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Oxidative Stress Induced Carbonyl Group Incorporation to ¹ Human RBC Membrane: Role in Vivo Senescence of Erythrocyte

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8 Abstract

6

This study was focused to identify the link between oxidative stress and senescence of 9 erythrocytes in vivo. To elucidate this mechanism, various modification of RBC membrane 10 proteins and lipids were analyzed invitro, exposing them to oxidative stress and the results 11 were compared with the changes observed in erythrocytes undergoing senescence in vivo. The 12 other objective was to confirm the mechanism of autoantibody mediated removal of aged 13 RBCs. Our results established that increased lipid peroxidation products, followed by the 14 enhanced damage of RBC membrane protein caused increased RBC membrane protein 15 carbonylation, to normal red cells exposed to the in vitro Fe2+ ascorbate induced oxidative 16 stress. It was presumable that these changes were mediated by hydroxyl (?H) radicals. 17 Further, similar changes were also seen in percoll gradient age fractionated high density aged 18 RBCs. 19

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Index terms— human RBC membrane, protein carbonylation, oxidative membrane damage, senescence of RBC, oxygen free radical, ROS, autoantibody to RBC.

23 **1** Introduction

24 he term 'senescence' in the context of normal blood cells implies that the cells are removed from the circulation 25 in an age dependent manner. Life span of erythrocyte varies between species, is exceptionally constant within a species ??1]. Over the last few decades several age related alterations of erythrocyte membrane and mechanisms 26 of damaged or aged RBCs have been investigated ??1, ??]. Out of these oxidative damage to erythrocyte 27 membrane proteins and lipids is presently thought to play key role during senescence of normal RBSs as well 28 as accelerated senescence of pathological red cells seen in thalassemia , sickle cell anaemia etc. The oxidative 29 damage is initiated probably by oxygen free radicals (ROS) and other oxidants produced endogenously [3, ??, 30 ??, ??, ??, ??]. In vitro characterization of density separated red cells also has provided cumulative oxidative 31 damage ??3]. 32

Dense human red cells show moderately increased amounts of methaemoglobin ??9]. Exposing intact 33 erythrocytes or ghosts to oxidizing systems in vitro showed various modifications e.g. lipid peroxidation, cross-34 35 linking and fragmentation of membrane cytoskeletal proteins, binding of hemichrome and heinz body to the inner 36 surface of the membrane, clustering of band-3 protein, loss of membrane free sulphydryl groups, incorporation of 37 carbonyl content in the membrane protein etc ??5, ??, ??0, ??1,12,13,14,15,16,17]. Some of these changes have also been studied in vitro density separated or biotinylated aged erythrocytes and also in pathological red cells of 38 accelerated ageing ??1, ??, ??,18,19,20,21]. Many of these changes are seen in stored RBC in blood bank [22,23]. 39 A possible reason for cumulative oxidative damage to the red cell membrane would be a reduction in the activity 40 of oxidative defense enzymes like, superoxide dismutase (SOD), glutathione peroxidase, glutathione reductase etc 41

42 ??24,25]. Superoxide radicals (? 2) undergo dismutation reaction spontaneously (2 ? 2 + 2H + ?H 2 O 2) or

 $_{\rm 43}$ $\,$ catalyzed by SOD, produces H 2 O 2 (2? $2+2{\rm H}+$ SOD $?{\rm H}$ 2 O 2) [26]. Again, $?\,$ 2 and H 2 O 2 in presence of Fe $\,$

2+ / Cu + / Mn 2+ can produce more reactive ? H radical by Haber Weiss reaction (? 2 + H 2 O 2 Fe2+ / Cu+ 44 /Mn2+?? H+OH-). In biological system, simple salt of Fe 2+ / Cu + / Mn 2+ and H 2 O 2 can also produce 45 ? H radical via Fenton reaction (Fe 2+ +H 2 O 2-?Fe 3+ +? H + OH -) [27] Fe 3+ to Fe 2+ may promote 46 metal catalyzed Haber Weiss reaction, to produce more damaging ?H radicals [28,29]. Hence, Ascorbate has 47 dual role -anti oxidant and pro-oxidant, where latter is exhibited in presence of transition metals [27,30,31,32,33]. 48 Again, it has been shown that membrane of high density red cell is highly susceptible to peroxidative damage 49 exclusively to poly unsaturated fatty acids of membrane in a chain reaction [27, ??,34,35,36,15]. The signals 50 that trigger the removal of aged erythrocytes and the actual mechanisms of such removal, i.e. elucidation of link 51 between oxidative membrane damage and removal of aged cells from circulation are still controversial. However 52 it has been suggested that recognition and removal of aged, infected or damaged RBC involve multiple pathways; 53 mainly antibody independent phagocytosis, antibody dependent phagocytosis and removal by splenic sinuses 54 [37]. In antibody independent removal, oxidatively damaged erythrocytes are phagocytosed by macrophages 55 through scavenger receptors for low density lipo protein (LDL) in absence of opsonizing antibodies [38,39]. In 56 antibody dependent removal of aged RBC, auto antibodies (anti band-3, antigalactosyl antibodies or antibodies 57 to malonaldehyde-protein adduct) bind to the surface of aged RBCs leading to phagocytic removal of aged, 58 59 oxidatively damaged and pathologically damaged red cells by macrophages [40,41,42, ??, ??,43,37,23]. But the 60 nature of such antibodies has been subjected to debate [40,41,42,37,18, ??].

It has been proposed that impaired deformability provides a major route of the destruction of senescent red cells [44,45]. Poor deformability or increased rigidity due to oxidatively modified red cells as well as aged RBC membrane proteins and lipids retard the red cells' movement in the spleen and they may be entrapped in splenic sinuses for destruction [14,18,46,47].

It was clear that although oxidative damage to RBC membrane protein and lipid play key role in the mechanism of senescence of erythrocyte; but there were important lacunae in our present state of knowledge. This investigation led to fill up that lacunae and which in the long run might lead to identification of suitable agents to prolong the survival of endogenous or transfused RBCs in the circulation.

⁶⁹ 2 II.

70 **3** Aims and Objectives

The specific objectives of this work were-a) study of various oxidative modifications of human erythrocyte membrane lipids and proteins exposed in vitro to physiologically relevant oxidizing system like iron and ascorbate. b) study of effects of free radicals and metal chelators on the above system. c) to investigate if oxidative damage to RBC membrane like increased carbonylation led to enhanced binding of immunoglobulins (IgG) to cell surface. d) to study whether the oxidative damage to RBC membrane observed in vitro were also present in aged erythrocytes obtained from the circulation by density gradient method.

78 4 Materials and Methods

Blood obtained from healthy human male and female volunteers of age group 20 to 50 years was collected in 3.2%
sodium citrate solution. Institute's research ethical committee approval was obtained. All the healthy blood
donors were informed the purpose of the study prior to drawing of blood.

82 5 Materials

Percoll, Alpha cellulose, Microcrystalline cellulose, Catalase, Superoxide dismutase (SOD), Diethylene triamine 83 penta acetic acid (DETAPAC), Butylated hydroxy toluene (BHT), Bovine serum albumin (BSA), Polyvinyl 84 difluride (PVDF) membranes, Alkaline phosphatase conjugated goat anti rabbit IgG, Polyclonal anti-DNP 85 antibody, 4-(2-hydroxy ethyl)-1-piperazine ethane sulphonic acid (HEPES), Phenyl methyl sulphonyl fluoride 86 (PMSF), Tween20(Polyoxyethylene sorbitan mono laurate, Sigma ultra) and Ferritin (Type1: from horse spleen) 87 were purchased from Sigma Chemical Co., USA., Biogel P-6 were obtained from Bio Rad, USA., 5-bromo-4 88 chloro-3-indolyl-phosphate/ nitro blue tetrazolium (BCIP/NBT), Tetra methyl benzidine (TMB), Protein-A-89 horse radish peroxidase conjugate, protein -A-agarose column and Human IgG purification kit were obtained from 90 Bangalore Geni Pvt. Ltd., India. Thiobarbituric acid (TBA), Trichloro acetic acid (TCA) were from E.Merck, 91 Germany. 2-Deoxy ribose (DR), Mannitol, Dimethyl sulphoxide (DMSO), L-Ascorbic acid, 2,4-dinitrophenyl 92 hydrazine (DNPH) and all chemicals for Sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-93 PAGE) were purchased from Sisco Research Laboratory, Mumbai, India. All other chemicals used were of 94 analytical grade. 95

⁹⁶ 6 Methods a) Preparation of human erythrocyte ghosts

Human blood collected in 3.2% sodium citrate solution made free from WBC and Platelets by passing through
a column bed of microcrystalline cellulose and alpha cellulose (1:3) as adopted from Beutler et al (1975) [48].
Briefly, microcrystalline cellulose (mean size 50?m,Sigma cell-50) and ?-cellulose were taken in 1:3 ratio by

weight and suspended in phosphate buffered saline (PBS), pH 7.4 and packed in a 10 ml plastic syringe with an

inner diameter of 1.5 cm, so that 5ml column was 1.5 cm by 5 cm. The packed column was washed with PBS, pH 101 7.4. The blood (approx 5 ml) was centrifuged at 1,000 x g for 5 min and the supernatant plasma was removed 102 along with the buffy coat. The packed red blood cells were poured over the cellulose column and washed down 103 with ice cold isotonic PBS, pH 7.4. The eluate was collected (approx. 40-50 ml for 5 ml of blood) and centrifuged 104 at 1,000 x g for 10 min to obtain the erythrocyte pellet. The erythrocytes were routinely recovered from the 105 column up to 80% and the preparations were practically free of leucocytes (> 99.5%) and platelets (> 95%) as 106 verified microscopically. The erythrocyte ghosts were prepared by hypotonic lysis [49]. The pelleted erythrocytes 107 were lysed in 60 volumes of 5 mM sodium phosphate buffer, pH 8.0 followed by centrifugation at 20,000 x g to 108 obtain haemoglobin free white ghosts. The creamy white ghosts were suspended in 50 mM phosphate buffer, pH 109 7.4 and kept frozen at -20° C for subsequent experiments. 110

111 7 b) Estimation of protein

Protein content of erythrocyte membrane ghosts was estimated by the method of Lowry et al (1951) [50] after 112 solubilizing in 1% SDS and using BSA as standard. c) 2-Deoxy ribose (DR) degradation assay Hydroxyl radical 113 114 generation in the incubated mixture of ascorbate and/or Fe 2+ has been measured by DR degradation assay [51]. 115 The reaction mixture in 50 mM phosphate buffer, pH 7.4 contained 1mM DR, 0.5 mM ascorbate, 0.2 mM FeSO 116 4 or 5.6 ?M Ferritin with or without other additions like mannitol (20mM) or 20 mM DMSO in a total volume of 600 ?l. Incubation was terminated after 1 h by addition of 1.4ml of 2.8% TCA to each tube followed by addition 117 118 of 0.6ml of 1% (w/v) TBA. Then all the tubes were heated for 10 min in a boiling water bath. The tubes were 119 cooled briefly and absorbance taken at 532 nm.

¹²⁰ 8 d) SDS-PAGE of erythrocyte membrane ghosts

121 Freshly prepared erythrocyte ghosts with Fe 2+ (0.2mM) and ascorbate (0.5mM) were incubated at 37°C for

122 2 h with or without other additions. Incubation was terminated by addition of electrophoresis sample buffer

123 containing 3% SDS, and 5% ?-mercapto ethanol followed by heating in boiling water bath for 3 min. The

samples were immediately applied for discontinuous SDS-PAGE in reducing condition using 10% separating gel following the method of Laemmli (1970) [52]. Slab gels were stained with coomassie brilliant blue-R and destained

126 in 5% methanol & 7.5% glacial acetic acid mixture.

¹²⁷ 9 e) Lipid peroxidation study

Lipid peroxidation in incubated ghosts was estimated by measuring the production of malonaldehyde (MDA) as 128 described by Ohkawa et al ??1979) ??53,51]. Erythrocyte ghosts incubated with or without ascorbate (0.5mM) 129 and /Fe 2+ (0.2mM) at 37°C up to 2 h with other additions like 0.5 mM BHT, mannitol (20mM), DMSO 130 (20mM) or catalase (50 ?g/ml) in a total volume of 200 ?l. To each 200 ?l of incubation mixture were added 100 131 ? I of 8.0% SDS, 750 ? I of 20% acetic acid (pH 3.5) and 750 ? I of 8.0% aqueous TBA. The samples were heated at 132 100° C for 15 min. The tubes were briefly cooled and 2.5 ml of a mixture of nbutanol and pyridine (15:1, v/v) 133 was added to each tube and vortexed thoroughly. The samples were centrifuged at 4,000 r.p.m. for 10 min. The 134 organic layer was collected and absorbance measured at 532 nm. Amount of MDA was expressed in n moles/mg 135 protein using a molar extinction co-efficient of 1.56 x 10 5 M -1. cm -1 [54]. 136

¹³⁷ 10 f) Estimation of carbonyl content

Erythrocyte ghosts were incubated with or without ascorbate (0.5mM) and Fe 2+ (0.2mM) at 37°C up to 2 h with other additions in a total volume of 200 ?l.

To each tube was added 500 ?l of TCA followed by centrifugation at 3,000 r.p.m. to obtain a precipitate. 140 To each precipitate 500 ?l of 10 mM DNPH in 2 M HCl was added. The tubes were allowed to stand at room 141 temperature for 10 min with occasional vortexing. Then to each sample 500 ?l of 10% TCA was added followed by 142 vortexing and centrifugation at 10,000 x g for 5 min. The supernatant was discarded and the pellet was washed 143 3 times with 1ml ethanol-ethyl acetate mixture (1:1) to remove excess reagent. Precipitated protein as pellet 144 was solubilized in 1ml protein dissolving solution containing 2% SDS, 0.05% EDTA in 0.8 M sodium phosphate 145 buffer, pH 8.0. Appropriate sample and reagent blanks were kept for this assay. Absorbance was taken at 370 146 nm and the carbonyl content of each sample was calculated using a molar absorption coefficient of 22,000 M -1 147 .cm -1 and the results were expressed in terms of n moles/mg protein [55,56,57,58]. 148

¹⁴⁹ 11 g) Estimation of sulphydryl group

Erythrocyte ghosts, in presence or absence of ascorbate (0.5 mM) and Fe 2+ (0.2mM) with other additions were incubated for 2 h at 37°C in 50 mM phosphate buffer, pH 7.4 in a total volume of 200 ?l. At the end of the incubation, erythrocyte membrane proteins were solubilized by the addition of 1 ml of protein dissolving solution containing 2% SDS, 0.05% EDTA in 0.8 M sodium phosphate buffer, pH 8.0. To each sample 50 ?l of DTNB solution (40mg DTNB in 10 ml of 0.1 M sodium phosphate buffer, pH 8.0) was added. A set of reagent blanks were made and absorbance taken at 412 nm after 15 min, The thiol content was estimated using

14 K) SEPARATION OF YOUNG AND AGED ERYTHROCYTES

a molar absorption coof thiols/mg of protein [13,57]. containing IgG was passed through a Bio-gel P-6 column equilibrated in PBS, pH 7.4. IgG was stored in aliquots in presence of 0.1% sodium azide at 4°C.

¹⁵⁸ 12 i) Binding of autologous IgG to oxidatively damaged red ¹⁵⁹ blood cells

Erythrocyte ghosts were incubated with or without Fe 2+(0.2 mM) /ascorbate (0.5mM) in presence or absence 160 of mannitol (20 mM), DMSO (20 mM) or catalase (50 ?g/ml). After the incubation at 37°C for 2h the red cell 161 ghosts were pelleted down. The supernatant from each tube was aspirated and the pellet suspended in PBS, 162 pH 7.4 containing 1% BSA. Samples (100 ?l) of red cell ghost suspension (control, oxidized and oxidized with 163 inhibitors) were taken in micro centrifuge tubes pre-coated with 1% BSA. To each tube was added 100 ?l of IgG in 164 PBS -1% BSA and the tubes were left overnight at 4°C for IgG binding. Appropriate blanks (without membrane 165 suspension) and negative controls (without added IgG) were kept. After the incubation, to each tube was added 166 0.8 ml of PBS-0.2% BSA followed by centrifugation at 10,000 x g for 10 min. The supernatant was removed and 167 the pellet was washed twice with 1 ml of PBS-0.2% BSA. 100 ?l of protein-A-HRP conjugate (diluted 1:10,000) 168 was added to each tube and kept at 4° C for 1 h followed by centrifugation at 10,000 x g for 10 min after addition 169 of 0.8 ml of PBS-0.2% BSA. The pellet was washed twice with PBS-0.2% BSA to remove completely the excess 170 unbound Protein-A-HRP. Subsequently, 100 ?l of the substrate solution TMB/H 2 O 2 was added to each tube 171 and mixed thoroughly. The tubes were kept in dark for 30 min during which a blue colour was developed. The 172 reaction was stopped by addition of 200 ?l of 2 M H 2 SO 4 which also changed the colour of the reaction mixture 173 from blue to yellow. The absorbance was taken at 450 nm. The absorbance of the blank and the appropriate 174 negative control were subtracted from the absorbance of each test sample. The net absorbance of each test 175 sample was a measure of IgG bound to red cell ghosts. IgG bound to oxidized ghosts with or without inhibitors 176 was expressed as ratio to IgG bound to control unoxidized ghosts after appropriate corrections for differences in 177 protein content in individual samples [13,43]. 178

¹⁷⁹ 13 j) Purification of band-3 protein from human erythrocyte

Human erythrocyte ghosts obtained were suspended in a solution containing 1 mM EDTA, 5 mM ?-mercapto 180 ethanol, 0.03 mM PMSF, which was adjusted to pH 7.5 with 1 M NaOH and stirred gently for 18 h at 4°C. The 181 membranes were recovered by centrifugation at $1,00,000 \ge 1$ h and re suspended in 50 mM sodium phosphate 182 buffer, pH 7.4. The membrane pellet was subjected to SDS-PAGE as described earlier. The protein visible in 183 the gel was predominantly band-3 and this alkali treated pellet was used as partially purified band-3 protein. 184 185 Autologous IgG (1,5 mg/ml) was mixed with an equal volume of suspension of partially purified band-3 protein 186 (alkali treated erythrocyte pellet, protein content 2 mg/ml) and kept at 4°C for 2 h followed by centrifugation at 1,00,000 x g for ½ h. The supernatant recovered was used as autologous IgG, depleted of anti band -3 antibodies 187 (IgG -depleted) and used for IgG binding assay [43]. 188

¹⁸⁹ 14 k) Separation of young and aged erythrocytes

Young and aged erythrocyte of human were isolated by isopyenic centrifugation through gradient of "Percoll". 190 a commercially available polyvinyl-pyrrolidonecoated colloidal silica following the method of Rennie et al (1979) 191 [59]. A continuous density gradient of "Percoll" (0-100%) was prepared with the help of a gradient maker and 192 a peristaltic pump and using two stock solutions: one containing 5.263% BSA in water and another containing 193 194 5.263% BSA (w/v) in percoll. 19 volumes of each of these solutions were separately mixed with 1 volume of solution (final pH 7.4) containing 2.66 M NaCl, 0.09 M KCl and 200 mM HEPES, pH 8.5. Thus, resultant 195 mixtures were isotonic containing 5% BSA. 0.8 ml of red cell suspension was brought to room temperature and 196 carefully layered on the top of the continuous percoll gradient (8 ml). Age fractionation of red cell was achieved 197 by centrifugation at 1,100 x g for 9 min at room temperature followed by slow deceleration of the rotor. The cell 198 fractions were then sequentially aspirated from the top of the gradient using a Pasteur pipette attached to the 199 peristaltic pump. Top 20% of the fractionated cells were taken as young cells and bottom 20% of the cells used as 200 aged red cells. The cells were finally washed to free percoll by 3 washes in PBS, pH 7.4 [59]. 1) After preparation 201 of young and aged erythrocyte membrane, protein carbonyl content, free thiol groups, protein cross-linking etc 202 were measured by previously mentioned methods. 203

m) Immunoblot analysis for protein carbonyls in young and aged erythrocyte membrane ghosts

205 Freshly prepared young and aged erythrocyte membrane ghosts were taken in two separate tubes each 206 containing 100 ?l of RBC membrane ghosts and to each of them was added 50 ?l of 18% SDS to solubilized 207 the protein followed by addition of 200 ?l 20 mM DNPH in 10% TFA. A yellow colour was developed after incubating each mixture for 10 min. The samples were then neutralized to orange red colour by slow addition 208 of 2 M Tris in 30% glycerol and 19% ?-mercapto ethanol. The samples were subjected to SDS-PAGE followed 209 by electroblotting in a semi dry electroblotter unit, using a constant current of 2 m.A./cm 2 for 45 min. The 210 transfer of proteins to PVDF membrane was confirmed by staining one portion of the blotted1 Volume XIV Issue 211 VII Version I (D D D D) 212

Year 2014 C containing1% BSA and kept for 2 h for blocking the nonspecific sites. The blocked membrane was 213 incubated at 37°C with shaking in presence of anti-DNP antibody (diluted 1:1000) for 1 h. After the incubation, 214 the PVDF membrane was washed in a washing solution containing 0.1% (v/v) Tween-20 in PBS followed by 215 further incubation for 1 h with alkaline phosphatase conjugated goat anti rabbit IgG. The bands in the PVDF 216 membrane were detected by the addition of BCIP/NBT [55,32]. 217

n) Quantitation of cell bound IgG of young and aged 15218 erythrocytes 219

The young and aged red cells were washed thoroughly in PBS followed by preparation of corresponding membrane 220 ghosts as described earlier. 221

Membranes of young and aged erythrocytes were taken in separate microcentrifuge tubes pre-coated with 1% 222

BSA. To each tube was added autologous IgG in PBS-1% BSA and the tubes were left overnight at 4°C for 223

224 IgG binding. Bound IgG was quantitated as mentioned earlier. IgG bound to aged RBC membrane ghosts was

225 expressed as a ratio of IgG bound to control young RBC membrane ghosts [43].

o) Statistical Analysis 16226

All experiments were checked for reproducibility and statistical significance worked out. All the values in Tables 227 and Charts of result section were presented as mean \pm standard error of mean (SEM). Significance of difference 228 229 between groups were determined following Students't' test for paired and unpaired observations [60].

IV. ascor bate (0.5 mM) for 2 h at 37°C. d) A significant increase in protein carbonyl content in red cell 230 membrane ghosts exposed to iron and ascorbate for a period of 2 h was observed (Fig-??). 231

Results 17 232

$\mathbf{18}$ a) Results presented in 233

The phenomenon was inhibited by hydroxyl radical scavengers like mannitol, DMSO and also by antioxidant 234 enzyme, catalase (Fig-??). The inhibition was statistically significant. e) When erythrocyte ghosts (protein 235 content 0.6 to 1.0 mg/ml were incubated with Fe 2+ (0.2 mM) and ascorbate (0.5 mM) at 37° C, MDA produced 236 per mg protein per 2 h were significantly higher with respect to control erythrocyte ghosts. Lipid peroxidation 237 238 was significantly inhibited (about 90%) by 0.5 mM of BHT, while mannitol (20 mM), DMSO (20 mM) and 239 catalase (50 ?g/ml) did not inhibit MDA production (Table-3). f) g) In order to elucidate the link between 240 oxidative stress and enhanced IgG binding to red cell ghosts several radical scavengers and antioxidant enzymes were used in the binding assay. Table-5 showed that catalase and hydroxyl radical scavengers like mannitol (20 241 242 mM) and DMSO (20 mM) prevented significantly enhanced binding of autologous IgG to oxidized erythrocyte ghosts. h) The parameters of oxidative stress were then measured in density separated erythrocytes in a percoll 243 gradient. There was a statistically significant rise (1.5 fold) in protein carbonyl content in aged RBC membrane 244 ghosts with respect to young red cell membrane ghosts (Fig-??). 245

(1971) [49]. Incubation of erythrocyte membranes with Fe 2+ (0.2 mM) and ascorbate (0.5 mM) did not 246 however produce any cross-linking or protein fragmentation (Lane-b, c & d). 247

i) Immunoblot analysis of young and aged erythrocyte ghosts using anti-DNP antibody also showed increased 248 249 carbonylation of proteins in aged erythrocyte membrane (Fig-??).

blotted membrane was incubated with PBS enzyme catalase (50 ?g/ml) upto 58.9%. Again, it 250

Table-4 showed that a very significant increase (about 4 fold) in the binding of auto logous IgG occurred 251 in oxidized ghosts compared to control. The binding of auto logous IgG to ghosts was quantitated by an 252 immunoassay using Protein-A-HRP conjugate and TMB (substrate). Since it was difficult to estimate the 253 number of IgG molecules bound per ghost cell, the net absorbance at 450 nm for each sample normalized to 254 a protein concentration of 1 mg/ml was calculated and taken as a measure of bound IgG. The net absorbance for 255 each sample was derived by subtracting from the total absorbance, the values for the blank and the appropriate 256 negative control (ghost incubated without IgG). This increased binding of oxidized ghosts was however not seen 257 when anti-band-3 depleted IgG was used for binding assay. 258

259 membrane after it was dried. The other half of the j) Fig- ?? showed that there was no significant decrease in 260 protein thiol content in aged RBC membrane ghosts compared to young RBC membrane ghosts. k) SDS-PAGE 261 of young and aged erythrocyte ghosts showed that there was no such remarkable difference in the band pattern 262 of membrane proteins (Fig- ??), and especially no cross-linked high molecular weight proteins or evidence of protein fragmentation was apparent in aged erythrocyte. 1) As presented in Table-6, autologous IgG binding to 263 aged erythrocyte ghosts was more than 2.5 fold higher than that of young erythrocyte ghosts, when incubation 264 was carried out with IgG concentration of 1 mg/ml. Raising IgG concentration to 2.5 mg/ml, IgG binding to 265 aged red cell ghosts was about 3.7 times more than that of young erythrocyte ghosts. 266 V.

267

268 19 Discussion

Erythrocyte membrane encounters oxidative stress both from cell interior and exterior [3]. Different investigators have studied red cell membrane alterations by subjecting intact RBC or RBC membrane ghosts to different oxidizing systems like ascorbic acid and iron /copper, H 2 O 2 /iron, ADP/Fe 3+, phenyl hydrazine, xanthine / xanthine oxidase etc have been reported [47,31, ??0,43, ??1, ??]. Many of these changes have also been noticed in aged red cells [1, 2, 3, 14, 15] or in damaged RBCs undergoing accelerated senescence in some pathological conditions [19, ??, ??,21,15,16] or in stored RBC in blood bank [22,23].

In this study, mixture of ascorbic acid and iron (Fe 2+) has been used to induce oxidative stress, because both these components are physiologically relevant in the context of oxidative injury to erythrocytes in vivo. Several discrete iron compartments in red cell have been suggested [21]. This iron is bioactive and can valance-cycle between Fe 3+ and Fe 2+ states is capable of generating hydroxyl radicals [31,33,27,29]. Reducing agents like ascorbate can potentiate this mechanism and thereby promote the formation of hydroxyl radicals via Fenton's mechanism [30,29]. The elaborate transport and recycling of ascorbate by red blood cells in vivo have been highlighted by many investigators [28,33].

Results from Table-1 indicate that a mixture of ascorbate and iron (Fe 2+) is an active source of hydroxyl 282 radicals generated by Fenton's reaction as evident from the inhibition of 2-deoyribose degradation by catalase 283 and hydroxyl radical scavengers like mannitol and DMSO. The oxidation of Fe 2+ to Fe 3+ produces ? 2 radicals, 284 285 which by dismutation produces H 2 O 2. H 2 O 2 produce H 2 O 2 [33]. This autoxidation of ascorbate is ? 2 mediated and metal dependent as evident from its inhibition by SOD and DETAPAC [33]. The fact that 286 ascorbate can interact with ferritin to generate ?H radicals as shown in Table-1 implies that a similar reaction 287 between ascorbate and native or denatured haemoprotein in red cells may also lead to the formation of toxic 288 reactive oxygen species(ROS). 289

ROS can induce various forms of protein damage such as cross-linking of polypeptide chains, oxidation of protein thiol (-SH) groups, incorporation of carbonyl groups into protein etc had been reported [57,43,55,22]. In hydroxyl radical generating system like iron/ascorbate, we have examined some of these modifications which are considered as hallmark for oxidative modification [55,57,32]. Under our experimental conditions protein cross-linking or fragmentation were not observed (Fig- 1). Some earlier studies however, showed cross-linking of isolated cytoskeletal proteins of erythrocyte membrane when incubated for prolonged time with haemoglobin /H 296 2 O 2 [46] or with xanthine/ xanthine oxidase [11] etc.

Oxygen free radicals also cause aggregation of proteins, converting the side chains of cysteine, methionine, 297 histidine and tyrosine and form disulphide bonds as the consequence of oxidation of free thiol groups [13,22]. The 298 present results showed no significant decrease in protein thiol content, which was in conformity with the result 299 of Beppu et al (1990) [43]. Beppu et al (1989) also observed from their result of SDS-PAGE of RBC membrane 300 301 proteins in reducing condition no significant cross-linking of RBC membrane protein exposed to oxidative stress 302 with respect to control ghosts [13] which was in conformity of our result (Fig- 1). However, it has been observed 303 by the same observers that SDS-PAGE of RBC membrane proteins in non-reducing condition formed disulphide bond and cross-linking of protein during oxidative stress [13,43]. 304

305 The metal catalyzed oxidation of proteins can introduce carbonyl group at lysine, arginine, proline and threenine residues in a 'site-specific' manner [61]. Free radical damage to amino acid residues and/or reaction with 306 aldehydes or both derived from lipid peroxides could contribute towards generation of protein carbonyl [61]. The 307 carbonyl bearing residues have not been completely identified but gamma-glutamyl semialdehyde appeared to be 308 the major residue [55]. In different purified and crude extracts of protein several workers have reported changes 309 in carbonyl content as a consequence of oxidative modification ??55, 62.57], but on erythrocyte membranes 310 311 such reports are scanty. The results obtained from this study (Fig- ?? further reacts with Fe 2+ to give rise to 312 ?H radicals by Fenton's mechanism [31,26,33]. Ascorbate potentiates the reaction by reductive cycling of iron from Fe 3+ to Fe 2+ [29]. Further ascorbate undergoes autoxidation to ascorbate induced oxidative stress to 313 RBC membranes caused a significant rise in red cell membrane protein carbonyl content which was inhibited by 314 catalase and hydroxyl radical scavengers-mannitol, and DMSO (Fig- ??) which indicated that the increase in 315 protein carbonyl content was mediated by hydroxyl radical attack. 316

In crude tissue preparation increased carbonyl content as a result of oxidant stress may also be secondary to 317 associated lipid peroxidation [57]. During lipid peroxidation, lipid peroxide or hydroperoxide or peroxyl radicals 318 are formed [27]. These preformed lipid peroxidation products which broke down to yet more radical species by 319 reacting with transition metal like iron [27] in turn caused oxidative damage to membrane proteins [57]. This type 320 of damage could not be inhibited by ?H radical scavengers like mannitol, DMSO or anti oxidant enzyme catalase; 321 322 but lipid soluble chain breaking antioxidants like ?-tocopherol and BHT were effective [57]. In this study however, 323 protein carbonylation has been inhibited significantly by hydroxyl radical scavengers like mannitol, DMSO and 324 catalase (Fig-??); but lipid peroxidation of RBC membrane was not inhibitible by these agents (Table -3). This 325 implied that in this system protein oxidation was not secondary to lipid peroxidative damage. The enhanced production of MDA in red cell membranes by Fe 2+ and ascorbate during in vitro incubation was presumably the 326 result of breakdown of pre-formed lipid peroxides or hydro peroxides by Fe 2+ (Table -3). Ascorbate potentiated 327 such iron catalyzed break down of peroxides and hydro peroxides keeping the iron in the reduced (Fe 2+) 328 state [57]. That has been observed in other systems also [27, ??, ??0, ??3]. However, the consequence of such 329 peroxidative damage to membrane was controversial ??8,34]. 330

Autologous IgG binding to red cell surface has been studied by many investigators [40,18,42], but there were several controversies regarding the binding site and antigenic specificity of bound autologous antibody [37,18,42]. The signal that led to enhanced IgG binding to red cell surface also were controversial.

[3]. Clustering of band-3 protein or proteolytic cleavage of band-3 have been suggested by different workers as trigger for enhanced IgG binding to RBC membrane [40,18,41,23,62]. Beppu et al (1990) [43] and later on other scientist ??7] have shown that in vitro enhanced binding of autologous Ig G to red cell surface occured following an oxidant stress. We have tried to confirm that finding. Our results confirmed that increased binding of anti band-3 immunoglobulins to red cell surface following an oxidative stress by iron and ascorbate . However that was prevented by hydroxyl radical scavengers like mannitol, DMSO and anti oxidant enzyme catalase (Table-5). Results presented in Fig- 1 indicated that oxidant stress to red cell ghosts under our experimental conditions

did not lead to protein cross-linking binding of autologous IgG to red cell surface following an oxidative stress .

That obviously implied that the two phenomena were inter-related. It was presumable that the bound IgG in our in vitro assay systems were auto antibodies generated earlier in vivo against oxidatively modified proteins with increased carbonylation.

Various types of membrane alterations have been reported to aged red cells, which were linked to oxidative 345 damage ??2, ??,18,19]. In the present study increased carbonyl content to aged RBC membrane, observed both 346 347 in spectrophotometric and immunedetection assay system (Fig- ??, Fig- ??) with respect to that of young red 348 cell membrane ghosts represented that the aged red cells have been subjected to oxidative stress in vivo which had been considered as the hall mark of oxidative protein damage [55]. That result seemed interesting, as no such 349 information was available in aged erythrocytes except for an isolated study on stored RBC in blood bank [22]. 350 Again, we have observed no significant alteration in membrane protein thiol content of aged red cells compared 351 to young cells (Fig-??). That was in conformity with our in vitro result, where oxidative stress and ascorbate 352 failed to cause measurable change in protein thiol content of RBC membrane ghosts after incubation up to 2 h 353 (Table-2). Our result [63]. 354

Increased binding of autologous IgG to aged red cell membrane have been well documented [40,18,37], although 355 disputes were there regarding precise nature of the auto-antibodies binding to RBC surface [37]. Our results 356 have confirmed this (Table -6 ??able-2). On the other hand ?H radical scavengers like mannitol, DMSO and 357 anti oxidant enzyme catalase prevented both the increased incorporation of carbonyl groups to membrane protein 358 (Fig- ??) and enhanced in our study supported the observation of Kannan et al (1991) [64] and the result was also 359 consistent with elevated IgG binding seen in red cells of other species like mouse, dog etc, using different methods 360 361 for No cross-linking or polymerization of red cell membrane polypeptides were noticed in reducing condition discontinuous SDS-PAGE pattern of aged red cell membranes in comparison to young red cell membranes (Fig-362 ??). These results were also in accordance with the in vitro data where oxidant treatment of RBC membrane 363 ghosts failed to produce any cross-linking or fragmentation of proteins (Fig-1). However Rettig et al ??1999) [2] 364 have reported a covalent, non-reducible, non disulphide cross-linking of globin subunits of denatured haemoglobin 365 with membrane at the end of the RBC's life span. Our failure to observe any cross-linking of membrane proteins 366 in SDS-PAGE of aged erythrocytes membrane might be related to running the electrophoresis of aged RBC 367 membrane ghosts in reducing condition [13] or species difference or method of isolation of aged red cells. by Fe 368 2+ was in agreement with the result of Piccinini et al ??1995) separating aged red cells from young red cells [65, 369 ??]. 370

Since oxidant stress in vitro led to enhanced binding of autologous IgG to red cell surface (Table-4), it might be 371 implicated that the increased binding of autoantibodies to aged erythrocyte membrane was the result of oxidative 372 injury to red cell membrane during in vivo ageing. This had also been suggested by Beppu et al (1990) [43] and 373 Fujino et al (2000) ???]. The nature of the auto antibody binding to aged red cells have not been elucidated in our 374 study. However, since increase in lipid peroxidation products followed by increased damage of RBC membrane 375 protein as increased carbonyl group incorporation was the only noticeable change in the membrane proteins of 376 aged erythrocytes compared to that of control young erythrocyte, it would be tempting to speculate that auto-377 antibodies had been directed against such altered membrane proteins. The antibody dependent phagocytosis 378 of red cells had been considered as an important mechanism of removal of old and damaged red cells from the 379 circulation [40,18, ??, ??]. The oxidative modification of membrane protein during ageing of erythrocytes was 380 therefore, directly linked with the subsequent removal of those cells from circulation. 381

382 **20 VI.**

383 21 Conclusion

384 From the results and discussion of our study, it can be concluded that in vitro oxidant stress to red blood 385 cell ghosts by iron and ascorbate, led to increased lipid peroxidation products followed by enhanced damage to 386 RBC membrane protein, caused increased membrane protein carbonylation; presumably mediated by hydroxyl radicals and that phenomenon was directly linked to enhanced binding of autologous IgG to oxidized ghosts 387 under similar condition of incubation. It was presumable that bound autologous immunoglobulins to oxidized 388 red blood cell membranes represented auto-antibodies generated in vivo against carbonylated proteins. Further, 389 in vivo senescence of red cells was associated with increased membrane protein carbonylation and enhanced 390 binding of autologous IgG to red cell surface. Since removal of aged red cells from the circulation in large 391

measure depended upon antibody dependent phagocytosis of red cells by macrophages, our results indirectly pointed out that oxidative stress induced modification of RBC membrane carbonylated proteins triggered the

³⁹⁴ latter mechanism for removal of RBC by subsequent formation of auto-antibodies.

395 **22** 6.53

Incubation of erythrocyte ghosts was carried out at 37°C for 2 h in phosphate buffer (50 mM, pH 7.4) with or 396 without addition of FeSO 4 (0.2 mM) and ascorbate (0.5 mM) in presence or absence of other additions like 397 BHT (0.5 mM), mannitol (20 mM), DMSO (20 mM) or catalase (50 ?g/ml) followed by measurement of lipid 398 peroxidation as described in the methods. Results presented were the mean \pm SEM of 5 observations (N=5), 399 followed by Student's 't' test (paired). Value marked '*' indicated significant rise of lipid peroxidation with 400 respect to control (erythrocyte ghosts incubated alone with buffer). *p < 0.001. Value marked '?' showed 401 significant inhibition by BHT. ? p < 0.001. Value marked '#' showed no significant inhibition when compared 402 with the value marked '*'. 403

Erythrocyte ghosts incubated with or without FeSO 4 (0.2 mM) and ascorbate (0.5 mM) for 2 h at 37°C and binding of unfractionated autologous IgG (1mg/ml) and anti band-3 depleted autologous IgG (0.8 mg/ml) were measured as described in the methods. Results presented were mean \pm SEM of 6 observations (N=6).



Figure 1: 25 h

406 407 ²

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Figure 2:



Figure 3: 29 /



Figure 4: Figure 1 :



Figure 5: Figure 2 : Figure 3 : 39 Figure 4 :) 2014 CFigure 6 :



Figure 6:



Figure 7:

catalase ? 2H 2 O +

O 2) or by the action of selenium dependent glutathione peroxidas (H 2 O 2 + 2GSH glutathione

Figure 8:

1 showed that the mixture of Fe 2+(0.2 mM) and ascorbate (0.5 mM)produced hydroxyl radicals by Fenton reaction as shown by 2-deoxyribose (DR) degradation assay. The phenomenon was inhibited by hydroxyl radical scavenger like mannitol (20 mM) upto 62.4% and DMSO (20 mM) up to 77.4% or by antioxidant was also observed that ferritin (5.6 ?M) can also produce ?H radical when incubated with 0.5 mM ascorbate (Table-1). Hence iron and ascorbate were physiologically relevant to erythrocyte in vivo and mixture of these components was used as a model system to oxidative stress in vitro. b) Fig-1 showed that SDS-PAGE analysis of membrane proteins of erythrocyte ghosts incubated with Fe

[Note: 2+ and ascorbate. In control sample (Lanea) different protein bands of red cell membranes were visible and named according to Fairbanks et al c)]

Figure 9: Table -

1

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Figure 10: Table 1 :

relationhincreased

 $\mathbf{2}$

Year 2014 1 33 Volume XIV Issue VII Version I D D D D) C (

[Note: Values were not significantly different from corresponding control values as observed from Student's 't' test (paired).]

Figure 11: Table 2 :

3

Incubation mixture

Figure 12: Table 3 :

 $\mathbf{4}$

YearVolume XIV Is-
sue VII VersionI()Incubation mixtureAutologous IgG binding
(ratio to control)Oxidized ghost + Unfractionated IgG4.26 \pm 0.175
Oxidized ghost + Anti band-3 depleted0.98 \pm 0.188
IgG

[Note: 2014 $C \odot$ 2014 Global Journals Inc. (US)]

Figure 13: Table 4 :

$\mathbf{5}$

Incubation mixture	Autologous IgG binding (ratio to control)
Oxidized $ghost + IgG$	4.26 ± 0.176
Catalase treated oxidized $ghost + IgG$	0.85 ± 0.155 *
Mannitol treated oxidized ghost + IgG	$0.91 \pm 0.159 *$
DMSO treated oxidized ghost $+$ IgG	0.85 ± 0.149 *

Figure 14: Table 5 :

6

	Year 2014
	1 35
Concentration Autologous IgG binding	Volume XIV Issue
of IgG 1 (ratio to control) $2.70 \pm$	VII Version I D D
mg/ml $2.5 \ 0.174$? 3.72 ± 0.183 *	D D) C
m mg/ml	
	(
	Concentration Autologous IgG binding of IgG 1 (ratio to control) $2.70 \pm$ mg/ml 2.5 0.174 ? 3.72 ± 0.183 * mg/ml

[Note: Year () 2014 $C \, \odot \,$ 2014 Global Journals Inc. (US)]

Figure 15: Table 6 :

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