The effect of erythropoietin on potassium levels during ischemia reperfusion injury in rats

Abstract

The aim of this experimental study was to examine the effect of erythropoietin on a rat model and particularly a hypoxia reoxygenation (HR) protocol. The effect of that molecule was studied biochemically using blood mean potassium (K) levels.

Materials and methods: 40 rats of mean weight 247.7g were used in the study. Potassium levels were measured at 60minute (groups A and C) and at 120minute (groups B and D) of reoxygenation. Erythropoietin (Epo) was administered only in groups C and D.

Results: Erythropoietin administration decreased the potassium levels insignificantly by 2.21%±3.66% (P=0.5134). Reoxygenation time increased the potassium levels insignificantly by 3.58%±3.63% (P=0.3375). The interaction of erythropoietin administration and reoxygenation time increased the potassium levels insignificantly by 0.18%±2.22% (P=0.9338).

Conclusion: Erythropoietin administration, re-oxygenation time and their interaction have miscellaneous insignificant effects on potassium levels in the narrow context of 2hours. Perhaps, a longer study time or a higher Epo dose may have clearer and significant effects.

Keywords: hypoxia, erythropoietin, potassium, re-oxygenation, ischemia reperfusion

Abbreviations: Epo, erythropoietin; rHuEPO, recombinant human erythropoietin; HPA, hypothalamo pituitary-adrenal

Introduction

Tissue hypoxia and reoxygenation (HR) remain the main causes of permanent or transient damage with serious implications on adjacent organs and certainly on patients health. Although important progress has been made regarding the usage of erythropoietin (Epo) in managing this kind of damages, satisfactory answers to fundamental questions have yet to be found. Fundamental questions such as by what velocity this factor should act at, when it should be administered and in which dosage.

Table 1 The erythropoietin (Epo) influence (±SD) on the levels of some seric variables concerning reperfusion (rep) time

<table>
<thead>
<tr>
<th>Variable</th>
<th>1h rep P-value</th>
<th>1.5h rep P-value</th>
<th>2h Rep P-value</th>
<th>Interaction of Epo and rep P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>White Blood Cells</td>
<td>+24.01%±13.38%</td>
<td>0.0121</td>
<td>+22.09%±9.11%</td>
<td>0.0351</td>
</tr>
<tr>
<td>Hematocrit</td>
<td>+0.14%±2.89%</td>
<td>0.9626</td>
<td>-0.61%±2.37%</td>
<td>0.8072</td>
</tr>
<tr>
<td>Mean Corpuscular Hemoglobin</td>
<td>+0.01%±1.29%</td>
<td>0.9904</td>
<td>+0.67%±0.80%</td>
<td>0.3549</td>
</tr>
<tr>
<td>Platelet Distribution Width</td>
<td>+1.60%±0.80%</td>
<td>0.0765</td>
<td>+1.36%±0.58%</td>
<td>0.0205</td>
</tr>
<tr>
<td>Platelet-Crit</td>
<td>-16.47%±10.40%</td>
<td>0.0921</td>
<td>-13.74%±7.01%</td>
<td>0.0158</td>
</tr>
<tr>
<td>Urea</td>
<td>+21.42%±7.84%</td>
<td>0.0115</td>
<td>+20.11%±7.25%</td>
<td>0.0059</td>
</tr>
<tr>
<td>Uric Acid</td>
<td>+10.13%±15.10%</td>
<td>0.4917</td>
<td>+15.86%±10.21%</td>
<td>0.1408</td>
</tr>
</tbody>
</table>
The effect of erythropoietin on potassium levels during ischemia reperfusion injury in rats

The aim of this experimental study was to examine the effect of Epo on a rat model and particularly on a HR protocol. The effect of that molecule was studied by measuring the blood mean potassium (K) levels.

Materials and methods

Animal preparation

This experimental study was licensed by Veterinary Address of East Attiki Prefecture under 3693/12-11-2010 & 14/10-1-2012 decisions. All settings needed for the study including consumables, equipment and substances used, were a courtesy of Experimental Research Center of ELPEN Pharmaceuticals Co. Inc. S.A. at Pikermi, Attiki. Accepted standards of humane animal care were adopted for Albino female Wistar rats. Normal housing in laboratory 7 days before the experiment included continuous access to water and food. The experiment was acute which meant that awakening and preservation of the rodents was not following the experiment. They were randomly delivered to four experimental groups by 10 animals in each one. Hypoxia for 45 minutes followed by reoxygenation for 60 minute (group A). Hypoxia for 45 minutes followed by reoxygenation for 120 minute (group B). Hypoxia for 45 minute followed by immediate Epo intravenous (IV) administration and reoxygenation for 60 minute (group C). Hypoxia for 45 minute followed by immediate Epo IV administration and reoxygenation for 120 minute (group D). The molecule Epo dosage was 10mg/Kg body weight of animals.

At first, the animals were submitted into prenarcosis followed by general anesthesia. The detailed anesthesiologic technique is described related.\(^2\) Oxygen supply, electrocardiogram and acidimetry were continuously provided during whole experiment performance.

The protocol of HR was followed. Hypoxia was caused by forceps clamping the inferior aorta over renal arteries for 45 minutes and after laparotomic access had been achieved. Re-oxygenation was induced by removing the clamp and there establishment of the inferior aorta patency. The molecules were administered at the time of re-oxygenation, through inferior vena cava after catheterization had been achieved. The K levels measurements were performed within 60 minutes of reoxygenation (for groups A and C) and at 120 minutes of re-oxygenation (for groups B and D). The mean weight of the forty (40) female Wistar albino rats used was 247.7g (Std.Dev: 34.99172g), with min weight ≥165g and max weight <320g. The rats weight could be potentially a confusing factor, e.g. the more obese rats appear to have greater K levels. This suspicion was investigated.

Model of hypoxia-re-oxygenation injury

Control groups: 20 control rats (mean mass 252.5g (Std. Dev: 39.31988g) suffered by hypoxia for 45 minutes and after laparotomic access had been achieved. Re-oxygenation was induced by removing the clamp and there establishment of the inferior aorta patency. The molecules were administered at the time of re-oxygenation, through inferior vena cava after catheterization had been achieved. The K levels measurements were performed within 60 minutes of reoxygenation (for groups A and C) and at 120 minutes of re-oxygenation (for groups B and D). The mean weight of the forty (40) female Wistar albino rats used was 247.7g (Std.Dev: 34.99172g), with min weight ≥165g and max weight <320g. The rats weight could be potentially a confusing factor, e.g. the more obese rats appear to have greater K levels. This suspicion was investigated.

Model of hypoxia-re-oxygenation injury

Control groups: 20 control rats (mean mass 252.5g (Std. Dev: 39.31988g) suffered by hypoxia for 45 minutes followed by re-oxygenation.

Group A: Re-oxygenation lasted for 60 minutes (n=10 controls rats) mean mass 243g (Std. Dev: 45.77724g), mean potassium levels 6.85mmol/L (Std. Dev: 0.8449194 mmol/L) (Table 2).

Table Continued

<table>
<thead>
<tr>
<th>Variable</th>
<th>1h rep</th>
<th>P-value</th>
<th>1.5h rep</th>
<th>P-value</th>
<th>2h Rep</th>
<th>P-value</th>
<th>Interaction of Epo and rep</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Protein</td>
<td>-0.02%±2.47%</td>
<td>0.9904</td>
<td>-1.27%±1.51%</td>
<td>0.3721</td>
<td>-2.52%±2.03%</td>
<td>0.1509</td>
<td>-0.68%±2.48%</td>
<td>0.443</td>
</tr>
<tr>
<td>Alkaline Phosphatase</td>
<td>+0.20%±18.57%</td>
<td>0.9904</td>
<td>+10.70%±12.78%</td>
<td>0.3549</td>
<td>+21.20%±17.11%</td>
<td>0.1509</td>
<td>+5.79%±7.72%</td>
<td>0.443</td>
</tr>
<tr>
<td>Acid Phosphatase</td>
<td>+0.06%±5.79%</td>
<td>0.9904</td>
<td>+3.11%±3.71%</td>
<td>0.3172</td>
<td>+6.16%±4.97%</td>
<td>0.1509</td>
<td>+1.68%±2.23%</td>
<td>0.443</td>
</tr>
<tr>
<td>CPK</td>
<td>+0.15%±14.09%</td>
<td>0.9904</td>
<td>+7.91%±9.44%</td>
<td>0.3549</td>
<td>+15.67%±12.65%</td>
<td>0.1509</td>
<td>+4.28%±5.70%</td>
<td>0.443</td>
</tr>
<tr>
<td>LDH</td>
<td>+0.08%±7.92%</td>
<td>0.9904</td>
<td>+4.48%±5.35%</td>
<td>0.3549</td>
<td>+8.89%±7.17%</td>
<td>0.1509</td>
<td>+2.42%±3.22%</td>
<td>0.443</td>
</tr>
<tr>
<td>Sodium</td>
<td>+0.72%±0.74%</td>
<td>0.3054</td>
<td>+0.21%±0.63%</td>
<td>0.7136</td>
<td>-0.29%±1.09%</td>
<td>0.767</td>
<td>-0.11%±0.38%</td>
<td>0.7531</td>
</tr>
<tr>
<td>Phosphorus</td>
<td>+1.92%±5.25%</td>
<td>0.6982</td>
<td>+3.95%±3.35%</td>
<td>0.21</td>
<td>+5.98%±4.81%</td>
<td>0.293</td>
<td>+2.45%±2.01%</td>
<td>0.2168</td>
</tr>
<tr>
<td>Progesterone</td>
<td>-0.20%±18.65%</td>
<td>0.9904</td>
<td>-8.86%±10.58%</td>
<td>0.3549</td>
<td>-17.53%±14.15%</td>
<td>0.1509</td>
<td>-4.79%±6.39%</td>
<td>0.443</td>
</tr>
<tr>
<td>Mean</td>
<td>+2.91%±9591%</td>
<td>0.6448</td>
<td>+4.39%±9.81%</td>
<td>0.2941</td>
<td>+5.88%±11.93%</td>
<td>0.2282</td>
<td>+2.93%±6.29%</td>
<td>0.3458</td>
</tr>
</tbody>
</table>
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Table 2 Weight and potassium mean levels and Std. of groups

<table>
<thead>
<tr>
<th>Groups</th>
<th>Variable</th>
<th>Mean</th>
<th>Std. Dev</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Weight Potassium</td>
<td>243 g</td>
<td>6.85mmol/l</td>
</tr>
<tr>
<td>B</td>
<td>Weight Potassium</td>
<td>262 g</td>
<td>6.85mmol/l</td>
</tr>
<tr>
<td>C</td>
<td>Weight Potassium</td>
<td>242.8 g</td>
<td>6.44mmol/l</td>
</tr>
<tr>
<td>D</td>
<td>Weight Potassium</td>
<td>243 g</td>
<td>6.94mmol/l</td>
</tr>
</tbody>
</table>

Group B: Re-oxygenation lasted for 120 minutes (n=10 controls rats) mean mass 262g (Std. Dev: 31.10913g), mean potassium levels 6.82mmol/L (Std. Dev: 0.9507891mmol/L) (Table 2).

Erythropoietin group

20 Epo rats (mean mass 242.9g (Std. Dev: 30.3105g) suffered by hypoxia for 45 minutes followed by reoxygenation in the beginning of which 10mg Epo/kg body weight were IV administered.

Group C: Re-oxygenation lasted for 60 minutes (n=10 Epo rats) mean mass 242.8g (Std. Dev: 29.33636g), mean potassium levels 6.44mmol/L (Std. Dev: 0.4742245mmol/L) (Table 2).

Group D: Re-oxygenation lasted for 120 minutes (n=10 Epo rats) mean mass 243g (Std. Dev: 32.84644g), mean potassium levels 6.94mmol/L (Std. Dev: 0.5581715mmol/L) (Table 2).

Statistical analysis: Weight comparison of everyone of 4 rats groups initially was performed with each other from 3 remaining groups applying statistical paired t-test (Table 3). Any emerging significant difference among K levels was investigated whether owed in the above-mentioned significant weight correlations. K levels comparison of every one the 4 rats groups was initially performed with each other from the above mentioned 3 remaining groups applying statistical paired t-test (Table 3). The application of generalized linear models (glm) with the dependent K level variable and independent variables namely, the Epo administration or not, the reoxygenation time and their interaction, was followed. Inserting the rats weight as an independent variable at glm, a non significant relation turns on potassium levels (P=0.0525), which justifies that further investigation is not needed.

Results

The glm application resulted in: Erythropoietin administration decreased the potassium levels insignificantly by 0.145mmol/L (+0.6168402 mmol/L/0.3268402mmol/l) (P=0.5376), also in accordance with paired t-test (P=0.4893).

Reoxygenation time increased the potassium levels insignificantly by 0.235mmol/l (+0.2329153 mmol/L/0.7029154 mmol/l) (P=0.3157), also in accordance with paired t-test (P=0.3592). The interaction of erythropoietin administration and reoxygenation time increased the potassium levels in significantly by 0.0118182 mmol/L/0.2741311mmol/L/0.2977674mmol/l (P= 0.9338). Reviewing the above and Table 3–5 sum up concerning the alteration influence of erythropoietin in connection with reoxygenation time. These observations indicate a short-term miscellaneous effect of Epo on potassium levels. That effect ought not to be into consideration for their biological or clinical impacts, since the results are non-significant. Another trial must be held providing pure alteration direction and significant results.

Discussion

Bibliography lacks references concerning whether hypoxia can influence the potassium levels. Potassium levels influence, a multiple physiological processes, including resting cellular-membrane potential and the propagation of action potentials in neuronal, muscular, and cardiac tissue. Also K influences the hormone secretion and action, the vascular tone, controls the systemic blood pressure and the gastrointestinal motility. It joins in acid–base homeostasis, in glucose and insulin metabolism, in mineralocorticoid action, in renal concentrating ability and in fluid and electrolyte balance. Isolated potassium administration is impossible due to its feature to have a single electron in its outer electron shell, which readily gives up to create an atom with a positive charge - a cation and oxidizes rapidly in air and reacts vigorously with water combining with anions to form salts. Potassium occurs only in ionic salts usually associated with another drug or a factor. This last chemical conjugate probably influences the potassium occurrence. So, the administration of potassium is by means of a salt.

Chiu et al.7 enhanced the protein kinase C-e potassium assisted translocation (PKC) from the cytosol to mitochondria in rat myocardium, with resultant inhibition of the mitochondrial permeability transition through the opening of mitochondrial K (ATP) channels, affording protection against myocardial ischemia-reperfusion (IR) injury by the herbal Danshen-Gegen decoction formulation. Kuhrt et al.9 found a decrease in the expression of inwardly rectifying potassium (Kir) currents, is a characteristic feature of retinal glial (Müller) cells after transient retinal ischemia. During 4 days in vitro, Müller cells displayed a decrease in Kir currents and an increase in transient A-type potassium currents, which was similar to IR in vivo. Nossaman et al.9 showed that the free radical peroxynitrite (ONOO−) has significant vasodilator activity in the hind limb vascular bed of the cat mediated by a cGMP- dependent mechanism.10 Pollesellop et al.10 have shown that levosimendan has vasodilatory and anti ischemic effects mediated via the opening of ATP-sensitive K channels in vascular smooth-muscle cells and also acts on mitochondrial ATP-sensitive K (mito KATP) channels, an action thought to protect the heart against IR damage. Chicco et al.11 abolished the sarcolemmal K (ATP) channel blockade training-induced cardio protection increasing infarct size to 47.5±3.5% of zone at risk (ZAR) in rats. This study demonstrates that resistance to myocardial IR injury is dependent on sarcolemmal K (ATP) activity during ischemia.

Garcia Gonzalez et al.12 have shown that levosimendan exerts a coronary and systemic vasodilatory effect through its ATP-dependent K channel K (ATP) opening properties, as was experienced in patients with cardiogenic shock.13 Bitner et al.14 used either Euro-Collins-aprotinin procurement solution (Apt-EC group) or aprotinin in combination with low -K dextran (LPD) flush solution (Apt-LPD group) for the procurement of donor lungs.13 The associated mortality rate was 40%. There was no mortality in the Apt-EC group and one patient died in the Apt-LPD group due to PTRI- induced graft failure. Severe PTRI increased short-term morbidity and mortality. Reines et al.14 isolated from rat cerebral IR cortex an
endogenous Na(+), K(+)-ATPase inhibitor, termed endothain E (approximately 80 mg original tissue). Its effect on synapticosomal membrane Na(+), and K(+)-ATPase activity binding to cerebral cortex membranes, was found that the endogenous modulator isolated from IR rats was able to inhibit both enzyme activity and ligand binding. Müllenheim et al.19 found that racemiz ketamine blocks K (ATP) channels in isolated cells and abolishes short-term cardio protection against prolonged ischemia. All rabbits were then subjected to coronary IR but controls without ischemic late preconditioning LPC and the drugs (10mg/kg) were given 10 minutes before IR. Racemiz ketamine, but not S (+)-ketamine, blocks the cardio protection induced by ischemic late preconditioning. Reshef et al.18 studied the effect of opening and blocking of ATP-sensitive potassium (K (ATP)) channels in a model of primary rat neuronal cultures, subjected to metabolic IR.18 The metabolic poisoning resulted in a marked decrease in cellular ATP content by 3-fold. In the neurons, the opening of the K (ATP) channels confers protection against an ATP-depleting crisis. Schmidt et al.17 measured Na, K-ATPase concentration more effectively from various parts stable during IR, of porcine and canine myocardium.17 A relationship between higher concentration of Na, K-ATPase and larger pressure work is suggested.

A majority of the following examples concern the influence of K levels fluctuation on Epo and a minority the reverse influence. Vac halp et al.18 found that the spiroxindoles optimized for PK/PD profile (short-acting prolyl hydroxylase (PHD1) inhibitors).18 K channel off-target activity (hERG) was successfully eliminated. Spiro hydantoins represent a class of short-acting PHD1-3 inhibitors causing a robust Epoup-regulation in vivo treating anemia. Zhang et al.19 carried out systematic optimization of parameters in the existence of cations.19 A convenient and sensitive determination of rHuEPO-alpha with a LOD of 0.4Nm was achieved. Tringalig et al.20 found that increased levels of Epo in the hypothalamus may play a role in the control of hypothalamo-pituitary-adrenal (HPA) function modulating CRH release. Epo effects were studied in short-term (1-3h) experiments under basal conditions or after stimulation with 56Mm KCl. Moreover, Epo, inhibited KCl. Epo effects were not mediated by modification of CRH gene expression, either in the absence or the presence of KCl; in this paradigm, KCl per se did not modify CRH gene expression. Epo contributes to the regulation of the HPA axis activation in pathological conditions such as brain ischemia. Casino et al.21 analyzed serum K with the following guideline based targets: 3.5-6.0mmol/L. They suggested a systematic search of the well-known factors that could affect each CPM, for each failed patient. As an example, they screened all patients associated with inadequate responses to epoetin treatments. Ksiazeb et al.22 found insignificant statistical differences in K clearance of intermittent peritoneal dialysis (IPD) patients before and after correction of anemia with (rHuEPO) therapy. Mohini et al.23 promoted an increase in ingestion of more K correcting anemia in (end-stage renal disease) ESRD patients.23 Experience with patients receiving rHuEPO demonstrates that high-flux short-time hemodialysis is effective without significant differences between K groups.

Bahoul et al.24 associated poor prognosis in the ICU with the presence of circulatory failure (shock) and thrombocytopenia.24 Hypoxemia with PaO (2)/FiO(2)< 200 is among the predictive factors of pulmonary embolism (PE) in post-traumatic critically ill patients. Prevention is highly warranted. Bahoul et al.25 also associated PE with a high ICU and in-hospital mortality rate, including hypoxemia with PaO (2)/FiO(2)< 300 among predictive factors of PE.25 Sun et al.26 found severe acute respiratory distress syndrome as the most common manifestation in critically ill patients with 2009 influenza A H1N1 infection in adult.26 Failure to obtain satisfactory oxygenation with high-level ventilation settings within the first 7-days, onset of acute kidney injury and barotrauma, and continuous need for vasopressors portend a poor prognosis. Diez-Tejedor et al.27 suggested treatment of increased plasma glucose levels, although a clinical trial of glucose-insulin-potassium infusions is ongoing.27 Moreover, insulin therapy in critically ill patients, including stroke patients, is safe and determines lower mortality and complication rates. Hypoxemia also worsens the stroke prognosis, and oxygen therapy in case of <92% O2 saturation is recommended. This could help to save more brain tissue to get the best conditions for further specific stroke therapies such as the use of neuroprotective or thrombolytic drugs in the hospital.

**Conclusion**

Erythropoietin administration, re oxygenation time and their interaction have miscellaneous insignificant effects on potassium levels on the narrow context of 2 hours. Perhaps, a longer study time or a higher Epo dose may have clearer and significant effects.

**Acknowledgments**

None.

**Conflicts of interest**

The author declares no conflict of interest.

**References**


