



Ukrainian Journal of Nephrology and Dialysis

Scientific and Practical, Medical Journal

Founders:

- State Institution «Institute of Nephrology NAMS of Ukraine»
- National Kidney Foundation of Ukraine

ISSN 2304-0238;

eISSN 2616-7352

Journal homepage: <https://ukrjnd.com.ua>

Research Article

doi: 10.31450/ukrjnd.3(59).2018.03

N. Stepanova¹, L. Korol¹, V. Novakivskyy², M. Kolesnyk¹

The relationship between the dose of continuous erythropoietin receptor activator and oxidative stress in hemodialysis patients

¹State Institute «Institute of Nephrology of the National Academy of Medical Sciences», Kyiv, Ukraine

²LLC “Fresenius Medical Care Ukraine” Medical Centre, Cherkasy, Ukraine

Citation:

Stepanova N, Korol L, Novakivskyy V, Kolesnyk M. The relationship between the dose of continuous erythropoietin receptor activator and oxidative stress in hemodialysis patients. Ukr J Nephrol Dial. 2018;3(59):18-24. doi: 10.31450/ukrjnd.3(59).2018.03

Abstract. *At present, there have been no reports on the dose-dependent effects of continuous erythropoietin receptor activator (CERA) therapy on oxidative stress and red blood cell membrane lipid peroxidation parameters in hemodialysis (HD) patients.*

The aim of our work was to evaluate whether the dose of CERA treatment affected lipid peroxidation and antioxidant system in HD patients.

Methods. *38 HD patients were included in this single-center cross-sectional observational study. The study protocol was approved by a local Ethics Committee and all patients provided signed informed consent. The patients were stratified into quartiles ($\leq 25\%$ and $\geq 75\%$) according to the average dose of continuous erythropoietin receptor activator (CERA) and grouped in the following way: Group I (> 6 months of CERA treatment in a low dosage ≤ 50 $\mu\text{g}/\text{month}$, $n = 20$) and Group II (> 6 months of CERA treatment in a high dosage ≥ 125 $\mu\text{g}/\text{month}$, $n = 18$). Along with the standard diagnostic methods, we defined the content of malondialdehyde levels in the serum (MDAs) and erythrocytes (MDAe) spectrophotometrically as an indicator of lipid peroxidation. Such parameters as the concentration of ceruloplasmin (CP) and transferrin (TR) in the blood and total peroxidase activity (TPA) in erythrocyte were studied as the indicators of antioxidant system. In addition, we calculated the percentage of hemolysis, the RBC membrane permeability and oxidation coefficient.*

Results. *We obtained heterogeneous results in assessing oxidative stress parameters. The significantly higher levels of CP ($p = 0.007$) and TR ($p = 0.0003$) were found in the patients treated with a high dosage CERA. TPA activity in erythrocyte in the patients of Group II was statistically higher compared to Group I ($p = 0.02$). Moreover, we determined a statistically high percentage of hemolysis ($p = 0.03$) and RBC membrane permeability ($p < 0.0001$) in the patients who were treated with CERA in a dose ≥ 125 $\mu\text{g}/\text{month}$ compared to other patients. Using the probit regression model, we established the dose-dependent effect of CERA on the level of RBC membranes permeability: $\chi^2 = 21$; $p < 0.0001$.*

Conclusions. *We demonstrated that administration of CERA in a dose more 125 $\mu\text{g}/\text{month}$ improved the antioxidant status in HD patients. But, at the same time, it increased the hemolysis and RBC membranes permeability. Our preliminary data pointing to the dose-dependent effect of CERA on the RBC membrane lipid peroxidation parameters require further confirmation.*

Key words: *hemodialysis patients, continuous erythropoietin receptor activator, oxidative stress.*

Conflict of interest statement: all the authors declared no competing interests.

© N. Stepanova, L. Korol, V. Novakivskyy, M. Kolesnyk, 2018. All rights reserved.

Correspondence should be addressed to Natalia Stepanova: nmstep@ukr.net

Article history:

Received 11 July 2018

Received in revised form
01 August 2018

Accepted 9 August 2018



© Степанова Н.М., Король Л.В., Новаківський В.В., Колесник М.О., 2018

УДК: 616.61-085.38-073.27:577.152.1

Н. Степанова¹, Л. Король¹, В. Новаківський², М. Колесник¹

Взаємозв'язок між дозою тривалого активатора рецепторів еритропоєтину та оксидативним стресом у хворих, які лікуються методом гемодіалізу

¹ДУ «Інститут нефрології НАМН України», м. Київ

²Медичний центр ТОВ «Фрезеніус Медикал Кер Україна» у м. Черкаси

Резюме. Наукові дані щодо дозозалежного впливу тривалого активатора рецепторів еритропоєтину (ТАРЕ) на показники оксидативного стресу та перекисного окислення ліпідів мембран еритроцитів у хворих на хронічну хворобу нирок V, які лікуються гемодіалізом (ГД), є обмеженими.

Метою нашої роботи було оцінити взаємозв'язок між застосованою дозою ТАРЕ та інтенсивністю оксидативних процесів у ГД пацієнтів.

Матеріал та методи. 38 ГД пацієнтів були включені до одномоментного обсерваційного дослідження. Протокол дослідження був схвалений локальною етичною комісією ДУ «Інститут нефрології НАМН України». Усі пацієнти надали письму інформовану згоду на участь у дослідженні. Пацієнти були стратифіковані за квартилями ($\leq 25\%$ та $\geq 75\%$) відповідно до середньої дози ТАРЕ і розподілені наступним чином: I групу склали хворі, які отримували ТАРЕ більше 6 місяців у дозі ≤ 50 мкг/міс ($n = 20$), до II групи увійшли хворі, у яких доза ТАРЕ складала ≥ 125 мкг/міс ($n = 18$). Поряд із стандартними діагностичними методами досліджували концентрацію малонового діальдегіду у сироватці (МДА) й еритроцитах (МДАе), вміст у крові церулоплазміну (ЦП) та трансферину (ТР). Крім того, визначали загальну пероксидазну активність (ЗПА) у еритроцитах, проникність еритроцитарних мембран (ПЕМ), підраховували відсоток гемолізу.

Результати. У ГД пацієнтів, які лікувались ТАРЕ у дозі ≥ 125 мкг/міс визначено підвищення концентрації ЦП ($p = 0,007$) та ТР ($p = 0,0003$), тоді як ЗПА була статистично значуще зниженою порівняно з II групою ($p = 0,02$). Крім того, у пацієнтів I групи встановлено вищий відсоток гемолізу ($p = 0,03$) та ПЕМ ($p < 0,0001$). Використовуючи модель пробіт-регресії, ми визначили дозозалежний вплив ТАРЕ на рівень ПЕМ: $\chi^2 = 21$; $p < 0,0001$.

Висновки. Таким чином, застосування ТАРЕ у дозі понад 125 мкг на місяць покращує антиоксидантний статус ГД хворих, але разом з тим, підвищує гемоліз еритроцитів та ПЕМ. Отримані нами дані потребують подальшого підтвердження.

Ключові слова: гемодіаліз, тривалий активатор рецепторів еритропоєтину, оксидативний стрес.

Introduction. Oxidative stress (OS) is a constituent of the inflammatory mechanisms that contributes to anemia in hemodialysis (HD) patients. The intensity of OS is closely correlated with anemia and associated with poor clinical outcomes [1]. One of the possible causes of anemia association with oxidative stress is considered red blood cell (RBC) membrane lipid peroxidation due to chronic hemolysis in HD patients [2].

Nowadays, the prescribing of erythropoiesis-stimulating agents (ESA) plays an indispensable role in clinical practice for the treatment of anemia. Today, regular supplements of intravenous iron and ESA are standard therapies in the treatment of anemia in HD patients [3, 4].

There are a lot of current studies devoted to the effects of ESA on oxidative status in HD patients [5-9]. But, the results of these studies are contradictory.

Several clinical reports have demonstrated the positive impact of ESA on OS status in HD patients [7, 8] while other scientists have found no antioxidant effects of ESA [9, 10].

Continuous erythropoietin receptor activator (CERA) is a newer, longer acting ESA based on its lower frequency of administration [11]. However, at the present time, there have been no reports on the dose-dependent effects of CERA therapy on oxidative stress and red blood cell membrane lipid peroxidation parameters in HD patients.

Objectives. Therefore, the aim of our work was to evaluate whether the dose of CERA treatment affected lipid peroxidation and antioxidant system in HD patients.

Materials and methods. Study Design and Subjects. 38 HD patients were included in this single-center cross-sectional observational study which was conducted at State Institution «Institute of Nephrology of the National Academy of Medical Sciences» in Kyiv, Ukraine. The study protocol was approved by a local Ethics Committee and all patients provided signed informed consent.

The enrolment criteria were: patients aged >18 years who were at least six months on HD treatment, with a

Natalia Stepanova
nmstep@ukr.net

stable clinical condition and normally functioning arteriovenous fistula. We excluded the patients with erythropoietin-resistant anemia, systemic disease, diabetes mellitus, malignancy, acute inflammation processes, immunosuppressive treatment and active hepatitis.

The patients were stratified into quartiles ($\leq 25\%$ and $\geq 75\%$) according to the average dose of CERA and grouped in the following way: Group I (> 6 months of CERA treatment in a low dosage $\leq 50 \mu\text{g}/\text{month}$, $n = 20$) and Group II (> 6 months of CERA treatment in a high dosage $\geq 125 \mu\text{g}/\text{month}$, $n = 18$).

Dialysis prescription. All patients were routinely dialyzed three times a week, 4 h per session with bicarbonate based dialysate, volumetric ultrafiltration control, single use synthetic (polysulphone) dialyzers and heparin as a standard anticoagulant. Dialysis prescription was guided by a goal of achieving a value of $Kt/V \geq 1.2$.

Anemia treatment. Treatment of anemia was carried out in accordance with the clinical protocol of medical care "Treatment of patients with chronic kidney disease stage V with anemia" approved by the Ministry of Health of Ukraine [12]. Erythropoietin was prescribed via a standardized algorithm. All of the 38 HD patients received subcutaneous CERA (methoxy polyethylene glycol-epoetin beta) and intravenous iron replacement therapy. The dose of CERA was adjusted to maintain the individual patient's Hb within a range of $\pm 10 \text{ g/L}$ of the reference hemoglobin (Hb) concentration to achieve a hemoglobin value of 110–120 g/L. The iron dose was adjusted to reach ferritin and transfer saturation (TSAT) levels of 300–400 ng/ml and 30–40%, respectively. 100 mg of iron sucrose was administered in a dilution with 100 ml saline as 30 minute intravenous infusion at the end of HD session. Iron supplementation was temporarily discontinued in the patients with serum ferritin $> 800 \text{ ng/mL}$ or TSAT $> 50\%$ until serum ferritin decreased to $< 800 \text{ ng/mL}$ and TSAT to $< 50\%$ [12].

Methods. All measurements were performed after an overnight fast between 7.00–9.00 a.m. during a midweek non-dialysis day. The blood samples were processed immediately after sampling. Along with the standard diagnostic methods, we defined the content of malondialdehyde levels in the serum (MDAs) and erythrocytes (MDAe) spectrophotometrically as an indicator of lipid peroxidation. Such parameters as the concentration of ceruloplasmin (CP) and transferrin (TR) in the blood and total peroxidase activity (TPA) in erythrocyte were studied as the indicators of antioxidant system. In addition, we calculated the percentage of hemolysis AND the RBC membrane permeability.

Preparation of red blood cells: red cell suspensions for blood were taken with an anticoagulant (heparin) and centrifuged for 10 minutes at 3000 g. Red cell mass was washed three times with isotonic sodium chloride and centrifuged for 10 minutes at 3000 g. Here is the list of the reagents used in this study: 2,4-dinitrophenylhydrazine, 1,2-phenylenediaminedihydrochloride, Indigo Carmine, (Sigma-Aldrich, USA), transferrin (Fluka), urea, sodium fluoride, sodium acetic, ammonium

iron(III) citrate, potassium iodide, tris, trichloroacetic and thiobarbituric acid (Merck, Germany).

In order to estimate MDA, we used serum and red cell suspensions for blood which was drawn using vials containing of 3.8% sodium citrate and centrifuged for 10 minutes at 3000 g [13].

TPA concentration in erythrocyte: the reaction mixture contained 0.5 mL hemolysate of erythrocyte hemolysate (1:1000) sample, 1 mL of 0.2 M acetic buffer solution (pH 4.9), 1 mL 0.05 mM solution of Indigo Carmine solution. After 5 minutes incubation at 30 C, it was added 0.5 ml 0.03 M solution hydrogen peroxide to the samples. The control samples were added 0.5 ml of distilled water. The reaction was stopped 2 min after the addition of 3 ml of 20% sulfuric acid. The absorbance was measured at 670 nm.

For the purpose of studying CP and TR, it was used 2.5 mL of whole blood which was taken from a vein and centrifuged at 2500 r.p.m. for 5 minutes. **CP concentration:** the reaction mixture contained 0.05 mL sample, 4 mL of 0.4M acetic buffer solution (5.5) and 0.5 mL of 0.5% aqueous solution of 1.2 phenylenediamine dihydrochloride. After 1h incubation at 37 C, it was added 1 mL 3% aqueous solution of sodium fluoride, and, absorbance was measured at 530 nm.

TR concentration: the reaction mixture contained 0.1 mL sample, 1 mL of 0.2% solution of ammonium iron (III) citrate (5.5–5.8). After 30 minutes incubation at room temperature, absorbance was measured at 440 nm. As a standard, a TR solution was used [13].

The rate of hemolysis was calculated based on the measurement of Hb released from the cells relatively to the total amount of Hb in the RBC suspension. Free Hb concentration was determined by cyanmethemoglobin method using Drabkin's reagent. The percentage of hemolysis was calculated using the formula described by K. Janatpour et al [14]. A supernatant volume was calculated from Ht.

Determination of RBC membrane permeability (%) was performed by the method, which is based on the detection of differences in the osmotic stability of erythrocytes in a mixture with different concentrations of isotonic solutions of sodium chloride and urea [15].

Statistical analysis. Analysis and all graphs were performed using MedCalc (Belgium). The average means (M) and standard deviations (SD) or the median (Me) and interquartile ranges [Q25 - Q75] were calculated according to a normal distribution. For the statistical analysis, we used the Student's t-test and nonparametric (U-test) Mann-Whitney. Categorical variables were expressed as proportions, and, chi-square tests were used for the comparison of 2 Groups.

The dose dependence assessment was performed using the probit regression model. Odds ratios (OR) and 95% confidence intervals (CI) were calculated using logistic regression. Correlation analysis was performed using the Spearman non-parametric criterion (r).

Results. The demographic, biochemical and clinical characteristics of the HD patients both with and without CERA treatment are presented in Table 1.

Table 1

Demographic and clinical characteristics of the HD patients

Parameter	Group I (n = 20)	Group II (n = 18)	p
Gender (m/f, %)	60/40	44/56	0.71
Age, years	52.6 ± 4.4	55.1 ± 3.3	0.07
Duration of HD (months)	46.8 ± 14.3	52.8 ± 16.2	0.23
Kt/V	1.31 ± 0.3	1.36 ± 0.8	0.8
Hb (g/L)	109.4 ± 6,8	112.2 ± 3,4	0.12
TSAT (%)	45.6 ± 15.5	52.4 ± 12.02	0.17
Ferritin (ng/mL)	695 [402-725]	723 [425-806]	0,76
CERA dose (µg/month)	50 [25-75]	125 [100-150]	0.0001
iPTH (pg/ml)	527.4 [380-750]	593.4 [390-810]	0.8
Albumins (g/L)	36.1 ± 3.8	37.8 ± 3.1	0.14
P (µmol/L)	1.82 ± 0.7	2.2 ± 1.1	0.2
Ca (µmol/L)	2.21 ± 0.3	2.11 ± 0,7	0.56
CRP (mg/L)	6.8 ± 2.9	6.1 ± 4.01	0.54

Data are presented as mean ± standard deviation, or median [25th–75th percentile], or proportions; Ca: calcium; CERA: continuous erythropoietin receptor activator; CRP: C-reactive protein; iPTH: intact parathyroid hormone; Hb: hemoglobin; P: phosphate; TSAT: transferrin saturation.

Accordingly, the data given in Table 1 represented the evidence that CERA therapy with different dosages showed non-significant difference in most of measured demographic and clinical parameters.

In assessing oxidative stress parameters, we obtained heterogeneous results. The evaluated parameters in the study groups are shown in Table 2.

Table 2

Oxidative stress parameters in the HD patients depending on the monthly overage dose of CERA

Parameter	Group I (n = 20)	Group II (n = 18)	p
MDAs, , µmmol/L	476 [411-565]	540 [437-604]	0.06
MDAe, , µmmol/L	604 [411-772]	617 [450-639]	0.27
CP, g/L	0.13 [0.11-0.17]	0.15 [0.16-0.2]	0.007
TR, g/L	2.3 [1.9-2.2]	2.7 [2.4-2.9]	0.0003
TPA, µmol/min/g Hb	196 [175-216]	204 [179-261]	0.02
Hemolysis (%)	4.79 ± 3.05	7.07 ± 4.3	0.03
RBC membrane permeability (%)	9 [4-13]	21.6 [16.6-28]	< 0.0001

Data are presented as mean ± standard deviation or median [25th–75th percentile] or proportions; CP: ceruloplasmin; MDAs: serum malondialdehyde; MDAe: erythrocytes malondialdehyde; RBC: red blood cells; TR: transferrin; TPA: total peroxidase activity.

The data displayed in Table 2 demonstrated the significantly higher levels of CP ($p = 0.007$) and TR ($p = 0.0003$) in the patients on high CERA dosage treatment. TPA activity in erythrocyte of Group II patients was statistically higher compared to Group I ($p = 0.02$).

But, the results of our study indicated not only the positive antioxidant effects of high doses of CERA in

the HD patients. We determined a statistically high percentage of hemolysis ($p = 0.03$) and RBC membrane permeability ($p < 0.0001$) in the patients who were treated with CERA in a dose ≥ 125 µg/month compared to other patients.

A direct correlation was observed between the rate of hemolysis and TSAT ($r = 0.43$, $p = 0.008$; Fig. 1).

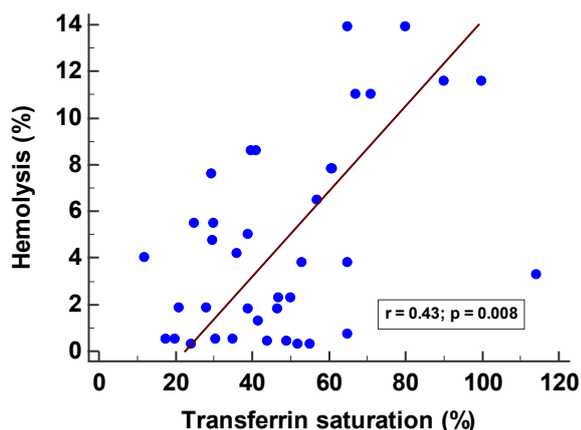


Fig. 1. The correlation between the rate of hemolysis and the serum transferrin saturation level in the HD patients.

Moreover, the level of TPA had a significant negative correlation with the RBC membrane permeability ($r = -0.4$; $p = 0.01$; Fig. 2).

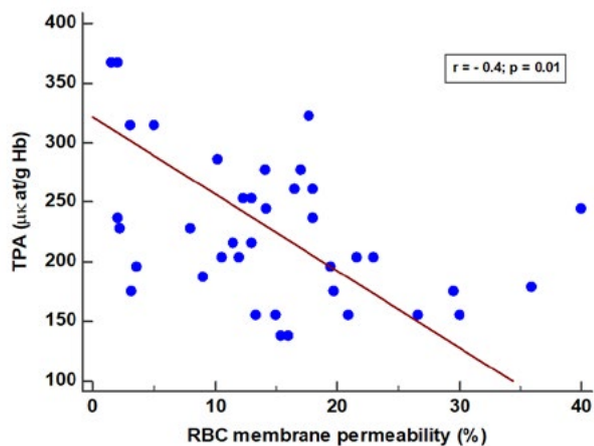


Fig. 2. The correlation between the TPA level and the RBC membrane permeability in the HD patients.

Using the probit regression model, we established the dose-dependent effect of CERA on the level of RBC membranes permeability: $\chi^2 = 21$; $p < 0.0001$ (Fig. 3).

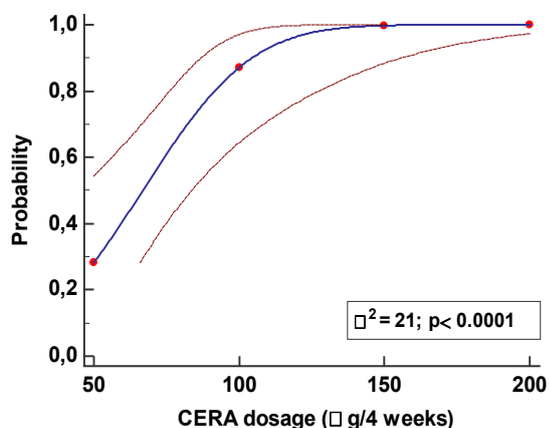


Fig.3. Dose-response plot of a CERA dosage and RBC membrane permeability in the HD patients.

That is, the higher a CERA dosage was applied to achieve the target Hb level in the HD patients, the higher RBC membrane permeability was observed: OR = 17.9, (95% CI 3.03 to 106).

Discussion. There are numerous evidence suggesting excessive oxidative stress in HD patients which can result from loss of antioxidants during dialysis procedures and accumulation of oxidative products [1, 16-18]. Focusing on the association between oxidative stress and anemia, it can be noted that increasing oxidative stress is frequently declared in HD patients who fail to respond to ESA administration [16, 19]. In accordance with these data, we confirmed this position in our study: a tendency of increasing serum MDA level was found in the patients requiring a higher dose of CERA.

Moreover, previous evidence has shown that ESA therapy has an antioxidant effects in HD patients [1, 7, 8, 20, 21]. In this study, we also showed and proved the positive association between antioxidant parameters and a CERA dosage. CP, TR and TPA blood concentration were significantly higher in the HD patients who were treated with a CERA dose ≥ 125 $\mu\text{g}/\text{month}$.

It is well known that the oxidative stress promotes ESA resistance by causing lipid peroxidation of RBC membranes [22]. Erythrocytes are particularly prone to the action of free radicals because they are a potential source of reactive oxygen species. In addition, in order to prevent peroxidation reactions, the oxidative stress depletes the protective mechanisms of RBC [23]. Based on these facts, we found it interesting to compare RBC membrane lipid peroxidation parameters in the HD patients depending on a CERA dose. To our knowledge, studies on the dose-dependent effect of CERA on RBC membrane lipid peroxidation parameters in HD patients are few. It should be noted that our study is not the first to observe the impact of ESA on erythrocyte's oxidative status. But, actually, we are the first to demonstrate a significantly negative effect of a higher CERA dose on RBC membrane lipid peroxidation parameters.

The main finding of the present study was the strong association of a higher CERA dose (≥ 125 $\mu\text{g}/\text{month}$) in the HD patients with the increasing level of RBC hemolysis and membrane permeability levels. As the patient groups with different CERA doses had similar demographic and clinical characteristics in our study, the differences encountered between them with respect to hemolysis and RBC membrane permeability levels could not be attributed to varying degrees of anemia, treatment duration, dialysis quality or inflammation.

In point of fact, there is no current consensus as to the impact of ESA in general and CERA specifically on the intensity of oxidative stress and RBC membrane lipid peroxidation parameters in HD patients. The results we obtained are contrary to the most of existing data. Thus, in a recent experimental study, Aizawa K et al has demonstrated that the intravenous administering of the adequate CERA dose (0.6 $\mu\text{g}/\text{kg}$ every 2 weeks during 5 weeks) improves erythrocyte quality (deform-

ability and life-span) in rats [24]. Zorica M. Dimitrijevic with her colleagues has suggested that long term of ESA administration attenuates the lipid peroxidation process and restores the levels of antioxidants [8]. In an earlier study, Galluci M et al has indicated that the ESA therapy is related to a decrease in RBC membrane oxidative damage [25].

On the other hand, Pawlak et al has showed no effect of one-year ESA therapy on oxidative stress markers in patients undergoing regular HD [9]. Moreover, E. Tatal et al has indicated a significantly higher intensity of oxidative stress in HD patients requiring high doses of EPO. The patients with poor ESA responses had a significantly higher level of MDA and lower levels of plasma superoxide dismutase [26]. The authors have concluded that increased oxidative stress has a strong influence on ESA response in HD patients, and, therefore, it can also be a potential explanation of the data we received.

Thus, according to our findings, we observed that the patients with higher CERA requirements showed a higher antioxidant status simultaneously with higher percents of hemolysis and RBC membrane permeability.

We acknowledged some limitations in our study. First, in this study, the oxidative stress markers were measured just a once. In this way, cross-sectional design of our study could not provide definite information about cause-effect relationship between CERA treatment and oxidative stress. Second, it was a small sample size study performed in a single center; therefore, our findings only revealed associations. Third, we

did not take into account the iron dose which could also affect the results. Finally, there was a high probability of changes in the intensity of oxidative processes in the HD patients with a high level of the comorbidity index.

Despite its limitations, the strong association observed in the present study has indicated the potential impact of a monthly CERA dose $\geq 125 \mu\text{g}$ in the violating of RBC membrane lipid peroxidation. The larger scale well-planned studies are needed for further confirmation of our findings.

Conclusions. In conclusion, we demonstrated that administration of CERA in a dose more $125 \mu\text{g}/\text{month}$ improved the antioxidant status in HD patients. But, at the same time, it increased the hemolysis and RBC membrane permeability. Our preliminary data pointing to the dose-dependent effect of CERA on the RBC membrane lipid peroxidation parameters require further confirmation.

Disclosure Statement. The authors declare no conflict of interest.

Authors' contributions.

N. Stepanova: analyzed and interpreted the patient data, a major contributor in writing the manuscript.

L. Korol: performed the biochemical examination in the blood samples, analyzed and interpreted the patient data.

V. Novakivskyy: interpreted the data.

M. Kolesnyk: idea and management of the research.

References:

1. Liakopoulos V, Roumeliotis S, Gorny X, Dounousi E, Mertens PR. Oxidative Stress in Hemodialysis Patients: A Review of the Literature. *Oxidative Medicine and Cellular Longevity*. 2017;2017:3081856. doi:10.1155/2017/3081856.
2. Tharmaraj D, Kerr PG. Haemolysis in haemodialysis. *Nephrology*. 2017;22: 838-847. doi:10.1111/nep.13119
3. KDIGO Clinical practice guideline for Anaemia in chronic kidney disease. *Kidney Int Suppl*. 2012;2:279-335. doi:10.1038/kisup.2012.37
4. Mikhail A, Brown C, Williams JA, et al. Renal association clinical practice guideline on Anaemia of Chronic Kidney Disease. *BMC Nephrology*. 2017;18:345. doi:10.1186/s12882-017-0688-1.
5. Khalil SKM, Amer HA, Behairy AM, Warda M. Oxidative stress during erythropoietin hyporesponsiveness anemia at end stage renal disease: Molecular and biochemical studies. *Journal of Advanced Research*. 2016;7(3):348-358. doi:10.1016/j.jare.2016.02.004.
6. Fassett RG, Driver R, Healy H, et al. Comparison of markers of oxidative stress, inflammation and arterial stiffness between incident hemodialysis and peritoneal dialysis patients – an observational study. *BMC Nephrology*. 2009;10:8. doi:10.1186/1471-2369-10-8.
7. Siems W, Carluccio F, Radenkovic S, Grune T, Hampl H. Oxidative stress in renal anemia of hemodialysis patients is mitigated by epoetin treatment. *Kidney & Blood Pressure Research*. 2005;28(5-6):295–301. doi: 10.1159/000090184.
8. Dimitrijevic ZM, Cvetkovic TP, Djordjevic VM, et al. How the Duration Period of Erythropoietin Treatment Influences the Oxidative Status of Hemodialysis Patients. *International Journal of Medical Sciences*. 2012;9(9):808-815. doi:10.7150/ijms.4910.
9. Pawlak KI, Pawlak D, Mysliwiec M. Long-term erythropoietin therapy does not affect endothelial markers, coagulation activation and oxidative stress in haemodialyzed patients. *Thromb Res*. 2007;120(6):797-803. doi: 10.1016/j.thromres.2007.02.004
10. Mircescu GI, Căpușă C, Stoian I, Mărăcine M, Muscurel C, Gârneată L, Bărbulescu C. Influence of epoietinum therapy on the oxidative stress in haemodialysis patients. *Nephron Clin Pract*. 2005;100(4):126-32. doi: 10.1159/000085441

11. *Saglimbene VM1, Palmer SC, Ruospo M, Natale P, Craig JC, Strippoli GF.* Continuous erythropoiesis receptor activator (CERA) for the anaemia of chronic kidney disease. *Cochrane Database Syst Rev.* 2017 Aug 7;8:CD009904. doi: 10.1002/14651858.CD009904.pub2.
12. Likuvannia khvorykh na z khronichnu khvorobu nyrok V HD stadii. Adaptovana klinichna nastanova, zasnovana na dokazakh ta unifikovani klinichni protokoly. – K. : «Polihraf plius», 2016. – 228 s.
13. *Korol LV, Mygal LYa, Stepanova NM.* Intensity of oxidative stress and activity of angiotensin converting enzyme in blood of patients with uncomplicated pyelonephritis. *Ukr.Biochem.J.* 2017;89(2):99-105. doi: 10.15407/ubj89.02.099
14. *Janatpour K, Denning L, Nelson K, et al.* Comparison of X-ray vs. gamma irradiation of CPDA-1 red cells. *Vox Sang.* 2005;89:215–219. doi: 10.1111/j.1423-0410.2005.00699.
15. *Kamyishnikov VS.* Spravochnik po klinicheskoy laboratornoy diagnostike: V 2t. Minsk: Belarus; 2002. 463 s.
16. *Bartnicki P, Fija kowski P, Majezyk M, Baszczyk J, Banach M, Rysz J.* Effect of methoxy polyethylene glycol-epoetin beta on oxidative stress in predialysis patients with chronic kidney disease. *Medical Science Monitor : International Medical Journal of Experimental and Clinical Research.* 2013;19:954-959. doi:10.12659/MSM.884024.
17. *Sosa M, Balk ME, Lau J, Liangos O, Balakrishnan VS, Madias NE, Pereira BJG, Jaber BL.* A systematic review of the effect of the Excebrane dialyser on biomarkers of lipid peroxidation. *Nephrology Dialysis Transplantation.* 2006; 21(10):2825–2833. <https://doi.org/10.1093/ndt/gfl376>
18. *Müller C, Eisenbrand G, Gradinger M, Rath T, Albert FW, Vienken J, Singh R, Farmer PB, Stockis J-P, Janzowski C.* Effects of hemodialysis, dialyser type and iron infusion on oxidative stress in uremic patients, free radical research. 2004; 38(10):1093-1100, doi: 10.1080/10715760400011452
19. *Tonon J, Guarnier FA, Cecchini AL, Cecchini R.* Anemia associated with extraerythrocytic oxidative stress damage mediated by neutrophil superoxide anion production in chronic renal failure patients undergoing hemodialysis. *Pathophysiology.* 2012; 19(4):261-8. doi: 10.1016/j.pathophys.2012.07.006.
20. *Katavetin PI, Tungsanga K, Eiam-Ong S, Nangaku M.* Antioxidative effects of erythropoietin. *Kidney Int Suppl.* 2007;(107):10-5. doi: 10.1038/sj.ki.5002482.
21. *Ahmadiasl N, Banaei S, Alihemmati A.* Combination Antioxidant Effect of Erythropoietin and Melatonin on Renal Ischemia-Reperfusion Injury in Rats. *Iranian Journal of Basic Medical Sciences.* 2013;16(12):1209-1216. <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3933796/>
22. *Bamgbola OF.* Pattern of resistance to erythropoietin-stimulating agents in chronic kidney disease. *Kidney International.* 201;80(5):64-474. doi:10.1038/ki.2011.179
23. *Mohanty JG, Nagababu E, Rifkind JM.* Red blood cell oxidative stress impairs oxygen delivery and induces red blood cell aging. *Frontiers in Physiology.* 2014;5:84. doi:10.3389/fphys.2014.00084.
24. *Aizawa K, Kawasaki R, Tashiro Y, Shimonaka Y, Hirata M.* Epoetin beta pegol for treatment of anemia ameliorates deterioration of erythrocyte quality associated with chronic kidney disease. *BMC Nephrology.* 2018;19:19. doi:10.1186/s12882-018-0818-4.
25. *Gallucci MT, Lubrano R, Meloni C, Morosetti M, Manca di Villahermosa S, Scoppi P, Palombo G, Castello MA, Casciani CU.* Red blood cell membrane lipid peroxidation and resistance to erythropoietin therapy in hemodialysis patients. *Clin Nephrol.* 1999;52(4):239-45.
26. *Tutal E, Sezer S, Bilgic A, Aldemir D, Turkoglu S, Demirel O, Ozdemir N, Haberal M.* Influence of oxidative stress and inflammation on rHuEPO requirements of hemodialysis patients with CRP values “in normal range”. *Transplant Proc.* 2007;39(10):3035-40. doi: 10.1016/j.transproceed.2007.06.090.