

Early Change In Platelet Count And MPV Levels Of Patients Who Received Hemodialysis For The First Time: Mogadishu Somalia Experience

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Research Article

Keywords: HD, hemodialysis, CKD, MPV, MPV/PLT

Posted Date: June 15th, 2022

DOI: <https://doi.org/10.21203/rs.3.rs-1520048/v2>

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Abstract

Introduction

Mean platelet volume (MPV) is a marker used to assess the platelet's size and is also an indicator of platelet reactivity and prothrombic status.

Objective

In this study, we aimed to determine the relationship between MPV and biochemical parameters in patients who had received hemodialysis (HD) for the first time and then in respect of those same patients after their fourth HD.

Method

151 HD patients were enrolled in this study. Patients were eligible for inclusion if they had received their first HD session during this study protocol. Pre-hemodialysis blood samples were taken. Most laboratory values, including mean platelet volume (MPV) level and Platelets (PLT) count, were measured before the first HD and after the fourth HD session for each patient.

Results

Among the patients in our study, the mean age profile of the male patients (n= 103; 68.2%) was found to be higher than that of the female patients (n = 48; 31.8%) (53.62±18.19 vs. 46.17±17.9 years) (p = 0.019). In the patients' laboratory results after the fourth HD session, MPV, MPV/Plt, and Na values had increased to those after the first HD session (p<0.001). When age and gender status were taken into account, the level of weak positive correlation with white blood cell count (WBC), neutrophil, and red cell distribution width (RDW) was found, while the weak negative correlation with platelet to lymphocyte ratio (PLR) was found (p<0.001).

Conclusions

In our study, we found that increase in MPV and MPV/PLT levels was significant in the fourth HD session of patients with CKD. It is also debatable that there are findings indicating an increase in platelet reactivity in the first weeks of the onset of HD. This could be an early indicator of the early prevention of cardiovascular diseases.

Introduction

Chronic kidney disease (CKD) is a recognised public health problem, with mortality rates exceeding those in the general population. For dialysis patients, annual mortality stands at 21.2 %. While many organ systems are affected in the uremic process, cardiovascular disease (CVD) mortality is responsible for 45% of all mortality in dialysis patients.²

The literature indicated that the start of dialysis and subsequent initial months are associated with increased mortality.³⁻⁶ For patients with chronic kidney disease (CKD), these first months on hemodialysis (HD) have been shown to be a high-risk period for mortality. However, there is a relative paucity of data regarding cardiovascular event (CVE) rates following HD initiation. In particular, it is unclear whether the time to event admits of any variation with regard to CVE risk factors give evidence of any difference for both early and later dialysis periods.

Mean platelets volume, which a hematological analyzers can easily evaluate, is a useful marker of both platelet functions and activation. As a parameter, MPV indicates platelet activation and has been the subject of research in various patient groups with atherosclerosis.^{7,8} Larger platelets with metabolic and enzymatic properties are more active, and in a recent meta-analysis, it has been argued that this increased activity serves as a predictive biomarker for CVD.⁹

However, the relative paucity of studies on the uremic population is problematic. In this study, we aimed to find out the platelet activities during the initial stages of hemodialysis and the relationship between MPV and MPV/PLT with biochemical parameters in patients who had received HD for the first time and then in respect of those same patients after their fourth HD. The increased platelets activity can be used as an early indicator for cardiovascular prevention.

Materials And Methods

Study design and setting

This cohort cross-sectional study was carried out at Mogadishu Somali-Turkey Training and Research Hospital in Mogadishu, the capital of Somalia, which is the largest training and research hospital in the country.

Inclusion Criteria

All adult patients who received their first hemodialysis session during this study protocol between October 2018 and July 2019. Total of 151 patients who fulfilled the eligibility criteria have been included in the study.

Exclusion Criteria

Patients with malignancy, on routine hemodialysis, with a history of recent trauma, surgery or burns, with the evidence of acute infection, with a history of chronic liver disease and history of aspirin use, were excluded from the study.

Data collection

The ethical approval was obtained from Institutional Review Board (IRB) of Mogadishu Somali Turkey, Recep Tayyip Erdogan, Training and Research hospital and informed consent was taken from the patients.

Demographics like age and gender were obtained from medical records. Routine biochemical tests were performed with an autoanalyzer (Mindray BS-400 Clinical Chemistry Analyzer). Parameters such as white blood cell (WBC) count, MPV, platelet count and hemoglobin (Hb) were obtained from blood samples with an autoanalyzer (Sysmex XN-1000 Sysmex Corporation, Kobe, Japan).

Prior to hemodialysis, blood samples were obtained for measurement. Again, prior to the first HD session, MPV-level and platelet count among other laboratory values were measured for each patient, with these repeated following the fourth HD session. The interval between the time blood samples were drawn and their subsequent analysis was less than 30 minutes.

Hemodialysis

All patients underwent native hemodialysis with a temporary jugular catheter using the Fresenius 4008S dialysis machine. The first HD session was 2 hours and the following HD sessions were 4 hours duration each. Each hemodialysis session utilized a dialyzer with a blood flow rate of 250 to 300 ml/min, where the dialysate flow was 500 ml/min.

Data Analysis

We used SPSS 25.0 (IBM Corporation, Armonk, New York, United States) and PAST 3 (Hammer, Ø., Harper, D.A.T., Ryan, P.D. 2001. Paleontological Statistics) programs with regard to variable analysis. The Shapiro-Wilk Francia test evaluated the conformity of the univariate data to normal distribution. In contrast, the Mardia (Doornik and Hansen omnibus) test evaluated the conformity of multivariate variables to normal distribution. Comparison of the two independent groups according to the quantitative data was conducted via the Independent Sample t-test in conjunction with the Bootstrap results. The Bootstrap results were also employed with the Paired-Sample t-test.

The Wilcoxon signed-rank test, however, was used with the Monte Carlo simulation with respect to the comparison of the two quantitative dependent variables' repetitive measurements. The analysis of the correlation between the variables after controlling for the effects of age and gender was undertaken via the Partial Correlation test, with the analysis of the correlation between the variables before controlling for the effects of age and gender conducted via Kendall's Tau-b test. The quantitative data were set out in the tables as Mean \pm SD (standard deviation) (Minimum/Maximum) and Median (1st quartile/3rd quartile), while the categorical variables were n (%). Analysis of the variables was conducted at the 95% confidence level and a p-value of less than 0.05 was held to be significant.

Results

Demographic variables and laboratory results

Among the patients in our study, the mean age profile of male patients (n= 103; 68.2%) was

found to be higher than that for female patients (n = 48; 31.8%) (53.62 ± 18.19 versus 46.17 ± 17.9 years) ($p = 0.019$) (**Table 1**).

Laboratory results of patients after the fourth HD session showed that MPV, MPV/PLT, MCV, Hb, HCT, and Na values had increased compared to before the first HD session. ($p < 0.001$) (**Table 2**). As expected, a significant decrease was noted in BUN, creatinine, and K values ($p < 0.001$). There was no significant change in Ca, LYM, RDW, and PDW values ($p > 0.05$) (**Table 2**).

Relationship with MPV change

Having controlled for the effects of age and gender, we found the level of positive correlation with WBC, NEU, and RDW to be low, while we found the level of negative correlation with PLR to be low ($p < 0.001$). Where age and gender effects were not controlled for, the level of weak positive correlation detected between MPV and RDW was found. A weak negative relationship was found between MPV and PLT, PDW, and PLR ($p < 0.001$) (**Table 3**).

Discussion

Mean platelets volume and various inflammatory clinical states have been the subject of a number of investigations. Significant changes in platelet reactivity and platelet levels are observed in patients with a diagnosis of CKD.¹⁰ In chronic inflammatory conditions such as chronic kidney disease, platelet parameters were seen to differ significantly and were thus more marked in the state of HD. In our study, a slight increase in MPV levels was noted in patients in the first week of HD contrary to what is expected in the literature.^{11,12}

The higher prevalence of CKD and HD in males is commonly observed, but its explanation is not fully understood. It could be due to the larger propensity for CKD to progress to ESRD in males than in females. In the United States, the prevalence of ESRD was highest among people aged 65 to 74 years old until 2010, when the margin between those aged 75 and older began to close. While those ≥ 75 showed the highest incidence rate, ESRD prevalence was somewhat lower as a result of increased mortality in this cohort.¹³ The region in which the patients included in our study live is among those countries where gross domestic product (GDP) levels are low according to World Bank criteria.¹⁴ This may be attributed to the fact that the rate of male patients in our study is high, while ESRD patients' average age is lower than

normal or high-GDP societies. Cultural differences in the Mogadishu community also contribute to difficulties experienced by women in gaining access to and benefitting from healthcare services.

However, the number of studies conducted regarding changes in MPV in patients with CKD is low. The relevant literature indicates the difficulties in elucidating the influence of multiple determinants on PLT and MPV in physiology and during disease state, principally as a result of the differing factors affecting platelet reactivity.¹⁰ Nevertheless, the evaluation of this aspect of MPV within clinical practice should be the subject of additional research. In a study conducted by Balcioglu and Kirlioglu¹⁵ found that there is a negative correlation between MPV and number of hospital stays and total duration of hospital stays.

Activated large platelets have an important role in the pathogenesis of atherothrombosis and cardiovascular events (CVE). Platelets are not only constituent elements of thrombosis formation, but contribute to the induction of inflammation.¹⁶ Recent research has also shown that with respect to larger platelets, platelet reactivity levels are greater.¹⁷ In *The Anglo-Scandinavian Cardiac Outcomes Trial* (ASCOT), patients with end-organ damage and hypertension (HT) showed higher levels of MPV.¹⁶ However, in a study by Bilen *et al.*¹⁸ it was seen that with respect to increased inflammation in CKD patients, MPV cannot be shown to have a predictive value. It has also been argued that the larger MPV seen in CKD can be attributed to platelet swelling rather than platelet reactivity. More recently, in HD patients, it has been observed that there is a negative correlation between plasma HCO₃⁻ and MPV.¹⁹ Turgutalp *et al.*²⁰ found in their preliminary study results that patient MPV values were higher in those diagnosed with diabetic nephropathy than those in the normal population.

Given the existence of a nonlinear inverse relationship between PLT and MPV, it has been suggested that the assessment of MPV should always be correlated with platelet count.²¹ Other factors, which may be germane include age, gender, race, ethnicity, and lifestyle (including diet), while evidence would suggest that both MPV and PLT are also strongly influenced by genetic factors^{22,23} However, research to date is sparse with respect to changes in platelet-size parameters in patients with renal failure and according to GFR. With respect to CKD patients, Ju *et al.* previously noted higher MPV values and a progressive decrease in both platelet count and PDW²⁴. Heritability was 84% and of 75% respectively for PLT and MPV, a higher rate, attributed to genetic variations.²⁵

We think that this study is unique because it includes the results of ESRD patients living in Somalia, where there are significant difficulties in accessing healthcare facilities. In our study, changes in platelet volume and number in patients in the first week of HD were similar to those in the literature. We would argue that further studies are required to evaluate the effect of race, ethnicity and demographic features on the results.

Conclusions

In our study, we found that increase in MPV and MPV/PLT levels was significant in the fourth HD session of patients with CKD. It is also debatable that there are findings indicating an increase in platelet

reactivity in the first weeks of the onset of HD. This could be used as an early alarming feature for cardiovascular disease. In particular, CKD patients have a risk for cardiovascular event, so we recommend monitoring closely with other risk factors too.

Limitations:

There are some limitations to our study. We could not obtain the demographic features, chronic diseases, and CKD etiologies from the file system because they had not been integrated into the registry system. For this reason, it was only possible to evaluate HD patients with measured laboratory tests. Another main limitation is that we didn't include the cardiometabolic parameters in our study. Socioeconomic difficulties and cultural constraints preclude a ready solution to this problem.

Declarations

Conflicts of Interest

Authors declare no conflict of interests.

Ethical Committee Approval

The ethical approval was obtained from Institutional Review Board (IRB) of Mogadishu Somali Turkey, Recep Tayyip Erdogan, Training and Research hospital and informed consent was taken from the patients.

Tables

Table I

Table 1. Age and gender distributions of hemodialysis patients.

	Female	Male	Total	P
	(n=48)	(n=103)	(n=151)	
	Mean±SD. (Min./Max.)	Mean±SD. (Min./Max.)	Mean±SD. (Min./Max.)	
Age (year)	46,17±17,90 (22 / 86)	53,62±18,19 (18 / 88)	51,25±18,37 (18 / 88)	0.019

Independent Samples t-test(Bootstrap); SD., Standard deviation; Min., Minimum; Max., Maximum

Table 2. Change in laboratory tests before the first hemodialysis and after the fourth hemodialysis.

	Before 1 st HD	After 4th HD	P
	(n=151)	(n=151)	
	Mean±SD.	Mean±SD.	
MPV (fL)	9,86±1,49	10,06±1,47	0.045
MCV (fL)	82,04±5,88	83±4,86	0.008
	Median (Q1 / Q3)	Median (Q1 / Q3)	
Urea (md/dL)	231 (180 / 309)	87 (62 / 112)	<0,001
Creatinine (mg/dL)	11,70 (7,53 / 18,26)	4,99 (3,43 / 6,96)	<0,001
Sodium (meq/L)	134 (128 / 139)	136 (133 / 140)	<0,001
Potassium (meq/L)	4,70 (3,99 / 5,59)	3,83 (3,32 / 4,33)	<0,001
Total Protein (g/dL)	6,40 (5,50 / 7)	5,80 (5,30 / 6,50)	<0,001
Albumin (g/dL)			<0,001
	3,60 (2,90 / 4,10)	3,20 (2,70 / 3,50)	
Calcium (mg/dL)	8,10 (7,60 / 9)	8,20 (7,50 / 8,90)	0.896
WBC (*10 ³)	10,77 (7,56 / 16,28)	8,25 (6,10 / 11,85)	<0,001
Neutrophil (*10 ³)			<0,001
	8,10 (5,25 / 13,85)	5,80 (3,90 / 9,31)	
Lymphocyte (*10 ³)	1,03 (0,76 / 1,58)	1,04 (0,71 / 1,52)	0.962
Hb (g/dL)	8,50 (7,30 / 9,90)	9,20 (8,30 / 10,10)	0.004
HCT (%)	24,90 (20,70 / 28,60)	26,70 (24,50 / 28,60)	0.002
RDW-CV (%)	14,60 (13,40 / 16,30)	14,60 (13,50 / 16,10)	0.052
Platelets (*10 ³)	231 (171 / 315)	193 (140 / 260)	<0,001
PDW (%)	12,60 (10,40 / 16,20)	12,40 (10,20 / 16)	0.760
NLR	7,93 (4,76 / 14,48)	5,64 (3,52 / 11,33)	<0,001
PLR	215,12 (134,21 / 355,77)	185,71 (128,10 / 272,86)	0.002
MPV/PLT	0,04 (0,03 / 0,06)	0,43 (0,33 / 0,58)	<0,001

Paired Sample t-Test (Bootstrap), Wilcoxon sign Test(Monte Carlo); Q1, First Quartile; Q3, Third Quartile; SD., Standard deviation; WBC, total leucocyte; Hb, hemoglobin; HCT, Hematocrit; RDW-CV, Red blood cell distribution width - coefficient variation; NLR, Neutrophil to lymphocyte ratio; PLR, Platelet to lymphocyte value, MPV, Mean platelet volume; PLT, Platelets; fL, femtoliters, mg, milligram; L, liter; g, gram

Table 3. Correlation of MPV exchange with other laboratory tests.

Difference MPV	Controlled for age and gender		Not controlled for age and gender	
	r	p ^{pc}	r	p ^{kc}
MCV	-0.020	0.811	-0.005	0.929
Urea (md/dL)	-0.054	0.510	-0.049	0.379
Creatinine (mg/dL)	0.032	0.701	-0.004	0.936
Sodium (meq/L)	0.007	0.936	0.059	0.296
Potassium (meq/L)	0.058	0.481	0.041	0.462
Total Protein (g/dL)	0.048	0.558	0.007	0.894
Albumin (g/dL)	-0.056	0.499	-0.087	0.118
Calcium (mg/dL)	-0.013	0.876	-0.002	0.968
WBC (*103)	0.173	0.035	0.105	0.058
Neutrophil (*103)	0.171	0.037	0.101	0.067
Lymphocyte (*103)	0.152	0.065	0.085	0.125
Hb (g/dL)	-0.095	0.248	-0.053	0.335
HCT (%)	-0.118	0.153	-0.056	0.311
RDW-CV (%)	0.274	0.001	0.126	0.024
Platelets (*103)	-0.111	0.176	-0.139	0.012
PDW (%)	0.098	0.235	0.326	<0,001
NLR	-0.024	0.776	-0.005	0.933
PLR	-0.197	0.016	-0.159	0.004
MPV/PLT	0.149	0.069	0.093	0.092

^{pc} Partial Correlation Test, ^{kc} Kendall's Tau-b Test, r: Correlation Coefficient; WBC, total leucocyte; Hb, hemoglobin; HCT, Hematocrit; RDW-CV, Red blood cell distribution width - coefficient variation; NLR, Neutrophil to lymphocyte ratio; PLR, Platelet to lymphocyte value, MPV, Mean platelet volume; PLT, Platelets; fL, femtoliters, mg, milligram; L, liter; g, gram

References

1. Sarı Ö, Bashir AM. Early Change In Platelet Count And MPV Levels Of Patients Who Received Hemodialysis For The First Time: Mogadishu Somalia Experience. *Res Sq*. Published online 2022:1-12.
2. Valdivielso JM, Rodríguez-Puyol D, Pascual J, et al. Atherosclerosis in Chronic Kidney Disease. *Arterioscler Thromb Vasc Biol*. 2019;39(10):1938-1966. doi:10.1161/ATVBAHA.119.312705
3. Noordzij M, Jager KJ. Increased mortality early after dialysis initiation: A universal phenomenon. *Kidney Int*. 2014;85(1):12-14. doi:10.1038/ki.2013.316

4. Bradbury BD, Fissell RB, Albert JM, et al. Predictors of early mortality among incident US hemodialysis patients in the Dialysis Outcomes and Practice Patterns Study (DOPPS). *Clin J Am Soc Nephrol*. 2007;2(1):89-99. doi:10.2215/CJN.01170905
5. Robinson BM, Zhang J, Morgenstern H, et al. Worldwide, mortality risk is high soon after initiation of hemodialysis. *Kidney Int*. 2014;85(1):158-165. doi:10.1038/ki.2013.252
6. de Jager DJ. Cardiovascular and Noncardiovascular Mortality Among Patients Starting Dialysis. *JAMA*. 2009;302(16):1782. doi:10.1001/jama.2009.1488
7. Korniluk A, Koper-Lenkiewicz OM, Kamińska J, Kemonia H, Dymicka-Piekarska V. Mean platelet volume (MPV): New perspectives for an old marker in the course and prognosis of inflammatory conditions. *Mediators Inflamm*. 2019;2019. doi:10.1155/2019/9213074
8. Chen Y, Xiao Y, Lin Z, et al. The Role of Circulating Platelets Microparticles and Platelet Parameters in Acute Ischemic Stroke Patients. *J Stroke Cerebrovasc Dis*. 2015;24(10):2313-2320. doi:10.1016/j.jstrokecerebrovasdis.2015.06.018
9. De Luca G, Santagostino M, Secco GG, et al. Mean platelet volume and the extent of coronary artery disease: Results from a large prospective study. *Atherosclerosis*. 2009;206(1):292-297. doi:10.1016/j.atherosclerosis.2009.02.008
10. Cozzolino M, Mangano M, Stucchi A, Ciceri P, Conte F, Galassi A. Cardiovascular disease in dialysis patients. *Nephrol Dial Transplant*. 2018;33:iii28-iii34. doi:10.1093/ndt/gfy174
11. Sarnak MJ, Amann K, Bangalore S, et al. Chronic Kidney Disease and Coronary Artery Disease. *J Am Coll Cardiol*. 2019;74(14):1823-1838. doi:10.1016/j.jacc.2019.08.1017
12. Subbiah AK, Chhabra YK, Mahajan S. Cardiovascular disease in patients with chronic kidney disease: A neglected subgroup. *Heart Asia*. 2016;8(2):56-61. doi:10.1136/heartasia-2016-010809
13. Tucker BM, Freedman BI. Need to reclassify etiologies of ESRD on the CMS 2728 medical evidence report. *Clin J Am Soc Nephrol*. 2018;13(3):477-479. doi:10.2215/CJN.08310817
14. World Bank. Retrieved 6 July 2015.
15. Balcioglu YH, Kirlioglu SS. C-reactive protein/albumin and neutrophil/albumin ratios as novel inflammatory markers in patients with schizophrenia. *Psychiatry Investig*. 2020;17(9):902-910. doi:10.30773/pi.2020.0185
16. Morrell CN, Aggrey AA, Chapman LM, Modjeski KL. Emerging roles for platelets as immune and inflammatory cells. *Blood*. 2014;123(18):2759-2767. doi:10.1182/blood-2013-11-462432

17. Karolczak K, Soltysik B, Kostka T, Witas PJ, Watala C. Platelet and red blood cell counts, as well as the concentrations of uric acid, but not homocysteinaemia or oxidative stress, contribute mostly to platelet reactivity in older adults. *Oxid Med Cell Longev*. 2019;2019. doi:10.1155/2019/9467562
18. Bilen Y, Cankaya E, Keles M, et al. Does decreased mean platelet volume predict inflammation in chronic renal failure, dialysis, and transplanted patients? *Ren Fail*. 2014;36(1):69-72. doi:10.3109/0886022X.2013.832310
19. Nasri H, Baradaran A. Platelet count and mean volume (MPV) in association with plasma HCO₃⁻ in regular hemodialysis patients. *Rev Bras Hematol Hemoter*. 2006;28(2):127-130. doi:10.1590/S1516-84842006000200012
20. Turgutalp K, Özhan O, Akbay E, et al. Mean platelet volume and related factors in patients at different stages of diabetic nephropathy: A preliminary study. *Clin Appl Thromb*. 2014;20(2):190-195. doi:10.1177/1076029612456734
21. Pafili K, Penlioglou T, Mikhailidis DP, Papanas N. Mean platelet volume and coronary artery disease. *Curr Opin Cardiol*. 2019;34(4):390-398. doi:10.1097/HCO.0000000000000624
22. Eicher JD, Chami N, Kacprowski T, et al. Platelet-Related Variants Identified by Exomechip Meta-analysis in 157,293 Individuals. *Am J Hum Genet*. 2016;99(1):40-55. doi:10.1016/j.ajhg.2016.05.005
23. Kunicki TJ, Williams SA, Nugent DJ. Genetic variants that affect platelet function. *Curr Opin Hematol*. 2012;19(5):371-379. doi:10.1097/MOH.0b013e3283567526
24. Ju HY, Kim JK, Hur SM, et al. Could mean platelet volume be a promising biomarker of progression of chronic kidney disease? *Platelets*. 2015;26(2):143-147. doi:10.3109/09537104.2014.890179
25. Shameer K, Denny JC, Ding K, et al. A genome- and phenome-wide association study to identify genetic variants influencing platelet count and volume and their pleiotropic effects. *Hum Genet*. 2014;133(1):95-109. doi:10.1007/s00439-013-1355-7