# ANALYSIS AND COMPARISON OF ARTERIAL BLOOD GASES (ABGS) COMPONENTS (pH, pO<sub>2</sub>, pCO<sub>2</sub>) WITH AND WITHOUT EXTENSIVE EXERCISE IN SELECTED GROUPS OF ATHLETES AND NON-ATHLETES

# Humayun Imran Azeemi<sup>1</sup>, Sarah Sughra Asghar<sup>2</sup>, Basit Ansari<sup>1</sup>, Junaid Mahmood Alam<sup>3</sup> and Syed Riaz Mahmood<sup>4</sup>

<sup>1</sup>Department of Health, Physical Education and Sports Science, University of Karachi, Pakistan.

<sup>2</sup>Department of Anatomy, Sir Syed Medical College for Girls, Karachi, Pakistan.

<sup>3</sup>Department of Biochemistry Lab Services and Chemical Pathology, Liaquat National Hospital and Medical College, Karachi, Pakistan.

<sup>4</sup>Department of Pathology, Govt. Lyari general hospital, Karachi. Pakistan.

#### ABSTRACT

Current research studied the changes and comparison in pH, pO<sub>2</sub>, pCO<sub>2</sub> and related parameters in set of 3 groups.

Group I = Non -athletes with no extensive exercise. Group II = Non- athletes with extensive exercise.

Group III = Athletes with extensive exercise.

Blood was collected from radial arteries 30 minutes prior to exercise or walk and 15 minutes after exercise in resting condition from subjects in all three groups. Blood was analyzed for pH, pCO<sub>2</sub>, pO<sub>2</sub>, HCO<sub>3</sub> using Nova Phox Pro Arterial blood gas analyzer (Nova Biomedical, Massachusetts, and USA). Data is presented as pre and post exercise Mean  $\pm$  Standard error of mean (SE) and compared statistically Pre and Post exercise using Students't-test for paired samples having P value < 0.05 as significant.

(Standard error is the standard deviation of sampling distribution)

Group I show mild, moderate and no significant difference when pre and post arterial blood gases (ABGs) parameters were compared with each other thus depicting non ius respiratory/Ventilatory distress. However, both group II and III exhibited one moderate and remaining very markedly significant difference amongst arterial blood gases (ABGs) parameters, Pre, and Post extensive exercises, suggesting activation of corrective, retaliatory and compensatory mechanisms.

Key words: ventilatory, arterial blood gases (ABGs), pCO<sub>2</sub> and pO2.

#### **INTRODUCTION**

It is documented that values of arterial pH, pressure of Oxygen ( $pO_2$ ) and of Carbon dioxide ( $pCO_2$ ) changed with minimal increments during exercise (Burton *et al.*, 2004). However, increased dependence on glycolysis during vigorous exercise leads to production of pyruvate, and then lactate, that eventually cause elevated  $pCO_2$  and declined pH. Besides, compensatory hyperventilation results in a decrease of  $pCO_2$ , decline in further lactic acid production and precluding decline in blood pH as well (Azizi, 2011; Martin, 2008; Litch 2006; Brown *et al*, 2008). In many cases during routine exercise, arterial oxygen and carbon dioxide pressures and ventilatory actions are not abnormal enough to stimulate hyper-respiration (Burton *et al*, 2004; Powers and Howley, 2004). However, it was reported that changes in ventilation is also a linear function of production of carbon dioxide ( $CO_2$ ) volume during light to heavy exercise (Arena *et al.*, 2003; Guazzi *et al.*, 2005; Phillips *et al.*, 2020). When exercise was performed above respiratory compensation threshold, ventilation volume becomes disproportional to carbon dioxide ( $CO_2$ ) volume due to excessive acidosis and lactic acid production (Phillips *et al.*, 2020).

Such changes in arterial blood components and elevation are mostly reported in conditions such as chronic heart failure (CHF), pulmonary hypertension (PH), interstitial lung disease (ILD), and chronic obstructive pulmonary disease (COPD). Significantly, such altered response contributes to dyspnea and exercise intolerance, which might lead to diminished endurance capacity of athletes (Phillips *et al.*, 2020). However, in some cases of compromised pulmonary functions, increasing ventilatory efficiency through anti-coagulant therapy does elevate exercise capacity of that individual (Huang *et al.*, 2020).

Therefore, current research studied the changes and comparison in pH, pO2, pCO2 and related parameters in set of 3 groups, Group I who were not athletes and did not perform any exercise, group II non athletes but went through exercise regiments and Group III, who were athletes and exercised.

#### MATERIALS AND METHODS

#### Selection of subjects:

Normal healthy adults, n = 60, aged 23-30 years, either gender was selected as per research criteria (Azizi, 2001) and classified in three groups as follows,

Group I = Non athletes, with no extensive exercise (n = 20) [control group]

Group II = Non Athletes, with extensive exercise (n = 20)

Group III = Athletes, with extensive exercise (n = 20)

Complete medical history was evaluated before inclusion of these subjects for research.

#### **Research Protocols:**

Group I was subjected to routine walk for 20 minutes, where as Group II and III were subjected to extensive aerobic exercise (Azeemi *et al*, 2020) for 20 minutes. Blood was collected from radial arteries 30 minutes prior to exercise or walk and 15 minutes after exercise in resting condition. Blood was analyzed for pH, pCO<sub>2</sub>, pO<sub>2</sub>, HCO<sub>3</sub> using Nova Phox Pro Arterial blood gas analyzer (Nova Biomedical, Massachusetts, USA).

#### Statistical analysis:

Data is presented as pre and post exercise Mean  $\pm$ Standard error of mean (SE) and compared statistically Pre and Post exercise using Students't-test for paired samples having P value < 0.05 as significant

## RESULTS

Group I show mild, moderate and no significant difference when pre and post ABGs parameters were compared with each other viz pO2 (mmHg) < 0.0019 [moderate], pCO2 (mmHg) P < 0.0418 [mild], HCO<sub>3</sub> (mM/L) P < 0.0009 and pH P < 0.2648 [NS], thus depicting non respiratory/ventilatory distress (Table 1). However, both group II (Table 2) and III (Table 3) exhibited one moderate and remaining very markedly significant difference amongst arterial blood gases (ABGs) parameters, Pre and Post extensive exercises (Table 2 and 3), therefore illustrating respiratory/ Ventilatory distress, acidosis and dawdling compensatory mechanism. Comparison of Blood Gasses changes after extensive exercise training in group II (Table 2), who were non athletes, significantly marked changes in pO<sub>2</sub> (mmHg) P < 0.00001, pCO<sub>2</sub> (mmHg) P < 0.00001, pH P < 0.00001 and moderate in HCO<sub>3</sub> (mM/L) P < 0.0017 were noted pre and post exercise. Similar findings were observed in Group III.

Table 1. Comparison of Blood Gases changes in group I with no extensive exercise Non athlete.							
Data presented as Pre and post Mean ± SE							
Parameters	Pre exercise	Post-exercise	P < 0.05	t-value			
pO <sub>2</sub> (mmHg)	$88.90 \pm 0.73$	$89.75 \pm 0.43$	< 0.0019	-3.32484			
pCO <sub>2</sub> (mmHg)	44.25 ± 1.99	$45.10 \pm 1.25$	< 0.0418	-2.11201			
HCO <sub>3</sub> (mM/L)	$25.50 \pm 1.21$	$27.00 \pm 1.16$	< 0.0009	-4.3589			
pH	$7.36 \pm 0.001$	$7.35 \pm 0.002$	< 0.2648	1.13177			

Table 2. Comparison of Blood Gases changes after extensive exercise training in group II non athletes. Data presented as Pre and post Mean  $\pm$  SE

Parameters	Pre exercise	Post-exercise	P < 0.05	t- value
pO2 (mmHg)	$88.35 \pm 0.77$	81.85 ± 1.93	< 0.00001	17.7253
pCO2 (mmHg)	44.15 ±3.5	$46.90 \pm 2.62$	< 0.00001	-4.9698
HCO3 (mM/L)	$25.35 \pm 1.50$	$26.6 \pm 0.99$	< 0.0017	-3.5411
pH	$7.38 \pm 0.02$	$7.26 \pm 0.03$	< 0.00001	5.97121

Table 3. Comparison of Blood Gases changes after extensive exercise training in group III athletes							
Data presented as Pre and Post Mean ± SE							
Parameters	Pre exercise	Post-exercise	P < 0.05	t- value			
pO2 (mmHg)	$88.75 \pm 1.04$	$98.15 \pm 0.56$	< 0.00001	-33.2888			
pCO2 (mmHg)	$44.85 \pm 1.50$	$58.45 \pm 1.35$	< 0.00001	-36.2454			
HCO3 (mM/L)	$25.75 \pm 2.09$	$27.05 \pm 0.58$	< 0.0019	-3.5590			
pН	$7.32 \pm 0.02$	$7.22 \pm 0.05$	< 0.00001	7.3697			
ND 0: 1 1				D 1 .			

N.B. Standard error of mean (SE) is used in dynamic studies where outcome is dependent on Pre-status compared with Poststatus in biological systems.

#### DISCUSSION

Our study showed elevated levels of  $pO_2$ ,  $pCO_2$  and  $HCO_3$  in post exercise state as compared to resting state before start of exercise, more markedly in athletes than non athletes and those who did not go for extensive exercise. Previously reported study also exhibited similar pattern where extensive aerobic exercise did influenced changes in arterial blood gases and pH (Azizi, 2011). It is well documented that apparent dyspnea that occurs during extensive exercise elevate demands for ventilatory accelerations, causing metabolic changes resulting in stress on dissolved gases oxygen (O<sub>2</sub>) and carbon dioxide (CO<sub>2</sub>) (Faisal *et al.*, 2015; Ofir *et al.* 2008, a,b; Phillips *et al.*, 2020). This strain due to reactionary ventilatory inefficiency on available metabolic stores cause onset of an early critical dynamic mechanical constraints and increase in respiratory neural drive (Phillips *et al.*, 2020). Both of these physiological alterations lead to sharp perceived dyspnea and resulting exercise intolerance, thus stirring modification to compensate respiratory chain reactions, resulting in high or low pCO<sub>2</sub>, pO<sub>2</sub>, pH and HCO<sub>3</sub>. Comparable interpretation can also be suggested for our findings as well where extensive exercise regiments caused marked changes in ABGs parameters, even in seasoned athletes.

It was argued that at peak of extensive exercise ventilation volume and carbon dioxide volume (VCO<sub>2</sub>) is often at much elevated level as compared to resting state because individuals were characteristically hyperventilating which was preceding excessive metabolic acidosis (Neder et al., 2017; Phillips et a., 2019). However, it was shown that individual who didn't possess the capability or potential to exercise above anaerobic threshold exhibited mild to moderate changes in dissolved gases, arterial blood gases (ABGs) or no change at all (Nader et al., 2015; Phillips et al., 2020), as we have witnessed in our case of non athletes with no exercise. Recent assessments contended that ability or potential to perform vigorous/extensive exercise depends on cardio-pulmonary system and ability of body physiology to deliver oxygen to exercising tissues is essentially required to eliminate metabolic by-products (Umapathi and Nyuagn, 2020). This phenomenon is coupled with movement of  $O_2$  and  $CO_2$  in and out of the human body to compensate any change sequentially. Interestingly, anticoagulant therapy in cardio-blood-pulmonary diseases does improve ventilatory efficiency to increase exercise capacity (Huang et al., 2020). Furthermore, plasma volume expansion was also considered as another therapy or option to increase and improve O<sub>2</sub> pressure in athletes (Zavorsky et al., 2003) to tackle exercise induced arterial hypoxemia. Conclusively, inducing acute hypervolaemia does improved blood gases status, it was shown, in athletes experiencing exercise induced arterial hypoxemia, and consequently arterial blood gases (ABGs) changes occurred for short period of time and not impaired further (Zavosreky et al., 2003).

## CONCLUSION

Research study presented here provide data about changes in pH, pO2, pCO2 and related parameters in set of 3 selected groups and compared pre and post exercise. Group I who were not athletes and did not perform any exercise exhibited no considerable difference in parameters whereas, group II who were non athletes but went through exercise regiments and Group III, who were athletes and exercised showed moderate to marked significant difference in pre and post exercise arterial blood gases parameters thus suggesting activation of compensatory, retaliatory and corrective cardio-pulmonary and metabolic mechanisms.

#### REFERENCES

Azizi, M. (2011). The comparison of aerobic training on blood gases in athlete and non-athlete high school girls. *Procedia Social and Behavioral Sciences*, 15: 1556–1560.

Arena, R., J. Myers., S.S. Aslam, E.B. Varughese, and M.A. Peberdy (2003). Peak VO<sub>2</sub> and VE/VCO<sub>2</sub> slope in patients with heart failure: a prognostic comparison. *Am. Heart J.*, 147: 354–360. doi: 10.1016/j.ahj.07.014

- Azeemi, H.I., S.S. Asghar, B. Ansari, J.M.Alam, and S.R. Mahmood (2020). Effects of exercise in fasting and postprandial conditions with metabolic profiling of fats and carbohydrates. *Int. J. Biol. Biotech.*, 17 (3): 583-587
- Brown, P.I., G.R. Sharpe., M.A. Johnson (2008). Inspiratory muscle training reduces blood lactate concentration during volitional hyperpnoea. *Eur. J. Appl. Physiol.*, 104(1): 111-7.
- Burton, D.A., K. Stokes and G.M. Hall (2004). Physiological effects of exercise. Continuing Education in Anaesthesia. *Critical Care & Pain*, 4 (6): 185-188.
- Faisal, A., K.A. Webb, J. A. Guenette, D. Jensen, J.A. Neder and D.E. O'Donnell (2015). Effect of age-related ventilatory inefficiency on respiratory sensation during exercise. *Respir. Physiol. Neurobiol.*, 205: 129–139. doi: 10.1016/j.resp.2014.10.017
- Guazzi, M., G. Reina, G. Tumminello and M.D. Guazzi (2005). Exercise ventilation inefficiency and cardiovascular mortality in heart failure: the critical independent prognostic value of the arterial CO<sub>2</sub> partial pressure. *Eur. Heart J.*, 26: 472–480. doi: 10.1093/eurheartj/ehi0
- Huang, D., J. Goa, W. Yang and J. Liu (2020) Exercise Capacity and Ventilatory Efficiency in Patients with Pulmonary Embolism After Short Duration of Anticoagulation Therapy. *Am. J. Med. Sci.*, 359(3): 140-146.
- Martin, L. (2008). The four most important equations in clinical practice. Global RxPh, Clinician's Ultimate Reference-COPYRIGHT -1999 PriorityDigital.com Prepared for: GlobalRPH Inc.
- Litch, P.M. (2006). Good breathing /, Bad breathing. Physiology and pathophysiology textbook/exercise, sports, and doping. Pharmpedia.
- Neder, J. A., F.E. Arbex, M. C. Alencar, C.D. O'donnell, J. Cory and K.A. Webb (2015). Exercise ventilatory inefficiency in mild to end-stage COPD. *Eur. Respir. J.*, 45: 377–387. doi: 10.1183/09031936.00135514
- Neder, J. A., D.C. Berton, F. F. Arbex, M. C. Alencar, A. Rocha and P.A. Sperandio (2017). Physiological and clinical relevance of exercise ventilatory efficiency in COPD. *Eur. Respir. J.*, 49: 1602036. doi: 10.1183/ 13993003.02036-2016
- Ofir, D., P. Laveneziana, K.A. Webb, Y. M. Lam and D.E. O'Donnell (2008a). Mechanisms of dyspnea during cycle exercise in symptomatic patients with GOLD stage I chronic obstructive pulmonary disease. Am. J. Respir. Crit. Care Med., 177: 622–629. doi: 10.1164/rccm.200707-1064OC
- Ofir, D., P. Laveneziana, K.A. Webb, Y. M. Lam and D.E. O'Donnell (2008b). Sex differences in the perceived intensity of breathlessness during exercise with advancing age. J. Appl. Physiol., 104: 1583–1593. doi: 10.1152/ japplphysiol.00079.2008
- Phillips, D. B., S.E. Collins, T. L. Bryan, E. Y. L. Wong, M. S. Mcmurtry, M. Bhutani and M.K. Stickland (2019). The effect of carotid chemoreceptor inhibition on exercise tolerance in chronic obstructive pulmonary disease: a randomized-controlled crossover trial. *Respir. Med.*, 160: 105815. doi: 10.1016/j.rmed.2019.10581
- Phillips, D.B., S.C. Collins and M.K. Stickland (2020). Measurement and Interpretation of Exercise Ventilatory Efficiency. *Front. Physiol.*, 11: 659. doi: 10.3389/fphys.2020.00659
- Powers, S,K. and E.T. Howley (2004). Exercise Physiology, 5th Edn. McGraw-Hill
- Umapathi, K.K. and H. Nguyen (2020) Cardio-pulmonary fitness. In: StatPearls [Internet]. Treasure Island (FL): StatPearls Publishing LLC. last update Jul 21, 2020.
- Zavorsky, G.S., K.R. Walley, G.S. Hunte, D.C. McKenzie, G.P. Sexsmith and J.A. Russell (2003) Acute hypervolaemia improves arterial oxygen pressure in athletes with exercise-induced hypoxaemia. *Exp. Physiol.*, 88 (4): 555-564.

(Accepted for publication June 2021)