A Practical Approach to Nutrition, Protein-Energy Wasting, Sarcopenia, and Cachexia in Patients with Chronic Kidney Disease

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Protein-energy wasting · Cachexia · Sarcopenia · KALANTAR score · Chronic kidney disease · End-stage renal disease

Abstract
Objectives of Review: Protein-energy wasting (PEW) is a state of disordered catabolism resulting from metabolic and nutritional derangements in chronic disease states. Patients with chronic kidney disease (CKD), and end-stage renal disease (ESRD) in particular, have muscle wasting, sarcopenia, and cachexia that contribute to frailty and morbidity. Moreover, reverse epidemiology findings have strongly linked PEW with mortality in CKD and ESRD. Updated Findings: The malnutrition-inflammation score (KALANTAR Score) provides a useful tool to predict nutritional risk. A stronger focus on renal nutrition in renal patients is needed to attenuate cachexia and muscle loss. Malnutrition is a far greater threat in patients with renal disease than obesity, which means dietary counseling needs to be tailored to reflect this observation. The need to achieve optimal caloric intake is compounded by the need to limit excess protein intake in CKD, resulting in the need for energy supplementation to avoid PEW. Preventing PEW is the most pressing clinical concern in CKD/ESRD. Other nutritional issues to reckon in renal disease include the need to normalize serum bicarbonate to manage acidosis, uric acid control, and phosphorous control in CKD and ESRD. Exercise maybe beneficial, but further work is needed to prove a conclusive benefit via a randomized trial. Summary: PEW prevention is an integral part of renal nutrition and is of paramount importance given the obesity paradox. Integrative approaches by physicians and dieticians are needed to take a holistic view of a patient’s diet beyond just control of particular laboratory parameters.

Introduction

Protein-energy wasting (PEW) was historically first introduced in 2007 by the International Society of Renal Nutrition and Metabolism [1, 2]. The definition codified was that PEW is a state of nutritional and metabolic de-rangements in patients with chronic kidney disease (CKD) and end-stage renal disease (ESRD) characterized by simultaneous loss of systemic body protein and energy stores. The reason this occurs is related to the hypercatabolic state induced by uremia, anorexia due to poor appetite, inflammation from systemic conditions (Diabetes), and auto immune conditions that generally lead to
CKD and ESRD [2]. The evidence tends to suggest that this develops more in patients with CKD stage 3b (estimated glomerular filtration rate < 45 mL/min) as defined by the Kidney Disease Improving Global Outcomes staging of CKD. While PEW and malnutrition can result in a similar patient phenotype, they should be distinguished from one another given the CKD-related factors are unique to the development of PEW in patients with renal disease. See Table 1 for diagnostic criteria for PEW.

This physiologically leads to cachexia, fat loss, and lean muscle mass loss [1, 2]. Further studies have revealed an alarmingly high rate of PEW in CKD, ESRD, and even acute kidney injury (AKI) patients [3]. Carrero et al. [3] revealed that studies among AKI patients showed 60–82% PEW prevalence, CKD 3–5 patients showed 11–54% PEW prevalence, and studies of transplant patients showed 28–52% PEW prevalence. Figure 1 demonstrates the relationship between Kidney Disease Improving Global Outcome staging of CKD and contribution of relative factors to nutritional dysfunction.

Etiologically, the unique mechanisms of PEW in CKD were described in Carrero et al. [4] 2014 [5]. These include uremia-induced alterations resulting in increased energy expenditure, persistent inflammation, acidosis, and multiple endocrine disorders. The hyper catabolism of muscle and fat combined with decreased nutritional intake, inflammation, nutrient loss, poor physical activity, frailty, and the dialysis procedure per se further contribute to PEW. See Figure 2 for a more precise etiological map of the causal factors of PEW in CKD and ESRD patients. In dialysis patients, slowing down or preventing PEW remains a challenge. The cornerstone of therapy is nutritional supplementation, optimization of dialysis regimen, management of volume overload, and management of endocrine disorders [6]. Our objective in this review is to define the diagnosis of PEW and connect it with its consequences and relationship with reverse epidemiology, muscle wasting, frailty, and ultimately morbidity and mortality outcomes. We aim to discuss nutritional assessment tools in CKD and ESRD, the state of the art in dietary counseling in renal patients, as well as practical therapeutic interventions to limit the effects of PEW on patients suffering from renal disease.

### PEW and Clinical Outcomes in CKD

Studies have shown that patients who develop ESRD and started on hemodialysis treatment are prone to progressive decline in nutritional parameters [7]. Weight, body mass index, and fat mass tend to decrease, while proinflammatory cytokine markers such as C-reactive protein and interleukin 6 (IL-6) increase progressively over time [8]. As albumin is negatively affected by inflammation as an acute phase marker (inversely related to inflammation), it unsurprisingly tends to decrease with a heightened proinflammatory state [9]. This phenomenon has been observed to increase with time on dialysis (previously titled dialysis vintage). The above factors are directly proportional to weakness, frailty, decreased response to erythropoiesis-stimulating agents, and low quality of life scores by patient self-assessment. More om-

<table>
<thead>
<tr>
<th>Table 1. Clinical criteria for PEW diagnosis in CKD</th>
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<tbody>
<tr>
<td><strong>Serum chemistry</strong></td>
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<tr>
<td>Serum albumin &lt; 3.8 g/dL, transthyretin (pre-albumin) &lt; 30 mg/dL (in dialysis patients)</td>
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<tr>
<td><strong>BMI</strong></td>
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<tr>
<td>&lt; 23 kg/m²</td>
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<tr>
<td>5% weight loss over 3 months or 10% over 6 months</td>
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<tr>
<td>Total body fat percentage &lt; 10%</td>
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<td><strong>Muscle mass loss</strong></td>
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<tr>
<td>5% reduction in muscle mass over 3 months or 10%</td>
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<tr>
<td>over 6 months</td>
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<tr>
<td>Reduced mid arm muscle circumference by 10% in relation to 50th percentile</td>
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<tr>
<td>Lower than expected serum creatinine</td>
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<tr>
<td><strong>Deficient dietary intake</strong></td>
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<tr>
<td>&lt; 0.8 g/kg/day of protein for at least 2 months in dialysis patients</td>
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<tr>
<td>&lt; 25 kcal/kg/day for at least 2 months</td>
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PEW, protein-energy wasting; CKD, chronic kidney disease; BMI, body mass index.
Protein-Energy Wasting

The changes of PEW due to inflammation also correspond to increased hospitalization and mortality [7–11]. Hypoalbuminemia remains one of the most potent risk factors for excess mortality in hemodialysis patients [10, 11].

**Methods used in the Assessment of PEW**

Besides the diagnostic criteria for PEW (body mass index, muscle mass loss, and deficient dietary intake), see Table 1, dedicated scoring systems have been developed...
to aid in assessing nutritional status. The measures of nutritional risk in turn correlate with the aforementioned poor outcomes and attendant higher risks of morbidity, hospitalization, frailty, and mortality [12–14]. The 2 scoring systems available are the Subjective Global Assessment and the KALANTAR score also known as the Malnutrition Inflammation Score (MIS). Table 2 contains the data collected in the SGA and the MIS/KALANTAR score [15, 16]. Moreover, the data track outcomes in patients with AKI and renal transplants, not just in patients with CKD and ESRD [12–16]. The MIS/KALANTAR score has also been associated with elevated IL-6 levels, providing a possible link to the observed increase in morbidity and mortality risk [12–16]. Recently a “simple objective nutritional tool” was developed for assessing PEW and nutritional risk in geriatric patients [2]. The geriatric nutritional risk index utilizes only serum albumin and actual/ideal body ratio. It was validated in Asian ESRD patients and had a higher interobserver agreement in prevalent hemodialysis patients than the MIS [12, 17–20]. Despite agreement of both scores with markers of iron metabolism (transferrin), lipid metabolism (cholesterol), muscle metabolism (creatinine), and cytokines (IL-6), the MIS was superior with regard to predicting death and associations with changes in daily energy and protein intake.

Table 2. Data collected for the SGA and MIS

<table>
<thead>
<tr>
<th>SGA</th>
<th>MIS/KALANTAR score</th>
<th>GNRI</th>
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<tbody>
<tr>
<td>Nutrient intake, weight change, symptoms (abdominal pain, dental pain, anorexia, early satiety, nausea, vomiting, constipation, dysphagia, diarrhea). Functional capacity, metabolic requirements, physical exam findings. These go into calculating final SGA rating</td>
<td>Change in dry weight, dietary intake, gastrointestinal symptoms, functional capacity, comorbidity, change in fat stores, signs of muscle wasting, BMI (kg/m²), serum albumin (g/L), serum total iron-binding capacity (mg/dL). Scored 0–3 (0 not present, 3 is severe)</td>
<td>Serum albumin (g/L), actual-to-ideal body weight ratio</td>
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</table>

SGA, subjective global assessment; MIS, malnutrition inflammation score; GNRI, geriatric nutritional risk index; BMI, body mass index.

The Reverse Epidemiology of Obesity and Hyperlipidemia in CKD

A key tenant to understanding the pathophysiology of increased mortality in patients with renal disease is the observation of the obesity paradox. The finding that obesity is protective in CKD and ESRD was unexpected given the deleterious role it played in classical epidemiological cohorts. Consistently, it has been noted that regardless of sex, age, and the severity of obesity, obesity has an unexpected protective effect in CKD and ESRD patients. While surprising at first glance, the effects of reduced renal clearance of waste and solutes, disordered IL signaling, dysregulation of endocrine pathways, and electrolyte homeostasis may well ultimately explain these observations. The finding that hyperlipidemia and obesity are protective again bears witness to the fact that the usual epidemiological rules do not apply in renal patients [21, 22]. Rather what is at play is the powerful effect of malnutrition and PEW as a primary determinant of patient morbidity and mortality in CKD and transplant listed ESRD patients [23, 24]. It is also quite possible that disorders of electrolyte homeostasis, particularly hyperphosphatemia and elevated fibroblast growth factor 23, may have more significant roles in cardiovascular disease in CKD and ESRD than traditional hyperlipidemia parameters [25].

The strength of signal obtained from the protective effect of high BMI was stronger in younger incident US hemodialysis patients (<65 years old) [26], and this was different than a U-shaped association reported in a similar population in the Netherlands [27]. There did not seem to be any apparent protective effect in incident dialysis patients older than 65 years old [28]. Differences in follow-up have made direct comparisons of the aforementioned studies between older and younger incident dialysis patients difficult [27, 28]. Further, cohorts enriched for peritoneal dialysis patients tended to show less consistent reverse epidemiology associations [23].

There appears to be a shift in the epidemiological risk factors for morbidity and mortality as patients progress through the stages of CKD and into ESRD. Lu et al. [28] demonstrated that BMI had a U-shaped association with mortality in patients with stage III CKD. The best out-
comes were in mildly overweight individuals with poor outcomes at the extremes of obesity and cachexia. As patients transition to stage IV CKD and ESRD the classic epidemiology risk factor of obesity disappeared and high BMI became protective [28]. Reverse epidemiology of obesity has also been observed in patients with normal renal function but ongoing proteinuria. In this case, the risk of obesity is modified, and it becomes less deleterious in patients with proteinuria [29].

**Frailty**

Frailty is defined by observations of weakness, slow gait, exhaustion, low tolerance for physical activity, unintentional weight loss, and muscle mass to the point of being <10 the percentile of age- and sex-matched population controls [2]. If 3 or more of the preceding are true, frailty can be diagnosed. It has a strong relationship with muscle wasting, PEW, and morbidity and mortality in CKD, maintenance dialysis, and even posttransplant [30–34]. The prevalence of frailty approaches 30% of incident hemodialysis patients [35], and this is a progressive change that has been noted to worsen as patients advance through the stages of CKD [36]. Worsening CKD stage and longer dialysis vintage also seemed to correlate with progressive loss of leg skeletal muscle mass as measured by computed tomography scans [37].

While research studies can take advantage of dual energy radiograph absorptiometry and magnetic resonance imaging to measure lean body mass, a 24-h urine creatinine excretion can be a surrogate for muscle mass. In particular, faster decline in urine creatinine can be an effective surrogate for muscle mass decline [38]. Serum creatinine can also provide an estimate for muscle mass procured diet and residual renal function are accounted for [39, 40]. Park et al. [41] verified that in 2 Korean Cohorts of hemodialysis patients and matched with US Caucasian and African American cohorts a higher BMI and serum creatinine was associated with better survival.

Conversely, it was noted that a decline in serum creatinine correlated with a higher risk of mortality than weight loss. This would make sense as declining serum creatinine would indicate a decrease of lean muscle tissue as described by Kalantar-Zadeh et al. [42, 43]. The result described suggests that mortality is linked with lean muscle mass rather than weight per se. This is likely since edema fluid is part of the component of a great deal of weight fluctuations, and more precise volume management with bioimpedance spectroscopy is not associated with worse outcomes but better outcomes [44]. It is notable that volume overload as estimated by bioimpedance spectroscopy and higher fat mass were also both associated with a higher risk of frailty [35], again this zeroes in on lean muscle mass rather than weight, BMI, or body fat composition as the strongest predictor of frailty risk and risk of poor outcomes in CKD, ESRD, and transplant patients.

**Dietary Therapy in Renal Disease**

In light of the above, dietary interventions to preserve lean muscle mass and provide adequate energy supplementation are paramount to the health of dialysis patients. Particularly hemodialysis patients who specifically show one or more surrogate measures concerning for low or decreased lean muscle mass. Dietary protein intake of 0.6–0.8 g/kg/day and dietary calorie intakes of 30–35 kcal/kg/day have been recommended for stage IIIb–V CKD. Despite the risks of negative protein balance, a low protein diet (LPD) is advocated in CKD given the risk of CKD progression with a higher protein diet [2, 6, 45, 46]. Wu et al. [45] suggested that supplementing a LPD improves adherence to this diet has beneficial effects on renal disease in addition to attenuating PEW.

Protein restriction can be used via a LPD with energy supplementation with fat and carbohydrates [45], but it is best used in younger CKD patients as the protective effects of a LPD are less pronounced in the elderly [47]. Thus, protein restriction to limit CKD prevention should be balanced against development of PEW, which would result in worse renal and mortality outcomes [2].

The recommendations for protein intake in ESRD patients reflect the concern for worsening PEW and need to prevent sarcopenia and cachexia that carry such devastating clinical and epidemiological risks. The typical recommendation is 1.2 g/kg/day protein intake in ESRD. This is since uremia is no longer a concern and can be cleared by efficient dialysis, but the appearance of declining muscle mass is an acute concern that augurs poorly for incident hemodialysis patients [2, 48–52].

**Hyperphosphatemia Management**

Elevated levels of phosphorous have been shown to increase cardiovascular and mortality risks in CKD and ESRD patients [2]. Non calcium-based binders reduce phosphorous and mitigate the risk of vascular calcifications, cardiovascular mortality [2, 53]. Despite the high
phosphorous content in animal protein-containing foodstuffs, high phosphorous negates any protective effect of protein and results in increased mortality risk [54–56].

The apparent mechanism is inflammation, and hyperphosphatemia induces dose-dependent oxidative stress independent of level of underlying renal function in animal models as shown by Yamada et al. [57]. In doing so hyperphosphatemia induces PEW, weight loss, hypalbuminemia, and sarcopenia reflected in reduced urinary creatinine clearance. Though further data in humans on hyperphosphatemia and PEW are needed, proteinuria is not advocated in ESRD patients, but rather phosphorous control to eliminate its unwanted inflammatory sequela. The effect of controlling protein to reduce serum phosphorous has produced results showing that hypoalbuminemia negates the benefits of controlling serum phosphorous. This suggests that dietary protein restriction in ESRD to control phosphorous can inadvertently increase the risk of mortality [58].

**Maintenance of Acid Base Homeostasis**

Acidemia is a commonly seen complication of CKD, ESRD, and in transplant patients with CKD. Acidosis can result in systemic symptoms, leukocytosis, disordered breathing, paradoxical hypothermia, and multisystem organ failure acutely [59]. Chronically, a less severe acidosis can result in a reduction of glomerular filtration rate independently of the cause of the acidosis or CKD [60]. An acidotic state can affect PEW by inducing muscle wasting via increases in glucocorticoid secretion [3]. Additionally, acidemia contributes to decreased muscle protein synthesis via decreased albumin synthesis caused by insulin resistance and other endocrine abnormalities [60, 61]. Muscle breakdown also increases via increased branched chain ketoacid dehydrogenase and ATP-dependent ubiquitin protease besides glucocorticoid increases [60, 61]. Other mechanisms increasing the PEW state include generalized inflammation due to acidemia and reductions in serum leptin leading to anorexia [60, 61].

Alkalization of serum-to-normal levels 23 meq/L but not over alkalization results in improved survival and decreased rate of progression of CKD [62–68]. Dialysis patients requiring high bicarbonate dialysate concentration especially if their serum HCO₃ remained < 23 meq/L were also found to have a high mortality risk in the Dialysis Outcomes and Practice Patterns Study. Despite possible confounding by indication as acidic patients tend to have higher dialysis bicarbonate prescriptions [69].

**Exercise**

Given the cascade of inflammatory and nutrient wasting in CKD and ESRD, renal patients often have decreased physical activity levels [70, 71]. The lack of physical activity exacerbates the pathophysiology of PEW directly. Strength training has been studied in dialysis patients and has been noted to induce skeletal muscle hypertrophy, increases muscle strength, and improves health-related quality of life in CKD and ESRD patients [72, 73]. Further studies are needed to confirm safety and efficacy in CKD, ESRD, and transplant patients. Findings from the transplant literature strongly suggest higher risks in transplanting frail patients [74], however, that suggest that exercise tolerance as a marker of frailty and exercise as a means to reverse frailty should be investigated.

**Nutrition on Dialysis**

Recent articles have demonstrated that the hemodialysis procedure is opportunities to improve renal nutrition in ESRD patients. Meals can be provided during hemodialysis as an effective strategy to help reduce the burden of PEW [3, 75], given the prevalence of PEW in ESRD. In patients unable to tolerate meals or with ongoing PEW despite oral meal supplementation, intradialytic and intraperitoneal parenteral nutrition can be used with efficacy [76]. Food frequency questionnaires are a simple descriptive tool to identify patients at risk for reduced nutrition intake given socioeconomic, psychological, renal/dialysis-related reasons, as well as systemic physiological causes [77].

Hemodialysis is often run with zero glucose baths in dialysate and induces a catabolic state similar to fasting with diminished gluconeogenesis [78]. This creates a situation for diabetic patients, or malnourished patients with PEW could be in significant risk of hypoglycemic events. Given known mortality risks of these events in diabetic patients, the concept of dialysis nutrition can have an immediate clinical benefit: reduction of hypoglycemia [78].

To reiterate the most important risk factor to control in dialysis patients is the PEW/inflammation complex, and the triad of nutrient loss, decreased nutritional intake, and breakdown of lean muscle mass [79–81]. This is in addition to the critical role dialysis nutritional counseling plays in control of hyperkalemia and hyperphosphatemia [75, 78]. We advocate for aggressive nutritional therapy and providing nutritional supplementation to di-
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This prevents losing a convenient opportunity to reverse the PEW pathology actively and control one of the most pressing risks for mortality in hemodialysis patients [46, 79–84].

Uric Acid Control

Though pertaining more to CKD than ESRD, it should be mentioned that uric acid control in CKD patients has been shown in multiple studies to reduce CKD progression [85]. Uric acid is an end product of protein metabolism; therefore, while a balanced approach to protein restriction can be used in early-stage CKD, pharmacological control may be necessary in later stage CKD given rising PEW risks. Please see Table 3 that summarizes the nutritional recommendations for patients with CKD and ESRD.

Conclusions

We have reviewed the pathophysiology of pertinent nutritional issues across the CKD, ESRD, and transplant CKD spectrum. New developments include nutritional benefits

Table 3. Summary of nutrition recommendations in CKD and ESRD

<table>
<thead>
<tr>
<th>Nutrition Category</th>
<th>CKD and ESRD Recommendations</th>
</tr>
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<tbody>
<tr>
<td>Protein intake</td>
<td>- CKD 0.6–0.8 g/kg/day protein restriction&lt;br&gt;- ESRD 1.2 g/kg/day increased protein intake due to PEW; don't restrict protein to control Po4</td>
</tr>
<tr>
<td>Weight, obesity, and lean muscle mass</td>
<td>- CKD III obesity behaves as classical risk factor&lt;br&gt;- CKD IV–V obesity's risks are reduced as malnutrition risks due to PEW increases, reverse epidemiology pattern becomes apparent in studies, particularly in older (&gt;65-year-old) patients&lt;br&gt;- ESRD PEW is a fearsome risk factor and obesity is protective against mortality in ESRD population&lt;br&gt;- Transplant patients similar to CKD (depends on estimated glomerular filtration rate)&lt;br&gt;- Proteinuria obesity is less deleterious risk factor in proteinuric patients than nonproteinuric patients&lt;br&gt;- Reduced lean muscle mass is most predictive of poor outcomes rather than BMI or weight per se. Anthropometric measures, urine creatinine clearance, and serum creatinine are surrogates of this&lt;br&gt;- Energy supplementation may reduce risks of PEW</td>
</tr>
<tr>
<td>Fluid overload</td>
<td>- CKD and ESRD maintenance of euvolemia associated with better outcomes</td>
</tr>
<tr>
<td>Frailty</td>
<td>- Worse outcomes at any stage of CKD, ESRD, peri and posttransplantation</td>
</tr>
<tr>
<td>Hyperphosphatemia</td>
<td>- CKD and ESRD use of non-calcium-based binders&lt;br&gt;- Goal Po4 5.5 and below but avoidance of low Po4</td>
</tr>
<tr>
<td>Alkalization</td>
<td>- CKD maintenance of serum bicarbonate at 23 meq/L without inducing supraphysiological bicarbonate levels&lt;br&gt;- ESRD goal is maintenance of normal serum bicarbonate, acidosis with greater dialysate bath associated with greater mortality risk</td>
</tr>
<tr>
<td>Exercise</td>
<td>- CKD and ESRD strength training is under study to reverse effects of PEW and frailty</td>
</tr>
<tr>
<td>Nutrition on dialysis</td>
<td>- ESRD supplementing with meals or intra dialytic parenteral nutrition in hemodialysis and intraperitoneal nutrition on PD when oral supplements are not effective&lt;br&gt;- ESRD eating on dialysis provides a convenient way to supplement nutrition and curb hypoglycemia</td>
</tr>
<tr>
<td>Uric acid</td>
<td>- CKD maintenance of normal serum uric acid which slows progression of CKD to ESRD</td>
</tr>
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</table>

CKD, chronic kidney disease; ESRD, end-stage renal disease; PEW, protein-energy wasting; Po4, phosphorous; BMI, body mass index.
of intradialytic meal replacement, scoring systems for PEW, and the emerging field of exercise therapy in CKD and ESRD to combat frailty and reverse the effects of PEW.

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Author Contributions

R.M.H. led writing of manuscript. L.G. contributed to text, editing, and compiling references. O.W. contributed to text, editing, and compiling references. C.M.R. contributed to editing text providing significant scientific input. K.K.-Z. led review of manuscript as senior author and expert on renal nutrition.

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References

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