Risk factors of one year increment of coronary calcifications and survival in hemodialysis patients

Giorgio Coen*1, Andrea Pierantozzi1, Daniele Spizzichino1, Daniela Sardella1, Daniela Mantella2, Micaela Manni3, Luigi Pellegrino4, Andrea Romagnoli4, Roberta Pacifici5, Piergiorgio Zuccaro5 and Salvatore DiGiulio2

Abstract

Background: Heart and coronary calcifications in hemodialysis patients are of very common occurrence and linked to cardiovascular events and mortality. Several studies have been published with similar results. Most of them were mainly cross-sectional and some of the prospective protocols were aimed to evaluate the results of the control of altered biochemical parameters of mineral disturbances with special regard to serum calcium, phosphate and Ca x P with the use of calcium containing and calcium free phosphate chelating agents. The aim of the present study was to evaluate in hemodialysis patients classic and some non classic risk factors as predictors of calcification changes after one year and to evaluate the impact of progression on survival.

Methods: 81 patients on hemodialysis were studied, with a wide age range and HD vintage. Several classic parameters and some less classic risk factors were studied like fetuin-A, CRP, 25-OHD and leptin. Calcifications, as Agatston scores, were evaluated with Multislice CT basally and after 12-18 months.

Results: Coronary artery calcifications were observed in 71 of 81 patients. Non parametric correlations between Agatston scores and Age, HD Age, PTH and CRP were significant. Delta increments of Agatston scores correlated also with serum calcium, C x P, Fetuin-A, triglycerides and serum albumin. Logistic regression analysis showed Age, PTH and serum calcium as important predictors of Delta Agatston scores. LN transformation of the not normally distributed variables restricted the significant correlations to Age, BMI and CRP. Considering the Delta Agatston scores as dependent, significant predictors were Age, PTH and HDL. A strong association was found between basal calcification scores and Delta increment at one year. By logistic analysis, the one year increments in Agatston scores were found to be predictors of mortality. Diabetic and hypertensive patients have significantly higher Delta scores.

Conclusions: Progression of calcification is of common occurrence, with special regard to elevated basal scores, and is predictive of survival. Higher predictive value of survival is linked to the one year increment of calcification scores. Some classic and non classic risk factors play an important role in progression. Some of them could be controlled with appropriate management with possible improvement of mortality.

Background

Coronary calcifications are of frequent occurrence in chronic renal failure both in the conservative [1,2] and hemodialysis stages [3]. They are considered to be responsible for increased cardiac events and mortality [4], since the extent of coronary calcium deposits are associated with decreased survival [5]. Calcifications are of two types, affecting media and intima. Medial artery calcifications are known to be more frequently found in diabetes mellitus and uremia. Intimal calcifications are located in the atherosclerotic plaques and denote the presence of atherosclerotic changes of the arteries [6]. However, with the currently available non invasive means, either Electron Beam or Multislice Computerized Tomography, there is no possibility to distinguish between intimal and medial calcifications. Most of the clinical studies of uremic vascular calcifications, mainly devoted to frequency of occurrence and to risk factors enhancing calcium deposits [7-10], were based on cross-sectional data. It has been found by several authors that most of patients on hemodialysis, about 80%, are affected
by coronary calcifications while only 20% or less are unaffected [3,8]. Patients showing unaffected arteries remain generally free of calcium deposits when followed for long periods of time. These patients are probably protected from calcifications thanks to inhibitory factors which have been in part identified [11,12]. Above all, age and HD age are important risk factors, together with serum phosphate, serum calcium and Ca × P product. Also PTH has been considered in many studies as a risk factor of coronary calcification [13] and mortality [14-16], with special regard to cases of severe hyperparathyroidism. As previously stated, most of the studies to identify risk factors published to date are of a cross-sectional type, not statistically strong enough to show a cause-effect relationship, while only a few were prospective [17-20]. In three of them mere progression of calcification scores was evaluated [17,19,20], while in another [18] the study was conducted to evaluate the effect on progression of phosphate chelating drugs. In the present study hemodialysis patients have been evaluated by Multislice CT basally and after 12-18 months. The relationship between clinical data, basal hormonal values and calcification scores, and progression of heart calcifications and mortality were evaluated in order to establish relevant risk factors and their linkage to fatal events.

Methods
Eighty-one unselected patients with end-stage renal disease on hemodialysis were studied to evaluate the evolution of coronary and heart calcification scores. This study, due to its observational nature, has not required ethical approval. Samples were collected as part of the standard care. The patients had a mean age of 59.2 ± 10.4 years and hemodialysis vintage of 82.5 ± 99.5 months. Sex distribution was 54 M/27 F. 8 patients had diabetes mellitus and 38 patients were affected by arterial hypertension. Causes of renal failure were: chronic glomerulonephritis in 19 patients; hypertension/ischemic nephropathy in 25 patients; polycystic kidney disease in 7 patients; diabetic nephropathy in 6 patients; renal malformations in 2 patients; and unknown in 22 patients. Exclusion criteria included ethanol or drug abuse, malignancy, clinical cardiovascular disease, chronic inflammatory disease, human immunodeficiency virus infection, use of steroids, antiepileptic drugs, non-steroidal anti-inflammatory drugs, estrogen and anticoagulants. All patients were treated with hemodialysis. Hemodialysis was based on standard bicarbonate dialysis using Cuprophan or Low-Flux-PolySulfone (LF-PS) membranes, 20% with acetate-free biofiltration (AFB) using Polycrylonitrile-AN69 (PAN-AN69 ST), or with hemodiafiltration using Helix- one (FX80) or Polycarbonate (Spiraflo SG 8 Plus). The dialysis sessions lasted 4 hours and were delivered at constant blood and dialysate flow rate values of 300 and 500 ml/min, respectively. Calcium concentration in the dialysate was 1.25, or 1.5 or 1.75 mmol/L (only 3 pts).

Most patients had been treated with relatively limited doses of calcitriol administered orally (1.5 μg/wk) or intravenously (calcitriol 4.5 μg/wk), but had discontinued this treatment at the time of the study. However, 17 patients were on IV calcitriol treatment with weekly doses ranging from 3 to 6 μg and 8 patients were treated with paricalcitol, 10-16 μg per week. Calcitriol/Paricalcitol doses were adjusted to comply with KDIGO guidelines. 81% of the patients were on phosphate binders, mainly calcium salts (calcium carbonate 1.9 ± 0.8 g/day) and sevelamer (6200 ± 2010 mg/day). In the majority of patients, regular intravenous erythropoietin treatment was underway.

Blood samples for biochemical evaluations were drawn prior to a dialysis session. Serum samples were stored at -30°C until the assays. The following assays were made: serum calcium, phosphorus, alkaline phosphatase, intact PTH, serum albumin, cholesterol, triglycerides, Fetuin-A, CRP, 25-OHD and Leptin.

Serum calcium, phosphorus, alkaline phosphatase and other biochemical parameters were measured by colorimetric methods using a Roche autoanalyzer or other standard laboratory techniques. Total calcium was corrected for serum albumin using the equation: Calcium = Ca + 0.8 (4.0 - albumin, g/dl). Normal values of these variables are reported in Table 1. Serum intact PTH was measured by an IRMA (Nichols Institute Diagnostic, San Juan Capistrano, CA). Both 1-84 and "7-84" PTH species are measured together. The normal range of values is 10 to 65 pg/ml. Fetuin-A was measured with an ELISA kit (Epitope Diagnostics Inc., San Diego). Leptin concentrations were determined with a radioimmunoassay for human leptin (Mediagnost, Reutlingen, Germany), which uses a high affinity polyclonal antibody specific for this protein. Serum 25-hydroxycholecalciferol was measured with a radioimmunoassay provided by IDS (Boldon, UK), following an extraction procedure with acetonitrile. The assay measures both 25-OHD$_3$ and 25-OHD$_2$.

Multislice computed tomography (MSCT) was performed basally and after 12-18 months, with a 64-channel multidetector scanner (LightSpeed VCT, General Electric Medical Systems, USA). A retrospective gating technique was used to synchronize the data reconstruction with the ECG signal. Mean heart rate during MSCT was 70 ± 8 beats/min (range 50 to 85). All scans were performed with the following parameters: detector collimation 64 × 2.5, reconstruction interval 160 mm, gantry rotation time 0.35 s, tube voltage 120 K, tube current 300 mAs effective, field of view (FOV) 25 cm, medium smooth convolution filter, acquisition volume 12 cm (i.e. pulmonary artery bifurcation to diaphragm), cranio-caudal scan direction. Images were obtained during a single breath-
**Table 1: Baseline Clinical, Biochemical Data, Calcification Scores and Normal Values**

| Parameter                  | Mean   | Median | SD    | Interquartile range | Normal value |
|----------------------------|--------|--------|-------|---------------------|--------------|
| Number                     | 81     |        |       |                     |              |
| Gender (M/F)               | 54/27  |        |       |                     |              |
| Age, years                 | 58,7   | 10,8   |       |                     |              |
| HD Age, months             | 45     | 99,5   |       |                     |              |
| BMI                        | 23,4   | 4      |       |                     | 18-24*       |
| Calcium, mg/dl             | 9,3    | 1      |       |                     | 8,5-9,5      |
| Phosphate, mg/dl           | 5,4    | 1,4    |       |                     | 3,5-5,5      |
| Albumin, g/dl              | 3,9    | 0,4    |       |                     | 3,5-5,2      |
| Ca, corrected, mg/dl       | 9,3    | 1,1    |       |                     | 8,5-9,5      |
| PTH, pg/ml                 | 300    | 399,9  | 10-65 |                     |              |
| Cholesterol, mg/dl         | 149    | 41,2   |       |                     | 150-210      |
| HDL, mg/dl                 | 43     | 14     |       |                     | 35-65        |
| Cholesterol/HDL            | 103,6  | 45,6   |       |                     | 3,1-4,7      |
| LDL, mg/dl                 | 82,6   | 32,4   |       |                     | 60-150       |
| Triglycerides, mg/dl       | 140    | 142    |       |                     | 50-180       |
| Haemoglobin, g/dl          | 11,7   | 1,2    |       |                     | 13,5-15      |
| Kt/V                       | 1,3    | 0,3    |       | ≥1,2                |              |
| Fetuin-A, g/L              | 0,5    | 0,2    |       |                     | 0,5 - 1,5    |
| CRP, mg/dl                 | 1      | 4,2    |       |                     | 0 - 3        |
| Alkaline Phosphatase, U/L  | 149    | 117,3  | 35 - 125 |                  |              |
| 25 OHD, ng/ml              | 19,3   | 9      |       |                     | 20-40        |
| Leptin, ng/ml              | 6,7    | 8,7    |       |                     | 2,5-15       |
hold of approximately 5 to 7 seconds. Image reconstruction was performed using three windows at 70, 75 and 80% of the time between two consecutive R-R intervals, which corresponded to the end-diastolic phase of the cardiac cycle. The phases with minimal motion were selected and used for calculating Calcium scoring.

All MSCT data were transferred to a GE AW Workstation (software version 4.2, General Electric Medical Systems, USA) for post-processing. Calcium scoring was performed on the reconstructed image sets with commercially available software (SmartScore™ 4.0, General Electric Medical Systems, USA) [21]. Calcium was scored according to the Agatston method to quantify the amount of calcification in the coronary arteries. A focus of coronary calcium was defined as the presence of four or more contiguous pixels with more than 130 Hounsfield units. The total calcium score was calculated from the sum of the individual scores of the four major epicardial coronary arteries. Informed consent was obtained from all patients.

The statistical evaluation was carried out using a personal computer equipped with a statistical package (SPSS for windows, Chicago, Illinois, release 13.0). In addition to descriptive statistics for the selected variables, Kolmogorov-Smirnov test was employed for the continuous variables to compare the observed cumulative distribution function with the normal distribution. Non parametric (Spearman) correlations were evaluated. Agatston score was measured basally and after 12-18 months. All values were normalized to 365 days, assuming a linear trend of calcification over time. The patient’s survival was followed for additional 2 years. The increment in Agatston score observed between baseline and one year was called delta. Logistic models, using tertiles of the dependent variables (coronary Agatston score), were performed to examine the relationship between the coronary calcium scores (baseline and delta) and clinical and biochemical parameters. In the logistic regression models, only those independent variables that showed to have statistically significant Spearman correlation with the dependent variable were included. Although statistically associated in bivariate analysis, some variables were excluded by logistic models because of collinearity problems. Cox regression analysis, stepwise method, has been used to investigate the association between Agatston score and mortality. The Cox regression was adjusted for age, gender, HD age, calcium and PTH. In case of OR well below unity, we used a scientific notation, reporting only the exponent (E) of base 10. A p-value < 0.05 was considered statistically significant.

Results
Clinical characteristics, serum biochemistry and calcification scores of the patients are reported in Table 1. In case of normally distributed variables, mean and standard deviation were shown, otherwise median and interquartile range were reported. Coronary calcifications were found in 71 of 81 patients. In table 2, left side, the non parametric correlations between the basal calcification scores and the basal clinical and humoral parameters are reported. Several parameters are found to be significantly correlated to the basal scores, like Age, HD Age, PTH and CRP. On the right side of Table 2, the logistic regression’s output adjusted for gender is reported, considering the tertiles of basal Agatston scores as dependent variables. The OR is referred to the high tertile compared to the low tertile as reference category. Cut-off values of Agatston score tertiles are: basal score, 157,6 and 1273; delta score, 12,2 and 239. Age (p = 0.001) and HD Age (p = 0.013) are found to be significant predictors of the basal Agatston scores. Which variables have been found to predict the value of the scores after one year? Firstly, we

Table 1: Baseline Clinical, Biochemical Data, Calcification Scores and Normal Values (Continued)

| Therapy: Calcium carbonate or Sevelamer | 81,4% |
|-----------------------------------------|-------|
| Hypertension, yes                       | 47,4% |
| SIST, mmHg                              | 122,4 | 19,2 |
| DIAST, mmHg                             | 71,2  | 11,1 |
| Coronary Agatston Score t0              | 481   | 1782,5 |
| Coronary Agatston Score t1              | 528,2 | 2405,9 |

T0, baseline scores; T1, one year scores
* Non CKD normal values
observed a strong association between basal calcification scores values and the delta increment after one year \((p = 0.001)\): a low basal score is associated with a low delta, while high basal score with a high increment. Then, the answer to the question is reported in Table 3 and 4, where correlations between basal values of parameters and delta scores are reported (left sides). The logistic regressions presented in those tables have been adjusted for HD age and gender. As shown in Table 3, age \((p = 0.042)\), calcium \((p = 0.047)\) and PTH \((p = 0.027)\) are significant predictors in calculating the risk of increased Agatston score after one year, while Fetuin \((p = 0.027)\) is a protective factor. In Table 4 the results of a logistic regression of three categorized delta Agatston score as dependent variable are reported: no delta coronary calcification (reference category), those with a medium increment (delta <1000) and those with severe (delta >1000) coronary calcification increment. In the model we considered as independent variables only those that resulted as statistically associated in logistic regression of table 3. Calcium \((p = 0.036)\) is a risk factor in moderate delta increment of Agatston score, but for severe calcification other variables are statistically significant: age \((p = 0.036)\), PTH \((p = 0.030)\) and, as protective factor, Fetuin-A \((p = 0.023)\).

Table 3: Non parametric significant correlations and logistics regressions, adjusted for gender, of basal Agatston score

| Variables  | Spearman | p-value | OR*  | C.I. 95%       | p-value |
|------------|----------|---------|------|----------------|---------|
| Age        | 0.358    | 0.001   | 1.136| 1.056 - 1.223  | 0.001   |
| HD Age     | 0.228    | 0.039   | 1.012| 1.002 - 1.021  | 0.013   |
| PTH        | 0.263    | 0.017   | 1.002| -              | 0.069   |
| CRP        | 0.326    | 0.018   | 1.400| -              | 0.106   |

* in logistic regression dependent variables are tertiles of basal Agatston score: high versus low (reference category). Pseudo R square (Cox and Snell) = 0.330

| Variables  | Spearman | p-value | OR*  | C.I. 95%       | p-value |
|------------|----------|---------|------|----------------|---------|
| Age        | 0.358    | 0.001   | 1.136| 1.056 - 1.223  | 0.001   |
| HD Age     | 0.228    | 0.039   | 1.012| 1.002 - 1.021  | 0.013   |
| PTH        | 0.263    | 0.017   | 1.002| -              | 0.069   |
| CRP        | 0.326    | 0.018   | 1.400| -              | 0.106   |

* in logistic regression dependent variables are tertiles of Delta Agatston Score: increment more than 239 versus increment up to 12 (reference category). Pseudo R square (Cox and Snell) = 0.505

**scientific notation
In Figure 1, a chart showing the surviving patients, stratified by tertiles of delta Agatston after one year is shown. There were 11 death among the patients, 5 due to cardiovascular events. 72.7% of the deaths were observed among those who showed a worsening of calcifications after one year. A Cox regression, stepwise method, has been carried out to evaluate the risk factors of mortality. Age, gender, HD age, PTH, calcium and Agatston score were considered as covariates. A statistically significant cumulative mortality hazard ratio for Agatston Score (p = 0.027) was found.

We have also performed parametric analysis and used statistical models, carried out using the transformation in LN for those variables not normally distributed. The results were similar to those of non-parametric analysis (not reported in the text).

Diabetic patients have higher levels Agatston scores after one year (p < 0.042), even if they do not differ from non diabetic patients in basal conditions. The same conclusions could be drawn for arterial hypertension. Delta Agatston score was significantly higher in hypertensive (p < 0.020) compared to non hypertensive HD patients.

**Discussion**

In the last 10 years intense scientific interest has been dedicated to the increased cardiovascular and all-cause mortality of patients with CKD, both on predialysis and dialysis stage [4]. It is known that most of the patients with renal insufficiency, at whatever stage, die mainly due to heart disease well before reaching the stage of hemodialysis [22]. There is an accelerated atherosclerosis in uremia [23] which is partly associated to calcification of the arteries, both in the intima and in the media. Calcification of the media is prevailing in uremia, as it is with diabetic patients. Calcification of arteries, with special regard to coronary arteries can be detected by radiological techniques, which however cannot distinguish between intimal and media calcifications. However, vascular calcification is very frequent in uremia and has a recognized association with worse survival prognosis [5]. In vivo studies have shown hyperphosphatemia to be an important risk factor of death [24], and in vitro observations have shown that increased phosphate concentrations are an important factor in the calcification of the smooth muscle cells of the media, which can be able to induce differentiation of these cells in osteoblast-like cells and to induce osteogenetic changes of arterial wall and calcification [25]. Many studies have evaluated the impact of appropriate phosphate control with chelating agents on the process of vascular calcification in hemodialysis patients and compared calcium containing chelating drugs to calcium-free drugs [18,26,27], like sevelamer and lanthanum carbonate. These studies have been conducted prospectively showing that calcium-free chelating

### Table 4: Logistics regressions, adjusted for gender and HD age, of categorized delta Agatston scores

| Variables | OR     | C.I. 95%  | p-value |
|-----------|--------|-----------|---------|
| delta increment < 1000 | | | |
| Age       | 1.044  | -         | 0.363   |
| Calcium   | 3.716  | 1.089 - 12.678 | 0.036 |
| PTH       | 1.003  | -         | 0.132   |
| Fetuin-A  | 0.015  | -         | 0.059   |
| delta increment > 1000 | | | |
| Age       | 1.178  | 1.010 - 1.373 | 0.036 |
| Calcium   | 1.550  | -         | 0.584   |
| PTH       | 1.005  | 1.001 - 1.010 | 0.030 |
| Fetuin-A  | 0.001  | 4.1 E-07** - 0.336 | 0.023 |

Reference category: delta increment equal to zero
Pseudo R square (Cox and Snell) = 0.499
**scientific notation
agents induce a lower increase of vascular calcification with time, compared to calcium containing chelating drugs [18]. Other studies have been planned and published to evaluate which are the major risk factors of vascular calcifications, mainly with cross-sectional protocols [9,10,28-30]. In addition to classical risk factors linked to deranged mineral metabolism, other non classical factors of vascular calcification and mortality have been identified, like Fetuin-A, Osteoprotegerin and FGF 23 [10,11,29]. Only a few studies have been conducted prospectively to evaluate progression of calcification in HD patients [17,19,20]. This study, made on a cohort of 81 hemodialysis patients, is a statistically more significant prospective analysis of the evolution of calcium deposits. The study strengthens the importance of several classical risk factors considered markers of mineral metabolism derangements, like serum calcium and PTH, and also Age, HD vintage and CRP. In addition, the study underlines the protective effect of Fetuin-A. Serum phosphate did not reach a statistically significant level, probably due to a limited range of values. In the evaluation of predictors of calcification progression, Age, serum calcium, PTH and Fetuin-A were selected as significant variables. The importance of PTH as a risk factor of vascular calcification is in apparent contrast with the histological findings of London et al [30], but in line with translational studies [31-33] and a clinical contribution [13]. In addition, the study highlights the increased mortality in the patients with higher calcification score progression and also validates earlier findings of a higher risk of progression in the patients with higher basal calcification scores [20]. Our results show that a higher increase of calcification after one year is associated to increased mortality. Monitoring progression of calcifications is of value in the survival evaluation of patients.

This study, in spite of the longitudinal approach, has some limitations mainly due to the relatively limited number of patients in the tertile groups. In addition the assumption was made that the progression of calcification deposits in the arteries was linear with time in the assessment of coronary calcium increment after one year. We used a retrospective gating in the evaluation of the score, due to frequent heart rhythm disorders in our patients, preventing the use of beta-blockers.

Conclusions
This study of 81 hemodialysis patients examines which factors are predictive of basal coronary calcification and of their progression. Age and HD vintage, PTH and CRP are significant factors of coronary calcifications on the bivariate analysis while regression analysis selected Age and HD age. The increment of calcifications after one year was predicted by Age, serum Calcium, PTH and Fetuin-A. Categorized delta Agatston scores selected Age, serum Calcium, PTH and Fetuin-A. The delta increment of coronary calcification was predictive of mortality. As a further deduction, there are factors of progression which cannot at present be controlled, like Age and Fetuin-A, and other risk factors which could be brought into control with appropriate treatment with a potential improvement of expected survival.
Competing interests
There are no conflicts of interest or financial or non-financial competing interests for this manuscript.

Giorgio Coen, MD

Authors' contributions
GC, design of protocol, evaluation of data, draft of the manuscript. AP, evaluation of data and statistical analysis. DS, contribution to study design and statistical analysis. LS, laboratory routine and immunoassays. DM, clinical investigation and collection of data, preparation of database. MM, clinical investigation and collection of data. LP, CardioTC x-ray specialist. AR, Cardio TC, calcium scoring. RP, laboratory biochemical assays. PZ, laboratory immunoassays. SD, contribution to study design, collaboration in drafting the manuscript. All authors read and approved the final manuscript.

Acknowledgements
This study has been supported by research funds of Ospedale Israelitico, Rome

Author Details
1Nephrology and Hypertension Unit, Ospedale Israelitico, Rome, Italy, 2Department of Nephrology and Dialysis, Forlanini-San Camillo Hospital, Rome, Italy, 3Department of Nephrology and Dialysis, San Giovanni Hospital, Rome, Italy, 4Department of Radiology, Policlinico Tor Vergata, Rome, Italy, and 5Istituto Superiore di Sanità, Rome, Italy

Received: 22 September 2009 Accepted: 21 June 2010
Published: 21 June 2010

References
1. Russo D, Palmiero G, De Blasio AP, Ballestra MM, Andreucci VE: Coronary artery calcification in patients with CRF not undergoing dialysis. *Am J Kidney Dis* 2004, 44:1024-1030.
2. Kramer H, Toto R, Peshock R, Cooper R, Victor R: Association between chronic kidney disease and coronary artery calcification: The Dallas Heart Study. *J Am Soc Nephrol* 2005, 16:S57-S53.
3. Goodman WG, Goldin J, Kuizon BD, Gales B, Sider D, Wang Y, Chung J, Emerick A, Greaser L, Elashoff RM, Salusky IB: Coronary artery calcification in young adults with end-stage renal disease who are undergoing dialysis. *N Engl J Med* 2000, 342:1478-1483.
4. Foley RN, Parfrey PS, Samak MJ: Clinical epidemiology of cardiovascular disease in chronic renal disease. *Am J Kidney Dis* 1998, 32(Suppl 3):S112-S119.
5. Blacher J, Guerin AP, Pannier B, Marchais SJ, Guerin AP: Arterial stiffness, and cardiovascular risk in end-stage renal disease. *Hypertension* 2001, 38:938-942.
6. London GM: Cardiovascular calcifications in uraemic patients: clinical impact on cardiovascular function. *J Am Soc Nephrol* 2003, 14:5305-5309.
7. Barreto DV, Barreto FC, Carvalho AB, Cuppari L, Cendoroglio M, Drabe SA, Moyes RM, Jorgetti V, Blair A, Guiberteau R, Fernandes Canziani ME: Coronary calcification in haemodialysis patients: the contribution of traditional and uremia related risk factors. *Kidney Int* 2005, 67:1576-1582.
8. Coen G, Manni M, Agnoli A, Balducci A, Desi M, De Angelis S, Jankovic L, Mantella D, Morosetti M, Naticchia A, Norfino I, Romagnoli A, Gallucci MT, Tomassini M, Simonetti G, Splendiani G: Cardiac calcifications: fetuin-A and other risk factors in hemodialysis patients. *ASAOJ* 2006, 50:150-156.
9. Krasniak A, Drozdz M, Pasowicz M, Chmiel G, Michalek M, Szumlak D, Podolec P, Klimeczek P, Konieczynska M, Wicher-Muniak E, Tracz W, Khoa TN, Souberbielle JC, Drueke TB, Sulowicz W: Factors involved in vascular calcification and atherosclerosis in maintenance haemodialysis patients. *Nephrol Dial Transplant* 2007, 22:S51-S52.
10. Baber U, de Lemos JA, Khera A, McGuire DK, Orland T, Toto RD, Hedaya SS: Non-traditional risk factors predict coronary calcification in chronic kidney disease in a population-based cohort. *Kidney Int* 2008, 73:615-621.
11. Ketteler M, Bongartz P, Westenfeld R, Wildberger JE, Mahnken AH, Böhm R, Metger T, Wanner C, Jhanneh-Dechent W, Fioege J: Association of low fetuin-A (AHSG) concentrations in serum with cardiovascular mortality in patients on dialysis: a cross-sectional study. *Lancet* 2003, 361:827-833.
12. Moe SM, Røsleva M, Ketteler M, O'Neill K, Duan D, Koczman I, Westenfeld R, Jhanneh-Dechent W, Chen NX: Role of calcification inhibitors, in the pathogenesis of vascular calcification in chronic kidney disease (CKD). *Kidney Int* 2005, 67:2295-2304.
13. Coen G, Manni M, Mantella D, Pierantozzi A, Balducci A, Condo S, DiGiulio S, Yancovic L, Lipp B, Marca S, Morosetti M, Pellegrino L, Simonetti G, Gallucci MT, Splendiani G: ArePTH serum levels predictive of coronary calcifications in haemodialysis patients? *Nephrol Dial Transplant* 2007, 22:S526-S527.
14. Ganesh SK, Stack AG, Levin NW, Hulbert-Sharon T, Port FK: Association of elevated serum PO4, Ca × PO4 product and parathyroid hormone with cardiac mortality risk in chronic hemodialysis patients. *J Am Soc Nephrol* 2001, 12:2131-2138.
15. Slinn Y, Foley RN, Collins AJ: Calcium, phosphorus, parathyroid hormone, and cardiovascular disease in hemodialysis patients: the USRDS waves 1,3, and 4 study. *J Am Soc Nephrol* 2005, 16:1788-1793.
16. Covic A, Kothavala P, Bernal M, Robbins S, Chalian A, Goldsmith D: Systematic review of the evidence underlying the association between mineral metabolism disturbances and risk of all-cause mortality, cardiovascular mortality and cardiovascular events in chronic kidney disease. *Nephrol Dial Transplant* 2009, 24:1506-1523.
17. Braun J, Oldendorf M, Moshage W, Heidler R, Zeitler E, Luft FC: Electron beam computed tomography in the evaluation of calcium calcification in chronic dialysis patients. *Am J Kidney Dis* 1996, 27:594-601.
18. Chertow GM, Burke SK, Raggi P: for the Treat to Goal Working Group: Sevelamer attenuates the progression of coronary and aortic calcification in hemodialysis patients. *Int Kidney Med 2002*, 62:245-252.
19. Moe SM, O'Neill KD, Røsleva M, Fineberg N, Persohn S, Meyer CA: Natural history of vascular calcification in dialysis and transplant patients. *Nephrol Dial Transplant* 2004, 19:2387-2393.
20. Bellasi A, Kooienga L, Block GA, Veledar E, Spiegel DM, Raggi P: How long is the warranty period for nil or low coronary artery calcium in patients new to hemodialysis? *J Nephrol* 2009, 22:255-262.
21. Knez A, Becker A, Becker C, Leber A, Boekstegers P, Reiser M, Steinbeck G: Detection of coronary calcinosis with multislice spiral computerized tomography: an alternative to electron beam tomography. *J Cardiovasc Med* 2002, 91:642-649.
22. Keith DS, Nichols GA, Guillon CM, Brown JB, Smith DH: Longitudinal follow-up and outcomes among a population with chronic kidney disease in a large managed care organization. *Arch Intern Med* 2004, 164:659-663.
23. Lindner A, Chirra B, Sherrard DJ, Scribner BH: Accelerated atherosclerosis in prolonged maintenance hemodialysis. *N Eng J Med* 1974, 290:697-701.
24. Block GA, Hulbert-Sharon TE, Levin NW, Port FK: Association of serum phosphorus and calcium × phosphate product with mortality in chronic hemodialysis patients: a national study. *Am J Kidney Dis* 1998, 31:607-617.
25. Giacchelli CM: The emerging role of phosphate in vascular calcification. *Kidney Int* 2001, 72:890-912.
26. Chertov GM: Slowing the Progression of Vascular Calcification in Hemodialysis. *J Am Soc Nephrol* 2003, 14:S310-S314.
27. Block GA, Spiegel DM, Ehrlich J, Mehta R, Lindbergh J, Dreisbach A, Raggi P: Effects of sevelamer and calcium on coronary artery calcification in patients new to hemodialysis. *Kidney Int* 2005, 68:1815-1824.
28. Stomporf T, Pasowicz M, Sulowicz W, Dembinska-Kiec A, Janda K, Wójcik K, Tracz W, Židenčička A, Klimpka K, Janusz-Grzybowska E: An association between coronary artery calcification score, lipid profile, and selected markers of chronic inflammation in ESRD patients treated with peritoneal dialysis. *Am J Kidney Dis* 2003, 41:203-211.
29. Jean G, Bresson E, Terrat JC, Vanet V, Hurot JM, Lonriaux C, Mayor B, Chazot C: Peripheral vascular calcification in long-haemodialysis patients: associated factors and survival consequences. *Nephrol Dial Transplant* 2009, 24:948-955.
30. London GM, Martí-C, Marchais SJ, Guerin AP, Metivier F, DeVernejoul MC: Arterial calcifications and bone histomorphometry in end-stage renal disease. *Am J Nephrol* 2004, 19:1543-1591.
31. Neves R, Grazzoli FG, Dos Reis LM, Grazzoli RG, Neves CL, Pararthy AO, Custodio MR, Jorgetti V, Moyes RMA: Vascular calcifications:...
contribution of parathyroid hormone in renal failure. Kidney Int 2007, 71:1262-1270.
32. Tomiyama C, Higa A, Dalboni MA, Cendoroglio M, Draibe SA, Cuppari L, Carvalho AB, Neto EM, Canziani ME: The impact of traditional and non-traditional risk factors on coronary calcification in pre-dialysis patients. Nephrol Dial Transplant 2006, 21:2464-2471.
33. Price PA, Roublick AM, Williamson MK: Artery calcifications in uremic rats is increased by a low protein diet and prevented by treatment with ibandronate. Kidney Int 2006, 70:1577-1583.

Pre-publication history
The pre-publication history for this paper can be accessed here: http://www.biomedcentral.com/1471-2369/11/10/prepub

doi: 10.1186/1471-2369-11-10
Cite this article as: Coen et al, Risk factors of one year increment of coronary calcifications and survival in hemodialysis patients BMC Nephrology 2010, 11:10