# PREDICTIVE UTILITY OF MONOCYTE CHEMOATTRACTANT PROTEIN-1 (MCP-1) AND D-DIMER IN RISK STR ATIFICATION OF SEPSIS: A PROSPECTIVE COHORT STUDY

Iskandar A. a,

Fathonah S. a,

Soraya M. a, b,

Lova N. N. c

<sup>&</sup>lt;sup>a</sup> Dr. Saiful Anwar General Hospital, Malang, East Java, Indonesia.

<sup>&</sup>lt;sup>b</sup> Ulin General Hospital, Banjarmasin, South Borneo, Indonesia.

<sup>&</sup>lt;sup>c</sup> Indonesian Doctor Association, Malang, East Java, Indonesia.

10.15789/2220-7619-PUO-17982 МОНОЦИТАРНОГО

ПРОГНОСТИЧЕСКАЯ ЗНАЧЕНИЕ ХЕМОАТТРАКТАНТНОГО ПРОТЕИНА-1 (МСР-1) И D-ДИМЕРА В ОЦЕНКЕ РИСКА СЕПСИСА: ПРОСПЕКТИВНОЕ КОГОРТНОЕ ИССЛЕДОВАНИЕ

Искандер А. <sup>1</sup>, Фатона C. <sup>1</sup>, Сорайя M. <sup>1, 2</sup>, Лова H. H. <sup>3</sup>

<sup>&</sup>lt;sup>1</sup> Больница общего профиля им. д-ра Сайфула Анвара, Маланг, Восточная Ява, Индонезия.

<sup>&</sup>lt;sup>2</sup> Больница общего профиля им. Улина, Банджармасин, Южный Борнео, Индонезия.

<sup>&</sup>lt;sup>3</sup> Индонезийская ассоциация врачей, Маланг, Восточная Ява, Индонезия.

PREDICTIVE BIOMARKERS FOR MORTALITY IN SEPSIS PATIENTS
ПРОГНОЗИРУЮЩИЕ БИОМАРКЕРЫ СМЕРТНОСТИ ПАЦИЕНТОВ С СЕПСИСОМ
10.15789/2220-7619-PUO-17982

**Abstract** 

Background: Sepsis is a severe and life-threatening clinical syndrome characterized by a dysregulated host immune response to infection. This maladaptive response promotes widespread endothelial injury and abnormalities in coagulation, often progressing to multi-organ dysfunction and death. Mortality rates remain high, highlighting the urgent need for reliable biomarkers to enable early identification of patients at high risk. Such tools are particularly valuable in low- and middle-income countries (LMICs), where access to advanced diagnostic and therapeutic resources is limited. Monocyte Chemoattractant Protein-1 (MCP-1) is a chemokine that reflects hyperactivation of the innate immune system, while D-dimer indicates activation of coagulation and fibrinolysis. Although both pathways are central to the pathophysiology of sepsis, data evaluating their combined prognostic value in LMIC settings remain scarce.

**Objectives:** This study aimed to assess the prognostic significance of MCP-1 and D-dimer, both individually and in combination, for predicting 28-day mortality in patients with sepsis.

**Methods:** We conducted a prospective cohort study involving 83 adult patients with newly diagnosed sepsis at Dr. Saiful Anwar General Hospital, Malang, Indonesia. Serum MCP-1 levels were measured using enzyme-linked immunosorbent assay (ELISA), and plasma D-dimer levels were determined by immunoturbidimetry at the time of diagnosis. Patients were followed for 28 days, and survival outcomes were evaluated using Kaplan–Meier survival analysis and Cox proportional hazards regression models.

**Results:** Among the 83 participants, 58 patients (70%) died within 28 days. Non-survivors demonstrated significantly higher MCP-1 and D-dimer levels compared with survivors. An MCP-1 concentration ≥123.03 pg/mL was strongly associated

with increased mortality (HR 2.664, p = 0.005). Elevated D-dimer ( $\geq$ 43.5 mg/L FEU) showed a weaker individual association, but when combined with MCP-1,

predictive accuracy for mortality was significantly enhanced (HR 3.986, p = 0.037).

Conclusion: The concurrent elevation of MCP-1 and D-dimer identifies patients with sepsis who are at markedly increased risk of death. These findings support the potential utility of integrating inflammatory and coagulation biomarkers for early risk stratification. Moreover, they highlight the central role of the inflammation—coagulation axis in sepsis pathophysiology, with particular relevance for clinical

practice in resource-limited settings.

**Keywords:** sepsis; mortality prediction; MCP-1; D-dimer; coagulation biomarkers; resource-limited settings.

#### Резюме

собой тяжёлый Введение: Сепсис представляет жизнеугрожающий клинический синдром, характеризующийся нарушением регуляции иммунного ответа организма на инфекцию. Такой дезадаптивный ответ способствует обширному повреждению эндотелия и нарушениям коагуляции, часто прогрессирующим в полиорганную дисфункции и летальный исход. Показатели смертности остаются высокими, что подчёркивает острую необходимость в выявлении надёжных биомаркеров для раннего выявления пациентов с высоким риском, особенно востребованных в странах с низким и средним уровнем дохода (СНСД), где доступ к передовым диагностическим и терапевтическим ресурсам ограничен. Моноцитарный хемоаттрактантный белок-1 (МСР-1) – это хемокин, отражающий гиперактивацию врождённого иммунитета, в то время как D-димер указывает на активацию коагуляции и фибринолиза. Хотя оба биологических каскада играют центральную роль в патофизиологии оценивающих совокупное сепсиса, данных, ИХ прогностическое значение в условиях СНСД, получено недостаточно. Цели: Целью данного исследования была оценка прогностической значимости определения уровня MCP-1 и D-димера, как по отдельности, так и в сочетании, для прогнозирования 28-дневной смертности у пациентов с сепсисом.

**Методы:** Было проведено проспективное когортное исследование с участием 83 взрослых пациентов с впервые диагностированным сепсисом в больнице общего профиля им. д-ра Сайфула Анвара в Маланге, Индонезия. На момент постановки диагноза уровень МСР-1 в сыворотке крови измерялся с помощью иммуноферментного анализа (ИФА), а уровень D-димера в плазме — методом иммунотурбидиметрии. Наблюдение за пациентами проводилось в течение 28 дней, а результаты выживаемости оценивались с помощью анализа выживаемости Каплана—Майера и регрессионных моделей пропорциональных рисков Кокса.

Результаты: Из 83 участников 58 пациентов (70%) умерли в течение 28 дней.

У невыживших пациентов наблюдались значительно более высокие уровни

MCP-1 и D-димера по сравнению с выжившими. Концентрация MCP-1 ≥123,03

пг/мл была выраженно связана с повышенной смертностью (ОР 2,664, р =

0,005). Повышенный уровень D-димера (≥43,5 мг/л ФЭЕ [фибриноген-

эквивалентная единица]) имел более слабую индивидуальную взаимосвязь,

однако в сочетании с уровнем МСР-1 прогностическая точность смертности

значительно повысилась (OP 3,986, p = 0,037).

Заключение: Одновременное повышение уровня МСР-1 и D-димера

позволяет выявить пациентов с сепсисом, имеющих значительно повышенный

риск смерти. Указанные данные подтверждают потенциальную применимость

сочетанной оценки воспалительных и коагуляционных биомаркеров для

ранней стратификации риска, а также подчеркивают центральную роль оси

воспаление-коагуляция в патофизиологии сепсиса, что особенно актуально

для клинической практики в условиях ограниченных ресурсов.

Ключевые слова: сепсис; прогнозирование смертности; МСР-1; D-димер;

биомаркеры коагуляции; страны с ограниченными ресурсами.

#### 1 Introduction

Sepsis is a life-threatening syndrome resulting from a dysregulated host response to infection, leading to systemic inflammation, microvascular injury, and organ dysfunction. Despite advances in critical care, sepsis continues to cause substantial morbidity and mortality worldwide, with icu death rates ranging from 30% to 50% [1-3]. Among the various pathological processes in sepsis, coagulopathy has emerged as a key driver of poor outcomes, contributing to disseminated intravascular coagulation (dic), microthrombi formation, and multiple organ failure [4-5]. Early identification of coagulation disturbances is therefore critical for timely intervention and prognostication.

monocyte chemoattractant protein-1 (mcp-1), also known as c—c motif chemokine ligand 2 (ccl2), is a pro-inflammatory chemokine that recruits monocytes and other immune cells to sites of infection and injury. In sepsis, excessive mcp-1 release can amplify endothelial activation and trigger procoagulant pathways, linking inflammation to thrombotic complications [4-5]. Elevated mcp-1 levels have been associated with increased disease severity and mortality, possibly through their role in immune—coagulation crosstalk.

similarly, d-dimer, a fibrin degradation product, serves as a marker of coagulation activation and fibrinolysis. In sepsis, elevated d-dimer levels are frequently associated with sepsis-induced coagulopathy (sic) and the development of disseminated intravascular coagulation (dic), conditions that exacerbate organ dysfunction and increase the risk of death [6-8]. As a marker of endothelial injury and clot formation, d-dimer has shown prognostic value in assessing the severity of sepsis.

in recent years, mcp-1 and d-dimer have emerged as promising prognostic biomarkers in sepsis, offering insights into immune and coagulation pathways involved in disease progression [9-11]. Given the intertwined roles of inflammation and coagulation in sepsis progression, evaluating both mcp-1 and d-

dimer may provide a more comprehensive understanding of the immunothrombotic pathways driving mortality. However, evidence on their combined prognostic utility, particularly in low- and middle-income countries, remains scarce. This prospective study aims to assess the individual and joint predictive performance of mcp-1 and d-dimer for 28-day mortality in sepsis patients in indonesia, with the goal of improving accessible, mechanism-based risk assessment in resource-limited settings.

#### 2 Materials and methods

#### Study design

This prospective observational cohort study was conducted at dr. Saiful anwar general hospital (rssa), malang, indonesia. It is part of a larger sepsis biomarker project designed to investigate multiple pathophysiological pathways in sepsis. Each analysis within the cohort focuses on distinct biomarkers and hypotheses. The present study specifically examines mcp-1, an inflammatory chemokine, and d-dimer, a coagulation marker, in relation to 28-day mortality. Other analyses from the same cohort that evaluated different biomarkers, such as presepsin, soluble urokinase plasminogen activator receptor (supar), and procalcitonin, have been published or are under review separately, and are cited accordingly.

sepsis diagnosis was made by the attending physicians based on sepsis-3 criteria (2016). Patients meeting the eligibility criteria were enrolled consecutively over a one-year period. Blood samples were collected on the day of sepsis diagnosis, using residual serum from routine venous blood draws to analyze mcp-1 and d-dimer levels. All patients received standard medical care and were followed for 28 days to assess clinical outcomes. The study protocol was approved by the ethics committee of rssa, malang (approval no. 400/235/k.3/302/2019).

#### Study population and eligibility criteria

10.15789/2220-7619-PUO-17982

The study included adult patients (≥18 years old) newly diagnosed sepsis based on sepsis-3 definitions. Exclusion criteria included chronic inflammatory diseases, malignancies, or other conditions that could confound biomarker levels.

#### **Mcp-1** measurement

Serum mcp-1 levels were measured on the first day of hospitalization using the enzyme-linked immunosorbent assay (elisa) method. The analysis was performed in accordance with the max<sup>TM</sup> human mcp-1/ccl2 protocol (catalog no. 430107, biolegend inc., usa). Results were expressed in ng/ml.

#### **D-dimer measurement**

D-dimer was quantified by immunoturbidimetric method on a sysmex cs2100i analyzer with siemens innovance d-dimer reagents. The assay is based on the aggregation of polystyrene particles coated with monoclonal antibodies (clone 8d3) that react with d-dimer present in the sample.

#### **Statistical analysis**

Comparative analysis between survivors and non-survivors was performed using the mann—whitney u test. Receiver operating characteristic (roc) curve analysis identified optimal cut-off values for mcp-1 and d-dimer. Survival analysis was conducted using kaplan—meier curves, stratified by biomarker levels based on the derived cut-offs. Hazard ratios (hrs) were calculated using cox proportional hazards regression. The proportional hazards assumption was confirmed before performing time-independent multivariate cox regression to assess the predictive value of mcp-1, d-dimer, and their combination.

A p-value <0.05 was considered statistically significant. All statistical analyses were performed using spss version 24.0 for windows (ibm corp., armonk, ny, usa).

#### 3 Results

#### **Baseline characteristics**

### PREDICTIVE BIOMARKERS FOR MORTALITY IN SEPSIS PATIENTS ПРОГНОЗИРУЮЩИЕ БИОМАРКЕРЫ СМЕРТНОСТИ ПАЦИЕНТОВ С СЕПСИСОМ

10.15789/2220-7619-PUO-17982

A total of 83 sepsis patients met the inclusion criteria and were enrolled. 82 Among them, 25 patients (30.1%) survived, while 58 (69.9%) died during 83 hospitalization. The baseline characteristics are summarized in table 1, with 84 comparative analysis between survivors and non-survivors is presented in **table 2**. 85 Gender distribution was equal in the non-survivor group (29 males, 29 females), 86 with no significant difference. The mean age was 52 years for survivors and 55 years 87 for non-survivors, showing no statistical significance. Median length of hospital stay 88 was 7 days for survivors and 10.22 days for non-survivors, also not significantly 89 different. 90

#### **Biomarker levels**

91

92

93

94

95

96

97

98

99

100

101

102

103

104

105

106

107

108

109

110

Both mcp-1 and d-dimer levels were significantly elevated in non-survivors compared to survivors. Median mcp-1 was 282.27 pg/ml in non-survivors and 75 pg/ml in survivors (p = 0.000). Likewise, median d-dimer levels were 6.03 mg/l feu in non-survivors, compared to 2.66 mg/l feu in survivors (p = 0.012).

#### Overall survival analysis

Kaplan—meier survival analysis revealed a median survival time of 9 days for the entire cohort, indicating that 50% of patients died within this period. The survival curve also demonstrated that fewer than 10% of patients remained alive beyond day 28 (**figure 1**).

### Survival analysis and risk stratification based on mcp-1

Survival analysis stratified by mcp-1 levels is shown in **figure 2 and table 3**. The kaplan–meier curves, based on the cut-off value of 123.03 pg/ml (determined via roc analysis), crossed, indicating a violation of the proportional hazards (ph) assumption. Patients with mcp-1 levels < 123.03 pg/ml had a longer median survival of 31 days, while those with levels  $\geq$  123.03 pg/ml had a median survival of 11 days. The hazard ratio of mcp-1  $\geq$  123.03 pg/ml was 2.664 (ci 1.341 – 5.29; p value 0.005).

#### Survival analysis based on d-dimer

As depicted in **figure 3 and table 4**, survival analysis using a d-dimer cut-off of 43.5 mg/l feu also yielded kaplan—meier curves that violated the ph assumption.

Patients with d-dimer levels < 43.5 mg/l feu had a median survival of 17 days, compared to 8 days for those with levels  $\ge 43.5$  mg/l feu. The hazard ratio of d-dimer  $\ge 43.5$  mg/l feu was 1.980 (ci 0.612 - 6.40; p value 0.264).

#### Survival analysis based on combination of mcp-1 and d-dimer

A combined analysis of mcp-1 and d-dimer levels was performed based on their respective cut-off values. Patients with both mcp- $1 \ge 123.03$  pg/ml and d-dimer  $\geq$  43.5 mg/l feu were compared to those with either or both biomarkers below the thresholds. The kaplan–meier curve (**figure 4**) again violated the ph assumption. The group with elevated levels of both biomarkers had a significantly shorter median survival of 5 days, compared to 20 days in the group with lower levels of either mcp-1 or d-dimer. Patients were divided into 3 groups based on the presence of risk factors (mcp-1 and/or d-dimer levels). Table 5 showed that patients with 1 risk factor had an hr of 2.605 (ci 1.306 - 5.196; p value 0.007), while patients with 2 risk factors had an hr of 3.986 (ci 1.084 – 14.652; p value 0.037). 

#### 4 Discussion

#### **Baseline characteristics**

This study included 83 patients diagnosed with sepsis, with an almost equal gender distribution (49.4% female, 50.6% male) and a mean age of  $53.89 \pm 15.21$  years. The median length of hospital stay was 8 days, and 67.5% of patients had a sofa score  $\geq$ 6, indicating high illness severity. Overall, 69.9% (58/83) of patients died during hospitalization.

respiratory tract infections (42.2%) and lower urinary tract infections (27.7%) were the most common sources of sepsis. This distribution aligns with common etiologies in clinical settings, where respiratory infections are frequently associated with complications such as ards and poorer outcomes [1; 12; 3]. Several characteristics of patients in this study can be seen in table 1 below.

as shown in table 2, d-dimer and mcp-1 levels significantly differed between survivors and non-survivors. Non-survivors had higher d-dimer levels (median: 6.03 mg/l feu) compared to survivors (2.66 mg/l feu, p = 0.012), reflecting Russian Journal of Infection and Immunity

ISSN 2220-7619 (Print) ISSN 2313-7398 (Online)

- possible sepsis-associated coagulopathy, which is known to be linked to mortality [13; 14]. Likewise, mcp-1 levels were significantly elevated in non-survivors (282.27 pg/ml vs. 75 pg/ml, p < 0.001), indicating a stronger inflammatory response and supporting its role as a marker of immune activation and tissue injury in sepsis [15-17].
- no significant age difference was found between groups (median age: 55 vs. 52 years, p = 0.335), consistent with prior findings suggesting that age may not be an independent predictor of mortality in all sepsis populations [18]. Length of hospital stay was also not significantly different between groups (p = 0.905), implying that other factors like biomarker levels and illness severity may better reflect prognosis.
  - crucially, 96% of survivors had sofa scores <6, while only 5.2% of non-survivors had scores in this range (p = 0.000). This finding highlights the prognostic value of the sofa score at admission and supports its use in risk stratification among sepsis patients [19].

### Survival analysis and risk stratification based on mcp-1

Kaplan-meier analysis showed a median survival of 9 days among 83 sepsis patients, with less than 10% surviving beyond 28 days, underscoring the high early mortality risk in sepsis (figure 1).

Stratification by mcp-1 levels (<123.03 pg/ml vs. ≥123.03 pg/ml) revealed that patients with lower mcp-1 had significantly better survival (median: 31 vs. 11 days; figure 2, table 3), suggesting a strong association between lower inflammatory response and improved outcomes.

However, in figure 2 the kaplan–meier curve for mcp-1 violated the proportional hazards assumption, indicating the effect may not be consistent over time. The cut-off of 123.03 pg/ml yielded an auroc of 89.2% (95% ci: 81.1%-97.3%, p = 0.000) for mcp-1 in predicting mortality, with 81% sensitivity and 80% specificity.

151

152

153

154

155

156

157

158

159

160

161

162

163

164

165

166

167

This aligns with bozza et al., who found that mcp-1 levels >1000 pg/ml were associated with increased mortality in severe sepsis [15]. Jansen et al. Reported that mcp-1 >1600 pg/ml upon icu admission predicted 28-day mortality [20], and yamamoto et al. Found that levels >1500 pg/ml were associated with worse outcomes [21]. Chen et al.'s meta-analysis of 805 patients also supported the prognostic utility of mcp-1, despite variability in cut-offs across studies [22].

in our cohort, survival rates on days 10, 20, and 28 were consistently higher in patients with mcp-1 <123.03 pg/ml. Mean survival time was significantly longer for this group (31.16 days) compared to those with higher levels (11.05 days). Median survival for the lower mcp-1 group was 21 days versus 8 days for the higher group, reinforcing the prognostic significance of mcp-1. Although a specific study by duan et al. Reporting median survival times based on mcp-1 levels was not found, other studies confirm its predictive value. Matsumoto et al. Observed significantly higher mcp-1 levels in non-survivors, with an auc of 0.763 for 28-day mortality prediction [23]. A retrospective study in elderly patients showed mcp-1 had comparable prognostic performance to sofa and apache ii scores [10; 24]. Zhu et al. Also reported a strong correlation between mcp-1 and 28-day mortality, particularly in septic shock cases [9].

although most studies use mcp-1 to assess survival at fixed time points (e.g., 28 days), fewer report median survival, likely due to survival beyond this window. Nevertheless, the consistent association between elevated mcp-1 and mortality highlights its utility as a prognostic biomarker in sepsis.

### Survival analysis and risk stratification based on d-dimer

Survival analysis based on d-dimer levels (cut-off at 43.5 mg/l feu) revealed a mean survival of 17 days for patients with d-dimer <43.5 mg/l feu, compared to 8 days for those with higher levels (figure 3).

This supports the use of d-dimer as a biomarker for coagulopathy in sepsis and its potential role in mortality prediction. Like mcp-1, d-dimer also violated the

proportional hazards (ph) assumption, indicating that the relationship between ddimer and survival may change over the course of sepsis.

in this study, the auc for d-dimer levels in determining mortality was 67.4% (95% ci 55.8%-79.1%, p = 0.012). This auc value falls in the weak category (>60% - 70%) but is statistically significant. The cut-off point was determined using the coordinates of the curve table, yielding a cut-off of 43.5 mg/l feu with a sensitivity of 67.2% and specificity of 60%. Several studies have highlighted the prognostic value of d-dimer in predicting mortality among sepsis patients. Yunus et al. Reported that a d-dimer cut-off of 57 ng/ml demonstrated good predictive performance for mortality [25].

in a cohort of 684 sepsis patients, d-dimer levels were significantly higher in non-survivors compared to survivors (2489 ng/ml vs 1475 ng/ml, p = 0.0001), and showed the strongest predictive value for mortality among tested biomarkers (auc 0.68). Multivariate analysis confirmed that d-dimer was the only biomarker with a linear association with mortality, with an odds ratio of 3.03 (95% ci: 1.38–6.62) for levels above 2409 ng/ml [6]. Similarly, schupp et al. Found that d-dimer levels and dic scores showed good diagnostic accuracy for distinguishing septic shock (auc 0.710 and 0.739), but their prognostic performance for 30-day mortality was moderate (auc 0.590–0.610). Very high d-dimer levels (>30 mg/l) and dic scores ≥3 were significantly associated with increased mortality risk and remained independent predictors after multivariable adjustment [26].

across these studies, d-dimer cut-off values typically ranged from 2.5 to 4.0 mg/l feu, with aucs between 0.71 and 0.77—indicating moderate to good discriminatory ability. Sensitivity values commonly fell between 70–80%, and specificity ranged from 65–70%. The variability in results likely reflects differences in patient populations, d-dimer assay methods, sepsis definitions, and outcome measures. Despite these variations, most evidence supports that d-dimer levels exceeding 3.0 mg/l feu are associated with an increased risk of mortality in sepsis patients.

## PREDICTIVE BIOMARKERS FOR MORTALITY IN SEPSIS PATIENTS ПРОГНОЗИРУЮЩИЕ БИОМАРКЕРЫ СМЕРТНОСТИ ПАЦИЕНТОВ С СЕПСИСОМ

it is important to emphasize that while d-dimer is a useful biomarker for mortality prediction, it should not be used in isolation. Combining d-dimer measurements with comprehensive clinical evaluation and other prognostic markers can enhance the accuracy of risk stratification in sepsis. Our findings indicate that sepsis patients with d-dimer levels ≥43.5 mg/l feu exhibited lower survival rates compared to those with levels <43.5 mg/l feu, particularly on days 10 and 20 of observation (figure 3). Patients with lower d-dimer levels (<43.5 mg/l feu) demonstrated better survival outcomes, with a mean survival of 16.58 days, significantly higher than the 8.33 days observed in those with elevated d-dimer levels (≥43.5 mg/l feu). The median survival times also differed between the two groups. However, the hazard ratio (hr) for mortality in patients with elevated d-dimer levels was 1.980 (95% ci: 0.612–6.40), with a p-value of 0.254, indicating a non-significant association in this study (table 4).

In a study by zhang et al., 343 hospitalized covid-19 patients were evaluated to assess the predictive value of d-dimer levels for in-hospital mortality. The optimal cutoff value for d-dimer was found to be 2.0  $\mu$ g/ml, with a sensitivity of 92.3% and specificity of 83.3%. Of the 13 deaths during hospitalization, 12 occurred in patients with d-dimer levels  $\geq$ 2.0  $\mu$ g/ml, compared to just one death in patients with d-dimer levels <2.0  $\mu$ g/ml. This difference was statistically significant (p < .001). The hazard ratio for mortality with elevated d-dimer levels was 51.5 (95% ci: 12.9–206.7). The study concluded that d-dimer levels above 2.0  $\mu$ g/ml on admission could serve as a strong predictor of in-hospital mortality, suggesting that it could be an effective early marker for improving the management of covid-19 patients [27].

similarly, rodelo et al. Found that the 28-day mortality rate was 77.8% in patients with d-dimer levels >4.2 mg/l, compared to just 25% in those with levels ≤4.2 mg/l, although they did not provide data on median survival times [6]. These results are consistent with our findings, further supporting the link between elevated d-dimer levels and higher mortality in sepsis patients. Notably, most existing studies assess mortality based on 28- or 30-day outcomes rather than median survival time,

# PREDICTIVE BIOMARKERS FOR MORTALITY IN SEPSIS PATIENTS ПРОГНОЗИРУЮЩИЕ БИОМАРКЕРЫ СМЕРТНОСТИ ПАЦИЕНТОВ С СЕПСИСОМ

possibly because a substantial number of patients survive beyond the observation period.

nevertheless, elevated d-dimer levels—generally above 2.0 to 4.0 mg/l feu—consistently correlate with poorer prognosis in sepsis patients. Differences in cut-off values and mortality rates across studies may reflect variations in patient populations, d-dimer assay techniques, and sepsis diagnostic criteria. These findings are in line with our study, which also observed significantly elevated d-dimer levels in non-survivors.

# Survival analysis and risk stratification based on combination of mcp-1 and d-dimer

In the combined survival analysis of mcp-1 and d-dimer levels, patients with both biomarkers elevated above their respective cut-offs (mcp-1  $\geq$  123.03 pg/ml and d-dimer  $\geq$  43.5 mg/l feu) had a significantly shorter median survival of 5 days, compared to 20 days in patients with lower levels of either or both biomarkers (figure 4).

This highlights the value of combining inflammatory and coagulation markers to predict sepsis outcomes. The results suggest that both inflammatory and coagulatory pathways contribute to poor prognosis in sepsis, and combining biomarkers from these systems may improve risk stratification.

table 5 showed that the survival rate of sepsis patients with one risk factor was higher than in those with two risk factors. The hazard ratio (hr) was 3.986 (95% ci 1.084 - 14.652, p = 0.037), indicating that sepsis patients with both elevated d-dimer  $\geq 43.5$  mg/l feu and mcp-1  $\geq 123.03$  pg/ml are 3.986 times more likely to die quickly than those with lower levels of both biomarkers.

mikuła et al. Combined d-dimer with other biomarkers, finding that the d-dimer cut-off of 3570 ng/ml (3.57 mg/l feu) had an auc of 0.731, which increased to 0.801 when combined with other biomarkers [28]. Similarly, innocenti et al. Combined d-dimer with lactate, achieving an auc of 0.74 [29]. Although no studies have reported combined mcp-1 and d-dimer cut-offs, combining biomarkers Russian Journal of Infection and Immunity

ISSN 2220-7619 (Print) ISSN 2313-7398 (Online)

generally improves diagnostic accuracy. Ding et al. (2021) showed that combining ischemia-modified albumin (ima), d-dimer, and mcp-1 improved diagnostic accuracy, with an auc of 0.9047 in acute myocardial infarction patients [30].

The potential benefit of combining multiple biomarkers in sepsis stratification has been increasingly supported by recent evidence. While the combination of mcp-1 and d-dimer has not been extensively studied, the rationale for such an approach is strong, given that mcp-1 reflects monocyte-driven inflammation while d-dimer reflects coagulopathy. This dual representation of inflammatory and thrombotic pathways aligns with the multifactorial pathophysiology of sepsis. In our previous study, presepsin—a soluble cd14 subtype—demonstrated excellent prognostic value in sepsis, with an auc of 0.939 for predicting 28-day mortality and a hazard ratio of 3.654 for patients with levels above 17,085 pg/ml [31]. Notably, its prognostic performance improved when interpreted in conjunction with clinical severity scores and other laboratory markers. These findings emphasize that no single biomarker sufficiently captures the complexity of sepsis. Therefore, incorporating mcp-1 and d-dimer, possibly alongside markers such as presepsin, may offer a more holistic risk assessment framework. A multimarker strategy, tailored to local resources and patient populations, could improve early risk stratification and inform timely, personalized interventions [31].

#### **5 Conclusion**

283

284

285

286

287

288

289

290

291

292

293

294

295

296

297

298

299

300

301

302

303

304

305

306

307

308

309

310

In conclusion, our study suggests that the combination of mcp-1 and d-dimer levels may serve as valuable biomarkers for predicting mortality in sepsis patients. Elevated levels of both biomarkers are associated with significantly shorter survival and higher mortality risk.

#### Acknowledgements

The author extends sincere thanks to colleagues from the department of anesthesiology and intensive care at saiful anwar general hospital for their invaluable collaboration. Special thanks are also due to the rector of brawijaya university for providing funding support through the applied research grant with contract number 697.13/un10.c10/pn/2019.

#### **Author contributions**

311

312

313

314

315

316

317

318

319

320

321

322

323

324

Ai contributed to the conceptualization of the study, methodology design, visualization of data, and writing of the original draft, also contributed to the review and editing of the manuscript. Ms was involved in data collection, analysis of the data, and contributed to the review and editing of the manuscript. Sf had a role in methodology development, database management, and contributed to the review and editing of the manuscript. Nnl was contributed to the review and editing of the manuscript. All authors have made significant contributions to the article, have reviewed and edited the manuscript, and have approved the final version for submission.

#### **Conflict of interest**

The authors report no conflict of interest.

### ТАБЛИЦЫ

Table 1. Characteristics of Research Subjects.

Characteristics	N = 83
	f (%) or median (Q1- Q3)
Gender	
Man	42 (50.6%)
Woman	41 (49.4%)
Age (in year, mean±SD)	53.89 ± 15.21
D-dimer (in mg/L FEU)	4.81 (1.85 – 13.67)
MCP-1 (in pg/mL)	205.61 (98.64 – 405.00)
Length of stay (day)	8 (4 — 14)
SOFA score	6 (4 — 8)
<6	27 (32.5%)
≥6	56 (67.5%)
Outcome	
Alive ( survivor )	25 (30.1%)
Died ( non-survivor )	58 (69.9%)
Source of infection	
Respiratory	35 (42.2%)
Skin or joints	8 (9.6%)

10.15789/2220-7619-PUO-17982

Gastrointestinal tract	13 (15.7%)
Upper Urinary tract	3 (3.6%)
Central Nervous System	1 (1.2%)
Lower Urinary tract	23 (27.7%)

**Table 2.** Subject Characteristics based on Outcomes.

Characteristics	Survivor (n=25)	Non survivor (n=58)	p value
Gender			
Man	13 (52%)	29 (50%)	0.867 a
Woman	12 (48%)	29 (50%)	
Age (years)			
median (Q1- Q3)	52 (38.5 – 64.5)	55 (44.75 – 67.25)	0.335 °
D-dimer (mg/L			
FEU)			
median (Q1- Q3)	2.66 (1.28 – 6.46)	6.03 (1.98 – 16.58)	0.012 °
MCP-1 (pg/mL)			
median (Q1- Q3)	75 (42,42 – 11 1	282.27 (146.06	-0.000 c
	.82)	651.59)	
SOFA scores			
<6	24 (96%)	3 (5.2%)	0.000 °

# PREDICTIVE BIOMARKERS FOR MORTALITY IN SEPSIS PATIENTS ПРОГНОЗИРУЮЩИЕ БИОМАРКЕРЫ СМЕРТНОСТИ ПАЦИЕНТОВ С СЕПСИСОМ 10.15789/2220-7619-PUO-17982

≥6	1 (4%)	55 (94.8%)	
Length of stay (day)			
median (Q1- Q3)	7 (4.5 – 11.5)	10.22 (4 – 15.75)	0.905 °

**Notes:** <sup>a</sup> p value based on *chi - square test;* 

Table 3. Survival Analysis and Risk Stratification Based on MCP-1.

MCP -1	Low (<123.03 pg/mL)	High (≥123.03 pg/mL)
Total subject	31	52
Amount event	11 (35.48%)	47 (90.38%)
Amount censored	20 (64.52%)	5 (9.62%)
Mean (95% CI)	31.16 (11.36 – 50.96)	11.05 (8.39 – 13.72)
Median (95% CI)	21 (14.61 – 27.39)	8 (6.27 – 9.73)
p value	0.002	
Hazard Ratio		2.664 (1.341 – 5.29)
(95% CI)		
p value		0.005

<sup>&</sup>lt;sup>b</sup> p value based on *Independent t-test*;

<sup>&</sup>lt;sup>c</sup>p value based on Mann-Whitney U test;

<sup>\*</sup>p<0.05; significant

Table 4. Survival Analysis and Risk Stratification Based on D-Dimer.

D- dimer	Low (<43.5 mg/L FEU)	High (≥43.5 mg/L FEU)
Total subject	80	3
Amount event	55 (68.75%)	3 (100%)
Amount censored	25 (31.25%)	0 (0%)
Mean (95% CI)	16.58 (10.48 – 22.68)	8.33 (0 – 19,892)
Median (95% CI)	9 (7,044 – 10,957)	4 (0 — 8,801)
p value	0.223	
Hazard Ratio		1.980 (0.612 - 6.40)
(95% CI)		
p value		0.254

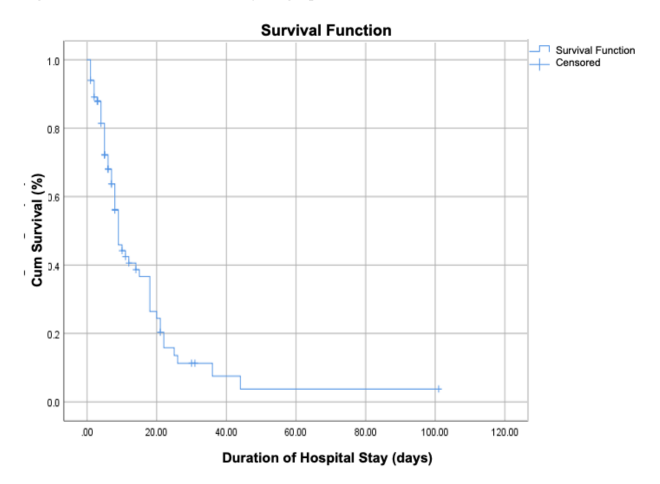
**Table 5.** Survival Analysis and Risk Stratification Based on Combination of MCP-1 and D-Dimer.

Variable	Group 1ª	Group 2 <sup>b</sup>	Group 3 <sup>c</sup>
Total subject	31	49	3
Amount event	11 (35.5%)	44 (89.8%)	3 (100%)
Censored amount	20 (64.5%)	5 (10.2%)	0 (0%)
Mean (95% CI)	31.16 (11.36 – 50.96)	11, 23 (8.5 – 14.0)	8.33 (0 - 19.89)
Median (95% CI)	21 (14.6 – 27.4)	8 (6.32 – 9.68)	4 (0 - 8.8)
p value		0.007	
Hazard Ratio (95% CI)	1	2.605 (1.306 – 5.196)	3.986 (1.084 – 14.652)
p value		0.007	0.037

**Notes:** a: Group with no risk factor (MCP-1 <123.03 pg/mL and D-Dimer <43.5 mg/L FEU); b: Group with 1 risk factors (MCP-1  $\geq$  123.03 pg/mL or D-dimer  $\geq$ 43.5 mg/L FEU); c: group with 2 risk factors (MCP-1  $\geq$  123.03 pg/mL and D-dimer  $\geq$ 43.5 mg/L FEU)

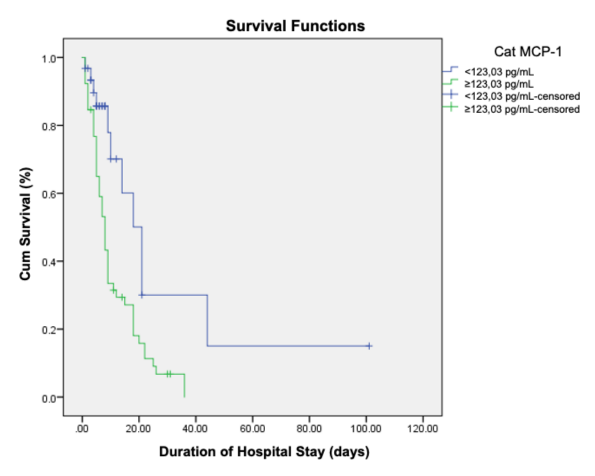
#### РИСУНКИ

**Figure 1.** Overall *survival* analysis graph.



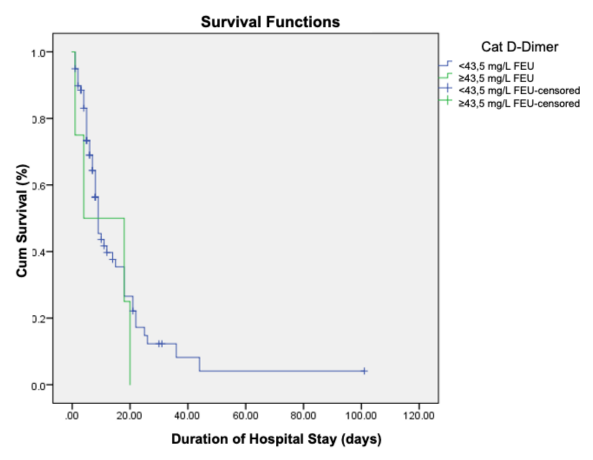
**Notes:** Kaplan–Meier survival analysis of 83 sepsis patients revealed a median survival time of 9 days. The survival rate sharply declined during the first two weeks of observation, with fewer than 10% of patients surviving beyond 28 days. This curve highlights the high early mortality associated with sepsis in the studied population.

**Figure 2.** Kaplan-meier survival analysis graph based on mcp-1 levels in sepsis patients.



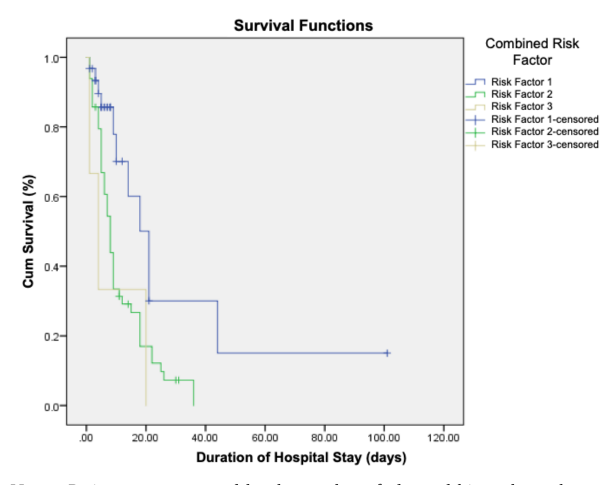
**Notes:** Survival analysis based on MCP-1 levels using a cut-off value of 123.03 pg/ml. Patients with MCP-1 <123.03 pg/ml (blue line) showed significantly improved survival compared to those with MCP-1 ≥123.03 pg/ml (green line). Median survival was 31 days in the lower MCP-1 group versus 11 days in the higher MCP-1 group. The survival curves crossed, indicating a violation of the proportional hazards assumption. This suggests that elevated MCP-1 levels are associated with a higher risk of mortality in sepsis.

**Figure 3.** Kaplan-meier survival analysis graphs based on d-dimer levels in sepsis patients.



**Notes:** Patients were divided by a D-dimer cut-off of 43.5 mg/L FEU. The blue line represents those with D-dimer < 43.5 mg/L FEU, and the green line those with D-dimer  $\ge 43.5$  mg/L FEU. Median survival was 17 days in the lower D-dimer group versus 8 days in the higher D-dimer group. The survival curves cross, indicating a violation of the proportional hazards assumption, but overall higher D-dimer levels are associated with increased mortality risk (AUROC 67.4%, p = 0.012).

**Figure 4.** Kaplan—meier survival analysis graph based on the combination of mcp-1 and d-dimer.



**Notes:** Patients were grouped by the number of elevated biomarkers: those with neither or only one elevated marker (MCP-1 < 123.03 pg/ml and/or D-dimer < 43.5 mg/L FEU; blue line) versus those with both MCP-1 ≥ 123.03 pg/ml and D-dimer ≥ 43.5 mg/L FEU (green line). Median survival was 20 days in the lower-risk group compared to 5 days in the high-risk group

### ТИТУЛЬНЫЙ ЛИСТ МЕТАДАННЫЕ

# Блок 1. Информация об авторе ответственном за переписку Agustin Iskandar,

Academic Degree : Medical Doctor, Clinical Pathologist, PhD

Academic status : Associate Profesor in the Medical Faculty Universitas

Brawijaya, Indonesia

Department of Clinical Pathology, Faculty of Medicine Universitas Brawijaya/ Dr. Saiful Anwar General Hospital, Malang, East Java, Indonesia;

Department of Clinical Pathology Study Programme, Faculty of Medicine Universitas Brawijaya/ Dr. Saiful Anwar General Hospital, Malang, East Java, Indonesia;

address: Jl. Jaksa Agung Suprapto No. 2, Malang 65112, East Java, Indonesia;

telephone: +62-8125298643;

e-mail: agustin\_almi@ub.ac.id

### Блок 2. Информация об авторах

#### Siti Fathonah,

Academic Degree : Medical Doctor, Clinical Pathologist

Academic status : Lecturer in the Department of Clinical Pathology at

Universitas Brawijaya in Malang, Indonesia

Department of Clinical Pathology, Faculty of Medicine Universitas Brawijaya/ Dr.

Saiful Anwar General Hospital, Malang, East Java, Indonesia;

Clinical Pathology Study Programme, Faculty of Medicine Universitas Brawijaya/

Dr. Saiful Anwar General Hospital, Malang, East Java, Indonesia;

# PREDICTIVE BIOMARKERS FOR MORTALITY IN SEPSIS PATIENTS ПРОГНОЗИРУЮЩИЕ БИОМАРКЕРЫ СМЕРТНОСТИ ПАЦИЕНТОВ С СЕПСИСОМ 10.15789/2220-7619-PUO-17982

#### Mira Soraya,

Academic Degree : Medical Doctor, Clinical Pathologist

Academic status : -

Clinical Pathology Study Programme, Faculty of Medicine Universitas Brawijaya/ Dr. Saiful Anwar General Hospital, Malang, East Java, Indonesia;

Department of Clinical Pathology, Ulin General Hospital, Banjarmasin, South Borneo, Indonesia;

### Novi Nirwanti Lova,

Academic Degree : Medical Doctor

Academic status : -

Indonesian Doctor Association, Malang, East Java, Indonesia;

#### Блок 3. Метаданные статьи

PREDICTIVE UTILITY OF MONOCYTE CHEMOATTRACTANT PROTEIN-1 (MCP-1) AND D-DIMER IN RISK STR

ATIFICATION OF SEPSIS: A PROSPECTIVE COHORT STUDY

Сокращенное название статьи для верхнего колонтитула:
PREDICTIVE BIOMARKERS FOR MORTALITY IN SEPSIS PATIENTS

**Keywords:** sepsis; mortality prediction; MCP-1; D-dimer; coagulation biomarkers; resource-limited settings.

Блок 1. Информация об авторе ответственном за переписку

Агустин Искандер,

Ученая степень: врач, клинический патолог, кандидат медицинских наук.

Ученый статус: Доцент медицинского факультета Университета Бравиджая,

Индонезия.

Кафедра клинической патологии, медицинский факультет Университета

Бравиджая/больница общего профиля доктора Сайфула Анвара, Маланг,

Восточная Ява, Индонезия;

Программа обучения кафедры клинической патологии, медицинский

факультет Университета Бравиджая / Больница общего профиля доктора

Сайфула Анвара, Маланг, Восточная Ява, Индонезия;

адрес: ул. Джакса Агунг Супрапто, 2, Маланг 65112, Восточная Ява,

Индонезия;

телефон: +62-8125298643;

электронная почта: agustin almi@ub.ac.id

Блок 2. Информация об авторах

Сити Фатона,

Ученая степень: врач, клинический патологоанатом.

Преподаватель Ученый статус: клинической кафедры

Университета Бравиджая в Маланге, Индонезия.

Кафедра клинической патологии, медицинский факультет Университета

Бравиджая/больница общего профиля доктора Сайфула Анвара, Маланг,

Восточная Ява, Индонезия;

Программа изучения клинической патологии, медицинский факультет

Университета Бравиджая / Больница общего профиля доктора Сайфула

Анвара, Маланг, Восточная Ява, Индонезия;

патологии

PREDICTIVE BIOMARKERS FOR MORTALITY IN SEPSIS PATIENTS
ПРОГНОЗИРУЮЩИЕ БИОМАРКЕРЫ СМЕРТНОСТИ ПАЦИЕНТОВ С СЕПСИСОМ
10.15789/2220-7619-PUO-17982

#### Мира Сорайя,

Ученая степень: врач, клинический патологоанатом.

Ученое звание: -

Программа изучения клинической патологии, медицинский факультет Университета Бравиджая / Больница общего профиля доктора Сайфула Анвара, Маланг, Восточная Ява, Индонезия;

Отделение клинической патологии, Больница общего профиля Улин, Банджармасин, Южный Борнео, Индонезия;

#### Нови Нирванти Лова,

Ученая степень: врач

Ученое звание: -

Индонезийская ассоциация врачей, Маланг, Восточная Ява, Индонезия

# PREDICTIVE BIOMARKERS FOR MORTALITY IN SEPSIS PATIENTS ПРОГНОЗИРУЮЩИЕ БИОМАРКЕРЫ СМЕРТНОСТИ ПАЦИЕНТОВ С СЕПСИСОМ 10.15789/2220-7619-PUO-17982

#### Блок 3. Метаданные статьи

ПРОГНОСТИЧЕСКАЯ ЗНАЧЕНИЕ МОНОЦИТАРНОГО ХЕМОАТТРАКТАНТНОГО ПРОТЕИНА-1 (МСР-1) И D-ДИМЕРА В ОЦЕНКЕ РИСКА СЕПСИСА: ПРОСПЕКТИВНОЕ КОГОРТНОЕ ИССЛЕДОВАНИЕ

#### Сокращенное название статьи для верхнего колонтитула:

ПРОГНОЗИРУЮЩИЕ БИОМАРКЕРЫ СМЕРТНОСТИ ПАЦИЕНТОВ С СЕПСИСОМ

**Ключевые слова:** сепсис; прогноз смертности; МКП-1; D-димер; биомаркеры свертывания крови; настройки с ограниченными ресурсами.

Оригинальные статьи.

Количество страниц текста -11,

количество таблиц -5,

количество рисунков -4,

29.07.2025

#### СПИСОК ЛИТЕРАТУРЫ

Referen			
ce			
Squence	Authors, title of a publication and source	Authors, title of a publication and source	
Number	where it was published, publisher's imprint	where it was published, publisher's imprint	Reference's URL
1.	Bozza FA, Salluh JI, Japiassu AM, Soares M,	Cytokine profiles as markers of disease	10.1186/cc5783
	Assis EF, Gomes RN, Bozza MT, Castro-	severity in sepsis: a multiplex analysis.	
	Faria-Neto HC, Bozza PT. Crit Care.	Crit Care, 2007; 11(2); R49	
		Monocyte chemoattractant protein-1 as a	
		potential marker for patients with sepsis: a	https://pubmed.ncbi.
		systematic review and meta-analysis.	nlm.nih.gov/377205
2	Chen Z, Li C, Yu J. Front Med (Lausanne).	Front Med (Lausanne) 2023, 1;10:1217784	14/
		Cytokine storm and sepsis disease	
		pathogenesis.	https://pubmed.ncbi.
	Chousterman BG, Swirski FK, Weber GF.	Semin Immunopathol, 2017Jul; 39(5), 517-	nlm.nih.gov/285553
3	Semin Immunopathol.	528	<u>85/</u>

		Monocyte chemoattractant protein-1 (MCP-	
		1): an overview.	https://pubmed.ncbi.
	Deshmane SL, Kremlev S, Amini S, Sawaya	J Interferon Cytokine Res, 2009, 29 (6): 313-	nlm.nih.gov/194418
4	BE. J Interferon Cytokine Res.	326	83/
		Clinical diagnostic value of combined	
		detection of IMA, D-D and MCP-1 in acute	https://pubmed.ncbi.
		myocardial infarction,	nlm.nih.gov/337471
5	Ding M, Li M, Yang H. Exp Ther Med.	Exp Ther Med, 2021; 21(5): 457	90/
		Association Between Plasma Levels of	
		Monocyte Chemoattractant Protein-1 (MCP-	
		1) and 28-Day Mortality in Elderly Patients	
		with Sepsis: A Retrospective Single-Center	https://pubmed.ncbi.
	Duan Y, Liu M, Wang J, Wei B. Med Sci	Study.	nlm.nih.gov/381694
6	Monit.	Med Sci Monit, 2024; 30, e942079	<u>64/</u>
	Evans L, Rhodes A, Alhazzani W, Antonelli	Surviving sepsis campaign: international	https://pubmed.ncbi.
	M, Coopersmith CM, French C, Machado FR,	guidelines for management of sepsis and	nlm.nih.gov/346057
7	Mcintyre L, Ostermann M, Prescott HC,	septic shock 2021.	81/

	Schorr C, Simpson S, Wiersinga WJ, Alshamsi	Intensive Care Med, 2021; 47(11): 1181-1247	
	F, Angus DC, Arabi Y, Azevedo L, Beale R,		
	Beilman G, Belley-Cote E, Burry L, Cecconi		
	M, Centofanti J, Coz Yataco A, De Waele J,		
	Dellinger RP, Doi K, Du B, Estenssoro E,		
	Ferrer R, Gomersall C, Hodgson C, Møller		
	MH, Iwashyna T, Jacob S, Kleinpell R,		
	Klompas M, Koh Y, Kumar A, Kwizera A,		
	Lobo S, Masur H, McGloughlin S, Mehta S,		
	Mehta Y, Mer M, Nunnally M, Oczkowski S,		
	Osborn T, Papathanassoglou E, Perner A,		
	Puskarich M, Roberts J, Schweickert W,		
	Seckel M, Sevransky J, Sprung CL, Welte T,		
	Zimmerman J, Levy M. Intensive Care Med.		
			https://pubmed.ncbi.
	Gao Q, Yang L, Teng F, Guo SB. BMC Emerg	Peripheral blood monocyte status is a predictor	nlm.nih.gov/367210
8	Med.	for judging occurrence and development on	90/

		sepsis in older adult population: a case control	
		study.	
		BMC Emerg Med, 2023, 23(1): 11	
		Sepsis: pathophysiology and clinical	https://pubmed.ncbi.
		management.	nlm.nih.gov/272170
9	Gotts JE, Matthay MA. BMJ.	BMJ, 2016, 353, i1585	<u>54/</u>
		Performance of D-dimer for predicting sepsis	https://pubmed.ncbi.
	Han YQ, Yan L, Zhang L, Ouyang PH, Li P,	mortality in the intensive care unit.	nlm.nih.gov/341408
10	Lippi G, Hu ZD. Biochem Med (Zagreb).	Biochem Med (Zagreb), 2021, 31(2): 020709	<u>32/</u>
			https://pubmed.ncbi.
	Iba T, Levy JH, Levi M, Connors JM, Thachil	Coagulopathy of Coronavirus Disease 2019,	<u>nlm.nih.gov/324674</u>
11	J. Crit Care Med.	Crit Care Med, 2020, 48, 9, 1358-1364	43/
	Iba T, Levy JH, Warkentin TE, Thachil J, van		
	der Poll T, Levi M; Scientific and	Diagnosis and management of sepsis-induced	
	Standardization Committee on DIC, and the	coagulopathy and disseminated intravascular	https://pubmed.ncbi.
	Scientific and Standardization Committee on	coagulation.	nlm.nih.gov/314109
12	Perioperative and Critical Care of the	J Thromb Haemost, 2019, 17(11): 1989-1994	83/

**Russian Journal of Infection and Immunity** 

ISSN 2220-7619 (Print) ISSN 2313-7398 (Online)

	International Society on Thrombosis and		
	Haemostasis. J Thromb Haemost.		
		Prognostic scores for early stratification of	
	Innocenti F, Bianchi S, Guerrini E, Vicidomini	septic patients admitted to an emergency	https://pubmed.ncbi.
	S, Conti A, Zanobetti M, Pini R. Eur J Emerg	department-high dependency unit.	nlm.nih.gov/239701
13	Med.	Eur J Emerg Med, 2014, 21(4): 254-9	00/
		Presepsin and Mortality Risk in Sepsis: A	
	Iskandar A, Prihastuti YA, Wulanda IA,	Valuable Tool for Predicting Patient Survival.	https://ojs.ukscip.co
	Aprilia A, Anshory, Muhammad. Trends in	Trends in Immunotherapy, 2025, 9 (2): 107-	m/index.php/ti/articl
14	Immunotherapy.	117	<u>e/view/1097</u>
		Mitochondrial DNA is Released in Urine of	
	Jansen MPB, Pulskens WP, Butter LM,	SIRS Patients With Acute Kidney Injury and	https://pubmed.ncbi.
	Florquin S, Juffermans NP, Roelofs JJTH,	Correlates With Severity of Renal	nlm.nih.gov/288375
15	Leemans JC. Shock.	Dysfunction. Shock. 2018; 49(3): 301-310	<u>26/</u>
		Diagnosis and Prognosis of Sepsis Based on	https://pubmed.ncbi.
	Jekarl DW, Kim JY, Ha JH, Lee S, Yoo J, Kim	Use of Cytokines, Chemokines, and Growth	nlm.nih.gov/315830
16	M, Kim Y. Dis Markers.	Factors.	25/

		Dis Markers, 2019, 8;2019:1089107	
	Kumar A, Roberts D, Wood KE, Light B,	Duration of hypotension before initiation of	
	Parrillo JE, Sharma S, Suppes R, Feinstein D,	effective antimicrobial therapy is the critical	https://pubmed.ncbi.
	Zanotti S, Taiberg L, Gurka D, Kumar A,	determinant of survival in human septic shock,	nlm.nih.gov/166251
17	Cheang M. Crit Care Med.	Crit Care Med, 2006, 34, 6, 1589-96	<u>25/</u>
	Liu VX, Fielding-Singh V, Greene JD, Baker	The Timing of Early Antibiotics and Hospital	https://pubmed.ncbi.
	JM, Iwashyna TJ, Bhattacharya J, Escobar GJ.	Mortality in Sepsis, Am J Respir Crit Care	<u>nlm.nih.gov/283459</u>
18	Am J Respir Crit Care Med.	Med, 2017, 196, 7, 856-863	<u>52/</u>
	Matsumoto H, Ogura H, Shimizu K, Ikeda M,	The clinical importance of a cytokine network	https://pubmed.ncbi.
	Hirose T, Matsuura H, Kang S, Takahashi K,	in the acute phase of sepsis, Sci Rep, 2018, 8,	nlm.nih.gov/302283
19	Tanaka T, Shimazu T. Sci Rep.	1, 13995	<u>72/</u>
		Significance of Heparin-Binding Protein and	
	Mikuła T, Sapuła M, Jabłońska J, Kozłowska	D-dimers in the Early Diagnosis of	https://pubmed.ncbi.
	J, Stańczak W, Krankowska D, Wiercińska-	Spontaneous Bacterial Peritonitis, Mediators	nlm.nih.gov/303639
20	Drapało A. Mediators Inflamm.	Inflamm, 2018, 2018, , 1969108	05/

	Rodelo JR, De la Rosa G, Valencia ML,	D-dimer is a significant prognostic factor in	https://pubmed.ncbi.
	Ospina S, Arango CM, Gómez CI, García A,	patients with suspected infection and sepsis,	nlm.nih.gov/227959
21	Nuñez E, Jaimes FA. Am J Emerg Med.	Am J Emerg Med, 2012, 30, 9, 1991-9	<u>96/</u>
	Rudd KE, Johnson SC, Agesa KM,		
	Shackelford KA, Tsoi D, Kievlan DR,		
	Colombara DV, Ikuta KS, Kissoon N, Finfer S,		
	Fleischmann-Struzek C, Machado FR,		
	Reinhart KK, Rowan K, Seymour CW, Watson	Global, regional, and national sepsis incidence	
	RS, West TE, Marinho F, Hay SI, Lozano R,	and mortality, 1990-2017: analysis for the	https://pubmed.ncbi.
	Lopez AD, Angus DC, Murray CJL, Naghavi	Global Burden of Disease Study	nlm.nih.gov/319544
22	M. Lancet.	Lancet, 2020, 395, 10219, 200-211	<u>65/</u>
	Schupp T, Weidner K, Rusnak J, Jawhar S,	D-Dimer Levels and the Disseminated	
	Forner J, Dulatahu F, Brück LM, Hoffmann U,	Intravascular Coagulation Score to Predict	https://pubmed.ncbi.
	Kittel M, Bertsch T, Akin I, Behnes M. Clin	Severity and Outcomes in Sepsis or Septic	nlm.nih.gov/371450
23	Lab.	Shock.Clin Lab, 2023, 69, 5,	<u>79/</u>

		D-dimer and histamine in early stage	
	Schwameis M, Steiner MM, Schoergenhofer	bacteremia: A prospective controlled cohort	https://pubmed.ncbi.
	C, Lagler H, Buchtele N, Jilma-Stohlawetz P,	study.	nlm.nih.gov/265862
24	Boehm T, Jilma B. Eur J Intern Med.	Eur J Intern Med, 2015, 26, 10, 782-6	<u>87/</u>
		D-dimer corrected for thrombin and plasmin	
	Semeraro F, Ammollo CT, Caironi P, Masson	generation is a strong predictor of mortality in	https://pubmed.ncbi.
	S, Latini R, Panigada M, Pesenti A, Semeraro	patients with sepsis.	nlm.nih.gov/318551
25	N, Gattinoni L, Colucci M. Blood Transfus.	Blood Transfus, 2020, 18, 4, 304-311	<u>52/</u>
	Singer M, Deutschman CS, Seymour CW,		
	Shankar-Hari M, Annane D, Bauer M,		
	Bellomo R, Bernard GR, Chiche JD,		
	Coopersmith CM, Hotchkiss RS, Levy MM,		
	Marshall JC, Martin GS, Opal SM, Rubenfeld	The Third International Consensus Definitions	https://pubmed.ncbi.
	GD, van der Poll T, Vincent JL, Angus DC.	for Sepsis and Septic Shock (Sepsis-3),	nlm.nih.gov/269033
26	JAMA.	JAMA, 2016, 315, 8, 801-10	38/

		The SOFA (Sepsis-related Organ Failure	
		Assessment) score to describe organ	
		dysfunction/failure. On behalf of the Working	
	Vincent JL, Moreno R, Takala J, Willatts S, De	Group on Sepsis-Related Problems of the	https://pubmed.ncbi.
	Mendonça A, Bruining H, Reinhart CK, Suter	European Society of Intensive Care Medicine.	nlm.nih.gov/884423
27	PM, Thijs LG. Intensive Care Med.	Intensive Care Med, 1996, 22(7): 707-10	9/
		The role of leukocyte emigration and IL-8 on	
		the development of lipopolysaccharide-	https://pubmed.ncbi.
	Yamamoto T, Kajikawa O, Martin TR, Sharar	induced lung injury in rabbits.	nlm.nih.gov/982055
28	SR, Harlan JM, Winn RK. J Immunol.	J Immunol, 1998, 161(10): 5704-9	2/
		The use of procalcitonin in the determination	
		of severity of sepsis, patient outcomes and	https://pubmed.ncbi.
		infection characteristics,	nlm.nih.gov/304278
29	Yunus I, Fasih A, Wang Y. PLoS One.	PLoS One, 2018, 13, 11, e0206527	<u>87/</u>
		D-dimer levels on admission to predict in-	https://pubmed.ncbi.
	Zhang L, Yan X, Fan Q, Liu H, Liu X, Liu Z,	hospital mortality in patients with Covid-19,	nlm.nih.gov/323064
30	Zhang Z. J Thromb Haemost.	J Thromb Haemost, 2020, 18(6): 1324-1329	92/

31	Zhu T, Liao X, Feng T, Wu Q, Zhang J, Cao X,	Plasma Monocyte Chemoattractant Protein 1	https://www.jstage.j
	Li H. Tohoku J Exp Med.	as a Predictive Marker for Sepsis Prognosis: A	st.go.jp/article/tjem/
		Prospective Cohort Study <sub>o</sub>	<u>241/2/241_139/_arti</u>
		Tohoku J Exp Med, 2017, Volum 241, Issue 2,	<u>cle</u>
		Pages 139-147	