NKF-DOQI CLINICAL PRACTICE GUIDELINES
FOR PERITONEAL DIALYSIS ADEQUACY
**Acronyms and Abbreviations**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Term</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACE</td>
<td>angiotensin-converting enzyme</td>
</tr>
<tr>
<td>BSA</td>
<td>body surface area</td>
</tr>
<tr>
<td>BUN</td>
<td>blood urea nitrogen concentration (an old term; SUN is technically correct, see below)</td>
</tr>
<tr>
<td>CANUSA</td>
<td>Canada/USA Peritoneal Dialysis Study</td>
</tr>
<tr>
<td>CAPD</td>
<td>continuous ambulatory peritoneal dialysis</td>
</tr>
<tr>
<td>CCPD</td>
<td>continuous cycling peritoneal dialysis</td>
</tr>
<tr>
<td>CCr</td>
<td>creatinine clearance</td>
</tr>
<tr>
<td>C&lt;sub&gt;r&lt;/sub&gt; Cr</td>
<td>residual renal creatinine clearance</td>
</tr>
<tr>
<td>DI</td>
<td>dialysis index</td>
</tr>
<tr>
<td>DPI</td>
<td>dietary protein intake</td>
</tr>
<tr>
<td>eC</td>
<td>effective clearance</td>
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<tr>
<td>ESRD</td>
<td>end-stage renal disease</td>
</tr>
<tr>
<td>GFR</td>
<td>glomerular filtration rate</td>
</tr>
<tr>
<td>HD</td>
<td>hemodialysis</td>
</tr>
<tr>
<td>K&lt;sub&gt;p&lt;/sub&gt; Cr</td>
<td>peritoneal creatinine clearance</td>
</tr>
<tr>
<td>K&lt;sub&gt;p&lt;/sub&gt;/V&lt;sub&gt;urea&lt;/sub&gt;</td>
<td>the peritoneal component of Kt/V&lt;sub&gt;urea&lt;/sub&gt;</td>
</tr>
<tr>
<td>K&lt;sub&gt;p&lt;/sub&gt;&lt;sub&gt;r&lt;/sub&gt;/V&lt;sub&gt;urea&lt;/sub&gt;</td>
<td>the sum of peritoneal and renal Kt/V&lt;sub&gt;urea&lt;/sub&gt;. These terms are interchangeable in that Kt/V&lt;sub&gt;urea&lt;/sub&gt; is total unless otherwise noted.</td>
</tr>
<tr>
<td>Kt/V&lt;sub&gt;urea&lt;/sub&gt;</td>
<td>urea clearance × time normalized by total body water, the volume of distribution of urea</td>
</tr>
<tr>
<td>K&lt;sub&gt;p&lt;/sub&gt;&lt;sub&gt;r&lt;/sub&gt;/V&lt;sub&gt;urea&lt;/sub&gt;</td>
<td>the renal component of Kt/V&lt;sub&gt;urea&lt;/sub&gt;</td>
</tr>
<tr>
<td>MTC</td>
<td>mass transfer coefficient</td>
</tr>
<tr>
<td>n</td>
<td>normalized</td>
</tr>
<tr>
<td>nBSA</td>
<td>normalized body surface area</td>
</tr>
<tr>
<td>NIPD</td>
<td>nightly intermittent peritoneal dialysis</td>
</tr>
<tr>
<td>nPCR</td>
<td>normalized protein catabolic rate</td>
</tr>
<tr>
<td>nPNA</td>
<td>normalized protein equivalent of total nitrogen appearance</td>
</tr>
<tr>
<td>nV</td>
<td>normalized volume</td>
</tr>
<tr>
<td>PCR</td>
<td>protein catabolic rate</td>
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<tr>
<td>PD</td>
<td>peritoneal dialysis</td>
</tr>
<tr>
<td>PET</td>
<td>peritoneal equilibration test</td>
</tr>
<tr>
<td>PNA</td>
<td>protein equivalent of nitrogen appearance</td>
</tr>
<tr>
<td>QOL</td>
<td>quality of life</td>
</tr>
<tr>
<td>RRF</td>
<td>residual renal function</td>
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<tr>
<td>SGA</td>
<td>subjective global assessment</td>
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<tr>
<td>SHR</td>
<td>standardized hospitalization rates</td>
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<tr>
<td>SUN</td>
<td>serum urea nitrogen concentration</td>
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<tr>
<td>t</td>
<td>time</td>
</tr>
<tr>
<td>UKM</td>
<td>urea kinetic modeling</td>
</tr>
<tr>
<td>UNA</td>
<td>urea nitrogen appearance</td>
</tr>
<tr>
<td>URR</td>
<td>urea reduction ratio</td>
</tr>
<tr>
<td>USRDS</td>
<td>United States Renal Data System</td>
</tr>
<tr>
<td>V</td>
<td>volume of distribution. When referring to urea, this is total body water</td>
</tr>
</tbody>
</table>
Introduction

This Work Group was charged with preparing practice guidelines for the "adequacy of peritoneal dialysis," a topic which could be defined broadly or narrowly. The Work Group elected to focus its guidelines on those areas of "adequacy" that needed the most urgent development, knowing that subsequent guidelines will be developed or that others were currently under development (e.g., for management of peritonitis). In addition, the Work Group focused on topics for which guidelines would likely have the greatest impact on patient outcomes. However, the Work Group's focus should not be construed to mean that areas not covered are unimportant.

Some external reviewers criticized these guidelines as too complex, while others wrote that they were not thorough enough. Some wanted guidelines merged, and others thought the guidelines were too dense. The Work Group considered all these issues.

We advise the reader to first become familiar with the Table of Contents, which provides a listing of the Clinical Practice Guidelines for Peritoneal Dialysis Adequacy; detailed rationales are provided for each guideline. Redundancies are often intentional because it is anticipated that a reader might review only selected topics. However, the Work Group considers these guidelines as best viewed in their entirety, rather than in their component parts.

There is a paucity of data on children in the areas covered by these guidelines. Pediatricians were represented on the Work Group, and outside pediatric consultations were obtained. Because some recommendations for adults do not apply to children, additional recommendations are included when appropriate for pediatric patients. For the purpose of these guidelines, a child was considered to be a patient less than 19 years of age.

An "effective dose" is that which achieves its stated goal. That goal is some form of outcome measure(s), and could be determined by patient, provider, payer, regulator or a combination of these parties. At the lower extreme is the "minimal effective dose." In certain circumstances this may be interpreted as "adequate." At the other extreme is the "maximal effective dose," the dose above which there are no additional benefits. For hemodialysis and peritoneal dialysis the maximal effective dose is not known. Somewhere between these extremes is the "optimal dose," the dose above which the additional derived benefit does not justify the additional cost or burden. If one accepts this definition, the Work Group intended to more precisely define "optimal dose" targets in a clinically relevant and quantitative fashion. It was the intention of the Work Group to bring "adequate dose" to the level of "optimal dose" by raising the outcome goals or expectations. The present guidelines attempt to make recommendations based on available scientific/medical evidence, resorting to expert opinion only when necessary. It is clearly stated in each guideline title when recommendations were based on evidence, opinion, or both. Even when guidelines were based on opinion, that opinion is supported by direct or extrapolated evidence.

These guidelines are intended for use by health care professionals trained to understand variations in the practice of medicine and the necessity for such variation. These guidelines are not intended for punitive use by any oversight official who does not understand the reasons or the necessity for practice variations including variations in societies different from that of the United States.
I. Initiation of Dialysis

BACKGROUND

Two clinical guidelines for when to initiate dialysis are provided because there appear to be two independent predictors of clinical outcome. The first guideline is based on the level of renal function (as measured by Kt/V urea per week); the second is based on the level of protein intake (as estimated from measured 24-hour urinary urea nitrogen output and estimated non-urea nitrogen losses).

Although less than 1% of American dialysis patients begin dialysis with a serum creatinine concentration <8.0 mg/dL or a Cr > 10 mL/min, approximately 60% suffer from nausea/vomiting at the time of dialysis initiation. Thus, the likelihood of malnutrition in this population is high. Evidence from the Modification of Diet in Renal Disease (MDRD) study and a recent large Australian study clearly show that when glomerular filtration rate (GFR) decreases to 25 to 50 mL/min, patients adapt by reducing their protein intake. Protein intake continues to decline as renal disease progresses to end stage. These observations have been corroborated in a prospective study. As renal function deteriorates, protein and energy intake decreases, leading to changes in body weight, fat mass, serum albumin, and transferrin concentrations. Earlier initiation of dialysis may prevent or perhaps even reverse this deterioration in nutritional status. Increased serum albumin concentration has been shown to parallel the increase in Kt/V urea for HD patients. In addition, an editorial review cited data suggesting that albumin level at initiation of dialysis is predictive of survival.

Adverse clinical and economic consequences of failure to properly manage patients as they approach ESRD and dialysis were first described in Britain. These observations have now been corroborated in other regions of Britain, the United States, and France. Specifically, costs, hospitalization, and morbidities decrease if attention is paid to nutrition, acid-base status, hypocalcemia, hyperphosphatemia, anemia, hypertension, volume status, and dialysis access (vascular or peritoneal). Hence, it is likely that delays in referral for initiation of dialysis result in unnecessary morbidity and potentially higher costs as well. While seeing a nephrologist does not guarantee that patients will be adequately prepared and referred for dialysis, it dramatically increases the likelihood that this will occur.

GUIDELINE 1

When to Initiate Dialysis—Kt/V urea Criterion (Opinion)

Unless certain conditions are met, patients should be advised to initiate some form of dialysis when the weekly renal Kt/V urea (Kt/V urea) falls below 2.0. The conditions that may indicate dialysis is not yet necessary even though the weekly Kt/V urea is less than 2.0 are:

1. Stable or increased edema-free body weight. Supportive objective parameters for adequate nutrition include a lean body mass >63%, subjective global assessment score indicative of adequate nutrition (see Guideline 12: Nutritional Status Assessment and Appendix B: Detailed Rationale for Guideline 2) and a serum albumin concentration in excess of the lower limit for the lab, and stable or rising; and
2. nPNA >=0.8 g/kg/d (see Guideline 2: When To Initiate Peritoneal Dialysis—nPNA Criteria and Appendix B: Detailed Rationale for Guideline 2); and
3. Complete absence of clinical signs or symptoms attributable to uremia.

A weekly Kt/V urea of 2.0 approximates a renal urea clearance of 7 mL/min and a renal creatinine clearance that varies between 9 to 14 mL/min/1.73 m³. Urea clearance should be normalized to total body water (V) and creatinine clearance should be expressed per 1.73 m² of body surface area. The GFR, which is estimated by the arithmetic mean of the urea and creatinine clearances, will be approximately 10.5 mL/min/1.73 m² when the Kt/V urea is about 2.0.

Rationale A detailed rationale is described in Appendix A. The following is a summary.

It is paradoxical that nephrologists have focused on optimizing urea clearance once patients are started on dialysis, but have accepted much lower levels of renal urea clearance during the pre-dialysis phase of patient management. For
example, a weekly total (residual renal plus peritoneal dialysis) \( Kt/V_{urea} (K_{pr}V_{urea}) \) of 2.0 or higher is associated with improved outcomes in patients on PD (see Guideline 15: Weekly Dose of CAPD), yet dialysis is usually not initiated until weekly \( Kt/V_{urea} \) falls to the range of 0.71 to 1.3. There is no definitive direct proof for the belief that a given level of urea clearance by the kidney is associated with better control of uremia than PD with this same urea clearance. In fact, recent studies suggest that the relationship between protein intake and weekly \( Kt/V_{urea} \) is nearly identical in patients with chronic renal failure not yet on dialysis and in patients on PD. Thus, until proven otherwise, residual renal and peritoneal clearances of small solutes should be considered equivalent.

Once \( Kt/V_{urea} \) falls below 2.0 per week, patients should be considered at increased risk for malnutrition and uremic complications. With further decreases in \( Kt/V_{urea} \) in the absence of renal replacement therapy, the risk increases. Dialysis, or some form of renal replacement therapy, should be strongly considered when \( Kt/V_{urea} \) falls below 2.0 (or \( C_{cr} \) falls in the range of 9 to 14 mL/min/1.73 m\(^2\)) and definitely implemented if:

1. There has been an unintentional decrease in edema-free body weight. Supportive objective parameters for malnutrition include a lean body mass <63%, subjective global assessment score indicative of malnutrition (see Guideline 12: Nutritional Status Assessment and Appendix B: Detailed Rationale for Guideline 2) and a serum albumin concentration that is decreasing even within the normal range or is less than the lower limit for normal for that lab; or
2. nPNA <0.8 g/kg/d due to spontaneous, unplanned, and unsupplemented protein restriction (see Guideline 2: When To Initiate Peritoneal Dialysis—nPNA Criteria and Appendix B: Detailed Rationale for Guideline 2); or
3. There are clinical signs or symptoms attributable to uremia.

If PD is initiated, the \( K_{pr}V_{urea} \) could be increased incrementally so the combined weekly value of \( K_{pr}V_{urea} + K_{pr}V_{urea} \) (\( K_{pr}V_{urea} \) or total \( Kt/V_{urea} \)) does not fall below the target level of 2.0. With the incremental initiation approach frequent measurement of residual renal function (RRF) will be necessary to assure that total delivered solute removal does not drop below targets (see Guidelines 3: Frequency of Delivered PD Dose and Total Solute Clearance Measurement Within Six Months of Initiation and Guideline 5: Frequency of Measurement of \( Kt/V_{urea} \), Total \( C_{cr} \), nPNA, and Total Creatinine Appearance). Alternatively, the initiation of a “full dose” of PD may be offered (equivalent of four 2 L exchanges per day, which may yield a weekly \( K_{pr}V_{urea} \) of 1.5 to 2.0, depending on transport characteristics, ultrafiltration, and body size). With initiation of “full dose” PD, frequency of measurement of RRF can be less intense.

The Work Group strongly supports the opinion that the PD outcome data for a weekly \( Kt/V_{urea} \) of ≥2.0 are so compelling that using the same figure for initiation of dialysis justifies the small risks of performing peritoneal dialysis. Those risks include infections and the possibility that increasing the length of time on PD contributes to eventual patient “burn-out.” If a patient is suspected to be at high risk for these complications, PD may not be the best choice for renal replacement therapy. The Work Group acknowledges that the risks of early initiation of PD are not clearly known, but that the risks of late initiation are known and are unacceptable. Furthermore, not knowing which initiation strategy (incremental versus full therapy initiation) is better, the Work Group recommends that either approach be used to reach or exceed targets.

Compared to CAPD, it is more complex to calculate the incremental dose of hemodialysis (HD) that would be needed such that the total continuous delivered weekly \( Kt/V_{urea} \) would be greater than 2.0. However, it can be estimated using the fundamental assumption underlying CAPD, that at the same protein catabolic rate, continuous renal replacement therapy must keep the steady state BUN equal to the average pre-hemodialysis BUN (see Appendix G for further discussion of this assumption). If weekly \( Kt/V_{urea} \) is 1.6 , for example, a one time per week HD treatment must deliver an equilibrated (double-pool) \( Kt/V_{urea} \) of 2.0 to achieve a total continuous weekly \( Kt/V_{urea} \) equivalent to 2.0. This is quite difficult to achieve, so two HD treatments per week may be more realistic for this level of RRF. If weekly \( Kt/V_{urea} \) is 0.5, two HD treatments must each deliver an equilibrated (double-pool) \( Kt/V_{urea} \) of 2.0 to achieve a total continuous weekly \( Kt/V_{urea} \) equivalent to 2.0. This is also quite difficult to achieve, so three HD treatments.
per week may be more realistic for this level of RRF. More technical details about intermittent HD are described in Appendix A, including the role of biocompatible membranes to help preserve RRF.

It is a general consensus that patients with diabetes should initiate dialysis at levels of RRF higher than in patients with causes of ESRD other than diabetes. That practice is not altered by this guideline.

The Work Group also recognizes that for many clinicians, initiating dialysis based on $Kt/V_{urea}$ is a new concept. Therefore, we have attempted to equate this to the traditional measure of urea clearance, $C_{Cr}$, and GFR (estimated by the arithmetic mean of urea and creatinine clearance).

The Work Group recognizes that the patient will play a major role in accepting the initiation of dialysis when the above conditions become applicable. In particular, the nephrologist must explain to the patient the risk of malnutrition with delayed initiation of dialysis and the strong inferential evidence that survival might be improved with an earlier start of dialysis. Thus, appropriate patient education regarding an informed decision about dialysis is necessary. Medical conditions that may explain why dialysis is not being initiated when weekly $Kt/V_{urea}$ is less than 2.0 need to be documented. These conditions are described above.

Some individuals have expressed concern that this guideline will run afoul of the Health Care Financing Administration (HCFA) regulations regarding the initiation of dialysis (e.g., form 2728, ESRD Medicare Medical Evidence Report). The leadership of the NKF-DOQI is working with HCFA to ensure that this will not be the case.

GUIDELINE 2

When to Initiate Dialysis—nPNA Criterion (Opinion)

It is recognized that an adequately supplemented, tightly monitored, low protein diet may slow the progression of renal failure in certain circumstances. When properly performed, such diets do not result in loss of lean body mass or other manifestations of malnutrition. However, progressive renal failure is associated with spontaneous anorexia and malnutrition. Under these circumstances and in the absence of comorbid causes of anorexia, and after unsuccessful intervention by a registered dietitian, dialysis should be started in adult patients when nPNA spontaneously falls below 0.8 g/kg/d.

Rationale A detailed rationale is presented in Appendix B. The following is a summary.

Decreasing renal function in chronic renal failure is associated with spontaneous reductions in protein intake and deterioration in nutritional status. Worse nutritional status at the initiation of chronic dialysis is predictive of poorer clinical outcomes on PD over the subsequent months. A weekly $K_{nt}V_{urea}$ of $\geq 2.0$ is typically required to maintain a nPNA of 0.9 g/kg/d in CAPD patients or those with chronic renal failure. In chronic renal failure with negligible urinary protein losses, many patients are at increased risk for protein malnutrition at a level of protein intake less than 0.8 g/kg/d. It is recognized that an adequately supplemented, tightly monitored, low protein diet may slow the progression of renal failure in certain circumstances and may delay the need for dialysis. When properly performed, such diets do not result in loss of lean body mass and are associated with other features of malnutrition. A registered dietitian should be involved in the care of these patients and caloric intake should exceed 35 kcal/kg/d if the diet was prescribed or designed specifically for a protein intake <0.8 g/kg/d (a planned low-protein diet). With urinary protein losses in the nephrotic range and/or PD dialysate protein losses, a nPNA of 0.9 g/kg/d or more may be necessary to maintain nitrogen balance. Comorbid causes of anorexia such as gastroparesis, infection, acidosis, and depression should be identified and treated. However, if these measures fail to restore nPNA to 0.8 g/kg/d or higher, the Work Group recommends that dialysis be initiated to maintain $Kt/V_{urea}$ targets that are usually associated with adequate levels of protein intake.

The recommendation to use an nPNA criterion for the initiation of dialysis does not imply that a broader view of nutritional status should be neglected. On the contrary, the presence of symptoms, such as the following, are all factors that should influence the clinical judgment regarding the need for dialysis:
• anorexia
• weight loss
• nausea
• vomiting
• falling caloric or protein intake by history
• decreased fat-free, edema-free (lean) body mass by creatinine kinetics (see Guideline 17: Determining Fat-Free, Edema-Free Body Mass) or other measures
• declining subjective global assessment of nutritional status (see Appendix F: Detailed Rationale for Guideline 15)
• decreasing serum albumin concentration

A critical value of nPNA below which dialysis should be started has not been identified in children. However, nutritional status, specifically as it pertains to growth, should be considered a mandatory part of each child’s evaluation.

The registered dietitian from the dialysis facility to whom the patient will be referred should be involved in the care of the patient at these early pre-dialysis stages. Doing so will provide continuity of nutritional care for the patient as he/she enters the dialysis program and assures that a registered nurse is actually available.

RECOMMENDATIONS FOR RESEARCH

Although data from available studies indirectly support our recommendation to initiate dialysis when residual renal Kt/V urea falls below 2.0 per week, this recommendation can only be validated by a prospective, randomized controlled trial in which selected weekly levels of Kt/V urea at the start of dialysis (e.g., Kt/V urea 1.0 and Kt/V urea 2.0) are related to clinical outcomes.

In addition, while available retrospective and prospective cohort studies strongly suggest that dialytic intervention should be initiated before nutritional status is compromised (i.e., when nPNA <0.8 g/kg/d), this issue will not be resolved until a prospective randomized trial is undertaken which allows comparison of the clinical outcomes of patients starting on dialysis at nPNA levels below 0.8 g/kg/d to those whose levels are significantly higher, e.g., ≥1.0 g/kg/d. Data from the CANUSA Study suggest that a follow-up period of about 2 years would be adequate for such a study, assuming a sufficient number of patients are enrolled in the trial.
II. Measures of Peritoneal Dialysis Dose

GUIDELINE 3

Frequency of Delivered PD Dose and Total Solute Clearance Measurement Within Six Months of Initiation (Opinion)

The total solute clearance (delivered PD dose plus residual renal function) should be measured at least twice and possibly three times within the first 6 months after initiation of PD. For patients initiating dialysis for the first time and/or patients with substantial residual renal function, the first measurement should be performed approximately 2 to 4 weeks after initiation of PD. For patients transferring from another renal replacement therapy to PD and/or for patients who do not have substantial residual renal function, the first measurement of delivered dose of PD should be made by 2 weeks after initiation of PD. To establish a baseline, at least one and possibly two additional measurements will need to be performed in the subsequent 5 months. The frequency of measurement of residual renal function depends on the PD prescription of incremental vs. full dose (see Table II-I).

Rationale

Adequate total solute clearance (delivered dose of PD plus residual renal function) will improve patient outcomes (see Guideline 15: Weekly Dose of CAPD). To assure delivery of adequate solute clearance, measurements for solute clearance are required. In dialysis, as in other human endeavors, continuous education and repetition of a process diminish the frequency of errors, or at least increase the likelihood of recognition of such errors and compensation for them. Furthermore, physiological variations occur which must be taken into account. These lines of reasoning apply to measurement of the dose of PD. Thus, measurement of delivered PD dose should be repeated periodically. The recommendation to measure $C_r$ and $K_t/V_{urea}$ three times within the first 6 months relates to items discussed in Guideline 7: PD Dose Troubleshooting, specifically, establishing a baseline creatinine excretion and following residual renal function (RRF). The rationale for three measurements in the first 6 months is to establish a more accurate baseline excretion of creatinine.

Two measurements within the first 6 months are probably sufficient if the results are similar. Based on our collective personal experience, the Work Group believes that patient compliance with a prescribed PD regimen is highest soon after initiation of PD, e.g., within the first 6 months; hence, this period is used to establish a baseline.

Delivered peritoneal dialysis dose depends on many factors, including the transport properties of the peritoneal membrane, assessed by the peritoneal equilibration test (PET). There is evidence that the PET performed within the first week after initiation of PD may yield higher transport results than a PET performed a few weeks later. This difference is statistically significant, but may not be clinically relevant. It may be more convenient to perform the first PET at the end of training, rather than at the end of the first month, and the Work Group thinks this is acceptable. However, the results after a month of PD may more accurately reflect peritoneal transport properties for the subsequent period.

Table II-1. Peritoneal Dialysis Dose and Total Solute Clearance Measurement Schedule: Initial 6 Months

<table>
<thead>
<tr>
<th>Month</th>
<th>PD Fluid</th>
<th>PET</th>
<th>Urine*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$K_t/V_{urea}$</td>
<td>$C_r$</td>
<td>$K_t/V_{urea}$</td>
</tr>
<tr>
<td>1†</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>2‡</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>3‡</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>4‡</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>5‡</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>6‡</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>

NOTE. X, measurement; Y, additional measurement if "incremental" PD utilized.
* For patients who void infrequently (<3 times in 24 hours), collect urine over a 48-hour period.
† If possible, at the end of month 1, but at the end of training if that is more convenient.
‡ The measurement interval in months 2 to 6 is flexible. At least one additional measurement after the first month’s measurement is necessary. If the results of the second measurement are similar to those of the first measurement, an adequate baseline is established, obviating the third measurement. If the result of the second measurement is discrepant, a third measurement is necessary to establish a more reliable baseline.

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MEASURES OF PERITONEAL DIALYSIS DOSE

In patients initiating ESRD therapy for the first time who have some RRF, delaying the PET and the first measurement of delivered dose for a month is safe and appropriate. However, for patients initiating PD because of transfer from HD and/or for patients who do not have substantial RRF, the first measurement of delivered dose of PD should be performed earlier. In the absence of substantial RRF, waiting a month to measure delivered dose may result in inadequate dialysis for a month. Thus, the Work Group recommends that these patients undergo measurement of delivered dose of PD at 2 weeks post-initiation, assuming maintenance exchange volumes have been achieved. Patient care technicians may be able to perform these measurements.

If "incremental" PD is initiated instead of "full dose" (see Guideline 1: When to Initiate Dialysis—Kt/V urea Criterion and Appendix A: Detailed Rationale for Guideline 1), RRF must be followed carefully and frequently such that PD dose can be increased as RRF deteriorates. While urine production rate is presumed to be a clue to deteriorating RRF, that is not always the case.21 Thus, for patients initiating PD with "incremental" PD, the Work Group recommends measuring RRF every 2 months. For patients on "full dose" PD, the Work Group recommends measuring RRF with total solute removal measurements every 4 months. If urine production rate is decreasing, measure RRF every 2 months or as often as needed and considered helpful, but at least every 4 months. Once weekly Kt/V falls to less than 0.1, RRF can be considered negligible and its routine measurement can be stopped. Guideline 11: Dialysate and Urine Collections, addresses this subject again.

GUIDELINE 4

Measures of PD Dose and Total Solute Clearance (Opinion)

Both total weekly creatinine clearance normalized to 1.73 m² body surface area (BSA) and total weekly Kt/V urea should be used to measure delivered PD doses.

Rationale A valid and reproducible measure of PD dose is essential to assess the quantity of dialysis delivered to an individual patient. The quantity of dialysis is an important component of the quality of dialysis. Of the few available measures of PD dose, total weekly Kt/V urea and total creatinine clearance normalized to 1.73 m² BSA are the best, because they are most strongly associated with mortality and morbidity (see Guideline 15: Weekly Dose of CAPD). Additionally, when properly performed, these measures are reproducible enough to be useful in routine clinical practice.

The urea-based measure, Kt/V urea, measures removal of the direct product of protein catabolism. The creatinine clearance (CCr) measures removal of a product of muscle metabolism which provides insight into lean (i.e., fat-free, edema-free) body mass and possibly into compliance (see Guideline 7: PD Dose Troubleshooting). In Guideline 1: When to Initiate Dialysis—Kt/V Criterion and Guideline 15: Weekly Dose of CAPD, there is a discussion of the comparison of these two different measures (see Guideline 6: Assessing Residual Renal Function, for a definition of total weekly CCr).

These two recommended measures have both been used to measure delivered dialysis dose. Since each measure provides slightly different information, the Work Group recommends that both measures be used. Both creatinine and urea concentration can be obtained on the same sample of urine, blood, and dialysate. No additional samples need to be collected to perform both, rather than one, of these measures. Most laboratories perform both measures simultaneously (e.g., 6/60, Chem 6, etc.) on automated equipment and the cost is the same for one or both measures.

GUIDELINE 5

Frequency of Measurement of Kt/V urea, Total CCr, PNA, and Total Creatinine Appearance (Opinion)

After 6 months, total Kt/V urea, total CCr, and PNA (with all its components), should be measured every 4 months unless the prescription has been changed or there has been a significant change in clinical status (see Table II-2).

Rationale Despite the establishment in 1993 of ESRD Network/Health Care Finance Administration guidelines that measurements of delivered PD dose and total solute clearance be performed twice yearly, a Network Core Indicator review of 1,208 patient charts in 1995 revealed
Table 11-2. Peritoneal Dialysis Dose and Total Solute Clearance Measurement Schedule After 6 Months

<table>
<thead>
<tr>
<th>Month</th>
<th>Kt/V urea</th>
<th>CCR</th>
<th>Kt/V urea</th>
<th>CCR</th>
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<tbody>
<tr>
<td>7</td>
<td></td>
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NOTE. X, measurement.

* If incremental PD is still being utilized at this point, the frequency of RRF testing applies as described in Table 1 of Guideline 3: Frequency of Delivered PD Dose and Total Solute Clearance Measurement Within Six Months of Initiation. For patients who void infrequently (<3 times in 24 hours), collect urine over a 48-hour period. Urine testing can cease when the residual renal function component is a weekly Kt/V urea < 0.1.

† For children who have greater difficulty with accurate urine collection than adults, this may be deferred until full urine and dialysate collections occur every 4 months (see Guideline 11, Dialysate and Urine Collections).

GUIDELINES FOR PERITONEAL DIALYSIS ADEQUACY

cause it strikes a balance: every 4 months is often enough to be clinically helpful, but not so often as to be intrusive into a patient's lifestyle or to create a burden for the dialysis facility. Awareness of loss of RRF must be paramount. If "incremental" PD is initiated instead of "full dose" (see Guideline 1: When to Initiate Dialysis—Kt/V Criterion and Appendix A: Detailed Rationale for Guideline 1), RRF must be followed carefully and frequently such that PD dose can be increased as RRF deteriorates. While urine production rate is presumed to be a clue to deteriorating RRF, that is not always the case. Thus for patients initiating PD with "incremental" PD, the Work Group recommends measuring RRF every 2 months (see Table II-2). For patients on "full dose" PD, the Work Group recommends measuring RRF with total solute removal measurements every 4 months. If urine production rate is decreasing, measure RRF every 2 months, or as often as needed and considered helpful, but at least every 4 months. Once weekly Kt/V falls to less than 0.1, RRF can be considered negligible and its routine measurement can be stopped. Guideline 11: Dialysate and Urine Collections, addresses this subject again.

The impact of a change in prescription should be assessed within 2 to 4 weeks in order to determine if the recommended change has actually been executed and if it has accomplished its goal. The promptness of the assessment is important because clinical events could occur in the interval which could postpone the measurement or confound the results (see Guideline 10: Timing of Measurement).

Some clinical events may impair the quality of delivered PD. A change in clinical condition which warrants measurement of delivered PD dose is defined as any serious problem which affects nutritional status, the ability of the patient to perform PD mechanically or technically (such as stroke or arthritis, loss of surface area from surgery, decreased exchange volumes due to hernias, etc.), or permanently affects the transport properties of the peritoneum (e.g., protracted peritonitis). Many conditions that lead to hospitalization fall into one of these categories. Any suggestion of exacerbation of uremia should prompt a measurement of delivered dose of PD.
The Work Group recognizes that technical problems in urine collections in some children may justifiably decrease the frequency of urine collections in selected cases.

**GUIDELINE 6**

**Assessing Residual Renal Function**

_Residual renal function (RRF)_ , which can provide a significant component of total solute and water removal, should be assessed by measuring the renal component of _Kt/V_urea (_KrtIV_ urea) and estimating the patient’s glomerular filtration rate (GFR) by calculating the mean of urea and creatinine clearance.

**Rationale** A detailed rationale is presented in Appendix C. The following is a summary.

During the first few years of dialysis therapy, residual renal function (RRF) contributes significantly to total solute and water removal. Preservation of RRF may be particularly important to the effectiveness of long-term PD. For solute removal targets to be met (see Guideline 15: Weekly Dose of CAPD and Guideline 16: Weekly Dose of NIPD and CCPD) in many patients without a possibly unacceptable dialytic burden, there must be a substantial contribution from RRF.

As GFR declines over time, the contribution of secreted creatinine to total creatinine clearance (CCr) rises disproportionately and CCr becomes an inaccurate marker of GFR. Since the peritoneal membrane does not secrete solute, the GFR measure that corrects for creatinine secretion is the preferred measure to add to peritoneal clearance. In the case of a low GFR, the measurement of GFR with endogenous solutes is best done by defining GFR as the arithmetic mean of urea and creatinine clearance. This arithmetic mean essentially corrects for secretion of creatinine. This GFR measure is added to peritoneal CCr, normalized to 1.73 m² of body surface area and is totaled for a week. This is the “total weekly creatinine clearance.”

\[
\text{GFR} = \left( \text{renal urea clearance (mL/min)} + \text{renal creatinine clearance (mL/min)} \right) / 2
\]

Total weekly CCr = GFR + Peritoneal CCr, normalized to 1.73 m² of Body Surface Area.

An alternative measure of RRF is residual renal urea clearance, normalized to total body water, _Ktu/V_urea. This measure can be directly added to the peritoneal urea clearance component, _Kpt/V_ urea, to create the total urea clearance normalized to total body water, _Ktu/V_urea (shortened to _Ktu/V_urea).

Creatinine clearance corrected for renal secretion and _Kt/V_urea are both valuable measures in the management of PD patients. Each measure offers different information. Since the dialysate and urine collections are being performed for either measure, the Work Group recommends that both measures be determined.

**GUIDELINE 7**

**PD Dose Troubleshooting**

In adult patients, a daily creatinine excretion in urine and dialysate that differs from the baseline rate (as determined during the first 6 months in Guideline 3, Table II-1) by >15% should prompt an investigation for noncompliance, improper collection of drained dialysate and/or urine, or altered peritoneal transport function. Compliance should not be assessed by comparing measured to predicted creatinine excretion.

**Rationale** Twenty percent of PD patients report some non-compliance with their dialysis prescription. Preliminary data suggest that total daily creatinine excretion or appearance can be used as an indicator of compliance in CAPD. The premise for such use is that, in noncompliant patients who perform the proper number of exchanges only during the day of the clearance measurement, the amount of creatinine excreted in 24 hours (equal to the daily amount of creatinine in the spent dialysate and urine plus an estimated amount of creatinine lost through other routes, primarily the gastrointestinal tract) will exceed the amount of creatinine produced daily. Essentially, noncompliance creates an unsteady state of recently accumulated creatinine. Thus, an increase in the daily excretion of creatinine in dialysate plus urine may indicate noncompliance just prior to the collection. Other potential causes of variation in the measured amount of creatinine excreted include changes in muscle mass (see Guideline...
13: Determining Fat-Free, Edema-Free Body Mass), improper collection of dialysate or urine due to timing errors, and inaccurate urine or dialysate creatinine measurement by the laboratory. Finally, another potential cause of change in total creatinine excretion may be peritoneal membrane transport dysfunction. Quantitatively, a very large transport defect must occur to result in 15% variation in the daily creatinine excretion. However, if this is suspected, a peritoneal equilibration test (PET) or its alternative should be performed.

A similar approach should be considered in children, although only a small number of pediatric patients have been studied in this manner. Furthermore, in growing children with increasing muscle mass, there will be an increase in total creatinine excretion over time.

The Work Group's decision to use a variance of >15% in creatinine appearance over the established baseline was based on convincing but indirect evidence in adult patients. There are no data to support a similar approach in children. In PD patients who appear to be stable, creatinine appearance may vary by up to 15%, depending on a variety of factors. This variance of >15% is simply a suggestion or warning to the clinician that the creatinine excretion data are not consistent with prior evaluations and suggests a need for further investigation. The intensity of the investigation that the variance triggers is a clinical decision that requires taking many issues into consideration.

The Work Group does not recommend assessing compliance by comparing measured creatinine excretion to predicted creatinine excretion. Our reasoning is as follows: The estimated amount of creatinine lost in the gut is equal to 0.036 × serum creatinine concentration in mg/dL × body weight. The predicted creatinine production, in mg/day, is calculated by the Cockroft-Gault formulae as follows:

For men: $(28 - 0.20 \times \text{Age}) \times \text{Weight}$

For women: $(24 - 0.17 \times \text{Age}) \times \text{Weight}$

where age is in years and weight is in kilograms. These formulae were derived in a nondialysis population. The discrepancy between measured and predicted creatinine generation is expressed as the ratio of measured/predicted creatinine generation.

The use of a cut-off value of measured/predicted creatinine generation to identify noncompliance is not warranted because the measured/predicted creatinine generation in compliant CAPD patients appears to vary widely. In addition, the increase in the amount of creatinine excreted during the clearance day in noncompliant patients is very small in most cases. Measured/predicted creatinine generation is better used sequentially in a patient after establishing baseline values during a period of close observation (see Table II-1). At least for short periods of time (days, weeks), the steady-state excretion of creatinine is constant in CAPD patients, although a variation of 15% may be essentially physiologic. However, since absolute creatinine excretion is being measured and compared to a previously established reliable baseline measurement, the Work Group considers the use of predicted creatinine production to be unnecessary.

In summary, the Work Group recommends establishing a baseline creatinine excretion on the basis of 2 to 3 measurements in the first 6 months of PD (see Guideline 3: Frequency of Delivered PD Dose and Total Solute Clearance Measurement Within Six Months of Initiation). The Work Group feels that following these measures longitudinally will be more helpful than comparing measured to predicted creatinine appearance. Causes for any subsequent deviation from the baseline total creatinine excretion over time should be sought, recognizing that noncompliance is only one of several possible explanations.

**RECOMMENDATIONS FOR RESEARCH**

Since preservation of RRF is important for solute removal and contributes to total renal replacement therapy, there is a need to identify contributors to loss of RRF. For example, do antibiotics play a role? Hypotension? Other factors? Does aggressive solute removal and/or highly efficient dialysis remove stimulatory factors favoring remnant kidney hyperfiltration?

Why does GFR vary so much on a day-to-day basis and how does one account for this in
adequacy studies? Is this a collection artifact or true physiologic variation? What factors alter the daily production and excretion of creatinine? Is it a function of creatinine production or simply excretion? If the latter, is it variation in renal secretion, filtration, or both? Urine output also varies dramatically on a day-to-day basis. Is this simply a volume phenomenon or is it a reflection of true clearance changes? Why does creatinine appearance vary even in compliant patients? In children who are growing, how often should total creatinine excretion be measured and is it useful as an assessment of compliance with dialysis prescription?

The recommendation that >10% variance in creatinine appearance be considered as indicative of a status change or noncompliance should be validated.

An accurate, reproducible and easy-to-perform method of measuring RRF should be developed.
III. Measurement of Peritoneal Dialysis Dose

GUIDELINE 8
Reproducibility of Measurement (Opinion)

Accurate measurement of total Kt/Vurea and total creatinine clearance (CrCl) requires collection and analysis of urine, dialysate, and serum in a way that yields reproducible and valid results. Dialysate creatinine concentration must be corrected for the presence of glucose in some assays. Peritonitis precludes reliable measurement of delivered PD dose for up to a month. Compliance with complete collections is mandatory. For patients who void ≥3 times per day, a 24-hour urine collection is sufficient. For patients who void less frequently, a 48-hour collection is recommended. For CAPD patients, the serum sample can be obtained at any convenient time. For NIPD patients, the serum sample should be obtained at the midpoint of the daytime empty period. For CCPD patients, the serum sample should be obtained at the midpoint of the daytime dwell(s).

Rationale

A detailed rationale is presented in Appendix D. The following is a summary. Measurement of PD dose must be performed in a valid and reproducible fashion. The measure of creatinine concentration in effluent dialysate must be corrected for the presence of glucose with some creatinine assays. Each facility must determine whether this is necessary by specifically inquiring of its laboratory whether the creatinine assay used by that lab is altered by high glucose concentrations. PD dose measures should not be made until a month after peritonitis resolves, because peritonitis causes residual effects on membrane transport. Either total dialysate collections or aliquots (i.e., samples of dialysate) can be used with proper patient training and compliance. For CAPD patients, the timing of the blood sample is not important. For patients on NIPD or CCPD, the blood sample must reflect the overall average for the entire 24 hours. For NIPD patients, the serum sample should be obtained at the midpoint of the daytime empty period. For CCPD patients, the serum sample should be obtained at the midpoint of the daytime dwell. For most NIPD and CCPD patients, these time points occur in the early afternoon.

GUIDELINE 9
Estimating Total Body Water and Body Surface Area (Opinion)

V (total body water) should be estimated by either the Watson or Hume method in adults using actual body weight, and by the Mellits-Cheek method in children using actual body weight.

Watson method:
For Men: \[ V = 2.447 + 0.3362 \times \text{Wt (kg)} + 0.1074 \times \text{Ht (cm)} - 0.09516 \times \text{Age (yrs)} \]
For Women: \[ V = -2.097 + 0.2466 \times \text{Wt} + 0.1069 \times \text{Ht} \]

Hume method:
For Men: \[ V = -14.012934 + 0.296785 \times \text{Wt} + 0.192786 \times \text{Ht} \]
For Women: \[ V = -35.270121 + 0.183809 \times \text{Wt} + 0.344547 \times \text{Ht} \]

Mellits-Cheek method for children:
For Boys: \[ V = -1.927 + 0.465 \times \text{Wt (kg)} + 0.045 \times \text{Ht (cm)}, \text{when Ht ≤ 132.7 cm} \]
\[ V = -21.993 + 0.406 \times \text{Wt} + 0.209 \times \text{Ht}, \text{when height is ≥ 132.7 cm} \]
For Girls: \[ V = 0.076 + 0.507 \times \text{Wt} + 0.013 \times \text{Ht}, \text{when height is ≤ 110.8 cm} \]
\[ V = -10.313 + 0.252 \times \text{Wt} + 0.154 \times \text{Ht}, \text{when height is ≥ 110.8 cm} \]

Body surface area, BSA, should be estimated by either the DuBois and DuBois method, the Gehan and George method, or the Haycock method using actual body weight. For all formulae, Wt is in kg and Ht is in cm:

DuBois and DuBois method:
\[ \text{BSA} (\text{m}^2) = 71.84 \times \text{Wt}^{0.425} \times \text{Ht}^{0.725} \]

Gehan and George method:
\[ \text{BSA} (\text{m}^2) = 0.0235 \times \text{Wt}^{0.51456} \times \text{Ht}^{0.43246} \]

Haycock method:
\[ \text{BSA} (\text{m}^2) = 0.024265 \times \text{Wt}^{0.5378} \times \text{Ht}^{0.3964} \]

Rationale

A detailed rationale is presented in Appendix E. The following is a summary. The Watson and Hume formulae were derived by comparing total body water measurements to simple anthropometric measurements (weight, height, age) in subjects without edema, volume deficit, or end-stage renal disease. In peritoneal dialysis patients, the Wat-
son and Hume formulae provide reasonable approximations of isotopic body water measurements. Volume abnormalities (edema) are apparently the major cause of discrepancy. The Mellits-Cheek formulae were derived from subjects aged 1 month to 34 years for males and 1 month to 31 years for females. In each case, the measurement of total body water was performed in normal subjects by the use of deuterium oxide distribution, with simultaneous measurement of weight and height.

The Work Group recommends the use of the Watson or Hume formulae in adults and the Mellits-Cheek formula in children as methods for estimating V. Attention should be paid to the presence of edema at the time of the clearance study. A special case is the underweight patient (see Table X-I, Appendix E for a definition). Successful efforts to restore weight to a normal level in such a patient will result in a rising V, and consequently in a proportionally declining $K_{pt}/V_{urea}$. This does not alter the methodology of estimating total body water using actual weight. It does affect target doses of dialysis, however. This issue is discussed again in Guideline 15: Weekly Dose of CAPD.

Like the formulae above for total body water, the formulae for BSA were determined in a normal population. Many of the disclaimers described for calculating V are less of an issue in calculating BSA, because the relationship to the defining simple anthropometric measurements is less influenced by clinical conditions accompanying ESRD. Historically, many nephrologists have utilized the method of DuBois and DuBois, and much of our data are from its application. Only nine subjects were used to define this formula. More than 400 subjects, including many children, were used to define the formula of Gehan and George, and in an independent comparison, the Gehan and George method was preferred. The Haycock formula is based on measurements of 81 subjects ranging from premature infants to adults.

Amputation alters the relationship between body height and weight. This causes a mathematical distortion of the calculation of both anthropometric V and BSA, because the calculation of each takes this relationship into account. Modification of V and BSA in patients with amputations are described in detail in Appendix E.

**GUIDELINE 10**

**Timing of Measurement (Opinion)**

Routine measurements of total $Kt/V_{urea}$ and total creatinine clearance should be performed when the patient is clinically stable (eg, stable weight, stable BUN and creatinine concentrations), and at least 4 weeks after resolution of peritonitis.

Following a change in prescription or a major change in clinical status (eg, hospitalization, weight loss), but in the absence of recent peritonitis, measurements of delivered weekly $Kt/V_{urea}$ and total weekly $C_{Cr}$, should be performed within the next 4 weeks and then at 4-month intervals.

**Rationale** The effect of body weight on the calculation of V is discussed in the rationale for Guideline 9: Estimating Total Body Water and Body Surface Area and in Appendix E: Detailed Rationale for Guideline 9. Variations in serum urea and creatinine concentration can potentially increase the error in the clearance calculations, and indicate that the patient is not in a steady state.

Peritonitis may, in some instances, affect peritoneal solute transport for long periods. The rationale for waiting 4 weeks after resolution of peritonitis to repeat the clearance studies is presented in Appendix D: Detailed Rationale for Guideline 8.

The Work Group recommends frequent assessment of delivered dose of PD and total solute clearance, specifically every 4 months after the first 6 months of PD (see Guideline 11: Dialysate and Urine Collections). Major changes in clinical status (eg, patient compliance, weight gain, weight loss, technical/mechanical complications, some causes of hospitalization) may alter PD dose requirements. For example, pneumonia may contribute to loss of residual renal function, which would be undetected unless measured. Therefore, in the absence of peritonitis, a major change in clinical status should prompt a re-evaluation of weekly $Kt/V_{urea}$ and total weekly $C_{Cr}$, and this should occur within one month following the change in clinical status. Within a month following a PD prescription change, weekly $Kt/V_{urea}$ and total weekly $C_{Cr}$ should be measured to demonstrate that the goals of the prescription change have been achieved. If the patient is not stable, all attention should be directed to determining the cause of the instability and to correcting it. This may or may not include measuring the delivered dose of PD. There will be circumstances in which the change in clinical status might only alter RRF (eg, exposure to
nephrotoxins), not delivered dose of PD. While one must be cautious in assuming that persistent urine flow rate implies stable RRF, a clinical clue that deterioration has occurred may be a decrease in what previously had been a stable daily urine volume. In those settings, only measurement of RRF is indicated.

GUIDELINE 11
Dialysate and Urine Collections (Opinion)

Two to three total solute removal measurements are required during the first 6 months of PD (see Guideline 3: Frequency of Delivered PD Dose and Total Solute Clearance Measurement Within Six Months of Initiation). After 6 months, if the dialysis prescription is unchanged:

1. Perform both complete dialysate and urine collections every 4 months; and
2. Perform urine collections every 2 months until the renal weekly Kt/Varea is <0.1.

Thereafter, urine collections are no longer necessary, as the RRF contribution to total Kt/Varea becomes negligible. In children, urine collections are recommended only with complete dialysate collections (see Table II-2 reproduced from Guideline 5.)

Rationale Loss of residual renal function is the major cause of decreasing clearance in PD subjects followed longitudinally. The CANUSA study demonstrated substantial loss of renal function at 6-month intervals. The Work Group concludes that measurements of urinary clearances should be performed at 2-month intervals to prevent long periods of underdialysis. For children in whom urine collections are difficult (requiring special collection apparatus, etc.), urine collections can be deferred until the next total solute removal measurement (see Guideline 3: Frequency of Delivered PD Dose and Total Solute Clearance Measurement within Six Months of Initiation and Guideline 5: Frequency of Measurement of Kt/Varea, Total Ccr, PNA, and Total Creatinine Appearance).

RECOMMENDATIONS FOR RESEARCH

The optimal timing of blood sampling for subjects on asymmetric PD (NIPD, CCPD) should be determined. The recommendations we have made are based on pharmacokinetic theory.

Comparison of the “batch” and “aliquot” methods of dialysate sampling should be studied.

Development of a clinically applicable method of assessing TBW in children undergoing PD is recommended.

Methodology to calculate renal urea and creatinine clearance and PNA from random urine samples should be developed.
IV. Assessment of Nutritional Status Specifically as it Relates to Peritoneal Dialysis

**GUIDELINE 12**

**Assessment of Nutritional Status (Opinion)**

Nutritional status of adult PD patients should be assessed on an ongoing basis in association with $Kt/V_{area}$ and $C_{cr}$ measurements using the Protein equivalent of Nitrogen Appearance (PNA) and Subjective Global Assessment (SGA). For pediatric PD patients, nutritional status should be assessed using the PNA and other standard nutritional assessments (see Guideline 14).

**Rationale**

A detailed rationale is presented in Appendix F. The following is a summary.

There is strong indirect evidence linking survival on dialysis with nutritional status both at initiation of dialysis (see Section I: Initiation of Dialysis) and during longitudinal follow-up. Better survival has been reported in PD patients with high normalized protein equivalent of nitrogen appearance (nPNA) (see Guideline 28: Measurement of Normalized PNA in PD Patients). Positive correlations between nPNA and clearance of urea or creatinine have been reported repeatedly in PD subjects.49-52 The correlation between nPNA and $Kt/V_{area}$ may indicate increased appetite and dietary protein intake as $Kt/V_{area}$ increases, but also may simply reflect the fact that nPNA and $Kt/V_{area}$ are mathematically linked.53 This mathematical linkage makes the correlation between nPNA and $Kt/V_{area}$ in cross-sectional studies of questionable clinical significance. However, nPNA tends to increase in the same subjects when $Kt/V_{area}$ and creatinine clearance ($C_{cr}$) are increased by increasing the dose of PD, especially if the increase in the dose of PD was prescribed because of inadequate clearances.54 In the latter instance, the association between $Kt/V_{area}$ and nPNA is not the result of a mathematical coupling. In addition, there is strong evidence suggesting that quality and quantity of dialysis influences nutrition.55,56 While the precise relationship between renal function (or renal replacement therapy) and nutrition is not yet adequately understood, it is the Work Group’s opinion that adequate renal replacement therapy is necessary for normal appetite and metabolism. Thus, nutritional problems may reflect inadequate dialysis which, if corrected, may lead to subsequent improved outcomes.

Although nutritional status is influenced by many nondialysis-related factors, appetite suppression, nausea, and vomiting are major clinical features of uremia and inadequate dialysis. Therefore, nutritional status is also an important measure of PD adequacy. Of the available measures of nutrition, PNA is recommended because it provides an estimate of protein intake and protein losses. The SGA is recommended because it is a valid clinical assessment of nutritional status and is strongly associated with patient survival.

**Protein equivalent of total nitrogen appearance (PNA)**

Nitrogen intake is almost entirely from protein. The final product of protein catabolism is urea. Therefore, if steady-state nitrogen balance conditions exist, one can work backward from urea excretion to determine what the protein intake was. Yet, other protein losses (urinary, peritoneal dialysate, diarrhea) also reflect the body’s turnover of protein. Studies in dialysis patients show that predictable mathematical relationships exist between urea excretion, protein catabolism, and dietary protein intake. If peritoneal protein losses are greater than 15 g/d, PNA should be calculated as protein catabolic rate plus protein losses. If dialysate protein losses are less than 15 g/d, the formula:

$$PNA(g/d) = 10.76*(0.69*UNA + 1.46)$$

can be used to calculate PNA where UNA is total urea nitrogen appearance in grams per day.57 PNA is an indirect method for estimating dietary protein intake, a key measure of nutritional status in dialysis patients. For pediatric patients, see Guideline 14: Use of the Modified Borah Equation to Assess Nutritional Status of Pediatric PD Patients.

**Subjective Global Assessment (SGA)**

The SGA is a simple assessment that uses the clinician’s experience to subjectively rate a patient’s nutritional status based on the medical history and physical exam. The SGA was modified for use in PD patients as described in Appendix F: Detailed Rationale for...
Guideline 12. This assessment is valuable because it does not focus on a single variable; rather, it forces the clinician to view the patient more broadly. SGA addresses four items (recent weight change, anorexia, subcutaneous tissue, and muscle mass) scored on a 7-point Likert scale (see Appendix B: Detailed Rationale for Guideline 2). It can be performed by physicians, nurses, or registered dietitians during routine clinic visits. Several studies have validated that the SGA accurately reflects nutritional status in dialysis patients and, in the CANUSA study, a higher SGA was associated with a lower risk of death.

The SGA is easy to perform and can be performed within minutes during a routine clinic visit. There are no data to dictate how often to perform the SGA, so the Work Group bases its opinion on the following: the SGA should be done often enough to detect changes and to intervene in a timely manner. It should be performed in association with measurement of Kt/V urea and Ccr, every 4 months after the initial 6 months (see Guideline 5: Frequency of Measurement of Kt/V urea, Total Ccr, PNA, and Total Creatinine Appearance).

The SGA has not been validated as a means of nutritional assessment in the pediatric PD population.

GUIDELINE 13

Determining Fat-Free, Edema-Free Body Mass (Opinion)

Total creatinine appearance should be used to determine fat-free, edema-free body mass.

Rationale Fat-free, edema-free body mass is probably a more accurate term for what had previously been called lean body mass. It is an important index of overall nutritional status. Total daily creatinine production, measured as the sum of creatinine excreted in dialysate and urine plus the estimated creatinine lost in the gut, can be used to calculate fat-free, edema-free body mass. Fat-free, edema-free body mass reflects somatic protein stores in the same way that serum albumin reflects visceral protein stores. In adults, fat-free, edema-free body mass in kilograms is computed by the equation:

\[
\text{Fat-free, edema-free body mass} = 0.029 \times \text{total creatinine production in mg/d} + 7.38.
\]

Norms vary by patient gender and size. A steady-state of creatinine excretion should exist for the equation result to be valid. Factors other than muscle mass can affect the fat-free, edema-free body mass calculation by creatinine kinetics. These factors include errors in the collection or measurement of creatinine in urine or dialysate, and large variations in the dietary intake of creatine plus creatinine (meat). Fat-free, edema-free body mass estimates by creatinine kinetics may be a better index of nutritional status in PD patients, because they reflect dry fat-free, edema-free body mass and changes in muscle mass better than dual-energy X-ray absorptiometry or bioimpedance. The day-to-day variability of total creatinine excretion is only 2% to 4% over a short time interval, but is up to 15% over much longer intervals.

Serum creatinine concentration is not, by itself, an index of adequacy of peritoneal dialysis because of the large variations in creatinine production between individuals. However, a change in serum creatinine concentration may indicate changes in creatinine and urea removal to a much larger extent than a change in serum urea concentration. A rising serum creatinine concentration is usually caused by a decrease in total creatinine clearance, often secondary to a loss of residual renal function, and much less frequently by an increase in muscle mass. A decreasing serum creatinine concentration is caused more often by progressive loss in muscle mass and less often by an increase in total clearance. In other words, increases in peritoneal solute transport or recovery of renal function do not occur very often.

GUIDELINE 14

Use of the Modified Borah Equation to Assess Nutritional Status of Pediatric PD Patients (Opinion)

Nutritional status of pediatric PD patients should be assessed at least every 6 months by standard clinical nutritional evaluations and by the modified Borah equation:

\[
\text{PNA (g/d)} = [6.49 \times \text{UNA}] + [0.294 \times \text{V}] + \text{protein losses (g/d)}
\]

Rationale The equation described in Guideline 12, Assessment of Nutritional Status, from the glossary is a further modification of the original Borah equation. Since this modification
has not been validated in children, the Work Group recommends using the modification above from Kopple et al.\textsuperscript{57} and Keshaviah et al.\textsuperscript{64}

Although not validated in children, this modified Borah equation contains a factor, V, that controls for patient size, and has been employed in pediatric studies. Furthermore, dialysate protein losses must be measured directly in the dialysis effluent and not estimated when using the modified Borah equation in children. The use of equations in which dialysate protein losses are estimated has been studied in very few pediatric PD patients.\textsuperscript{65}

The dialysate protein measurement is the only additional laboratory determination from standard total solute removal measurements, as described in Guideline 4: Measures of PD Dose and Total Solute Clearance. Thus, this nutritional assessment could be easily merged to accompany total solute removal measurements.

**RECOMMENDATIONS FOR RESEARCH**

The precise relationships among PNA, dialysis dose, and outcome are unclear. Although these relationships are currently being studied in hemodialysis patients, they should also be examined in PD patients. There are probably limits, for example, to the role of increasing delivered dialysis dose in order to improve appetite and nutritional parameters. Other questions of interest include: How does improved dialysis affect nutrition, and above what delivered dose does that effect dissipate, if at all? What interventions act synergistically with improved dialysis to improve nutritional parameters? What complications interfere with nutrition, in what manner, and can the interference be overridden by another type of intervention?

A standardized nutritional assessment tool for children analogous to the SGA should be developed.
V. Adequate Dose of Peritoneal Dialysis

GUIDELINE 15
Weekly Dose of CAPD (Evidence)

For CAPD, the delivered PD dose should be a total Kt/V_{urea} of at least 2.0 per week and a total creatinine clearance (C_{Cr}) of at least 60 L/wk/1.73 m².

Rationale A detailed rationale is presented in Appendix G. The following is a summary.

Theoretical constructs predict that a weekly peritoneal Kt/V_{urea} between 2.0 and 2.25 will provide adequate dialysis. These constructs assume no residual renal function, full equilibration of plasma and dialysate urea, a target serum urea nitrogen concentration between 60 and 80 mg/dL, and nPCR between 1.0 and 1.2 g/kg/d.

Clinical studies addressing the validity of these predictions can be divided into those using univariate and those using multivariate statistical analyses. The former are methodologically weaker. Four studies which used univariate analysis suggest that total (renal and peritoneal) weekly Kt/V_{urea} values greater than 1.5, 1.89, 2.0, and 2.0, respectively, are associated with better patient survival than lower values.

Three studies from France, Italy, and North America (CANUSA) have used multivariate statistical analysis. The French study found better survival among patients with an initial weekly Kt/V_{urea} > 1.7 but did not evaluate changes in Kt/V_{urea} associated with loss of residual renal function. The Italian study evaluated prevalent CAPD patients with minimal residual renal function. Improved patient survival was observed with a weekly Kt/V_{urea} > 1.96. Values higher than 1.96 were not associated with increased survival but the statistical power to detect this association was low. The CANUSA study of 680 incident continuous peritoneal dialysis patients reported a 5% decrease in patient survival in association with every 0.1 decrease in total weekly Kt/V_{urea}, for Kt/V_{urea} between 1.5 to 2.3. There was no association between Kt/V_{urea} and technique failure or hospitalization. The predicted 2-year survival associated with a constant total Kt/V_{urea} of 2.1 was 78%. These predictions assume that renal and peritoneal Kt/V_{urea} are equivalent.

Clinical experience suggests that a total weekly creatinine clearance >50 L/1.73 m² is required for adequate dialysis. Among patients with minimal residual function in the Italian study, a weekly Kt/V_{urea} of 1.96 correlated with a weekly creatinine clearance (C_{Cr}) of 58 L. The CANUSA study reported a 7% decrease in patient survival in association with a 5 L/1.73 m²/wk decrease in C_{Cr}. Unlike the situation for Kt/V_{urea}, both technique failure and hospitalization were worse with decreased weekly creatinine clearance. The predicted 2-year survival of 78% was associated with a weekly Kt/V_{urea} of 2.1 or a weekly C_{Cr} of 70 L.

There are insufficient data to address the issue of adequate compared to optimal dialysis (see Introduction.) The latter is in part defined as the dialysis dose above which the incremental clinical benefit does not justify the patient burden or financial costs. Nor are there sufficient data to evaluate the relative importance of renal and peritoneal clearances. The recommendations assume equivalence but this requires further study. The correlation between Kt/V_{urea} and C_{Cr} will vary with residual renal function (see Figure X-2, Appendix A). The higher C_{Cr} observed in the CANUSA study compared to the Italian study was due to the greater residual renal function in the former.

Based on the available evidence, the minimum delivered dialysis dose target Kt/V_{urea} should be 2.0 per week; the minimum weekly target C_{Cr} should be 60 L/1.73 m². If there is discordance in achieving these targets, the Kt/V_{urea} should be the immediate determinant of adequacy because it directly reflects protein metabolism and is less affected by extreme variations in residual renal function (see Appendix A and Figure II-2.) However, a cause for the discrepancy should be sought and the patient followed closely for signs of underdialysis.

A special case is the underweight patient, defined in Appendix E: Detailed Rationale for Guideline 9. Successful efforts to restore weight to a normal level in such a patient will result in a rising V, and consequently in a proportionally declining K_{pt}/V_{urea}. To achieve a weekly K_{pt}/V_{urea} of 2.0 at the increased weight, the weekly target K_{pt}/V_{urea} provided during
the malnourished state must be greater than 2.0. The Work Group recommends that the target K\textsubscript{V}\textsubscript{urea} should be raised in a malnourished CAPD patient to the level that would provide a weekly K\textsubscript{V}\textsubscript{urea} of 2.0 for that patient if he or she were at the desired weight. That level is calculated by multiplying the target of 2.0 for CAPD times the ratio of V\textsubscript{desired}/V\textsubscript{actual}. This is described in detail in Appendix E: Detailed Rationale for Guideline 9, and discussed in Guideline 17: PD Dose in Subpopulations. The same upward target adjustment should be made in creatinine clearance. The target creatinine clearance should be adjusted upward by multiplying the target for that therapy (CAPD or APD) by the ratio of BSA\textsubscript{desired}/BSA\textsubscript{actual}.

Clinical judgment suggests that the target doses of PD for children should meet or exceed the adult standards. However, there are currently no definitive outcome data in pediatric patients to suggest that any measure of dialysis adequacy is predictive of well-being, morbidity, or mortality. There also are no data regarding the real protein needs of children, especially young children, on dialysis. It is the opinion of the Work Group that the nutritional requirements per kilogram of body weight are higher in children than in adults. Therefore, PD doses in children, and especially small infants who have very high protein intakes, may have to be higher than PD doses in adults.

**GUIDELINE 16**

**Weekly Dose of NIPD and CCPD (Opinion)**

For NIPD, the weekly delivered PD dose should be a total Kt/V\textsubscript{urea} of at least 2.2 and a weekly total creatinine clearance of at least 66 L/1.73 m\textsuperscript{2}.

For CCPD, the weekly delivered PD dose should be a total Kt/V\textsubscript{urea} of at least 2.1 and a weekly total creatinine clearance of at least 63 L/1.73 m\textsuperscript{2}.

**Rationale** In the absence of data that relate delivered dose of automated PD (APD) to patient outcomes, targets for NIPD and CCPD are based on opinion.

Theoretically, there is an 8% difference in clearance between CAPD and NIPD. This difference is based on calculations which describe a 200% increase in the intermittent HD clearance required to achieve the same solute removal as in continuous dialysis, (Kt/V\textsubscript{urea} of 4.0 in HD and 2.0 in CAPD) holding protein intake constant.\textsuperscript{66,67} The Work Group assumed that the delivered dose of NIPD would need to be 8% higher than the CAPD dose (108% of 2.0 = 2.16, rounded up to 2.2). The Work Group assumed that the requisite delivered dose of CCPD would be intermediate between those for CAPD and NIPD. Some variations of CCPD with diurnal exchanges of less duration than the nocturnal exchange of CAPD may be considered equal to CAPD. However, in order to simplify recommendations, the target weekly total dose for CCPD is 2.1. The recommendations for creatinine clearance are percentage adjustments corresponding to the changes in Kt/V\textsubscript{urea} targets for these groups.

Clinical judgment suggests that the target doses of PD for children should meet or exceed the adult target doses. There are no definitive outcome data in pediatrics to suggest that any measure of dialysis adequacy is predictive of well-being, morbidity, or mortality.

**GUIDELINE 17**

**PD Dose in Subpopulations (Opinion)**

There is no adequate basis for recommending any change in the target doses of dialysis discussed in Guidelines 15: Weekly Dose of CAPD, and 16: Weekly Dose of NIPD and CCPD, for various patient subpopulations (eg, patients with diabetes or who are elderly), with the exception of the malnourished patient, whose target dose is increased by the ratio of the V\textsubscript{desired}/V\textsubscript{actual} for Kt/V\textsubscript{urea}. For creatinine clearance, the target dose in a malnourished patient is increased by the ratio BSA\textsubscript{desired}/BSA\textsubscript{actual}.

**Rationale** There are no data available in the literature on which to base a recommendation for different adequacy targets for patients with diabetes or for the elderly. However, it must be remembered that malnourished patients may appear to have an adequate Kt/V\textsubscript{urea} due to calculation of V from the actual or malnourished body weight. If V were calculated from an estimate of desired body weight, the target would reflect that target body weight. This is discussed in Guidelines 9: Estimating Total Body Water and Body Surface Area, and 15: Weekly Dose of CAPD, and in detail in Appendix E, Detailed Rationale for Guideline 9.
In the absence of data in these subpopulations, if no other cause of malnutrition is discovered, the target delivered dose of dialysis should be increased by multiplying the target \( Kt/V \) urea for a normally nourished patient by the ratio of the \( V_{desired} / V_{actual} \), and the target creatinine clearance should be increased by the ratio of \( BSA_{desired} / BSA_{actual} \). These modifications are described in Guideline 15: Weekly Dose of CAPD, and Appendices E and G.

**GUIDELINE 18**

**Use of Empiric and Computer Modeling of PD Dose (Evidence)**

Both empiric and computer modeling methods can be used to estimate adequate doses of PD. Specific prescriptions are described below.

**Rationale** The Work Group has elected to describe these two empiric and computer modeling approaches in detail. They are by no means mutually exclusive.

**EMPIRIC APPROACH FOR DETERMINATION OF DOSE OF PERITONEAL DIALYSIS**

**A. General Pre-ESRD Evaluation**

1. Explain all options (transplant, HD, and PD) to patients/parents/caregivers in a non-biased manner.
2. Review medical condition/comorbidities to determine if contraindications, relative or absolute, exist for any modality (see Section VIII, Suitable Patients for PD).
3. If no medical contraindications exist and the patient is a candidate for self therapy, allow patient to choose a modality.
4. Place the chronic dialysis access (PD or HD). The Vascular Access Work group recommends that vascular accesses be placed in patients on PD. The PD Adequacy Work Group feels that this decision should be made on an individual patient basis, but our position does not necessarily disagree with the recommendations of the Vascular Access Work Group.
5. If dialysis is needed at the time of presentation, place the temporary HD access, or after placing the PD catheter, initiate therapy as suggested under point B.2, below.

**B. Initiation of Peritoneal Dialysis**

1. If possible, wait 10 days to 2 weeks after catheter placement to start PD.
2. If PD must be started in less than 10 days following catheter placement, do low-volume, supine dialysis.
3. Obtain baseline 24-hour urine collection for urea and creatinine clearance (see Guideline 6: Assessing Residual Renal Function.) These collections are for solute clearance calculations, assessment of creatinine generation, and PNA determinations.
4. Note patient’s weight and the presence or absence of edema.
5. At initiation of dialysis, explain to patient/patients/caregivers that the patient’s prescription will be individualized. Specifically, state that their instilled volume almost certainly will need to increase over time. For patients who choose Automated Peritoneal Dialysis (APD), one or more daytime dwells will be needed in approximately 85% of patients. Patients should know from the start of PD that their total solute clearance will be monitored and that, if their residual renal function or peritoneal transport changes over time, their prescription may need to change as well.

**C. Initial Dialysis Prescription for Adults**

Initial dialysis can be prescribed empirically based on patient’s weight, amount of residual renal function, and lifestyle constraints. These empiric recommendations should be implemented prior to peritoneal equilibration testing.

PD may be initiated incrementally, or as full therapy, depending on RRF at the time of initiation (see Guideline 1: When to Initiate Dialysis—\( Kt/V \) urea Criterion). For example, if \( K_{t/V} \) \( V_{area} \) is 1.8 per week, only 0.2 \( K_{t/V} \) \( V_{area} \) is needed per week. Assuming complete urea equilibration (serum to dialysate) at 6 hours, a single 2-L overnight exchange would contribute 14 L per week. If \( V \) is 40 L, this contributes a \( K_{p/V} \) \( V_{area} \) of 14/40 or 0.35 per week. Any ultrafiltrate would add further to total solute removal. That, plus the \( K_{t/V} \) \( V_{area} \) of 1.8, brings the \( K_{p/V} \) \( V_{area} \), to at least 2.15, satisfying the target requirement. This approach uses basic principles of dialysis prescription development. Thus, the dose of \( K_{p/V} \) \( V_{area} \) depends on the \( K_{t/V} \) \( V_{area} \) as the Work Group has empha-
ADEQUATE DOSE OF PERITONEAL DIALYSIS

sized throughout these guidelines. Keeping in mind that the weekly $K_{t}/V_{urea}$ goal is 2.0, the following more intense empiric approach is reasonable:

1. Patients with an estimated underlying GFR $>2$ mL/min
   a. If patient’s lifestyle choice is CAPD:
      - BSA $<1.7$ m² $\rightarrow 4 \times 2.0$ L exchanges/day
      - BSA $1.7$ to $2.0$ m² $\rightarrow 4 \times 2.5$ L exchanges/day
      - BSA $>2.0$ m² $\rightarrow 4 \times 3.0$ L exchanges/day
   b. If patient’s lifestyle choice is CCPD:
      - BSA $<1.7$ m² $\rightarrow 4 \times 2.0$ L (9 hours/night) + 2.0 L/d
      - BSA $1.7$ to $2.0$ m² $\rightarrow 4 \times 2.5$ L (9 hours/night) + 2.0 L/d
      - BSA $>2.0$ m² $\rightarrow 4 \times 3.0$ L (9 hours/night) + 3.0 L/d
   c. If patient’s lifestyle choice is NIPD:
      - Specific attention to certain details will be required. Nightly intermittent peritoneal dialysis (NIPD) is not a therapy that is typically used at the initiation of dialysis. It has been reserved for high or rapid transporters. However, in patients with significant RRF (and ability to diurese), they may initially do well on nightly exchanges only (dry day) because of the supplemental clearance provided by the patient’s RRF. See further comments on NIPD under point 2.c, below.

2. Patients with an estimated underlying GFR $\approx 2$ mL/min
   a. If patient’s lifestyle choice is CAPD:
      - BSA $<1.7$ m² $\rightarrow 4 \times 2.5$ L/d
      - BSA $1.7$ to $2.0$ m² $\rightarrow 4 \times 3.0$ L/d
      - BSA $>2.0$ m² $\rightarrow 4 \times 3.0$ L/d (consider use of a simplified nocturnal exchange device to achieve optimal dwell times and to augment clearance).
   b. If patient’s lifestyle choice is CCPD:
      - BSA $<1.7$ m² $\rightarrow 4 \times 2.5$ L (9 hours/night) + 2.0 L/d
      - BSA $1.7$ to $2.0$ m² $\rightarrow 4 \times 3.0$ L (9 hours/night) + 2.5 L/d
      - BSA $>2.0$ m² $\rightarrow 4 \times 3.0$ L (10 hours/night) + 2 × 3.0 L/d (consider combined HD/PD or transfer to HD if clinical situation suggests need).
   c. If patient’s lifestyle choice is NIPD:
      - Many of the issues discussed above for patients with an estimated underlying GFR $>2$ mL/min still apply to urine volume. Namely, if RRF provides enough diuresis, NIPD may provide enough solute removal for a while. This should be tested early on. If during training, it is noted that a patient has very low drain volumes with no apparent mechanical problem or leak, a PET should be done to determine if the patient is a rapid transporter. If so, NIPD can be prescribed using kinetic modeling.

D. Initial Dialysis Prescription for Children

In view of the close, age-independent relationship between peritoneal surface area and body surface area (BSA), the use of BSA as a normalization factor for the prescribed exchange volume in children is preferred. An instilled volume of at least 1100 mL/m² is recommended for most pediatric patients, although individual tolerance must be considered. It should be emphasized that the preceding prescriptive guidelines are general empiric guidelines for patients initiating PD, generally as first renal replacement therapy. For patients transferring from HD with minimal RRF, prompt adequacy testing is required. The above empiric recommendations must be individualized and guided by documentation that the delivered dose equals the prescribed dose. Furthermore, the instilled volumes are ones that theoretically will result in a weekly target $K_{t}/V_{urea}$ of greater than 1.9 for the average patient. High or low transporters may be below target if RRF is low. Finally, although most patients tolerate instilled volumes of greater than 2.0 liters, this needs to be evaluated for each patient.

E. Observations Needed During Training

1. Determine 4-hour drain volumes during training. This is to note if drain volumes are as expected for typical 4-hour dwells with 1.5%, 2.5%, or 4.5% dextrose exchanges. This is not a formal peritoneal equilibration test (see below), but is done to determine if the patient’s peritoneal membrane transport characteristics are markedly different from the mean.
2. Monitor for evidence of leakage in the vicinity of the catheter.
3. Complete laboratory studies.
a. Delay baseline peritoneal equilibration test (PET) until after training (see F below).

b. Perform serum chemistries and complete blood count.

c. If a computer-assisted kinetic modeling system is available, enter preliminary data to predict if the current prescription will be adequate.

F. Early Follow-Up

1. Perform 24-hour dialysate and urine collection for \( K_t/V_{\text{urea}} \), creatinine clearance, PNA calculation, creatinine generation, and \( D/P_{\text{creatinine}} \) and \( D/P_{\text{urea}} \) values. These should be done 2 to 4 weeks following initiation (see Table II-1 and Guideline 3: Frequency of Delivered PD Dose and Total Solute Clearance Measurement Within Six Months of Initiation).

2. Perform peritoneal equilibration testing (PET) approximately 1 month following initiation of PD, an appropriate time physiologically. This baseline PET could be performed at the end of a prolonged (>1 week) training period (see Guideline 3: Frequency of Delivered PD Dose and Total Solute Clearance Measurement Within Six Months of Initiation). This PET (1 month) is used as the baseline measure of peritoneal membrane transport characteristics, not to determine total solute clearance. This PET is done to rule out unsuspected problems or deviation from mean transport characteristics. Low transporters will probably require high-dose CAPD or CCPD. High transporters will eventually have ultrafiltration problems (when RRF diuresis fails) and will need short-dwell therapy such as NIPD. Average transporters will have the most flexibility (i.e., all options will be feasible).

3. Perform serum chemistries and complete blood count.

4. If a computer-assisted modeling program is available, enter baseline data. Actual data from 24-hour collection can be compared.

5. If clearances are at or above target, continue routine monitoring on a regular basis. Look for changes in 24-hour urine studies and PET data. Kinetic modeling can be used to guide future therapy.

6. If clearance is below target at 1 month, a change in prescription may be needed. Compliance issues and collection procedures should be evaluated for abnormalities.

G. Adjusting Dialysis Prescription

If kinetic modeling is not available, unless PET has changed, dialysis dose is most effectively increased by increasing the instilled volume, therefore maximizing mass transfer and dwell time. Another option would be to increase the number of exchanges/day while maintaining maximum dwell time, i.e., by using a single nighttime exchange to increase to 5 equal dwells per day. To this end, simplified mechanical exchange systems have been developed to perform a nocturnal exchange.

If kinetic modeling is available, use these programs to tailor a new prescription to meet adequacy target goals and patient lifestyle issues. This is discussed in the next section.

COMPUTER-ASSISTED KINETIC MODELING APPROACH TO ACHIEVING TARGET DOSES OF PERITONEAL DIALYSIS

As mentioned above, the availability of computer-assisted kinetic modeling to tailor PD prescriptions to transport type, body size, lifestyle, etc. may have distinct advantages. Much of what is described in the preceding discussion of the empirical approach has a mathematical basis. Computer-assisted kinetic modeling is a logical extension of the empirical approach in that it uses computer calculations to speed and assist the physician in PD prescription development. PD solution manufacturers, such as Baxter and Fresenius, provide urea kinetic modeling (UKM) models without charge.

The major advantage of this approach is the flexibility and speed of the calculations of solute clearance. Recent studies have shown that certain models very accurately predict dialysis delivery.\(^{69-71}\) Kinetic modeling is especially important for APD therapies because the dwell times are so variable and may not approach the optimal for many patients. Since the models accurately and reliably predict the delivered dose, the patients and caregivers can discuss options in a timely manner. The trial and error empirical approach discussed above moves at a much slower rate. Even with computer-assisted UKM, actual measurements are still necessary to confirm adequate dose delivery. But with computer-assisted UKM, the process is accelerated. The only theoretical disadvantage of a computer kinetic model-
Adequate dose of peritoneal dialysis

ing approach is that there might be a tendency for the caregivers not to learn the principles behind the modeling program which form the basis for the prescription strategies.

The use of computer-based modeling to achieve target PD doses has been applied successfully to a small number of children.\(^7\)\(^2\)

The dose of PD is defined as the sum of the total daily or weekly renal plus peritoneal urea clearance normalized to the total body water, the \(K_{\text{ur}}/V_{\text{area}}\), which is a dimensionless parameter expressed as either the total daily or weekly fractional clearance of body water for urea.\(^7\)\(^3\)\(^,\)\(^4\) It is substantially more difficult to compute the appropriate prescribed dialysis dose in PD than in HD. In HD, \(K_t/V_{\text{area}}\) is delivered in a single dialysis session and can be precisely calculated from the dialyzer mass transfer coefficient (MTC), constant blood and dialysate flows, ultrafiltration rate, and treatment time.\(^6\)\(^6\)\(^,\)\(^7\)\(^5\) The delivered \(K_t/V_{\text{area}}\) and PCR (or PNA) can both be calculated from the pre- and post-dialysis BUNs from a single dialysis.\(^6\)\(^6\)\(^,\)\(^7\)\(^5\)

In PD, however, the total daily peritoneal clearance, \(K_t\), is comprised of the sum of the clearances provided by several discrete exchanges. This therapy consists of several batch exchanges during which clearance and ultrafiltration are not constant (unlike the situation in HD) but fall exponentially to zero over the course of each exchange. The peritoneal mass transfer coefficient (MTC), which controls the rate of solute transport between blood and dialysate (and hence clearance), is an individual patient characteristic,\(^6\)\(^7\)\(^,\)\(^6\) which must be determined and which can clearly vary as a function of exchange volume and body position.\(^7\)\(^7\)\(^,\)\(^7\)\(^8\) In HD, ultrafiltration contributes minimally to urea clearance, while in PD it contributes up to 25% or more of total clearance and must be included in calculation of the dose. The net ultrafiltration can be precisely controlled in HD, while in PD it is a complex function of glucose absorption, membrane water permeability and lymphatic flow. The PD prescription variables include MTC, which is dependent on patient, body position, and exchange volume; distribution of exchanges between ambulatory and supine cycler exchanges; exchange volume(s); exchange times; and the osmotic gradient or percent dextrose in each exchange. Additionally, residual renal urea clearance must be measured and included in the prescription and body water or \(V\) estimated from age, gender, and surface area.\(^3\)\(^7\)\(^,\)\(^7\)\(^9\)

In current clinical practice, the peritoneal dialysis prescription is usually based on transport categorization using the peritoneal equilibration test (PET) and subsequently more finely tuned through empirical prescription changes guided by clinical experience, as described in the preceding section of this guideline.

With computerized UKM many possible PD regimens with variable exchange schedules and volumes can be tailored to be compatible with individual patient lifestyle preferences and to minimize total dialysate volume relative to required \(K_{\text{ur}}/V_{\text{area}}\). A PD prescription can be quickly and rigorously evaluated mathematically using programs written for the personal computer.\(^6\)\(^9\)\(^-\)\(^7\)\(^1\)\(^,\)\(^8\)\(^0\) These programs all require baseline transport characterization using either the PET or peritoneal function test data.

The most common method to monitor delivered \(K_{\text{ur}}/V_{\text{area}}\) is measurement of total daily peritoneal and renal urea clearance by analysis of blood, total drained dialysate, and 24-hour urine for urea nitrogen. Although highly reliable for determination of \(K_{\text{ur}}/V_{\text{area}}\) and PCR/PNA, batch analyses do not provide data to distinguish between noncompliance and/or possible changes in MTC when the delivered \(K_{\text{ur}}/V_{\text{area}}\) deviates from that expected with the current prescription. There is further discussion of this subject in Guideline 7: PD Dose Troubleshooting and Guideline 8: Reproducibility of Measurement.

An alternative technique is measurement of BUN, urine volume, and urea nitrogen in aliquots of each exchange \(^8\)\(^1\) combined with the patient’s report of the exchange time and volume for each exchange. A further discussion of aliquot methodology is in Guideline 8: Reproducibility of Measurement. Although this technique, based on the peritoneal function test approach, requires substantially more dialysate urea nitrogen measurements in addition to measurement of \(K_{\text{ur}}/V_{\text{area}}\) and PCR (or PNA), it permits calculation of a MTC for each exchange. If there are important deviations from the reported exchange schedule, the deviant exchanges will be identified by markedly deviant MTCs.

There are advantages and disadvantages of both the PET and PFT measurements. The wide range of exchange times, including nearly complete equilibration in long exchanges and the assumption of constant BUN, will result in some
variability in the MTCs measured with the PFT which do not reflect true differences. On the other hand, the MTC calculated with a single 2 L exchange under carefully controlled conditions will not always accurately reflect the MTC under clinical exchange conditions with variable exchange volumes and body position.

A simple kinetic technique for routine monitoring of delivered $K_{pt}/V_{area}$ and PCR/PNA in established patients for whom the MTCs for urea and creatinine and 24-hour dialysate creatinine have been previously established, is measurement of BUN and serum creatinine and dialysate urea and creatinine from an aliquot of one exchange. This data combined with the number of exchanges, exchange times, and exchange volumes reported by the patient permits calculation of $K_{pt}/V_{area}$, PCR/PNA and expected total dialysate creatinine content. The validity of the data can be assessed by comparison of the calculated dialysate creatinine content to the measured historical value for the patient (see Guideline 7: PD Dose Troubleshooting). In this way, both $K_{pt}/V_{area}$ and PCR/PNA can be estimated from measurements of urea nitrogen and creatinine in a blood sample and a small aliquot of one exchange using computerized UKM. When there is deviation of more than 10% in the expected creatinine excretion, more complete dialysate collections would be indicated for analysis of therapy.

**RECOMMENDATIONS FOR RESEARCH**

The amount of dialysis required for malnourished patients is not known. While there probably is consensus that such patients need extra dialysis, the requisite increase is unclear and should be studied. Other malnutrition-related questions of interest include: Can aggressive dialysis delivery reverse malnutrition? What $V$ is to be used in malnourished patients?

Can increasing dialysis dose improve outcomes in a linear manner, or is there a dose above which no benefit is noted, or complications or costs outweigh the benefits?

A multi-centered study of pediatric patients to evaluate clinical outcome as a function of delivered PD dose should be initiated. Urea kinetic modeling computer programs specifically designed for children should be developed and validated in a prospective trial.

Although there is a database documenting the validity of UKM to describe transport in PD, UKM has not been widely used to prescribe and control the delivered dose. Since the risk of mortality may be highly non-linear with increased dose of delivered dialysis, it is reasonable to assume that the coefficient of variation on mean $K_{pt}/V_{area}$ should not exceed 10% to 15% for individual patients. A multi-center clinical trial to study clinical outcome as a function of $K_{pt}/V_{area}$, using UKM to control $K_{pt}/V_{area}$ prospectively in individual randomized patients is recommended.

Do patients with diabetes need higher targets for delivered dose of PD? Should PD delivered dose be increased in hospitalized patients during acute illness or stress?
VI. Strategies for Increasing the Likelihood of Achieving the Prescribed Dose of Peritoneal Dialysis

GUIDELINE 19
Identify and Correct Patient-Related Failure to Achieve Prescribed PD Dose (Opinion)

Potential patient-related causes of failure to achieve prescribed peritoneal dialysis dose should be investigated and corrected. These include:

- Failure to comply with the prescription
- Lack of understanding of the importance of adherence to the full prescription
- Sampling and collection errors.

**Rationale** A detailed rationale is presented in Appendix H. The following is a summary.

Preliminary data from the USRDS DMMS Wave II project show that 487 CAPD patients self-report full compliance with 82.8% of their exchanges. One exchange per week is missed by 11.5% of patients and 2 to 3 exchanges per week are missed by 4.5% of patients, all self-reported. The American Association of Kidney Patients completed a patient self-reported survey about the impact of the NKF-DOQI guidelines and came up with a very similar number for frequency of missed exchanges.

Conditions causing noncompliance in PD patients have not been adequately analyzed. From studies on compliance with chronic drug regimens, it is known that patients are more compliant when they are convinced about the appropriateness and beneficial effects of the prescribed treatment and that frequent reinforcement of the importance of the treatment is associated with better compliance. Therefore, it is the opinion of the Work Group that, in addition to giving careful consideration to the selection of medically appropriate candidates for PD, as detailed in Section VIII: Suitable Patients for Peritoneal Dialysis, special emphasis should be placed on education of PD patients about the importance and technique of the PD prescription. Instructions should be repeated at least every 6 months, and patients should be monitored for signs of change in compliance. Monitoring the output of creatinine in the dialysate plus urine, as detailed in Guideline 7: PD Dose Troubleshooting, is recommended by the Work Group as a method for measuring compliance.

GUIDELINE 20
Identify and Correct Staff-Related Failure to Achieve Prescribed PD Dose (Opinion)

Potential staff-related causes of failure to achieve prescribed peritoneal dialysis dose should be investigated and corrected. These include:

- Errors in prescription
- Inadequate monitoring of delivered dose
- Inadequate patient education.

**Rationale** To increase the likelihood of achieving a prescribed dose of PD, it is necessary to elucidate the staff-related causes of failure to achieve a prescribed dose of peritoneal dialysis. The Work Group found no reports addressing this issue in PD; the following discussion represents the opinion of the Work Group members.

Inadequate understanding of the physiology and kinetic principles of PD by the physicians and nursing staff may result in:

- Errors in patient selection
- Errors in the prescription of the PD dose
- Errors in monitoring whether the prescribed dose is delivered
- Errors in PD dose modification to achieve the prescribed goal
- Inability to test for and recognize patient noncompliance
- Inadequate patient education.

The impact of patient education on patient compliance with the PD prescription was discussed in Guideline 19: Identify and Correct Patient-Related Failure to Achieve Prescribed PD Dose. The chance of inappropriate prescription of the PD dose is enhanced when the prescribing physician has a sketchy knowledge of the principles of clearance studies in PD. It has recently been recognized that nephrology fellowship curricula lack emphasis on training in dialysis. To prescribe and deliver the proper dose of PD, nephrologists must ensure adequate education and training in PD.

The use of computer modeling in PD may help...
achieve the prescribed dose by suggesting various options to alter the PD dose. This approach may assist in avoiding unrealistic PD dose schedules for certain patients (see Guideline 18: Use of Empiric and Computer Modeling of PD Dose).

The use of total creatinine appearance/output data in detecting noncompliance is important as discussed in Section II: Measures of Peritoneal Dialysis Dose.

Inadequate education may be a key factor in the patient non-adherence to the prescribed dose of therapy resulting in the above-mentioned shortcuts. The American Association of Kidney Patients suggests that PD patients are willing to increase the frequency and/or volumes of exchanges, if necessary, and that explanations (education) and participation in decision making are good incentives. Inadequate education may stem from both poor educator understanding of the principles of clearance and lack of proper teaching technique. Staff responsible for patient education should be trained and competent in both the principles of clearance and the technique of patient instruction.

**RECOMMENDATIONS FOR RESEARCH**

Validate methods of assessing compliance.

Evaluate the association between patient understanding of PD techniques and compliance. Specifically, what is the role of inadequate patient knowledge in noncompliance?

Evaluate effect of staff's knowledge of clearance principles and teaching techniques, and repetition frequency of patient instruction in proper delivery of PD dose.

Is there a psychological profile which is predictive of noncompliance? If so, what is the best method to characterize this profile?
VII. Clinical Outcome Goals for Adequate Peritoneal Dialysis

BACKGROUND

Throughout these Guidelines, the Work Group has focused on patient outcomes. Improving patient outcomes is the primary objective of the DOQI (Dialysis Outcomes Quality Initiative). The Work Group realizes that definitions of goals regarding patient outcomes are needed. As stated in the Introduction to these guidelines, the goals are integral to the definitions of adequate, optimal, and effective dialysis.

GUIDELINE 21
Measurement of PD Patient Survival (Opinion)

Survival of PD patients should be quantitated serially as an outcome measure. 

Rationale  Patient survival is an objective outcome that is dependent upon many variables, some controllable and some uncontrollable. Sub-optimal doses of delivered dialysis will adversely affect patient survival in adults (data unavailable in children), as discussed in detail in Guideline 15: Weekly Dose of CAPD. A primary goal of ESRD therapy is to prolong life while minimizing uremic symptoms. United States Renal Data Systems (USRDS) data from hemodialysis patients have demonstrated an association between low Kt/V and increased incidence of death from coronary artery disease, other cardiac disease, cerebrovascular accidents, and other conditions. There is also evidence that underdialysis adversely affects mortality in PD patients with ischemic cardiac disease or left ventricular dysfunction. Thus patient survival is a measure of renal replacement program effectiveness. Case mix must be factored into survival analysis, however. The USRDS case mix analysis is an excellent starting point, addressing age, race, primary renal disease, and presence or absence of diabetes. The USRDS is attempting to refine case mix further by addressing other underlying comorbidities. At dialysis centers with a small number of patients, survival may need to be evaluated over many years to obtain a reliable estimate.

GUIDELINE 22
Measurement of PD Technique Survival (Opinion)

PD technique survival, both dependent and independent of peritonitis, should be quantitated serially in PD patients as an outcome measure. 

Rationale  It is common for ESRD patients to change renal replacement therapy modalities during the course of their treatment. Reasons for transfer include: complications of the therapy, inability to perform the therapy (lack of suitable access, no partner to do self care, medical contraindication), and patient request/lifestyle issues. Patient case mix, geographical location, and experience with PD in complicated cases are factors affecting transfer. For some patients, for optimal outcome, it may be medically appropriate to transfer from PD to HD; this does not imply failure of the therapy or the dialysis facility.

Peritonitis remains the primary cause of transfer from PD. It is acknowledged that at times peritonitis is the "precipitating" event for transfer, while the real underlying reason is patient burnout, non-compliance, inadequate dialysis, a request based on lifestyle, or an underlying exit site infection. Overall peritonitis rates can be influenced by the center. An association between malnutrition and frequency of peritonitis has been reported. Inadequate dialysis may lead to inadequate dietary protein intake and malnutrition as described in detail in Section IV: Assessment of Nutritional Status as it Relates to Peritoneal Dialysis. The relationship between solute clearance and frequency or severity of peritonitis has not been adequately studied. For example, it is not clear if inadequate dialysis or malnutrition directly predispose the patient to peritonitis. However, peritonitis is an important outcome and its frequency and severity is an index of the overall suitability of PD. It is, therefore, the opinion of the Work Group that peritonitis should be monitored.

Inadequate dialysis is directly responsible for at least 10% of transfer to HD. There is an association between PD technique failure and total solute clearance, and one possible reason for poor technique survival rates may be underlying inadequate dialysis. While the
CANUSA study found a relationship between creatinine clearance and PD technique survival, the investigators suspected this was more related to RRF than delivered dose of PD. In the CANUSA study, technique survival was approximately 75% at 2 years in North America. However, peritonitis was not analyzed as a variable in the multivariate analysis. No difference was found in technique survival between patients from Canada and the United States. Average $\text{Kt/V}_{\text{urea}}$ started at 2.38 and decreased to 1.99 over 2 years. The risk of technique failure increased by 5% for every 5-L/wk decrease in creatinine clearance. These findings corroborated data from Tattersall et al who found that patients with a lower $\text{Kt/V}_{\text{urea}}$ had a lower rate of technique survival. The lower rate of technique survival was related to underdialysis, not to peritonitis or hernia development. The nutritional parameter of nPCR has been shown to be predictive of CAPD technique failure in a multivariable analysis.

It is a common perception that patients transferring from any renal replacement therapy are at increased risk for death in the immediate post-transfer period. However, indirect evidence from the USRDS does not support that perception.

GUIDELINE 23

Measurement of Hospitalizations (Opinion)

ESRD-related and ESRD-unrelated hospitalizations (admissions/year, hospitalized days/year) in PD patients should be quantitated as an outcome measure.

Rationale  Hospitalization is an indicator of the overall effectiveness of treatment of chronic conditions and therefore constitutes an important outcome for dialysis patients. The number of admissions per year and total number of hospital days per year are two separate, but related, serial measures of outcome. Hospitalizations of PD patients can be related or unrelated to ESRD. An association between low creatinine clearance and increased overall hospitalization rate has been reported. Based on this evidence, the Work Group recommends that cause, frequency, and length of hospitalizations of PD patients be monitored. Categorizing hospitalizations according to whether they are related to ESRD or not offers certain advantages for analysis and, therefore, it is the opinion of the Work Group that this type of stratified analysis should be performed.

According to the USRDS, PD patients are hospitalized an average of 1.8 times per year. The CANUSA study found, by multi-factorial analysis, an association between prolonged hospitalization and low creatinine clearance. Some admissions are specific to PD, ie, not seen with other dialysis therapy, such as elective abdominal wall herniography. As larger instilled volumes are administered to maintain target doses of dialysis, there is an increased risk of leaks and hernia formation, both of which can lead to hospitalization. Admissions unrelated to ESRD are important indicators of morbidity (cardiac disease, infections, etc.) in PD patients. Since hospitalization data are important outcome parameters for all dialysis patients and can reflect solute clearance, the Work Group recommends that they should be monitored.

Although it is uncertain whether inadequate dialysis is directly related to an increased risk of peritonitis and catheter infections, inadequate dialysis is related to uremic symptoms such as nausea, vomiting, and gastrointestinal bleeding. It is acknowledged that some centers may treat all episodes of peritonitis in the hospital, while others only admit those with severe or refractory perito-
CLINICAL OUTCOME GOALS FOR ADEQUATE PERITONEAL DIALYSIS

Therefore, in addition to tracking hospitalization rates, centers should also monitor reasons for hospitalization (related vs. unrelated to ESRD, and specific reason for admission). The use of ICD-9 (International Classification of Diseases, 9th revision) or similar codes may be valuable in this process. The USRDS is attempting to categorize causes of hospitalizations as infectious, cardiovascular, dialysis access-related, and all other.

Hospitalizations from causes unrelated to ESRD may be related to inadequacy of PD. As discussed in Guideline 25: Measurement of PD Patient Survival, disease-specific (particularly cardiac) mortality is related to the dialysis dose for both HD and PD. Therefore, although studies of disease-specific hospitalizations and their relation to the dose of PD have not been reported, it is reasonable to monitor this relationship. Each outcome measure should be adjusted as well as possible for case mix. The USRDS is attempting to do so with Standardized Hospitalization Rates (SHRs).

GUIDELINE 24
Measurement of Patient-Based Assessment of Quality of Life (Opinion)

Patient-based assessment of quality of life (QOL) in PD patients should be evaluated serially as an outcome measure.

A patient-based quality of life instrument should have both generic and disease/treatment-specific measures of health-related quality of life, and should be shown to be valid, reliable, and responsive prior to use. Once such an instrument is available, it should be administered at initiation of dialysis and at intervals determined to be appropriate by its validation studies.

**Rationale** Quality of life (QOL) can be assessed with generic or disease-specific measures. Many quality of life measures have been used in dialysis patients. However, fewer measures have been used for peritoneal dialysis than for hemodialysis patients. Measures used in peritoneal dialysis patients and reported in the literature include:

- Medical Outcomes Study Short Form 36 (SF-36)
- Sickness Impact Profile (SIP)
- Index of Well Being, Index of Overall Life Satisfaction
- Index of Psychological Affect
- General Health Questionnaire
- Simmons Self Esteem Scale
- Profile of Mood States
- Multidimensional Health Locus of Control
- Modality Specific Stresses Scale
- General Treatment Stress Scale
- Global Illness Stress on Self and Others
- Global Adjustment to Illness Scale
- Quality of Life (QL 100 mm) Analogue Scale
- Dialysis Relationship Quality Scale
- Social Leisure Activities Index, Social Support Satisfaction Scale
- General Well Being Index
- Index of General Affect, Overall Life Satisfaction
- Katz Activities of Daily Living
- Time Tradeoff Measures.

Unfortunately, many of these instruments do not have published data indicating that reliability (test-retest, inter-rater), validity (content, construct, internal consistency), and responsiveness-to-change have been rigorously tested. For this reason, no particular instrument can be strongly recommended over another. Furthermore, many instruments developed for research purposes may be burdensome for patients or facilities, eg, require interviewer assistance or have complicated scoring algorithms. Nevertheless, generic and disease-specific measures hold promise as useful clinical tools.

A popular generic measure used in peritoneal dialysis patients is the Medical Outcomes Study Short Form-36 (SF-36). Promising self-administered instruments used in peritoneal dialysis patients include the CHOICE Health Experience Questionnaire and the Kidney Disease Quality of Life (KDQOL).

The Work Group recommends that each facility keep abreast of future developments regarding these instruments. As experience increases and one or more instruments are clearly established as useful in PD patients, standardized QOL measurement should be integrated into the routine care and evaluation of patients, programs, and facilities.

GUIDELINE 25
Measurement of School Attendance, Growth, and Developmental Progress in Pediatric PD Patients (Opinion)

School attendance (in the absence of other comorbidities precluding school attendance),
growth, and developmental progress should be measured serially in pediatric PD patients.

**Rationale** The ability of pediatric PD patients to attend school is an important measure of the success of PD.

Underdialysis may affect the cognitive development and statural growth of children. However, the exact relationship between dialysis dose and normal growth and development in children is not clear. Nonetheless, the Work Group believes that cognitive development and statural growth should be monitored serially in children and charted in relation to patient age.

**GUIDELINE 26**

**Measurement of Albumin Concentration in PD Patients (Opinion)**

A stable or rising serum albumin concentration that is greater than or equal to the lower limit of normal for each laboratory should be used as an outcome goal.

**Rationale** In PD patients, as with HD patients, there is strong evidence to suggest that a low serum albumin level is associated with an increased risk of technique failure and death. In the CANUSA study, a difference of 0.1 g/dL of serum albumin concentration was associated with a 5% change in the risk of technique failure, a 5% change in days hospitalized, and a 6% change in the risk of death. Therefore, the Work Group recommends monitoring serum albumin concentration in PD patients because of its association with important outcomes.

Although the significance of serum albumin as a predictor of outcomes in adults is undisputed, its relationship to overall nutrition and, to a larger extent, to the levels of urea or creatinine clearance is unclear. That albumin synthesis depends on dietary protein intake is well known. However, catabolic illness can reduce albumin synthesis, and probably increase albumin degradation, even when dietary protein intake is not low. Observations in PD patients have provided indirect support for this effect of catabolic illness. Although serum albumin concentration is an important predictor of outcome, it was not found to be significant in another study when comorbid conditions were entered as covariates in their model. In this last study, the presence of comorbid conditions was associated with low serum albumin. Several cross-sectional studies have identified a positive correlation between serum albumin concentration and solute clearance. However, urea and creatinine clearance were not identified as predictors of serum albumin by multivariate analysis. Age, comorbid factors (diabetes) and peritoneal solute transport were the major predictors of serum albumin in these multi-factorial analyses.

Normal serum albumin concentrations vary by laboratory methodology; hence local standards should be used. If the serum albumin level is below normal for the laboratory, but is increasing, this suggests that the patient is anabolic and is increasing protein stores. Conversely, a low albumin or decreasing albumin level is likely to be associated with malnutrition or decreasing protein stores. Although there are no published data specifically addressing this point, it is the Work Group's opinion that a patient whose serum albumin has decreased 0.1 g/dL/month from a baseline of 4.0 g/dL to 3.7 g/dL may be at higher risk than a patient with a stable serum albumin concentration of 3.7 g/dL.

Taken together, the data discussed above suggest to the Work Group that:

- Serum albumin concentration should be monitored on a regular basis and a stable or rising value is desirable. It should be measured at least every 4 months.
- Serum albumin levels should be evaluated in the context of the patient's overall clinical status including comorbid diseases, peritoneal transport type, delivered dose of PD, and quality-of-life issues.
- The highest albumin level possible should be the goal for each patient.

It is the Work Group's opinion that an optimal serum albumin level can be obtained by adequate nutrition monitored frequently by the renal dietitian, prevention and treatment of catabolic illness, and maintenance of $Kt/V_{\text{urea}}$ and creatinine clearance at or above the levels recommended in Section V: Adequate Dose of Peritoneal Dialysis.

In summary, low serum albumin is a strong predictor of mortality and morbidity in PD patients. Therefore, serum albumin is an important outcome measure in PD patients and should be
monitored, although an association between serum albumin and urea or creatinine clearance has not been convincingly shown. Efforts to maintain serum albumin in the normal range should include adequate nutrition, adequate clearances, and prevention and treatment of catabolic illness.

GUIDELINE 27
Measurement of Hematocrit in PD Patients
(Evidence)

Providers should strive to achieve a hematocrit level of 33% to 36% in 75% of PD patients.

Rationale See NKF-DOQI’s Clinical Practice Guidelines for the Treatment of Anemia of Chronic Renal Failure.

GUIDELINE 28
Measurement of Normalized PNA in PD Patients (Opinion)

Providers should strive to achieve a normalized PNA (nPNA) of greater than or equal to 0.9 g/kg/day in PD patients.

Rationale The role of PNA is discussed in Guideline 12: Assessment of Nutritional Status. Maintenance of positive nitrogen balance and the prevention of underlying malnutrition is important because of the documented detrimental impact of hypoalbuminemia and low SGA scores on patient survival. From nitrogen balance studies, it has been estimated that PD patients need to ingest at least 1.2 g/kg/d of protein to maintain positive nitrogen balance. This is higher than the recommended daily protein intake for healthy individuals, but not surprising due to the significant amount of protein known to be lost in dialysate. Despite these recommendations from balance studies, it has been reported that many patients are in positive nitrogen balance with protein intakes of 0.9 to 1.0 g/kg/d. Cross-sectional studies would suggest that in the absence of significant comorbid diseases, patients with PD doses in the range of Kt/V urea of 2.0 spontaneously ingest at least 0.9 g/kg/d of protein. Total solute clearance and nPCR

* Studies that used the term PCR are cited in this rationale. Since the original study authors used the term PCR, this rationale will use the term PCR when specifically describing results from studies which used that term. However, the Work Group favors the term PNA.

are strongly correlated in cross-sectional studies. However, it has been suggested that this is due in part to mathematical coupling of data. Three studies have investigated the effect of an increased Kt/V urea on nutritional status (nPNA and serum albumin concentration) in a limited number of subjects. While nPCR increased as Kt/V urea increased, an increase in serum albumin concentration did not occur. A reasonable conclusion from these data would be that in the absence of significant comorbidity, an increase in delivered dialysis dose should result in a corresponding increase in nPNA. In patients who show signs of malnutrition, their dialysis prescription should be closely evaluated with consideration to increasing their dose of dialysis if significantly below target. This is discussed in Guideline 15: Weekly Dose of CAPD.

No study of PD patients has demonstrated that nPNA is an independent predictor of outcome when a multiple regression model is used. Correlation coefficients relating DPI to nPCR are on the order of 0.6. However, PD patients who are neither anabolic nor catabolic tend to demonstrate higher correlation coefficients between nPNA and DPI. Also, there are little data to show a significant relationship between nPCR and serum albumin levels. Age, peritoneal transport type, presence of diabetes, and other comorbid diseases have a greater effect on albumin than does nPCR. This subject is discussed in further detail in Appendix B.

Despite these concerns, the Work Group recommends that PNA should be monitored. Low nPNA values in nonanabolic PD patients indicate a low DPI regardless of the values of other nutritional indices. One should strive to achieve an nPNA of at least 0.9 g/kg/d in adult PD patients. PNA values at this level or higher are likely to be associated with neutral or positive nitrogen balance in the absence of significant comorbidity or dialysate protein losses.

While the recommended dietary allowance for normal children is known, there are no definitive data regarding the real protein needs of children, especially young children on dialysis. However, clinical practice suggests that the protein needs of children on PD are greater than the recommended dietary allowance, in part related to dialysate protein losses. Current recommended protein intakes for children receiving PD, referenced for age are as follows:

111;
<table>
<thead>
<tr>
<th>Age</th>
<th>Protein Intake</th>
</tr>
</thead>
<tbody>
<tr>
<td>Birth to 3 years</td>
<td>2.5-3.0 g/kg/d</td>
</tr>
<tr>
<td>3 years to puberty</td>
<td>2-2.5 g/kg/d</td>
</tr>
<tr>
<td>Pubertal patients</td>
<td>2.0 g/kg/d</td>
</tr>
<tr>
<td>Post-pubertal</td>
<td>1.5 g/kg/d</td>
</tr>
</tbody>
</table>

Because dialysate protein losses may vary widely in children, individualized recommendations for dietary protein intake may benefit from measurement of dialysate protein losses. An equivalent amount of protein to replace dialytic losses must be added to the recommended normal daily allowance for normal children.112

RECOMMENDATIONS FOR RESEARCH

The relationship between solute clearance and the frequency and/or severity of peritonitis has not been adequately studied. Specifically, does inadequate PD dose contribute to peritonitis?

Studies to assess whether an increase in nPNA is associated with an increase in serum albumin levels, nutritional status, or improved survival would be valuable.

Studies to further define the relationship between nPCR and outcome are needed. For example, the longitudinal follow-up of nutritional status (determined by a variety of methods) will be more influential in improving understanding of nutritional outcomes. Longitudinal studies should be emphasized over cross-sectional studies.

The relationship between PD dose and outcome parameters in children needs definition. Studies of nutritional interventions are lacking and are encouraged.
VIII. Suitable Patients for Peritoneal Dialysis

GUIDELINE 29

Indications for PD (Opinion)

Indications for PD include:

- Patients who prefer PD or will not do hemodialysis (HD)
- Patients who cannot tolerate HD (e.g., some patients with congestive or ischemic heart disease, extensive vascular disease, or in whom vascular access is problematic, including the majority of young children)
- Patients who prefer home dialysis but have no assistant for HD, or whose assistant cannot be trained for home HD

Rationale There is a rapid change in solute transport as well as rapid shifting of volume within compartments during HD. Some patients with severe cardiac disease may be better managed on PD since these acute changes are avoided. PD has been proposed as a method of managing refractory heart failure even in patients without renal failure.

Advantages of PD in patients with cardiovascular disease include: better hemodynamic control, less acute hypokalemia (or electrolyte shifts) which could result in arrhythmia, and better control of anemia (important in patients with coronary artery disease). Although a comparison of PD to HD for patients with severe heart failure has not been published, there are several reports of successful PD performance in subjects with severe heart failure. Tolerance of the procedure (PD), fluid management, prevention of arrhythmias, and patient survival were satisfactory in these reports.

Extensive peripheral or central venous occlusive disease prohibits surgical placement of some types of hemodialysis access. Manifestations of severe ischemia, even gangrene, of the hands follow placement of vascular access in the same wrist or forearm in a few patients with severe peripheral vascular disease, particularly diabetics. Marginal vascular beds are at risk for ischemia or reduced perfusion during hypotension, which is frequent in some HD patients. These patients benefit from increased vascular stability, which can be achieved with PD.

Over a period of time, vascular accesses fail and revisions are no longer able to restore adequate blood flow. As a result, the patient receives inadequate hemodialysis and should be evaluated for PD.

Home hemodialysis requires an assistant. For patients who prefer dialysis at home, the lack of a hemodialysis assistant may mandate PD. In addition, patients who have transportation problems to a hemodialysis center or live a great distance from a center may prefer home PD.

The decision to initiate PD rather than HD in children is influenced by a variety of factors. Because of the difficulties in maintaining vascular access in infants and small children, PD is usually the modality of choice when weight is <20 kg. Regular school attendance by children of all ages can best be achieved with a home dialysis procedure. PD is typically preferred over HD. Finally, renal replacement therapy can also best be provided by PD when the child lives a long distance from a pediatric ESRD center.

GUIDELINE 30

Absolute Contraindications for PD (Opinion)

Absolute contraindications for PD include:

- Documented loss of peritoneal function or extensive abdominal adhesions that limit dialysate flow
- In the absence of a suitable assistant, a patient who is physically or mentally incapable of performing PD
- Uncorrectable mechanical defects that prevent effective PD or increase the risk of infection (e.g., surgically irreparable hernia, omphalocele, gastroschisis, diaphragmatic hernia, and bladder extrophy)

Rationale Documented Loss of Peritoneal Function. PD efficiency relies on effective peritoneal blood flow, dialysate flow, sufficient peritoneal surface area, and permeability to allow adequate solute and fluid removal. Any compromise in these functions may result in inadequate peritoneal dialysis and thus the failure of PD.

It should not be assumed that children who have previously undergone extensive abdominal
surgery will not achieve successful PD. A trial of PD is warranted in such children and adequate dose delivery must be documented.

*Psycho-neurological Problems.* The optimal performance of PD requires certain physical and intellectual capabilities of the patient or caregiver. With major loss of mechanical function or eye-hand coordination, PD becomes difficult to perform. Patients or caregivers are responsible for problem identification and problem solving during PD. If the patient is deemed psychologically incompetent, these tasks and decisions may not be reliably or safely executed.116,127

*Abdominal Mechanical Problems.* The dialysate in the abdomen must be accessible to the vascular bed of the peritoneal membrane. Any mechanical problem that prevents this (eg, herniap sack, subcutaneous leak) will impair the efficiency of PD. Intra-abdominal pressure increases with dialysate infusion and during the ultrafiltration process, thereby exacerbating any structural defect such as hernia. Some of these abdominal defects are not surgically correctable.116,117

**GUIDELINE 31**

**Relative Contraindications for PD (Opinion)**

Relative contraindications for PD include:
- Fresh intra-abdominal foreign bodies (e.g., 4-month wait after abdominal vascular prostheses, recent ventricular-peritoneal shunt)
- Peritoneal leaks
- Body size limitations
- Intolerance to PD volumes necessary to achieve adequate PD dose
  - Inflammatory or ischemic bowel disease
  - Abdominal wall or skin infection
  - Morbid obesity (in short individuals)
  - Severe malnutrition
  - Frequent episodes of diverticulitis.

*Rationale* Fresh Intra-abdominal Foreign Bodies. Newly implanted abdominal prostheses must be allowed sufficient time for healing to avoid leakage or possible dialysis-related peritonitis with potential spread to the prosthetic device or material. The time required for healing may vary from 6 to 16 weeks.129-131 The bacterial seeding of any vascular prosthesis during hemo-dialysis is also a risk. The best type of dialysis in this setting is unclear.

*Peritoneal Leaks.* Peritoneal leakage into subcutaneous tissues, pleural space, or genitalia can be painful and cause local problems. Leaking into the vagina or rectum increases the risk of contamination. Unsatisfactory drainage and clearance, as well as medical complications, such as respiratory compromise in the case of diaphragmatic leak, can occur as a result of such leakage.114,117,127

*Body Size Limitations.* Body size can be a relative contraindication to PD when the patient is either too small to tolerate the prescribed dialysate volume or too large to achieve adequate dialysis. For patients with little or negligible RRF, there are definite size limitations for adults on CAPD with 4 daily exchanges.152 However, even larger individuals can achieve acceptable clearances if they are treated with a combination of daily CAPD and nocturnal automated PD.133 In large individuals, increase in the exchange volume is more efficient than increase in the number of daily exchanges. However, the patient with the increased exchange volume may experience abdominal pain or discomfort, shortness of breath, or loss of appetite as a result of abdominal pressure.134

*Intolerance to PD Volumes Necessary to Achieve Adequate PD Dose.* Intolerance to a PD volume is generally not known until it is attempted. Frequent exchanges with small volumes, as observed during automated PD, may not be able to provide an adequate delivered dose of PD. Raising volumes to the limit of tolerance may be problematic in patients with advanced lung disease, or patients with recurrent hydrothorax. Infrequently, this may be applicable to some patients with polycystic kidney disease or severe lumbo-sacral disk disease.

*Inflammatory or Ischemic Bowel Disease.* Inflammatory or ischemic bowel disease or frequent episodes of diverticulitis are relative contraindications to peritoneal dialysis. It is reasonable to assume that there may be increased risk for transmural contamination by enteric organisms in these circumstances.127

*Abdominal Wall or Skin Infection.* Abdominal wall or skin infection can lead to contamination of the catheter exit site, tunnel, and peritoneal cavity through touch and cross contamination.116 The decision to use PD in patients with a colostomy or ileostomy must be individualized, since successful application of PD has been described in such patients.
**Morbid Obesity.** Morbid obesity can pose special dilemmas in peritoneal catheter placement, the healing process, and in providing adequate dialysis. The possibility that increased caloric absorption from the dialysate could lead to further weight gain should also be considered.

**Severe Malnutrition.** Wound healing is compromised in severely malnourished patients. Self-dialysis such as PD may not be suitable for many severely malnourished patients because of inability to comply with the dialysis regime. Furthermore, peritoneal protein losses may not be tolerated.

**Frequent Episodes of Diverticulitis.** Diverticulitis during peritoneal dialysis often results in peritonitis. Peritoneal dialysis in patients with frequent episodes of diverticulitis places these patients at higher risk for peritonitis.

**GUIDELINE 32**

**Indications for Switching from PD to HD (Opinion)**

The decision to transfer a PD patient to HD should be based on clinical assessment, the patient’s ability to reach HD dose target levels, and the patient’s wishes. In particular, these patients should have vascular access addressed as advised by the NKF-DOQI Vascular Access Work Group.

Indications for switching from PD to HD include:

- **Consistent failure to achieve target \( \text{Kt/V}_{\text{area}} \) and \( C_{\text{Cr}} \) when there are no medical, technical, or psycho-social contraindications to HD**
- **Inadequate solute transport or fluid removal.** High transporters may have poor ultrafiltration and/or excessive protein losses (relative contraindication, obviously discovered after initiation and the first PET)
  - Unacceptably frequent peritonitis or other PD-related complications
  - Development of technical/mechanical problems
  - Severe malnutrition resistant to aggressive management (relative)

Patients should be informed of the risks of staying on PD at a level of adequacy below that recommended by their physician.

**Rationale** The above recommended indications for switching a patient from PD to HD are based on the following considerations:

**Consistent Failure to Achieve Target \( \text{Kt/V}_{\text{area}} \) and \( C_{\text{Cr}} \).** Consistent failure to achieve the target total solute removal with proper PD prescription management should lead to evaluation of compliance issues and deterrents to appropriate performance of peritoneal dialysis exchanges. After all avenues have been explored, if social or physical issues cannot be overcome, transfer to HD may be necessary as long as the same issues do not deter appropriate therapy (e.g., adequate ultrafiltration, single pool delivered \( \text{Kt/V}_{\text{area}} \) of 1.2 thrice weekly, etc.) on this modality.\(^{128,135}\)

**Inadequate Solute Transport or Fluid Removal.** Peritoneal solute transport determined by PET affects both solute and fluid removal by PD. Obviously, peritoneal transport type is discovered after initiation of PD by the first PET. High transporters may have poor ultrafiltration and/or excessive protein losses (relative contraindication). Excessive protein losses are those that exceed the patient’s ability to compensate by an increase in dietary protein consumption. However, peritoneal urea and creatinine clearances tend to be adequate in high transporters. Many high transporters with poor ultrafiltration can be effectively dialyzed with short dwell periods and daytime exchanges, but such a regimen may become too burdensome for the patient’s lifestyle.\(^{114,128,135}\)

Low transporters usually have adequate ultrafiltration, but, when they are relatively large, may have inadequate peritoneal clearance of creatinine, but not necessarily a decreased clearance of urea.\(^{61}\)

Excessive protein losses can occur if the patient’s underlying disease includes active nephrosis, if the patient is a high transporter, or if frequent peritonitis occurs. The resulting malnutrition will increase the patient’s mortality and morbidity. In some children who are actively nephrotic, protein losses may be successfully replaced by supplemental (e.g., nasogastric, gastrostomy) tube feedings.

There are medical complications that may develop or have been present prior to initiation of dialysis, but these may become apparent only after peritoneal equilibration testing and adequacy studies.

Inadequate solute transport documented by measures of \( \text{Kt/V}_{\text{area}} \) and creatinine clearance
must be evaluated. If the maximum PD prescription has been reached (increases in volumes and frequency of exchanges including use of nocturnal cycling) or the procedure is no longer achievable due to lifestyle complications, hemodialysis as an alternative should be explored.\textsuperscript{114}

These guidelines have defined adequate solute transport with regard to urea and creatinine. However, the failure to adequately remove other solutes such as potassium may require switching to another form of renal replacement therapy.

Inadequate ultrafiltration is usually secondary to high transport characteristics or a mechanical defect hampering catheter patency or drainage.\textsuperscript{136} In rare instances, inadequate ultrafiltration is associated with low peritoneal transport characteristics, probably due to a significant reduction in the area of the peritoneal membrane, or secondary to increased peritoneal lymphatic flow.\textsuperscript{137}

**Unacceptably Frequent Peritonitis.** The definition of unacceptably frequent peritonitis has to be individually determined for each patient. Such considerations as the availability of hemodialysis facilities will inevitably play a role.

**Unmanageably Severe Hypertriglyceridemia.** Unmanageably severe hypertriglyceridemia, resulting from, or exacerbated by, the dextrose load intrinsic to the dialysate, may increase the risk for cardiovascular disease.

**Development of Technical/Mechanical Problems.** Irreparable technical or mechanical defects, such as catheter malposition, resulting in access failure.

**Severe Malnutrition Resistant to Aggressive Management (Relative).** Due to the continuous protein loss associated with PD, malnourished patients must be aggressively evaluated and treated. If treatment of malnutrition is not successful, transfer to HD is indicated.

**RECOMMENDATIONS FOR RESEARCH**

The topic of suitability of patients for PD or HD has not been thoroughly investigated. Prospective comparisons of PD and HD for specific ESRD patient categories (eg, those with severe heart failure, those with advanced malnutrition, those with large body size, etc.) are needed to define the subsets of ESRD patients which are most suitable or unsuitable for PD.
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Appendix A: Detailed Rationale for Guideline 1

GUIDELINE 1
When to Initiate Dialysis—Kt/V_{urea}
Criterion (Opinion)

Unless certain conditions are met, patients should be advised to initiate some form of dialysis when the weekly renal Kt/V_{urea} (Kt/V_{urea}) falls below 2.0. The conditions that may indicate dialysis is not yet necessary even though the weekly Kt/V_{urea} is less than 2.0 are:

1. Stable or increased edema-free body weight. Supportive objective parameters for adequate nutrition include a lean body mass >63%, subjective global assessment score indicative of adequate nutrition (see Guideline 12: Nutritional Status Assessment and Appendix B: Detailed Rationale for Guideline 2) and a serum albumin concentration in excess of the lower limit for the lab, and stable or rising; and

2. nPNA \approx 0.8 \text{ g/kg/d} (see Guideline 2: When To Initiate Peritoneal Dialysis—nPNA Criteria and Appendix B: Detailed Rationale for Guideline 2); and

3. Complete absence of clinical signs or symptoms attributable to uremia.

A weekly Kt/V_{urea} of 2.0 approximates a renal urea clearance of 7 mL/min and a renal creatinine clearance that varies between 9 to 14 mL/min/1.73 m². Urea clearance should be normalized to total body water (V) and creatinine clearance should be expressed per 1.73 m² of body surface area. The GFR, which is estimated by the arithmetic mean of the urea and creatinine clearances, will be approximately 10.5 mL/min/1.73 m² when the Kt/V_{urea} is about 2.0.

Rationale
In patients with chronic renal disease, progression of renal failure should be monitored by following total weekly renal urea nitrogen clearance (Kt_{urea}) normalized to urea volume of distribution (V), i.e., Kt/V_{urea}. This does not imply that a weekly collection of urine is necessary. A daily collection multiplied by seven yields a reasonable approximation of weekly clearance. The knowledge of Kt/V_{urea} is especially important when glomerular filtration rate (GFR) falls below 25 to 50 mL/min, at which time spontaneous decrease in dietary protein intake is commonly observed. The blood urea nitrogen (BUN) and serum creatinine values should not be used to monitor progression of renal failure, particularly in patients with diabetes. BUN may be low secondary to low protein intake and may not adequately reflect the degree of the renal functional impairment. Serum creatinine may be low due to decreased muscle mass as seen in some women, the elderly, and in malnourished patients. Hence, serum creatinine concentration may not adequately reflect the degree of the renal functional impairment.

The estimation of V (total body water) by any formulae has not been validated in children with renal failure. Thus, the use of Kt/V_{urea} as an indication for the initiation of PD is recommended considering this caveat. Creatinine clearance as a means of assessing RRF for purposes of initiation of dialysis should be normalized to body surface area (BSA).

An increasing body of evidence suggests that Kt/V_{urea} is a reliable predictor of outcome in PD and that weekly values in the range of 2.0 provide adequate therapy (see Guideline 15: Weekly Dose of CAPD). Although the CANUSA data indicated a linear decrease in modeled mortality rate with increasing Kt/V up to 2.3, there is some uncertainty about the significance of the high Kt/V levels achieved in this study.

It has always been a paradox that nephrologists have insisted on optimal therapy once patients are started on dialysis but have accepted much lower levels of renal function, defined as Kt/V_{urea}, during the pre-dialysis phase of patient management. For example, while we recognize that a weekly Kt/V_{urea} of 2.0 or higher is associated with improved outcome on PD, dialysis is usually not initiated until weekly Kt/V_{urea} is in the range of 0.71 to 1.3. It is possible that the consequences of delaying initiation of PD may be analogous to the experience of the National Cooperative Dialysis Study, wherein the mortality rate in the year after the study ended was more than twice as high in those randomized to the low dose dialysis protocols, even though they
were returned to standard dialysis after completing 24 weeks in the high BUN (low clearance) arm of the study. The Work Group feels that the data of Ikizler et al., McCusker et al., and Pollock et al. strongly demonstrate the linkage between decreasing renal function and worsening nutritional status. In the CANUSA study, multiple estimates of nutritional status were associated with two estimates of RRF. Patients who started PD at lower levels of RRF had a worse nutritional status than those who started at a higher level of RRF. CANUSA further demonstrated an association between the relative risk of death and worse baseline serum albumin concentration, worse time-dependent SGA, and worse time-dependent percent lean body mass.

While an association between risk of death and nPCR could not be demonstrated in the multivariate analysis, several univariate analyses did demonstrate an association of individual estimates of baseline nutritional status with survival. These data are consistent with the observations of Bonomini et al who found that patients starting dialysis with a residual renal creatinine clearance of <5 mL/min had a worse long-term outcome than patients starting incremental hemodialysis with a mean residual renal creatinine clearance of 11 mL/min.

From these observational data, it seems reasonable to draw the following conclusions:

Once $K_t/V_{area}$ falls below 2.0 per week or creatinine clearance falls into the range of 9 to 14 mL/min/1.73 m², initiation of dialysis or transplantation should be strongly advised. The patient should be considered to be at increasing risk with any further decreases in $K_t/V_{area}$ in the absence of renal replacement therapy intervention. If dialysis is not instituted when $K_t/V_{area}$ falls below 2.0, it is mandatory to document: (1) nPNA ≥0.8 g/kg/d (see Guideline 2 and Appendix B), (2) stable or increased body weight without edema, and (3) a negative inquiry for clinical signs or symptoms of uremia. When PD is initiated, the $K_t/V_{area}$ could be increased incrementally such that the combined value of $K_t/V_{area}$ + $K_p/V_{area}$ does not fall below the target level of 2.0 (see Figure II-1 and Guideline 15: Weekly Dose of PD). Alternatively, the initiation of “full dose” PD may be offered. Since residual renal function (RRF) is such a crucial component of total solute removal utilizing the incremental initiation approach, more intense scrutiny of RRF is necessary. For initiation with full dose PD, less intense scrutiny of RRF is indicated. This is discussed in Guideline 3: Frequency of Delivered PD Dose and Total Solute Clearance Measurement Within Six Months of Initiation and Guideline 5: Frequency of Measurement of $K_t/V_{area}$, Total $C_r$, PNA, and Total Creatinine Appearance, which address frequency of measurements.

In the CANUSA study, the weekly $C_r$ equivalent to a $K_p/V_{area}$ of 2.0 was 70 L/wk/1.73 m². As will be clear later in this discussion, a $C_r$ level this high is indicative of residual renal function, which was clearly present in the CANUSA patients at initiation of PD.

CAPD is the only continuous chronic renal replacement therapy with which to quantitatively compare continuous residual renal solute clearance. The Work Group strongly supports the opinion that the outcome data for a weekly $K_t/V_{area}$ of ≥2.0 are so compelling that using the same figure for initiation of dialysis justifies the unknown but presumably small risks of performing peritoneal dialysis. Those risks include infections and the possibility that increasing the length of time on PD contributes to eventual patient “burn-out.” If a patient is suspected to be at high risk for these complications, PD may not be the best choice for renal replacement therapy.

The Work Group recognizes that the patient will play a major role in accepting the initiation
of dialysis based on a certain "laboratory value." It is the responsibility of the care providers to make clear to the patient the rationale for initiating dialysis when the above conditions become applicable. Reasons to justify a delay in initiating dialysis are listed above. These reasons should be documented, if present.

The Work Group also recognizes that for many clinicians, initiating dialysis based on Kt/V urea is a new concept. Therefore, we have attempted to equate this to the traditional measure of urea clearance, creatinine clearance, and GFR (estimated by the arithmetic mean of urea and creatinine clearance).

What follows is an explicit quantitative approach to the concept of renal urea clearance (Kt/V urea, mL/min). We recommend that adequate PD be considered to require:

\[ K_{pt/V} = 2.0 \text{ per week where } K_{pt} \text{ is total weekly peritoneal plus renal urea clearance.} \]

**Guideline 15: Weekly Dose of CAPD explains the rationale for recommending that Kpt/V be 2.0 per week.** (1)

For the purpose of this discussion, Kt/V urea is considered equivalent to Kt/V urea and, therefore,

\[ K_{pt/V} = 1.44 \times K_{t/V} \text{ urea/V, where 1.44 converts mL/min to L/d, when Kt/V is 2.0.} \]

\[ 2.0 = 1.44 \times K_{t/V} \text{ urea/V and } \]

\[ K_{t/V} \text{ urea} = 0.20 \times V \text{ (4)} \]

Equation 4 shows that Kt/V urea must equal 0.2 times V when Kt/V = 2.0. It is of interest to compare this criterion to those developed independently for hemodialysis (HD). For twice weekly (biw) HD, a coefficient has been developed from urea kinetic modeling to convert Kt/V urea, mL/min, to L of equivalent urea clearance during each twice weekly dialysis. The twice weekly adequate level of Kt/V urea in HD is 1.85 (double pool) and the coefficient to convert Kt/V urea equivalent urea clearance during dialysis is 9.0 with units of L/treatment/mL Kt/V urea. Therefore,

\[ 1.85 = 9.0 \times K_{t/V} \text{ urea/V or } \]

\[ K_{t/V} \text{ urea} = 0.20 \times V \text{ (6)} \]

Equation 6 for biw HD is identical to Equation 4 for PD. For thrice weekly (tiw) HD the (double pool) coefficient previously developed is 5.0, therefore,

\[ 1.0 = 5.0 \times K_{t/V} \text{ urea/V or } \]

\[ K_{t/V} \text{ urea} = 0.20 \times V \text{ (8)} \]

Since Equations 4, 6, and 8 are identical, it is apparent that PD, biw HD, and tiw HD should all be started when Kt/V urea = 0.20*V. For an average patient with V = 35 L, this defines a level of Kt/V urea = 7.0 mL/min. There are constraints on the lower level of Kt/V urea for biw HD. As Figure II-1 suggests, treatment could be started incrementally once weekly Kt/V urea falls below 2.0. Typically, this would involve a single overnight exchange, intended to restore Kt/V urea to 2.0 per week. Ultrafiltration would not be needed since at this level of Kt/V urea, urine volumes are usually adequate.

**Levels of Residual Renal Creatinine Clearance, C_{Cr}, mL/min at Which Dialysis Should Be Initiated.** There are no K_{Cr} criteria for HD, and the criteria defining the contribution of residual renal function (K_{Cr}) to therapy are different from the criteria defining the contribution of peritoneal creatinine clearance (K_{PCR}) to the dose of therapy. The problem arises from the argument that tubular secretion of creatinine should be subtracted from the total renal creatinine clearance, and renal function with respect to creatinine clearance is best expressed as "GFR" as developed below. Therefore, the definitions used with respect to K_{Cr} will each be considered separately and related to the level of Kt/V urea for which K_{t/V} urea and K_{PCR} are considered simply additive.

In all instances, the total weekly C_{Cr} is normalized to 1.73 m² of BSA. In order to compare this to the Kt/V urea, BSA must be normalized relative to V, e.g., based on the Hume equations and taking the mean of the genders, 1.73 m² \approx 35 L. It is also reasonable to extend this as a linear relationship over the domain of patient size although it should also be considered a gender-dependent relationship.

**Consideration of Renal Contribution to Dose Expressed as C_{Cr}**. As noted above, in all cases the dose of total creatinine clearance is expressed as L/wk/1.73 m² of BSA. The following will normalize total creatinine clearance to 1.73 m² (nBSA) and Kt/V urea to a standard V = 35 L corresponding to 1.73 m² of BSA in order to develop constants relating the creatinine and urea-based
dosage parameters. To the extent that BSA and V increase and decrease at the same ratio (reasonably valid), the constants developed are generalizable, and therefore, the relation between \( K_r \) and \( nBSA \) can be expressed as follows:

\[
[K_r t/nBSA] = 1.44*7*K_r t/nBSA = 10.10*K_r (9)
\]

Equation 9 simply describes the total weekly liters of renal creatinine clearance as a function of \( K_r \) and our normalized BSA of 1.73 m².

For the \( K_t/V \) normalized to \( V = 35 \), therefore,

\[
K_t/V = 1.44*7*K_t/V = 0.29*K_t \text{ and, therefore,} \quad \frac{K_t}{nV} = 3.47*K_r t/nV (10)
\]

Assuming

\[
K_r = 2*K_u, \text{ we can substitute Equation 11 into Equation 12 to show} \quad (12)
\]

\[
K_r = 2*3.47*K_t/nV = 6.94*K_t/nV (13)
\]

Substituting now Equation 13 into Equation 9 yields the following equation:

\[
[K_r t/nBSA] = 10.10 [K_r t/nV] = 70 (K_t/nV) (14)
\]

Equation 14 shows that if we define the renal contribution to PD therapy by \( K_r \), the level of weekly \( K_r \) per 1.73 m² must be 70 times the \( K_t/nV \) so at \( K_t/nV \) of 2.0, \( K_r \) t/1.73 m² is 140. For the average patient with \( V = 35 \) L and BSA = 1.73 m², the required level of \( K_r \) = 14 mL/min.

Consideration of Renal Contribution to Dose Expressed as "GFR." In this case, the effective renal creatinine clearance, \( eK_r \), is defined as:

\[
eK_r = GFR = [K_r + K_u]/2 (15)
\]

Therefore,

\[
eK_r t/nBSA] = 1.44*7 (K_r + K_u)/2
\]

\[
= 5.0*K_r + 5.0*K_u (16)
\]

Substituting from Equations 11 and 13, yields

\[
[eK_r t/nBSA] = 5*6.94 (K_t/nV)
\]

\[
+ 5*3.47 (K_t/nV) = 52 (K_t/nV) (17)
\]

Equation 17 shows that if the renal contribution to PD therapy is defined as GFR, which is equivalent to "effective" creatinine clearance or \( eK_r \), the total weekly \( eK_r \) required relative to \( K_t/V \) is 52 L/wk/1.73 m² per unit of \( K_t/V \). It can also be noted that in all instances these equations also relate to \( K_r t/V \) and \( K_r t/V \) since we have defined \( K_r = K_r \).

Consideration of Peritoneal Creatinine Clearance to PD Dose. In this case, the dose must be expressed directly in terms of peritoneal creatinine clearance (\( K_p \)) and by definition,

\[
[K_p t/nBSA] = 1.44*7*K_p = 10.10 K_p (18)
\]

The average relationship between \( K_p \) and \( K_r u, K_r u/t/nV, \) Equations 18, 19, and 11 can be combined to derive

\[
[K_p t/nBSA] = 28 [K_r u, t/nV]
\]

\[
= 28 [K_r u, t/nV] (20)
\]

Equations 14, 17, and 20 show that the Creatinine Dose Equivalency with respect to the single urea \( K_p t/nV \) criterion will vary widely depending on how RRF is defined. The relationships are plotted in Figure II-2, Dose of PD With Respect to
Weekly Creatinine Clearance Relative to Weekly 
K_{pt}/V, where the weekly creatinine clearance re-
quired per 1.73 m² corresponding to a weekly K_{pt}/V_{urea} of 2.0 ranges from 140 to 56 L, depending on the definitions used. There is no problem in either the case of pure residual renal function with no PD therapy or the case of pure peritoneal dialysis with no residual renal function present. The problem arises when one attempts to sum residual renal creatinine clearance with peritoneal creatinine clearance. In this common circumstance, the dialysis dose relationship will be bounded by regression lines 2 and 3 in Figure II-2.

For line 1, which defines creatinine clearance as the uncorrected (for secretion) creatinine clearance, the creatinine clearance that is equivalent to a K_{t/V_{urea}} of 2.0 is 140 L/wk/1.73 m². Line 2 defines creatinine clearance as the mean of urea and creatinine clearance, and the equivalence to a K_{t/V_{urea}} of 2.0 is 104 L/wk/1.73 m². Line 3 represents all clearance from PD (complete absence of residual renal function). Under this condition a K_{t/V_{urea}} of 2.0 is equivalent to a creatinine clearance of 56 L/wk/1.73 m².

This creates an irreconcilable ambiguity with respect to the creatinine and urea dosage criteria for defining optimal dialysis and the study of outcome as a function of dose. Because the practice of PD has used both C_{Cr} and K_{t/V} to quantify delivered dose and there is a large body of literature describing outcomes related to C_{Cr}, the Work Group recommends continuing to use both measures (see Guideline 4: Measures of PD Dose and Total Solute Clearance.) However, in view of this ambiguity, the Work Group recommends that if only one measure is to be utilized, use K_{pt}/V_{urea} rather than K_{Cr} (see Guideline 15: Weekly Dose of PD.) Nonetheless, creatinine kinetics as discussed in Guidelines 4, 6, and 17 are useful for estimating edema-free, fat-free body mass, compliance with dialysis prescription, and some programs may prefer it for quantification of delivered dose of PD. Thus, total creatinine excretion is valuable.

Finally, it is worth emphasizing that the basic relationship between the level of residual C_{Cr} to K_{t/V_{urea}} for initiation of dialysis will be the same for all of these creatinine dosage criteria. They are related to K_{pt}/V since K_{Cr} = 2 K_{t/V_{urea}} and all of the expressions ultimately reduce to this relationship.

Another way to view the creatinine clearance at which to initiate dialysis is to extrapolate backward from the CAPD target of 60 L/week/1.73 m² (see Guideline 15: Weekly Dose of PD.) This approximates a purely filtered only creatinine clearance of 6 mL/min. Since at this level of residual renal function much of the creatinine appearing in the final urine is from tubular secretion, 60 L/wk/1.73 m² approximates a total residual renal creatinine clearance of 9 to 14 mL/min.

**Peritoneal Dialysis and Residual Renal Function Equivalency.** Quantitative replacement of renal urea clearance by peritoneal clearance is based on the assumption that the two clearance parameters confer equal clinical benefit with respect to control of uremic morbidity. Thus, we can write the relationship

\[
K_{pt}/V_{urea} = K_{pr}/V_{urea} + K_{t}/V_{urea},
\]

where K_{pt}/V_{urea} is total daily or weekly peritoneal urea clearance normalized to V; K_{t}/V_{urea} is total daily or weekly renal urea clearance normalized to V. (21)

Solution of Equation 21 for K_{pt}/V_{urea} with the assumption that adequate weekly K_{pt}/V_{urea} is 2.0 results in

\[
K_{pt}/V_{urea} = 2.0 - K_{t}/V_{urea}
\]

Equation 22 provides a quantitative guideline for replacing residual renal urea clearance by peritoneal clearance such that the sum of weekly K_{pt}/V_{urea} and K_{t}/V_{urea} remains 2.0.

**Hemodialysis and Residual Renal Function Equivalency.** Compared to CAPD it is more complex to calculate incremental doses of HD such that the sum of intermittent dialyzer clearance (K_{d}/V_{urea}) and continuous K_{t}/V_{urea} remain constant at a level equivalent to a weekly K_{t}/V_{urea} of 2.0. However, the dose and frequency of HD which provide therapy equivalent to continuous K_{pt}/V_{urea} can be calculated using the fundamental assumption underlying CAPD therapy: the level of continuous K_{pt}/V_{urea} required for treatment which is clinically equivalent to intermittent HD is that K_{pt}/V_{urea} which results in a steady state BUN equal to the average predialysis BUN with any specific intermittent HD treatment schedule at the same nPCR. From this basic assumption and the urea kinetic model the dose and frequency of HD required for incremental replacement of K_{t}/V_{urea} as it falls below 2.0 can be readily calculated as depicted in Figure II-3.
Fig 11-3. Equivalent total dialysis doses for incremental replacement of $K_t/V_{urea}$ calculated with the assumptions that (1) $K_t$, $K_p$, and $K_d$ are clinically equivalent clearance terms and (2) the intermittent dialysis dose schedule is equivalent to continuous dialysis when average pre-dialysis BUN equals steady state BUN with continuous therapy at equal nPCR. Note that $eK_d/V_{urea}$ equals the equilibrated (double-pool), delivered, and normalized hemodialysis dose.

The dose of intermittent HD is expressed in Figure II-3 as $eK_d/V_{urea}$, which is the equilibrated, delivered and normalized hemodialysis dose (see the NKF-DOQI Clinical Practice Guidelines for Hemodialysis Adequacy for further discussion of $eK_d/V_{urea}$). The equilibrated measure is utilized here because in peritoneal dialysis, transcellular urea equilibration is achieved, and therefore, it makes conceptual sense to think in terms of equilibrated values. From preliminary data of the HEMO study, $eK_d/V$ is approximately 0.21 lower than that computed from immediate post-HD BUN sampling, using single-pool, variable volume kinetic modeling. The dashed line depicts incremental increase in daily $K_d/V_{urea}$ as $K_d/V_{urea}$ falls in accordance with Equation 22. Model solutions are shown for once, twice and thrice weekly hemodialysis ($N = 1$, 2, and 3, respectively). The model solutions are limited to $eK_d/V_{area} \leq 2.0$ since it is unrealistic to prescribe $eK_d/V_{area} > 2.0$. Such a dose would correspond to single pool $K_d/V_{urea}$ values of 2.8 and 2.3 with treatment times (t) of 2.0 and 4.0 hours, respectively. It can be seen that when $N = 1$, $eK_d/V_{urea} = 2.0$ when $K_d/V_{urea} = 1.6$. Thus, the option for once weekly hemodialysis is limited to a weekly $K_t/V_{urea} \geq 1.6$. If $K_t/V_{urea} = 0.5$ and $N = 2$, the $eK_d/V_{area}$ for each HD treatment must be 2.0 to achieve a weekly continuous $K_t/V_{urea}$ equivalent to 2.0. Therefore, for a weekly $K_t/V_{urea} < 0.5$, more than twice weekly HD will be necessary. Finally, in the case of $N = 3$, $eK_d/V_{urea}$ increases linearly to 1.05 as $K_t/V_{urea}$ falls to zero. The $eK_d/V_{urea}$ of 1.05 corresponds to single pool $K_d/V_{area}$ values of 1.46 and 1.20 at treatment times of 2.0 and 4.0 hours, respectively.

There is emerging evidence that RRF is better preserved in patients undergoing HD with the use of more biocompatible membranes.

APPENDIX A REFERENCES

APPENDICES


Appendix B: Detailed Rationale for Guideline 2

GUIDELINE 2
When to Initiate Dialysis—nPNA Criterion (Opinion)

It is recognized that an adequately supplemented, tightly monitored, low protein diet may slow the progression of renal failure in certain circumstances. When properly performed, such diets do not result in loss of lean body mass or other manifestations of malnutrition. However, progressive renal failure is associated with spontaneous anorexia and malnutrition. Under these circumstances and in the absence of comorbid causes of anorexia, and after unsuccessful intervention by a registered dietitian, dialysis should be started in adult patients when nPNA spontaneously falls below 0.8 g/kg/d.

Rationale The extrapolation of data from dialysis patients to pre-dialysis patients and vice versa may be appropriate, if cautiously applied. The Work Group attempts in this rationale to address the pre-dialysis patient population. It is recognized that an adequately supplemented, tightly monitored, low protein diet may slow the progression of renal failure in certain circumstances⁴ and may delay the need for dialysis. Patients being treated by such diets do not demonstrate findings of malnutrition or loss of lean body mass. Clearly though, energy intake must be increased to perhaps as high as 40 kcal/kg/d,² and these patients must be monitored very closely. This is particularly relevant since the USRDS reports that 45% of incident PD patients in 1996 had never visited with a dietitian prior to dialysis.³

However, often there is a spontaneous reduction in dietary protein intake as renal failure progresses. The USRDS reports that at least 40% of incident PD patients in 1996 began dialysis after months of nausea/vomiting.³ Estimating PNA from renal urea nitrogen kinetics will provide an estimate of protein intake. Malnutrition often follows prolonged decreases in protein intake, unless adequate energy intake is added to compensate. PNA measurements can be used to assist in documenting malnutrition. Malnutrition present at the initiation of dialysis has an adverse effect on survival. Thus, monitoring nutritional indices in pre-dialysis patients provides a means of early recognition so that appropriate interventional measures may prevent further deterioration of nutrition. The initiation of dialysis is one such appropriate measure, when causes of malnutrition other than uremia have been ruled out.

The level of spontaneous protein intake (as reflected by urea nitrogen appearance) prior to the initiation of dialysis has been shown in some studies under certain circumstances to be predictive of outcome.⁴⁵ A spontaneous decrease in protein intake occurs once the GFR falls below
50 mL/min\textsuperscript{5,10} and may fall below 0.7 g/kg/d when the GFR is <25 mL/min.\textsuperscript{7} In addition, it has been shown that serum albumin concentration\textsuperscript{7,11-15} and the subjective global assessment (SGA) of nutritional status at initiation of PD are both predictive of subsequent patient outcome.\textsuperscript{6} The method of SGA of nutritional status used in the CANUSA Study was a modification of the original method\textsuperscript{16,17} (see Appendix F: Detailed Rationale for Guideline 12). This technique, initially described in surgical patients, has been validated in ESRD patients.\textsuperscript{18} Four items (weight loss during a predetermined interval, anorexia, subcutaneous fat, and muscle mass) are scored on a 7-point Likert scale. Likert scales are responses to a question in which the answers have multiple discrete response options and directionality. Values as high as 6 or 7 reflect normal
nutrition and values as low as 1 or 2 reflect severe malnutrition. Intermediate values such as 3, 4, or 5 reflect moderate to mild malnutrition. Further confirmation of the relation between RRF and nutritional status has been presented. Using univariate analysis, a lower level of renal function at the time of initiation of PD (measured by $K_{\text{t/V}}$, $C_\text{Cr}$, or urea clearance) was associated with worse nutritional status (serum albumin concentration, lean body mass, SGA, and nPNA), which, in turn, was predictive of a poor clinical outcome on PD over the subsequent 24 months.

While an optimal level of protein intake for PD patients has not been rigorously defined, most would agree that neutral nitrogen balance could be maintained at an intake of 1.0 to 1.2 g/kg/d, and that negative nitrogen balance is likely at levels of protein intake below 0.8 g/kg/d. There is a marked difference between nPNA and weekly $K_{\text{t/V}}$ values in patients with continuous clearance (CAPD and pre-dialysis chronic renal failure patients) versus patients on chronic hemodialysis. In the former groups, $K_{\text{t/V}}$ values of 2.0 or more were required to achieve a nPNA value of 0.9 g/kg/d or greater.

Therefore, the Work Group recommends that patients with progressive renal failure should have quarterly estimates of nPNA (from 24-hour urine collections) once GFR falls below 25 mL/min. When nPNA is below 0.8 g/kg/d, vigorous nutritional intervention should be undertaken. Some patients may be specifically advised to maintain protein intake below 0.8 g/kg/d by design, with the close involvement of a registered dietician, a caloric intake of at least 35 kcal/kg/d, and careful follow-up with the nephrologist. However, a spontaneous reduction of protein intake to below 0.8 g/kg/d is suggestive of uremia. Comorbid conditions which may cause anorexia such as gastroparesis, infection, acidosis and depression should be identified and treated.

A registered dietician must be involved by this stage. It is appropriate to utilize the registered dietician at the dialysis facility to whom the patient will be referred. However, if these measures fail to restore nPNA to 0.8 g/kg or higher, dialysis should be initiated and increased incrementally until a satisfactory level of nPNA is achieved.

The recommendation to use an nPNA criterion for the initiation of dialysis does not imply that a broader view of nutritional status should be neglected. On the contrary, the presence of symptoms, such as the following, should influence the clinical judgment regarding the need for dialysis:

- anorexia
- weight loss
- nausea
- vomiting
- falling caloric or protein intake by history
- decreased fat-free, edema-free (lean) body mass by creatinine kinetics (see Guideline 13: Determining Fat-Free, Edema-Free Body Mass)
- declining subjective global assessment of nutritional status (see Appendix F)
- decreasing serum albumin concentration, etc.

These topics are discussed from a slightly different perspective in Guideline 28: Measurement of Normalized PNA in PD Patients. A detailed review of nutrition in ESRD and pre-dialysis patients has recently been presented.

APPENDIX B REFERENCES


9. Kopple JD, Chumlea WC, Gassman JJ, Hotlinger DL,
Appendix C: Detailed Rationale for Guideline 6

Assessing Residual Renal Function (Evidence)

Residual renal function (RRF), which can provide a significant component of total solute and water removal, should be assessed by measuring the renal component of Kt/V urea (Kt/V urea) and estimating the patient’s glomerular filtration rate (GFR) by calculating the mean of urea and creatinine clearance.

Rationale

The contribution of RRF to total solute and water clearance is significant (30% to 50%), especially during the first few years of dialysis therapy. Assessment of RRF is important for several reasons. A substantial fraction (30%) of the total renal replacement therapy may be provided by RRF when a patient begins PD. After 2 years of PD, the RRF may still contribute about 15% of the total Kt/V urea. Since the RRF contribution will be added to that of PD, it will be measured in the same units and for the same solutes.

Preservation of RRF may be of particular importance to the effectiveness of long term PD therapy. There is a progressive decline of RRF over time with both HD and PD. Several studies have compared the rate of decline of RRF with the two dialytic modalities and demonstrated that RRF is preserved better in patients undergoing PD therapies compared to HD. In a study of 25 CAPD patients and 25 HD patients, the rate of decline of creatinine clearance was significantly slower over the first 18 months of dialysis in the CAPD patients. The PD patients started dialysis with an uncorrected CCr of 4.4 mL/min, and after 18 months it was 4.0 mL/min. In the HD patients, CCr at initiation was 4.3 mL/min, and after 18 months it was 1.3 mL/min (P <0.01 compared to PD).

Similar differences were observed in patients with diabetes. In another study comparing the urine output and CCr, the urine output dropped significantly in HD patients at the end of one year compared to 3 years in CAPD patients. The mean annual decline of CCr was identical in HD and CAPD for patients with primary glomerulopathy. However, in the groups with nephrosclerosis and interstitial nephropathy, the rate of decline of CCr was significantly slower in CAPD compared to HD patients. In another retrospective study of 4 years duration which compared 55 CAPD patients to 57 HD patients, the rate of...
decline in the HD group was twice that of the CAPD group. This difference persisted after adjustment for age, gender, hypertensive status, and the use of ACE inhibitors. Children have a better preservation of urinary volume, but not GFR, in those receiving PD.

Despite their limitations, these studies generally demonstrate a slower rate of decline of RRF in patients on PD compared to HD. They also demonstrate that the rate of the decline varies from patient to patient. The faster rate of decline of RRF in HD is speculated to be due to repetitive hypotensive episodes, possibly complement activation and cytokine release, and the possibility that the more efficient HD may remove GFR stimulatory factors.

Several methods for measuring the Ccr component of RRF are available. These include the uncorrected Ccr, a flat percentage of uncorrected Ccr as an estimate of GFR, or the average of creatinine and urea clearance also as an estimate of GFR.

While each method has its particular merits, the Work Group recommends using the arithmetic mean of creatinine and urea clearances to determine the RRF component to Ccr, and as an estimate of GFR. Therefore, the Ccr component of RRF will subsequently refer to residual renal Ccr, corrected for secretion by taking the arithmetic mean of urea and creatinine clearances. This method was selected for several reasons. First, it was used in some of the major outcome studies used in establishing these guidelines (see Guideline 15, Weekly Dose of CAPD.) Second, the corrected Ccr correlates better with Kt/Vurea than the uncorrected Ccr.9 Third, it makes conceptual sense because the peritoneal transport of creatinine is by diffusion and convection, not secretion. The correction process addresses this.

However, the Work Group recommends using urea clearance, normalized to total body water, i.e., Kt/Vurea, as the key measure to follow serially to determine whether urine collections need to continue (see Guideline 11: Dialysate and Urine Collections.) This is termed the renal Kt/Vurea or Kt/Vurea. This recommendation was made to simplify the concept of a residual renal component to the total renal replacement dose. Kt/Vurea is believed to be the more valuable measure of renal replacement therapy and the Work Group carried this thinking through to using Kt/Vurea in both initiation of dialysis and for follow-up RRF changes over time.

Kt/Vurea as a measure of RRF is recommended because total Kt/Vurea is associated in a clinically important and statistically significant way with patient survival11,10 (see Guideline 15: Weekly Dose of CAPD). The peritoneal clearance of creatinine is about 80% of the urea clearance, while at end stage the renal clearance of creatinine is about 1.5 to 2 times that of urea. Perhaps as a consequence of this physiological phenomenon or for other reasons, there is a discrepancy between total Kt/Vurea and total Ccr normalized to 1.73 m2 BSA (see paragraphs below). In the case of discrepancy, Kt/Vurea is preferentially recommended to determine PD adequacy, because it is more predictable and reproducible, and is independent of the confounding effects of renal secretion of creatinine. A retrospective study of PD adequacy demonstrated an association between Kt/Vurea and outcomes.12

This emphasis on using Kt/Vurea is not intended to detract from the utility of Ccr. In terms of validity, total Ccr normalized to 1.73 m2 BSA is predictive of patient survival, technique survival, and hospitalization. The creatinine generation rate is useful for assessment of nutritional status, in particular, in measuring fat-free, edema-free body mass. Total Ccr may also be useful for assessment of compliance.

Ccr as an index of PD adequacy is associated statistically with both morbidity and mortality and correlates with urea clearance.13 Discrepancies between the two clearances may be found in approximately 20% of PD subjects.14,15 The main reasons for the discrepancies are the presence of substantial RRF, which tends to cause disproportionately high Ccr values, and low peritoneal solute transport type, which tends to cause disproportionately low Ccr values.14,15 Ccr values corresponding to a weekly Kt/Vurea of 2.0 differ between CAPD subjects with and without RRF (see Figure X-2, referenced previously in Appendix A: Detailed Rationale for Guideline 1). In patients with RRF, the mean Ccr corresponding to a Kt/Vurea of 2.0 weekly is between 60.513 and 67.615 L/wk/1.73 m2. In anuric CAPD subjects, the mean Ccr corresponding to a Kt/Vurea of 2.0 weekly is 52.1 L/wk/1.73 m2.12,15

In the case of a discrepancy, the Work Group recommends the use of Kt/Vurea as an immediate guide of dialysis adequacy because it directly
relates to protein metabolism. However, if there is a discrepancy between $C_{Cr}$ and $K_t/V_{urea}$, the patient must be observed closely because initially it may not be clear why the discrepancy exists and the reason may be important. This is discussed in Guidelines 1, 7, and 15.

**APPENDIX C REFERENCES**


**Appendix D: Detailed Rationale for Guideline 8**

**GUIDELINE 8**

**Reproducibility of Measurement (Opinion)**

Accurate measurement of total $K_t/V_{urea}$ and total creatinine clearance ($C_{Cr}$) requires collection and analysis of urine, dialysate and serum in a way that yields reproducible and valid results. Dialysate creatinine concentration must be corrected for the presence of glucose in some assays. Peritonitis precludes reliable measurement of delivered PD dose for up to a month. Compliance with complete collections is mandatory. For patients who void ≥3 times per day, a 24-hour urine collection is sufficient. For patients who void less frequently, a 48-hour collection is recommended. For CAPD patients, the serum sample can be obtained at any convenient time. For NIPD patients, the serum sample should be obtained at the midpoint of the daytime empty period. For CCPD patients, the serum sample should be obtained at the midpoint of the daytime dwell(s).

**Rationale**

To be clinically useful, measurement of PD dose must be performed in a valid and reproducible fashion. The following factors influence the validity and reliability of $K_t/V_{urea}$ and total $C_{Cr}$ as measures of PD dose.

**Dialysate glucose.** Dialysate creatinine concentration should be corrected for the presence of glucose, which interferes with some creatinine measurement methodologies. Each facility must determine this by specifically inquiring of its laboratory whether the creatinine assay used by that lab is altered by high glucose concentrations. Each laboratory should establish its own correction factor and should reestablish the correction factor if the laboratory’s methodology changes. The Work Group does not recommend using correction factors from the literature.
Peritonitis. Peritoneal solute transport increases during peritonitis and usually recovers some time after resolution of peritonitis, with a reported recovery time between 3 days\(^2\) and 1 month.\(^3\)

Patient compliance with the dialysis prescription. Following creatinine appearance in dialysate and urine longitudinally is an objective method to evaluate the degree of compliance (see Guideline 7: PD Dose Troubleshooting). Clinical tools for evaluating compliance are in the process of development.\(^4\)

Variability of residual renal function (RRF). Day-to-day total clearances can vary greatly in PD. The major portion of this variance is caused by changes in measured RRF,\(^5\) although creatinine generation may vary in apparently stable PD patients.\(^6\)

Completeness of urine collection. To avoid sampling errors, urine should be collected over 48 hours in patients who void infrequently (<3 times in 24 hours). A 24-hour urine sample can be used for all other patients. The urine collection should be performed on the same day as the dialysate collection. In children, the urine collection period may be reduced to a minimum of 12 hours.

Dialysate and urine collection for PD adequacy studies. Two methods of dialysate sampling predominate. In the first method, for CAPD, all effluent bags in a 24-hour period are brought to the center. While this “batch” method is simple in concept, it is difficult to carry out because it means transporting all the dialysate bags, which are heavy and bulky.

The second method is referred to as the “ aliquot” method. In this approach, each bag of effluent dialysate is shaken vigorously for a few seconds, then is emptied into a measuring container accurate to an error of <50 mL per 2000 mL. The volume for that bag is recorded in mL and the decimal point is moved three places to the left. The resulting figure is the number of mL which must be drawn from the dialysate effluent in the measuring container and placed in the laboratory red top test tube, provided by the dialysis center. For example, if the effluent volume for the CAPD bag is 2450 mL, moving the decimal point three places to the left means that 2.45 mL of this fluid is put in the test tube (or other small collection container). Each bag for the 24-hour interval is handled this way.

The aliquots are measured by syringes; usually a 5-mL syringe is accurate enough. The aliquots can be mixed in the same container, because the sampling proportion from each original bag is constant at 1/1000. The total effluent is recorded, and that figure plus the small container with all the collected aliquots are brought to the dialysis center. Some dialysate manufacturers have developed special aliquoting exchange bags that separate an aliquot as part of the exchange.\(^7\)

The collection of effluent dialysate from automated PD is conceptually similar to that described above. The effluent drained via a cycler is quantified automatically by the cycler and generally pools in one collection container, or if in several containers, free mixing is possible. Since the effluent volume is known and the containers are allowed to mix freely, a sample of any reasonable volume (e.g., 10 mL) can be brought to the dialysis unit. If several containers are used with equal filling, an equal volume aliquot from each container can be pooled. The total effluent volume must be known and recorded. If the effluent bags are not freely mixing, then a sample from each bag is required, as well as the exact volume of the container from which the sample was drawn. One cannot extrapolate from one container (bag) to the next.

No matter what method is used for dialysate, a complete and accurately timed urine collection is necessary. The urine volume is more easily managed since it is much smaller than the dialysate volume. The longer the collection interval, the more reliable are the collections, assuming patient compliance. A timed collection of 12 to 48 hours is recommended, depending on how frequently the patient voids. Polyuric patients, particularly children, or patients with a short attention span, may void frequently enough that a supervised 12-hour collection is accurate. As for any urine collection, the bladder should be emptied and the urine discarded moments before the start of the collection period. Then, moments before the end of the collection period, the patient empties the bladder, and this urine, plus all that has been collected in the interval, completes the collection. The patient should try to delay the final voiding until just before the interval ends. Three or more bladder voidings generally are necessary for urine collections. For patients who make little urine and hence void infre-
sequently, 48-hour collections may be more informative.

Serum samples. In CAPD, serum concentrations of urea and creatinine are relatively constant, and thus blood samples can be drawn at any convenient time for clearance determinations. For the asymmetric therapies (NIPD and CCPD), blood concentrations are lowest at the end of the cycling period and highest prior to the next cycling period. Theoretically, the best time to draw these blood samples is halfway between the lowest and highest times. For NIPD patients, the serum sample should be obtained at the midpoint of the daytime empty period. For CCPD patients, the serum sample should be obtained at the midpoint of the daytime dwell. For most NIPD and CCPD patients, this time point conveniently occurs in the early afternoon.

Appendix E: Detailed Rationale for Guideline 9

GUIDELINE 9

Estimating Total Body Water and Body Surface Area (Opinion)

V (total body water) should be estimated by either the Watson¹ or Hume² method in adults using actual body weight, and by the Mellits-Cheek method³ in children using actual body weight.

Watson method¹:

For Men: \( V = 2.447 + 0.3362 \times Wt \) (kg) + \( 0.1074 \times Ht \) (cm) - \( 0.09516 \times Age \) (yrs)

For Women: \( V = -2.097 + 0.2466 \times Wt + 0.1069 \times Ht \)

Hume method²:

For Men: \( V = -14.012934 + 0.296785 \times Wt + 0.192786 \times Ht \)

For Women: \( V = -35.270121 + 0.183809 \times Wt + 0.344547 \times Ht \)

Mellits-Cheek method for children³:

For Boys: \( V = -1.927 + 0.465 \times Wt \) (kg) + \( 0.045 \times Ht \) (cm), when \( Ht \leq 132.7 \) cm

\( V = -21.993 + 0.406 \times Wt + 0.209 \times Ht \), when height is \( \geq 132.7 \) cm

For Girls: \( V = 0.076 + 0.507 \times Wt + 0.013 \times Ht \), when height is \( \leq 110.8 \) cm

\( V = -10.313 + 0.252 \times Wt + 0.154 \times Ht \), when height is \( \geq 110.8 \) cm

Body surface area, BSA, should be estimated by either the DuBois and DuBois method⁴, the Ghan and George method⁵ or the Haycock method⁶ using actual body weight.

For all formulae, Wt is in kg and Ht is in cm:

DuBois and DuBois method:

\( BSA \ (m^2) = 71.84 \times Wt^{0.425} \times Ht^{0.725} \)

Gehan and George method:

\( BSA \ (m^2) = 0.0235 \times Wt^{0.51456} \times Ht^{0.42246} \)

Haycock method:

\( BSA \ (m^2) = 0.024265 \times Wt^{0.5378} \times Ht^{0.3964} \)

Rationale The practical methods described in the literature to estimate V include a fixed fraction of body weight (0.60 in males and 0.55 in females, or 0.58 in all subjects) and anthropometric formulae based on sex, age, height, and weight.¹-³ The fixed fraction method is inaccurate, as it overestimates total body water even in overhydrated PD subjects.⁷ Therefore, the Work Group recommends that this method not be used.

Both the Watson¹ and Hume² formulae were derived by comparing anthropometric measurements to measurements of body water by indicator dilution techniques. An advantage of these formulae is that they were derived in populations which included obese subjects and, therefore, can account for obesity. Estimates from the Watson and Hume formulae are, in general, close to isotopic body water measurements in PD patients.⁷ The error of the estimates based on these formu-
inadequate. Therefore, it is recommended in such individuals to provide a dose of PD which will result in adequate $K_{pt}/V_{area}$ when they reach their desired weight without changing the dialysis prescription. For malnourished patients defined by SGA or Table II-3 below, provide a PD dose to achieve a weekly $Kt/V_{area}$ of 2.0 for the volume of the patient at desired weight. The calculation of the target $K_{pt}/V_{area}$ in malnourished CAPD subjects equivalent to a weekly $K_{pt}/V_{area}$ of 2.0 at desired weight is as follows: If $V_{actual}$ is body water obtained from the Watson or Hume formulae using the actual weight and $V_{desired}$ is body water obtained by the same formulae using the desired weight, then for malnourished subjects $V_{actual} < V_{desired}$. The target CAPD weekly $K_{pt}/V_{desired}$ is 2.0 (for CCPD 2.1 and NIPD 2.2). If $K_{pt}/V_{actual} = x$, then for the CAPD patient:

\[
\frac{(K_{pt}/V_{desired})}{(K_{pt}/V_{actual})} = 2.0/x \quad (1)
\]

\[
2.0 * \frac{V_{desired}}{V_{actual}} = x, \quad (2)
\]

and $x$ is the new target $Kt/V_{area}$.

Equation 2 can be used to calculate the target $K_{pt}/V_{area}$ in malnourished CAPD subjects. For example, if $V_{actual}$ is 35 L and $V_{normal}$ is 40 L, then the weekly target $K_{pt}/V_{actual}$ is 2.0 times 40/35 or 2.29. In essence, the target of 2.0 is modified upward by a factor of $V_{desired}/V_{actual}$. The recommendation to increase the target $K_{pt}/V_{area}$ in malnourished PD subjects is based on indirect evidence.

The above water volume estimations are used in the $Kt/V_{area}$ measure. Target $Kt/V_{area}$ is modified (Equation 2). The same principle applies if $C_Cr$ is used. The normalization of $C_Cr$ by BSA should correct $weight_{actual}$ for $weight_{desired}$ in the formula used to determine BSA. This is further discussed in Guideline 15: Weekly Dose of CAPD.

The concept above is intended to deliver a dose of PD considered adequate for the patient at a “desired” weight. Defining a “desired” weight can be subjective, but objective definitions are available. One preferable method is that proposed by a broad collaborative “glossary” group where the “desired” weight described in this rationale is the glossary group’s “normal weight,” defined as the median body weight of normal Americans with the same age, height, sex, and skeletal frame as the patient in question. Table II-3 from the glossary details these weight
Table II-3. Median (50th Percentile) Body Weight (kg)\(^{11}\)

<table>
<thead>
<tr>
<th>Age Range (yr)</th>
<th>Males</th>
<th>Females</th>
</tr>
</thead>
<tbody>
<tr>
<td>18.0-24.9</td>
<td>&lt;38.3</td>
<td>&gt;41.6</td>
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<td>68.3</td>
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<td>&lt;38.6</td>
<td>&gt;42.1</td>
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<td>&lt;35.7</td>
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<td>57.6</td>
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<td>61.8</td>
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<td>&gt;42.5</td>
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<td>Median body weight (kg)</td>
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<td>62.8</td>
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<td>63.4</td>
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<td>63.7</td>
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</tr>
<tr>
<td>Median body weight (kg)</td>
<td>72.6</td>
<td>62.9</td>
</tr>
</tbody>
</table>


Frame Index has no units and is calculated as follows: [Elbow Breadth (mm)/Height (cm)] × 100.


Alterations Caused by Amputation. The anthropometric formulae for total body water calculate that as obesity develops and body weight increases, \(V\) also increases, but body water content (the ratio \(V/\text{weight}\)) decreases. This is consistent with the known fact that water content of fat tissue is low. Calculations of \(V\) in amputees by uncorrected anthropometric formulae (using the actual post-amputation weight and height in the calculations) distorts the relationship between \(V\) and weight. In this case, body water content is not consistent with the degree of obesity.\(^{12,13}\) The anthropometric formulae can be corrected in a way that restores the relationship between body water content and degree of obesity in three steps:

Step A: The fraction of body weight lost to amputation (\(f_w\)) is obtained from a normogram\(^{14}\) (see Table II-4). The hypothetical non-amputated weight at the same body composition would be equal to actual weight/(1 – \(f_w\)).

Step B: \(V\) at the hypothetical non-amputated weight \((V_{\text{non-amputated}})\) is calculated from the Watson formula. Body water content is \(V_{\text{non-amputated}}/\text{Weight}_{\text{non-amputated}}\).

Step C: Actual \(V\) is calculated by multiplying the actual post-amputation weight by \(V_{\text{non-amputated}}/\text{Weight}_{\text{non-amputated}}\). This correction makes the as-
Table II-4. Fraction of Weight and BSA Corresponding to Amputated Limbs

<table>
<thead>
<tr>
<th>Body Part</th>
<th>% Loss in Weight</th>
<th>% of BSA to Subtract</th>
</tr>
</thead>
<tbody>
<tr>
<td>Foot</td>
<td>1.8</td>
<td>3.5</td>
</tr>
<tr>
<td>Leg below knee</td>
<td>6.5</td>
<td>10.0</td>
</tr>
<tr>
<td>Leg above knee</td>
<td>8.0</td>
<td>12.5</td>
</tr>
<tr>
<td>Leg at hip</td>
<td>18.5</td>
<td>18.0</td>
</tr>
<tr>
<td>Hand</td>
<td>0.8</td>
<td>2.5</td>
</tr>
<tr>
<td>Arm at elbow</td>
<td>3.1</td>
<td>6.0</td>
</tr>
<tr>
<td>Arm at shoulder</td>
<td>6.6</td>
<td>10.0</td>
</tr>
</tbody>
</table>

Assumption that amputation per se will not change body water content. The calculations for body surface area (BSA) in amputees should be also corrected, because of inconsistent results obtained with the uncorrected calculation of BSA. The correction requires three steps:

Step A: Same as step A in the correction of V in amputees.

Step B: BSA at the hypothetical non-amputated weight is calculated by one of the three BSA formulae above.

Step C: The fraction of BSA corresponding to the amputated limb(s) \( f_{BSA} \) is obtained from Table II-4 derived from Herndon. Corrected BSA is \( BSA_{Non-amputated}/(1 - f_{BSA}) \).

APPENDIX E REFERENCES


Appendix F: Detailed Rationale for Guideline 12

GUIDELINE 12
Assessment of Nutritional Status (Opinion)

Nutritional status of adult PD patients should be assessed on an ongoing basis in association with \( Kt/V_{urea} \) and \( CCr \) measurements using the Protein equivalent of Nitrogen Appearance (PNA) and Subjective Global Assessment (SGA). For pediatric PD patients, nutritional status should be assessed using the PNA and other standard nutritional assessments (see Guideline 14: Use of the Modified Borah Equation to Assess Nutritional Status of Pediatric PD Patients).

Rationale Although nutritional status depends on many nondialysis-related factors, appetite suppression, nausea, and vomiting are major
clinical features of inadequate dialysis. Therefore, nutritional status is also an important measure of PD adequacy. Of the available measures of nutrition, PNA is recommended because it provides an estimate of protein catabolic rate (PCR) and other protein losses. The SGA is recommended because it is a clinical assessment of patient nutritional status and is strongly associated with patient survival. Both measures are discussed in detail below.

Protein Equivalent of Nitrogen Appearance (PNA). PNA is a useful tool for monitoring the absolute level of changes in dietary protein intake. Nitrogen intake is almost entirely (95%) in the form of protein. Therefore, total nitrogen excretion in stable humans multiplied by 6.25 (there are approximately 6.25 grams of protein per gram of nitrogen) should be a good estimate of protein intake. This relationship does not hold true for: individuals in a state of catabolism where body cell mass may be contributing to nitrogen excretion; conditions of anabolism where the opposite occurs; and inconsistencies in absolute or time-averaged blood concentrations of BUN or creatinine.

In normal humans and in dialysis patients in nitrogen balance who have no direct protein losses in urine, dialysate, or feces, the total daily excretion of nitrogen in urine, dialysate, feces, breath, and skin losses is in the form of low molecular weight nitrogenous metabolites (such as urea, creatinine, urate, amino acids, ammonia, and peptides).

The excretion of nitrogen as low molecular weight metabolites multiplied by 6.25 approximates the amount of nitrogen in ingested protein. This calculation has been termed the protein catabolic rate (PCR). PCR actually represents the net amount of protein catabolism exceeding protein synthesis required to generate an amount of nitrogen equal to that excreted. The nitrogen in ingested protein enters the body nitrogen pools; nitrogen excreted in urine, feces, breath, skin, and dialysate represents the metabolism of a variety of body substances in these pools, such as creatine and purine, in addition to body proteins. Thus, although PCR is a reasonable estimate of protein intake, not all excreted nitrogen comes directly from protein. The protein catabolic equivalent of nitrogen excretion is actually a net catabolic equivalent, rather than an absolute. It relates directly to the contribution of protein catabolism to uremic toxicity.

In patients on hemodialysis, nitrogen balance studies have been performed to estimate total nitrogen output or appearance (skin losses were estimated and breath losses were ignored). The relationship of urea nitrogen appearance to total nitrogen output was assumed to be fixed and a formula was developed, known as the Borah equation, to calculate the PCR directly from urea nitrogen appearance:

\[ \text{PCR (g/d)} = 6.49 \ast \text{UNA} + 0.294 \ast \text{V} \]  

where UNA represents the net production or appearance of urea nitrogen in body fluids (any increase in body fluid nitrogen concentration times body water volume) and all measurable outputs in g/d; V is the volume of distribution of urea in liters. In hemodialysis patients with no direct protein losses in dialysate or urine, this PCR also represents an estimate of dietary protein intake and is the protein equivalent of total nitrogen appearance (PNA).

In dialysis patients with substantial urinary or dialytic protein losses (>0.1 g/kg), the direct protein losses must be added to the PCR to yield the true PNA as an estimate of dietary protein intake.

\[ \text{PNA = PCR + protein losses} \]  

Nitrogen balance studies have also been performed in PD patients; the measured total nitrogen output (appearance) included estimates of skin and fecal nitrogen losses plus measurement of all nitrogen (including protein nitrogen) in dialysate and urine. The average daily dialysate protein loss in the CAPD patients was 7.3 g. Urine protein losses were <1 g/24 hrs. A formula for the calculation of PNA from UNA was developed:

\[ \text{PNA (g/d)} = 10.76 \ast (0.69 \ast \text{UNA} + 1.46) \]  

This calculation incorporates the average dialysate protein loss of 7.3 g/d.

Calculations of PNA in PD patients with Equations 2 and 3 have been shown to yield nearly identical results. Also, subtracting protein losses in dialysate from Equation 3 yields values nearly identical to the PCR calculated by Equation 1, as developed in HD patients.
If daily peritoneal dialysate protein losses exceed 15 g, PNA calculated from Equation 2 will exceed PNA calculated from Equation 3 by approximately 0.1 g/kg standard body weight.\(^6\) High transporters lose more protein into effluent dialysate than other PD patients.\(^5,7\) Therefore, in high transport patients it is best to measure protein losses in dialysate directly, and if dialysate protein losses exceed 15 g/d (found in <10% of peritonitis-free patients), calculate PNA from Equation 2.\(^5\) For patients who lose large amounts of protein from any nonperitoneal source (e.g., nephrotic syndrome), Equation 2 should be used.

Equations 2 and 3 have been validated with nitrogen balance studies only in CAPO and not in other therapies such as NIPD.\(^1,5\) However, since CAPD and HD patients have similar PCR values (PNA – protein losses) at any given UNA, the intermittent nature of NIPD would seem unlikely to alter the relationships. Furthermore, the daily protein losses on NIPD are similar to those of CAPD.\(^5\) A recent study of seven patients suggests that the PNA formula of Bergstrom et al.\(^9\) may be superior.\(^10\) However, this must be corroborated in a study with a larger number of patients. Bergstrom supports the use of Equation 3.\(^1\)

In summary, the most accurate determination of PNA in patients undergoing PO uses Equation 2, but this requires measurement of UNA and dialysate protein losses. Equation 3 is a suitable substitute and requires only measurement of UNA. However, if dialysate protein exceeds 15 g/d (many high transporters may fall into this category) and in all pediatric patients, Equation 2 is preferred.

Methods of normalizing PNA are still under debate. The Work Group recommends normalization by standard weight, which has been applied extensively. Standard weight is equal to V/0.58.\(^11\) PNA normalized by either standard weight or actual weight tends to be high in malnourished, underweight PD subjects.\(^12\) Normalization of PNA to fat-free, edema-free body mass provides appropriately low nPNA values in underweight individuals.\(^13\) The Work Group recommends that fat-free, edema-free body mass, estimated from creatinine kinetics (see Section II: Measures of PD Dose) should be used, in addition to standard weight, to normalize PNA in underweight PD subjects, defined by Table X-1 in Appendix E. Normalizing is important for patient-to-patient comparisons and to follow PNA measurements serially in an individual patient whose weight may change. If weight is stable, normalization is less important in serial measurements for an individual patient (see Guideline 14 for PNA discussion on pediatric patients).

**Subjective Global Assessment (SGA).** The SGA is a valid estimate of nutritional status for patients treated with PD.\(^13\) Furthermore, it is associated with the probability of patient survival.\(^14\) The SGA was developed as a clinical estimate of pre-operative nutritional status.\(^15\) For two physicians, the inter-observer agreement was 72% greater than would have been predicted by chance alone. Validity was based on correlations with three measures of post-operative hospital morbidity (incidence of infection, use of antibiotics and length of stay).\(^15\) A detailed description of the SGA was provided by Detsky in 1987.\(^16\)

The SGA was originally developed as a clinical assessment of pre-operative nutritional status for patients prior to gastrointestinal surgery.\(^16\) When applied to CAPD patients,\(^13\) validity testing reduced the number of items to four (weight loss, anorexia, loss of subcutaneous tissue, and muscle mass). To increase the ability of the SGA to detect a change in nutritional status, the scoring scale was increased from a 3-point to a 7-point scale. During the development phase, the SGA was determined by physicians, research nurses, and nurse clinicians\(^16\) but was determined by dialysis nurses and dietitians when used in the CANUSA study.\(^14\)

The SGA, as modified for use in CAPD patients,\(^13\) uses a 7-point scale\(^14\) which any healthcare professional can apply following a short training period.

The four items used to assess nutritional status in CAPD patients are: weight change, anorexia, subcutaneous tissue, and muscle mass.

**Weight change** is addressed by the question, “What was the patient’s weight change over the past 6 months?” Ideally, this should be documented by the actual weights, but historical information from the patient is acceptable. A loss of >10% is severe, 5% to 10% is moderate while 5% is mild. This is rated subjectively on a scale from 1 to 7, where 1 or 2 is severe malnutrition, 3 to 5 is moderate to mild malnutrition and 6 or 7 is mild malnutrition to normal nutritional status. If the weight change
was intentional, the weight loss would be given less subjective weight while edema might obscure greater weight loss.

Anorexia is addressed by the question, "Has the patient’s dietary intake changed?" Supplemental questions determine whether a decrease in dietary intake is by prescription or due to decreased appetite. Nausea and vomiting are adverse factors for this item. Again, the interviewer will rate intake on the 7-point scale with higher scores indicative of better dietary intake, better appetite, and the absence of nausea and vomiting.

Subcutaneous tissue (fat and muscle wasting) can be examined in many areas. A very detailed and illustrated brochure is available from Baxter Healthcare (publication #BRU-008-312-2000). Although the history-taking format is more detailed than required, the description of how to determine muscle wasting and subcutaneous tissue is excellent.

Subcutaneous fat can be assessed by examining the fat pads directly below the eyes and by gently pinching the skin above the triceps and biceps. The fat pads should appear as a slight bulge in a normally nourished person but are "hollow" in a malnourished person. When the skin above the triceps and biceps is gently pinched, the thickness of the fold between the examiner’s fingers is indicative of the nutritional status. The examiner then scores the observations on a 7-point scale.

Muscle mass and wasting can be assessed by examining the temporalis muscle, the prominence of the clavicles, the contour of the shoulders (rounded indicates well-nourished; squared indicates malnutrition), visibility of the scapula, the visibility of the ribs, and interosseous muscle mass between the thumb and forefinger, and the quadriceps muscle mass. These are scored on a 7-point scale.

The four-item scores are then aggregated into a global score. The global score is not a simple arithmetic average of the four items. The examiner can apply different weights to the items. For example, if the physical examination items clearly indicate severe malnutrition, but the patient indicates only a moderate decrease in weight and a good appetite, the examiner might weight the physical examination items higher than the historical items.

In 23 CAPD patients, four items were statistically associated with the SGA: weight loss, anorexia, loss of subcutaneous tissue, and loss of muscle mass (muscle wasting). Evidence for validity was provided by the correlations with serum albumin concentration, bioelectrical impedance, anthropometric measurements and normalized protein catabolic rate.

Using the SGA as originally described, 59% of prevalent CAPD patients were well-nourished. Mild and severe malnutrition was reported in 33% and 8% of the patients, respectively. In 263 hemodialysis patients and 224 CAPD patients, the SGA covaried with low visceral (i.e., serum) protein concentrations, midarm muscle circumference (somatic protein mass), and body fat stores.

In the CANUSA study of peritoneal dialysis, weight loss, anorexia, loss of subcutaneous tissue, and loss of muscle mass (muscle wasting), as identified above as being statistically associated with the SGA, were used to generate the SGA for the CANUSA study. To make the scale more discriminative, the 3-point scale was expanded to a 7-point scale, with 1 and 2 corresponding to severe malnutrition, 3-5 corresponding to mild to moderate malnutrition and 6-7 corresponding to mild malnutrition to normal nutritional status. In a multivariate analysis, a higher SGA was associated with a lower relative risk of death. A one unit increase on the 7-point scale was associated with a 25% decline in the relative risk of death (relative risk 0.75). During the first 6 months of dialysis, the mean SGA increased 0.72 units. There was a statistically significant correlation between the increment in adequacy due to the addition of peritoneal clearance (Kt/V urea and CCr) to RRF. Over the next 12 months, there was a small decrease in SGA and this correlated with loss of RRF estimated by CCr, but not with Kt/V urea.

The SGA has not been validated as a means of nutritional assessment in the pediatric PD population. Thus, the SGA cannot be recommended for use in children.

APPENDIX F REFERENCES

2. Borah MF, Schoenfeld PY, Gotch FA, Sargent JA, Wolfson M, Humphreys MH: Nitrogen balance during inter-


Appendix G: Detailed Rationale For Guideline 15

GUIDELINE 15

Weekly Dose of CAPD (Evidence)

For CAPD, the delivered PD dose should be a total Kt/Vurea of at least 2.0 per week and a total creatinine clearance (Ccr) of at least 60 L/week/1.73 m².

Rationale: The evidence supporting this guideline is derived from theoretical constructs and cohort studies which use either univariate or multivariate statistical analyses.

The original description of CAPD suggested that an anephric 70-kg patient with a total body water of 42 L would remain in nitrogen balance at a daily dialysis prescription of 10 L given as five 2-L exchanges. Full equilibration of urea between plasma and dialysate and 2 L/d of net ultrafiltration were assumed. This would produce a daily urea clearance of 12 L or a weekly urea clearance of 84 L. For a patient with a total body water of 42 L, this corresponds to a weekly Kt/Vurea of 2.0. Others, using the concept of the Dialysis Index, suggested that a similar patient would require 13.5 L daily of equilibrated drained dialysate to maintain nitrogen balance, and this would produce a weekly Kt/Vurea of 2.25. The difference between these two projections is due to the higher target protein intake used in the latter calculation. Using the peak urea concentration hypothesis, a weekly Kt/Vurea of 2.0 is equivalent to a single pool hemodialysis Kt/Vurea of 1.3 for patients receiving thrice weekly dialysis. These theoretical constructs suggest that a weekly Kt/Vurea of 2.0 to 2.25 would be appropriate.

Validation of these theoretical constructs requires clinical study. A series of cohort studies addressed this issue. Initially, no relationship was found between urea clearance and patient
survival, but a re-analysis, using an anthropometric estimate for total body water, found that patients with a weekly $Kt/V_{urea} < 1.5$ had an increased risk of death compared to patients with a weekly $Kt/V_{urea} = 1.5$. In another study, a mean weekly $Kt/V_{urea} > 1.89$ was associated with a decreased risk of death compared to patients with less dialysis, while yet another reported that patients surviving for a 12-month follow-up had a mean weekly $Kt/V_{urea}$ of 2.0 compared to a mean of 1.7 among those who did not survive for 12 months. A Belgian group reported that 16 patients surviving 5 years on CAPD had a mean weekly $Kt/V_{urea}$ of 2.0. These studies all used univariate analysis and therefore did not simultaneously evaluate the association between other important variables (eg, age, diabetes, cardiovascular disease) and patient survival.

Several studies have used multivariate statistical analysis to evaluate the association between adequacy of PD and survival while controlling for other variables. In one such study a lower serum albumin concentration, increased age, greater time on dialysis, and lower weekly $Kt/V_{urea}$ were associated with a decreased probability of patient survival. A French group reported that patients with a weekly $Kt/V_{urea} > 1.7$ and a weekly $C_{Cr}$ of $> 50 \, \text{L}/1.73 \, \text{m}^2$ at initiation of dialysis had better survival than those with lower values at initiation. However, these investigators did not evaluate the effect of changes in adequacy over time due to loss of RRF, nor did they attempt to evaluate any association of higher weekly $Kt/V_{urea}$ or $C_{Cr}$ with survival. An Italian group evaluated the association between estimates of adequacy and patient survival in a cohort of 68 prevalent continuous PD patients followed over 3 years. A mean weekly $Kt/V_{urea}$ of 1.96 was associated with better survival than lower values. No further benefit was observed with a $Kt/V_{urea}$ higher than 1.96. Among these patients, a weekly $Kt/V_{urea}$ of 1.96 corresponded to a weekly $C_{Cr}$ of 58 liters/1.73 m$^2$.

The CANUSA study evaluated the association between adequacy of PD and patient survival, technique survival, and hospitalization among 680 incident patients (new to starting PD) treated with continuous PD. A decrease of 0.1 in weekly $Kt/V_{urea}$ was associated with a 5% increase in the relative risk of death, and a decrease of 5 L/1.73 m$^2$/wk in $C_{Cr}$ was associated with a 7% increase in the risk of death. The risk of technique failure increased with decreased creatinine clearance, but was not associated with $Kt/V_{urea}$. Hospitalization increased with decreased $C_{Cr}$. Using data derived from the multivariate analysis, the predicted 2-year survival associated with a constant weekly $Kt/V_{urea}$ of 2.1 was 78%. The corresponding weekly $C_{Cr}$ was 70 L/1.73 m$^2$.

Thus, there is both a theoretical rationale and convincing evidence supporting an association between greater clearance of urea and creatinine and better patient survival. There is also evidence supporting an association between greater $C_{Cr}$ to longer technique survival and less hospitalization. In summary, theoretical constructs suggest that the minimum weekly $Kt/V_{urea}$ should be 2.0. Cohort studies using univariate statistical analysis support this "target." The CANUSA study predicts, among North American patients, a 78% two-year survival with a weekly $Kt/V_{urea}$ of 2.1.

There are no theoretical data to support a specific $C_{Cr}$ target. The $C_{Cr}$ which corresponds to a weekly $Kt/V_{urea}$ of 2.1 in the CANUSA study was 70 L/1.73 m$^2$/wk. The Italian group found that a weekly $Kt/V_{urea}$ of 1.96 corresponded to a weekly $C_{Cr}$ of 58 L. The CANUSA study involved incident patients with significant RRF, while the Italian study evaluated prevalent patients with much less RRF. The target of 60 L/1.73 m$^2$/wk was selected by the Work Group because it is more relevant to patients with diminished renal function.

There are few data to address the issue of adequate compared to optimal dialysis. The latter is defined in part as the dialysis dose above which the incremental clinical benefit is not justified by the social cost to the patient or the financial cost to society. Whether or not increased weekly $Kt/V_{urea}$ greater than 2.0 will be associated with improved clinical outcomes requires further study.

The relative importance of RRF compared to peritoneal clearance and the relative importance of urea compared to $C_{Cr}$ are important and interrelated issues. The convention has been to consider RRF and peritoneal clearance to be equivalent and therefore additive. Some believe that renal clearance is more important, but in the absence of data establishing the magnitude of that difference, the assumption of equivalence was adopted by the Work Group. $C_{Cr}$ appeared more
important than urea clearance in the CANUSA study. The former was associated with patient survival, technique survival, and hospitalization, while the latter was associated only with patient survival. One potential explanation for this finding is that $C_{Cr}$ is more strongly associated with better RRF than was $Kt/V_{urea}$. This explanation is based on the assumption that RRF is better than peritoneal clearance, an opinion not yet supported by evidence.

Until evidence to the contrary is available, the Work Group recommends that renal and peritoneal clearances be considered equivalent. If there is discordance between achieving the target $Kt/V_{urea}$ and $C_{Cr}$, the $Kt/V_{urea}$ should be the immediate determinant of adequacy since it reflects protein catabolism. However, the reason for the discrepancy should be sought and the patient monitored closely for clinical signs of underdialysis.

A special case is the underweight patient, defined in Table II-3, Appendix E. Successful efforts to restore weight to a normal level in such a patient will result in a rising $V_{r}$ and consequently in a proportionally declining $K_{pr}t/V_{urea}$. To provide a weekly $K_{pr}t/V_{urea}$ of 2.0 at the final increased weight, the weekly target $K_{pr}t/V_{urea}$ provided during the malnourished state must be greater than 2.0. The Work Group recommends that the target $K_{pr}t/V_{urea}$ should be raised in a malnourished CAPD patient to the level that would provide a weekly $K_{pr}t/V_{urea}$ of 2.0 for that patient if he or she was at normal weight. That level is calculated by multiplying the target of 2.0 for CAPD times the ratio of $V_{desired}/V_{actual}$. This is described in detail in Appendix E: Detailed Rationale for Guideline 9 and discussed in Guideline 17: PD Dose in Subpopulations. The same upward target adjustment would be made in $C_{Cr}$. The target $C_{Cr}$ should be increased by a factor of $BSA_{desired}/BSA_{actual}$.

Clinical judgment suggests that the target doses of PD for children should meet or exceed the adult standards. However, there are currently no definitive outcome data in pediatrics to suggest that any measure of dialysis adequacy is predictive of well-being, morbidity, or mortality. There are no data regarding the real protein needs of children, especially young children, on dialysis. It is the opinion of the Work Group that the nutritional requirements per kilogram of body weight are higher in children than in adults. Therefore, PD doses in children, and especially small infants who have very high protein intakes, may have to be higher than PD doses in adults.
Appendix H: Detailed Rationale for Guideline 19

GUIDELINE 19

Identify and Correct Patient-Related Failure to Achieve Prescribed PD Dose (Opinion)

Potential patient-related causes of failure to achieve prescribed peritoneal dialysis dose should be investigated and corrected. These include:

- Failure to comply with the prescription
- Lack of understanding of the importance of adherence to the full prescription
- Sampling and collection errors

Rationale

It is the opinion of the Work Group that to increase the likelihood of achieving a prescribed dose of PD, it is necessary to elucidate the patient-related causes of failure to achieve a prescribed dose. Selection of inappropriate candidates for PD may result in failure to achieve a prescribed dose due to medical, technical, and/or psycho-social reasons. The issue of medically appropriate patient selection is dealt with at length in Section VIII, Suitable Patients for PD. In addition to the medical reasons for selecting patients for PD or HD discussed in Section VIII, patient compliance is of paramount importance and should be explored.

Failure to Comply With the Prescription. Patients may decrease the delivered dose of PD in several ways. Some of the ways are listed below:

- Skipping exchanges
- Shortened exchange times
- Dialysate dumping: too much flushing resulting in too little fill
- Delayed dumping: This is achieved by partially draining before the dwell is completed.
- Reduction of total cycle time
- Unscheduled dry days on CCPD

A validated method to measure patient compliance is not currently available. Methods proposed for evaluating compliance include monitoring for variations in creatinine output in dialysate and urine as detailed in Guideline 7: PD Dose Troubleshooting. Evidence for this recommendation is currently not available. The recommendation represents, therefore, the opinion of the Work Group members.

In the absence of a validated method to measure patient compliance, its prevalence in the PD population is not known. Preliminary data from the USRDS DMMS Wave II project show that 487 CAPD patients self-report full compliance with 82.8% of their exchanges. One exchange per week is missed by 11.5% of patients and 2 to 3 exchanges per week are missed by 4.5% of patients, all self-reported. Other estimates vary between 5% and 38%. Thus, noncompliance is a major cause of a delivered PD dose being less than the desired dose and is potentially preventable.

Lack of understanding of the importance of adherence to the full prescription. Medical literature about conditions associated with noncompliance in PD is inadequate. The Work Group reviewed published information on compliance in hemodialysis and in drug treatment for chronic illness. The Work Group believes that some of the conclusions in this literature may be applicable to PD. An important conclusion of the studies on drug compliance is that lack of education regarding the importance of adherence to the full prescription partially contributes to compliance failure. Compliance with the drug prescription improves when the patient is convinced that the diagnosis is accurate, the reasons for the prescribed treatment are correct, and the prescribed treatment is beneficial. Some contend that patients on dialysis are more likely to follow the prescribed treatment if they can be convinced that adherence to the prescription is in their own interest. By not understanding the significance, importance, value, or relevance of the collections or the exchanges, noncompliance could occur without patient concern. Therefore, proper education about the treatment may increase compliance in many PD patients. Patients should be educated that dialysis prescription may change over time (different modality and/or increase in the number or volume of exchanges) due to loss of residual renal function (see Guideline 6: Assessing Residual Renal Function). The method of education should emphasize the expected positive results (improved survival, well-being) of adherence to the PD prescription, rather than the negative outcomes (morbidity, mortality) of non-adherence, to prevent the development of excessive anxiety, which has adverse effects on compliance.

Patient education should be continuous throughout the course of PD. Patients should be told the results of the repeated clearance measurements and should be aware of the target values for Kt/V and Cr, and of the clinical significance...
of these clearances. Prevention of non-compliance should include monitoring the patient’s psychological status. In studies on compliance to drugs, certain psychiatric conditions, such as hostility toward authority, depression and memory impairment, financial problems, impaired mobility, and language or ethnic barriers, have been associated with poor compliance. In addition, complexity of the prescription and chronicity of the treatment increased noncompliance. In the case of prolonged treatment, repetition of the teaching at 6-month intervals improved compliance. In studies on compliance in HD, male gender and young age were predictors of poor compliance with different aspects of HD prescription. Preliminary information suggests that a general negative attitude of the patients predicts noncompliance in PD. Finally, drug compliance improves with better education of the providers about compliance issues. The Work Group thinks that all of these issues are relevant to PD. The psychological profile which is predictive of noncompliance and the best method of characterizing this profile should be a subject for research in the future. For the present, the Work Group’s opinion is that monitoring of patients’ psychological status should be aimed at detecting conditions associated with increased risk of noncompliance, and particularly at detecting a negative patient attitude towards PD. Teaching patients about their PD prescription should be repeated at intervals of 6 months or less.

Sampling and collection errors. Sampling and collection errors committed by patients during the clearance study preclude accurate measurement of clearance. Such errors include:

- Batch method: the patient may not recognize the importance of accidentally spilled dialysate.
- Aliquot method: Inaccurate weighing of drain volumes which may be the result of inaccurate scales or misreading. Disproportionate filling of the syringe with dialysate.
- Errors in weighing bags for variable fill volumes: for example, when (due to cost issues) 3-liter bags are being used and only 2.5 liters are exchanged.
- Incomplete urine collections for the RRF determination.

Many of these errors can be prevented by careful patient instruction about the details and significance of the clearance procedure.

APPENDIX H REFERENCES

2. Keen M, Lipps B, Gotch F: The measured creatinine generation rate in CAPD suggests only 78% of prescribed dialysis is delivered. Adv Perit Dial 9:73-75, 1993

IMPORTANT NOTICE

Articles which were selected for structured review will be listed separately in a specially designed NKF-DOQI Program Caddy which will contain Executive Summaries in addition to the Clinical Practice Guidelines. The articles and guidelines will also be available on the World Wide Web on NKF’s cyberNephrology web site at www.kidney.org.
XI. Biographical Sketches of the NKF-DOQI Peritoneal Dialysis Adequacy Work Group Members

The following are brief sketches that describe the professional training and experience, as well as principal business affiliations of the Work Group members. All Work Group members completed a disclosure statement certifying that any potential conflict of interest would not influence their judgment or actions concerning the NKF-DOQI.

Thomas A. Golper, MD, FACP (Work Group Chair), is Medical Director, Director of Clinical Research, and Professor of Medicine for the Division of Nephrology at the University of Arkansas for Medical Sciences. Dr. Golper has been working to improve patient outcomes for the past 23 years. He serves as a CQI advisor to industry and has participated in guideline development for the management of peritonitis since 1986. Dr. Golper has published 136 articles and currently directs research on topics such as peritoneal fluid-drug interactions, peritonitis in PD, and atherosclerotic/thrombotic risk factors in ESRD. Active in many professional societies, at present, he serves on the Board of Directors for the Renal Physicians Association as well as the American Association of Kidney Patients. Dr. Golper reported an affiliation with the University of Arkansas for Medical Sciences Kidney Center and QualChoice, a managed care organization.

David Churchill, MDCM, FRCPC, FRCP (Edin) (Work Group Vice-Chair), is Professor of Medicine and Director, Division of Nephrology, at McMaster University in Hamilton, Ontario, Canada. His funded clinical research activities have included urolithiasis research, polycystic kidney disease, quality of life in ESRD, economic analysis, erythropoietin in ESRD, and adequacy of peritoneal dialysis. He is co-principal investigator of the CANUSA on the adequacy of peritoneal dialysis. He has over 100 peer-reviewed publications. Dr. Churchill has served on the scientific review panels for the NIH, Kidney Foundation of Canada, and the Ontario Ministry of Health, as well as on advisory committees for erythropoietin and dialysis services in Ontario. He has a cross-appointment in the Department of Clinical Epidemiology and Biostatistics, with a focus on evidence-based medicine. Dr. Churchill reported an affiliation with Althin Med Inc.

John Burkart, MD, is Associate Professor of Internal Medicine/Nephrology and Director of Outpatient Dialysis Services (CAPD and HD) for the Section of Nephrology at Wake Forest University/Bowman Gray School of Medicine in Winston-Salem, North Carolina. Dr. Burkart has been an active clinical nephrologist for 11 years and manages a large nephrology and dialysis practice. He currently cares for 70 end-stage renal disease patients and participates in a group practice serving more than 250 dialysis patients. He serves as a consultant to industry and has actively participated in clinical trials and educational workshops. Dr. Burkart has written or co-authored more than 90 publications, and is currently involved in research related to adequacy of peritoneal dialysis, improving outcomes on peritoneal dialysis, and alternative osmotic agents for peritoneal fluid. He is a co-investigator in the Hemodialysis Study and serves on the editorial boards of *Peritoneal Dialysis International* and *Advancement of Renal Replacement Therapy*. Dr. Burkart reported an affiliation with Baxter Healthcare.

Catherine Firanek, RN, CNN, MBA, is Peritoneal Dialysis Nurse Manager at Circle Medical Management Dialysis Services in Chicago, Illinois. Ms. Firanek has been involved with peritoneal dialysis program establishment, patient care, and education for the past 14 years. She is an advocate of patients and self-care dialysis through patient education programs. Nationally, Ms. Firanek has been actively involved in nursing educational program planning for the National Kidney Foundation, and serves on a nursing advisory board for industry for creation of nursing tools for CQI programs and nursing education. Ms. Firanek has published several articles in the areas of peritoneal dialysis in an urban population, peritonitis incidence, and adequacy. She is the past Chairperson of the Illinois Council Of Nephrology Nurses and Technicians, a member of CNNT, ANNA, the National Kidney Foundation of Illinois Board of Directors, and the International Society of Peritoneal Dialysis Subcommittee on International Studies.
Denis Geary, MB, MRCP(UK), FRCPC, is Associate Professor, Department of Pediatrics, at the University of Toronto, and Chief of Nephrology at the Hospital for Sick Children in Toronto, Canada. Dr. Geary’s clinical training took place in the United States, United Kingdom, and Ireland. He has practiced as a staff physician in pediatric nephrology for 15 years. Dr. Geary’s publications involve the subjects of dialysis, and the growth and development of children with chronic renal failure.

Frank Gotch, MD, is Medical Director of Dialysis Treatment and Research at Davies Medical Center in San Francisco, California. Dr. Gotch has worked in clinical dialysis and dialysis research, particularly quantification of therapy, for 30 years. He chaired the NIH Hemodialyzer Evaluation Study Group in 1972 and the National NIH Conference on Adequacy of Hemodialysis in 1975. He served on the planning committee and as kinetic consultant to the National Cooperative Dialysis Study. Dr. Gotch presently serves on the steering committee of the current HEMO study and is Co-Principal Investigator of a cooperative study of randomized peritoneal dialysis prescriptions and clinical outcome. He has more than 100 publications and provides consultation in dialysis kinetics and dialysis systems development to industry. Dr. Gotch serves as a consultant to Fresenius Medical Care.

Linda W. Moore, RD, is currently Manager, Clinical Marketing for SangStat Medical Corporation, a company focused solely on transplantation, applying a disease-management approach for improving outcomes and decreasing the costs of therapy. She has worked in dialysis and transplantation for many years at the University of Tennessee–Memphis where she focused on clinical outcomes in developing research protocols and patient care protocols. Ms. Moore has published more than 40 articles on renal nutrition and metabolism and consults both for industry and private, not-for-profit organizations. She has served on many local and national committees committed to defining and improving patient outcomes. Ms. Moore is currently Chair-Elect of the Council on Renal Nutrition for the National Kidney Foundation and also serves as a workgroup member of the ESRD Core Indicators for the Health Care Financing Administration.

Karl Nolph, MD, FACP, FRCPS, is Professor of Medicine and Director of the Nephrology Division at the University of Missouri–Columbia. Dr. Nolph has served as President of the American Society for Artificial Internal Organs and as President of the International Society for Peritoneal Dialysis. He has written more than 500 scientific publications and has edited four major textbooks on peritoneal dialysis. Dr. Nolph was Director of the Clinical Coordinating Center of the National Institutes of Health USA CAPD Registry from 1981-1988. He has been the program chairman of the Annual Peritoneal Dialysis Conference for 17 years. Dr. Nolph serves as a consultant to Baxter Healthcare, Inc., and is affiliated with Dialysis Clinics, Inc.

Neil Powe, MD, MPH, MBA, FACP, is Associate Professor of Medicine at the Johns Hopkins University School of Medicine, with joint appointments in the Departments of Epidemiology and Health Policy & Management at the Johns Hopkins University School of Hygiene and Public Health. Dr. Powe is Director of the Dialysis Care Patient Outcomes Research Team at Johns Hopkins University, a national study of patient outcomes in peritoneal dialysis and hemodialysis. He has trained in internal medicine, epidemiology and health services research. He has extensive experience in developing and measuring outcomes in dialysis patients using data from prospective studies, the USRDS, Medicare records, and patient surveys. Dr. Powe has published more than 70 articles on such topics as the effectiveness of recombinant human erythropoietin therapy in dialysis patients, treatment choices of peritoneal dialysis versus hemodialysis patients, vascular access management, and cost-effectiveness analysis of alternative treatments for ESRD. Dr. Powe was a member of the Institute of Medicine Committee on Measuring, Managing and Improving Quality of Care in the ESRD Treatment Setting. He has testified before the U.S. Congress on the role of patient outcomes research in improving the quality of care in the Medicare ESRD program.

Harmeet Singh, MD, is currently in private practice with Western Nephrology and Metabolic Bone Disease P.C., in Denver, Colorado. He finished a clinical and research fellowship in nephrology at the University of Colorado Health Sciences Center and was on the clinical faculty in the nephrology division at the University of Arkansas for Medical Sciences prior to his cur-
rent appointment. In his limited academic tenure, he has written on a variety of clinical nephrology topics, including sodium and potassium disorders and contrast nephrotoxicity.

**Brendan Teehan, MD, FACP**, is Clinical Professor of Medicine at Jefferson Medical College in Philadelphia and has been Chief of the Division of Nephrology at Lankenau Hospital/Lankenau Medical Research Center from 1992-1996. He has served as principal investigator at the NIH-sponsored National Cooperative Dialysis Study and as principal investigator for the ongoing Morbidity and Mortality in Hemodialysis Study. Dr. Teehan was also the Principal Investigator at the Lankenau site for the CANUSA Adequacy of Peritoneal Dialysis Study. He has written numerous articles and chapters dealing with kinetic modeling, nutrition, and adequacy of peritoneal dialysis. Dr. Teehan is the Past President of the American Society for Artificial Internal Organs and is presently a member of the finance committee of the Internal Society of Peritoneal Dialysis. He reported an affiliation with R.C.G.I. and Main Line Suburban Dialysis Centers.

**Antonios Tzamaloukas, MD, FACP**, is Acting Chief of the Renal Section of the Albuquerque VA Medical Center and Professor of Medicine at the University of New Mexico School of Medicine. Dr. Tzamaloukas has been practicing and teaching clinical nephrology for 21 years and has been responsible for the peritoneal dialysis program of the Albuquerque VA for a number of years. He has more than 170 scientific publications. His current research is directed toward issues related to adequacy of peritoneal dialysis. He is active in many professional societies.

**Bradley Warady, MD**, is Director of Dialysis and Transplantation and Chief of Nephrology at The Children’s Mercy Hospital, and Professor of Pediatrics at the University of Missouri–Kansas City School of Medicine. Dr. Warady’s clinical and research focus is end-stage renal disease, with particular emphasis on peritoneal dialysis. He established the Pediatric Peritoneal Dialysis Study Consortium and currently directs research projects on a number of topics including: the impact of “Flush Before Fill” on peritonitis in patients receiving automated peritoneal dialysis; long-term function of the peritoneal membrane; and growth hormone usage in pediatric dialysis patients. Dr. Warady has published more than 100 articles and currently serves on the executive committee of the Pediatric Nephrology and Urology Council of the National Kidney Foundation and the scientific advisory committee of the North American Pediatric Renal Transplant Cooperative Study. He reported receiving research grant funding from Baxter Healthcare Inc.