Klotho and calciprotein particles as therapeutic targets against accelerated ageing

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The klotho gene, named after a Greek goddess who spins the thread of life, was identified as a putative ‘ageing-suppressor’ gene. Klotho-deficient mice exhibit complex ageing-like phenotypes including hypogonadism, arteriosclerosis (vascular calcification), cardiac hypertrophy, osteopenia, sarcopenia, frailty, and premature death. Klotho protein functions as the obligate co-receptor for fibroblast growth factor-23 (FGF23), a bone-derived hormone that promotes urinary phosphate excretion in response to phosphate intake. Thus, Klotho-deficient mice suffer not only from accelerated ageing but also from phosphate retention due to impaired phosphate excretion. Importantly, restoration of the phosphate balance by placing Klotho-deficient mice on low phosphate diet rescued them from premature ageing, leading us to the notion that phosphate accelerates ageing. Because the extracellular fluid is super-saturated in terms of phosphate and calcium ions, an increase in the phosphate concentration can trigger precipitation of calcium-phosphate. In the blood, calcium-phosphate precipitated upon increase in the blood phosphate concentration is adsorbed by serum protein fetuin-A to form colloidal nanoparticles called calciprotein particles (CPPs). In the urine, CPPs appear in the renal tubular fluid when FGF23 increases phosphate load excreted per nephron. CPPs can induce cell damage, ectopic calcification, and inflammatory responses. CPPs in the blood can induce arteriosclerosis and non-infectious chronic inflammation, whereas CPPs in the urine can induce renal tubular damage and interstitial inflammation/fibrosis. Thus, we propose that CPPs behave like a pathogen that accelerates ageing and should be regarded as a novel therapeutic target against age-related disorders including chronic kidney disease.

Introduction
Phosphorus is one of the six elements essential for life (hydrogen, carbon, nitrogen, oxygen, sulfur, and phosphorus) [1]. In vivo, phosphorus exists in the form of phosphate, which is not only a major constituent of cell membrane (e.g. phospholipids) and nucleic acids but also a critical functional molecule for energy metabolism (e.g. ATP) and cell signaling (e.g. protein phosphorylation). Phosphate is so fundamental in the structure and function of organisms that a behavior termed ‘phosphate appetite’ has evolved. For instance, animals under phosphate-deficient state instinctively seek for phosphate-rich foods, leading to osteophagic behavior (lick/eat bones of dead animals) even in herbivores [2]. The instinctive behavior to seek selectively for a lacking nutrient is termed specific appetite or specific hunger, and observed only for a limited number of essential nutrients such as sodium, indicating how important phosphate is for life. However, unlike the wild life, we barely fall into a situation of phosphate deficiency. The phosphate content in diet is known to correlate positively with the protein content [3]. Therefore, the western-style diet characterized by high intake of meats and dairy products is rich in phosphate. In addition, some food additives and preservatives contain a large amount of phosphate [4,5]. Consequently, most people on the western diet take in far more phosphate than necessary. How does the excessive phosphate intake affect our health?
Physiological responses to dietary phosphate intake

Oral gavage of phosphate in mice quickly raises their blood phosphate levels, whereas their blood calcium levels are decreased reciprocally [6], although the mechanism is not clear. However, because the blood is a supersaturated solution regarding phosphate and calcium ions, even a slight increase in the phosphate concentration can trigger precipitation of calcium-phosphate and contribute to the decrease in blood calcium levels upon phosphate intake. A decrease in the blood concentration of calcium ions inactivates calcium-sensing receptor (CaSR) expressed on the surface of parathyroid chief cells and induces secretion of parathyroid hormone (PTH) [7]. PTH has the activity that increases blood calcium levels and urinary phosphate excretion, through which blood calcium and phosphate levels are restored to the basal levels. This negative feedback to maintain the phosphate homeostasis takes place within a few hours after dietary phosphate intake [6].

When mice are placed on high phosphate diet for a few days or longer, circulating levels of fibroblast growth factor-23 (FGF23) are increased [6]. FGF23 is a hormone secreted from osteocytes and osteoblasts when they sense phosphate intake [8]. FGF23 circulates in the blood and acts on the kidney to increase urinary phosphate excretion through suppressing phosphate resorption in renal proximal tubules, thereby increasing phosphate excretion per nephron [9]. Thus, two phosphaturic hormones, PTH and FGF23, promote urinary phosphate excretion in response to phosphate intake to maintain the phosphate homeostasis in a short-term (hours) and in a long-term (days), respectively.

Pathology induced by excess phosphate intake

It has been known for decades that mice and rats develop renal tubular damage and interstitial fibrosis when placed on high phosphate diet for a few months [10,11]. In rats, the severity of the kidney damage was shown to correlate with phosphate excretion per nephron [10]. However, it should be noted that the kidney damage was not observed when the phosphate excretion per nephron was below \( \sim 1.0 \) \( \mu \)g per day, indicating that a threshold exists for induction of the kidney damage [10,12].

In animals that maintain phosphate homeostasis, the amount of phosphate adsorbed from the gastrointestinal tract should be equal to the amount of phosphate excreted into urine. Therefore, even when placed on diet containing the same amount of phosphate, animals with lower nephron number should have higher phosphate excretion per nephron and thus develop severer kidney damage. Vice versa, the amount of phosphate intake that induces kidney damage should be different from individual to individual, depending on their residual nephron number.

Although there is a large individual and species difference in the nephron number and the amount of urinary phosphate excretion, normal young adults have \( \sim 1 \) million nephrons per kidney and excrete \( \sim 1 \) g of phosphate into urine on average per day [13,14]. Thus, phosphate excretion per nephron is roughly estimated as \( \sim 0.5 \) \( \mu \)g per day in normal adults. The nephron number is decreased progressively during the course of ageing. People in their sixties or seventies are reported to have \( \sim 50\% \) less nephrons than people in their twenties [14]. Assuming that elderly people whose nephron number has been decreased to \( \sim 0.5 \) million per kidney are on the same diet when they were young, their phosphate excretion per nephron may reach 1.0 \( \mu \)g per day, the level that potentially causes renal tubular damage and interstitial fibrosis. These histological changes are observed not only in patients with kidney diseases but also in elderly people without any particular disorders. The renal pathology universally observed in the aged is composed of renal tubular damage/atrophy, interstitial fibrosis, arteriosclerosis, and glomerulosclerosis among others, which are collectively designated as ‘ageing kidney’ [15,16]. Hence, dietary phosphate ingested on a daily basis may contribute to acceleration of kidney ageing in individuals with reduced nephron number such as elderly people and patients with chronic kidney disease.

Unlike humans, mice barely develop ageing kidney even when they reach an advanced age. Mice have \( \sim 10,000 \) nephrons per kidney on average [17] and excrete \( \sim 0.4 \) mg phosphate on standard chow diet containing 0.35% inorganic phosphate [12]. Thus, the phosphate excretion per nephron is estimated as \( \sim 0.02 \) \( \mu \)g per day, which is far below the threshold (1.0 \( \mu \)g per day) to develop kidney damage. However, when mice were placed on diet containing 2.0% inorganic phosphate, their phosphate excretion per nephron reached \( \sim 2.0 \) \( \mu \)g per day and developed renal tubular damage and interstitial fibrosis [11,12], recapitulating the features of ageing kidney at least in part.

Mechanism of phosphate-induced kidney damage

The molecular mechanism of the chronic kidney damage induced by dietary phosphate load has recently been elucidated [12]. An increase in phosphate intake must be associated with the corresponding increase in urinary phosphate excretion in order to maintain the phosphate balance. This demand is met mainly by increasing circulating FGF23.

\[ \text{FGF23} \]
levels. Because FGF23 suppresses phosphate reabsorption at proximal tubules, it should increase phosphate concentration in the proximal tubular fluid. Once the phosphate concentration exceeds a threshold, calcium-phosphate microcrystals are precipitated in the proximal tubular fluid. These microcrystals bind to toll-like receptor-4 (TLR4) expressed on the apical membrane of proximal tubular cells and removed from the tubular fluid by endocytosis to be transferred to lysosome. However, overload of calcium-phosphate microcrystals elevates the luminal pH of the lysosomes and disturbs lysosomal function and autophagy, which eventually induces proximal tubular cell damage [12,18]. In addition, binding of calcium-phosphate microcrystals to TLR4 activates the p38 and NFκB signaling pathways to induce expression of cytokines and chemokines, including interleukin-6, monocyte chemotactic protein-1, tumor necrosis factor-α, and osteopontin, which potentially contribute to interstitial inflammation [12]. Tubular cell damage and interstitial inflammation eventually lead to fibrosis and kill nephrons. The decrease in the nephron number further boosts FGF23 secretion to maintain the phosphate homeostasis unless phosphate intake is reduced, thereby triggering a deterioration spiral leading to progressive nephron loss and acceleration of kidney ageing (Figure 1).

It should be noted that the serum phosphate level does not contribute to this mechanism. The critical factor is not the phosphate level in the blood but the phosphate level in the proximal tubular fluid. In clinical settings, however, it is virtually impossible to harvest the renal tubular fluid from proximal tubules by micropuncture and measure the
phosphate concentration. Therefore, we have established a method for estimating the phosphate concentration in the renal tubular fluid at the distal portion of proximal tubules from the data obtained by regular blood and urine tests [12]. The phosphate concentration in the glomerular filtrate should be equal to the serum phosphate concentration. Phosphate is then reabsorbed while the glomerular filtrate flows down in the proximal tubule. Phosphate reabsorption takes place almost exclusively at proximal tubules [19]. Thus, the phosphate concentration in the tubular fluid at the distal portion of proximal tubules should be equal to the product of serum phosphate concentration and fractional excretion of phosphate (FEp) unless water is not reabsorbed. However, ~70% water is actually reabsorbed at proximal tubules [20], resulting in concentrating all the solutes by 3.33-fold. Therefore, the estimated phosphate concentration at the distal portion of proximal tubules (ePTFp) can be calculated by the following equation:

\[
ePTFp = Sp \times FEp \times 3.33 = Sp \times \frac{Up \times Scr}{Ucr \times Sp} \times 3.33 = \frac{Up}{Ucr} \times Scr \times 3.33
\]

where \( Sp, Up, Scr, \) and \( Ucr \) represent the concentration of serum phosphate, urine phosphate, serum creatinine, and urine creatinine, respectively. We confirmed that ePTFp indeed served as approximation of the actual phosphate concentration in the proximal tubular fluid harvested by micropuncture in rats [12,21].

Relation between ePTFp, renal tubular damage, and FGF23 was explored in mice (Figure 2A) [12]. Double logarithmic plots of FGF23 and osteopontin against ePTFp revealed the presence of a threshold (Figure 2B). Namely, neither renal tubular damage (i.e., increase in osteopontin expression) nor increase in FGF23 was observed when ePTFp was below ~5 mg/ml, but once ePTFp exceeded this threshold, renal tubular damage was induced, probably because calcium-phosphate microcrystals were precipitated in the tubular fluid. In addition, the fact that a linear regression between ePTFp and tubular damage became evident in double logarithmic plot is consistent with the fact that calcium-phosphate microcrystals are the true culprit of tubular damage, because it indicates the power function. Specifically,

\[
Ca_m Pi_n \rightleftharpoons mCa + nPi
\]

\[
K_d = \frac{[Ca]^m \cdot [Pi]^n}{[Ca_m Pi_n]}
\]

\[
\log([Ca_m Pi_n]) = m \cdot \log([Ca]) + n \cdot \log([Pi]) - \log(K_d)
\]

where \( K_d \) indicates the dissociation constant. \( Ca, Pi, \) and \( Ca_m Pi_n \) indicate calcium, phosphate, and calcium-phosphate, respectively. The equation above shows that the regression between the calcium-phosphate concentration \([Ca_m Pi_n]\) and the phosphate concentration \([Pi]\) should be linear in the double logarithmic plot.

The similar correlation between ePTFp and FGF23 was observed in CKD patients as well (Figure 3A) [12]. The threshold values of ePTFp and FGF23 in humans were 2.32 and 53 pg/ml, respectively. A prospective cohort study (5039 participants) indicated that patients with FGF23 \( \geq 53 \) pg/ml developed kidney events (initiation of chronic dialysis or serum creatinine doubling) within 5 years more frequently than patients with FGF23 \(< 53 \) pg/ml, independently of serum creatinine levels (Figure 3B) [12]. It should be noted that approximately 1/4 of ‘healthy’ adults aged 45 years or older and 2/3 of stage 3 CKD patients are estimated to have higher serum FGF23 levels than 53 pg/ml [22,23] and thus may have already developed renal tubular damage caused by calcium-phosphate microcrystals in the renal tubular fluid.

**The phosphate-centric view of chronic kidney disease**

Chronic kidney disease (CKD) is a relatively new disease entity established a few decades ago and defined as a state of any abnormality in renal function and/or structure persisting for 3 months or longer [24]. In most cases, CKD ensues when the age-associated decrease in the nephron number is accelerated by kidney diseases (e.g. acute kidney injury, glomerulonephritis, polycystic kidney) and disorders causing renal complications, most notably hypertension and diabetes mellitus. Thus, CKD is very prevalent in the ageing society and affects more than 10% of the total population [25,26]. CKD is classified into five stages based on the estimated glomerular filtration rate (eGFR) that can be calculated from the serum creatinine level [26]. In order of progression, Stage 1: eGFR 90 ml/min/1.72m² or greater, Stage 2: eGFR between 60 and 89, Stage 3: eGFR between 30 and 59, Stage 4: eGFR between 15 and 29, Stage 5: eGFR less than 15. Once CKD advances to stage 5, renal replacement therapy (dialysis or renal transplantation) should be considered.

One of the earliest signs of CKD includes an increase in FGF23 [23]. Serum FGF23 levels start increasing as early as stage 2-3, which is regarded as a physiological response to compensate for a decrease in the nephron number by
increasing phosphate excretion per nephron by FGF23 to maintain phosphate homeostasis. However, the increase in FGF23 can increase the risk for formation of calcium-phosphate microcrystals in the tubular fluid and induce renal tubular damage and interstitial fibrosis [12]. Once these renal damages reduce the functional nephron number, FGF23 should be further increased to trigger the deterioration spiral towards acceleration of CKD progression as shown in Figure 1 [12]. Besides functioning as a phosphaturic hormone, FGF23 functions as a counter-regulatory hormone for vitamin D [9]. FGF23 lowers circulating levels of active vitamin D (1,25-dihydroxyvitamin D₃) by down-regulating expression of 1α-hydroxylase necessary for synthesis of active vitamin D and up-regulating expression of 24-hydroxylase necessary for inactivation of active vitamin D in renal proximal tubular cells. A decrease in active vitamin D increases PTH, because a potent negative feedback loop exists between PTH and active vitamin D, in which PTH raises serum active vitamin D levels, whereas active vitamin D lowers serum PTH levels [27]. Thus, the increase in FGF23 induces secondary hyperparathyroidism through suppressing serum levels of active vitamin D. Indeed, increase in FGF23, decrease in active vitamin D, and increase in PTH occur in this order during the course of CKD progression [23]. Once the residual nephron number is decreased to the level that cannot maintain the phosphate balance by increasing FGF23 any longer, phosphate retention and hyperphosphatemia ensues. Accordingly, increase in serum phosphate is observed only in advanced CKD patients at stage 4-5 (end-stage renal disease; ESRD) (Figure 4). Besides hyperphosphatemia, hypocalcemia ensues due to low active vitamin D [28]. The disturbed phosphate/calcium homeostasis and the changes in the hormone levels (high FGF23, low active vitamin D, and high PTH) that occur during CKD progression are usually associated with bone mineral loss and reciprocal mineralization in extra-osseous tissues (e.g. vascular calcification), and designated as CKD-MBD (mineral and bone disorder) [29].

**Phosphate restriction**

Phosphate restriction by reducing dietary phosphate intake and/or by taking phosphate binders is currently applied only for ESRD patients with hyperphosphatemia (stage 4-5), aiming at lowering serum phosphate levels [30]. The expected clinical outcomes include suppression of vascular calcification and cardiovascular events. However, as discussed above, the deterioration spiral leading to progressive nephron loss has already been launched in early stage CKD patients (stage 2-3) when serum FGF23 levels start increasing. Thus, reduction of phosphate excretion per nephron by phosphate restriction should be beneficial not only for ESRD patients with hyperphosphatemia but also for early stage CKD patients with hyper-FGF23-emia but without hyperphosphatemia [8,12,31]. The expected clinical outcome is suppression of nephron loss or CKD progression, but not suppression of vascular calcification and cardiovascular events.
Figure 4. Progression of CKD-MBD
See the text. Regardless of the underlying disorders, progression of CKD can be viewed as the progressive loss of the functional nephron number. Modified from [8].

Figure 5. Phytate as a source of phosphate
See the text. Phytase is necessary to release phosphate from phytate.

The current diet therapy for hyperphosphatemia is to avoid phosphate-rich foods based on the food composition table, such as meat, fish, and dairy products [32]. Because the phosphate content in foods is positively correlated with the protein content [3], phosphate-restricted diet inevitably means protein-restricted diet. However, recent clinical studies have shown that strict restriction of dietary protein intake can lead to a malnutrition state termed protein-energy wasting (PEW), which is associated with increased mortality [33]. Therefore, ‘phosphate restriction without protein restriction’ is desirable. To attain this goal, the following two approaches may be practical and effective.

First, it is necessary to pay attention not only to the phosphate content but also to the absorption efficiency, namely, how much percent of phosphate in each food ingredient is actually absorbed from intestine. For example, soybeans are a major plant source of protein and listed as an ingredient rich in phosphate in the food composition table. Thus, ESRD patients with hyperphosphatemia may have been instructed to avoid soybean products. However, phosphate in soybeans exists primarily in the form of phytate (inositol hexakisphosphate; IP6), which is not absorbed from the intestine and excreted into feces as is [34]. In plants, phosphate exists mainly as phytate. In order for phosphate in phytate to be absorbed from the gastrointestinal tract, inorganic phosphate must be released from phytate by hydrolyzation of the inositol-phosphate linkages with phytase (Figure 5). Phytase is an enzyme produced by enteric bacteria that reside in the gut of ruminant animals [35]. Therefore, cows and sheep can hydrolyze phytate and utilize soybeans as a source of phosphate but monogastric animals such as pigs and chickens cannot. The fact that pigs and chickens are unable to absorb phosphate from phytate raises two problems [36]. First, inorganic phosphate must be added to their feed to support their growth. Second, phytate excreted into feces can cause soil pollution with phosphorus. To solve these problems, phytate has been added to the feed containing soy flour for pigs and chickens. Because humans cannot utilize phytate as a source of phosphate either, replacement of animal-based protein with plant-based protein is considered as an effective way to reduce phosphate absorption without restricting protein intake [37].
Second, it is of critical importance for dietary phosphate restriction to avoid food additives containing inorganic phosphate [5,32]. Unlike phosphate in foods, phosphate in food additives is inorganic and thus adsorbed nearly 100%. Although various kinds of food additives are used in processed foods, it is not compulsory under the current regulations to indicate the amount of phosphate added to the processed foods in the nutrition facts label. Therefore, it is virtually impossible for consumers to know how much phosphate-containing additives has been used. However, it was reported that instructions how to read the nutrition facts labels and avoid phosphate-containing additives when purchasing groceries or visiting fast food restaurants lowered serum phosphate levels in ESRD patients [38]. Substantial contribution of food additives to daily phosphate intake is also endorsed by the fact that the amount of phosphate intake is associated with socio-economic status. Specifically, people in a low socio-economic status may have insufficient money and time to prepare well-balanced homemade meals from fresh ingredients without food additives, and tend to purchase processed foods and cheap junk foods, which in most cases contain a large amount of food additives. Indeed, the socio-economic status has been reported as an independent determinant of serum phosphate levels (i.e. a low socio-economic status is associated with high serum phosphate levels) [39,40].

In summary, a practical way to attain ‘dietary phosphate restriction without protein restriction’ would be to replace animal-based protein with plant-based protein and to reduce intake of phosphate-containing food additives.

### Phosphate and ageing

Thirty years ago, we reported an obscure mutant mouse strain that exhibited complex phenotypes resembling human ageing [41]. The founder of this strain was a transgenic mouse carrying an insertional mutation caused by transgene integration on the chromosome 5. Homozygotes for the transgene grew normally until weaning, but thereafter developed multiple ageing-like phenotypes including growth arrest, multiple organ atrophy (gonads, thymus, skin), vascular calcification, cardiac hypertrophy, sarcopenia, osteopenia, emphysematous lung, hearing disturbance, cognition impairment, frailty, and died around 2 months of age. The mutant was named klotho after a Greek goddess who spins the thread of life. The gene disrupted by the transgene insertion (the klotho gene) encoded a ~130 kD type-I single-pass transmembrane protein with a large extracellular domain and a very short intracellular domain composed of only 11 amino acids. It was expressed in limited cell types including renal tubular epithelial cells, choroid plexus epithelial cells, and parathyroid chief cells. The extracellular domain of klotho was composed of two homologous domains. Each domain had weak homology to the family 1 glycosidases. It is still controversial whether Klotho protein has enzymatic activity as a glycosidase [42–46], because two conserved catalytic glutamate residues essential for the glycosidase activity and thus conserved in all the family 1 glycosidases are replaced with other amino acids in Klotho protein.

The clue for the Klotho protein function was a report on FGF23 knockout mice [47]. As predicted, FGF23 knockout mice failed to increase urinary phosphate excretion in response to phosphate intake and exhibited phosphate retention phenotypes, including hyperphosphatemia and vascular calcification. In addition, they unexpectedly displayed complex ageing-like phenotypes, including growth arrest, osteopenia, and shortened life span. These phenotypes were reminiscent of those seen in klotho mice [41]. The remarkable similarity between FGF23 knockout mice and klotho mice led us to hypothesize that FGF23 and Klotho might function in the same signaling pathway. At that time, FGF23 was supposed to function through binding to fibroblast growth factor receptors (FGFRs). FGFRs are receptor tyrosine kinases encoded by 4 distinct genes (FGFR1–4). The FGFR1, FGFR2, and FGFR3 genes generate multiple isoforms through alternative splicing [48]. However, the affinity of FGF23 to any FGFR isoforms are too low (KD = 200–700 nM) for FGF23 to function at its physiological concentration (1–2 pM) [49]. The answer was that FGF23 requires Klotho to bind to FGFRs. We found that Klotho protein formed constitutive complexes with FGFR1c, FGFR3c, or FGFR4. The FGFR-Klotho complexes had much higher affinity for FGF23 than FGFR alone [48]. The same finding was confirmed later independently by another laboratory [50]. The fact that Klotho functions as the obligate co-receptor for FGF23 clearly explained why FGF23 knockout mice and klotho mice developed very similar phenotypes. In 2018, crystal structure of the Klotho-FGFR1c-FGF23 ternary complex was solved [45]. Like its namesake who spins the thread of life, Klotho protein turned out to have a long ‘thread’ termed ‘receptor binding arm (RBA)’. Although the RBA appeared to have an intrinsically disordered structure, it took a solid structure once it captured FGFR1c to generate a groove into which FGF23 fit perfectly [51].

The true nature of the problem in FGF23 knockout mice and klotho mice is phosphate retention caused by defects in the FGF23-Klotho endocrine axis that lead to impaired phosphate excretion into urine. Indeed, restoration of the phosphate balance by placing FGF23 knockout mice and klotho mice on low phosphate diet rescued them from ageing-like phenotypes [52,53]. Based on these observations, we have reached the notion that phosphate accelerates ageing [31,54].
### Mechanism by which phosphate accelerates ageing

Consistent with the notion that life is derived from the sea, composition of the sea water and the human body is similar (Table 1) [55]. Indeed, 9 out of the top 10 abundant elements are identical between them. The exceptions are magnesium and phosphorus. Magnesium is among the top 10 elements in the sea water, whereas magnesium is replaced with phosphorus in the human body [55]. This fact may imply that organisms that accumulate phosphorus have emerged at some time point during evolution. It occurred about 400 million years ago when the bony fish appeared. Organisms evolved thereafter accumulate phosphorus in the bone in the form of calcium-phosphate. Some organisms before the bony fish have cartilage or skeletons made of calcium-carbonate. Because calcium-phosphate is physically much harder than calcium-carbonate, the bone made of calcium-phosphate is stronger than the skeletons made of calcium-carbonate. Thus, acquisition of the strong bone made of calcium-phosphate may be a prerequisite for evolution of terrestrial vertebrates, which are required to support their own body weight and move around without the help of water buoyancy [8,56]. The reason that magnesium has been replaced with phosphorus may lie in the fact that magnesium can inhibit formation of calcium-phosphate crystals.

To create the bone made of calcium-phosphate, terrestrial vertebrates maintain the extracellular fluid in a super-saturated condition regarding calcium and phosphate ions and provide a cue for precipitation when and where they want to make the bone. To secure this strategy, they have acquired two systems. First, the FGF23-Klotho endocrine system has evolved to strictly control the extracellular phosphate concentration. It should be noted that the klotho gene orthologs exist only in organisms that have bones made of calcium-phosphate [56], suggesting that the klotho gene may have evolved to maintain the phosphate homeostasis. Second, they have acquired defense systems to prevent growth of calcium-phosphate precipitations if it should occur in extraosseous tissues and extracellular fluid. Formation of calciprotein particles (CPPs) is one of such defense mechanisms.

### Calciprotein particles

CPPs are defined as mineral-protein complexes containing solid-phase calcium-phosphate and serum protein fetuin-A [57,58]. Fetuin-A has an ability to adsorb calcium-phosphate precipitates and prevent them from growing to large crystals. The process of CPP formation has been investigated in vitro using a solution containing calcium, phosphate, and serum (Figure 6) [31,59–62]. Once the concentration of phosphate and calcium ions exceeds the solubility limit, tiny amorphous calcium-phosphate precipitates appear in the solution. These precipitates are adsorbed immediately by fetuin-A. As a result, a fetuin-A molecule laden with amorphous calcium-phosphate are generated, which are the most primitive CPPs called calciprotein monomers (CPMs). CPMs have diameter of ~9 nm and spontaneously undergo aggregation to become primary CPPs with a diameter of less than 100 nm. Primary CPPs undergo further aggregation and transition of the calcium-phosphate from the amorphous phase to the crystalline phase. CPPs containing crystalline calcium-phosphate are called secondary CPPs with a diameter usually larger than primary CPPs [6,58]. This series of events is not a biological process but a physicochemical phenomenon that progresses spontaneously over time.

There is a significant difference in the activity between primary CPPs and secondary CPPs. Primary CPPs function as a physiological regulator of FGF23 production and secretion [6]. Regarding the mechanism by which osteoblasts and osteocytes sense phosphate intake, it has been postulated that these cells may express a putative
‘phosphate-sensing receptor’ through which they sense a transient increase in blood phosphate levels after phosphate intake (postprandial hyperphosphatemia) and secrete FGF23. This notion is obviously prompted by the fact that parathyroid chief cells express CaSR through which they sense changes in blood calcium levels and regulate PTH secretion. However, the putative phosphate-sensing receptor has not been identified. Rather, several lines of evidence suggest that FGF23 secretion may not necessarily be increased in response to an increase in serum phosphate levels. For instance, an increase in serum phosphate levels failed to induce FGF23 secretion in the presence of hypocalcemia in mice and rats [63,64]. In addition, an increase in serum calcium levels induced FGF23 secretion, but it did not in the presence of hypophosphatemia [64]. These observations indicate that both calcium and phosphate are necessary for inducing FGF23 secretion, suggesting that it may not be phosphate but CPPs that can induce FGF23 secretion and production. In fact, primary CPPs, but not phosphate, were found as a potent inducer of FGF23 production and secretion in cultured osteoblastic cells [6]. An increase in FGF23 production and secretion by CPPs induces phosphaturia and increases phosphate outflow from the blood. It also lowers the serum active vitamin D level. Because the primary activity of active vitamin D is to increase calcium absorption in the gut, a decrease in the serum active vitamin D level reduces calcium absorption from the intestine and thus calcium inflow into the blood. The increase in phosphate outflow and the decrease in calcium inflow by FGF23 should suppress formation of CPPs in the blood, thereby closing a negative feedback loop (Figure 7) [6]. Thus, CPPs have emerged as a physiological regulator of the FGF23-Klotho endocrine axis. Furthermore, CPPs may function as a carrier that delivers calcium and phosphate absorbed from the gastrointestinal tract directly to the bone. In fact, an in vivo imaging of mice after intravenous injection of fluorescently labeled CPPs detected accumulation of the CPPs on the inner bone surface [6].

On the other hand, secondary CPPs have been reported to exert pathogenic activity. Secondary CPPs can induce calcification in cultured vascular smooth muscle cells and innate immune responses in culture macrophages as if they were a pathogen [65–69]. In clinical studies, circulating levels of secondary CPPs were reported to correlate with parameters for inflammation (high sensitive CRP), vascular stiffness (aortic pulse wave velocity), vascular calcification (coronary artery calcification score), and coronary artery plaque thickness [70–72]. A recent clinical study using a novel high sensitive assay for quantification of secondary CPPs in serum and plasma samples identified the serum phosphate level and the age as two independent determinant of plasma secondary CPP levels [73]. Namely, high serum phosphate levels and old age are independently associated with high plasma secondary CPP levels. Based on these observations, we hypothesize that secondary CPPs may be a pro-ageing factor that induces non-infectious chronic inflammation and arteriosclerosis.
Figure 7. The FGF23-Klotho endocrine axis
See the text. Klotho protein has a long amino-acid stretch designated as the receptor binding arm (RBA) that directly interacts with FGFRs. Modified from [6,8,51].

Table 2 Colloids in the blood

<table>
<thead>
<tr>
<th>Insoluble materials</th>
<th>Lipids</th>
<th>Calcium-phosphate</th>
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</thead>
<tbody>
<tr>
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<td>Fetuin-A</td>
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<td>Lipoprotein</td>
<td>CPP</td>
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<td>LDL receptor</td>
<td>Toll-like receptor-4</td>
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<tr>
<td>Storage</td>
<td>Fat</td>
<td>Scavenger receptor-A</td>
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<td></td>
<td>Metabolic syndrome</td>
<td>Ageing</td>
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CPPs were reported to bind not only to TLR4 but also to scavenger receptor-A. Modified from [8].

Concluding remarks
In mammals, insoluble materials are adsorbed by specific serum proteins and become colloidal particles to be dispersed in the blood and transferred between organs. Lipids and calcium-phosphate are the two major insoluble materials in mammals. Lipids are adsorbed by apoproteins and become colloidal particles called lipoproteins. The activity of colloidal particles depends not only on their composition but also on their colloidal properties, including particle size and density. In fact, low-density lipoprotein (LDL) is pro-atherogenic, whereas high-density lipoprotein (HDL) is anti-atherogenic. Lipids should be eventually stored in adipose tissues, but when targeted ectopically in arteries, arteriosclerosis (atherosclerosis) ensues. When stored in the liver and skeletal muscles, fatty liver and insulin resistance ensues, leading to the metabolic syndrome. Likewise, calcium-phosphate is adsorbed by fetuin-A and become CPPs. The activity of CPPs depends on not only their composition (whether or not they contain crystalline calcium-phosphate) but also their particle size (secondary CPPs are larger than primary CPPs). Calcium-phosphate should be eventually stored in the bone, but when targeted ectopically in arteries, arteriosclerosis (vascular stiffness and calcification) ensues. When targeted in white blood cells, non-infectious chronic inflammation is induced, which may accelerate ageing (Table 2) [8,61]. Although research on CPPs is still in its infancy, we expect that CPPs may become a subject of study as important as lipoproteins in the future.

Data Availability
All supporting data are included within the main article.

Competing Interests
The authors declare that there are no competing interests associated with the manuscript.
Abbreviations

CKD, chronic kidney disease; CPM, calciprotein monomer; CPP, calciprotein particle; HDL, high-density lipoprotein; LDL, low-density lipoprotein; RBA, receptor binding arm.

References


