Regimen-related Mortality Risk in Patients Undergoing Peritoneal Dialysis Using Hypertonic Glucose Solution: A Retrospective Cohort Study

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\textbf{Objectives:} The main purpose of this study was to quantify the risk of mortality linked to various regimens of hypertonic peritoneal dialysis (PD) solution.

\textbf{Methods:} A retrospective cohort study of patients using home-based PD was carried out. The prescribed regimen of glucose-based PD solution for all patients, determined on the basis of their individual conditions, was extracted from their medical chart records. The primary outcome was death. The treatment regimens were categorized into 3 groups according to the type of PD solution used: original PD (1.5% glucose), shuffle PD (1.5 and 2.5% glucose), and serialized PD (2.5 and 4.5% glucose). Multivariate analysis (using the Weibull model) was applied to comprehensively examine survival probabilities related to the explanatory variable, while adjusting for other potential confounders.

\textbf{Results:} Of 300 consecutive patients, 38% died over a median follow-up time of 30 months (interquartile range: 15-46 months). Multivariate analysis showed that a treatment regimen with continued higher-strength PD solution (serialized PD) resulted in a lower survival rate than when the conventional strength solution was used (adjusted hazard ratio, 2.6; 95% confidence interval, 1.6 to 4.6, \(p<0.01\)). Five interrelated risk factors (age, length of time on PD, hemoglobin levels, albumin levels, and oliguria) were significant predictors contributing to the outcome.

\textbf{Conclusions:} Frequent exposure to high levels of glucose PD solution significantly contributed to a 2-fold higher rate of death, especially when hypertonic glucose was prescribed continuously.

\textbf{Key words:} Death, Glucose, Kidney diseases, Peritoneal dialysis, Survival rate, Thailand

\section*{INTRODUCTION}

Preserving the integrity of the peritoneal membrane for a longer time is one of the core goals for enhancing survival in patients undergoing long-term peritoneal dialysis (PD) [1]. Unfortunately, most of these patients encounter unavoidable and deleterious effects of glucose-based dialysate, including structural changes (deterioration of the peritoneal membrane over time), functional changes (e.g., an increased risk of developing insulin resistance and elevated blood glucose) [2-6], and even possibly mortality [5]. These negative effects have been reported to be a consequence of prolonged exposure to higher glucose concentrations [7-9]. However, although adverse effects from high glucose-based dialysis solutions are known to exist, it is still necessary in certain clinical conditions to use a conventional PD prescription, particularly in regions...
where non-glucose-based PD solutions are difficult to obtain or unaffordable due to high costs. Moreover, the difference in survival outcomes is still uncertain [10]. In keeping with the latest guidelines of the International Society for Peritoneal Dialysis (ISPD) [11,12], cautious short-term use of high-tonicity solutions is acceptable, but unambiguous and detailed dosing recommendations are not available for when use of a high-osmolarity agent is necessary. In the absence of a clearly identified preference, we assume that a wide range of prescriptions and regimens for high-osmolarity agents are used. In addition, adherence to daily adjustment of the dialysis prescription seems to be an extraordinary challenge for the at-home patient, particularly in remote areas, and most of the published work on this topic does not seem to reflect a routine clinical setting. While the diverse regimens may address the benefit-risk balance in routine clinical practice, to date, it is unclear whether there are differences between the various regimens of glucose-based PD solutions in terms of survival outcomes.

Rapid adverse events involving the peritoneal membrane may be prevented by exposure to lower doses of glucose-based PD solution. Measuring the predicted risk with various hypertonic solutions and regimens may help support clinicians’ decision-making by providing robust and rigorous evidence. Thus, this study attempted to establish the regimen-related mortality risk for alternative regimens in clinical practice, with the goal of improving survival outcomes. Therefore, the main purpose of this study was to determine the probability of risk of death in patients using different regimens of PD solution.

**METHODS**

**Study Design and Site**

This study was a retrospective cohort chart review of patients undergoing continuous ambulatory peritoneal dialysis (CAPD) who were registered with a PD facility between October 1, 2011 and September 30, 2016 and followed until January 31, 2017 at a single government-run general hospital in Thailand.

**Inclusion and Exclusion Criteria**

The eligible criteria included patients with newly initiated PD who were undergoing CAPD covered under the government’s universal coverage scheme for renal dialysis (a fairly homogeneous socioeconomic group), were aged 20 years or older, and were on a home-based PD exchange regimen. Patients were excluded from the study if they had the following conditions: polycystic kidney disease, malignancy, liver disease, were in a bedridden condition, or died within 3 months of starting PD. Patients who were unable to attend the PD center for regularly scheduled follow-up, regardless of the center protocol, were also excluded. The recorded information included demographics, prescribed PD solutions, and duration of therapy (start date, event date). A particular focus was placed on data about PD prescription regimens. Subsequently, the treatment regimens were collected and categorized according to the course of prescriptions of solutions for dialysis performed independently at home, which were used to define 3 prescribed PD regimens: the original PD group comprised patients who only underwent PD with conventional-strength glucose solution (1.5% PD solution); the shuffle PD group comprised patients who were treated with a 1.5% solution that was changed to a 2.5% solution, or cycled back and forth between the 1.5% PD and the 2.5% PD solutions for daily PD (regimen continuously used for longer than 90 consecutive days); and the serialized PD group comprised patients who serially underwent continuous dialysis with either 2.5% PD or alternated between 2.5 and 4.2% PD (similarly, with the regimen continuously used for longer than 90 consecutive days). For all regimens, the origin time was marked as the time point when exposure to the solution commenced, and the patient was followed until the outcome of interest. The primary outcome was death, and this endpoint was defined as the time of failure. Observations were censored if patients discontinued PD therapy for any reason (e.g., switching modes to either hemodialysis or transplantation, were lost to follow-up, or were alive at the end of the follow-up period).

Of the 330 patients who initiated CAPD at our study site between October 2011 and September 2016, 30 were excluded: 12 patients were younger than 20, 16 died or had a treatment change within less than 3 months, and 2 had pre-existing conditions preventing eligibility. The remaining 300 patients were enrolled and followed until January 2017, with 146 (48.7%) categorized into the original PD group, 101 (33.7%) categorized into the shuffle PD group, and 53 (17.7%) categorized into the serialized PD group.

**Clinical Data**

Either paper or electronic medical records were the primary sources of data information. Consequently, mortality status was extracted and linked with multiple data sources. According to several previous studies [13-16], age, gender, diabetes mellitus (DM), cardiovascular disease (CVD), history of peritonitis, residu-
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adjusting for other potential confounders. A backward selection procedure was employed to fit the final analysis, with the results expressed as adjusted hazard ratios (aHRs) with 95% confidence intervals (CIs), and the significance level was 0.05. We used Stata version 15 (StataCorp., College Station, TX, USA).

Ethical Statements
The study protocol was approved by both the Khon Kaen University ethics committee and the institutional review board for data collection (HE 602253 and EC 008/2016).

RESULTS

Patient Characteristics
In total, 300 consecutive patients with documented CAPD prescriptions during 6604 person-months of observation were eligible for analysis. The distribution of person-months of observation the three categories was 3673, 1823, and 1108 person-months for the original, shuffle, and serialized PD groups, respectively. The final status of patients at the end-point of the study was 62% alive and 38% who had died throughout a median follow-up time of 30 months (interquartile range, 15-46 months), based on the reverse Kaplan-Meier method [19]. The renal output (RUO), length of time on PD, and hemoglobin (Hb) and albumin (Alb) levels are well documented as potential risk factors for mortality among PD patients. Thus, these factors were gathered for use as covariates. Urine volume was measured by daily self-reporting on estimated urine output. Since these patients’ blood chemistry levels fluctuated throughout the time of the study, both serum Alb and blood Hb concentrations were collected longitudinally at regular clinic visits (typically, the follow-up visits were scheduled at 3-month intervals). Clinical presentation and positive results from dialysate fluid culture were defined as peritoneal dialysis-related peritonitis.

Based on findings from previous studies [5,17], we applied the formula used by Schoenfeld [18] to calculate sample size for both exposed and non-exposed outcomes with a significance of 0.05 and a power of 0.8; a total of 300 consecutive patients were required.

Statistical Methods
The demographic results were summarized as percentages for dichotomous variables and mean values and standard deviations for continuous variables. Multivariate analysis (using the Weibull model) was applied to comprehensively examine survival probabilities related to the explanatory variable, while adjusting for other potential confounders. A backward selection procedure was employed to fit the final analysis, with the results expressed as adjusted hazard ratios (aHRs) with 95% confidence intervals (CIs), and the significance level was 0.05. We used Stata version 15 (StataCorp., College Station, TX, USA).

Table 1. Patient characteristics according to the prescription course of PD

| Characteristics | Overall (n = 300) | Type of treatment regimen (PD) | Original (n = 146) | Shuffle (n = 101) | Serialized (n = 53) |
|-----------------|------------------|-------------------------------|-------------------|------------------|-------------------|
| Follow-up (mo)  |                  |                               |                   |                  |                   |
| Med (IQR)       | 30 (15-46)       | 32 (15-51)                    | 25 (9-35)         | 37 (26-43)       |
| Age (y)         |                  |                               |                   |                  |                   |
| Mean ± SD       | 56.3 ± 10.4      | 57.2 ± 9.8                    | 56.1 ± 10.6       | 53.9 ± 11.3      |
| Med (Min-Max)   | 57 (23-81)       | 58 (23-75)                    | 56 (26-79)        | 54 (26-81)       |
| Gender (men)    | 154 (51.3)       | 76 (52.1)                     | 54 (53.4)         | 24 (45.2)        |
| RUO (mL)        |                  |                               |                   |                  |                   |
| Mean ± SD       | 366.2 ± 316.8    | 411.9 ± 317.2                 | 366.3 ± 327.8     | 239.6 ± 260.4    |
| Med (Min-Max)   | 300 (0-1000)     | 400 (0-1000)                  | 300 (0-1000)      | 200 (0-1000)     |
| Hb (g/dL)       |                  |                               |                   |                  |                   |
| Mean ± SD       | 9.8 ± 1.5        | 9.9 ± 1.5                     | 9.6 ± 1.4         | 9.6 ± 1.5        |
| Med (Min-Max)   | 9.8 (4.2-16.0)   | 9.8 (4.2-16.0)                | 9.6 (4.2-12.6)    | 9.6 (6.7-13.0)   |
| Alb (g/dL)      |                  |                               |                   |                  |                   |
| Mean ± SD       | 2.3 ± 0.6        | 2.4 ± 0.6                     | 2.4 ± 0.6         | 2.1 ± 0.7        |
| Med (Min-Max)   | 2.4 (0.4-3.9)    | 2.4 (0.7-3.9)                 | 2.4 (0.7-3.9)     | 2.1 (0.4-3.4)    |
| DM (yes)        | 112 (37.3)       | 50 (34.2)                     | 43 (42.5)         | 19 (35.8)        |
| CVD (yes)       | 18 (6.0)         | 6 (4.1)                       | 7 (6.9)           | 5 (9.4)          |
| PDitis (yes)    | 157 (52.3)       | 60 (41.1)                     | 58 (57.4)         | 39 (73.5)        |
| TiPD (mo)       |                  |                               |                   |                  |                   |
| Mean ± SD       | 26.5 ± 16.2      | 25.1 ± 17.1                   | 27.1 ± 14.9       | 29.0 ± 15.8      |
| Med (Min-Max)   | 24 (3-64)        | 19.5 (3-60)                   | 27 (4-62)         | 25 (5-64)        |

Values are presented as number (%).
PD, peritoneal dialysis; Med, median; IQR, interquartile range (25th-75th); SD, standard deviation; Min, minimum; Max, maximum; RUO, residual urine output; Hb, hemoglobin; Alb, albumin; DM, diabetes mellitus; CVD, cardiovascular disease; PDitis, history of peritoneal dialysis-related peritonitis; TiPD, length of time on PD.
leading causes of death were sepsis, CVD, electrolyte imbalance, and pneumonia. Table 1 shows the characteristics of the study population. In all 3 groups, the age profiles and gender distribution were quite similar. Risk factors, including the prevalence of CVD and DM, and clinical data, such as Alb and Hb levels, were similar among the 3 groups. However, a history of peritonitis was more common in the shuffle PD group than the original PD group, and still higher in the serialized PD group (overall presence in more than 50% of the participants). Around one-third of the patients were diagnosed with pre-existing DM. The majority of patients underwent PD treatment for longer than 2 years in all 3 groups, with an average duration of PD treatment of 25.1, 27.1, and 29.0 months in the original, shuffle, and serialized PD groups, respectively.

**Univariate Analysis of Risk Factors**

In the univariate analysis shown in Table 2. Among the confounding factors, higher age, lower Hb levels, lower Alb levels, lower RUO, and the presence of DM and CVD were significantly associated with an increased risk of all-cause mortality. Hb levels were a risk factor because many enrolled patients developed chronic anemia. Every 1 g/dL increase in Hb reduced risk by 20%. For the analysis, we dichotomized Hb concentrations as lower or higher than 10 g/dL. We found that patients who had Hb levels ≥10 g/dL had a 53% lower risk of death. When we compared Alb levels, there was an almost 75% decreased risk for every increase of 1 g/dL. For analysis, we dichotomized participants based on Alb levels, regardless of persistent hypalbuminemia, using a cut-off point of 2.5 g/dL. We found patients with Alb levels >2.5 g/dL had an hazard ratio (HR) of 0.2. Survival time significantly increased by 3% for every 1 mL increase in urine output per day; however, the 95% confidence limit was extremely narrow. Thus, patients were categorized as having either oliguria or anuria (urine volume between 0-200 mL/d) or a urine output ≥200 mL/d. Patients with anuria/oliguria had a higher risk of death than patients with a urine volume ≥200 mL/d. The presence of CVD in PD patients was associated with a 2.7 times higher risk of mortality. For each additional month of the patient’s length of time undergoing PD, the risk of death decreased by 17% (HR, 0.8).

As the treatment regimen was the main point of interest, the median survival times were 48.9, 43.7, and 25.1 months, and the mortality rate for each regimen was 1.3, 1.8, and 2.7 per 100 person-months in the original, shuffle and serialized PD groups, respectively. The crude HRs associated with the

### Table 2. Univariate analysis of potential factors affecting all-cause mortality in PD patients

| Variables                        | Median time (mo) | Person-time (mo) | HR (95% CI)  |
|----------------------------------|-----------------|-----------------|--------------|
| Treatment regimen (PD)           |                 |                 |              |
| Original                         | 48.9            | 3673            | 1.0 (reference) |
| Shuffle                          | 43.7            | 1823            | 1.5 (0.9, 2.3) |
| Serialized                       | 25.1            | 1108            | 2.2 (1.3, 3.4)** |
| Gender                           |                 |                 |              |
| Men                              | 44.0            | 3545            | 1.0 (reference) |
| Women                            | 35.0            | 3059            | 1.1 (0.8, 1.6)** |
| Age (each year increase, y)      |                 |                 |              |
| <60                              | 44.0            | 4064            | 1.0 (reference) |
| ≥60                              | 25.0            | 2540            | 1.3 (0.9, 1.9)** |
| RUI (mL/d)                       |                 |                 |              |
| <200                             | 26.3            | 2013            | 1.0 (reference) |
| ≥200                             | 48.6            | 4591            | 0.4 (0.3, 0.7)** |
| Hb (g/dL)                        |                 |                 |              |
| ≤10                              | 29.0            | 3306            | 1.0 (reference) |
| ≥10                              | 56.7            | 3298            | 0.4 (0.3, 0.6)** |
| Serum albumin (g/dL)             |                 |                 |              |
| ≤2.5                             | 22.7            | 2672            | 1.0 (reference) |
| >2.5                             | >60.0           | 3932            | 0.2 (0.1, 0.3)** |
| DM (%)                           |                 |                 |              |
| No                               | 43.9            | 4387            | 1.0 (reference) |
| Yes                              | 30.2            | 2217            | 1.5 (1.0, 2.2)** |
| CVD                              |                 |                 |              |
| No                               | 38.5            | 6463            | 1.0 (reference) |
| Yes                              | 31.8            | 141             | 2.7 (1.7, 5.0)* |
| PDitis                           |                 |                 |              |
| No                               | 54.0            | 2768            | 1.0 (reference) |
| Yes                              | 28.0            | 3836            | 1.4 (0.9, 2.0)** |
| TIPD (each month increase, mo)   |                 |                 |              |
| No                               | 0.8             | 0.8             | 0.8 (0.8, 0.9)** |
| p<0.1, *p<0.05, **p<0.01.       |                 |                 |              |

PD, peritoneal dialysis; HR, hazard ratio; CI, confidence interval; RUO, residual urine output; Hb, Hemoglobin; DM, diabetes mellitus; CVD, cardiovascular disease; PDitis, history of peritoneal dialysis-related peritonitis; TIPD, length of time on PD.

Multivariate Analysis

For the multivariate analysis (using the Weibull model), we used a backward selection strategy to comprehensively examine survival probabilities related to the intervening variable while adjusting for other potential confounders; the final model is shown in Table 3. The selected covariates that were employed to probe the interrelationships of potential risk factors were age, gender, peritonitis, DM, RUO, Hb, and Alb levels, and length of time on PD.

In the final model (Table 3), patients with PD who were pre-
Prescribed PD Dialysate

scribed a continuous exchange of hypertonic glucose solutions were 2.6 times (95% CI, 1.6 to 4.6; \(p < 0.01\)) more likely to die than patients who did not receive hypertonic exchanges. Similarly to the serialized, the shuffle PD regimen was found to pose a significant risk (aHR, 2.2; 95% CI, 1.4 to 3.4; \(p < 0.01\)) when compared with the original PD group. Figure 1 shows the survival curves of patients undergoing different PD regimens according to the Weibull model, indicating that after patients received treatment with either intermittent (shuffle) or continuous (serialized) high-glucose dialysate regimens for around 20 months, the probability of survival was markedly lower than among those undergoing treatment with low-concentration glucose dialysate (original).

Additionally, for all treatment regimens, 5 interrelated risk factors emerged as significant predictors contributing to the outcomes: low Hb concentration, low Alb level, anuria or oliguria, advanced age, and short duration of receiving PD.

**DISCUSSION**

This investigation focused on survival according to treatment regimens. Multivariate analysis showed that treatment regimens using either a combination of concentrations (1.5 and 2.5% PD) regimen (shuffle) or a continuous higher-concentration PD solution for exchange (serialized PD) was associated with an approximately 2-fold greater attenuation of survival than the conventional concentration regimen. The percentage of overall deaths in patients undergoing PD throughout the study period is similar to that reported in a previous longitudinal study [20]. We found that sepsis was the leading cause of death. Sepsis resulting from peritonitis is a possible explanation for this finding, because more than half of the patients in the current study had a history of PD-related peritonitis.

The etiological mechanism of increased mortality in patients prescribed high-glucose dialysate is still uncertain. Previous work [17,21,22] has only proposed theoretical possibilities for factors influencing patient survival, and some of these studies did not yield convincing conclusions regarding the heightened risks of negative outcomes. We suggest that a plausible causal pathway was described by Hassan et al. [23]. It begins with peritoneal glucose dialysate-induced inflammation of the peritoneum, which may even progress to fibrosis [2], and metabolic effects from glucose loading, leading to a reduction in ultrafiltration and clearance, which is then dynamically linked to the occurrence of cardiovascular problems [24]. In light of our main findings, the following points briefly clarify our hypothesis. First, the destruction of peritoneal mesothelial cells has been reported in previous studies, and in particular, it has found to be higher after exposure to 4.2% dextrose [3,25]. Kang et al. [26] found that various incremental ranges in glucose concentrations, including 30, 60, and 90 mmol/L, caused 2.3-, 3.6-, and 4.0-fold increases in the induced fibrosis factor, respectively. Combining these findings with other theoretical suggestions, it can be inferred that the proportional increase was induced by the incremental changes in concentration [27,28]. Likewise, previous studies [4,29] highlighted destruction caused by glucose toxicity. Second, several studies have

**Table 3. Potential variables fit to a Weibull distribution according to treatment regimen**

| Treatment variables | aHR (95%CI)\(^1\) |
|---------------------|--------------------|
| **Treatment regimens (PD)** |                   |
| Original            | 1.0 (reference)    |
| Shuffle             | 2.2 (1.4, 3.4)**   |
| Serialized           | 2.6 (1.6, 4.6)**   |
| Hb (each unit increase, g/dL) |               |
| <10                 | 0.8 (0.7, 0.9)*    |
| ≥10                 | 1.0 (reference)*   |
| Alb (each unit increase, g/dL) |               |
| <2.5                | 0.6 (0.4, 0.8)**   |
| ≥2.5                | 1.0 (reference)**  |
| RUO (mL/d)          |                   |
| <200                | 0.6 (0.4, 0.8)*    |
| ≥200                | 1.0 (reference)    |
| Age (each year increase, y) |       |
| 1.0 (1.0, 1.1)*     |
| TiPD (each month increase, mo) |        |
| 0.8 (0.8, 0.9)**    |

\(\text{aHR, adjusted hazard ratio; CI, confidence interval; PD, peritoneal dialysis; Hb, hemoglobin; Alb, albumin; RUO, residual urine output; TiPD, length of time on PD.}\)

\(\text{1Adjusted for gender, peritonitis, diabetes mellitus, and cardiovascular disease.}\)

\(\text{*p < 0.05, **p < 0.01.}\)
pointed out that hypertonic PD solutions also affect systemic hemodynamic status [30]. Patients whose peritoneal physical characteristics changed as a result of glucose uptake showed poor mortality outcomes in an observational cohort study [31]. Our finding is also in agreement with a number of clinical research studies, including those by Krediet et al. [32], and Li and Chow [33]. They asserted that exposure to glucose-based PD significantly increased the risk of mortality by CVD. Unfortunately, the cause of at-home deaths could not be determined in our study, because a category of assumed general-cause mortality was mostly used for deaths outside the hospital; therefore, the number of CVD deaths may have been underestimated. In addition, we further discovered that the median survival time of the original PD group was around 48 months, which was longer than that of the high-glucose-concentration group by 25 months. This finding is in agreement with the previous study by Korbet and Rodby [34]. In their work, peritoneal transport properties during the first 2 to 4 years of PD normally remained stable. Later, patients who had undergone PD treatment for more than 4 years had the potential to experience a change in peritoneal solute transport and a concomitant loss of ultrafiltration. Similarly, a population-based cohort study [35], despite a relatively small sample size, found that it was possible to preserve peritoneal function for more than 5 years in patients who were given only low-strength glucose solutions, whereas there was a negative outcome when higher-tonicity solutions were used. To summarize, these facts support and could theoretically explain why continuous exposure to the highly hypertonic solution (serial-ized PD) was the factor most closely associated with an increased risk of death in our analysis.

Nonetheless, avoiding the use of highly hypertonic glucose solutions is not possible for glucose-based PD modalities; however, the shuffle PD regimen, despite its increased mortality risk, may be an alternative choice considering the recommendations in both ISPD guidelines [11,12]. These recommendations for the temporary use of hypertonic solutions are in line with the shuffle PD regimen, in which patients were able to survive longer than those who received a highly hypertonic glucose solution. Many regimens attempt to slow the progressive deleterious effects from higher-concentration glucose solutions, such as the combination exchange regimen, which alternates 1.5% dextrose for 3 exchanges with 1 exchange with 4.2% dextrose, as described in the study of Holmes and Shockley [36]. Regarding this, our results suggest that the shuffle regimen, with alternations between 1.5 and 2.5% PD solution, yielded a lower risk than 4.2% dextrose. This regimen may be an effective choice to reduce adverse outcomes. Therefore, our finding can bridge the gap between clinical practice and theoretical suggestions. The continuous use of a long-term highly hypertonic solution is a modifiable factor that affects PD survival, and continuous exchange at higher concentrations should be cautioned against as a first regimen of treatment.

This study has some limitations. First, only one commercial PD solution was available for this study, and it was carried out at a single center; therefore, the possible effects of divergent dialysate compositions and between-center differences cannot be dismissed. Second, we were unable to quantify the exact ultrafiltration loss and severity of peritoneal transport deterioration because of the absence of complete data from the peritoneal equilibration test. Third, data regarding the underlying causes of chronic kidney disease were not available at sufficient levels for analysis, because most of the causes were classified as unknown disease and DM. Therefore, this parameter was not useful for interpreting our findings. Finally, accounting for competing risks, such as switching treatment to hemodialysis, should be considered for further analyses to assess the accuracy of the current results.

In conclusion, evidence from the current analysis affirmed that frequent and continuous exposure to high levels of glucose PD solution significantly contributed to the risk of mortality, with a 2-fold increased risk of survival attenuation. No standards for prescriptions exist, and the higher-concentration solutions are often ordered. These main findings are widely applicable and are indicative of the benefit-risk balance of this issue in many parts of the world. The present study also proposes a simple alternative strategy for avoiding this effect by interrupting continuity in daily exchanges. We urge caution in the use of high-glucose-based solutions in clinical practice.

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**CONFLICTS OF INTEREST**

The authors have no conflicts of interest associated with the material presented in this paper.
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