KOAH Akut Alevlenmesi ve Copeptin / Acute Exacerbation of COPD and Copeptin

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Özet
Amaç: Copeptin son yıllarda inflamasyon belirteci olarak tanımlanmaktadır. Copeptin’in akut KOAH alevlenmesinde inflamasyon belirteci olarak rolü henüz tam olarak gösterilememiştir. Bu çalışmamızda copeptin’in akut KOAH alevlenmesi nedeni ile hastaneye yatış sürecini ve diğer inflamasyon belirteçleri ile ilişkisini araştırmayı amaçladık. Gereç ve Yöntem: Akut KOAH alevlenmesi olan 46 hasta (42 erkek, 4 kadın) çalışmaya dahil edildi. Kanda lökosit sayısı, C-Reaktif Protein (CRP), Beyin Natriüretik Protein (BNP) ve copeptin düzeyleri hastaneye yatışın 0, 3, 7 ve 14. günlerinde ölçüldü. Bulgular: Hastaların ortalama yaşı 65±10 yıl bulundu. Copeptin median değeri sırası ile 0, 3, 7 ve 14. günlerde sırası ile 129.8, 170.8, 235.6, 338.4 pmol/L ölçüldü. CRP median değeri sırası ile 3.42, 1.65, 0.73, 1.15 mg/L düzeyinde idi. Serum BNP median değeri ise sırası ile 97, 88, 43, 2.5 U/L idi. Çalışma süresi boyunca ölçulen copeptin düzeyi ile CRP, lökosit ve BNP arasında anlamlı ilişki bulunmadı. Tartışma: Bu çalışmada akut KOAH alevlenmesinde serum copeptin düzeylerinde artış görülmekte iken CRP, lökosit ve BNP düzeylerinde azalma gözlemlendi. Copeptin ve diğer belirteçler arasında ilişki saptanmadı. Copeptin’in literatürde sunulan diğer bazı hastalıkların aksine copeptin alevlenmenin bir saptanmadığı düşünülüyor.

Anahtar Kelimeler
Copeptin; KOAH; Akut Alevlenme

Abstract
Aim: Copeptin has been introduced as an inflammation marker in recent years. The role of copeptin as an inflammation marker in short term period of Acute Exacerbation of COPD (AECOPD) has not yet been well demonstrated. We aimed to investigate the course of copeptin and it’s correlation with other inflammatory markers in AECOPD during hospitalization. Material and Methods: Forty-six AECOPD patients (42 male, 4 female) were included in the study. Blood leukocyte count, C- Reactive Protein (CRP), Brain Natriuretic Protein (BNP) and copeptin levels were measured on days 0, 3, 7 and 14, respectively. Results: Mean age of the patients was 65±10 years. Copeptin median levels were 129.8, 170.8, 235.6, 338.4 pmol/L on days 0, 3, 7 and 14, respectively. CRP median levels were 3.42, 1.65, 0.73, 1.15 mg/L, respectively. Serum BNP median values were 97, 88, 43, 2.5 U/L, respectively. Copeptin and CRP, leukocyte count and BNP during the study period were not significantly associated. Discussion: In this study we observed that serum copeptin levels were increasing in contrast to decrease in leukocyte, CRP and BNP levels during AECOPD. As no correlation was observed between copeptin and other markers we think that copeptin may not be an early inflammation marker of AECOPD contrary to other reports in literature.

Keywords
Copeptin; COPD, Acute Exacerbation
Introduction

Chronic Obstructive Pulmonary Disease (COPD) is a condition associated with an increasing rate of morbidity and mortality [1]. The clinical prognosis is worse in patients with COPD during the exacerbation period versus the stable period. Exacerbations are the most common indicators of treatment failure and disease progression [2].

With reference to both clinical findings and laboratory results, differential diagnosis of the stable and exacerbated period at a late stage of the disease is very difficult. Lower Respiratory Tract Infection (LRTI) is the most common cause of exacerbations [2,3]. However, heart failure is also common among patients with Acute Exacerbations of COPD (AECOPD) (20–40%) and for this reason; it is generally problematic to differentiate pure AECOPD from the heart failure. Traditional biomarkers, including fever, white blood cells count, and serum C-Reactive Protein (CRP) level, have been reported to be unreliable for assessment of the mortality rate and severity of the disease [4]. B-Type natriuretic peptide (BNP) is a neurohormone elevated in states of increased ventricular wall stress. While it has been established to be a good indicator of heart failure, it can also be elevated in a wide variety of clinical settings with or without Congestive Heart Failure (CHF), and in non-cardiac diseases such as chronic obstructive pulmonary disease and pulmonary embolism. Most dyspnea patients with heart failure have BNP values higher than 400 pg/mL, whereas left ventricular dysfunction without exacerbation, pulmonary embolism, or cor pulmonale should be considered in dyspnea patients with plasma BNP concentrations between 100 and 400 pg/mL [5].

It has been shown that BNP levels in AECOPD can be directly correlated with common markers of infection, such as CRP and procalcitonin. BNP levels may, therefore, accurately reflect the severity of the most prominent prognostic factors in AECOPD [6].

Thus, one approach in estimating the severity and predicting the outcome of the exacerbation of COPD is the use of novel biomarkers [2].

Arginine Vasopressin (AVP) is a hormone increasing in hypotensive, hypoxic, hyperosmolar or acidic stimuli and infectious conditions. Its instability makes its difficult to routinely take reliable measurements. Copeptin is a more stable C-terminal part of AVP and directly reflects levels of vasopressin. It can be easily measured during clinical routine and effectively mirrors vasopressin levels [7-10]. It then seems reasonable to speculate that the diverse secretion sites and the smaller amount of AVP and directly reflects levels of vasopressin. It can be reliably measured. Copeptin is a more stable C-terminal part of AVP and directly reflects levels of vasopressin. It can be easily measured during clinical routine and effectively mirrors vasopressin levels [7-10]. It then seems reasonable to speculate that the diverse secretion sites and the smaller amount of AVP and directly reflects levels of vasopressin. It can be reliably measured. Copeptin is a more stable C-terminal part of AVP and directly reflects levels of vasopressin. It can be easily measured during clinical routine and effectively mirrors vasopressin levels [7-10]. It then seems reasonable to speculate that the diverse secretion sites and the smaller amount of AVP and directly reflects levels of vasopressin. It can be reliably measured. Copeptin is a more stable C-terminal part of AVP and directly reflects levels of vasopressin. It can be easily measured during clinical routine and effectively mirrors vasopressin levels [7-10]. It then seems reasonable to speculate that the diverse secretion sites and the smaller amount of AVP and directly reflects levels of vasopressin. It can be reliably measured. Copeptin is a more stable C-terminal part of AVP and directly reflects levels of vasopressin. It can be easily measured during clinical routine and effectively mirrors vasopressin levels [7-10]. It then seems reasonable to speculate that the diverse secretion sites and the smaller amount of AVP and directly reflects levels of vasopressin. It can be reliably measured. Copeptin is a more stable C-terminal part of AVP and directly reflects levels of vasopressin. It can be easily measured during clinical routine and effectively mirrors vasopressin levels [7-10]. It then seems reasonable to speculate that the diverse secretion sites and the smaller amount of AVP and directly reflects levels of vasopressin. It can be reliably measured. Copeptin is a more stable C-terminal part of AVP and directly reflects levels of vasopressin. It can be easily measured during clinical routine and effectively mirrors vasopressin levels [7-10].

A recent study reported copeptin to be of interest as a prognostic biomarker for short-term and long-term prognoses in patients with AECOPD requiring hospitalization [11]. The aim of the current study was thus to investigate the usefulness of copeptin in assessing hospital prognosis of AECOPD and its correlation with serum CRP and BNP levels during hospitalization period of time.

Material and Method

Study Design and Patient Selection

Patients (age range: 40-75) consecutively admitted to Yedikule Hospital For Chest Diseases and Thoracic Surgery Training and Research between 01.09.2008-31.12.2008 with a preliminary diagnosis of AECOPD were included in the study. The diagnosis of COPD and exacerbation was based on the definition of the American Thoracic Society [1]. Inclusion criteria at entry were a change in the patient’s baseline dyspnea, cough, and/or sputum that is beyond normal day-to-day variations which is acute in onset, and a change in regular medication in a patient with underlying COPD and additionally smoking history more than 20 pack-years and no history suggestive of asthma. Patients with a history of renal failure, hepatic failure, recent Acute Myocardial Infarction (AMI) (in the past one year), active tuberculosis or cancer, immunodeficiency, suspected asthma or Cystic Fibrosis (CF), or large infiltrations in the x-ray were excluded from the study.

Patients were evaluated in terms of age, sex, dyspnea, cough, sputum characteristics, the amount of sputum, cigarette smoking status (smokers, non-smokers, quitters), number of smoking pack-years, biomass exposure status and duration, number of referrals to an emergency department, length of stay in the intensive care unit, physical examination findings, axillary body temperature, and presence of Anthonisen criteria [12]. Chest x-ray, Arterial Blood Gas (ABG) analysis, serum hemoglobin and hematocrit levels, platelet and eosinophil count, and biochemical parameters (urea, creatinine, ALT, AST, total protein, albumin, total bilirubine, direct and indirect bilirubine, total cholesterol, LDL-cholesterol, HDL-cholesterol, CRP, Brain Natriuretic Peptide (BNP), sedimentation rate, d-dimer), sputum gram staining and history of antibiotic use due to acute exacerbation at baseline or in the past six weeks were all noted. Blood complete blood counts, serum Copeptin, CRP, and BNP levels were evaluated on days 0, 3, 7 and 14 post-admission. Patient’s sputum was analyzed by gram staining and nonspecific culture. Signed informed consent forms were obtained from all patients.

Copeptin

Copeptin was analyzed by commercial ELISA kits (Phoenix Pharmaceuticals Inc, Belmont, CA, USA).

Preparation of Serum

A-4 mL blood samples were collected in tubes containing EDTA and shaken a few times for anticoagulation. Samples were transferred into biochemistry tubes containing Aprotinin (0.6 TIU/mL) to inhibit proteinase activity and shaken. Tubes were centrifuged at 1600 x g for 15 minutes. Plasma was stored at -20°C for two months and analyzed in the laboratory supplying the analysis kit.

Elsa

The immuno plate in this kit is pre-coated with a secondary antibody with nonspecific binding sites blocked. The secondary antibody is able to bind to the Fe fragment of the primary antibody (peptide antibody) whose Fab fragment will be competitively bound by both biotinylated peptide and peptide standard or targeted peptide in samples. The biotinylated peptide interacts with streptavidin-horseradish peroxidase (SA-HRP), catalyzing the substrate solution. The intensity of the yellow is directly proportional to the amount of biotinylated peptide
SA-HRP complex, but inversely proportional to the amount of peptide in standard solutions or samples due to competitive binding of the biotinylated peptide with the standard peptide or samples to the peptide antibody (primary antibody). A standard curve of known concentration can be established accordingly, with an unknown concentration in samples determined by extrapolation to this standard curve.

CRP was analyzed using the standard turbulometric method. BNP was analyzed using standard electro-chemiluminescence immunoassay.

**Radiological analysis**

digital chest x-ray was performed for all patients.

**Data Analysis**

Statistical analyses were performed using a statistical software program (SPSS for Macintosh, Gradpack Rel. 17.0.0, SPSS Inc., Chicago, IL, USA). Qualitative measurements were defined in real numbers and percentages. Quantitative variables were defined in mean ± standard deviation (min-max) for normal distribution, but in median values in the absence of normal distribution. The ANOVA test was performed to compare groups in normal distribution, while the Kruskal-Wallis test was used in the non-normal distribution. Paired t-tests were used to compare the dependent variables in normal distribution, whereas the Mann Whitney and Wilcoxon Rank sum tests were used in the non-normal distribution. (Chi-square test was also used to compare qualitative variables between the groups). Pearson’s correlation test was performed to analyze the correlation between the variables. p<0.05 was considered to be statistically significant.

**Results**

A total of forty-six patients were included in the study: forty-two (91.3%) males and four (8.7%) females. All male patients had a >20 pack-year history of cigarette smoking. Eleven patients (23.9%) had biomass exposure. A total of twenty-five (54.3%) had presented to emergency departments due to acute exacerbation of COPD within the past year, while six (13%) had been admitted to the Intensive Care Unit (ICU) within the past year. Upon admission, all patients (100%) had baseline dyspnea, 14 (30.4%) had type I exacerbation, fourteen patients (30.4%) had type II exacerbation, and eighteen (39.1%) had type III exacerbations. The patients’ characteristics are presented in Table I.

Median CRP level was 3.42 mg/L during hospitalization. Median CRP level was 0.70 mg/L during hospitalization.

**Data**

| Characteristics | Data* |
|-----------------|-------|
| Sex (M/F)       | 42(91.3)/4(8.7) |
| Age             | 65±10 (42-83) |
| Smoking history | 39 (84.7) |
| Smoking pack-year | 53.6±25.6 (14-80) |
| Biomass history | 7 (15.3) |
| AECOPD visit in the past one year | 25(54.3) |
| ICU Hospitalization due to AECOPD | 6 (13) |
| Dyspnea         | 46 (100) |
| Cough           | 39 (84.8) |
| Discolored sputum | 26 (56.5) |
| Chest examination | 29 (63) |
| Ronchi          | 4 (8.7) |
| Rales           | 2 (4.3) |
| Decreased respiratory sounds | 2 (4.3) |
| Normal sounds   | 2 (4.3) |
| Minimal radiological abnormality | 39 (84.8) |
| Type of AECOPD using Anthonisen criteria | |
| I. Increased dyspnea, sputum purulence, sputum volume | 14 (30.4) |
| II. Two of the above | 18 (39.1) |
| III. One of the above and one or more one minor findings | |
| Baseline pH     | 7.35±0.04 (7.23-7.42) |
| Baseline PaO2, mm Hg | 54.8±17.73 (27-95) |
| Baseline PaCO2, mm Hg | 50.05±14.94 (25.3-86.9) |
| Baseline HCO3, mg/L | 28.9±5.46 (19.5-43.3) |
| Baseline O2 saturation (%) | 82±13.18 (51-98) |
| Visiting emergency service median number | 2.5 |
| Visiting physician median number | 4 |
| Sputum volume(cc.) median number | 10 |
| Sedimentation rate (mm/h) median number | 25.5 |
| d-Dimer (mg/L) median number | 0.70 |

* Data are presented as an number (%), mean±SD (min-max) or median (IQR).

**Table 1. Baseline clinical characteristics of 46 patients presenting with AECOPD.**

| Characteristics | Data* |
|-----------------|-------|
| Sex (M/F)       | 42(91.3)/4(8.7) |
| Age             | 65±10 (42-83) |
| Smoking history | 39 (84.7) |
| Smoking pack-year | 53.6±25.6 (14-80) |
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| Sedimentation rate (mm/h) median number | 25.5 |
| d-Dimer (mg/L) median number | 0.70 |

Dimer value was 0.70 mg/L during hospitalization.

Mean leukocyte value was 10.50/mm³ on the first day of hospitalization, and 10.25/mm³, 10.05/mm³, 10.50/mm³ on days 3, 7, and 14, respectively.

Mean serum hematocrit value was 43.11%±5.62% (min:29.1-max:58.8) on the first day of hospitalization, 42.77%±5.44% (min:28- max: %55) on day 3, 42.66%±5.53% (min:27.4-max: 54.3) on day 7, and 43.12%± 4.36% (min: 35.2- max:51.5) on day 14.

The mean pH of the baseline arterial blood gas was 7.35 ± 0.04 (min:7.23-max: 7.42), mean PaO2 was 54.82 ± 17.73 mmHg (min:27- max:95), mean PaCO2 was 50.05 ± 14.94 mm. Hg (min:25.3- max: 86.9), mean HCO3 was 28.9± 5.46 mmol/L (min:19.5- max:43.3), mean O2 saturation was 82.20%±13.18% (min: 51 - max: 98).

The median CRP level was 3.42 mg/L on the first day of hospitalization, 1.65 mg/L on day 3, 0.73 mg/L on day 7, and 1.15 mg/L on day 14 (Figure 1).

The median BNP values were 97 pg/ml on day 0, 88 pg/ml on day 3, 343 pg/ml on day 7, and 2.5 pg/ml on day 14 (Figure 2).

The median copeptin levels was 129.80 pmol/L on the first day, 368.3 pmol/L on day 3 and 89.40 pmol/L on day 14.

| Variable                  | Day 0 | Day 3 | Day 7 | Day 14 |
|---------------------------|-------|-------|-------|--------|
| Copeptin (pmol/L)         | 0.17±0.04 | 0.17±0.04 | 0.17±0.04 | 0.17±0.04 |
| BNP (pg/ml)               | 97 ± 88 | 88 ± 98 | 88 ± 98 | 88 ± 98 |
| CRP (mg/L)                | 3.42±1.65 | 1.65±0.73 | 0.73±0.15 | 0.11±0.01 |
| D-Dimer (mg/L)            | 0.70 ± 0.70 | 0.70 ± 0.70 | 0.70 ± 0.70 | 0.70 ± 0.70 |

**Table 2.** Median hormone levels in patients with AECOPD.

**Table 3.** Median hormone levels in patients with AECOPD.
pmol/L on day 14 (Figure 3). A significant difference was found between copeptin levels on day 0, day 3, day 7, and day 14 (p<0.01).

While there was no statistically significant difference between median copeptin levels on day 0 and day 3 (p>0.05), a statistically significant difference was found between the levels on day 0, 7 and 14 (respectively, p<0.05 and p<0.05).

A significant difference was obtained between CRP levels on days 0, 3, 7, and 14 (p<0.001).

A statistically significant difference was found between median CRP values on days 0, 3, 7, and 14 (respectively, p<0.001, p<0.001 and p<0.01).

A significant difference was found between BNP levels on days 0, 3, 7, and 14 (p<0.001).

There was no statistically significant difference between median BNP values on day 0 and day 3 (p>0.05), whereas a statistically significant difference was found between day 0 and day 14 (respectively, p<0.001 and p<0.01).

No statistically significant correlation was found between CRP and copeptin levels (days 0, 3, 7, and 14) (p>0.05).

No statistically significant relationship was found between ABG PaO2 and copeptin levels on the first day of hospitalization (p<0.05).

Discussion

Although patients are diagnosed with COPD in clinical practice, no consensus has yet been reached on a gold standard marker for the prognosis and survival of COPD. Leucocytes, CRP and procalcitonin levels associated with the severity of disease are used for diagnosis during acute exacerbation of COPD [3]. However, there has been an increase in the previously reported use of Copeptin for prognostic monitoring in clinical settings. This diagnostic use of copeptin, which has been found to be the precursor of AVP, is an important finding particularly in terms of heart failure and septic shock. A recent study has suggested that in comparison to CRP and procalcitonin, copeptin is more effective in determining the survival rate in acute exacerbation of COPD [11].

To the best of our knowledge, no other study is currently available determining the course of copeptin measurement in acute exacerbation of COPD. Our aim was to investigate changes in the rate of copeptin at the early stage of exacerbation, investigate a possible relationship between copeptin and CRP in acute exacerbation of COPD, as well as between copeptin and pro-BNP, an indicator of heart failure.

It is known that infections often account for AECOPD. Studies have shown that the Arginine Vasopressin (AVP) level increases in the presence of infections and febrile conditions [13,14]. AVP level has also been shown to be associated with hypoxia-related vasoconstriction in severe COPD patients [15,16]. AVP is considered to exert adverse inotropic effects on the left ventricle in pulmonary hypertension [17-19]. AVP has also been shown to increase pulmonary vasoconstrictor response in endotoxemia [15]. These findings may be related to increased copeptin levels in poor prognosis of COPD [20].

In our study, similar to the study conducted by Stolz et al., no statistically significant relationship was found between copeptin levels with PaO2 levels during hospitalization. On the other hand, copeptin levels were significantly higher in our study, in comparison to other values reported in the literature.

Previous studies have shown increased CRP levels during acute
Exacerbations of COPD and that these levels were associated with mortality on day 14 and at 6 months [21-23]. Data has established increased plasma copeptin levels in patients with LRTI and pneumonia [24]. Krüger et al reported significantly increased copeptin levels in patients who died on day 28, compared to those of patients who survived pneumonia [25]. Studies have also showed that elevated CRP levels provided evidence of side effects in cardiovascular diseases and worsened lung functions in COPD [26,27]. CRP may increase rapidly in the presence of infection, and it may decrease quickly once the factor is excluded. However, unlike the study conducted by Stolz et al., we did not find a significant difference between the groups, although CRP levels were higher in Group III compared to Group I based on the Anthonisen criteria (p>0.05).

Copeptin levels were higher in our study in comparison to values reported in the literature. This may be explained by our use of the ELISA technique and its unique scale and cut-off values, in comparison to the approaches applied by others in the literature. In addition, we believe that such a comparison is not appropriate, since the majority of the studies in the literature were conducted by AVP.

To the best of our knowledge, no study is available investigating changes in copeptin levels in the first two weeks of acute exacerbation of COPD. The results of our study demonstrated that CRP levels increased steadily between day 0 and 14 and the increase was statistically significant (p<0.05). In unison to previous studies, this may be because COPD is a systemic disease, and it affects the whole cardiovascular system [28].

An increase in CRP may not be analogous to increased copeptin level in the acute exacerbation of COPD. Furthermore, Stolz et al. also did not find a significant relationship between CRP and copeptin [11]. Widespread inflammation occurs in cases of acute exacerbation of COPD. Also, copeptin levels are considered to be increased through stress factors within a short period of time following acute exacerbation of COPD.

Throughout the disease’s progression, complications may be also seen in the clinical presentation of COPD. Chronic airway obstruction leads to hypoxia in the late stages of disease. Increased resistance and, eventually, pulmonary hypertension develop due to pulmonary hypoxic vasoconstriction. The term “cor pulmonale” is used to refer to pulmonary hypertension related to the underlying COPD. Increased pulmonary artery pressure generally results in right ventricular hypertrophy (RVH). In addition, increased jugular venous pressure and hepatomegaly present with cor pulmonale, leading to right heart failure [29].

Acute exacerbation of COPD presents with right heart load. As a result, the increase and gradual decrease in BNP levels in our study may be explained by the right heart load in the patients, although heart failure is one of the exclusion criteria at baseline. In a recent study including 786 patients with chronic heart failure, Neuhold et al. found that in comparison to BNP and NT-proBNP, copeptin was more effective in determining the severity of heart failure in clinical settings. However, these markers were related to each other. On the other hand, no relationship of copeptin to BNP and NT-proBNP was found in the study. The authors found a significant correlation between heart failure and the New York Heart Association (NYHA) Disease Severity Class.

In the study, the authors found high copeptin levels to be associated with high mortality risk at 6 months during follow-up [30]. In another study, Jochberger et al. demonstrated that serum copeptin and AVP levels returned to normal ranges in patients who underwent cardiac surgery, when cardiovascular functions improved [31]. Stoiser and Gegenhuber et al. also reported that increased AVP concentration was associated with poor prognosis and the difference was statistically significant in patients with chronic heart failure. The authors suggested that copeptin was more effective in evaluating prognosis [10,32]. Stoiser et al. also suggested that plasma copeptin level measurement and BNP concentration, a standard biomarker of heart failure, were more effective in the evaluation of prognosis [10].

A high level of copeptin during baseline may indicate that it functions as an acute phase reactive against hypoxia and microembolism. However, review of the literature revealed no clinical study available to support this finding.

Within the aims of our study, we evaluated copeptin levels for a short period of time at the early stage of the disease in acute exacerbation of COPD. We found that there was a statistically significant difference in copeptin levels between the groups in the short-term. However, the small sample size used in our study should be taken into consideration.

The results of our study might demonstrate that COPD is a systemic disease with a variety of phenotypic characteristics and biomarkers used to definitively determine the presence of inflammation, and which has the potential to be influenced by many factors, although acute exacerbation of COPD has similar definitions in the clinical setting.

Conclusion

In conclusion, copeptin, which has similar characteristics to other inflammation markers, is considered to be a novel marker for the prognostic evaluation of acute exacerbation of COPD. We have found no significant correlation between copeptin and WBC, CRP or BNP levels in the short term for AECOPD. However, we did find that copeptin levels did increase gradually from day 1 to day 14. In contrast to previous studies, we conclude that copeptin levels might not be an early inflammation marker in AECOPD. Further large-scale studies with similar phenotypic characteristics are required to define the course and prognostic value of copeptin in AECOPD.

Competing interests

The authors declare that they have no competing interests.

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