial peritonitis were not significantly different between the biocompatible and control groups, although the biocompatible group did experience a significantly lower rate of non-pseudomonal gram-negative peritonitis. The initial antibiotic regimens used to treat peritonitis episodes in the intervention and control groups were similar. Compared with conventional PD fluid use, biocompatible PD fluid use was also associated with a shorter duration of peritonitis-associated hospitalization (\(p = 0.05\)) and a higher likelihood of peritonitis severity being rated as mild by attending clinicians. However, peritonitis-related technique failure and peritonitis-related death were comparable between the groups.

The foregoing results are in keeping with an earlier 3-year nonrandomized study of incident PD patients in which 50 patients using bicarbonate-buffered biocompatible fluid (BicaVera: Fresenius Medical Care) experienced a peritonitis rate of 0.34 episodes per patient–year compared with 0.60 episodes per patient–year in 50 patients using conventional PD fluid (StaySafe), despite the fact that patients in the biocompatible group had a higher score on the Charlson comorbidity index and a higher frequency of diabetes mellitus. Using negative binomial regression to adjust for age, sex, nasal staphylococcal carriage, cause of end-stage renal failure, weekly Kt/V, and a 3-point subjective nursing assessment of the patients’ PD skills and ability, biocompatible fluid use was associated with a significant reduction in the risk of peritonitis (adjusted odds ratio: 0.64; 95% CI: 0.43 to 0.98) (33). Similar findings were observed when the results were expressed as patient–exchanges (0.30 vs 0.50 peritonitis episodes per 1000 patient–exchanges; adjusted odds ratio: 0.65; 95% CI: 0.43 to 0.98). Of peritonitis episodes in both groups, the greater proportion were caused by gram-positive organisms. Moreover, compared with PD patients receiving conventional dialysate, those receiving biocompatible PD fluid experienced shorter peritonitis durations (based on peritoneal white cell counts).

Furkert and colleagues (34) similarly observed that, in 53 patients receiving biocompatible fluid between 2000 and 2005, peritonitis rates were appreciably lower than rates in 67 patients receiving conventional dialysates.
between 1990 and 1999 (0.24 episodes vs 0.60 episodes per year respectively, \( p = 0.002 \)), despite the fact that patients in the biocompatible group were older and more likely to be diabetic. However, this retrospective analysis relied on historical controls and may have been affected by co-intervention bias, because overall peritonitis rates have generally improved in most PD centers since the early 1990s.

In contrast to those nonrandomized studies, a number of small randomized controlled trials have failed to observe a difference in peritonitis rates between patients receiving biocompatible and conventional dialysis fluids (35–43). In a 6-month extension substudy of one of those trials (35), which involved 57 of the original 106 participants, peritonitis rates were significantly lower in the biocompatible group than in the control group (0.24 episodes vs 0.63 episodes per patient–year respectively, \( p < 0.05 \)). However, longer-term follow-up of an augmented number of trial participants in another randomized controlled trial reported by Srivastava et al. (44) did not demonstrate a significant difference in peritonitis rates between the biocompatible and control groups (0.52 episodes vs 0.45 episodes per patient–year respectively, \( p = 0.82 \)). In that trial, 86% of patients in the intervention arm used Physioneal (Baxter Healthcare Corporation, Deerfield, IL, USA), which contains appreciably higher levels of GDPs than Balance does (28,43,45,46) and therefore may have been less effective at preserving peritoneal host defenses against infection. Other potential explanations for the apparent disparity in results between the balANZ trial and earlier randomized controlled trials may be insufficient statistical power, short-term follow-up, low event rates, high drop-out rates, enrollment of prevalent PD patients, and single-center designs in the earlier studies. Alternatively, the possibility of a type 1 statistical error (that is, chance finding of a significant result) in the balANZ trial cannot be excluded.

Nevertheless, the marked beneficial effect of biocompatible fluid on peritonitis occurrence and resolution in the present investigation is biologically plausible insofar as experimental evidence has shown that, compared with the effects of conventional acidic-pH, high-GDP dialysates, the effects of neutral-pH, low-GDP “biocompatible” fluids on the peritoneal membrane lead to improved local immune defenses. Numerous in vitro (17,18,18–23) and animal experimental studies (47,48) have demonstrated that conventional PD fluids impair the viability and function of peritoneal mesothelial cells, leukocytes, and macrophages, and that those adverse effects are largely abrogated by exposure to biocompatible fluids. Several clinical studies have further observed that compared with the use of conventional dialysates, the use of biocompatible fluids is associated with improved levels of peritoneal biomarkers such as cancer antigen 125 (36,38), suggesting possibly enhanced preservation of the peritoneal mesothelium. Moreover, a randomized controlled trial (38) reported significantly lower levels of C-reactive protein at 52 weeks of treatment in 25 incident PD patients using biocompatible solution than in 25 patients using conventional PD fluid, suggesting that biocompatible fluids may also exert favorable effects on systemic inflammatory responses. Any ensuing improvement in local peritoneal host defenses may have accounted for the observations in the present study of a generalized reduction in peritonitis rates across gram-positive and gram-negative organisms (although the reduction was statistically significant only for non-pseudomonal gram-negative organisms).

The balANZ trial has also previously reported that patients using biocompatible fluids (compared with conventional dialysate) experienced a significantly delayed onset of anuria (27). Because the development of anuria is associated with increased mortality, technique failure, and peritonitis episodes (49,50), it is possible that a renoprotective benefit of biocompatible fluids may also have enhanced general resistance to infection, such as peritonitis. In keeping with that possibility, the balANZ trial previously demonstrated a lower risk of non-PD-related infections in the biocompatible group compared with the control group (27).

Although the initial empiric antibiotic regimens used and the microbiology of peritonitis episodes were comparable between the two groups, another novel finding of the balANZ study is that patients in the biocompatible group who experienced peritonitis spent a shorter period of time in hospital (\( p = 0.05 \)). That finding may have been related to the initially less severe peritonitis episodes in the biocompatible group, because compared with peritonitis episodes in control patients, such episodes were significantly more likely to be rated as mild in severity by attending clinicians. No interaction was observed between biocompatible fluid use and causative organism with respect to the duration of peritonitis-associated hospitalization. The effects of center-specific practices (“center effect”) on peritonitis outcomes were balanced between the intervention and control groups by stratification of randomization according to center and were additionally adjusted for in the multivariate Cox proportional hazards model analyses.

The present study had a number of limitations, the principal one being that the trial was not formally powered for peritonitis outcomes, which were secondary analyses of the balANZ trial. The possibility of type 2 statistical errors for some of the outcomes (particularly...
ACKNOWLEDGMENTS

CONCLUSIONS

ACKNOWLEDGMENTS

REFERENCES


