INTRODUCTION

Pakistan has been ranked among highly polluted countries of the world causing drastic effects on health, posing economic burden on the country.\(^1,2\) World air quality index 2019, reported that Lahore is the second and Karachi is the seventh most polluted cities of the world, recorded particulate matter (PM) concentration in Lahore and Karachi were 188 \(\mu g/m^3\), 153 \(\mu g/m^3\) respectively. Approximately seven million early deaths have been reported around due to air pollution. Smog
is the worst form of pollution comprised of fog, smoke of burning crops and traffic generated pollutants; every year Lahore is submerged with smog with first winter spell.2,3 Exposure to traffic pollutants such as PM$_{2.5}$, PM$_{10}$ are associated with rise in mortality rate due to lung cancer and cardiovascular diseases.4 Study reported that prolong exposure to PM$_{10}$, nitrogen dioxide (NO$_2$), sulphur dioxide (SO$_2$), carbon monoxide (CO) is related with dyslipidemia, cardiac diseases (ischemic heart diseases, myocardial infarction) and diabetes mellitus.5 PM is one of the causative agents of neurological diseases by potentiating the oxidative damage in neural vasculature.6 Maternal Contact to PM$_{2.5}$, CO, SO$_2$ and oxides of nitrogen NO$_x$ is associated with preterm deliveries and low birth weight of infants and congenital abnormalities.7,8 Moreover prolong exposure to PM$_{10}$ and NO$_2$ is a causative factor of deaths in middle aged women due to cardiopulmonary diseases.9 It is reported by study that exposure of diesel exhaust potentiate the inflammatory process in human body which results in chronic diseases like asthma, cardiopulmonary disorders and cancers specially in susceptible population (children and elderly).10,11 Exposure of diesel exhaust fumes in infancy causes changes in gene expression of TNF, IL-10 and IL-13.12 As the traffic fume pollution is a global health concern, drastically affecting developing countries including Pakistan. however limited data is available till date to look into the pattern of markers of inflammation in apparently healthy population. Thus, this study was designed to evaluate the pattern of inflammatory markers in apparently healthy drivers who were exposed to traffic fumes. To the best of our knowledge, this is preliminary study in Pakistan that is addressing the effects of traffic exhaust fumes on human health.

**METHODS**

This cross-sectional study conducted from June 2016 to January 2017 at Liaquat University of Medical & Health Sciences (LUMHS), Jamshoro. It looked into the effects of traffic pollutants on markers of inflammation of apparently healthy automobile vehicle drivers. Markers of inflammation including: C-reactive protein (CRP), Leukocytes count, interleukin-6 (IL-6), tumor necrotic factor-α (TNF-α) and tumor necrotic factor-β (TNF-β), total leukocyte and differential leukocytes count (neutrophils, lymphocytes, eosinophils, monocytes and basophils). For this study eighty-seven, apparently healthy, non-smoking automobile vehicle drivers, having daily contact of traffic exhaust for at least six hours, aged between 18-40 years were recruited. Non-smoking, apparently healthy, not suffering from any systemic disease or autoimmune diseases volunteers were included in this study. While volunteers suffering from any systemic disorders, autoimmune diseases, overweight person (BMI ≥30 kg/m$^2$) were not considered.

**Methods of Inflammatory markers analysis:** 3 ml of whole blood was drawn from all the volunteers for analysis of inflammatory markers. CRP analyzed by “C-reactive protein Hitachi 902 turbidometry” and total leukocytes count by “Automated Analyzer” (sysmex). While IL-6, TNF-α and TNF-β were analyzed by “human instant ELISA (enzyme linked immunosorbent assay) (KOMA Biotech)® kits for analysis of markers. The standard laboratory method was followed, briefly 200 ul of washing solution added to each well. Wells were aspirated and excessive liquid removed 100 ul of standard (sample) then incubated at room temperature for two hours. Well aspirated and washed. 100 ul of diluted detection antibody (0.4 ug/ml for TNF-α, 0.1ug/ml for TNF-β and for IL-6 0.25ug/ml) to each well covered with the plate sealer then incubated for two hours then diluted Color Development Enzyme (1:20 dilute) each well was added. Incubated at room temperature until the appropriate color development at least for 17-27 minutes, plate read at 450 nm wavelength.

**Exposure analysis:** Measurement of pollutant (P.M$_{2.5}$, P.M$_{10}$ and NOx) carried out at seventeen busy areas of Hyderabad (Sindh), where traffic flow is usually heavy. P.M Meter (Model Aerocet: 531) used for the analysis of PM$_{2.5}$ and P.M$_{10}$ and NO$_x$ Meter (Model: AC32M) for NO$_x$. Level of pollutants compared with National Environmental Air Quality Standards (NEAQS). Concentration of each pollutant recorded at each location on three consecutive days of six months. Short duration exposure cut off value was five years ($\leq$ 5), while uppermost limit was twenty-five years ($\geq$25).

**Statistical analysis:** Data analyzed by using IBM statistical program for social sciences (SPSS), (IBM Corp. Released 2012. IBM SPSS Statistics for Windows, Version 23.0. Armonk, NY: IBM Corp. version) Spearman rank correlation used to determine the correlation of traffic pollutant with markers of inflammation. The results of all analyses evaluated for statistical significance using p-value < 0.05 and the 95 % confidence intervals (CI).
Ethical consideration: The project was approved by the Ethical Committee of Liaquat University of Medical and Health Sciences (Ref No. LUMHS/REC/94, dated Oct. 8, 2013) under the title: Detrimental Effects of Air Pollution on systemic health, Lung Volumes and Capacities of Young, Healthy Pakistani Volunteers.

RESULTS

During study period, 87 apparently healthy subjects included in the study. Mean age of subjects was 31(±7.3) years. While mean height and mean weight were, 174cm (±4.8) and, 75 kg (±6.3) respectively. Mean Body mass index (BMI) was, 24.3 (±2.4). Mean concentration of PM$_{2.5}$ (39.38), PM$_{10}$ (254.18) and NOx (37.26) recorded at different busy locations of city where traffic flow is quite high, compared with standard NEQS, as shown in Table-I.

Table-I: Mean concentration of pollutants of busy areas of Hyderabad City.

| Pollutants          | PM$_{2.5}$ ug/m$^3$ | PM$_{10}$ ug/m$^3$ | NOx ug/m$^3$ |
|---------------------|---------------------|---------------------|--------------|
| NEAQS Levels        | 35                  | 150                 | 80           |
| Jamshoro level crossing | 38.16             | 240.7               | 32.56        |
| Qasim Intersection  | 37.8                | 363.4               | 27.8         |
| Qasimabad           | 40.16               | 263.6               | 40.5         |
| Civil Hospital      | 43.4                | 234.1               | 35.6         |
| Hyder Intersecion  | 42.3                | 257.3               | 28.5         |
| Chandni market      | 44.2                | 233.5               | 41.23        |
| Giddu intersection  | 38.96               | 299.50              | 40.23        |

Mean levels of markers of inflammation are mentioned in Table-II.

Table-II: Mean levels of markers of inflammation.

| Markers of Inflammation | Mean |
|-------------------------|------|
| TNF-α ng/ml             | 7.7  |
| IL-6 pg/ml              | 60   |
| TNF-β pg/ml             | 29   |
| CRP mg/dl               | 0.66 |
| Leukocytes              | 9.1  |
| Neutrophils %           | 54   |
| Eosinophils %           | 5.0  |
| Basophils %             | 2.1  |
| Lymphocytes %           | 28   |
| Monocytes %             | 6.1  |

Table-III: Spearmen Correlation in markers of inflammation and PM$_{2.5}$

| Markers of Inflammation | PM$_{2.5}$ ug/m$^3$ |
|-------------------------|---------------------|
| r  | p-value          |
|-------------------------|---------------------|
| TNF-α pg/ml             | 0.41                | 0.001               |
| IL-6 pg/ml              | 0.99                | 0.001               |
| TNF-β pg/ml             | 0.47                | 0.66                |
| CRP mg/dl               | 0.99                | 0.001               |
| Leukocytes              | 0.03                | 0.73                |
| Neutrophils             | 0.29                | 0.06                |
| Eosinophils             | 0.20                | 0.006               |
| Basophils               | 0.10                | 0.14                |
| Lymphocytes             | 0.31                | 0.003               |
| Monocytes               | 0.42                | 0.001               |

$r = 0.31$, p-value = 0.003, eosinophils $(rs = 0.20, p-value = 0.06)$, monocytes $(rs = 0.42 p-value = 0.001)$ and basophils $(rs = 0.16, p-value = 0.14)$. While, there were no correlations present with total Leukocytes count $(rs = 0.03, p-value = 0.73)$, and TNF-β $(rs = 0.47 p-value = 0.66)$ as shown in, Table-III.

There was a positive correlation present among IL-6 $(rs = 0.21, p = 0.04)$, TNF-α $(rs = 0.49, p = 0.001)$, CRP mg/dl $(rs = 0.22, p = 0.03)$, Leukocytes $(rs = 0.14, p-value = 0.17)$ neutrophils % $(rs = 0.31, p-value = 0.003)$, lymphocytes % $(rs = 0.21, p-value = 0.042)$, monocytes % $(rs = 0.50, p-value = 0.001)$, basophils % $(rs = 0.17, p-value = 0.11)$ with PM$_{2.5}$. Whereas there was no correlation present with TNF-β $(rs = 0.01, p-value = 0.66)$ and eosinophils % $(rs = 0.09, p-value = 0.37)$, as shown in, Table-IV.

Table-IV: Spearmen Correlation among markers of inflammation and PM$_{10}$

| Markers of Inflammation | PM$_{10}$ ug/m$^3$ |
|-------------------------|---------------------|
| r  | p-value          |
|-------------------------|---------------------|
| TNF-α pg/ml             | 0.49                | 0.001               |
| IL-6 pg/ml              | 0.21                | 0.04                |
| TNF-β pg/ml             | 0.01                | 0.66                |
| CRP mg/dl               | 0.22                | 0.03                |
| Leukocytes              | 0.14                | 0.17                |
| Neutrophils             | 0.31                | 0.003               |
| Eosinophils             | 0.06                | 0.37                |
| Basophils               | 0.17                | 0.11                |
| Lymphocytes             | 0.21                | 0.042               |
| Monocytes               | 0.50                | 0.001               |
According to findings of our study, most of markers of inflammation showed positive correlation with pollutants. Our study results have elaborated the Th1/Th2 derived pro inflammatory cytokines (TNF-α, TNF-β and IL-6), since the study population was healthy subjects, increase in serum concentration at subclinical level is a frightening condition given that constant rise of inflammatory markers can lead to systemic diseases including autoimmune disease and even cancer. PM is highly hazardous for human health it is a combination of sulfate, sodium chloride, ammonia, mineral dust and black carbon. Due to its micro size it get lodge deep into lungs, along with NOx it causes irritation of respiratory mucosa and initiate local inflammation (TNF-α, IL-6), diffusion of markers of inflammation transported into circulation and mediate systemic inflammatory cascade. Effects of traffic pollutants on respiration already discussed in a separate chapter of thesis. In our study CRP showed positive significant correlation with pollutants PM2.5 and PM10, these findings are consistent with the results of Pilz (2018) study, in which PM2.5, PM10, NO2 and NOx showed positive association with CRP on long term exposure.13 Similar results by Lee (2018), reported that short term exposure of pollutant like SO2, NO2 and CO causes increase in fibrinogen level in non-smokers while, long term exposure causes rise in fibrinogen white blood cell (WBC) concentration.14 Study also revealed that short term exposure NO2 of non-smoking, healthy subject showed weak association with of markers of inflammation (IL-6, CRP, TNF-α) while strong positive association found on long term exposure among non-smokers and physically healthy subjects.15 In our study monocytes showed significant positive correlation with PM, NOx and CO while increase in monocytes concentration at subclinical level after exposure to traffic related NO2 and PM as reported by another study.16 Another study reported that both coarse PM10 and fine PM2.5 enhanced monocytes antigen presenting capacity.17 PM10 exposure to monocytes triggers increase intracellular calcium.18 Exposure of monocytes to nano particle of black carbon causes release of pro inflammatory cytokines (TNF-α, IL-6, IL-8) and increase in phagocytic capability of monocytes.19

Table-V: Spearman rank Correlation among markers of inflammation and NOx.

| Markers of Inflammation | NOx ug/m3 |
|-------------------------|-----------|
|                         | rs        |
| TNF-α pg/ml             | 0.48      |
| IL-6 pg/ml              | 0.22      |
| TNF-β pg/ml             | 0.02      |
| CRP mg/dl               | 0.22      |
| Leukocytes              | 0.14      |
| Neutrophils%            | 0.31      |
| Eosinophils %           | 0.10      |
| Basophils%              | 0.17      |
| Lymphocytes             | 0.13      |
| Monocytes               | 0.48      |

(rs = 0.31, p = 0.003), lymphocytes (rs = 0.13, p = 0.22), eosinophils (rs = 0.10, p = 0.34), basophils (rs = 0.17, p = 0.10), lymphocytes and monocytes (rs = 0.48, p = 0.001). While no correlation was found with TNF-β as shown in, Table-V.

**DISCUSSION**

According to findings of our study, most of markers of inflammation showed positive correlation with pollutants. Our study results have elaborated the Th1/Th2 derived pro inflammatory cytokines (TNF-α, TNF-β and IL-6), since the study population was healthy subjects, increase in serum concentration at subclinical level is a frightening condition given that constant rise of inflammatory markers can lead to systemic diseases including autoimmune disease and even cancer. PM is highly hazardous for human health it is a combination of sulfate, sodium chloride, ammonia, mineral dust and black carbon. Due to its micro size it get lodge deep into lungs, along with NOx it causes irritation of respiratory mucosa and initiate local inflammation (TNF-α, IL-6), diffusion of markers of inflammation transported into circulation and mediate systemic inflammatory cascade. Effects of traffic pollutants on respiration already discussed in a separate chapter of thesis.

In our study CRP showed positive significant correlation with pollutants PM2.5 and PM10, these findings are consistent with the results of Pilz (2018) study, in which PM2.5, PM10, NOx and NOx showed positive association with CRP on long term exposure.13 Similar results by Lee (2018), reported that short term exposure of pollutant like SO2, NO2 and CO causes increase in fibrinogen level in non-smokers while, long term exposure causes rise in fibrinogen white blood cell (WBC) concentration.14 Study also revealed that short term exposure NO2 of non-smoking, healthy subject showed weak association with of markers of inflammation (IL-6, CRP, TNF-α) while strong positive association found on long term exposure among non-smokers and physically healthy subjects.15 In our study monocytes showed significant positive correlation with PM, NOx and CO while increase in monocytes concentration at subclinical level after exposure to traffic related NO2 and PM as reported by another study.16 Another study reported that both coarse PM10 and fine PM2.5 enhanced monocytes antigen presenting capacity.17 PM10 exposure to monocytes triggers increase intracellular calcium.18 Exposure of monocytes to nano particle of black carbon causes release of pro inflammatory cytokines (TNF-α, IL-6, IL-8) and increase in phagocytic capability of monocytes.19

Furthermore, diabetogenic biomarkers (adiponectin, interleukin-1 and CRP) found to be elevated in healthy non diabetic persons on short and long duration exposure of PM2.5, PM10 and NO2.20 In addition, one more study documented that PM2.5 exposure causes increase in serum concentration of CRP and neutrophils in non-smoking healthy subjects.21 Likewise a Nepali study revealed positive association of CRP with PM2.5 while, negative association reported with TNF-α and IL-6.22 On the contrary a Belgian study reported no effects on total leukocytes concentration of healthy subject on exposure of traffic fumes and traffic related benzene causes decrease in WBC count, lymphocytes, eosinophils and platelets.23,24 Yet another study reported that when healthy female subjects exposed to pollutants for short duration CRP and leukocytes did not show positive association.25 A German study also stated prolong residential exposure of pollutant (PM2.5), showed weak association with CRP and no any association on short duration exposure in non-smoking volunteers.26 In the same way a Californian study reported association of TNF-α, IL-6 and CRP with NOx and CO.27 Association of TNF-α with NOx documented by another study.28 In our CRP, IL-6 and TNF-α showed positive correlation with PM10 on the other hand Tsai (2019) conducted study in Switzerland exposed the general population to PM10 at low concentration; reported positive association of similar marker of inflammation while no significant association with CRP.29
Inflammatory markers in healthy automobile vehicle drivers

Limitations of the Study: It included unavailability of continuous air quality data, small sample size and cross-sectional study design. However, prospective nature of the study is a major strength besides having assessed environmental pollution at different areas of the study in order to confirm the exposure of the pollutant.

CONCLUSION

Findings of our study suggest that almost all markers of inflammation are positively correlated with traffic generated pollutants, this condition might raise the risk of systemic diseases and causing deterioration health status of apparently healthy subjects.

Recommendation: Pakistan has been ranked top most polluted countries of the world; at present country is facing distressing air quality index. Quite a few studies have reported air quality status of major cities of Pakistan but still lacking consistent data of country. There is critical need of such research projects on broad scale and large sample size. Joint projects with collaboration of environmental protection department should be conducted on urgent basis.

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Conflicts of interest: None.

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Author’s Contribution:

BMS, ZL and SP are supervisors of HR.

HR conceived, designed, data collection and did statistical analysis & editing of manuscript, will be responsible and accountable for the accuracy or integrity of the work.

BMS supervised project, did review and final approval, editing of manuscript.

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