Renal osteodystrophy is a significant complication in chronic kidney disease. This condition is referred to as mineral and bone disorders in chronic kidney disease, mainly because of its wider ranging impact, including an association with increased mortality and non–bone-related morbidity. Because most of the abnormalities that characterize mineral and bone disorders in chronic kidney disease (e.g., hyperphosphatemia, secondary hyperparathyroidism) are amenable to therapeutic interventions, this field has also been in the cross-hairs of many pharmaceutical companies. The advent of a number of new therapeutic options for mineral and bone disorders in chronic kidney disease has broadened our armamentarium but has also resulted in an intense marketing battle between pharmaceutical companies. The paucity of randomized, controlled trials in this field has allowed the various companies to promote unilaterally data that fit their needs and to attempt to discredit data that support their competitors’ products. Although this attitude is expected and regarded as acceptable in a consumer society, on a scientific level, it has resulted in a polarized and often confused audience: The practicing nephrologists. This article provides a historical overview of how the field of mineral and bone disorders in chronic kidney disease has evolved from a pharmaceutical standpoint, with a critical emphasis of the key moments that resulted in the current acrimonious climate. Also assessed is what the key unanswered questions are in this field, and practical solutions to the discussed issues are provided.

Evolution of Uremic Osteodystrophy
Renal osteodystrophy has long been recognized as a prominent feature of the uremic syndrome. Until recently, SHPT has been considered to be the central abnormality that leads to the bone disease associated with uremia. In the previous three decades, experts have argued about the relative importance of hyperphosphatemia, hypocalcemia, and vitamin D deficiency in the genesis of this SHPT (4–8). Thus, SHPT has been managed by the provision of supplemental calcium (in the form of oral supplementation and through dialysate calcium concentrations as high as 3.5 mEq/L) and active vitamin D in the form of synthetic calcitriol (in the United States) and alfacalcidol (in Europe) (9–11). Until the mid-1980s, hyperphosphatemia was managed by aluminum-containing phosphorus binders and dietary phosphorus restriction (8). Emerging data on the unto-
ward consequences of aluminum-based binders (dementia, refractory anemia, and osteomalacia [12,13]) made calcium-based medications the binders of choice by the early 1990s with the added purported benefit of further suppressing PTH production (14).

Paradigm Shift in the Management of Renal Osteodystrophy

Two major events in the mid-1990s led to a drastic paradigm shift in the foregoing management regimen. First, the advent of the so-called “low-turnover (adynamic) bone disease,” which was attributed to an oversuppression of PTH secretion by administration of calcitriol and hypercalcemia and/or “calcium-loading” from the high calcium intake and the supraphysiologic dialysate calcium concentration (15). Second, vascular calcification was found to be highly prevalent in patients with CKD and associated with poor survival (16,17); this, too, was found to be associated with higher serum calcium level and higher calcium intake in some (17–19) but not all studies (20–24). As a result of the consequent calcium-loading, dialysate calcium concentration was decreased from 3.5 to 2.5 mEq/L by the turn of the millennium, and a significant move toward using non–calcium-based binders has arisen (25).

These events coincided with several major observational studies showing that hyperphosphatemia and SHPT were associated with increased mortality (3,26), thus raising the stakes for therapeutic interventions that target the abnormalities of CKD-MBD. Although no clinical trials have examined the impact of lowering serum phosphorus or PTH on mortality, the confluence of observational data created a market that was ready for the emergence of new pharmaceutical products that could lower serum phosphorus and PTH without raising serum calcium, including sevelamer-hydrochloride (Renagel; Genzyme, Cambridge, MA) as the first Food and Drug Administration (FDA)-approved noncalcium nonaluminum binder (27) and the newer generations of active vitamin D with less productive (K/DOQI) guidelines on bone and mineral metabolism in contemporary dialysis patients; short of such studies, it is very difficult to make assertions about what “too much” or “too little” calcium intake means. The route of intake may be an-

**Figure 1.** Timeline of various treatments for mineral and bone disorders in chronic kidney disease (CKD-MBD).

Calculations of what the ideal amount of calcium intake might be is far from settled. Calcium mass balance studies are missing in contemporary dialysis patients; short of such studies, it is very difficult to make assertions about what “too much” or “too little” calcium intake means. The route of intake may be another important question, given that any proposed detriment of calcium intake may to some extent be counterbalanced by the benefits of phosphate lowering with oral intake but not with parenteral administration (such as with dialysate).

During this current phase of the battle, several competing pharmaceutical industry-sponsored studies have presented results that supported the arguments presented by the relevant manufacturer (32,36,38–40). The battle became ferocious with opposing mass flyers and letters to nephrologists by competing companies discrediting each other’s data, especially during the period after the public release of the main results of the DCOR trial (34). These activities instigated an increasing degree of confusion among both patients with CKD and nephrologists, who were sometimes not sure how to translate all of the data into clinical practice. Nevertheless, the new anticalcemic paradigm seemed to become the dominating trend, enjoying the blessings of the first Kidney Disease Outcomes Quality Initiative (K/DOQI) guidelines on bone and mineral metabolism in...
2003, and has had a major impact on our current practice pattern (25).

Confounded Battle Ground
Recently, the once focused and straightforward war against calcium has become confounded as several other pharmaceutical products have found their way into the CKD market. On the binder front, newer non–calcium-based binders have been approved or are awaiting approval, such as lanthanum carbonate (Fosrenol; Shire Pharmaceuticals, Hampshire, UK) (41) or other metal-based products (42) (Figure 1). This has opened up new battle fronts among the noncalcium binders themselves, shifting the debate toward issues such as the long-term safety of lanthanum carbonate (43) or the worsening acidemia seen with sevelamer hydrochloride (44). Efforts are under way to obtain FDA approval for sevelamer carbonate (Renvela; Genzyme). Additional binders with diverse features are expected to come out of the pipeline in the near future (Figure 1).

On the vitamin D front, the arrival of doxercalciferol (Hectorol) (45) to the US market has disrupted the previously focused message that the third-generation active vitamin D paricalcitol is the best of all. A somewhat artificial debate ensued about doxercalciferol’s belonging to an evolved generation of “active” vitamin D compounds surpassing calcitriol versus its being a mere prohormone below its rank; this has bred more confusion among the rank-and-file nephrologists and dietitians. The battle has become even more complicated as the calcium-sensing receptor agonist cinacalcet (Sensipar) has emerged as a promising alternative therapy for SHPT (30). Hence, the once unified anticalcium alliance seems to be suffering from infighting among its own members.

Several recent observational studies have shown that the administration of any dosage of any active vitamin D is associated with significantly better survival independent of the level of calcium, phosphorus, or PTH; these studies, in turn, might have de-emphasized the dangers of calcium loading (3,46,47). As part of the same trend, more recent epidemiologic analyses have reopened the question regarding the optimal range of serum calcium (3). Moreover, studies in non–dialysis-dependent patients with CKD have found that vascular calcification may be independent of high serum calcium level (20,22–24). Clearly, there is also evidence to implicate hypercalcemia and calcium intake in coronary calcification (17–19,37), and the previously mentioned studies merely complement the picture of a complex disease, yet it is still very common to see only one side of the story included in presentations dealing with the consequences of CKD-MBD, and it is easy to see how this would breed confusion or resentment in the audience.

What to Do Next?
That the field of CKD-MBD has become a complicated battleground in the past few years cannot be denied. That the pharmaceutical sector cherry-picks from emerging data also should not surprise nephrologists. The paucity of clinical trials in this field will unfortunately allow an ongoing emphasis on studies using observational designs or surrogate end points. Because none of these offers answers that are regarded as final, we will likely see a continuation of the current style of debate. To make matters more complicated, it is widely alleged that financial conflicts of interest have cast a shadow on expert recommendations; this will likely remain a lingering problem that could offer ways to try to discredit otherwise scientifically sound opinions or research findings (48). This has culminated in a recent series of open debates about the National Kidney Foundation’s (NKF) K/DOQI guidelines, where the independence from pharmaceutical influence of the process and even the experts involved in the process were questioned (48,49). The substantial involvement by pharmaceutical companies in the funding process makes it conceivable that similar criticism will continue to be leveled at future revised CKD-MBD guidelines from the NKF or the Kidney Disease: Improving Global Outcomes (KDIGO) (1,25).

So what is the best course of action for the management of CKD-MBD? There is no clear answer to this question. In a perfect world, we could be devoid of pharmaceutical bias by having independently sponsored RCT examining hard end points related to all of the potential interventions to establish the best treatment strategy. Unfortunately, this head-to-head comparison of the numerous management strategies is unlikely to happen in the near future. On the bright side, the various abnormalities characterizing CKD-MBD are treatable, and there is an ever-growing armamentarium available to us to facilitate these treatments (Figure 1). There are still a number of questions that will have to be answered in order for us to have a clear understanding of what the end points of therapy should be and what the best treatment regimens are (Table 1).

Finally, is there a viable solution to the problems that we mentioned? The first step toward a solution is to synthesize what we know and, even more important, to recognize the deficiencies in our knowledge (Table 1). The treatments that we prescribe are borne out of a desire to do what is best for our patients and are based on our knowledge of medicine. Many of the questions that we face when treating CKD-MBD will never have definitive answers (defined as proof through RCT), but this is not a reason for therapeutic nihilism. Knowledge can be meaningful even without RCT and should allow for sound decision-making with an eye toward the patient’s best interest. We need to use the limitations in our knowledge to limit the scope of our decision-making to the extent that we follow first and foremost the basic principle of “do no harm.” Thus, our opinion is that the solution to the listed problems is not a straightforward clinical practice guideline but rather the more universal principles of conscientiousness and studiousness. As far as studiousness is concerned, we recognize the everyday pressures that busy clinicians face, which makes in-depth studying of the many fields of nephrology on an ongoing basis a daunting task. Thus, from a practical standpoint, the NKF-sponsored K/DOQI and the upcoming KDIGO guidelines on CKD-MBD may probably be the best tools available as long as the practitioners understand the many areas of uncertainty and controversy. Discrediting these guidelines in their entirety because of their funding by the pharmaceutical industry may be unfair and inappropriate. However, the proponents of clinical
<table>
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<th>Question</th>
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<tr>
<td>1. What is the benefit of treating the abnormalities of CKD-MBD?</td>
<td>No RCT was ever done to show that interventions to lower PTH or phosphorus improve mortality, even though there is biological plausibility and observational studies show a strong association. Beneficial impact on various morbid conditions is established.</td>
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<td>2. What are the ideal calcium and phosphorus levels in patients with various levels of CKD?</td>
<td>Observational studies in dialysis patients are not unanimous on the ideal serum levels of calcium and phosphorus. No RCT has ever addressed this issue. Data on ideal levels in non–dialysis-dependent patients with CKD is scarce.</td>
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<td>3. Should diet modification be advocated as a treatment for hyperphosphatemia?</td>
<td>It remains unclear whether strict restriction of dietary phosphorus can be achieved without compromising dietary protein intake; protein restriction may worsen malnutrition and increase mortality.</td>
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<td>4. What is the best phosphate binder?</td>
<td>Head-to-head comparisons using relevant end points such as mortality or coronary calcification are available only for sevelamer hydrochloride and calcium-based binders. Cost considerations, comorbidity profiles, and adverse effects are often limiting factors and make the availability of multiple binders helpful in individualizing therapy.</td>
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<td>5. Should calcium intake be minimized?</td>
<td>This involves dietary, medication, and dialysate-associated calcium intake. Observational studies suggest an association between higher calcium intake and coronary calcification in patients on dialysis. Available RCT were not designed to study outcomes as a function of the amount of ingested calcium (but rather to compare the impact of competing phosphate binders). No contemporary studies have examined calcium mass balance in patients on dialysis; such studies are needed before we can fully determine what &quot;too much calcium intake&quot; means.</td>
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<td>6. What is the role of vitamin D in the treatment of patients with CKD?</td>
<td>The various activated vitamin D products are used as a means to treat SHPT and are often limited by adverse effects such as hypercalcemia and hyperphosphatemia. Observational studies show a survival benefit from treatment with activated vitamin D in dialysis patients that is independent of PTH, calcium, or phosphorus level, suggesting that all patients with CKD could benefit from some form of vitamin D replacement therapy. No RCT are available to prove this hypothesis.</td>
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<td>7. What is the best vitamin D product for patients with CKD?</td>
<td>The main advantage of more modern (and expensive) activated vitamin D analogues is their lesser hypercalcemic and hyperphosphatemic effect. If the benefit of vitamin D replacement is indeed universal and independent of PTH, calcium, and phosphorus levels, then this may not be as important after all. Head-to-head comparisons of various products (including generic ones, not just &quot;competitors&quot;) using relevant &quot;hard&quot; end points are needed to clarify this.</td>
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<tr>
<td>8. What is the role of calcium-sensing receptor agonists in the treatment of CKD-MBD?</td>
<td>There are no data on the impact of calcium-sensing receptor management on patient survival.</td>
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<td>9. What is the most cost-effective way to manage CKD-MBD?</td>
<td>The high price of several medications represents a serious limitation in everyday practice. Many of these products have cheaper generic alternatives, but both the economic and the biologic advantages of one versus the other need to be better studied.</td>
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*CKD-MBD, mineral and bone disorders in chronic kidney disease; PTH, parathyroid hormone; RCT, randomized, controlled trial; SHPT, secondary hyperparathyroidism.
practice guidelines also need to acknowledge that collaboration with the pharmaceutical sector has undeniably become the Achilles heel of these processes. A balanced, transparent, and hands-off collaboration wherein industry supports the process rather than the specific set of guidelines that involve their products may diminish the perception of conflict of interest.

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C.P.K. has received honoraria from Genzyme, Inc., the manufacturer of Sevelamer hydrochloride (Renagel), and from Amgen, Inc., the manufacturer of cinacalcet hydrochloride (Sensipar), and has received grant support from Abbott laboratories, the manufacturer of Paricalcitol (Zemplar). R.M. has received honoraria and grants from Genzyme, Inc., and has received grants and honoraria from and serves as a consultant for Shire Pharmaceuticals, the manufacturer of lanthanum carbonate (Fosrenol). K.K.-Z. has received honoraria or grants from Genzyme, Inc., Shire Pharmaceuticals, and Abbott Laboratories.

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23. Russo D, Palmiero G, De Blasio A, Ipp E, Takasu J, Sevelamer hydrochloride (Renagel), and has received grant support from Abbott laboratories, the manufacturer of Paricalcitol (Zemplar). R.M. has received honoraria and grants from Genzyme, Inc., and has received grants and honoraria from and serves as a consultant for Shire Pharmaceuticals, the manufacturer of lanthanum carbonate (Fosrenol). K.K.-Z. has received honoraria or grants from Genzyme, Inc., Shire Pharmaceuticals, and Abbott Laboratories.


