Correlation between Serum Uric Acid Level and Severity of Coronary Artery Stenosis in Patients with Acute Coronary Syndrome

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Abstract

Acute coronary syndrome (ACS) is a life-threatening disease which remains a source of high morbidity and mortality despite advances in treatment. The relationship between serum uric acid (SUA) level and ischemic heart disease abides controversial and still has not been established as a cardiovascular risk factor. The cooperative interaction between those two factors is not fully understood. Prior epidemiological evidences of the causal relationship is still argumentative. There were various studies using the same methods yet the outcome were different. This study aims to conduct a meta-analysis to synthesize the results of recent studies in order to obtain data quantitatively and also accurately. Systematic study follows the guidelines for preferred reporting items for systematic reviews and meta-analysis (PRISMA), tracing studies published in vulnerable periods from January 2010 to May 2020. The Cochrane Library, Ebsco, Medline/PubMed, ProQuest and Science Direct are sources of published studies. Meta-analysis was conducted to synthesize the associations between SUA level and severity of coronary artery stenosis, using random effect model to account for possible study heterogeneity. Heterogeneity was assessed using I², and the meta-analysis was performed using Comprehensive Meta Analysis Version 3 (CMA3) software. Five studies (n=601 patients) identified a correlation between serum uric acid levels and Gensini scores (r=0.548; p<0.001) in ACS patients. Heterogeneity bias was found in the analysis, whereas publication bias was not found. Conclusion, severity of coronary artery stenosis in patients with ACS is positively correlated with serum uric acid levels.

Keywords: Acute coronary syndromes, Gensini score, uric acid

Hubungan Kadar Asam Urat Serum dengan Keparahan Stenosis Arteri Koroner pada Pasien Sindrom Koroner Akut

Abstrak

Sindrom koroner akut (SKA) adalah gangguan yang mengancam jiwa yang tetap menjadi sumber morbiditas dan mortalitas yang tinggi meskipun ada kemajuan dalam pengobatan. Hubungan antara asam urat serum dengan penyakit jantung iskemik masih kontroversial dan belum ditetapkan sebagai faktor risiko kardiovaskular. Interaksi koperatif antara keduanya tidak sepenuhnya dipahami. Beberapa bukti epidemiologis hubungan kausal tersebut masih kontroversial. Sering sekali penelitian dengan kasus yang sama dan menggunakan metode yang sama tetapi hasilnya berbeda. Penelitian ini bertujuan melakukan meta analisis untuk mensintesis hasil-hasil penelitian yang berbeda tersebut agar diperoleh data baru yang bersifat kuantitatif dan lebih akurat. Telaah sistematis mengikuti pedoman preferred reporting items for systematic reviews and meta-analyses (PRISMA), dengan menelusuri studi yang dipublikasikan dalam rentan waktu dari Januari 2010 hingga Mei 2020. Cochrane Library, Ebsco, Medline/PubMed, ProQuest, dan Science Direct adalah sumber dari studi yang dipublikasikan. Meta-analisis dilakukan untuk mensintesis korelasi antara kadar asam urat serum dan keparahan stenosis arteri koroner menggunakan model efek acak untuk menjelaskan kemungkinan heterogenitas penelitian. Heterogenitas dinilai menggunakan I²; dan meta analisis menggunakan perangkat lunak Comprehensive Meta Analysis Version 3 (CMA3). Lima studi (n=601 pasien) diidentifikasi didapatkan korelasi antara kadar asam urat serum dan skor Gensini (r=0.548; p<0.001) pada pasien SKA. Bias heterogenitas ditemukan dalam analisis, sedangkan bias publikasi tidak ditemukan. Simulan, keparahan stenosis arteri koroner pada pasien dengan SKA berkorelasi positif dengan kadar asam urat serum.

Kata kunci: Asam urat, sindrom koroner akut, skor Gensini

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Introduction
Cardiovascular disease (CVD) is a global primary cause of death which accounts for 4.1 million deaths in Europe alone. According to the WHO in 2019, 1.7 million of CVD-related deaths are caused by coronary artery disease (CAD). Acute Coronary Syndrome (ACS) has become the primary cause of deaths and disabilities in Asia-Pacific, with more than 5% mortality in hospitalized patients. The region still lacks consensus on the best approach to overcoming its specific challenges in reducing mortality from ACS. The Asia-Pacific Real world evidence on Outcome and Treatment of ACS (APRICOT ACS) covers a wide variety of clinical conditions, such as angina pectoris (UAP), non-ST elevation myocardial infarction (NSTEMI), dan ST-elevation myocardial infarction (STEMI). Increased serum uric acid (SUA) level is a risk factor for cardiovascular disease (CVD). An increase in the SUA level is a common finding in patients with hypertension, insulin resistance, obesity, and CVD. However, it is still under debate whether such elevation is an independent predictor of cardiovascular risk or not. Nevertheless, the hypothesis that a reduction in the SUA level could prevent CVD has not been tested. A meta-analysis study reported that an increase in the SUA level significantly increases mortality and risk of major adverse cardiovascular events (MACE) and severe acute coronary artery diseases. Escalation in the SUA level correlates with cardiometabolic risks, which play a principal role in the pathophysiology of atherosclerotic formation. Increased uric acid level will increase the occurrence of endothelial dysfunction, oxidative stress, local inflammation, and insulin resistance, as well as inducing the proliferation of vascular smooth muscle cells that leads to vasoconstriction. This condition will trigger the development of atherosclerotic plaques, emphasizing its important role in the pathophysiology of cardiovascular disease.

Acute coronary syndrome often reflects a degree of damages to the coronary arteries by atherosclerosis, plaque rupture, thrombosis, and inflammation. The SUA level in patients with ACS is higher than those with chronic coronary syndrome (CCS). As a result, many studies show a positive correlation between an increment in SUA levels and severity of coronary artery disease by comparing them with normouricemia in ACS patients. However, the hyperuricemia terms have various boundaries and interpretations, causing differences in the results. Until recently, no systematic review on the correlation between SUA level and coronary artery stenosis severity in patients with ACS has been conducted. Hence, this meta-analysis aimed to systematically analyze the correlation between SUA level and coronary artery stenosis severity in patients with ACS.

Methods
This study is registered with PROSPERO, CRD42020210948 and reported by following the recommendation and guideline of the Preferred Reporting Items of Systematic Reviews and Meta-analyses (PRISMA). Two authors independently extracted data from published studies and meta-analysis was conducted using the Comprehensive Meta Analysis version 3 (CMA 3) software.

The authors conducted literature search in Cochrane Library, Ebsco, Medline/PubMed, ProQuest, and Science Direct. Where available, both controlled vocabulary terms and text words (MesH and keywords) were used in the subject component blocks. The search was conducted on studies published in English during the period of 2010 to May 2020. This period was selected to provide a contemporary reflection of relationship between SUA and severity of coronary artery stenosis in ACS patients. Additional studies were identified by searching article reference lists and including grey literature from the first 200 hits on Google Scholar after entering the index term combinations.

Sample was collected using PRISMA approach by applying the following inclusion criteria: (i) literature observed a correlation between two variables or more; (ii) used quantitative approach; (iii) included serum uric acid and coronary artery stenosis severity as research variables; (iv) subjects were patients with ACS; (v) and distinct diagnostic criteria for coronary artery stenosis severity measured with Gensini score was applied. Literature was excluded if it took the form of abstract, a piece from


**Figure 1 PRISMA Flow Diagram on Study Inclusion and Exclusion and Their Rationale**

- Records identified through database searching (n=1,216)
  - Cochrane Library (n=45)
  - EBSCO (n=273)
  - Proquest (n=51)
  - Medline/PubMed (n=179)
  - ScienceDirect (n=668)
- Duplication (n=268)
- 948 title were identified
- 870 article were excluded
- 78 abstract were reviewed
- 6 full text were not available
- 72 full text were analysed
- 67 full text were excluded
  - Irrelevant (n=52)
  - Chronic coronary syndrome (n=14)
  - Chinese language (n=1)
- 5 Full text meets the criteria for analysis (n=5)

Results

The literature search, reviews, and reasons for study exclusion are described in Figure 1. Critical analysis of five studies showed good quality reports and study designs. Those studies showed no bias in initial data collection; thus, would be considered to have a high degree of heterogeneity if the value was >75%.

**Table 1 Description of Studies Included in Analysis**

| Study  | Year | Number of Patients | Population | ACS Type | Study Type |
|--------|------|--------------------|------------|----------|------------|
|        |      |                    |            | STEMI    | NSTEMI     | UAP        |                |
|        |      |                    |            | (            | (            | (            |                |
| Duran  | 2011 | 246                | ACS        | 97 (39.4%) | 94 (38.2%) | 55 (22.4%) | Cross sectional |
| Qureshi| 2013 | 100                | Male-ACS   | 46 (46%)   | 50 (50%)   | 4 (4%)     | Cross sectional |
| Pramanik| 2015 | 82                 | ACS        | 47 (57.3%) | 24 (29.3%) | 11 (13.4%) | Cross sectional |
| Ma     | 2016 | 93                 | ACS        | 34 (36.5%) | 29 (31.2%) | 30 (32.3%) | Cross sectional |
| Gaubert| 2018 | 80                 | NSTEMI     | -         | 48 (60%)   | 32 (40%)   | Cross sectional |
they were included in this systematic review and meta-analysis. Details on these studies are described in Table 1.

The systematic review of the five articles showed a positive correlation between uric acid level and severity of coronary artery stenosis (Gensini score) in ACS patients.\textsuperscript{6,9–12} The relationship between SUA and severity of coronary atherosclerosis evaluated via endothelial dysfunction using peripheral arterial tone (PAT) in ACS patients was associated with severity, wound area, and total occlusion number.\textsuperscript{6,10–12} Number of diseased vessels, critical lesions and total occlusions on coronary angiogram. Results: Mean age of normouricemic group (n=59) Results of the systematic review presented that the five studies were qualified to be analyzed using the CMA3 program to evaluate the correlation between uric acid level and Gensini score. The total number of subjects included in these studies was 601 and the data characteristics of each study are presented in Table 2.

Variance analysis was performed to determine whether the studies were heterogeneous or homogenous. Studies were classified as heterogeneous if p-value <0.05 or I\textsuperscript{2} >75% in the heterogeneity test. After testing, studies included in this meta-analysis were found to be heterogeneous, with I\textsuperscript{2} =90.5% and p <0.001 (95% CI). Therefore, the model used to quantify pooled effect was the random effect model.

The Forest plot (Figure 2) showed the correlation between each study (black box) and its confidence interval (horizontal line). In addition, pooled correlations were depicted as diamond (red). A p-value of the forest plot was <0.05, resulting in the null hypothesis being rejected. Therefore, there was a positive correlation between uric acid level and severity of artery stenosis, with a correlation coefficient of 0.584, which could be categorized as moderate correlation. Also, the p-value in the significance test was 0.001, implying that the association between the two variables was significant.

The quality assessment of the studies also considered the risk of bias. Report and design quality of all studies were classified as good according to the AXIS tool and all studies showed no initial bias. In terms of publication bias, because there were fewer than ten included studies, the utility of funnel plots to assess publication bias was limited as the power of the tests would be considered too low to distinguish chance from the actual asymmetry. The regression method (p=0.568) and rank correlation method (p=0.462) of this meta-analysis provided a more objective means to

### Table 2 Data Characteristics of Studies on Correlation between Uric Acid and Gensini Score

| Study                  | Sample size | r    | 95% CI         | Z    | p      | Weight (%) |
|------------------------|-------------|------|----------------|------|--------|------------|
|                        |             |      |                |      |        |            |
| Duran et al., 2011     | 246         | 0.452| 0.347—0.546    | 7.595| 0.000  | 41.47      |
| Gaubert et al., 2018   | 80          | 0.738| 0.619—0.824    | 8.302| 0.000  | 13.14      |
| Ma et al., 2016        | 93          | 0.760| 0.658—0.835    | 9.451| 0.000  | 15.36      |
| Pramanik et al., 2015  | 82          | 0.418| 0.221—0.582    | 3.958| 0.000  | 13.48      |
| Qureshi et al., 2013   | 100         | 0.215| 0.019—0.394    | 2.147| 0.032  | 16.55      |
| Fixed effects          | 601         | 0.519| 0.458—0.576    | 13.931| <0.001| 100.00    |
| Random effects         | 601         | 0.548| 0.328—0.711    | 4.397| <0.001| 100.00    |
Table 3 Rank Correlation and Regression of Funnel pPot

| Rank Correlation Method | Regression Method |
|-------------------------|-------------------|
| p-value | Rank Correlation | p-value | Regression coefficient |
| 0.462 | 0.735 | 0.568 | 0.639 |

identify the presence of publication bias than a highly subjective visual inspection of the funnel plot (Table 3). There was no publication bias identified in this meta-analysis.

Discussion

This systematic review and meta-analysis is able to prove the correlation and comparability of uric acid and severity of coronary artery stenosis (Gensini score). Gensini score assessment gives weight to the closeness of the lesion in the left main coronary artery in addition to the stenosis degree, which leads to the maximum score. Therefore, the final score reflects the severity of coronary artery arteriosclerosis since it summarizes the cumulative effects of all lesions. The results of this systematic review and meta-analysis are obtained from SKA patients with higher mean uric acid levels and in line with the findings of Fromonot et al. stating that the SUA levels are higher in ACS in comparison to chronic coronary syndrome (CCS). Elevated SUA levels are associated with side effects and mortality in patients with IMA, as well as with the severity and mortality of acute CAD. Also, increased levels of SUA are associated with an increased risk of subsequent clinical events and mortality in patients with CCS. Increase in the SUA level, even when it is only in a slight increase out of the normal range (hyperuricemia), has started to cause a significant increase in the short-term mortality of patients with STEMI and patients undergoing PCI. The study of Ma et al. reported an increase in SUA levels related to the clinical type of CAD. These results are in line with the research from Fromonot et al., reporting that patients with CAD have increased SUA levels compared to healthy people.

This systematic evaluation and meta-analysis also described the correlation coefficients of SUA levels with moderate significant association to the severity of coronary artery stenosis. The magnitude of the determination coefficient (R square) is 0.300 or equal to 30%. This means that the SUA level variable simultaneously influences the severity of coronary artery stenosis by 30%, while the remaining 70% is processed by other variables aside from other risk factors for atherosclerosis, both modified (such as dyslipidemia, smoking, Diabetes Mellitus/insulin resistance, hypertension, chronic kidney disease (CKD), metabolic syndrome, systemic inflammation) and unmodified (such as age, genetic and hereditary, and male gender/male). Thus, increasing level of SUA is only considered as an adaptation response to try to prevent atherosclerosis due to the antioxidant properties of gout. Therefore, the SUA level can be considered a risk factor for atherosclerosis. This supports several studies that have revealed that increased levels of SUA are only considered an adaptation response as trial prevention against atherosclerosis due to the antioxidant properties of gout.

Uric acid (UA) is the end product of purine metabolism in humans and great apes. The UA acts as an antioxidant and accounts for 50% of the total antioxidant capacity of biological fluids in humans. When it presents in the cytoplasm or acidic/hydrophobic milieu in atherosclerotic plaques, UA converts into a pro-oxidant agent and promotes oxidative stress. Through this mechanism, it participates in the pathophysiology of human disease, including CVD. Although the causality in the relationship between UA and CVD remains unproven, UA may be pathogenic and participates in the pathophysiology of CVD by serving as a bridging mechanism, mediating (enabling) or potentiating the deleterious effects of cardiovascular risk factors in vascular and myocardium tissue.

The correlation between SUA and the severity of stenosis is made possible through the ACS pathomechanism process, which causes ischemia and acute myocardial necrosis due to severe coronary artery stenosis and even blockage. This ischemic state will induce anaerobic metabolism, which leads to low ATP production and the failure of ion exchange channels that disrupt enzymatic activity in the cytoplasm. Mitochondrial damage and electrolyte imbalance in the state of reperfusion increase oxidative stress through three primary systems, namely nicotinamide adenine
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