Pregnancy-associated plasma protein A as a predictor of all-cause mortality and cardiovascular events in patients with chronic kidney disease: a meta-analysis of prospective studies

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Abstract

Introduction: The aim of the study was to assess the association of elevated serum pregnancy associated plasma protein A (PAPP-A) and the risk of all-cause mortality, cardiovascular events and mortality due to infection in patients with chronic kidney disease (CKD).

Material and methods: We systematically searched the Medline database up to March 2017. A random effects model was used to pool the relative risks (RRs) and their 95% confidence intervals (CIs). Sensitivity analysis and subgroup analysis were performed to explore the potential sources of heterogeneity.

Results: Six studies involving 2034 subjects were included. The pooled RRs for the risk of all-cause mortality and cardiovascular events were 1.50 (95% CI: 1.17–1.92), 1.26 (95% CI: 0.95–1.69), respectively. Sensitivity analysis by excluding each individual study showed no influence on the main results. Subgroup analysis showed that age, male proportion, follow-up term, and assay methods were not modifiable factors.

Conclusions: Our study suggests that elevated serum PAPP-A is associated with the risk of all-cause mortality in patients with CKD.

Key words: pregnancy-associated plasma protein A, all-cause mortality, chronic kidney disease, meta-analysis.

Introduction

Chronic kidney disease (CKD) is very prevalent, affecting 8–16% of the population globally. Patients with CKD suffer from high mortality and morbidity [1]. Previous studies have indicated that cardiovascular events are the leading cause of death for patients with CKD [2–4]. Current data show that CKD or end-stage renal disease (ESRD) and cardiovascular diseases share some common mechanisms, such as accelerated inflammation and atherosclerosis [5–7].

Pregnancy-associated plasma protein A (PAPP-A) is a newly discovered zinc-binding metalloprotein, which has been suggested to be
involved in the processes of inflammation and atherosclerosis [8–10]. PAPP-A primarily acts as a protease cleaving inhibitory binding proteins of insulin-like growth factors [11]. In vitro studies have indicated that proinflammatory cytokine tumor necrosis factor α and interleukin 1β are powerful stimulators in inducing the expression of PAPP-A [8, 10, 12]. In vivo studies have suggested that a mouse model with PAPP-A gene knock-out could resist the accelerated progression of atherosclerosis, whereas over-expression of the gene of PAPP-A in the mouse model could accelerate progression of atherosclerosis [8, 13]. Clinical studies have shown that PAPP-A is not only an acute and sensitive biomarker for the diagnosis of acute coronary syndrome [14–16], but also acts as a prognostic indicator of all-cause mortality and combined cardiovascular events for patients with coronary heart diseases [17–25], which was systematically reviewed in our previous meta-analysis [24].

It remains unclear whether elevated PAPP-A was a predictor in patients with CKD. Some previous studies have shown that PAPP-A is an independent risk factor associated with increased mortality and cardiovascular events, while others have indicated no association in patients with CKD [26–32]. So we combined all available prospective studies and conducted a meta-analysis to assess the association of elevated serum PAPP-A and the risk of all-cause mortality and cardiovascular events in patients with CKD.

Material and methods

Search strategy
We performed this meta-analysis according to the Meta-analysis of Observational Studies in Epidemiology (MOOSE) guidelines [33]. Two authors (Yuehua Li and Xv Meng) identified articles through searching MEDLINE from 2000 to Mar 2017. The key word used in the search was “PAPP-A”. No language restriction was applied for searching and study inclusion.

Study selection and outcomes
The inclusion criteria were determined as follows: (1) prospective studies design; (2) provided the referent (the lowest) and highest levels of serum PAPP-A, or the cut-off value of PAPP-A; (3) provided the unadjusted or multivariable adjusted relative risks (RRs) and their corresponding 95% confidence intervals (CIs).

The primary outcome was all-cause mortality. The second outcome was cardiovascular events, which were defined by each individual study, including cardiovascular death, myocardial infarction, revascularization, sudden death, and so on.

Data extraction
Data extraction was conducted by two independent authors (Yuehua Li and Xv Meng). Discrepancies were resolved by group discussion. The extracted data included the name of the first author, publication year, sample size, number of events, male proportion, mean age, duration of follow-up, assay methods for measuring serum PAPP-A, cut-off value of serum PAPP-A, outcomes, adjusted covariates and RRs and their corresponding 95% CIs. We extracted the most fully adjusted RRs for the highest level of PAPP-A compared with the lowest one. If the multivariable adjusted RRs were not provided, we extracted the unadjusted ones.

Statistic analysis
We considered the hazard ratios or odds ratios as RRs in prospective studies. Compared with the lowest category of PAPP-A, the pooled RRs and their 95% CI of the highest category were estimated by a fixed-effect model to incorporate the inter-study variability. We extracted data from the highest quartile if provided by > 2 quartiles. If there was significant heterogeneity among studies, we used the random-effect model. The heterogeneity was assessed by the Q statistic, I² and p-value. I² > 50% was considered to indicate significant heterogeneity among the trials [34]. We tried to explore the potential source of heterogeneity by conducting sensitivity analysis and subgroup analysis. We performed the sensitivity analysis by removing each individual study to evaluate the study’s effects on the summary estimate. The subgroup analysis was performed by specified study characteristics, such as age (≥ 65 or < 65 years), male proportion (≥ 60% or < 60%), the follow-up term (≥ 5 or < 5 years), and assay methods (Time Resolved Amplified Cryptate Emission (TRACE) or others).

We assessed publication bias by Begg’s funnel plot and Egger’s regression test [35]. A two-sided p-value < 0.05 was considered to be significant. All of these analyses were completed by STATA software (10.0 version, Stata Corporation, TX, USA).

Results

Search results
We identified 1866 articles in the initial research. Among these studies, 1838 were excluded based on the title and abstract due to being experimental studies, reviews and non-relevant. Twenty-eight potential studies were selected for detailed accession. We further excluded 22 for the following reasons: not prospective design (n = 3), not relevant to the outcomes (n = 18), did not provide RRs or their 95% CI (n = 1). Finally, 6 potential prospective studies concerning 2034 participants.
and 792 events were included in our meta-analysis [26, 28–32]. Among them, 5 trials have reported the endpoint of all-cause mortality [26, 28–30, 32], and 4 have reported the endpoint of cardiovascular events [26, 29–31].

**Study characteristics**

The characteristics of the included studies are presented in Table I. All the studies were conducted in Europe. The sample size ranged from 130

| Study                  | Year | Country          | Men % | Mean age | Follow-up term | No. of subjects | No. of death | No. of CV events | Type of CKD            |
|------------------------|------|------------------|-------|----------|----------------|-----------------|--------------|------------------|-----------------------|
| Lauzurica et al. [31]  | 2005 | Spain            | 65.7  | 53.14    | 49.3 months    | 178             | NA           | 27               | Renal transplant      |
| Astrup et al. [26]     | 2007 | Denmark          | 70.88 | 61       | 10.1 years     | 197             | 48           | 41               | Diabetic nephropathy  |
| Etter et al. [32]      | 2010 | Switzerland      | 65.3  | 70       | 17.5 months    | 170             | 23           | NA               | Haemodialysis         |
| Kalousova et al. [29]  | 2012 | Czech Republic   | 54    | 63.2     | 5 years        | 261             | 146          | 71               | Haemodialysis         |
| Kalousova et al. [30]  | 2014 | Germany          | 55    | 66       | 4 years        | 1098            | 534          | 398              | Diabetic haemodialysis|
| Jefferies et al. [28]  | 2015 | Finland, UK      | 67    | 65.3     | 407 days       | 130             | 25           | NA               | Haemodialysis         |

| Study                  | Assay method                      | Cut-off value [mIU/l] | Outcomes                                           | Adjusted variables                                                                 |
|------------------------|-----------------------------------|-----------------------|----------------------------------------------------|-------------------------------------------------------------------------------------|
| Lauzurica et al. [31]  | DSL ELLISA                         | 1.82                  | Cardiovascular event, chronic allograft nephropathy| Age, sex, donor age and sex, type of dialysis, time in dialysis, number of HLA mismatched antigens, cold ischemia, immunosuppressive treatment, acute rejection, chronic allograft nephropathy |
| Astrup et al. [26]     | Biotintyram-amplified enzyme immunoassay | 12.6                | All-cause mortality                                | Age, cholesterol, HbA$_{1c}$, Cr                                                   |
| Etter et al. [32]      | TRACE                              | 24                    | All-cause mortality                                | Age, sex, number of comorbidities, dialysis vintage, K/V, IL-6, CRP, PTH, Ca$_{3}$PO$_{4}$ product, cholesterol                  |
| Kalousova et al. [29]  | TRACE                              | 30.8                  | All-cause mortality, infectious mortality          | Age, sex, hypertension, DM, smoking, BMI, CVD, IGF, MMP, transplantation, cTnI, BNP, albumin, cholesterol, CRP, orosomucoid, retinol, phosphate, PTH |
| Kalousova et al. [30]  | TRACE                              | 20.9                  | All-cause mortality, cardiovascular events, infectious mortality | Age, sex, smoking, BMI, atorvasatin, CVD, SBP, dialysis vintage, cholesterol, Cr, albumin, phosphate, hemoglobin    |
| Jefferies et al. [28]  | Immunofluorometric assays          | 3.45                  | All-cause mortality                                | Age, cTnT, dialysis vintage                                                        |

BMI – body mass index, BNP – B type natriuretic peptide, CKD – chronic kidney disease, Cr – creatinine, CRP – C-reactive protein, cTnI – cardiac troponin T, CVD – cardiovascular disease, DM – diabetes mellitus, DSL – Diagnostic Systems Laboratory, F – female, M – male, ELLISA – enzyme-linked immunosorbent assay, HLA – human leukocyte antigen, IGF – insulin-like growth factor, IL – interleukine, MMP – matrix metalloproteinase, NA – not available, PAPP-A – pregnancy-associated plasma protein A, PTH – parathyroid hormone, SBP – systolic blood pressure, TnT – troponin T, TRACE – Time Resolved Amplified Cryptate Emission.
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The average age range was from 53 to 70 years. The male proportion ranged from 46.7% to 70.9%. Four studies were performed in dialysis patients [28–30, 32], 1 study was performed in patients with diabetic nephropathy [26] and 1 study was performed in patients with ESRD waiting for kidney transplantation [31]. The duration of follow-up were varied from 3 months to 5 years. Serum PAPP-A was measured by TRACE in 3 studies [29, 30, 32], and by other methods in 3 studies [26, 28, 31]. PAPP-A values were obtained by a cut-off value in 4 studies [26, 29, 31, 32], by quartiles in 1 study [30] and the median value in 1 study [28] (Figure 1).

Main analysis

Comparing the group with lowest levels of serum PAPP-A, the pooled RR for all-cause mortality was 1.50 (95% CI: 1.17–1.92, \( p = 0.001; I^2 = 76.2\%\), \( p\) for heterogeneity = 0.002) (Figure 2 A). For the endpoint of cardiovascular events, the pooled RR was 1.26 (95% CI: 0.95–1.69, \( p = 0.113; I^2 = 65.8\%\), \( p\) for heterogeneity = 0.032) (Figure 2 B).

To clarify the heterogeneity, we performed sensitivity analysis by excluding each individual study at a time (Table II). The sensitivity analysis also showed that the positive relationship between el-

### Table I

| Study ID | RR (95% CI) | Weight (%) |
|----------|-------------|------------|
| Astrup et al. (2007) | 1.92 (0.96–3.82) | 9.38 |
| Etter et al. (2010) | 3.20 (1.06–9.58) | 4.42 |
| Kalousova et al. (2012) | 1.24 (1.06–1.44) | 29.99 |
| Kalousova et al. (2014) | 1.98 (1.35–2.20) | 25.57 |
| Jefferies et al. (2015) | 1.19 (1.03–1.37) | 30.64 |
| Overall (\( I^2 = 76.4\%\), \( p = 0.002\)) | 1.50 (1.17–1.92) | 100.00 |

### Table II

| Study ID | RR (95% CI) | Weight (%) |
|----------|-------------|------------|
| Lauzirica et al. (2005) | 6.40 (1.24–33.11) | 2.93 |
| Astrup et al. (2007) | 2.50 (1.20–5.50) | 11.24 |
| Kalousova et al. (2012) | 1.07 (0.84–1.36) | 37.36 |
| Kalousova et al. (2014) | 1.11 (1.02–1.20) | 48.47 |
| Overall (\( I^2 = 65.8\%\), \( p = 0.032\)) | 1.26 (0.95–1.69) | 100.00 |

Note: Weights are from random effects analysis.
Table II. Sensitivity analysis excluding one study at a time (all-cause mortality)

| Excluding study     | Year | Pooled RR (95% CI) | I² (%) | Pa   | Pb   |
|---------------------|------|--------------------|--------|------|------|
| Astrup et al.       | 2007 | 1.31 (1.20–1.45)   | 81.1   | 0.001| 0.000|
| Etter et al.        | 2010 | 1.32 (1.20–1.47)   | 79.3   | 0.002| 0.000|
| Kalousova et al.    | 2012 | 1.38 (1.22–1.55)   | 81.0   | 0.001| 0.000|
| Kalousova et al.    | 2014 | 1.23 (1.11–1.37)   | 36.2   | 0.20 | 0.000|
| Jefferies et al.    | 2015 | 1.47 (1.27–1.64)   | 76.8   | 0.005| 0.000|

Table III. Subgroup analysis

| Characteristic                  | Data points, N | Pooled RR (95% CI) | P for heterogeneity | P for subgroup difference |
|---------------------------------|----------------|--------------------|---------------------|---------------------------|
| All studies                     | 5              | 1.50 (1.17–1.92)   | 0.002               |                           |
| Age [years]:                    |                |                    |                     |                           |
| < 65                            | 2              | 1.34 (0.96–1.90)   | 0.223               | 0.44                      |
| ≥ 65                            | 3              | 1.68 (1.04 to 2.70)| 0.001               | 0.95                      |
| Male proportion:                |                |                    |                     |                           |
| < 60%                           | 2              | 1.60 (0.94–2.71)   | 0.001               | 0.15                      |
| ≥ 60%                           | 3              | 1.55 (0.97–2.46)   | 0.001               | 0.95                      |
| Follow-up term [years]:         |                |                    |                     |                           |
| < 5                             | 3              | 1.68 (1.04–2.70)   | 0.001               | 0.44                      |
| ≥ 5                             | 2              | 1.34 (0.96–1.90)   | 0.223               |                           |
| Assay for PAPP-A:               |                |                    |                     |                           |
| TRACE                           | 3              | 1.69 (1.09–2.64)   | 0.183               | 0.06                      |
| Other                           | 2              | 1.34 (0.89–1.99)   | 0.223               |                           |

PAPP-A – pregnancy-associated plasma protein A, RR – relative risk, TRACE – Time Resolved Amplified Cryptate Emission.

Figure 3. Begg’s funnel plot (with pseudo 95% CIs) of the studies in the meta-analysis. Studies that reported the elevated serum PAPP-A and the risk of all-cause mortality were plotted with lnRR on the vertical axis and the SEs of the lnRR along the horizontal axis.

PAPP-A – pregnancy-associated plasma protein A, RR – relative risk, SEs – standard errors.

Discussion

To the best of our knowledge, this is the first meta-analysis which has systematically reviewed
the association between circulating elevated PAPP-A and the risk of all-cause mortality and cardiovascular events in patients with CKD. In the meta-analysis of prospective studies, we found that elevated serum PAPP-A was associated with 1.50-folder higher risk of all-cause mortality for the patients with renal dysfunction. The relationship of serum PAPP-A and the risk of all-cause mortality was not modified by age, male proportion, follow-up term or the assay methods of serum PAPP-A. There seemed to be a positive association between elevated serum PAPP-A and cardiovascular events, but it was not statistically significant.

Previous studies about the relationship between circulating PAPP-A and prognosis of CKD patients are inconsistent. Some trials have reported a positive association between serum PAPP-A and the risk of death, while others have reported no association [26–32]. We combined all available trials and found a positive association of circulating PAPP-A and the risk of all-cause mortality. Our findings supported the previous meta-analysis which reported that elevated serum PAPP-A was an independent risk factor for all-cause mortality in patients with coronary heart disease [24]. Furthermore, our sensitivity analysis showed that the positive association of the serum PAPP-A and the risk of all-cause mortality was not influenced by excluding each individual study at a time, which strengthened the statistical power. CKD, especially ESRD, are known as a heavy burden with high mortality, morbidity and economic cost. Therefore, it is helpful to identify the biomarkers for risk stratification and prognosis analysis. All of the included trials have controlled numerous factors such as age, gender, serum cholesterol, creatinine and dialysis parameter. We used the most fully adjusted results and found that elevated serum PAPP-A was an independent risk factor for all-cause mortality in patients with CKD.

Heterogeneity is often a concern of a meta-analysis. Although there was significant heterogeneity among the included studies, the sensitivity analysis showed that the main results were not influenced. Furthermore, our subgroup analysis indicated that age, male proportion, follow-up term and assay methods for measuring PAPP-A were not modifiable factors. Nevertheless, large-scale prospective studies are needed to verify the potential factors influencing the association between circulating PAPP-A and all-cause mortality.

It was not fully understood about the elevated serum PAPP-A in predicting adverse events in CKD patients. First, clinical studies have reported that serum PAPP-A is not only correlated with serum level of creatinine, a marker of renal function, but is also correlated with other inflammatory factors, such as C-reactive protein, cytokine tumor necrosis factor α and interleukin 1β in patients with renal disease [31, 36–39]. Second, in vivo studies have indicated that the animal model with PAPP-A gene knock-out was less likely to suffer from nephropathy compared with the wild type in old age [40, 41]. The mouse model without PAPP-A gene expression could resist the development of diabetic nephropathy [42]. Third, in vitro studies have suggested that inflammation cytokine tumor necrosis factor α and interleukin 1β could stimulate PAPP-A gene expression in human mesangial cells, indicating that the kidney may be another target organ of PAPP-A [43]. Two included trials have reported that serum PAPP-A was associated with high risk of infectious mortality in CKD patients, which may be an important cause of death [29, 30]. Fourth, PAPP-A is an important metalloprotein involved in the progression of atherosclerosis [8]. Elevated serum PAPP-A was correlated with the severity of the plaque load [44]. Accelerated atherosclerosis is an important cause of the adverse prognosis in patients with CKD [2, 4, 45, 46]. Our meta-analysis indicated that circulating PAPP-A seemed to be associated with the risk of cardiac events in the CKD patients, but it was not statistically significant. Restricted by the number of included trials and heterogeneity, it needs further exploration in experimental studies and large-scale prospective studies for PAPP-A in predicting cardiovascular events in CKD patients in future.

Our study has a few limitations. First, the residual confounders may be potential confounders. Most included studies have adjusted for a wide range of potential confounders. However, we could not exclude other confounders which might have influenced the results. For example, serum PAPP-A level might be affected by heparin administration. Three included trials [28–30] have indicated that blood samples were collected before heparin administration while others did not. So it needs further evaluation whether heparin administration is a modifiable factor for the predictive value of serum PAPP-A. Previous history of coronary artery disease may also be a modifiable factor for the prognosis. We could not perform a subgroup analysis according to the percentage of previous history of coronary artery disease because only three studies have provided the relative data [26, 29, 30]. Second, restricted by the number of included trials, the predictive value of serum PAPP-A in cardiovascular events and all-cause mortality needs further exploration in future studies. Third, the results may be influenced by study design, but we did not find that the predictive value of PAPP-A was influenced by age, male proportion, duration of follow-up or assay methods. Fourth, all of the included studies had different cut-off values of serum PAPP-A. Given the nature of a meta-analysis, we could not provide the cut-off value of serum PAPP-A. Fifth,
necrosis factor \( \alpha \) and interleukin 1\( \beta \) are stimulators of PAPP-A. However, we could not compare serum PAPP-A with tumor necrosis factor \( \alpha \) and interleukin 1\( \beta \) about the predictive value because only two included trials have provided the relative data about tumor necrosis factor \( \alpha \) and interleukin 1\( \beta \). Finally, publication bias could be a concern, but we found no evidence of publication bias.

In conclusion, elevated serum PAPP-A was associated with higher risk of all-cause mortality in patients with CKD. The independent predictive value of PAPP-A was not affected by age, male proportion, duration of follow-up term or assay methods. How PAPP-A was involved in kidney disease and whether serum PAPP-A is a candidate biomarker for prognosis of renal dysfunction needs further exploration in numerous experimental and large-scale clinical trials.

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Conflict of interest

The authors declare no conflict of interest.

References

1. Lozano R, Naghavi M, Foreman K, et al. Global and regional mortality from 235 causes of death for 20 age groups in 1990 and 2010: a systematic analysis for the Global Burden of Disease Study 2010. Lancet 2012; 380: 2095-128.
2. Thompson S, James M, Wiebe N, et al. Cause of death in patients with reduced kidney function. J Am Soc Nephrol 2015; 26: 2504-11.
3. Tonelli M, Mintzer P, Lloyd A, et al. Risk of coronary events in people with chronic kidney disease compared with those with diabetes: a population-level cohort study. Lancet 2012; 380: 807-14.
4. Sud M, Tangri N, Pittilie M, Levey AS, Naimark D. Risk of end-stage renal disease and death after cardiovascular events in chronic kidney disease. Circulation 2014; 130: 458-65.
5. Schefold JC, Filippatos G, Hasenfuss G, Anker SD, von Haehling S. Heart failure and kidney dysfunction: epidemiology, mechanisms and management. Nat Rev Nephrol 2016; 12: 610-23.
6. Sud M, Naimark DM. Cardiovascular disease in chronic kidney disease in 2015. Curr Opin Nephrol Hypertens 2016; 25: 203-7.
7. Rubenstein MH, Harrell LC, Sheynberg BV, Schunkert H, Bazer H, Palacios IF. Are patients with renal failure good candidates for percutaneous coronary revascularization in the new device era? Circulation 2000; 102: 2966-72.
8. Harrington SC, Simari RD, Conover CA. Genetic deletion of pregnancy-associated plasma protein-A is associated with resistance to atherosclerotic lesion development in apolipoprotein E-deficient mice challenged with a high-fat diet. Circ Res 2007; 100: 1696-702.
9. Resch ZT, Chen BK, Bale LK, Oxvig C, Overgaard MT, Conover CA. Pregnancy-associated plasma protein A a gene expression as a target of inflammatory cytokines. Endocrinology 2004; 145: 1124-9.
10. Conover CA, Bale LK, Harrington SC, Resch ZT, Overgaard MT, Oxvig C. Cytokine stimulation of pregnancy-associated plasma protein A expression in human coronary artery smooth muscle cells: inhibition by resveratrol. Am J Physiol Cell Physiol 2006; 290: C183-8.
11. Boldt HB, Conover CA. Pregnancy-associated plasma protein A (PAPP-A): a local regulator of IGF bioavailability through cleavage of IGFBPs. Growth Horm IGF Res 2007; 17: 10-8.
12. Conover CA, Harrington SC, Bale LK. Differential regulation of pregnancy associated plasma protein-A in human coronary artery endothelial cells and smooth muscle cells. Growth Horm IGF Res 2008; 18: 213-20.
13. Conover CA, Mason MA, Bale LK, et al. Transgenic overexpression of pregnancy-associated plasma protein-A in murine arterial smooth muscle accelerates atherosclerotic lesion development. Am J Physiol Heart Circ Physiol 2010; 299: H284-91.
14. Iversen KK, Teisner AS, Teisner B, et al. Pregnancy associated plasma protein A, a potential marker for vulnerable plaque in patients with non-ST-segment elevation acute coronary syndrome. Clin Biochem 2009; 42: 828-34.
15. Bayes-Genis A, Conover CA, Overgaard MT, et al. Pregnancy-associated plasma protein A as a marker of acute coronary syndromes. N Engl J Med 2001; 345: 1022-9.
16. Heeschen C, Dimmeler S, Hamm CW, Fichtlscherer S, Simoons ML, Zeiher AM. Pregnancy-associated plasma protein-A levels in patients with acute coronary syndromes: comparison with markers of systemic inflammation, platelet activation, and myocardial necrosis. J Am Coll Cardiol 2005; 45: 229-37.
17. Iversen KK, Teisner B, Winkel P, et al. Pregnancy-associated plasma protein A as a marker for myocardial infarction and death in patients with stable coronary artery disease: a prognostic study within the CLARICOR Trial. Atherosclerosis 2011; 214: 203-8.
18. Kavsak PA, Wang X, Henderson M, Ko DT, MacRae AR, Jaffie AS. PAPP-A as a marker of increased long-term risk in patients with chest pain. Clin Biochem 2009; 42: 1012-8.
19. Lund J, Wittfooth S, Qin QP, et al. Free vs total pregnancy-associated plasma protein A (PAPP-A) as a predictor of 1-year outcome in patients presenting with non-ST-elevation acute coronary syndrome. Clin Chem 2010; 56: 1158-65.
20. Lund J, Qin QP, Ilva T, et al. Circulating pregnancy-associated plasma protein a predicts outcome in patients with acute coronary syndrome but no troponin I elevation. Circulation 2003; 108: 1924-64.
patients with coronary heart disease: a systematic review and meta-analysis. Arch Med Sci 2013; 9: 389-97.
25. Li Y, Zhou C, Zhou X, Song L, Hui R. PAPP-A in cardiac and non-cardiac conditions. Clin Chim Acta 2013; 417: 67-72.
26. Astrup AS, Tarnow L, Christiansen M, Hansen PR, Parving HH, Rossing P. Pregnancy-associated plasma protein A in a large cohort of Type 1 diabetic patients with and without diabetic nephropathy - a prospective follow-up study. Diabet Med 2007; 24: 1381-5.
27. Bayes B, Granada ML, Pastor MC, et al. Obesity, adiponectin and inflammation as predictors of new-onset diabetes mellitus after kidney transplantation. Am J Transplant 2007; 7: 416-22.
28. Jefferies HI, Terti T, Witthooth S, et al. Elevated serum free pregnancy-associated plasma protein-A independently predicts mortality in haemodialysis patients but is not associated with recurrent haemodialysis-induced ischaemic myocardial injury. Nephron 2015; 129: 171-8.
29. Kalousova M, Benakova H, Kubena AA, Dusilova-Sulkovska S, Tesar V, Zima T. Pregnancy-associated plasma protein A as an independent mortality predictor in long-term haemodialysis patients. Kidney Blood Press Res 2012; 35: 192-201.
30. Kalousova M, Zima T, Krane V, et al. Pregnancy-associated plasma protein A associates with cardiovascular events in diabetic hemodialysis patients. Atherosclerosis 2014; 236: 263-9.
31. Lauzurica R, Pastor C, Bayes B, Hernandez JM, Romeiro R. Pretransplant pregnancy-associated plasma protein-a as a predictor of chronic allograft nephropathy and posttransplant cardiovascular events. Transplantation 2005; 80: 1441-6.
32. Etter C, Straub Y, Hersberger M, et al. Pregnancy-associated plasma protein-A is an independent short-time predictor of mortality in patients on maintenance haemodialysis. Eur Heart J 2010; 31: 354-9.
33. Stroup DF, Berlin JA, Morton SC, et al. Meta-analysis of observational studies in epidemiology: a proposal for reporting. Meta-analysis Of Observational Studies in Epidemiology (MOOSE) group. JAMA 2000; 283: 2008-12.
34. DerSimonian R, Laird N. Meta-analysis in clinical trials. Control Clin Trials 1986; 7: 177-88.
35. Egger M, Davey Smith G, Schneider M, Minder C. Bias in meta-analysis detected by a simple, graphical test. BMJ 1997; 315: 629-34.
36. Fialova L, Kalousova M, Soukupova J, et al. Relation of pregnancy-associated plasma protein-a to renal function and dialysis modalities. Kidney Blood Press Res 2004; 27: 88-95.
37. Coskun A, Duran S, Apaydin S, Bulut I, Sariyar M. Pregnancy-associated plasma protein-A: evaluation of a new biomarker in renal transplant patients. Transplant Proc 2007; 39: 3072-6.
38. Lukaszyk E, Lukaszyk M, Koc-Zorawska E, Bodzenta-Lukaszyk A, Malezsko J. Fibroblast growth factor 23, iron and inflammation – are they related in early stages of chronic kidney disease? Arch Med Sci 2016; 13: 845-50.
39. Fadel FI, Elshamaa MF, Elghoroury EA, et al. Usefulness of serum procalcitonin as a diagnostic biomarker of infection in children with chronic kidney disease. Arch Med Sci Atheroscler Dis 2016; 1: 23-31.
40. Conover CA, Bale LK, Mader JR, Mason MA, Keenan KP, Marler RL. Longevity and age-related pathology of mice deficient in pregnancy-associated plasma protein-A. J Gerontol A Biol Sci Med Sci 2010; 65: 590-9.
41. Swindell WR, Masternak MM, Bartke A. In vivo analysis of gene expression in long-lived mice lacking the preg-