FGF23: a key player in mineral and bone disorder in CKD

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RESUMEN

El FGF23 es una hormona de reciente identificación que regula el metabolismo de los minerales y de la vitamina D. En pacientes con insuficiencia renal crónica (IRC), los niveles circulantes de FGF23 se elevan de forma progresiva para compensar la retención renal de fosfato persistente, lo cual provoca una producción renal reducida de 1,25-dihidroxivitamina D y estimula, por tanto, la secreción de la hormona paratiroidea. Este hecho sugiere que su papel es crucial en la patogénesis de la homeostasis mineral alterada en la IRC. Asimismo, se ha demostrado recientemente que el FGF23 actúa directamente en la glándula paratiroidea y media en la secreción de la hormona paratiroidea en presencia del Klotho como cofactor, aunque hasta el momento dichos efectos no se han confirmado en pacientes con IRC. El FGF23 también puede utilizarse como predictor de la mortalidad así como de un futuro desarrollo de hipertiroídisco refractario en pacientes sometidos a diálisis, en los que los niveles de FGF23 son realmente elevados como reacción al tratamiento de hiperoxofatemia y a la actividad de la vitamina D. En este resumen breve se incluyen las aproximaciones más recientes en cuanto al papel del FGF23 en la patogénesis de las alteraciones del metabolismo óseo-mineral en la IRC.

Palabras clave: Role of FGF23 in CKD.

INTRODUCTION

Disorders of mineral and bone metabolism are common complications of chronic kidney disease (CKD). Abnormal mineral metabolism occurs early in the course of CKD, which can result in significant consequences even in patients not yet on dialysis. Traditionally, these abnormalities have been investigated mainly in association with the development of secondary hyperparathyroidism, where phosphate retention, hypocalcemia, and a progressive decline in 1,25-dihydroxyvitamin D [1,25(OH)₂D], have been considered to be the main factors for abnormal parathyroid hormone (PTH) secretion.

Recently, a novel phosphaturic hormone FGF23 has been identified, initially as a pathogenic factor in rare hypophosphatemic syndromes disorders. Studies since then have shown that this hormone plays an important role in...
normal physiology as well as in the pathogenesis of alterations in mineral metabolism such as that seen in patients with CKD. In this brief review, we summarize recent findings about the role of FGF23 in the pathogenesis of mineral and bone disorders in CKD.

**ROLE OF FGF23 IN PHOSPHATE AND VITAMIN D HOMEOSTASIS**

FGF23 is a 32-kDa protein with 251 amino acids that is secreted mainly by osteocytes in bone and tumor-induced osteomalacia (TIO) that are characterized by severe hypophosphatemia, inappropriate phosphaturia, low levels of 1,25(OH)2D, and rickets or osteomalacia. Dysregulated secretion of FGF23 is also involved in a number of other diseases with abnormal phosphate and vitamin D homeostasis, such as X-linked hypophosphatemia (XLH), autosomal recessive hypophosphatemic rickets/osteomalacia (ARHR), and McCune-Albright syndrome.

In accordance with human diseases, functional in vivo studies have shown that FGF23 is one of the most potent phosphonotins that induces renal phosphate wasting and reduction of 1,25(OH)2D. Administration of recombinant FGF23 results in phosphaturia and vitamin D homeostasis, such as X-linked hypophosphatemia (XLH), autosomal recessive hypophosphatemic rickets/osteomalacia (ARHR), and McCune-Albright syndrome.

In keeping with these observations, transgenic mice that overexpress either wild-type or a mutant form of FGF23 have shown that FGF23 results in phosphaturia and vitamin D homeostasis by suppressing the expression of sodium-phosphate cotransporter that mediates physiological phosphate uptake in proximal tubular epithelial cells. Excess FGF23 also suppresses 1,25(OH)2D via inhibition of 1α-hydroxylase (CYP27B1) which converts 25-hydroxyvitamin D [25(OH)D] to 1,25(OH)2D and stimulation of 24-hydroxylase (CYP24) which converts 1,25(OH)2D to more hydrophilic metabolites with lesser biological activity.

In keeping with these observations, transgenic mice that overexpress either wild-type or a mutant form of FGF23 that is resistant to cleavage developed hyperphosphatemia, low serum 1,25(OH)2D levels, and rickets and osteomalacia. Conversely, targeted ablation of FGF23 leads to the opposite renal phenotype, consisting of hyperphosphatemia and elevated production of 1,25(OH)2D. Subsequent studies highlighted the physiologic role of FGF23 in maintaining normal serum phosphate levels in the setting of dietary phosphate variation, although the precise mechanism by which phosphate loading mediates FGF23 production remains unknown.

**FGF23-KLOTHO AXIS**

Another unique characteristic of FGF23 is that this molecule derives from bone and exerts its hormonal effects in the kidney despite the ubiquitous presence of its receptors (FGFRs). This is in sharp contrast to other FGF family members that are thought to regulate various cell functions at a local level. This mystery has been progressively unraveled by a recent major breakthrough that FGF23 requires Klotho, an anti-aging protein, as a cofactor in FGF23-FGFR1c interaction. This fact clearly explains why Klotho mutant mice display a phenotype identical to that of FGF23 null mice, both of which are characterized by premature aging-related phenotypes associated with hyperphosphatemia and paradoxically high 1,25(OH)2D levels.

Of note, Klotho is expressed in limited tissues such as the kidney, parathyroid gland, and pituitary gland. It is intriguing that such limited expression pattern corresponds to the target tissues for FGF23 as functionally defined by the induction of early growth-responsive 1 (Egr-1) expression after intravenous administration of recombinant FGF23 to rats. It is, however, still unclear how in the kidney FGF23 exerts its physiological effects on the proximal tubule despite the highest expression of Klotho-FGFR complexes in the distal tubule. Several investigators hypothesize that FGF23 actions on the proximal tubule may be indirectly mediated by FGF23 stimulation of the distal tubule and subsequent release of paracrine factors, but further researches are needed to confirm this attracting hypothesis.

**ELEVATED LEVELS OF FGF23 IN CKD**

Insights into the role of FGF23 in mineral homeostasis have launched a new field of clinical research in CKD patients. Several studies have measured circulating FGF23 levels in predialysis and dialysis patients using an enzyme-linked immunosorbent assay (ELISA) that detects the carboxyl-terminal fragment of FGF23, and reported progressively elevated FGF23 levels as serum creatinine or phosphate levels increase, suggesting its physiologic response to chronic phosphate retention. However, the possibility of accumulation of carboxyl-terminal fragments due to decreased renal function cannot be excluded in these studies.

Accordingly, a subsequent study measured intact FGF23 levels in CKD patients using a sandwich ELISA system that exclusively detects full-length human FGF23, and found similar increase in FGF23 levels along with decline in glomerular filtration rate (GFR). Furthermore, serum FGF23 levels were negatively associated with 1,25(OH)2D levels and maximal tubular reabsorption of phosphate (TmP/GFR) correlated negatively with serum FGF23 levels, consistent with the physiological action of FGF23 to inhibit phosphate reabsorption in the proximal tubule. However, patients with more advanced CKD exhibited impaired urinary phosphate excretion despite extremely high FGF23 levels.

Taken together, it is suggested that in early stage of CKD, serum FGF23 is elevated to maintain normal serum
phosphate levels by promoting urinary phosphate excretion, but in advanced stage, overt phosphate loading may overcome such compensation for decreased GFR despite markedly elevated FGF23 levels, which in turn results in decreased renal production of 1,25(OH)_{2}D, possibly thereby worsening secondary hyperparathyroidism. Another group further supported this hypothesis by elegantly showing that increased FGF23 was an independent predictor of decreased 1,25(OH)_{2}D levels and the effects of renal function and hyperphosphatemia on serum 1,25(OH)_{2}D levels were completely extinguished by adjusting for FGF23, suggesting that FGF23 is a central factor in the early pathogenesis of secondary hyperparathyroidism.

Of note, a recent observational study have shown that in patients who are beginning hemodialysis treatment high FGF23 levels were associated with mortality independently of serum phosphate levels and other known risk factors. Supposing that FGF23 may indicate phosphate retention even in patients with normophosphatemia, its measurements may be useful to identify which of those patients might benefit from more aggressive phosphate management. Whether such strategies for the control of phosphate homeostasis would prolong survival of CKD patients is worthy of further investigation.

**FGF23 AND SECONDARY HYPERPARATHYROIDISM IN CKD**

Serum FGF23 levels are progressively increased as kidney function declines and are markedly elevated once on dialysis therapy. Such high levels of FGF23 may be due to persistent phosphate retention or hyperphosphatemia, while active vitamin D therapy has also been shown to increase serum FGF23 levels in dialysis patients. This observation was further supported by in vivo and in vitro studies showing that 1,25(OH)_{2}D directly increases the production
of FGF23 by osteoblasts through the vitamin D-responsive elements present in the FGF23 promoter

In this context, it is an interesting finding that in dialysis patients with secondary hyperparathyroidism high FGF23 levels may predict the future development of refractory hyperparathyroidism

Another issue of concern is a recent finding that PTH retention as reflected by elevated FGF23 levels may contribute to further progression of parathyroid hyperplasia, because high phosphate directly stimulates PTH secretion and parathyroid cell proliferation. Another possibility is that high levels of FGF23 in uremia may be a consequence of prolonged active vitamin D administration for severe hyperparathyroidism, which may be related to future resistance to vitamin D therapy.

Besides the above-mentioned indirect effect of FGF23 on parathyroid function via inhibition of 1,25(OH)2D production, the abundant expression of Klotho in the parathyroid suggests that FGF23 may directly affect parathyroid function. In fact, a recent study using rats with normal renal function has shown that FGF23 suppresses secretion of PTH in vivo and in vitro. FGF23 also increases parathyroid 1α-hydroxylase expression and partly thereby decreases secretion of PTH in primary cultures of bovine parathyroid cells. Thus, it is likely that FGF23 is a negative regulator of parathyroid function, at least in normal physiology. However, in CKD patients with secondary hyperparathyroidism, PTH secretion remains stimulated despite extremely high FGF23 levels. Such resistance of the parathyroid to high FGF23 levels in uremia should be investigated in future studies.

Another issue of concern is a recent finding that PTH secretion is regulated in a Klotho- and Na+,K+-ATPase-dependent manner. It is proposed that when extracellular calcium is low, Na+,K+-ATPase is quickly recruited to the plasma membrane and an electrochemical gradient created by increased Na+,K+-ATPase may cause PTH release. However, it remains unclear how such a Klotho- and Na+,K+-ATPase-dependent PTH regulation interacts with the inhibitory effect of FGF23 on PTH secretion through Klotho-FGFR complexes. Furthermore, it is also a matter of concern whether such a complex mechanism is modulated in dialysis patients in whom phosphate retention is prevalent and hypocalcemic state is rare due to treatment with calcium-based phosphate binders and active vitamin D analogs. Future studies should investigate the possible complex interaction between FGF23-, Klotho- and Na+,K+-ATPase-dependent pathways regulating PTH secretion and whether these complex mechanisms are modulated in the setting of CKD.

**CONCLUSION**

The recent identification of FGF23 and Klotho as a physiological regulator of phosphate and vitamin D metabolism has considerably advanced our understanding of the mineral and bone disorder in CKD. It is now clear that FGF23 plays a central role in the pathogenesis of altered mineral metabolism and secondary hyperparathyroidism in CKD patients. FGF23 can be used not only as a biomarker for assessing phosphate retention but also as a predictor of mortality and future development of refractory hyperparathyroidism. However, the precise role of extremely elevated FGF23 levels in uremia still remains unclear, especially as to its direct effect on parathyroid function. Further elucidation of the FGF23-Klotho axis will help us to establish a more rational approach for the management of mineral and bone disorder that is associated with high burden of morbidity and mortality in CKD patients.

**REFERENCES**


