

Search of new pharmaceuticals on the basis of darbepoetin in the treatment of ischemic stroke (review of literature) / K.M. Reznikov, N.S. Gorbunova, P.D. Kolesnichenko, A.V. Tverskoy, D.A. Kostina, D.A. Bashkatova, V.A. Nikitina // Research result: pharmacology and clinical pharmacology. – 2017. – Vol. 3, Nº1 – P. 125-136.

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SEARCH OF NEW PHARMACEUTICALS ON THE BASIS OF DARBEPOETIN IN THE TREATMENT OF ISCHEMIC STROKE (REVIEW OF LITERATURE)

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Abstract. The article contains the analysis of medical and biological publications from the global database created by the National Centre for Biology Information (NCBI), an intramural biotechnological division of the US National Library of Medicine. The authors have analyzed publications of the recent ten years. Major results of study of erythropoietins and their recombinant analogues have been generalized and systematized. There has been revealed the significant potential of the preparations of this group to be studied and used. Major advantages and drawbacks of erythropoietins and their recombinant analogues have been described. It has been pointed out that a great variety of erythropoietins and darbepoetins speaks to the fact that there is lack of "universe" erythropoietin meeting all the requirements. Genetically modified erythropoietin having both - pharmacokinetics convenient for clinical application and all properties of the natural analogue - is considered to be the most successful darbepoetin. This property has been resulted from the fact that comparing to standard erythropoietin darbepoetin has bigger molecular weight due to introduction of 2 complementary sites of glycosylation. This, in turn, results in the increase of half-life period and, consequently, decreases application frequency of the preparation that makes it more convenient to use comparing to erythropoietin. Application of erythropoietin and its derivatives in stroke therapy in experimental animal models undoubtedly has positive impact on the reduction of the infarction size and dynamics of recovery of neurological status. Results of the analysis demonstrate that the studied preparations are more effective in the early period following stroke than being applied in the later hours. However, there has been revealed some insufficient knowledge in treatment of brain ischemic lesions and ischemic heart disease.

Key words: erythropoietin, darbepoetin, stroke.

Introduction. Medicine is considered to be actively developing science judging from the dynamics of information accumulation.15% of medical database is renewed every year. This means that in the period less than 7 years practically all medical paradigms are being revised. The fact should be taken into consideration that medical practitioners are being trained for 7-8 years and 2-3 years are necessary to revise, write and publish learning material. Thus, it appears that young health care workers start their practice having knowingly obsolete knowledge and to be updated a young specialist should spend a lot of time studying incoming information. Needless to say, young medical workers have a lot of difficulties learning all new publications appearing during a year. Thus,

currently publication of review articles is essential to guide in the actual information flow of periodic.

The objective of this article is to summarize the most interesting experimental models and research results carried out for the last ten years to search new preparations on the basis of darbepoetin for the ischemic stroke therapy.

Materials and methods. To analyze research articles the authors used global database of medical and biological publications in English language created by the National Centre for Biotechnology Information (NCBI), an intramural biotechnological division of the US National Library of Medicine, which houses the most updated and complete articles on the subject of the review. There have been collected 2386 publications regarding erythropoietin in the text database on



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biotechnological information (NCBI) for the last ten years, almost a thousand more - regarding its recombinant analogue darbepoetin. This statistical fact alone supports essential medical significance of erythropoietins. Notable interest has been registered in the study of neuroprotective effect of erythropoietin and its analogues for the recent decades [1]. For more suitable analysis the authors of the article have designed a table, where methods and results of the investigational trials are presented concisely and comprehensibly. The Table contains research results of 11 studies demonstrating various models of brain damages, describes in details ways and methods of introduction of the investigated preparations and shows data that prove efficiency of application of erythropoietin, as well as its analogues, in various forms [2].

The advantage of application of erythropoietin and its derivatives in the therapy of brain damages of diverse genesis is considered to be the fact that erythropoietin is a cytokine, which is produced inside the body and its production is genetically determined. A stimulus for erythropoietin synthesis is reported to be decrease of the oxygen content in cells resulting in the formation of hypoxia- inducible factor-1(HIF-1) followed by mRNA and, actually, erythropoietin production [3, 4]. An erythropoietin gene contains a complementary HIF-1a region; binding to it triggers transcription of mRNA erythropoietin. HIF-1 expression is defined in cells just in 30 minutes after hypoxia has started. The major application point of erythropoietin action is granulocytic-monocyticmegakaryocytic-erythrocytic burst and colony forming units that have specific receptors. Erythropoietin is responsible for proliferation, differentiation and inhibition of apoptosis in these cells; decreased apoptosis of bone marrow erythroid progenitor cells being the major effect of erythropoietin in these conditions. Erythropoietin takes an effect through surface receptors that are amounted no more than 1000 per one cell. Receptors to erythropoietin are found in the cells of the nerve tissue, ovaries and testes, uterus, in the vascular smooth muscle cells, cardiomyocytes, endotheliocytes, lung and renal tubules epithelium. These cells are not only able to express erythropoietin receptors; some of them are capable to synthesize erythropoietin itself. Presence of these potencies allows assuming that erythropoietin performs functions different some from hematopoietic function [5]. Due to this fact, a number of investigational trials on animals, which support positive effect of erythropoietin and its analogues on the recovery of the brain functions after damages of various etiologies, have increased. The objective of this article is to review the most interesting

experimental models and results of the research studies carried out for the last ten years.



Figure 1. Amino acid sequence of the darbepoetin alpha molecule (replacement of 5 amino acids in polypeptide chain by the method of site-directed mutagenesis allowed creating 5 sites of glycosylation, two of which are in positions 30 and 80). Adapted from Lin F-K et al. Proc Natl Acad Sci USA. 1985; 82: 7580-7584 and Elliot S. et al. Nature Biotechnol. 2003; 21: 414-421.

Erythropoietins of the first generation, recombinant human erythropoietin (rHu-EPO) alpha and beta, were identical to native erythropoietin (EPO) in their chemical structure and represented glycoproteid with molecular weight 30.4 kDa. The structure of an EPO molecule includes a single polypeptide chain comprising 165 aminoacids that is subjected to glycosylation with complementation of 4 complex N-linked hydrocarbon chains (replacement of 3 asparaginic and 1 serine regions); they have several anionic free sialic acid residues determining EPO activity. It appears that not a single homogeneous molecule but a mixture of various isoforms specified on the number of free sialic acid residues is developed. This is associated with the fact that glycosylation of rHu-EPO is a posttranscriptional process and is not under the same strict genetic control as rHu-EPO mRNA translation. N-linked carbon side-chains pre-synthesize with various enzymes and are available for post-translational complementation to polypeptide rHu-EPO. Each isoform has its own bioactivity. Isoform 14 has the most erythropoietic activity. On the other hand, isoforms with the less number of sialic acid residues has more EPO receptor (EPOR) affinity, but a shorter period of circulation. Purified alpha and beta EPO



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consist of a mixture of isoforms from 9 to 14. Alpha and beta erythropoietins have a relatively short halflife period that necessitates their introduction into the correction phase as often as 3 times a week, and subcutaneous introduction once a week is allowed only in the maintenance phase of treatment [6, 7].

It is not unexpected that for a long time efforts of clinicians and pharmacologists were aimed at the development of preparations of the new generation with a longer half-life period; that would allow application of more convenient schemes of their introduction (once a week and even once in two weeks). A usual strategy of inhibiting rate of elimination of biomolecules lies in their pegylation, dimerization or synthesis of protein and polypeptide elements [8, 9].

When creating darbepoetin there was applied a novel approach generally aimed at the increase of activity and reduce of clearance including directed reglycosylation-attachment of 2 complementary Nlinked hydrocarbon regions with active sialic residues

base EPO molecule, termed а so to "glycoengineering", or site-directed mutagenesis. As a result there was developed a principally new darbepoetin alpha molecule with a weight up to 37.1 kDa having 5 glycosylation regions, and a number of free sialic groups were adjusted to 22. Darbepoetin alpha has less receptor affinity than EPO alpha and beta that is outweighed by the significantly bigger activity and long half-life period. As EPO alpha and beta preparations darbepoetin is produced by ovarian cells of the Chinese hamster subjected to the incorporation of a darbepoetin gene. Amino acid sequence of darbepoetin differs from that of human EPO in 5 positions, the fact that allows attaching complementary hydrocarbon branches to asparaginic residues in positions 30 and 88 without the destruction of total molecule conformation. Thus, darbepoetin differs from EPO by the high content of carbons and sialic residues, higher molecular weight and an increased negative charge [5, 10, 11].



- 30.400 daltons
- Hydrocarbon content up to 40%
- up to 22 sialic acid residues
- 37.100 daltons
- hydrocarbon content up to 52

Figure 2. Comparing structures of darbepoetin alpha (on the right) and rHuEPO (on the left) molecules. Arrows show 2 complementary tetra-antenna N-linked carbon chains that led to more erythropoietic activity of darbepoetin due to prolongation of half-life period (explanations are given in the text). Adapted from Sinclair AM, Elliot S, 2005

This provides proliferation, differentiation and survival of cells of erythroid lineage. In spite of the decreased receptor affinity darbepoetin has bigger biological activity and approximately tripled half-life EPO alpha 12]. period than [5, As stated above, darbepoetin has a prolonged period of half-life and inhibited clearance comparing to rHuEPO. Exact mechanisms explaining inhibited clearance of darbepoetin alpha and its metabolism are not completely investigated. Decreased EPO level in blood is described by a double decaying exponential; the first phase of rapid decrease may be conditioned



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by binding with endothelial and erythroid cells. When liver, kidneys and bone marrow are considered as possible sites of EPO and darbepoetin degradation, the primary element of its metabolism is erythropoietic tissue through the mechanism of EPO- receptor-inducible uptake. In this case difference between rapid elimination of de-sialylated EPO and inhibited clearance of darbepoetin is caused by the difference in EPO-receptor affinity [13].

Table 1

Comparative characteristic of rHuEPO alpha and beta and darbepoetin alpha (adapted from Deicher R, Horl WH, 2004)

	Epoetin-α	Epoetin-β	Darbepoetin-a	
Hydrocarbons content, %	40	40	52	
Number of N-linked carbon chains	3	3	5	
Half-life period, hours:intravenously,	4-11	8,8-10,4	18-25,3	
subcutaneously	19-25,3	24	48,8	
Bioavailability	30.36	15 50	37	
(subcutaneously), %	50-50	15-50	51	
Clearance (intravenously),	8186	7.0	2.0	
ml/hour \times kg	0,1-0,0	1,9	2,0	
Frequency of introduction	13	053	0.25.1	
(number of times a week)	1-3	0,5-5	0,23-1	

Table 2

Application of erythropoietin and its analogues in various models of the brain damage.

		Refer-			
Author and	Medici-	ence		Method, dosage and	
date of pub-	nal	me-	Model of stroke	time of introduction	Research results
lication	product	dicinal		of medicinal product	
	I	product		· · · · · · · · ·	
1	2	3	4	5	6
Ludmila	Dar-	Human	Right middle	Darbepoetin alpha	Animals were divided into four groups:
Belayeva;	bepoetin	serum	cerebral artery	dosed 10mkg/ kg	1. three-days survival value
Larissa	alpha	albumin	occlusion was	was introduced	(darbepoetin alpha n=8)
Khoutorova	•		performed by	intraperitoneally at	2. three-days survival value
;			ligation of this	the moment of re-	(reference medicinal product n=6)
Weizhao			vessel during	perfusion, i.e. in two	3. two-weeks survival value
Zhao;			two hours; the	hours after the onset	(darbepoetin alpha n=8)
Alexey			method of in-	of middle cerebral	4. two-weeks survival value
Vigdorchik;			traluminal in-	artery occlusion.	(reference medicinal product n=6)
Andrey			troduction of	Human serum	Neurologically significant improvement
Belayev;			poly-L-lysin	albumin (0.25%)	was in animals receiving reference
Raul Busto;			coated suture	10mkg/ kg	medicinal products during the first hour
Ella Magal;			was used. Su-		after re-perfusion.
Myron D.			ture material		Histologically areas of cerebral ischemia
Ginsberg			was introduced		were significantly less in animals
2006 [<u>14</u>]			retrogradely into		receiving darbepoetin comparing to
			the right exter-		animals receiving reference medicinal
			nal carotid ar-		products:
			tery, then into		• In group 3 (two-weeks survival value)
			the internal ca-		$-28,5\pm14,1$ against 68,0±4,5 mm ³
			rotid artery and		• In group 1 (three-days survival value)
			middle cerebral		$-28,9\pm5,3$ against 46,4 $\pm5,3$ mm ³
			artery at the		At that no significant difference in
			distance of 20-		animals between groups were registered.
			22 mm from		
			bifurcation.		



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Table 2 (continued)

1	2	3	4	5	6
Wang Y.;	Recom-	Car-	Middle cerebral	Recombinant human	Neurological status of the studied animals
Zhang	binant	bamylat	artery occlusion	erythropoietin was	was evaluated on the 7^{th} , 14^{th} , 21^{st} and 28^{th}
Z.G.:	human	ed re-	was performed	introduced in doses	days. Histological examination of the brain
Rhodes K.:	ervthro-	com-	by vessels em-	500: 1150 and	was performed in 28 days. Neurological
Renzi M ·	poietin	binant	bolization	$5000 \text{IU} / \text{kg}^2 \text{ in } 6.24$	status carbamylated recombinant human
Zhang R L	poietin	human	conzution.	and 48 hours after	erythropoietin was introduced in doses 50
Kanke Δ ·		eryth_		middle cerebral	mkg/kg^2 of animals receiving for 28 days
$I n M \cdot$		ropoi-		artery occlusion	was better comparing to the group of
Pool C ·		etin		Carbamylated	animals receiving recombinant human
Hoovpor G :		cum		racombinant human	armthropointin was introduced in doses 500:
Chopp M				arythropointin was	$1150 \text{ and } 5000 \text{II}/\text{kg}^2$ Histologically the
2007 [15]				introduced in deses	size of ischamia in the cortical area was loss
2007 [15]				50 mkg/kg ² in 6.24	size of ischemia in the contral area was less
				ond 48 hours often	in annuals receiving recombinant numan
				and 48 nours after	$5000 \text{ H} \text{ J} 150^2 \text{ and } 1000 \text{ H} 1500 1000 \text{ H} 10000 1000 10000 10000 10000 100000 100000 100000 1000000 100000 100000 100000000 10000000 10000000 10000000000000$
					$\frac{500010}{\text{ kg}}$ or carbanylated recombinant
				artery occlusion.	numan erythropoletin in doses 50 mkg/ kg
					(26% and 30% for recombinant human
					erythropoletin in doses 500; 1150 and $\frac{1}{2}$
					500010/ kg respectively, and 36% for
					carbamylated recombinant human
					erythropoletin in doses 50 mkg/ kg ⁻).
					recombinant human erythropoietin in a dose
					50001U/ kg ² significantly decreases
					ischemia area not only in cortical, but also
					in sub-cortical layer by 22% and 36%
					respectively.
Chrystal D	Asi-	Physi-	Middle cerebral	Rats from the experi-	The size of cerebral ischemia was assessed
Price;	aloeryth	ological	artery occlusion	mental group were	by the amount of apoptotic cells and
Zhongjin	ropoi-	saline	was performed	introduced	concentration of activated caspase-3 and 9
Yang;	etin	solution	by vessels em-	asialoerythropoietin	in the area of penumbra on the 4 th day.
Rachel			bolization.	in a dose 20 mkg/ kg,	The size of ischemia significantly
Karlnoski;				1 mcl/ hour for 24	decreased in rats of the experimental
Dipak Ku-				hours.	group comparing to rats from the group of
mar;				Rats from the group	comparison (168±19 mm ³ against 249±28
Raphael				of comparison were	mm ³), the amount of apoptotic cells and
Chaparro;				given physiological	concentration of activated caspase-3 and 9
Enric M				saline solution in a	was also significantly less in the
Gampores				dose 1 mcl/ hour for	experimental group.
2009 [16]				4 days.	
Reitmeir	Recom-	Physi-	Middle cerebral	The animals of the	Functional neurological tests performed
R.;	binant	ological	artery occlusion	experimental group	on the 3^{rd} , 14^{th} and 42^{nd} days after middle
Kilic E.;	human	saline	was performed	were dosed recombi-	cerebral artery occlusion demonstrated
Kilic U.;	erythro-	solution	by vessels em-	nant human erythro-	significant improvement of motor abilities
Bacigaluppi	poietin		bolization.	poietin through a	and coordination on the 14 th and 42 nd days
M.;	ſ			catheter to the left	after ischemia in the animals receiving 10
El Ali A.;				lateral ventricle of	IU of recombinant human erythropoietin a
Salani G.:				the brain in doses 1	day. Immuno-histochemical examination
Pluchino S.:				IU/ day and 10 IU/	performed on the 14 th , 30 th and 52 nd days
Gassmann				day; the medication	after ischemia allowed concluding that a
M.:				was diluted with	high dose of recombinant human
Hermann				0.9% solution of	erythropoietin increased neurons
D.M.				NaCl in the volume	"survival value" on the 52 nd day of



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Table 2 (continued)

1	2	3	4	5	6
1 2010 [<u>16</u>].	2	3	4	5 of infusion 0.25 mcL/ hour. The correctness of ischemia modeling was assessed by the Doppler sonography examination.	6 examination, decreased progressive cerebral atrophy having no affect on the callosal thickness, reduced diffuse astrocytosis and glial scars formation. De- creased level of inflammatory markers (IL-1β, IL-6, leukolysis inhibiting factor, transforming growth factor, tumor necrosis factor, glial fibrillary acid protein) in animals receiving 10 IU of recombinant human erythropoietin per day, which were examined on the 3 rd , 14 th and 30 th day after ischemia also
Morous	Emuthac	Dhuai	I Itanina antany	Decembinent humen	supported positive impact of high doses of recombinant human erythropoietin.
Mazur; Robert H.; Miller Shenandoah Robinson 2010 [17]	poietin	ological saline solution	occlusion during 60 min. on the 18 th day of em- bryogenesis (in- fant rats were born at term, i.e. on the 22 nd day of the embryo- nal develop- ment).	erythropoietin was introduced intraperitoneally starting with the 1 st day of post-natal period. Animals of group 1 were intro- duced 500 IU/kg a day. Animals of group 2 were introduced 1000 IU/kg a day for 3 days. Animals of group 3 were intro- duced 2000 IU/kg a day for 5 days.	performed on the 2 nd , 5 th and 9 th days after the birth. During 2 weeks after intrauterine ischemia there was increased activity of caspase-3 and increased amount of apoptotic cells in the animals receiving erythropoietin in a dose 1000 IU/kg a day for 3 days. Significantly less amount of immune-positive oligodendrocytes was found in alba of the animals receiving physiological saline solution in contrast to the animals receiving erythropoietin in any dosage. Physiological tests showed that animals receiving 2000 IU/kg a day for 5 days did the tasks better comparing to animals from other groups.
Lella Cherian; J. Clay Goodman; Claudia Robertson 2011 [<u>18</u>]	Dar- bepoetin alpha	Physi- ological saline solution	Brain trauma was performed using a crani- otome 8 mm in diametre, which was introduced into the right area of the skull above the pari- etal bone. Then the injury in the form of 8 mm hole was per- formed in a certain position using a "striking tool" and a stem was introduced through this hole; then 3 mm deformation	Darbepoetin was in- troduced by 2.5; 5; 10; 25 and 50 mkg/kg subcutaneously in 5 min., 1 hour, 3 hours, 6 hours, 9 hours 12 hours and 24 hours after the trauma.	 Histological examination of the size of the brain damage showed significant effect of the darbepoetin therapy in doses 25 and 50 mkg/kg 5 min after brain trauma (8.1±3.1 and 11.1±6 mm³, respectively), than in animals treated with physiological saline solution (39.1±6.7 mm). There was also significant effect depending on the time of preparation introduction: When injecting 25 mkg.kg of darbepoetin the size of the damaged area reduced up to 10.5±5 mm³ when introduced after 5 min after trauma; When introduced in 1 hour – up to 9.2±3.6 mm³; When introduced in 3 hours – up to 11.3±2.4 mm³;



Table 2 (continued)

1	2	3	4	5	6
			of the brain was		The size of the damaged area amounted to
			done using this		42.9 ± 11 mm ³ in the animals that were
			stem.		applied physiological saline solution.
Elizarova	Nano-	Native	Localized brain	Nanosomal form of	Dynamics of intracerebral post-traumatic
O.S.;	somal	low co-	hemorrhage	low co-sialyated re-	hematoma development was studied on
Balaban'ya	form of	sial-	(auto-hemor-	combinant human	the 1st, 3rd and 7th day with death
n V.YU.;	low co-	yated	rhagic left brain	erythropoietin and	registration. Study of survival dynamics
SHipulo	sial-	recom-	stroke) was	native low co-	in rats showed that up to the 7th day of
E.V.;	yated	binant	simulated in the	sialyated	observation all false-operated rats
Maksi-	recom-	human	area of internal	recombinant human	survived; in the group of animals with
menko	binant	eryth-	capsule (capsule	erythropoietin were	intracerebral post-traumatic hematoma the
0.0.;	human	ropoi-	interna, coordi-	injected	survival value amounted to 40%.
Vanchugo-	erythro-	etin	nates H=5mm,	intravenously in a	On the background of the repeated 3-days
va L.V.;	poietin	(000	L=3.5 mm, A=2	dose 0.05 mg/kg;	injection of low co-sialyated recombinant
Litvino-va	on the	"Prote1-	mm from	prior to injecting na-	human erythropoletin incorporated into
S.A.;	basis of	novy	bregma)	noparticles	nano-particles from polylactic-co-glycolic
Garibova	nano-	Kontur,		preparation was	acid, the survival value in rats was 77.8%
L.L.;	from	Russia)		lution of Divropio	40% more then in rate of the control
	nolvlac			F68 The first	40% more than in fats of the control
L.A., Gel'ne rine	tic co			introduction was	hematoma
S F	alveolie			nerformed in 3-3.5	Native low co-siglyated recombinant
2012 [19]	acid			hours after the	human erythropoietin did not practically
2012 [17]	stabi-			operation and	influence the survival value in rats with
	lized by			recovery of an ani-	intracerebral post-traumatic hematoma:
	1% hu-			mal after narcosis	this fact might indirectly prove
	man			The repeated	insufficient dose of the introduced
	serum			application was	preparation for producing therapeutical
	albu-			performed on the	concentration in the brain and neuro-
	min.			second and third	protective effect.
				days after the	^
				operation. False-	
				operated rats and rats	
				from the control	
				group with hem-	
				orrhagic stroke were	
				introduced	
				physiological saline	
				solution according to	
	T 1	D1		the same scheme.	
Alexander	Erythro-	Physi-	Ligation of the	Animals were	Statistical analysis of behavioral tests and
M.L.;	poietin	ological	right common	aivided into 2 groups	nistological examination was performed
HIII C.A.;		saime	carotic artery	and o sub-groups:	using dispersion method. Current study
rosenkrant		solution	of two nours;	Ammais of the group	snowed that there was practically no ther-
LI.S.; Fitch DU			hours hypovie	1000 IU/kg of om th	apeutical effect of the filloduction of erythropoietin in 60 or 180 min offer
2012 [1]			under humidi	ropoietin right after	ischemia Assessment of the ventriouler
2012 [1]			fied 8% ovvgen	ischemia	nathology revealed the fact that
			and 97% nitro-	animals of the group	significant swelling of the brain ventricles
			gen	1b were introduced	on the right was registered in the group of
			5~11.	1000 III/kg of	animals receiving physiological saline
			1	1000 10/18 01	anninais receiving physiological sainte



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					Table 2 (continued)
1	2	3	4	5	6
				erythropoietin in 60 min. after ischemia; animals of the group 1c were introduced 1000 IU/kg of eryth- ropoietin in 180 min. after ischemia; animals of the groups 2a, b and c were introduced physiological saline solution in volumes and with time intervals equivalent to each of sub- groups receiving erythropoietin.	solution, and also in animals that were injected erythropoietin in 180 min. after ischemia comparing to the other groups of animals that did not manifested apparent ventricular pathology. In both – animals receiving erythropoietin in 60 and 180 min. after ischemia and in animals receiving physiological saline solution the amount of apoptotic cells and the size of ischemia histologically significantly increased. There was also histologically proved therapeutical effect of erythropoietin in animals receiving erythropoietin right after ischemia. Thus, the study supported the inefficiency of the delayed introduction of erythropoietin.
Carin Sjolund; John-Kalle Lansberg; Tadeusz Wieloch; Karsten Ru- scher; Bertil Rom- ner 2013 [20]	Erythro- poietin	Physi- ological saline solution	Two vessels ten minute ligation with suture ma- terials (right and left carotid ar- teries)	Animals of the first group were single- dosed 80 IU of eryth- ropoietin multiplied by the volume of distribution (VD; 0.057 ml/g BW), intravenously (Neorecormon Roshe, Switzeland) right after the operation; 160 IU/hour in succeeding 72 hours. Animals of the second group were introduced physiological saline solution according to the same scheme.	Examination of sensor-motor functions and memory tests performed on the 3 rd day of the experiment showed that the group of animals receiving erythropoietin did tasks better than the group of animals receiving physiological saline solution. Neuro-protective effect of erythropoietin was also determined by the histological examination of the brain in which neurons of the control animals were specified as big violet cells 30-50 mm in diametre with a large sub-circular nucleus; damaged neurons were specified as red- rosy patches triangle in form with a shrunken dark nucleus. In general histo- logical examination did not reveal significant changes in the brain of the animals receiving erythropoietin as well as in animals receiving physiological sa- line solution. The results of the experiment demonstrated that treatment with erythropoietin did not affect the amount of apoptotic cells, did not protect from the ischemic damage, but preserved synaptic membrane function; behavioral tests and memory tests proved this.
Sheng-Kai Wu; Ming-Tao Yang; Kai-Hsiang Kang; Houng-Chi Liou;	Recom- binant erythro- poietin (Merck KGa, Darm- stadt,	-	Three vessels ligation with suture materials (right and left carotid arteries, middle cerebral artery)	In the study animals were divided into groups depending on the way of the prepa- ration introduction: Group A (control) – 50 min. ischemia; Group B – 50 min.	Histological picture of the brain showed significant decrease of the brain ischemia size in animals receiving erythropoietin using phonophoresis. Neurological status examination was performed in 24 hours after ischemic reperfusion and it was registered that therapy with the help of phonophoresis significantly improved



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Table 2 (continued)

1	2	3	4	5	6
Dai-Hua	Ger-	-		ischemia, animals	neurological functions and reduced
Lu;	many)			were twice	neurological assessment scores.
Wen-Mei	-			introduced 5000	Immuno-histochemical staining of the
Fu;				IU/kg of recom-	brain performed in 24 hours after
Win-Li Lin				binant erythropoietin	ischemia/ reperfusion showed evident
2014 [<u>21</u>]				in 5 hours after re-	neuron death in groups A and C, while in
				perfusion using	groups B and D phonophoresis had fa-
				phonophoresis;	vourable effect on the neurons "survival
				Group C – 50 min.	value'. Thus, the authors concluded that
				ischemia, animals	introduction of recombinant
				were single-dosed in-	erythropoietin using phonophoresis
				travenously 5000	increased penetration of the brain vessels
				IU/kg of	and improved neuroprotective effect of
				recombinant	this preparation.
				erythropoietin in 5	
				hours after re-perfu-	
				sion;	
				Group D – 50 min.	
				ischemia, animals	
				were single-dosed in-	
				travenously 5000	
				IU/kg of	
				recombinant	
				erythropoietin in 5	
				hours after re-perfu-	
				sion using	
				phonophoresis.	
Haiping	Erythro-	-	Occlusion of the	Animals of the first	Neurobehavioral deficiency and the brain
Zhao;	poietin		middle cerebral	group were	ischemia size was less in the animals
Rongliang			artery was per-	introduced 800 IU of	receiving 800 IU of erythropoietin per kg
Wang;			formed for 2	erythropoietin per kg	in the middle cerebral artery intravas-
Xiaoning			hours by	in the middle	cularly. Erythropoietin also suppressed
Wu;			embolization of	cerebral artery in-	expression of stress glucose dependant
Jia Liang;			vessels followed	travascularly.	protein 78 of the endoplasmatic reticulum,
Zhifeng Qi;			by 24 hours	Animals of the	activation of the tumor necrosis factor and
Xiangrong			reperfusion.	second group were	reduced level of pro-apoptotic caspase-3
Liu;				introduced 5000 IU	in micro-vessels of the brain in these
Lianqui				of erythropoietin per	animals. Research results support
Min;				kg subcutaneously.	neuroprotective effect of low doses of
Xunming					erythropoietin (800 IU/kg) when
Ji;					introducing intravascularly in the middle
Yumin Luo					cerebral artery after experimental acute
2015 [21]			1		ischemic brain damage.

Results. As the given Table shows, the most popular model of stroke is considered to be twoor three vessels ligation of arteries; simulation of occlusion of uterine arteries was used for the study of erythropoietin effect on the treatment of intra-uterine fetal hypoxia. The attention should be also paid at the interesting and technically complicated method of localized brain hemorrhage (auto-hemorrhagic left brain stroke) in the area of the internal capsule. All the researchers tended to compare low- and high dose effect of erythropoietin and its analogues and to study their effect in various dosages and using various methods of introduction. The most popular method of assessment of research results was histological validation of ischemia and



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assessment of neurological status performing static and dynamic tests. various When summarizing all the research results it is evident that neurological symptoms in the animals receiving high doses of erythropoietin and its analogues, especially during the first hours after damage, improved significantly; neurobehavioral deficiency reduced; functions of the limbs and memory functions recovered significantly more rapidly; inefficiency of the delayed introduction of erythropoietin and its derivatives was also registered. The brain ischemia size and the amount of apoptotic cells histologically reduced, concentration levels of inflammatory markers reduced. Analyzing the data obtained the authors may conclude that ways of introduction do not affect the outcome of the studied preparations.

Conclusions. The results of the analysis performed support efficiency of application of erythropoietin in various forms, as well as its genetically-modified analogue - darbepoetin. It has been demonstrated in vitro and in vivo that erythropoietin is considered to be a strong inhibitor of neuron apoptosis induced by deficiency. ischemia and oxygen However. erythropoietin blood-forming activity has unfavourable side effect - increased arterial pressure and risk of blood clot formation - that is, in case of ischemic stroke, strongly counter-indicative even if erythropoietin is applied for a very short period. Related to this fact there are known attempts to develop modified erythropoietin having no blood-forming activity but preserving cytoprotective properties. One of such modified erythropoietin variants is reported to be its de-sialyated form, which has high affinity to classical forms of erythropoietin receptors, but fails to reveal bloodforming activity in vivo due to short half-life period in blood plasma. Another variant of modified erythropoietin represents carbamylated erythropoietin. Protein carbamylation is widely known to be a sideeffect of urea application in purification of proteins and as a result of high urea level in the serum. In such cases carbamylation results from urea decomposition into cyanates. Cyanate is responsible for carbamylation of the primary amines of protein in the N-terminal end and amino acid residues of lytic protein subjected to carbamylation. Other amino acid residues possibly subjected to carbamylation are argentine, cysteine, tyrosine, aspartic acid, glutaminic acid, histidine; however, the reaction depends on pH and does not go as rapidly as with the N-terminal end and amino acid residues of lytic protein. Carbamylation of erythropoietin on 7 available lysine residues replaces them on the residues of homocitrullin not involving the profile of glycosylation of the entire molecule. It is demonstrated that carbamylated erythropoietin does not

interact with classical erythropoietin receptors but preserves cytoprotective properties. The major advantage of carbamylated erythropoietin comparing to de-sialyated form of erythropoietin lies in the fact that carbamylation in contrast to de-sialyation does not significantly change kinetic profile. Half-life period of carbamylated erythropoietin in the blood plasma, as is has been showed in rats, is the same as of erythropoietin -3-6 hours; this is caused by preservation of sialic acid residues. Darbepoetin is reported to be geneticallymodified erythropoietin and has all properties of the natural analogue. Comparing to standard erythropoietin darbepoetin has bigger molecular weight (37.1 kDa and not 30.4 kDa) and maximally possible amount of sialic acid residues (22 against 14 in erythropoietin) due to introduction of 2 complementary sites of glycosylation. This results in the increase of half-life elimination period and, consequently, reduces the application frequency of the preparation [5, 23]. There exists one more form of darbepoetin containing carbamylated groups of all eight amino acid residues of lysine included in a darbepoetin molecule and carbamylated amino acid residue of alanine in the N-terminal end of this protein – carbamylated darbepoetin, which does not affect hemopoetic activity but preserves cytoprotective properties. Carbamylated darbepoetin has more prolonged half-life period comparing to Carbamylated erythropoietin and, consequently, and may be prospective when applying in vivo as a cytoprotective medicinal product in case of disorders resulting in cell death due to hypoxia. There is lack of information in literature about the effect of this substance as a medication with cytoprotective action, and the number of investigational pre-clinical trials is not sufficient; this may become a pre-requisite to new research studies.

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