REVIEW

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Coronary artery calcification and end-stage renal disease: Vascular biology and clinical implications

KEY POINTS

Coronary artery calcifications appear to be an indicator of total atherosclerotic disease burden, but their relation to the stability of individual atherosclerotic plaques is not well understood.

Plaque rupture and acute coronary syndromes can occur without plaque calcification and may be more often associated with noncalcified, soft plaque.

Pharmacologic control of calcium and phosphate metabolism should be guided by the nephrologic and endocrinologic needs of the renal failure patient and not withheld out of concern about coronary calcifications.

Efforts to slow coronary artery disease progression in patients with end-stage renal disease should emphasize aggressive control of recognized cardiovascular risk factors.

T HE METABOLIC CHANGES associated with end-stage renal disease (ESRD) and its treatment may accelerate development of cardiovascular disease. In fact, cardiovascular disease, particularly coronary artery disease and chronic heart failure, is the leading cause of death in patients with ESRD.^{1,2} Yet the pathophysiology of this association is complex and only partially understood.³

Increased coronary artery calcification is one of the metabolic changes related to ESRD. This review summarizes current knowledge on the clinical significance of coronary calcification in patients with ESRD, the role of pharmacologic therapies to prevent skeletal bone loss, and their impact on calcium deposition in the vascular wall.

CALCIFICATION AND ESRD: A CONNECTION TO CARDIOVASCULAR DISEASE?

The strong association between coronary artery disease and ESRD may be partly explained by the many risk factors shared by the two conditions, such as advanced age, hypertension, diabetes mellitus, hyperhomocysteinemia, and hyperlipidemia.^{3,4} As reviewed separately in this supplement,⁵ recent attention has focused on disorders of calcium and phosphate metabolism and their treatments as potential accelerants of cardiovascular disease in ESRD. Briefly, decreased phosphate excretion and hypovitaminosis D cause hyperphosphatemia and subsequent hyperparathyroidism.^{6–8} These changes cause altered bone metabolism with skeletal bone resorption (renal osteodystrophy) and extraosseal calcifications (FIGURE 1). Extraosseal cal-

Both authors have indicated that they have no affiliation with or financial interest in a commercial organization that poses a potential conflict of interest with their article. cifications apparently result from passive precipitation of calcium if the level of the calcium/phosphate product in blood increases above local conditions of saturation.⁶ Calcifications are prominent in the kidney but have also been described in various cardiovascular tissues, such as heart valves, myocardium, and coronary arteries.^{9–12} Clear and compelling data have shown an increased prevalence of cardiac calcifications in ESRD, especially after long-term dialysis, but the pathogenesis and clinical significance of these calcifications are incompletely understood.

CORONARY CALCIFICATIONS AND ATHEROSCLEROTIC PLAQUE

Two seemingly discordant principles must be understood about the role of calcification in the development of atherosclerotic coronary plaques and their relation to coronary risk:

- The prevalence of coronary atherosclerosis and calcifications is high in persons who do not have clinically evident coronary artery disease.
- Plaque rupture and acute coronary syndrome can occur without calcification and, in fact, may be more frequently associated with noncalcified, soft plaque.

Biology of coronary calcifications

Coronary artery calcifications occur almost exclusively at sites of atherosclerotic lesions.¹³ Calcification in the development of these plaques is a complicated, actively regulated process of mineralization that is similar to bone formation and remodeling.^{14–17} Coronary artery calcification is found in small amounts in early lesions and more extensively in advanced lesions (**FIGURE 2**).¹⁸

Calcium phosphate (hydroxyapatite) is formed in vesicles that pinch off from arterial wall cells, analogous to the way that matrix vesicles pinch off from chondrocytes in developing bone. A close spatial association exists between cholesterol deposits and hydroxyapatite. Atherosclerotic lesions in younger adults reveal small aggregates of crystalline calcium among the lipid particles of the necrotic plaque core. It has been postulated that membrane vesicles derived from apoptotic foam cells within extracellular, lipid-rich necrotic

Calcium/phosphate metabolism in ESRD

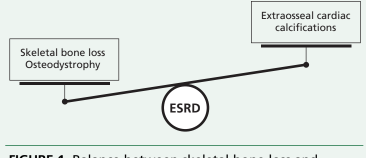


FIGURE 1. Balance between skeletal bone loss and extraosseal calcification in patients with end-stage renal disease (ESRD).

plaque cores may serve as the sites of calcium deposits. Macrophages in atherosclerotic lesions express 1-alpha-hydroxylase activity, producing 1,25-vitamin D,¹⁹ and also have osteoclastic capacity for phagocytic removal of calcium mineral from the artery wall.

Epidemiology of coronary calcifications

There are many risk factors for coronary artery calcifications besides ESRD,^{20–23} including advanced age,²² male gender,²⁴ elevated plasma cholesterol,^{25,26} diminished high-density lipoprotein cholesterol,²⁶ cigarette smoking,^{20,21} elevated blood pressure,²¹ obesity,²⁶ diabetes,²⁰ and elevated triglycerides.²⁶

Coronary calcifications and atherosclerotic plaque are much more common than clinically symptomatic coronary artery disease, positive stress tests, or angiographic stenosis. Coronary calcification is present in 50% of persons 40 to 49 years old and 80% of those 60 to 69 years old.^{21,23,24,27–30} Similarly, histologic studies and intravascular ultrasonography show that the prevalence of atherosclerotic plaque rises from 40% to 50% among persons 20 to 29 years old to 60% to 80% among those 30 to 39 years old.^{31,32} However, results from the Framingham study³³ indicate that the expected 8-year incidence of coronary events ranges from less than 1% for persons younger than 40 to 15% for those older than 80, and significant angiographic stenoses are present in 30% of persons 60 to 69 years old.³⁴ Thus, the prevalence of coronary calcifications correlates better with the prevalence of atherosclerotic plaque than with coronary events^{35,36} The prevalence of cardiac calcifications is increased with ESRD, especially after long-term dialysis

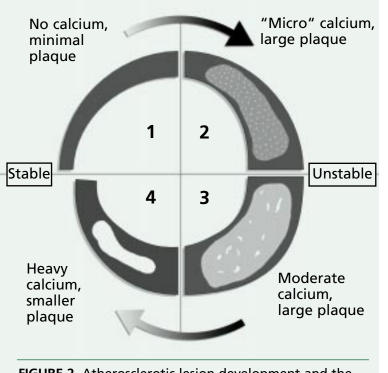


FIGURE 2. Atherosclerotic lesion development and the role of calcification. Different phases of plaque development are shown in the four quadrants, indicating temporal development. Calcium's role in lesion instability is complex and incompletely understood.

or angiographically severe stenosis.

Atherosclerotic plaque and coronary calcifications are frequently present in asymptomatic persons. While the overall plaque burden may predict cardiovascular risk, only a small proportion of persons with atherosclerosis and detectable coronary calcium will eventually experience clinical coronary events.

Plaque vulnerability and acute coronary syndromes

Traditional models of coronary artery disease described slow, progressive plaque growth with increasing passive calcification, eventually leading to vessel occlusion and acute coronary syndromes. According to these models, the amount of calcification in individual lesions should be directly related to the risk of these lesions causing ischemic events. These advanced, calcified plaques were often compared to "rusty pipes."

More recent vascular biology studies show

that this analogy is incomplete and even misleading.^{37,38} Several angiographic studies show that the progression of coronary artery disease in humans is neither linear nor predictable.^{39–42} It has become apparent that sudden, episodic changes of mildly stenotic coronary plaques residing in the vessel wall are most important in disease progression.⁴³ Most acute coronary events result from rupture of these "vulnerable" plaques, which often accompany more advanced atherosclerotic lesions, and subsequent thrombosis.³⁷ These vulnerable lesions may account for as many as two thirds of cases of unstable angina or other acute coronary syndromes.

"Plaque vulnerability" describes the tendency of atherosclerotic lesions to cause acute coronary syndromes. Vulnerable lesions are characterized by an accumulation of inflammatory cells and the formation of a lipid-rich, necrotic core separated from the lumen by a fibrous cap.^{43–45} The relatively large size of these atheromas is not well reflected by luminal stenosis because adaptive arterial enlargement maintains lumen size in spite of increasing plaque burden. This compensatory vessel enlargement in response to plaque growth is termed positive arterial remodeling^{46–48} and appears to be associated with development of acute coronary syndromes.^{49–52}

The junction between the necrotic core of the plaque and the normal vessel wall (plaque shoulder) is a location of high stress that is predisposed to rupture.^{53,54} Local secretion of proteolytic enzymes (such as matrix metalloproteinases and myeloperoxidase) by smooth muscle cells and macrophages contributes to degradation of the intercellular matrix of the fibrous cap, initiating plaque rupture.^{55,56}

As described above, coronary artery wall calcification is part of the development of atherosclerosis. The relation of coronary calcifications to the probability of plaque rupture is unknown, and plaques vulnerable to rupture or erosion are frequently not calcified.^{17,57} In fact, intravascular ultrasonography indicates that vulnerable plaques are most often not calcified.^{49,50,58,59} and that calcification is associated with plaques causing stable rather than unstable coronary syndromes.

It has been hypothesized that early micro-

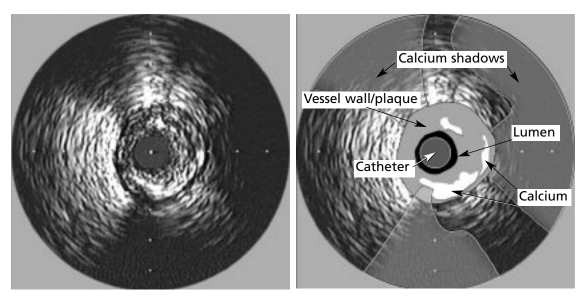


FIGURE 3. Intravascular ultrasound image of a highly stenotic, calcified coronary lesion.

calcifications near the junction of the plaque and the adjacent normal intima may lead to increased stress at the interface between calcified and noncalcified atherosclerotic sections, which could facilitate plaque rupture. However, more extensive calcification and fibrosis of the vessel could eventually eliminate these weak points and reduce the risk of rupture. Biomechanical data support the concept that calcified lesions are much stiffer than cellular lesions and are unlikely to be associated with sites of plaque rupture.⁶⁰ According to these concepts, calcifications could in fact represent an attempt to stabilize weakened atherosclerotic plaque prone to rupture.

METHODS FOR IDENTIFYING LESION CALCIFICATIONS

Intravascular ultrasonography

Intravascular ultrasonography, performed during cardiac catheterization, provides tomographic images of the vessel wall that demonstrate vessel size, plaque size, and plaque morphology.^{61,62} A miniature ultrasound catheter is placed beyond the target lesion site and is then withdrawn during continuous imaging, resulting in a series of cross-sections. The vessel wall of each cross-section can be described by its signal characteristics on a continuum from echodense (bright echo signal) to echolucent (faint echo signal). Several studies demonstrate the reliability of ultrasound imaging in predicting the composition of atherosclerotic plaque relative to histology.^{58,59} Calcified tissues are recognized as bright echoes with a characteristic signal shadow (**FIGURES 3 AND 4**).

Ultrasound imaging shows significant superiority over fluoroscopy or angiography in detecting coronary calcification.⁶³ The severity of calcification has been quantified according to the angle subtended by the calcified arc of the vessel wall.^{64,65} When calcium was detected angiographically, the calcification detected by ultrasound was likely greater than 90 degrees.⁶⁴ The image characteristics of microcalcifications, as described above, are incompletely understood.⁵⁷

Computed tomography

Computed tomography (CT) is very sensitive in detecting and quantifying coronary artery calcifications and can survey the entire coronary tree noninvasively. Computed tomography techniques are described more fully in an accompanying article in this supplement.⁶⁶ Briefly, different calcium scoring algorithms, including the traditional Agatston score,⁶⁷ the total calcium volume score,^{68,69} and calcium mass,^{70,71} can be applied to either electronbeam CT or mechanical CT images and provide a measure of total coronary plaque burden.^{72,73} The prognostic value of this informaOnly a small share of patients with atherosclerosis and coronary calcium will experience coronary events

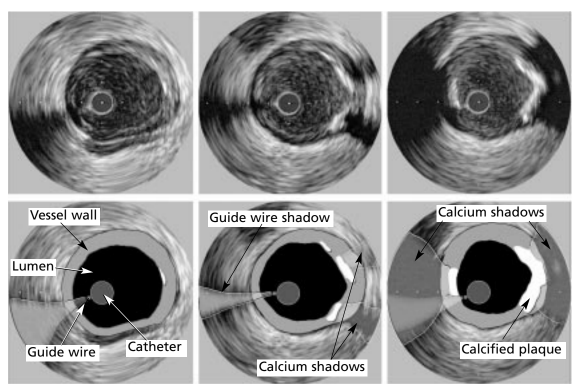


FIGURE 4. Intravascular ultrasound images of a mildly stenotic plaque with calcification. Three adjacent images of the same lesion are shown.

Reliable identification of vulnerable plaques is currently not possible

tion has been examined in several studies⁷⁴ in individuals without chronic renal failure, as discussed in detail elsewhere in this supplement.⁶⁶ Because compensatory vessel enlargement (positive remodeling) allows plaque accumulation without stenosis, the correlation of calcium area with luminal dimension is only moderate.^{75,76}

Coronary angiography

Coronary angiography is not a sensitive method for evaluating calcification compared with intravascular ultrasound and CT.^{63,64,77–79} The coronary angiogram shows a silhouette of the vessel lumen but not the vessel wall and plaque.^{38,80} Information on calcifications of the vessel wall is limited, although the extent of fluoroscopic calcifications is a marker of the overall atherosclerotic disease burden and does have prognostic value.^{81–83}

Magnetic resonance imaging

Magnetic resonance imaging (MRI) is a noninvasive technique that differentiates tissue structures on the basis of their proton magnetic properties. A wide range of image contrast can be obtained with different pulse sequences, which is more helpful for differentiation of soft-tissue structures than for calcifications.^{84–86} Shinnar et al⁸⁶ described the diagnostic accuracy of MRI for plaque characterization and emphasized the significant technical improvements in resolution and gating that are needed before this technique can be used to examine coronary arteries in clinical settings.

CLINICAL IMPLICATIONS OF CORONARY CALCIFICATION IN ESRD

Uncertain value for risk assessment

Noninvasive quantification of coronary calcium with electron-beam CT calcium scores has been shown to reflect the total atherosclerotic plaque burden,^{17,35,36} at least in individuals without chronic renal failure. For this reason, CT can help to predict future risk and offers the potential, through serial imaging, to follow disease progression, stabilization, and possible regression.^{87,88} This information can help guide the aggressiveness of risk-factor modification.

However, according to a current consensus statement of the American Heart Association and American College of Cardiology,⁸⁹ the incremental value of calcium scores over "traditional" multivariate risk-assessment models has not yet been established. This consensus statement does not recommend CT screening of coronary calcification for asymptomatic individuals, but it concludes that such screening may be justified in select patient groups with intermediate risk (ie, those in whom CT results could change the aggressiveness of risk-factor modification).

Applying these consensus recommendations to patients with chronic renal failure or ESRD is particularly difficult. As discussed above, these patients are already in a high-risk group and aggressive risk-factor modification may be justified independent of the CT result. It is conceivable that asymptomatic young patients with moderate renal failure could fall into an intermediate-risk group. This needs to be examined in further studies, especially since current guidelines for lipid management do not provide specific recommendations for patients with ESRD.

Vulnerable plaques hard to pinpoint

It is important to understand that calcium scoring has prognostic value but does not localize areas of vulnerable plaque. Studies in individuals without chronic renal failure show that coronary calcium scores correlate with atherosclerotic plaque burden. However, while more calcifications (ie, a higher calcium score) may be associated with a greater number of vulnerable plaques overall, coronary calcifications of individual lesions are not markers of lesion vulnerability and are often found in stable patients.⁹⁰ Currently, reliable identification of vulnerable plaques is not possible,⁹¹ but preliminary results with intravascular ultrasound⁹² and contrast-enhanced multislice CT93,94 demonstrate the potential role of these tomographic imaging techniques.

Role of medications

The derangements of calcium, phosphate, vitamin D, and parathyroid hormone in patients with chronic renal failure or ESRD are characterized by extraosseal calcifications (FIGURE 1), including cardiac calcifications. The high prevalence of coronary calcification in ESRD occurs in a clinical context far removed from the extensive studies done on coronary calcifications and cardiac risks in persons without chronic renal failure. This context includes the common use of calcium, calcium-containing phosphate binders, and pharmacologic doses of vitamin D in a setting where renal clearance of calcium and phosphate is markedly reduced or absent.

As a result, questions arise about the effect of these medications on coronary calcifications and how to interpret calcifications in this markedly different patient population. For example, excessive intake of vitamin D or its metabolites and analogues may lead to arterial calcifications,^{95,96} but appropriate doses of vitamin D metabolites given to control secondary hyperparathyroidism might actually reduce the propensity for vascular calcification.⁹⁷

A direct interaction between these medications and coronary calcifications has not been consistently shown. In addition, there is no evidence of an increased risk of coronary calcifications independent of atherosclerotic atheroma burden. Therefore, the pharmacologic control of calcium and phosphate metabolism should be directed by the patient's nephrologic and endocrinologic needs and not be withheld for concern about coronary calcifications. The cardiologist should emphasize aggressive cardiovascular risk-factor modification to slow progression of coronary artery disease in patients with ESRD. This includes tight control of hypertension, diabetes mellitus, and lipid abnormalities. Unfortunately, current lipid management guidelines do not specify a particular strategy for patients with ESRD.

CONCLUSION

Studies in subjects without chronic renal failure indicate that coronary calcifications are a manifestation of coronary atherosclerosis and develop in an actively controlled mineralization process. Their quantity correlates with the overall extent of atherosclerotic disease burden. The role of coronary calcifications in Guidelines for lipid management do not specify a strategy for patients with ESRD the development of acute coronary syndromes is complex and incompletely understood. Calcification of individual lesions may not be a marker of lesion instability. However, the presence of calcified lesions implies the likely association of lipid-rich and possibly unstable plaque.

Although calcified coronary lesions are more common in patients with ESRD, no independent increase in cardiovascular risk

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has been associated with these calcifications. Current evidence does not support the concept that specific attempts to reduce coronary calcium may benefit patients with renal failure. On the other hand, these patients have a high risk of developing coronary artery disease and are candidates for aggressive risk-factor modification, including control of hypertension, diabetes mellitus, and hypercholesterolemia.

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