

# Comparative Studies on the Antioxidant Enzymes in Bone Marrow Precursor Cells and Mature Rat Polymorphonuclear Leukocytes

Ms Swechha Mishra<sup>1</sup>

<sup>1</sup> Indian Institute of Information Technology Allahabad

*Received: 15 December 2017 Accepted: 5 January 2018 Published: 15 January 2018*

---

## Abstract

Continuous blood cell production throughout the lifetime of an individual is ensured by the hematopoietic stem cells (HSCs), which are bone marrow cells that possess extensive self renewal capacity and ability to differentiate to all the blood cell lineages. The redox status of the bone marrow stroma is an important factor determining whether the HSCs would differentiate or remain quiescent. Lower levels of ROS (reactive oxygen species) are required to maintain HSC quiescence, whereas higher ROS levels drive the HSCs out of quiescence and reduces their self renewal capacity. The ROS levels in the bone marrow should therefore be regulated to maintain the self renewal capacity of HSCs. Endogenous antioxidants play an important role in regulation of the ROS levels. The present study was therefore undertaken to examine the changes in the cellular detoxifying defences against superoxide and H<sub>2</sub>O<sub>2</sub> i.e. superoxide dismutase (SOD), catalase and glutathione peroxidase (GPx), in the precursor cells from the rat bone marrow and in the rat blood polymorphonuclear leukocytes. An increase in the Catalase and GPx activities in the precursor cells over the differentiated cells was seen. However, the SOD levels in the differentiated cells were greater than in the precursor cells. The levels of Nitric Oxide (NO)- an antioxidant and a regulator of superoxide generation- in the precursor cells were also found to be greater than those in the differentiated cells. The results of the present study thus indicate an up regulation of the endogenous antioxidants in hematopoietic precursor cells, which might be related to the maintenance of their self-renewal capacity.

---

**Index terms**— bone marrow precursor cells, polymorphonuclear leukocytes, antioxidants, nitric oxide.

Abstract-Continuous blood cell production throughout the lifetime of an individual is ensured by the hematopoietic stem cells (HSCs), which are bone marrow cells that possess extensive self renewal capacity and ability to differentiate to all the blood cell lineages. The redox status of the bone marrow stroma is an important factor determining whether the HSCs would differentiate or remain quiescent. Lower levels of ROS (reactive oxygen species) are required to maintain HSC quiescence, whereas higher ROS levels drive the HSCs out of quiescence and reduces their self renewal capacity. The ROS levels in the bone marrow should therefore be regulated to maintain the self renewal capacity of HSCs. Endogenous antioxidants play an important role in regulation of the ROS levels. The present study was therefore undertaken to examine the changes in the cellular detoxifying defences against superoxide and H<sub>2</sub>O<sub>2</sub> i.e. superoxide dismutase (SOD), catalase and glutathione peroxidase (GPx), in the precursor cells from the rat bone marrow and in the rat blood polymorphonuclear leukocytes. An increase in the Catalase and GPx activities in the precursor cells over the differentiated cells was seen. However, the SOD levels in the differentiated cells were greater than in the precursor cells. The levels of

## 6 B) ANIMAL SACRIFICE AND SAMPLE COLLECTION I. ISOLATION OF PMNS FROM BLOOD

---

42 Nitric Oxide (NO)-an antioxidant and a regulator of superoxide generation-in the precursor cells were also found  
43 to be greater than those in the differentiated cells. The results of the present study thus indicate an up regulation  
44 of the endogenous antioxidants in hematopoietic precursor cells, which might be related to the maintenance of  
45 their selfrenewal capacity.

### 46 1 Introduction

47 he metabolic activities have variant degree of differences on the basis of tissue or organ where they are expressed,  
48 which leads to differences in their energy requirement. ROS are implicated in expression of several diseases affecting  
49 tissues and systems. ROS are mainly Free radicals, considered to have atoms having lone pair of electrons, which  
50 is formed via breaking of covalent bond [1]. Ros can be categorized in two broad groups Oxygen derived free  
51 radicles and Nitrogen derived free radicals. These reactive entities are produced outside as well as inside of  
52 cell and cell organelles, such as mitochondria [2]. Reactive nitrogen species (RNS) are a family of antimicrobial  
53 molecules derived from nitric oxide (?NO) and superoxide (O 2 - . ) produced via the enzymatic activity of nitric  
54 oxide synthase (NOS) and NADPH oxidase respectively. Nitrosative stress is one of the mechanism where (ROS)  
55 and (RNS) works together to damage microbial cell. Therefore, these two species are often collectively referred to  
56 as ROS/RNS. Functions of various aminoacids can be modified by ROS that may leads to inappropriate actions  
57 of proteins like dimerization and interaction with Fe-S [3]. It may also causes impairment of cell function and  
58 development of morbit condition [4]. Inspite of all the anomiles it causes studies have also showed that ROS are  
59 required in a certain amount to maintain proper functioning of signalling pathways and metabolism [5]. Change  
60 in redox state by these reactive species might be responsible for proper defferentiation of cell. Changes in the  
61 redox balance during differentiation appear to be due to an increase in the rate of O 2 ? generation. The effective  
62 concentration of oxidants depends on cellular levels of antioxidants-molecules which counteract the oxidants the  
63 cells possess several antioxidant enzymes such as superoxide dismutase (which reduces O 2

### 64 2 ?

65 ? to H 2 O 2 ), catalase, and glutathione peroxidase (which reduces H 2 O 2 to H 2 O). In current work we  
66 have indirectly investigated the role of ROS and RNS via doing comparative analysis of antioxidant enzymes in  
67 the precursor cells of bone marrow and differentiated cells of blodod. Since ROS and RNS play a role in cellular  
68 proliferation and differentiation during hematopoiesis, our aim was to compare the level of antioxidant enzymes  
69 in the precursor cells of bone marrow and the differentiated cell of blood.

### 70 3 II.

### 71 4 Materials and Methods

#### 72 5 a) Experimental animal

73 Closed-colony-bred male rats, 4-6 weeks of age and weighing 80-100 g were used. They were kept in plastic cages  
74 with paddy husk bedding in a temperature between 25-28 o C.

#### 75 6 b) Animal sacrifice and sample collection i. Isolation of PMNs 76 from Blood

77 Isolation of PMNs was carried out by the method of Boyum, 1976 [6], Rat blood was collected under ether  
78 anaesthesia by cardiac puncture in sodium citrate (0.129 M, pH 6.5, 9:1 v/v). Blood was centrifuged at 1500  
79 rpm for 20 min at 20 o C and the buffy coat was carefully layered onto HiSep gradient and centrifuged at 2,000  
80 rpm for 20 min at 20 o C .The leukocyte rich layer was transferred to a fresh tube and centrifuged at 2,000g for  
81 20 min at 20 o C . Pellet was washed and suspended in 2 ml PBS {Composition (mM): NaCl 138; KCl 2.7; Na  
82 2 HPO 4 8.1; KH 2 PO 4 1.5; pH 7.4}. PMNs were counted in Neubaur's chamber and the viability of the cells  
83 was determined by the trypan blue dye exclusion test.

84 ii. Isolation of Precursor Cells from Rat Bone Marrow (BMCs) Both the femurs were surgically removed.  
85 Ends of the femur bones were snipped open with fine scissors. A 23-gauge needle fitted on a 5 ml sterile plastic  
86 disposable syringe (Dispovan, Hindustan Syringes & Medical Devices, Faridabad, India) containing 2 ml of  
87 Dulbecco's Modified Eagles Medium (DMEM, GIBCO, USA) was inserted inside the femur from one end and its  
88 content was forcefully ejected into the femoral cavity. As a result, the marrow plug was flushed from the femoral  
89 shaft. The sample was collected in graduated plastic centrifuge tubes. The procedure was repeated two times to  
90 ensure collection of all the cells from the femur. Precursors from bone marrow were isolated according to method  
91 described by [7].Following sedimentation of debris (5-10 min), the clear supernatant containing leukocytes was  
92 removed and the cells were recovered by centrifugation (2000rpm for 10 min at room temperature). The pellet  
93 was washed and then resuspended in PBS. Cell counting was done using the Neubaur's chamber and the viability  
94 of the cells was determined using the trypan blue dye exclusion test.

95 iii.

---

## 96 7 Isolation of Platelets

97 The platelets were obtained by centrifuging whole blood at 1,500 rpm for 20 min at 20 °C. The platelet rich  
98 plasma so obtained was used as the source of platelets. Cell counting was done using the Neubaur's chamber and  
99 the viability of the cells was determined using the trypan blue dye exclusion test.

100 iv. Nitrite Estimation Nitrite content in the rat platelets, neutrophils and precursors (BMCs) was measured by  
101 using the Griess reagent. Hypotonic lysis of the cells was carried out for 5 mins on ice followed by centrifugation  
102 at 2,000rpm for 20 min at 4 °C. The supernatant thus obtained was used for nitrite estimations using the  
103 Griess reagent (1% sulphanilamide, 0.1% N-1 naphthylethylene diamine and 2.5% orthophosphoric acid. Briefly,  
104 the samples were treated with Griess reagent and incubated for 30 min at 37 °C. Concentration of nitrite was  
105 estimated by measuring the absorbance at 548 nm using sodium nitrite as standard [8].

106 v. Myeloperoxidase Activity MPO activity was evaluated following the method of Pember et al, 1983  
107 [9], Neutrophils/ BMCs were freeze-thawed consecutively for three times and then they were hypotonically lysed  
108 for 5 mins. Cetyltrimethyl ammonium bromide (0.3%) was incubated with the cell lysate at 37 °C for 10 min  
109 followed by centrifugation at 3,000rpm for 20 min at 20 °C. Supernatant was taken for evaluation of enzymatic  
110 activity. Enzyme kinetics was run for 3 min at 15 s intervals using TMB (tetra methyl benzidine 0.5%) and H<sub>2</sub>O<sub>2</sub>  
111 (0.2M) at 37 °C in the presence of 1.0 M citrate buffer pH 6.0). One unit of enzyme activity is defined as  
112 the amount of enzyme which causes an increase in absorbance of TMB by 0.4 in 2 mins.

## 113 8 c) Antioxidant enzyme Assays i. Superoxide Dismutase 114 (SOD)

115 The activity of superoxide dismutase (SOD) was assayed by the method of Kakkar, et al, 1984 [10]. Briefly the  
116 assay mixture consisted of sodium pyrophosphate buffer (0.052 M, pH 8.3), phenazine methosulphate (6.5M),  
117 NBT (30M), NADH (52M) and lysed neutrophil, BMC or platelet supernatant. SOD activity was estimated  
118 as the inhibition of chromogen formation due to NADH mediated PMS dependent reduction of NBT and is  
119 expressed in terms of U/ml/10<sup>6</sup> cells. One unit of enzyme is defined as the amount of enzyme, which causes  
120 50% inhibition of chromogen formation under the experimental conditions.

## 121 9 ii. Catalase

122 Catalase enzyme activity was estimated by the method of Aebi et al, 1974 [11] Briefly 100% of the lysed neutrophil,  
123 BMC or platelet supernatant was added to 2.9 ml of buffered substrate (50mM phosphate buffer, pH 7.0 containing  
124 10 mM H<sub>2</sub>O<sub>2</sub>). The decrease in absorbance was monitored at 37 °C for 3mins at intervals of 15 secs. The  
125 activity was calculated using the coefficient for H<sub>2</sub>O<sub>2</sub> at 240 nm (0.041 M<sup>-1</sup> cm<sup>-1</sup>) and is reported as M of  
126 H<sub>2</sub>O<sub>2</sub> utilized per minute per 10<sup>6</sup> cells.

## 127 10 iii. Glutathione Reductase (GR)

128 The activity of glutathione reductase in the neutrophil, BMC, or platelet lysate was carried out by the method  
129 of Theokand et al., 2008 [12] Incubation mixture for GR enzyme assay consisted of 0.1 M phosphate buffer (pH  
130 7.5), 5 mM oxidized glutathione (GSSG), 0.2 mM NADPH and 100 μL enzyme extract in a final volume of 1.5  
131 mL. Addition of GSSG, initiated the enzyme reaction. The decrease in absorbance at 340 nm due to oxidation  
132 of NADPH was monitored. The enzyme activity was calculated by using the extinction coefficient value of 6.2  
133 mM<sup>-1</sup> cm<sup>-1</sup> for NADPH. One unit of enzyme activity was equivalent to one nmol of NADPH oxidised during  
134 the reaction.

## 135 11 III.

## 136 12 Results

### 137 13 a) Catalase

138 In bone marrow cells amount of enzyme activity is 0.933 Unit/10<sup>6</sup> cells and in neutrophils it is 0.365 Unit/10<sup>6</sup>  
139 cells and 0.232 Unit of enzyme/10<sup>6</sup> cells in platelets. The finding shows that the activity of catalase is much  
140 more in the precursor cells and it decreases in differentiated cells.

### 141 14 b) Nitrite (stable end product of nitric oxide synthase)

142 Enzyme activity in bone marrow cells is 0.509, which is decreasing in neutrophil and platelets as 0.112 and 0.009  
143 M/10<sup>6</sup> cells.

### 144 15 c) Myeloperoxidase

145 In case of bone marrow cells myeloperoxidase activity is 1.6894 units/10<sup>6</sup> cells and in neutrophil it is 1.0717  
146 units/10<sup>6</sup> cells. Amount of enzyme activity is decreasing in mature cells.

**16 d) Superoxide dismutase**

Superoxide dismutase activity in bone marrow cells was 0.071units/10<sup>6</sup> cells and in neutrophil it was 0.104 units/10<sup>6</sup> cell. But in platelet the level again decreased to 0.004 units/10<sup>6</sup> cells.

**17 e) Glutathione reductase**

Activity of enzyme in bone marrow cell is 0.066 and it is 0.0001 and 0.0003 in neutrophil and platelets respectively. Enzyme activity of glutathione reductase is decreasing in mature cells.

**18 IV.****19 Discussion**

This study has demonstrated a comparison of the antioxidant enzymes in precursor cells of bone marrow and the differentiated neutrophils and platelets. The level of the antioxidants like catalase, nitric oxide, and glutathione was greater in the precursor cells than in the differentiated cells. In contrast, superoxide dismutase level was much elevated in the differentiated neutrophils than in the precursor bone marrow cell population. The difference in the antioxidant profiling can be attributed to the varying ROS and RNS concentration in the bone marrow microenvironment where precursor cells reside. Hematopoietic stem cells in bone marrow are maintained in the quiescent stage in hypoxic condition in the osteoblast niche [13] whereas in the relatively more oxygenic vascular niche, due to the proximity to blood circulation, stem cells actively proliferate and differentiate [14,15,16]. Which might increase the intracellular ROS level [17] However, increased ROS may result in accumulation of DNA damage and unscheduled activation of senescence mechanisms in the stem cell compartment in the long term. Unregulated accumulation of ROS in hematopoietic stem cells leads to abnormal hematopoiesis (Yalcin et al, 2008) [18]. Thus, tight regulation of oxidative stress in hematopoietic stem cells is essential for normal control of homeostasis in hematopoietic tissues and this may be obtained by regulating the level of antioxidant enzymes. Catalase enzyme is compartmentalized into small granules in both bone marrow precursor cells and peripheral blood neutrophils [19]. Myeloid, erythroid, and megakaryocytic cells all contain substantial amounts of catalase [20] [21]. The number and size of the diaminobenzidine (DAB)-reactive catalase-containing compartments decreased with differentiation of immature myeloid, erythroid, and megakaryocytic cells progressed, [19][22]. This supports our finding of reduced catalase activity in mature peripheral blood neutrophils and platelets than in precursor cells of bone marrow. Similar trend of reduction was found for the level of nitric oxide which was measured in the form of nitrite level in bone marrow cells and peripheral blood neutrophils and platelets. The level was several times elevated in the bone marrow precursor cells probably because in the bone marrow it may have a role in normal and malignant hematopoietic cell differentiation. Earlier studies have also reported that NO can modulate cellular differentiation [23].The enzyme myeloperoxidase (MPO) is synthesized only in myeloid and monocytic cells, making it an important marker of the myeloid lineage. The elevated MPO level in bone marrow precursor cell from our present study is supported by the earlier finding that transcription of the MPO gene is turned on early during the myeloblast stage of myeloid differentiation and is turned off when myeloid precursors are induced to differentiate along any one of a number of pathways [24] Superoxide dismutase (SOD) is responsible for dismutation of superoxide radicals into molecular oxygen and hydrogen peroxide. Catalase, the subsequent enzyme in this pathway, converts hydrogen peroxide into water and molecular oxygen. High levels of SOD1 activity in neutrophils as reported from our study, without a concomitant increase in catalase, could cause escalation of the Fenton reaction producing hydroxyl radical from hydrogen peroxide [25][26]  $H_2O_2$  produces hydroxyl radicals (OH<sup>•</sup>) were generated by a Fenton reaction, involving an ADP-Fe<sup>2+</sup> (or ATP-Fe<sup>2+</sup>) complex. It has been reported that a portion of the OH<sup>•</sup> free radicals derived from  $H_2O_2$ , produced by the action of SOD, play a role in differentiation [27] This may account for the increased SOD level in differentiated cells revealed from our study. Our study also has shown a decline in the glutathione reductase activity in the peripheral blood neutrophils and platelets. Previous studies have reported that glutathione reductase activity decrease as erythroid cell differentiate [28] The higher level of the enzyme in bone marrow precursor cell indicates the greater importance of this enzyme in the precursor cell.

V.

**20 Conclusion**

Cellular oxidants, called reactive oxygen species (ROS), are constantly produced in animal and human cells. Excessive ROS can induce oxidative damage in cell constituents and promote a number of degenerative diseases and aging. Cellular antioxidants protect against the damaging effects of ROS. However, in moderate concentrations, ROS are necessary for a number of protective reactions. ROS are essential mediators of antimicrobial phagocytosis, detoxification reactions carried out by the cytochrome P-450 complex, cellular differentiation and apoptosis which eliminates cancerous and other life-threatening cells this effect of ROS is dose dependent. Endogenous antioxidants do not completely remove ROS in animal and human cells. This raises the question of why, despite the existence of a powerful cellular system of antioxidants, the shortlived ROS are not removed entirely and are permanently present in cells. The reasonable explanation for this phenomenon is that continuously produced ROS are needed to perform some important biological functions. Seemingly, the cells

---

205 are tuned to remove excessive ROS and to leave the required level of oxidants. Thus, an interplay of oxidants and  
206 antioxidants decides the role of free radicals beneficial or deleterious. This depends on the basal concentration  
207 of free radicals and antioxidants in the cells. With this aim in mind, the present study was undertaken to find  
208 out the oxidant and the antioxidant status in precursor and differentiated cells. It was found that with the  
209 exception of SOD, basal level of all other antioxidants-catalase, NO and GR levels were higher in the precursor  
210 cells than in the differentiated cells, pointing towards the role of cellular antioxidants in maintaining the self-  
211 renewal characteristics of the bone marrow cells. Current study is undertaking the elaborative role of ROS in  
212 keeping the naive undifferentiated blood cell confined to that realm itself. On the contrary antioxidants are helping  
213 in differentiation of the blood cells.

## 214 **21 Declaration of competing interest**

215 The authors declare that they have no competing interests.

## 216 **22 Contribution of authors**

217 Swechha Mishra carried out all the experiment and also contributed in designing this whole experiment. Sonia  
218 Chadha, Sayali Mukherji, Sangeeta singh carried out formulation of the methods used in this experiment and  
also helped in drafting the manuscript via reviewing it.

Ms Swechha Mishra

*[Note: ? , Dr. Sonia Chadha ? , Sayali Mukherjee ? & Sangeeta Singh ?]*

Figure 1:

219



## .1 Acknowledgement

220  
221  
222  
223  
224  
225  
226  
227  
228  
229  
230  
231  
232  
233  
234  
235  
236  
237  
238  
239  
240  
241  
242  
243  
244  
245  
246  
247  
248  
249  
250  
251  
252  
253  
254  
255  
256  
257  
258  
259  
260  
261  
262  
263  
264  
265  
266  
267  
268  
269  
270  
271  
272  
273  
274  
275

- This research was supported by Amity University. We are thankful to the institute for believing in us and providing with all the best facilities to work.
- We thank our colleagues Eshan khan, Yashika Aggarwal, Agnivesh sharma, Pranav Tripathi, from Amity University (bio-tech department) who provide insight and expertise that greatly assisted research, although may not agree with all the conclusion of this paper.
- We thank Vaibhav Gupta for assistance with drafting and finalizing and also for the valuable comments that greatly improved the manuscript.
- [Kakkar et al. ()] ‘A modified spectrophotometric assay of superoxide dismutase’. P Kakkar , B Das , P N Viswanathan . *Indian J Biochem Biophys* 1984. 2 p. .
- [Greenberg et al. ()] ‘An in vivo cytokine and endotoxinindependent pathway for induction of nitric oxide synthase II mRNA, enzyme, and nitrate/nitrite in alveolar macrophages’. S S Greenberg , J Xie , X Zhao , O Jie , T D Giles . *Biochem Biophys Res Commun* 1996. 227 p. .
- [Denton et al. ()] ‘Biochemical and enzymatic changes during erythrocyte differentiation. The significance of the final cell division’. M J Denton , N Spencer , Hrv Arnstein . *Biochem J* 1975. 146 p. 205.
- [Aebi ()] ‘Catalase in vitro’. H Aebi . *Meth. Enzymol* 1984. 105 p. .
- [Boon et al. ()] ‘Catalase Structural Tutorial Text. Boyum’. E M Boon , A Downs , D Marcey . *Scand J Immunol* 2007. 5 p. .
- [Sheokand et al. ()] ‘Effect of nitric oxide and putrescin on antioxidative responses under NaCl stress in chickpea plants’. S Sheokand , A Kumari , V Sawhney . *Physiol. MoBiol. Plant* 2008. 14 p. .
- [Yim et al. ()] ‘Enzyme function of copper,zinc superoxide dismutase as a free radical generator’. M B Yim , P B Chock , E R Stadtman . *J Biol Chem* 1993. 268 p. .
- [Russanov et al. ()] ‘Enzymes of oxygen metabolism during erythrocyte differentiation’. E M Russanov , M D Kirkova , M S Setchenska , Hrv Arnstein . *Bioscience Reports* 1981. 1 p. .
- [Breton-Gorius and Vainchenker ()] ‘Expression of platelet proteins during the in vitro and in vivo differentiation of megakaryocytes and morphological aspects of their maturation’. J Breton-Gorius , W Vainchenker . *Semin Hematol* 1986. 23 p. 43.
- [Breton-Gorius and Guichard ()] ‘Fine structural and cytochemical identification of microperoxisomes in developing human erythrocytic cells’. J Breton-Gorius , J Guichard . *Am J Pathol* 1975. 79 p. 523.
- [Yalcin et al. ()] ‘Foxo3 is essential for the regulation of ATM and oxidative stress-mediate homeostasis of hematopoietic stem cells’. S Yalcin , X Zhang , J P Luciano , D Marinkovic , C Vercherat , A Sarkar , M Grisotto , R Taneja , GhaffariS . *J Biol Chem* 2008. 283 (37) p. .
- [Lin and Austin ()] ‘Functional activity of three distinct myeloperoxidase (MPO) promoters in human myeloid cells’. K M Lin , G E Austin . *Leukemia* 2002. 16 (6) p. .
- [Nagy et al. ()] ‘Induction of Granulocytic Maturation in HL-60 Human Leukemia Cells by Free Radicals: A Hypothesis of Cell Differentiation Involving Hydroxyl Radicals’. K Nagy , G Pásti , L Bene , I Z Nagy . *Free Radical Research* 1993. 19 (1) p. .
- [Karlsson ()] ‘Introduction to Nutraology and Radical Formation’. J Karlsson . *Antioxidants and Exercise* 1997. Human Kinetics Press. p. .
- [Cowland and Borregaard ()] ‘Isolation of neutrophil precursors from bone marrow for biochemical and transcriptional analysis’. J B Cowland , N Borregaard . *J Immunol Methods* 1999. 232 p. .
- [Pember et al. ()] ‘Multiple forms of myeloperoxidase from human neutrophilic granulocytes: Evidence for differences in compartmentalization, enzyme activity, and subunit structure’. S O Pember , R Shapira , J M KinkadeJr . *Arch Biochem Biophys* 1983. 221 p. .
- [Pember et al. ()] ‘Multiple forms of myeloperoxidase from human neutrophilic granulocytes: Evidence for differences in compartmentalization, enzyme activity, and subunit structure’. S O Pember , R Shapira , J M KinkadeJr . *Arch Biochem Biophys* 1983. 221 p. .
- [Magrinat et al. ()] ‘Nitric oxide modulation of human leukemia cell differentiation and gene expression’. G Magrinat , S N Mason , P J Shami , J B Weinberg . *Blood* 1992. 80 p. .
- [Aims et al. ()] ‘Oxidants, antioxidants and the degenerative diseases’. B N Aims , M K Shinegava , T M Hagen . *Proc Natl Acad Sci* 1993. 90 p. .
- [Peled-Kamar et al. ()] ‘Oxidative stress mediates impairment of muscle function in transgenic mice with elevated levels of wild-type Cu/Zn-superoxide dismutase’. M Peled-Kamar , J Lotem , I Wirguin , L Weiner , Hermalin A Groner , Y . *Proc Natl Acad Sci* 1997. 94 p. .
- [Cross et al. ()] ‘Oxygen radicals and human disease’. C E Cross , B Halliwell , E T Borish , W A Pryor , B N Ames , R L Saul , J M Mccord , D Harman . *Ann Intern Med* 1987. 107 p. .

## 22 CONTRIBUTION OF AUTHORS

---

- 276 [Ueda et al. ()] 'Redox control of cell death'. S Ueda , H Masutani , H Nakamura , T Tanaka , M Ueno . *Antioxid*  
277 *Redox Signal* 2002. 4 p. .
- 278 [Suda et al. ()] 'Regulation of stem cells in the niche'. T Suda , F Arai , S Shimmura . *Cornea* 2005. 24 p. .
- 279 [Saretzki et al. ()] 'Stress defense in murine embryonic stem cells is superior to that of various differentiated  
280 murine cells'. G Saretzki , L Armstrong , A Leake , M Lako , T Von Zglinicki . *Stem Cells* 2004. 22 p. .
- 281 [Marcus et al. ()] 'Studies on human platelet granules and membranes'. A J Marcus , D Zucker-Franklin , L B  
282 Safier , H L Ullman . *J Clin Invest* 1966. 45 p. 14.
- 283 [Kopp et al. ()] 'The bone marrow vascular niche: home of HSC differentiation and mobilization'. H G Kopp , S  
284 T Avecilla , A T Hooper , S Rafii . *Physiology* 2005. 20 p. .
- 285 [Muller ()] 'The nature and mechanism of superoxide production by the electron transport chain: Its relevance  
286 to aging'. Florian Muller . *AGE* 2000. 23 (4) p. .
- 287 [Li and Li ()] 'Understanding hematopoietic stemcell microenvironments'. Z Li , L Li . *Trends Biochem Sci* 2006.  
288 31 p. .