

1997

Effects of ambient air quality on children's lung function in urban and rural Iran

Maryam Mandana Asgari
Yale University

Follow this and additional works at: <http://elischolar.library.yale.edu/ymtdl>

Recommended Citation

Asgari, Maryam Mandana, "Effects of ambient air quality on children's lung function in urban and rural Iran" (1997). *Yale Medicine Thesis Digital Library*. 2355.
<http://elischolar.library.yale.edu/ymtdl/2355>

This Open Access Thesis is brought to you for free and open access by the School of Medicine at EliScholar – A Digital Platform for Scholarly Publishing at Yale. It has been accepted for inclusion in Yale Medicine Thesis Digital Library by an authorized administrator of EliScholar – A Digital Platform for Scholarly Publishing at Yale. For more information, please contact elischolar@yale.edu.

MED
T113
+.Y12
6448



EFFECTS OF AMBIENT AIR QUALITY
ON CHILDREN'S LUNG FUNCTION
IN URBAN AND RURAL IRAN

Maryam Mandana Asgari

Yale University

1997

YALE
UNIVERSITY



CUSHING/WHITNEY
MEDICAL LIBRARY

Permission to photocopy or microfilm processing of this thesis for the purpose of individual scholarly consultation or reference is hereby granted by the author. This permission is not to be interpreted as affecting publication of this work or otherwise placing it in the public domain, and the author reserves all rights of ownership guaranteed under common law protection of unpublished manuscripts.

Maryam M. Asyari


Signature of Author

3/10/97

Date

YALE MEDICAL LIBRARY

AUG 04 1997



Digitized by the Internet Archive
in 2017 with funding from
The National Endowment for the Humanities and the Arcadia Fund

<https://archive.org/details/effectsofambient00asga>

EFFECTS OF AMBIENT AIR QUALITY ON
CHILDREN'S LUNG FUNCTION
IN URBAN AND RURAL IRAN

A Thesis Submitted to the
Yale University School of Medicine
in Partial Fulfillment of the Requirements for the
Degree of Doctor of Medicine

by

Maryam Mandana Asgari

1997

Med Lib.

T 113

+ Y 12

6448

ABSTRACT

EFFECTS OF AMBIENT AIR QUALITY ON CHILDREN'S LUNG FUNCTION IN URBAN AND RURAL IRAN.

Maryam M. Asgari. (Sponsored by William S. Beckett). Yale University School of Medicine, New Haven, CT.

A cross-sectional epidemiological study comparing the pulmonary function of children in Tehran with those in a rural town in Iran was conducted in the summer of 1994. Four hundred children ages 5-11 were studied. Daytime ambient NO_2 , SO_2 , and PM_{10} were measured by portable devices in the children's neighborhoods on the days they were studied. Levels of these ambient substances were markedly higher in urban Tehran. Children's parents were questioned about home environmental exposures (including heating source and environmental tobacco smoke) and children's respiratory symptoms. Pulmonary function was assessed both by spirometry and peak expiratory flow meter. FEV_1 and FVC as a percent of predicted for age, sex and height were significantly lower for urban children compared with rural children. Both measures showed significant associations with levels of SO_2 , NO_2 and PM_{10} . Differences in spirometric lung function were not explained by nutritional status as assessed by height and weight for age, nor by home environmental exposures. Reported airway symptoms (cough, phlegm, and wheeze) were higher among rural children, while reported physician diagnosis of bronchitis and asthma were higher among urban children. This association of higher pollutant concentration with lower pulmonary function in this urban rural comparison suggests an effect of urban air pollution on short-term lung function and/or lung growth and development during the pre-adolescent years.

ACKNOWLEDGMENTS

This work was supported by the Wilbur Downs Fellowship and the NHLBI/DLD Preventative Pulmonary Academic Award. Invaluable support was given by the staff at the John B. Pierce Laboratory including Dr. Brian Leaderer for his recommendations and loaning of air quality measuring devices, Cindy Toth for assistance in preparing the air quality measuring devices, Dr. Vahid Mohsenin, Dr. Ethan Nadel and Fred Strumpf. We appreciated the collaborative efforts of Jim Sullivan at Harvard University for analyzing the Palmes tubes, as well as the University of Tehran School of Public Health, and the local public health professionals at Neeknejad and Lavasan. The Iranian government was especially helpful in allowing access to subjects as well as to air pollution data in the Department of Environmental Health.

All of the gratitude that I could express would not suffice to thank the efforts of my thesis advisor, Dr. William Beckett and my project advisor, Dr. Arthur DuBois without whose guidance, dedication and vision this project would not be possible. A special and heartfelt thanks to my father who encouraged, and at times, even pushed me. He pulled every string he could reach to make this project come to fruition. Any task at hand, from rescuing the spirometer at Mehrabad airport to securing the cooperation of the Iranian department of Environmental Health was performed with patience, tact, and zeal. Lastly, many thanks to my mother for loving, sister Haleh for advising, brothers Farid and Saied for supporting, Marc and Dr. Carrie Redlich for editing and all YMS friends for commiserating.

TABLE OF CONTENT

INTRODUCTION	Page 1
FACTORS INFLUENCING TEHRAN'S AIR QUALITY	Page 1
Cleansing Sources: Geographic Location and Meteorologic	
Factors	Page 2
Pollutant Sources: Heavy Industry and Motor Vehicles ...	Page 3
AIR QUALITY: WHAT IS BEING MEASURED AND WHY	Page 6
Nitrogen Dioxide	Page 7
<i>Sources</i>	Page 7
<i>Kinetics and Toxicity</i>	Page 8
<i>Alterations in Pulmonary Structure</i>	Page 9
<i>Alterations in Pulmonary Function</i>	Page 10
<i>Effects on Respiratory Symptoms</i>	Page 11
Sulfur Dioxide	Page 13
<i>Sources</i>	Page 13
<i>Kinetics and Toxicity</i>	Page 14
<i>Alterations in Pulmonary Structure</i>	Page 14
<i>Alterations in Pulmonary Function</i>	Page 15
<i>Effects on Respiratory Symptoms</i>	Page 15
Particulate Matter	Page 16
<i>Sources</i>	Page 16
<i>Kinetics and Toxicity</i>	Page 16
<i>Alterations in Pulmonary Structure</i>	Page 18
<i>Alterations in Pulmonary Function</i>	Page 18
<i>Effects on Respiratory Symptoms</i>	Page 19
AIR QUALITY STUDIES IN TEHRAN	Page 21
STATEMENT OF PURPOSE	Page 21
METHODS	Page 22
STUDY SITES	Page 22
STUDY POPULATION	Page 23
RESPIRATORY HEALTH QUESTIONNAIRE	Page 24
ANTHROPOMETRIC MEASUREMENTS	Page 24
PULMONARY FUNCTION MEASUREMENTS	Page 25
AIR POLLUTION MEASUREMENTS	Page 26
METEOROLOGY	Page 27
RESULTS	Page 28
AIR QUALITY	Page 28
LUNG FUNCTIONS	Page 30
DISCUSSION	Page 34
TABLES, FIGURES, AND APPENDIX	Page 42
REFERENCES	Page 59

INTRODUCTION

As the population of developing nations continues to increase at an alarming rate, so does their consumption of energy. The majority of the energy for industry, domestic use and transport is provided by fossil fuel combustion. Combustion of fossil fuels releases a variety of pollutants into the atmosphere, including sulfur oxides (SO_2 and SO_3), nitrous oxides (NO_2 and NO_3), ozone (O_3), carbon monoxide (CO) and particulate matter (PM). These pollutants can accumulate to reach toxic levels in cities within developing nations with rapid rates of industrialization and population growth. For Tehran, one of the world's megacities with the eighth highest population¹, air quality has been especially problematic. Within the past decade, the levels of major air pollutants have exceeded three to four fold the internationally adopted World Health Organization standards.²

FACTORS INFLUENCING TEHRAN'S AIR QUALITY

Levels of air pollutants are determined by a balance between the types and quantities of pollutants produced by various sources and the amount cleared by winds and precipitation. As Iran's capital city, Tehran is the financial, commercial and industrial center of the country. The principal sources of air pollution, namely heavy industry and motor vehicle traffic, are concentrated within the city's boundaries. Within the past twenty years, the city's

population has grown at a tremendous rate leading to further increases in industrial and motor vehicle activity. Tehran's geographic location, however, is such that dispersion of pollutants via strong winds and precipitation occurs infrequently. This imbalance of production versus clearance of air pollutants has led to Tehran's current poor air quality.

Cleansing Sources: Geographic Location and Meteorologic Factors

Tehran is located 1,200 meters above sea level in northern Iran on a plain south of the Alborz mountains (Figure 1). With an average annual precipitation between 200 to 1400 mm/year, Tehran's climate is arid. The "rainy season" encompasses three months starting in late fall and ending in early spring. The precipitation, being brief in both duration and quantity, is insufficient to cleanse the air of pollutant gases and precipitate suspended particles which are emitted on a year-round basis.

Another climatic variable that influences concentration of pollutants is wind-speed and direction. The Alborz mountains north of Tehran set up wind patterns dominated by winds that travel from mountain to plain and plain to mountain. These winds can be categorized into two types based on their speed: strong (> 17 knots) and weak (< 17 knots). The strong winds, which are capable of blowing the polluted air over and beyond the city boundaries, occur very rarely and comprise approximately 5% of the total winds.³ The weak

winds merely function to sweep the polluted air back and forth between the north and south end of the city. The lack of strong winds prevents dispersion and dilution of air pollutants.

Not only is horizontal circulation of air limited by lack of strong winds, but frequent temperature inversions prevent vertical air circulation. Under normal conditions, pollutants emitted from combustion processes are hotter than ambient air. Upon contact with the cooler ambient air, the gaseous pollutants rise. During temperature inversions, however, masses of cold air get trapped underneath warm air. Within the cold air mass, temperature paradoxically increases with altitude as the cold air rises to meet the warm air mass above it. This so called "inversion layer" traps pollutants close to the emission source. Thus, air pollution tends to accumulate waiting to be dispersed by an alternate airflow pattern or by strong winds.

The combination of frequent inversions, weak winds, and scarce precipitation which result from Tehran's unique geographical location thus contribute to the poor air quality within the city.

Pollutant Sources: Heavy Industry and Motor Vehicles

The livelihood of most urban centers depends upon the production and consumption of energy. Energy is required for industrial processes, motorized transport, as well as domestic activities such as heating and cooking. In

developing nations, fossil fuels provide the principal source of energy. Iran, having access to one of the world's richest oil reserves in the Persian Gulf, relies on crude oil as a source of energy. As the most industrialized urban center in Iran, Tehran is responsible for over 50% of the country's total oil consumption. The city harbors thirty seven percent of the nation's industrial activity including oil refineries, cement plants, sulfuric acid plants, plaster works and foundries.⁴ These industries lack stringent emission standards.

Industrial processes are not, however, the main culprit of the air pollution problem in most urban centers. That distinction goes to motor vehicles. According to Zerbonia et al.,⁴ motor vehicles account for between 60 and 90 percent of most air pollutants in Tehran. A recent report documented that 7.5 million liters of gas are consumed on a daily basis by automobiles within the city. The automobiles disperse a combined annual total of 3,000 tons of CO, 120 tons of nitrogen dioxide, 30 tons of sulfur dioxide, and 30 tons of particulates into the air.⁵ The amount of pollution is determined by the sheer number of vehicles on the road at any given time (estimated at about 2 million)², but also with the year and make of the vehicle. Old vehicles tend to be less fuel efficient and have poorer emission controls. They also tend to be more poorly maintained. In Tehran, however, automobiles are a near necessity due to the lack of an effective means of public transportation within the city.

Deterioration of the air quality in Tehran over the past few decades is primarily due to increasing industrial productivity and motor vehicle traffic in response to a several-fold rise in population. Iran has one of the highest population growth rates of any country in the world. From 1980 to 1990, the city's population increased by 40% from approximately 5.55 million to 9.21 million. The rate of growth is anticipated to continue, resulting in an estimated population of 13.78 million by the year 2000.¹ The growing urban population has led to increasing energy consumption, and hence, more emissions.

The resulting population boom has also led to unrestricted urban sprawl. Since 1956, Tehran, which used to encompass an area of approximately 80 km², has expanded to cover over 600 km². The average distance traversed to get from one point to another within the city has increased. On any given day, it is estimated that 10,000,000 trips are made, each lasting 45 minutes and covering a distance of 60 km.² Such heavy motor vehicle usage results in more emissions.

The fast-paced urban spread did not allow for implementation of strict planning on housing or industry to address overcrowding. Poor urban planning meant that new residential communities could spring up in close proximity to stationary sources of air pollution. It also led to a dearth of

protected areas such as parks and open green spaces. Thus more people were at risk for exposure to the increasingly polluted air.

In summary, rapid expansion, urbanization and overpopulation, combined with the unique features of the geographic location of Tehran have led to the high air pollution levels seen today. The major sources of pollutants within the city are motor vehicles with contributions from heavy industry.

AIR QUALITY: WHAT IS BEING MEASURED AND WHY

Recognition of an association between ambient air quality and respiratory health dates back to the 14th century when coal was first explored as a fuel source.⁶ However, it wasn't until several centuries later that the adverse health effects of air pollution were brought to public attention. Between the 1930s to the 1950s, several air pollution catastrophes brought about increases in mortality among inhabitants of heavily industrialized areas. Among the most notable were epidemics in the Meuse Valley of Belgium in 1930, Donora, Pennsylvania in 1948 and London in 1952. These epidemics, the worst of which occurred in London and resulted in an excess mortality greater than 4,000 people, focused international attention of the scientific and legal communities on air quality.⁷

In response to growing concern about the adverse effects of air pollution, the World Health Organization set out in 1972 to establish international

guidelines outlining the amount and the duration of exposure to various air pollutants at which no adverse health effects would be expected to occur (WHO, 1976). The resulting WHO guidelines, which were last revised in 1987, are now the most widely used international guidelines.

In the United States, the guideline of choice is the National Ambient Air Quality Standards (NAAQS) established by the Environmental Protection Agency (EPA). The NAAQS provide criteria for six pollutants identified by the 1990 Amendment of the Clean Air Act including nitrogen dioxide, sulfur dioxide, respirable particulate matter, carbon monoxide, ozone and lead. Assessment of air quality involves the comparison of levels of air pollutants against these established guidelines.

The health effects of inhaling air pollutants have been extensively studied, both experimentally and epidemiologically. In experimental studies, many of these pollutants have been correlated with adverse changes in pulmonary function. Three of these pollutants, nitrogen dioxide, sulfur dioxide and respirable particulate matter, will be the focus of this study.

Nitrogen Dioxide

Sources

Nitrogen dioxide forms in the process of high temperature combustion, where oxygen (O_2) combines with nitrogen (N_2) to make nitrogen oxide (NO),

nitrogen dioxide (NO_2) and other nitrogen oxides (NO_x). NO_2 can react with aerosols to form nitrous and nitric acids or alternatively, react with oxygen and hydrocarbons in sunlight to form ozone and other photochemical oxidants. The primary source of nitrogen dioxide in urban air is motor vehicle emissions. Other contributors include fossil fuel burning industries as well as chemical and nitration industries.

The National Ambient Air Quality Standard (NAAQS) recommends an annual arithmetic mean of NO_2 less than 0.053 ppm as a guideline (Table 2). The concentration of NO_2 in most urban centers world-wide ranges from an annual mean of 0.01 to 0.06 ppm.⁸ The range reflects differences in fossil-fuel consumption. Within cities, ambient levels vary with traffic density. During rush hour traffic, there is a corresponding peak in NO_2 emissions. Heavily trafficked roads can average hourly concentrations of NO_2 as high as 0.5 ppm.⁸ These levels far exceed the WHO guideline of 0.21 ppm as the recommended one-hour maximum level (Table 2).

Unlike other ambient air pollutants, NO_2 can occur in high concentrations indoors as well as outdoors. In homes containing sources of NO_2 such as unvented gas appliances and oil stoves, the concentration of NO_2 indoors can often exceed the outdoor concentration.⁹ For example, during cooking, maximum hourly concentrations of indoor NO_2 can reach levels as high as 1.0

ppm.⁸

Kinetics and Toxicity

Due to its low water solubility, NO₂ has the capacity to penetrate into the lower airways, where more than 60% is deposited.¹⁰ The pattern of deposition is predominantly centri-acinar. During normal breathing, the nasopharynx prohibits a significant portion of NO₂ from reaching the lower airways.

However, during exercise when breathing is shifted through the mouth, a greater amount of NO₂ reaches the lung periphery. Bauer et al.¹¹ conducted a study where they noted a 17% increase in the deposition of NO₂ with exercise, suggesting that exposure during exercise may potentiate NO₂'s adverse health effects. NO₂ is thought to produce adverse pulmonary health effects through oxidative capacities.

Alterations in Pulmonary Structure

Animal studies have shown that pulmonary epithelium can undergo irreversible morphologic changes after prolonged NO₂ exposure including hypoplasia of terminal bronchi as well as shortening and decrease in number of cilia.¹² Type I pneumocytes and ciliated epithelium, which are most sensitive to NO₂ exposure, are often replaced by less sensitive Type II cells.

Cytoarchitectural changes can also be induced in these substitute cells after acute exposures to NO₂ levels above 0.5 ppm.¹³ Emphysema-like changes, such

as loss of alveolar structure and enlargement of air-spaces, were noted in the lungs of animals that were exposed to high levels of NO₂ (20 ppm for 30 days).¹⁴ The mechanism by which chronic high-dose NO₂ exposure leads to these structural changes is thought to be mediated by inactivation of lung protease inhibitors. In a study conducted on human subjects, bronchoalveolar lavage fluid demonstrated a 45% decrease in activity of the alpha-1 protease inhibitor after exposure to 3 or 4 ppm NO₂ for 3 hours.¹⁵ Deficiency in lung proteases has been linked with the occurrence of emphysema in humans.

Alterations in Pulmonary Function

Controlled human studies have failed to show significant changes in the pulmonary function of normal subjects in response to acute NO₂ exposures. Exposures of 2.0 to 4.0 ppm NO₂ for periods of time ranging from 75 minutes to three hours have not shown decrements in pulmonary function as measured by spirometry or flow resistance.¹⁶⁻¹⁷ Although pulmonary function does not appear to be directly affected, bronchial reactivity, as measured by bronchoconstricting agents, does increase with acute NO₂ exposures at levels as low as 1.5-2.0 ppm for 3 hours.¹⁷⁻¹⁸

Asthmatics may be more sensitive to NO₂. The data on asthmatic subjects is controversial. Bauer et al, 1986¹¹ studied 15 asthmatics who were exposed to 0.3 ppm NO₂ for 30 min (20 minutes of rest followed by 10 minutes of

exercise), after exercise, noted a significant reduction in forced expiratory volume in one second. Morrow and Utell were unable to confirm the decrement in lung function in a study of 20 asthmatics exposed to the same level of pollutant for 4 hours.¹⁹ No pulmonary function changes were found for asthmatics exposed to 0.15 to 0.6 ppm NO₂ for 75 minutes,²⁰ 0.2 ppm for 2 hours,²¹ or 0.3-0.4 ppm for 75 minutes.²² The inconsistency of these results may be attributable to differing experimental protocols such as exercising vs. non-exercising or subject selection, such as mild vs. severe asthmatics.

Effects on Respiratory Symptoms

Epidemiologic studies suggest that a positive correlation exists between respiratory symptoms and levels of NO₂ exposure. In the Harvard Six Cities study, respiratory symptoms including shortness of breath with wheeze, chronic wheeze, chronic cough, phlegm, and bronchitis increased in prevalence with increasing indoor NO₂ concentrations.²³ Melia et al. also noted an association between living room NO₂ levels and respiratory symptoms.²⁴ With regard to the effects of ambient NO₂ on respiratory symptoms, several studies in Chattanooga, Tennessee have revealed an association with respiratory illness, especially amongst children. Chattanooga is home to a munitions factory which emits NO₂ into surrounding areas. Residents who are in close proximity to the factory are exposed to higher levels of ambient NO₂ than

those who reside further away. However, both communities are exposed to similar levels of SO₂ and particulate matter. Comparison of the rates of respiratory illness amongst the two communities has shown that the rates of respiratory illness are higher among families, and bronchitis episodes higher among school children in the high-exposure area.²⁵⁻²⁶ Meta-analysis of the epidemiological studies shows that an long-term increase in exposure to NO₂ of 0.015 ppm is associated with an increase odds of respiratory illness of approximately 20% in children.¹⁰

The increase in respiratory symptoms may be due, in part, to increasing susceptibility to bacterial and viral lung infections. Animal studies suggest that exposure to NO₂ at levels approximately an order of magnitude above ambient levels impairs immune responses to some bacteria and viruses. The experimental protocol employed in the majority of these studies involved exposure to NO₂ followed by inhalation of a pathogenic organism. These studies detected increased mortality from a bacterial challenge following NO₂ exposure.²⁷ With respect to viruses, Henry et al.²⁸ showed that prolonged exposure to 1.0 ppm NO₂ in squirrel monkeys was associated with enhanced serologic evidence of influenza virus. In mice, intermittent exposure to 5 ppm for 6 days decreased by nearly 100-fold the amount of viral innoculum needed to induce infection.²⁹

Alveolar macrophages appear to be the target for NO_2 's immune effects. In a study of bronchoalveolar lavage fluid derived from human lungs after exposure to 2.0 ppm NO_2 for 4 hours, alveolar macrophages displayed impaired phagocytic activity and decreased superoxide anion production.³⁰ The macrophage's capacity to defend against pathogens in vitro also appears to be diminished with NO_2 exposure. After 3 hours of exposure to 0.60 ppm NO_2 , alveolar macrophages obtained by bronchoalveolar lavage from human subjects inactivated influenza virus less effectively than macrophages obtained after exposure to air.³¹ These studies suggest that NO_2 has a detrimental effect on the ability of alveolar macrophages to perform their immune function.

Sulfur Dioxide

Sources

Sulfur dioxide (SO_2) is a clear, highly soluble gas that originates from the burning of sulfur containing compounds such as coal and crude oil. Sulfur dioxide may be further oxidized in the atmosphere to sulfur trioxide which can be converted to sulfuric acid in the presence of moisture. SO_2 can also react with metals and other pollutants to produce metallic acids and ammonium sulfates.

Major industrial sources of SO_2 include electrical power plants, oil refineries and smelters. The concentration of SO_2 in urban centers world-wide

ranges from an annual mean of 0.035 to 0.070 ppm,⁸ reflecting differences in fossil-fuel consumption. Peak hourly concentrations of SO₂ can rise as high as 0.35 to 0.70 ppm. Indoor concentration of SO₂ tends to be low as a result of absorption of the gas by walls, furniture and fabric.

Kinetics and Toxicity

Due to its high water solubility, SO₂ is effectively absorbed by mucous membranes lining the upper respiratory airways under resting conditions. With increased mouth ventilation, more SO₂ penetrates into the deeper portion of the lungs. SO₂ interacts with the mucus layer of the upper airways resulting in the production of hydrogen ions, bisulfate, and sulfate ions, all three of which have the potential for local irritation. These molecules are thought to cause bronchoconstriction by acting on sensory afferent nerve fibers in pulmonary smooth muscle.³²

Alterations in Pulmonary Structure

At high concentrations (>300 ppm), animal studies have shown that SO₂ can potentiate bronchitis-like pathology with sloughing of ciliated epithelium in the trachea and proximal airways and subsequent replacement by secretory cells.³³ Mice exposed for 72 hours to 10 ppm SO₂ had subsequent edema, necrosis and desquamation of their respiratory and olfactory epithelium in mice.³⁴ At ambient SO₂ levels, however, animal studies fail to indicate acute or

chronic toxicity.

In human subjects, ciliary abnormalities were discovered in the nasal epithelial cells after a two hour exposure to 0.75 ppm SO₂.³⁵ Altered mucociliary clearance is postulated to underlie the reduction in tracheobronchial clearance rate after exposure to acid sulfates. The suppressive action of the acid aerosol increases with rising acidity and appears to be cumulative.³⁶

Alterations in Pulmonary Function

Experimental studies on healthy humans have failed to demonstrate an acute effect of SO₂ on pulmonary function at levels which typically occur during air pollution episodes.³⁷⁻³⁹ Asthmatics, however, are sensitive to even low levels of SO₂. Exercising asthmatics can experience bronchoconstriction at concentrations of 0.25 ppm SO₂.⁴⁰⁻⁴¹ Asthmatic subjects exposed to 0.25 ppm SO₂ for brief periods lasting 5-10 minutes can develop significant increases in airway resistance, and decreases in FEV₁, especially with oral breathing.⁴² Although these changes in pulmonary function are rapid and remain stable with lengthening exposure, they are readily reversible and resolve within an hour after the end of exposure.

Effects on Respiratory Symptoms

In epidemiological studies, exposure to levels of SO₂ as low as 0.5 ppm has

been shown to increase mortality.⁸ One such study, which followed 5,500 women in western Pennsylvania, noted an increased frequency of respiratory symptoms associated with levels of ambient SO₂.⁴³ Euler et al.⁴⁴ studied Seventh-Day Adventists in California and found that their incidence of respiratory symptoms such as chronic cough/phlegm, diagnosis of asthma with wheeze or emphysema was associated with cumulative SO₂ and particulate exposures. The strongest association was found with particulates. SO₂ and particulates often occur together as components of a complex mixture of emissions from primary combustion. Increasing epidemiological evidence points to particulates as being the more toxic of the mix.

Particulate Matter

Sources

Particulate matter is produced by incomplete combustion of fossil fuels. Inefficient combustion of carbon containing fuels releases a mixture of solid and liquid particles that are collectively termed total suspended particles (TSP). Constituent particles vary in size and composition and include carbon, lead, vanadium, bromine, sulfur oxides, nitrogen oxides, silicon, aluminum, iron, sodium and chloride. The concentration and composition of particulate air pollution varies with the source, the dispersion by air motion, and the amount of time available for sedimentation. There is a wide range of

concentrations of particulate matter throughout urban centers of the world. In developing countries, the particulate levels can range as high as 100 times the current U.S. standards.⁷

Kinetics and Toxicity

The toxicity of particulates varies with the size and the physical composition of the mixture. Particulates can be classified on the basis of their size. Particles greater than 10 microns in diameter tend to be deposited above the epiglottis, adhering to nasal hairs and mucous membrane of the nose. Particles up to 10 microns in diameter (PM_{10}), otherwise referred to as respirable particles, tend to be deposited in the upper and lower airways. Particles less than 5 microns in diameter (PM_5) can be inhaled and can penetrate to the alveoli. Exercise increases oral breathing and therefore bypasses the nasal clearance mechanism. Consequently, the deposition pattern changes with more particulates being deposited in the lower airways.

Particle clearance depends on the site of deposition. Particles deposited in the anterior nasal cavity are cleared by sneezing or wiping whereas those in the posterior cavity are cleared to the posterior pharynx and swallowed. Particles in the trachea, bronchi and bronchioles are moved up the tracheobronchial tree by the sweeping action of the cilia to be expelled by coughing or swallowing. In the terminal bronchioles and alveoli, particles are cleared by lung

macrophages. Fine particles < 5 microns in diameter may pass through the alveolar tissue and move into the lymphatic circulation.

The chemical composition and kinetics of many of the components of particulate matter has not been fully elucidated. Toxicologic studies have focused primarily on the effects of single components of particulate matter or on simple mixtures. Yet many of the toxic effects of particulates are thought to derive from the combination of the complex mixtures which includes SO_x as well as acid aerosols. The lack of standardization of dosimetry for different particulates in complex mixtures makes for comparison of most toxicologic studies difficult. In preliminary studies, particulate complexes appear to be far more toxic to macrophage defense capabilities than classic particles such as asbestos and silica.⁴⁵

Alterations in Pulmonary Structure

In animal studies, edema and pathologic changes were noted to occur in animals within 24 hours of exposure to particulates from space heaters, burning wood, coal, and automotive waste oil.⁴⁶

Alterations in Pulmonary Function

Numerous studies have documented an association between particulate pollution and decreased pulmonary function both in healthy volunteers as well as those with preexisting respiratory conditions. In a study of 335 healthy

children, a decrease in $FEV_{0.75}$ and FVC was noted after an air pollution episode where the concentration of TSP reached a 24 average of $312 \mu\text{g}/\text{m}^3$.⁴⁷ Dassen et al.⁴⁸ reported a decrement in FEV_1 and FVC in children in Ijmond, Netherlands associated with TSP levels of 200-250 $\mu\text{g}/\text{m}^3$. Focusing on a population more prone to respiratory illness, Pope and Kanner⁴⁹ found that in over 6,000 cigarette smokers, changes in FEV_1 and FEV_1/FVC were inversely associated with changes in levels of PM_{10} . They noted that a 100 $\mu\text{g}/\text{m}^3$ increase in PM_{10} levels resulted in a 2% drop in FEV_1 .

Effects on Respiratory Symptoms

Epidemiologic studies indicate a link between particulate air pollution and increased respiratory morbidity and mortality. One study of patient records in Ontario indicated a marked association between admissions for acute respiratory disease and levels of SO_2 and floating particulate matter.⁵⁰ Pope demonstrated a statistical association between respiratory hospital admissions and particulate matter in Utah, Salt Lake, and Cache valleys.⁵¹ In a health interview survey of 50,000 households over a five year period, Ostro et al.⁵² found an association between fine particulate matter and respiratory conditions severe enough to result in work loss and disability in adults. Total number of emergency room visits were associated with ambient levels of TSP in Steubenville, Ohio.⁵³ In a multiple regression analysis, an overall increase of

100 $\mu\text{g}/\text{m}^3$ over the base-line 24 hour mean PM_{10} increased emergency room visits for respiratory disease by 3%.⁵⁴

Particulates have been singled out as the major cause of mortality in the 1952 epidemic in London.⁵⁵ More recently, Schwartz and Dockery⁵⁶ associated daily mortality counts for a seven year period in Philadelphia with daily concentration of particulates. In examining the mortality data, they found significant correlation with daily mortality counts of death and particulate and SO_2 concentrations for the current and preceding day. Controlling for season, temperature and humidity, total mortality increased by 7% per 100 $\mu\text{g}/\text{m}^3$ increase in total suspended particulates. Mortality data from Steubenville, Ohio also showed a significant association between daily total mortality and TSP concentration on the previous day.

In a review of the epidemiologic literature, Dockery and Pope⁵⁷ concluded that for each 10 $\mu\text{g}/\text{m}^3$ rise in levels of PM_{10} , there is a corresponding 1% increase in total mortality, a 1.4% increase in cardiovascular mortality and a 3.4% increase in respiratory mortality. With regard to morbidity, there was a corresponding rise of 1% in the incidence of all respiratory symptoms requiring medical attention with 2-3% increase in asthma alone. By demonstrating quantitatively similar adverse effects of acute PM_{10} exposure, these results suggested coherence across morbidity and mortality data.

AIR QUALITY STUDIES IN TEHRAN

In Tehran, the effects of air quality on respiratory morbidity were examined by Shamsi² who studied 10,720 children in 3 different areas of Tehran. He selected the relatively unpolluted area in northern Tehran (Area 1), and two highly polluted areas in City Center (Area 2) and southern Tehran (Area 3). He measured SO₂ and TSP in these areas and found annual mean concentration (in µg/m³) of SO₂=53.06 and TSP=12.5 in Area 1, SO₂=198.48 and TSP=442.28 in Area 2, and SO₂=227.04 and TSP=309.08 in Area 3. He administered a questionnaire examining respiratory symptoms and found that the percentages of children with asthma and bronchitis in Areas 1, 2, and 3 were 1.9, 5.6, and 4.4 respectively. TSP appeared to be a stronger factor in predicting respiratory symptoms among children living in Tehran.

STATEMENT OF PURPOSE

The goal of this project was to assess how levels of air pollutants, namely SO₂, NO₂ and PM₁₀, affect pulmonary function and respiratory symptoms of children in Tehran. Children were chosen as the subjects of the study because according to statistics of the Department of Public Health in Tehran, the rate of respiratory disease is high and the majority of the city's patients consist of children.² Children's lungs may be more vulnerable to the effects of air

pollution. To assess the degree to which lung function of the indigenous population of Tehran is altered as a result of exposure to high levels of air pollutants, the lung functions of children in Tehran were compared to those of an unpolluted nearby rural area.

Lung function measurements consisted of peak expiratory flow readings (PEF) and spirometric tracings of forced expiratory volume at one second (FEV_1) and forced vital capacity (FVC). PEF, FEV_1 and FVC for the two sites were compared and subsequently correlated with levels of air pollution as measured by portable air quality monitoring devices. To assess respiratory symptomatology as well as the contribution of possible confounding variables, the parents or guardians of each subject completed a detailed questionnaire at the time of testing regarding parameters such as indoor air quality, parental smoking patterns, socioeconomic status and nutrition.

METHODS

The study was conducted from June to August of 1994. The summer months were chosen to minimize the incidence of upper respiratory infections which could potentially confound the results of pulmonary function studies.

STUDY SITES

The study sites were selected to represent areas of high and low exposure to

SO₂, NO₂ and particulate matter. The highly polluted district of Neeknejad in the inner city of Tehran (henceforth designated “urban”) was chosen because of its proximity to mobile sources of air pollution such as heavily congested thoroughways and railroads as well as stationary sources such as a nearby cement factory. The district of Lavasan (“rural”), located approximately 30 km Northeast of Tehran, was selected because of its geographical location beyond the Alborz mountain range which served to block the polluted air from entering the area (Figure 2).

STUDY POPULATION

The study population consisted of two hundred children from each site between the ages of 5 and 11 years. There were 107 females and 93 males in the urban and 101 females and 99 males in the rural subgroups. In the urban site, an area of approximately 2 miles x 2 miles was chosen on the basis of its proximity to major roadways. All houses located on streets adjoining the roadways were considered eligible. Each street was canvased by knocking on doors, and those which were inhabited and which responded were invited to participate in the study. With the exception of two households, who refused to participate, all other households that were approached consented.

Children from rural areas included Lavasan and smaller surrounding villages within an approximately fifteen mile radius. In Lavasan, children were

recruited by the door-to-door method described above. In the smaller villages, where possible, organized gatherings of children were arranged by the Department of Public Health in nearby health centers. The response rate for participation was greater than 99% in both areas.

RESPIRATORY HEALTH QUESTIONNAIRE

In both regions, parents or guardians were approached about the study and told its purpose. Having received written consent, a respiratory health questionnaire was administered to parents or guardians of subjects. This questionnaire, a modified version of the Recommended Respiratory Disease Questionnaire for Use with Adults and Children in Epidemiological Research⁵⁸ translated into Farsi, was used to obtain information on the child's respiratory illness history and respiratory symptoms, as well as potential confounding variables that may have influenced respiratory health such as home cooking and heating sources, smoking patterns within household, and parental education (sample questionnaire included in appendix).

ANTHROPOMETRIC MEASUREMENTS

Following completion of the questionnaire, the subject's erect standing height, without shoes, was measured using measuring tape to the nearest 0.25 inches. Next, the child's weight, without shoes, was measured to the nearest pound using a portable scale.

PULMONARY FUNCTION MEASUREMENTS

Two tests of pulmonary function were performed. First, the child's peak expiratory flow rate was measured by asking the child to perform forcible maximum exhalation from total lung capacity into a Personal Best peak flow meter. The readings from this instrument have been shown to have a linear relationship to the expected standard for both ATS waveform 24 as well as the Fleisch pneumotachometer, and thus to fulfill the National Asthma Education Program's technical standards.⁵⁹ Each subject was fitted with a nose clip, instructed to stand up and inhale as deeply as possible, place their lips firmly around the mouthpiece and blow out as fast and as hard as he or she could. The subject was told that their goal was to get the red indicator on the peak flow meter to the highest possible point on the scale. The test was repeated three times. The highest value was defined as the child's PEF.

The second pulmonary function test used a portable Breon Spirometer (Model 2400) to obtain a tracing of their forced expiration at total lung capacity. Each subject was fitted with a nose clamp and was instructed to inhale as deeply as possible, seal their lips tightly around the mouthpiece, exhale as forcefully as possible and maintain expiration until the pen on the spirometer reached the end of the tracing. Each subject attempted a trial run using only the mouthpiece unattached to the tube to acquaint him or her with

the procedure and to make sure that he or she could perform the task correctly. Evidence of acceptable performance included an unhesitating start, apparent maximal effort with smooth continuous exhalation, absence of cough, glottis closure or mouthpiece obstruction by tongue or teeth. Once the child felt comfortable, the mouthpiece was attached to the spirometer, and the child was given at least three test trials, or as many as needed to obtain two consistent readings. Each child's spirometric tracings were analyzed for FEV₁ and FVC. The largest acceptable FEV₁ and FVC volumes were recorded, even if the two values did not originate from the same curve.

AIR POLLUTION MEASUREMENTS

Portable air monitoring devices to measure NO₂, SO₂, and PM₁₀ were used in both urban and rural areas. Palmes tubes were used to measure the amount of NO₂ and SO₂ in ambient air. For each day of pulmonary function testing, two Palmes tubes, one for NO₂ and one for SO₂, were placed outside the home environment of one of the subjects who was tested during that day. The tubes were exposed to ambient air by removing the protective cap and hanging them near subjects' home environments, either in their yard or outside their doors or windows. Devices were hung approximately six to nine feet above ground level to minimize tampering. Approximately two weeks later (number of exposure hours ranging from 285-438), the tubes were retrieved, capped, and stored in

an air-tight plastic bag at room temperature until analysis. Analysis of the Palmes tubes was performed by Jim Sullivan at Harvard University.

PM₁₀ was measured with a MINIRAM (Miniature Real-time Aerosol Monitor) Model PDM-3, a light scattering airborne particulate monitor whose operating principle is based on the detection of scattered electromagnetic radiation in the near infrared. The scattering sensing parameters were designed for preferential response to particles whose diameter ranges between 0.1 and 10 microns. During operation, a liquid-crystal display indicated the aerosol concentration in units of $\mu\text{g}/\text{m}^3$ which was updated every 10 seconds. PM₁₀ levels were measured in the mornings (and sometimes in the afternoons) on each day of testing. Obtaining afternoon PM₁₀ measurements was temperature dependent. On afternoons where the temperature exceeded 100 °F, the equipment had a high rate of failure. Daily PM₁₀ values were calculated by averaging the morning and afternoon levels as measured over a one to two minute period. If afternoon measurements were unobtainable, only the morning levels were used in calculating daily PM₁₀ levels. The error rate for a 1 minute averaging is approximated at $0.02 \mu\text{g}/\text{m}^3$. The monitor was placed in a plastic bag and calibrated with filtered air prior to each reading.

METEOROLOGY

Meteorological data on local temperature and barometric pressure were

collected for each day of testing using hand held portable monitoring devices. Temperature was measured with a mercury thermometer, placed in the shade, at the start and completion of every day of testing. The daily temperature was calculated as the average of the two readings. Barometric pressure was measured with an aneroid barometer at the start of each day of testing.

RESULTS

AIR QUALITY

A total of 17 NO₂ and 15 SO₂ Palmes tubes in the urban and 8 NO₂ and 6 SO₂ Palmes tubes in the rural areas were retrieved for analysis. The exposure time of each Palmes tube was factored into calculating the ppm concentration of air pollutants in the subjects' ambient environment. Five NO₂ and nine SO₂ monitors were lost to theft or tampering and were therefore not included in the analyses. Within the urban area, the mean concentration of NO₂ was 0.0247 ppm (46.44 µg/m³), SO₂ 0.0158 ppm (45.19 µg/m³), and PM₁₀ 99.99 µg/m³. In contrast, the mean concentrations for the rural site were NO₂ of 0.00985 ppm (17.39 µg/m³), SO₂ of 0.00651 ppm (18.62 µg/m³) and PM₁₀ of 0.01 µg/m³ (Figure 3). The differences in concentrations for all three pollutants were significant at the 0.001 level (Table 1).

To assess the validity of the air pollution data, pollutant levels were

compared to those gathered during the same time period by the Iranian Department of Environmental Health (IDEH) in Tehran. For the months of June-July of 1994, the IDEH reports a mean NO₂ concentration of 0.052 ppm, SO₂ of 0.038 ppm and TSP of 159 µg/m³ (no PM₁₀ levels were measured). The IDEH levels of NO₂ and SO₂ are higher than those recorded by our portable monitoring devices (PMDs). Several factors could explain this discrepancy. The central IDEH collection site is located on a major thoroughfare in a non-residential area in mid-town Tehran, approximately 6 km north of the urban site. The proximity of the central monitoring station to mobile sources of air pollutants would allow for detection of more concentrated levels of such pollutants. In contrast, the PMDs in this study were placed within the yards and surrounding vicinity of the children's homes, further away from the direct influence of mobile sources (as there were no residential areas situated on the major thorough ways). The lower values measured by the PMDs may be more representative of the actual exposure to pollutants in the ambient air. Alternatively, the IDEH levels may be more accurate given the small sample size of the PMDs or may be discrepant based on differences in instrumentation.

In comparing the IDEH values to the WHO guidelines⁸ and the National Ambient Air Quality Standards (NAAQS) for annual arithmetic means of these

air pollutants,⁶⁰ it is evident that the IDEH levels of SO₂ and TSP exceed both the WHO and NAAQS guidelines (Table 2). IDEH measured NO₂ levels border the NAAQS established standard of <0.053 ppm but fall well below the WHO guideline of <0.080 ppm. Although the PMD levels of SO₂ and NO₂ did not exceed the established guidelines, the PM₁₀ did exceed the WHO standard of <70 µg/m³ as well as that of the NAAQS of <50 µg/m³.

LUNG FUNCTIONS

Using the meteorologic data collected for each day, the measured lung volumes were corrected to BTPS standards. The following formula was used to calculate corrected lung volumes:

$$\text{Corrected lung volumes} = \text{Uncorrected lung volumes} \times (P_a \times T_b) / (P_b \times T_a)$$

where:

$$P_a = [\text{Barometric pressure (inches Hg)} \times 25.4 \text{ mm/inch}] - \text{water vapor pressure at ambient temperature}$$

$$P_b = [\text{Barometric pressure (inches Hg)} \times 25.4 \text{ mm/inch}] - \text{water vapor pressure at 37 degrees Celsius}$$

$$T_a = \text{Ambient temperature (degrees Celsius)} + 273.18$$

$$T_b = 37 + 273.18$$

The BTPS corrected FEV₁s and FVCs were then compared to the predicted values. Predicted values were calculated for each child based on the

child's age, sex and height using a formula derived from studies of healthy American children.⁶¹ Similarly, predicted values for PEF were calculated based on the child's height using established norms from studies of healthy British children.⁶²

The mean percent predicted FEV₁ of children in the urban area was 89.0% (SD=23.7%) whereas that of the rural area was 99.0% (SD=18.9%). Similarly, the mean percent predicted FVC in the urban area was 89.7% (SD=22.9%) whereas the rural percent predicted FVC was 106.3% (SD=17%). For both FEV₁ and FVC, this difference was significant at the $p < 0.001$ level as measured by Levine's Test. The mean percent predicted PEF of the urban area (82.4%, SD=13.5%) did not differ significantly from that of the rural area (81.8%, SD 16.9%--Table 3 and Figure 4).

To assess if other variables contributed to the lower lung function of the urban children, indoor air quality, socioeconomic status, and nutritional status were explored. With regard to indoor air quality, three potential sources of pollutants were examined: cooking sources, heating sources and tobacco smoke (Table 4). Cooking and heating sources were categorized based on their fume-generating capability. Cooking fuels such as fuel oil, kerosene and coal burners were categorized as fume-generating sources whereas central gas stoves were categorized as a non-fume generating source. Rural households used more

fume-generating cooking sources (17 rural vs. 2 urban, $p < 0.001$). A similar trend held true for heating sources. Rural households used more fume generating heating sources such as petroleum and charcoal furnaces and kerosene burners as compared to urban homes which relied more heavily on central gas (171 rural vs. 2 urban, $p < 0.001$). With regard to smoking patterns, more fathers smoked in the rural area ($p < 0.016$). Maternal smoking was approximately equal in incidence in both areas (4 in rural, 3 in urban, $p < 0.70$).

Socioeconomic status (SES) is considered a risk factor for respiratory disease.⁶³ The indices that were used to assess SES in this study were the size of the household, the number of rooms in the house, and the educational level of the parents. The number of rooms per household and the number of people in the household did not differ significantly between the rural and urban areas ($p < 0.20$ and $p < 0.40$ respectively). However, the mean number of years of education of the father (urban mean = 8.8 years, rural mean = 6.4 years) as well as the mother (urban mean = 6.9 years, rural mean = 5.2 years) were significantly greater in the urban area (Table 5).

The measures of nutritional status included children's height for age, weight for age, and body mass index (weight in grams/[height in cm]²). The two populations did not differ significantly with respect to age ($p < 0.12$),

weight ($p < 0.07$), or body mass index ($p < 0.53$). The urban children were slightly taller than rural children overall ($p < 0.05$). The urban/rural comparison of these anthropomorphic variables is illustrated in table 6. The similarity in nutritional status is illustrated in figures 5 and 6 which show an urban-rural comparison of age vs. weight and age vs. height for both urban and rural children.

The respiratory symptoms that were investigated in the questionnaire were coughing, phlegm and wheezing. In addition, the incidence of physician-diagnosed asthma, bronchitis, pneumonia and sinusitis were assessed. A summary of the frequency of reported respiratory symptoms is presented in table 7. Although the rural and urban children did not differ in their episodes of coughing apart from colds, the rural children had more episodes of cough with colds ($p < 0.025$). Rural children also had more phlegm with colds ($p < 0.009$), and phlegm apart from colds ($p < 0.042$) with cough or phlegm that lasted for more than one week ($p < 0.001$). Rural children had a greater number of chest colds per year when compared to urban children ($p < 0.028$) though chronic chest illnesses defined as those lasting 7 or more days did not differ in prevalence between the two groups. With regard to wheezing, rural children wheezed more while having a cold ($p < 0.001$), apart from having a cold ($p < 0.011$), had more episodes of wheezing leading to shortness of breath

($p < 0.051$) and wheezed more during exercise ($p < 0.014$). No children in the rural area were diagnosed with asthma, whereas three in the urban area carried the diagnosis ($p < 0.082$). More rural children had pneumonia ($p < 0.010$) and sinusitis ($p < 0.030$) whereas more urban children had episodes of bronchitis ($p < 0.008$). Thus, where differences in frequency of reported symptoms exist, rural children reported having more respiratory symptoms overall.

DISCUSSION

Tehran, the most industrialized urban center in Iran, harbors thirty-seven percent of the nation's industrial activity. Both light and heavy industries, such as oil refineries, cement plants, sulfuric acid plants, plaster works and foundries are contained within the city's boundaries.⁴ Tehran's rapid population growth has led to an increase in both mobile and stationary pollutant sources. These sources, lacking stringent emission standards, have resulted in poor ambient air quality within the city. The aim of this study was to determine how levels of these ambient air pollutants affected children's lung functions.

The results of this study indicate that spirometric lung function measurements of urban children were consistently lower than those of rural children. The average percent predicted FVC and FEV₁ in the urban area were

16.6% and 10.0% less than those of the rural area. This difference could not be explained by differences in anthropomorphic measurements of the children, in exposures to differing sources of indoor air quality such as cooking, heating, or paternal smoking, or by socio-economic factors. The decrements in lung function were, however, associated with differences in ambient levels of SO₂, NO₂, and PM₁₀ amongst urban and rural areas.

Ambient air quality was measured using portable monitoring devices (PMDs). The PMD-measured levels of NO₂ and SO₂ were slightly less than half of the values measured by the Iranian Department of Health's (IDEH) central monitoring station for the same time period in central Tehran. This difference in levels is most likely attributable to the location on the monitoring stations. The IDEH monitoring station was located in a heavily trafficked thorough way in a non-residential area whereas the PMDs were placed in back yards of subjects which were not in as close a proximity to direct exposure from mobile sources. With regard to NO₂, the monthly mean as measured by IDEH was 0.052 ppm, barely under the NAAQS established guideline of 0.053 ppm. For SO₂, the IDEH measurement of the monthly mean exceeded both the NAAQS and WHO guidelines by 20% and 60% respectively. The IDEH monthly mean of total suspended particles exceeded the NAAQS and WHO by two-fold, whereas the PMD measured value for PM₁₀ exceeded the NAAQS

by two fold and the WHO guideline by 43% (table 2).

These high levels of air pollutants are not unusual for Tehran. Data on TSP and SO₂ levels over the past decade, as measured by the IDEH, indicate that levels of these pollutants regularly exceeded the WHO and NAAQS guidelines by several fold (Table 8). The most significant air pollutant in Tehran that has consistently violated established air quality guidelines is suspended particulate matter. As outlined in the introduction, several studies have recently pointed to particulates as the main culprit for the morbidity and mortality associated with air pollution episodes,⁵⁰⁻⁵⁷ as well as decrements in pulmonary function associated with air pollution.⁴⁷⁻⁴⁹ Shamsi's study of children residing in Tehran² also implicated TSP levels as the strongest predictor of respiratory illness.

As table 8 shows, Tehran experiences the least amount of air pollution during the summer. This study was conducted during the relatively unpolluted time period of summer. Therefore, the documented decrements in lung function may be more reflective of cumulative exposures.

This study has several strengths. For one, it is, to date, the only published study of the effects of ambient air quality on children's lung function in Iran. Shamsi² studied three areas within the city of Tehran with differing levels of air pollutants and found a positive correlation between the

frequency of respiratory symptoms as reported by students in a respiratory questionnaire and the amount of TSP and SO₂ in the ambient air. However, Shamsi's study did not include any measure of pulmonary function. Despite these trends, few studies have examined the physiological effects of poor air quality on respiratory function in Tehran. Other strengths include significant air quality differences between the urban and rural sites, good exposure data, a well-selected control group, and an extensive search for possible confounding variables.

This study also has several weaknesses, one of which being that the mean percent predicted PEF did not differ significantly between the urban and rural groups ($p=0.472$). The similarity in PEFs may be due to the fact that the peak flow meter is a less precise instrument for measuring pulmonary function. Although for the purposes of this analysis, the highest value of three efforts was defined as the child's peak flow value, there was a wide degree of variability within subjects. In contrast, the lung volume measurements demonstrated far less within-subject variability. Another concern is that not all of the spirometers met the ATS criteria for reproducibility, indicating variable effort. Also, some of the total measured lung volumes were lower than expected for height and weight, thus indicating submaximal effort. However, because this was observed in both urban and rural spirometers with similar

frequency, there was no bias in urban-rural comparisons.

Another weakness in the data is that although rural children had better overall lung function, they reported having more respiratory symptoms. Several factors may explain this discrepancy. For one, rural children reported having a borderline significant higher incidence of allergies to pollen and dust (11 rural vs. 4 urban, $p = 0.065$). Therefore, the increase in respiratory symptoms among the rural population may be due to differential exposure to allergens that may provoke coughing, phlegm and wheezing but not necessarily lead to a decrement in pulmonary function. Lavasan is one of the prime fruit-producing areas of Iran. The fruit trees, which blossom on an annual basis, can serve as prime pollen producing sources. This is in sharp contrast to the urban area, where few green spaces exist.

Alternatively, the differing incidences of symptomatology between the rural and urban areas may reflect differences in indoor air quality. As stated earlier, rural children tended to live in homes with more fume generating heating and cooking sources. These sources may have emitted more NO_2 into the home, especially if they were unvented. As highlighted in the introduction, NO_2 has been shown in epidemiological and animal studies to compromise immune response leading to an increase in bacterial and viral infections. However, exposure to NO_2 has not been linked with decrements in

lung function. Therefore, high indoor exposures of NO_2 could explain the increase in respiratory symptoms in rural children, as it would render them more susceptible to pulmonary pathogens, without necessarily causing decreases in FEV_1 and FVC.

Another factor that could be postulated to explain this difference is varying incidences of upper respiratory tract infection. The rural area is situated above the Alborz mountain range, lending it to a colder climate. Also, the rural area has inferior quality of sanitation. Some of the smaller villages in the rural area lacked a clean water supply. The combination of colder climate and poor sanitation may have contributed to higher incidences of respiratory infection especially during the winter months. When parents were asked to report their child's symptoms, their answers reflected the child's symptoms throughout the duration of the year. Thus, an increased amount of respiratory symptoms may have reflected increased incidence of upper respiratory tract infection occurring during the winter months with little sequelae on lung function during the summer months.

The findings of lower FEV_1 and FVC in the more polluted urban environment are in agreement with the current literature. The association between decrements in pulmonary function and $\text{TSP}/\text{PM}_{10}$ levels suggested in this study has also been documented in other international studies. Dassen et

al.⁴⁸ noted declines in FEV₁ and FVC in children in Ijmond, Netherlands with TSP levels in the range of 200-250 µg/m³. He et al⁶⁴ studied the effects of pollution on children's pulmonary function in urban and suburban areas of Wuhan, China and noted that decrements in FEV₁ and FVC were associated with high annual mean levels of SO₂ (59 µg/m³), NO₂ (47 µg/m³) and TSP (251 µg/m³). They found that the average FVC and FEV₁ in children in the suburban area was 6.7 and 3.8 percent lower, respectively, as compared to the suburban area. Further analysis indicated that the suburban-urban difference in both FVC and FEV₁ increased with increasing height, suggesting a failure on the part of the urban children to maintain lung function growth. They suggested that exposures in childhood may slow lung maturation and growth with volume related indices, such as FVC being more severely effected than flow-related indices, such as FEV₁.

Similarity, Xu et al⁶⁵ in their study of adult pulmonary function in relation to TSP and SO₂ levels, also proposed that FVC may be better indicator for assessing the effects of long-term cumulative exposure to air pollutants. In this study, comparing the spirometric lung functions of urban children to rural children reveals a similar trend-- a larger decrement of FVC (16.6%) than FEV₁ (10.0%). This is consistent with Xu et al's, and He et al's hypothesis^{64,65} that changes in FVC may be more reflective of long-term,

cumulative exposures.

In summary, in an urban-rural comparison of lung function and symptoms in Tehran, higher pollutant levels in the urban area were associated with lower lung function among children ages 5-11. This difference was not explained by indoor air quality, environmental tobacco smoke, differences in socioeconomic status, or by nutritional status. We believe that further research is warranted since the pollutant levels and health effects measured are consistent with studies in other parts of the world ^{46, 64,66,67} and could indicate an important and potentially irreversible response to ambient pollutants.

TABLE 1: Urban-Rural comparison of ambient air quality as measured by portable monitoring devices for June-August, 1994

Pollutants	URBAN		RURAL		URBAN vs. RURAL
	Mean	SD	Mean	SD	p values
NO ₂ (ppm)	0.0247	0.0108	0.0099	0.0048	0.001
SO ₂ (ppm)	0.0158	0.0042	0.0065	0.0033	<0.001
PM ₁₀ (µg/m ³)	99.9	72.4	0.01	0.01	0.001

TABLE 2: Comparison of PMD* and IDEH† measurements of ambient air pollutant levels in Tehran for June-July of 1994 to NAAQS‡ and WHO¶ Guidelines for Annual Arithmetic Means

Pollutants	PMD	IDEH	NAAQS	WHO
	One Month Mean	One Month Mean	Annual Arithmetic Mean	Annual Arithmetic Mean
NO ₂ (ppm)	0.025	0.052	<0.053	<0.080 ^h
SO ₂ (ppm)	0.016	0.038	<0.030	<0.015-0.023
PM ₁₀ (µg/m ³) ^g	100	not measured	<50	<70
TSP (µg/m ³) ⁱ	not measured	159	<75	<60-90

Portable Monitoring Devices

Iranian Department of International Health

U.S. National Ambient Air Quality Standards

World Health Organization

This value is the mean 24 hour guideline for NO₂. WHO has not established a guideline for annual arithmetic mean for NO₂.

^g Particulate Matter less than or equal to 10 microns in diameter

ⁱ Total Suspended Particulates

TABLE 3: BTPS corrected pulmonary function tests as a percent of predicted for age, height and sex

Pulmonary Function Parameters	URBAN		RURAL		URBAN vs. RURAL
	Mean	SD	Mean	SD	p values
FEV ₁	89.0%	23.7%	99.0%	18.9%	<0.001
FVC	89.7%	22.9%	106.3%	17.3%	<0.001
PEF	82.4%	13.5%	81.8%	16.9%	0.472

TABLE 4: Urban-Rural Comparison of Home Environmental Exposure

Source	Urban	Rural	Urban vs. Rural p value
Fume generating cooking sources ¹ (Number of households)	2	17	<0.001
Fume generating heating sources ² (Number of households)	2	171	<0.001
Smoking (paternal)	72	94	0.015
Smoking (maternal)	3	4	0.698
Mean Number of cigarettes/day smoked in household	5.54	8.02	0.021

¹ Includes fuel oil, kerosene, and coal

² Includes petroleum furnace, charcoal furnace and kerosene burners.

TABLE 5: Urban-Rural Comparison of Socio-Economic Status (SES)

SES Measures	Urban	Rural	Urban vs. Rural p value
Rooms/household	2.74	2.66	0.467
People/household	5.60	5.76	0.173
Paternal education (yrs)	8.84	6.36	<0.001
Maternal education (yrs)	6.93	5.19	<0.001

TABLE 6: Urban vs. Rural Comparison of Anthropomorphic Data

Anthropometric Parameters	URBAN		RURAL		URBAN vs. RURAL
	Mean	SD	Mean	SD	p values
Age (years)	8.012	1.154	7.824	1.308	0.115
Height (inches)	48.874	3.359	48.217	3.288	0.049
Weight (lbs)	49.975	9.592	48.295	9.175	0.074
BMI (wt gm/[ht cm]²)	1.461	0.171	1.451	0.152	0.534

TABLE 7: Urban-Rural Comparison of Respiratory Symptoms

Symptom	Number of Children with Respiratory Symptoms		Urban vs. Rural p value
	Urban	Rural	
Cough with cold	175	188	0.025
Cough apart from cold	15	12	0.550
Phlegm with cold	82	108	0.009
Phlegm apart from cold	4	12	0.041
Cough & phlegm for 1+ wk	20	50	<0.001
Wheeze with cold	37	66	<0.001
Wheeze apart from cold	1	9	0.010
Wheeze causing SOB	9	19	0.050
Wheeze during exercise	5	16	0.014
Asthma	3	0	0.082
Bronchitis	7	0	0.009
Pneumonia	11	26	0.010
Sinusitis	3	11	0.030

TABLE 8: IDEH mean monthly levels of TSP and SO₂ averaged from 1980 - 1989

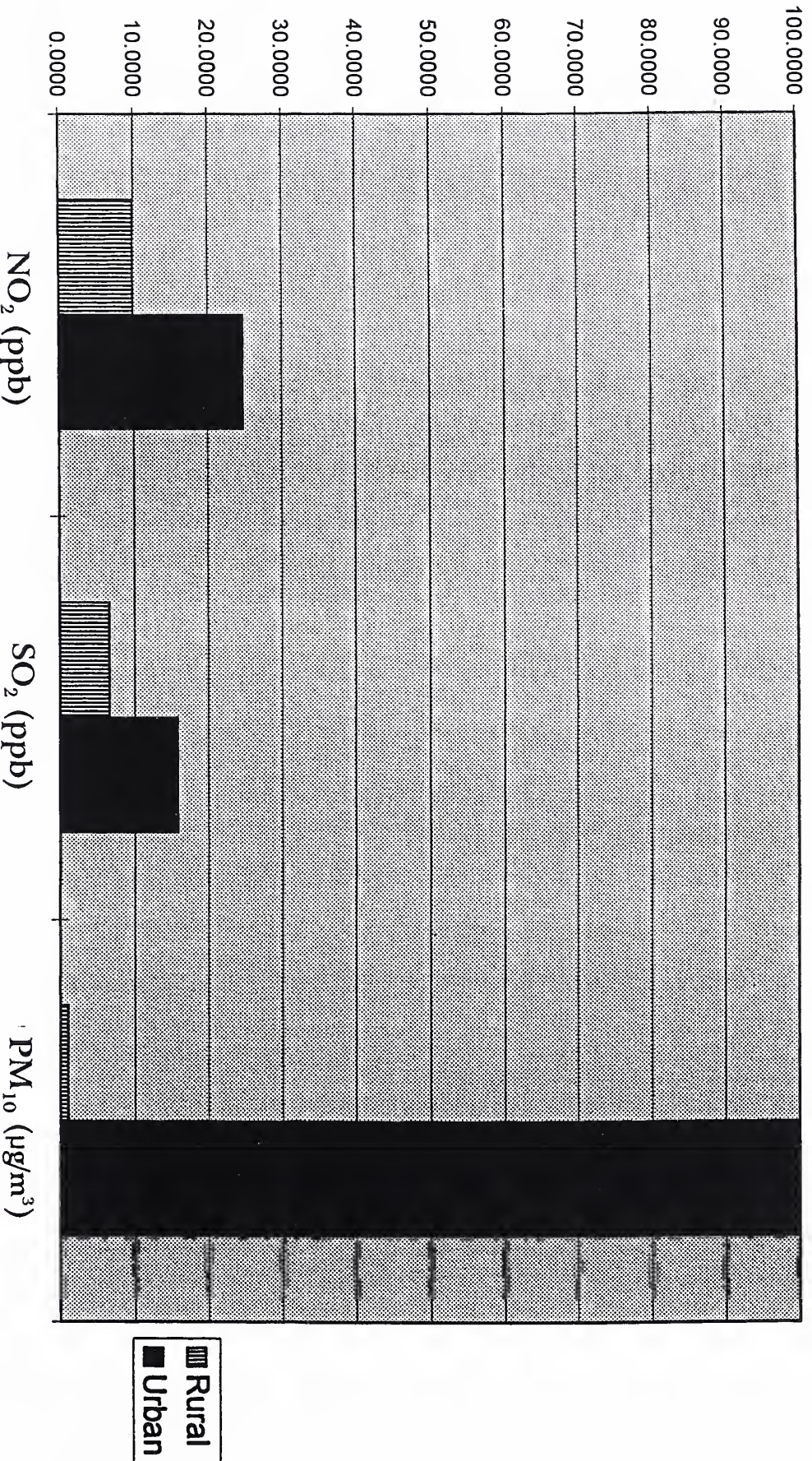
MONTH	TSP ($\mu\text{g}/\text{m}^3$)	SO ₂ (ppm)
January	294	0.179
February	302	0.165
March	262	0.106
April	242	0.075
May	242	0.077
June	246	0.077
July	271	0.088
August	280	0.080
September	293	0.094
October	304	0.081
November	291	0.106
December	274	0.159

FIGURE 1
Map of Iran



FIGURE 3

Urban-Rural Comparison of SO₂, NO₂ and PM₁₀ Levels as Measured by Portable Monitoring Devices (PMDs)



* : SO₂ and NO₂ levels are shown in ppb for ease of display. To convert values to ppm, multiply by 10⁻³.

FIGURE 4

Urban-Rural Comparison of Percent Predicted Pulmonary Function Tests

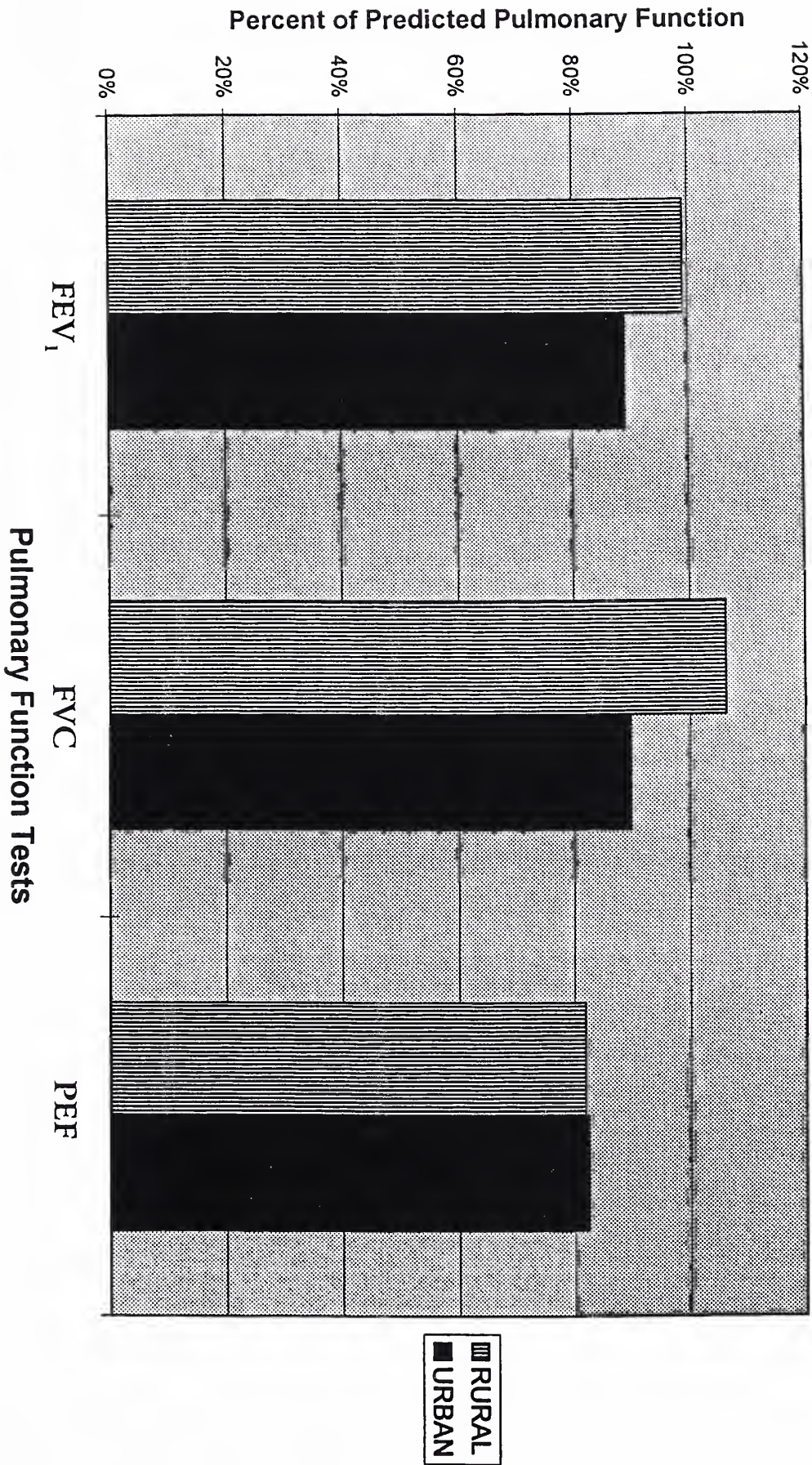


FIGURE 5

Age vs. Weight for Urban and Rural Subjects

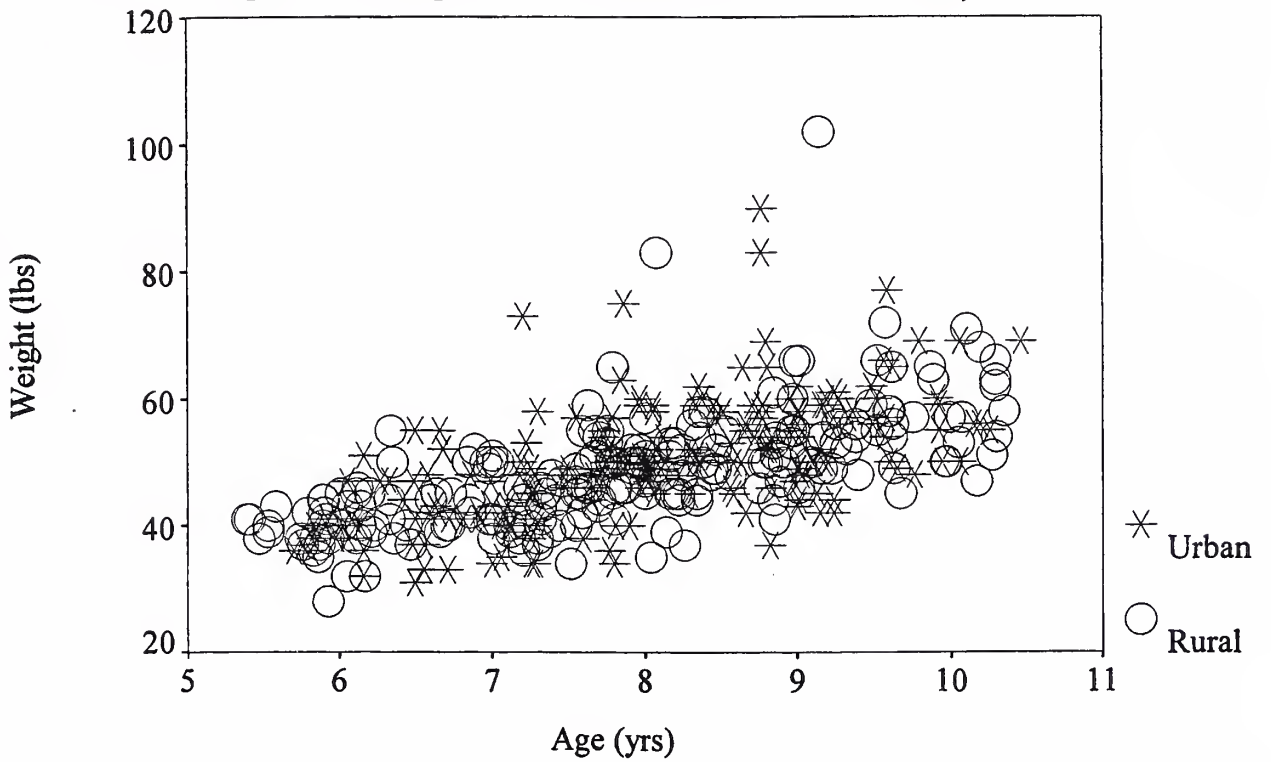
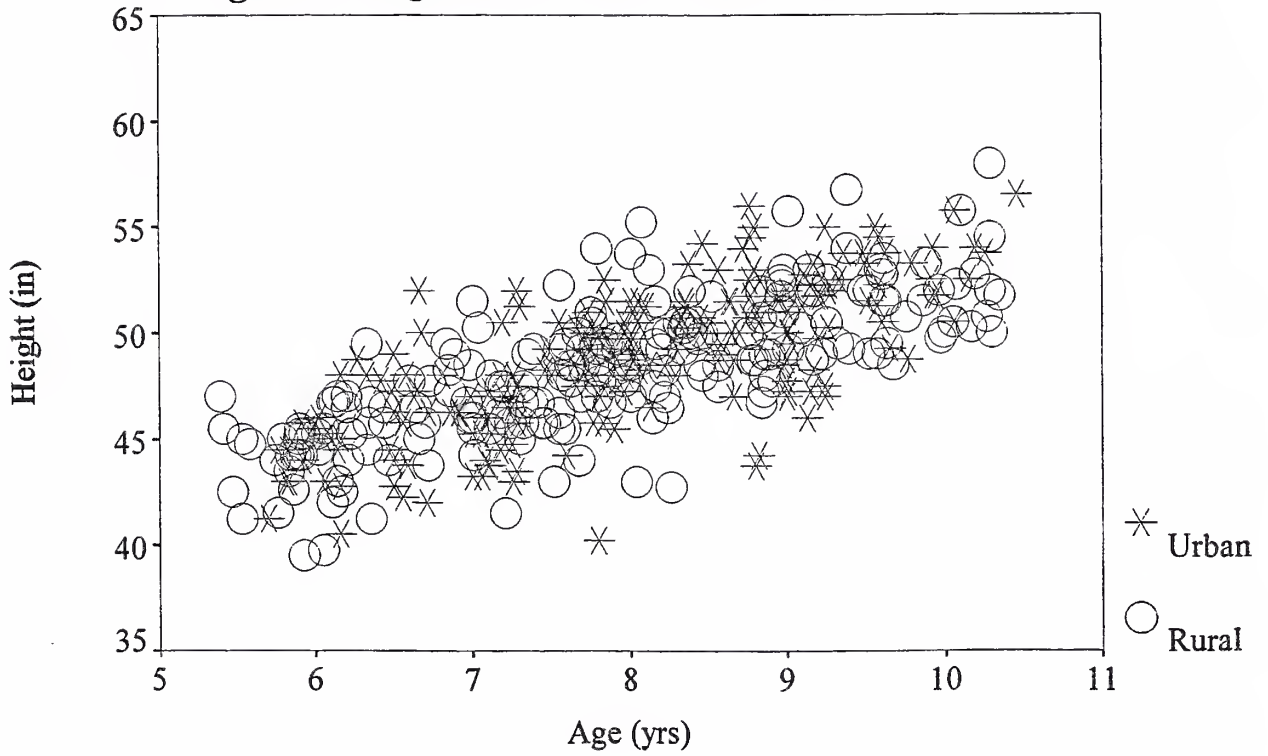


FIGURE 6

Age vs. Height for Urban and Rural Subjects



APPENDIX

Respiratory Health Questionnaire

1. ID _____ 2. Date _____ 3. Sex _____ 4. Location _____
5. Relation of person completing questionnaire to child _____
6. Date of child's birth: _____
7. Where was this child's mother living when this child was born? _____
8. Please list all places where s/he lived for 6 months or longer, from birth to present: _____
9. Does s/he attend school? _____ if so, what grade is s/he in? _____
10. What is the age of the youngest child living in this child's home? _____
11. A. How many people share his/her bedroom? _____
 B. Does this child have his/her own bed? _____
 C. Do any of the people that share his/her room smoke? _____
12. How many rooms are in your house (not incl. bathrooms & kitchens)? _____
13. How many people live in your home? _____
14. How is your home heated? _____
15. What fuel is used most for cooking in your home? _____
16. How is your home cooled? _____
17. A. Do you have any pets? _____ if so, what kind? _____
 B. Do the pets reside mostly indoor or outdoor? _____

COUGH

18. A. Does s/he usually have a cough with colds? _____
 B. Does s/he usually have a cough apart from colds? _____
 IF YES (to part B):
 C. Does s/he cough on most days (4+ days/week) for as much as 3 months/year? _____
 D. For how many years has s/he had this cough? _____

PHLEGM

19. A. Does s/he usually bring up phlegm with colds? _____
 B. Does s/he usually bring up phlegm apart from colds? _____
 IF YES (to part B):
 C. Does s/he bring up phlegm, sputum, or mucus from his/her chest on most days (4+ days/week) for as much as 3 months per year? _____
 D. For how many years has s/he raised phlegm, sputum, or mucus from his/her chest? _____
20. A. Does this child get attacks of cough, or phlegm lasting for 1 week or more each year? _____
 IF YES:
 B. For how many years? _____
 C. On average, how many chest colds per year does s/he get? _____

WHEEZING

21. A. Does this child's chest ever sound wheezy or whistling when s/he has a cold? _____
 B. Does this child's chest ever sound wheezy or whistling apart from colds? _____
 C. Does this child's chest sound wheezy or whistling most days or nights? _____
 IF YES (to parts B or C):
 D. For how many years _____
22. A. Has this child ever had an attack of wheezing that has caused him/her to be short of breath? _____
 IF YES:
 B. Has s/he had 2 or more such episodes? _____
 C. Has s/he ever required medicine or treatment for the(se) attack(s)? _____
 D. How old was this child when s/he had his/her first such attack? _____
 E. Is his/her breathing completely normal between attacks? _____

23. Does this child ever get attacks of wheezing after s/he has been playing hard or exercising? _____

CHEST ILLNESS

24. During the past 3 years, has this child had any chest illness that kept him/her from his/her usual activities for as much as 3 days? _____

IF YES:

A. Did s/he bring up more phlegm or seem more congested than usual with any of these illnesses? _____

B. How many illnesses like this has s/he had in the past 3 years? _____

D. How many of these illnesses have lasted for as long as 7 days? _____

25. Was s/he ever hospitalized for a severe chest illness/chest cold before the age of 2 years? _____

26. Did this child have any other chest illness or chest cold before the age of 2 years? _____

OTHER ILLNESSES

27. Has this child had any of the following diseases?

A. Measles _____ B. Sinus Trouble _____ C. Bronchitis _____

D. Croup _____ E. Whooping Cough _____ F. Pneumonia _____

28. A. Has this child had any ear infections between 0-2 years of age? _____

B. Has this child had any ear infections between 2-5 years of age? _____

C. Has this child had any ear infections over age 5? _____

29. Did this child require tubes to be placed in his/her ears to drain them? _____

30. Did this child ever have an operation on his/her tonsils? _____

31. A. Has a doctor ever said that this child had asthma? _____

IF YES:

B. At what age did the asthma begin? _____

C. Does s/he still have asthma? _____

IF YES (to part C):

D. Is s/he currently taking any medication? _____

IF NO (to part C):

E. When did it his/her asthma stop? _____

32. Has this child ever had an operation on his/her chest? _____ if so, specify _____

33. Does this child have heart disease? _____ If so, specify _____

34. When s/he was born, was s/he kept in the hospital after the mother went home? _____

if so, specify reason _____

ALLERGY

35. Does this child have an allergic reaction to food or medicine? _____

36. Does this child have an allergic reaction to pollen or dust? _____

37. Does this child have an allergic skin reaction to detergents or other chemicals? _____

38. Did this child ever receive allergy shots? _____

FAMILY HISTORY

MALE PARENT

39. Is he the natural father? _____ If no, specify the relationship: _____

40. What is the highest grade of school he completed? _____

41. What is his present job? _____

42. A. Does he smoke regularly (at least 1 cigarette per day)? _____

IF YES:

B. With what frequency? _____

43. A. Does he smoke any other type of tobacco, ie: cigars, pipe, etc.? _____

IF YES:

B. Specify type and frequency of usage _____

44. A. Has he ever smoked regularly while living at home with this child? _____

IF YES:

B. With what frequency? _____

45. Has a doctor ever said he had:
- A. Bronchitis _____ B. Emphysema _____ C. Asthma _____
- D. Any respiratory condition (specify) _____

FEMALE PARENT

46. Is she the natural mother? _____ If no, specify the relationship: _____

47. What is the highest grade of school she completed? _____

48. What is her present job? _____

49. A. Does she smoke regularly (at least 1 cigarette per day)? _____

IF YES:

B. With what frequency? _____

50. A. Does she smoke any other type of tobacco, ie: cigars, pipe, etc.? _____

IF YES:

B. Specify type and frequency of usage _____

51. A. Has she ever smoked regularly while living at home with this child? _____

IF YES:

B. With what frequency? _____

52. Has a doctor ever said she had:

A. Bronchitis _____ B. Emphysema _____ C. Asthma _____

D. Any respiratory condition (specify) _____

OTHER HOUSEHOLD MEMBERS

53. A. Are there any other members of the household who currently smoke regularly? _____

IF YES:

B. Specify relation _____

C. With what frequency? _____

REFERENCES

1. World Health Organization and United Nations Environmental Programme. 1992. Urban air pollution in megacities of the world. Blackwell Publishers, Oxford.
2. Shamsi, A. 1990 Air pollution and its relation with prevalence of respiratory diseases among primary school students of three educational areas of Tehran. MSPH Thesis. University of Tehran School of Public Health.
3. Azimi, A.A. 1979. Evaluation of SO₂ levels in two regions of the city of Tehran. MSPH Thesis. University of Tehran School of Public Health.
4. Zerbonia R., and Soraya, B. 1978. Air pollution control in Iran. *J. Air Pollut. Control Assoc.* 28: 34-337.
5. *Hamshahri* Newspaper. Published in Tehran, Iran. March 1996. 4(932). p. 3.
6. Ayres, S.M., Evans, R.G., and Buehler, M.E. Air pollution: A major public health problem. 1972. *CRC Critical Review of Laboratory Science.* 3: 1-40.
7. Chivian, E., McCally, M., Hu, H., and Haines, A. 1993. Critical condition, human health and the environment. MIT Press. Cambridge, MA, p. 13-29.
8. World Health Organization. 1987. Air Quality Guidelines for Europe. WHO Regional Publications, European series no. 23. World Health Organization, Copenhagen.
9. Leaderer, B. 1982. Air pollutant emissions form kerosene space heaters. *Science.* 218: 1113-1115.
10. Committee of the Environmental and Occupational Health Assembly of the American Thoracic Society. 1996. Health effects of outdoor air pollution. Part 2. *Am. J. Resp. Crit. Care Med.* 153: 477-498.

11. Bauer, M.A., Utell, M.J., Morrow, P.E., Speers, D.M. and Gibbs F.R. 1986. 0.30 ppm nitrogen dioxide inhalation potentiates exercise-induced bronchospasm in asthmatics. *Am Rev. Resp. Dis.* 134:1203-1208.
12. Bouhuys, A. 1974. Breathing: Physiology, environment and lung disease. Grune and Stratton. New York.
13. U.S. Environmental Protection Agency. 1982. Air quality criteria for oxides of nitrogen. Research Triangle Park, N.C. EPA-600/8-82-026F.
14. Evans, M.J., Johnson, L.V., Stephens, R.J., & Freeman, G. 1976. Renewal of the terminal bronchiolar epithelium in the rat following exposure to NO₂ or O₃. *Lab. Invest.* 35: 246-257.
15. Mohsenin, V., and Gee, B.L. 1987. Acute effects of nitrogen dioxide exposure on the functional activity of alpha-1-protease inhibitor on brochoalveolar lavage fluid of normal subjects. *Am. Rev. Resp. Dis.* 136:646-650.
16. Mohsenin, V. 1987. Effects of vitamin C on NO₂-induced airway responsiveness in normal subjects: A randomized double-blind experiment. *Am. Rev. Resp. Dis.* 136: 1408-1411.
17. Mohsenin, V. 1988. Airway responses to 2.0 ppm nitrogen dioxide in normal subjects. *Arch. Environ. Health.* 43: 242-246.
18. Frampton, M.W., Morrow, P.E., Cox, C., Gibb, F.R., Speers, D.M., and Utell, M.J. 1991. Effects of nitrogen dioxide exposure on pulmonary function and airway reactivity in normal humans. *Am. Rev. Resp. Dis.* 143:522-527.
19. Morrow, P.E. and Utell, M.J. 1989. Responses is susceptible sub-populations to nitrogen dioxide. *Res. Rep. Health Eff. Inst.* 23:1-44.
20. Roger, L.J., Horstman, D.H., McDonnell, W., Kehrl, H., Ives, P.J., Seal, E., Chapman, R., and Massaro, R.J. 1990. Pulmonary function, airway responsiveness, and respiratory symptoms in asthmatics following exercise in NO₂. *Toxicol. Ind. Health.* 6:155-171.

21. Kleinman, M.T., Balley, R.M., Linn, W.S., Anderson, K.R., Whynot, J.D., Shamoo, D.A., Hackney, J.D. 1983. Effects of 0.2 ppm nitrogen dioxide on pulmonary function and response to bronchoprovocation in asthmatics. *J. Toxicol. Environ. Health.* 12:815-826.
22. Linn, W.S., Shamoo, D.A., Avol, E.L., Whynot, J.D., Anderson, K.R., Venet, T.G. and Hackney, J.D. 1986. Dose-response study of asthmatic volunteers exposed to nitrogen dioxide during intermittent exercise. *Arch. Environ. Health.* 41:292-296.
23. Neas, L.M., Dockery, D.W., Ware, J.H., Spengler, J.D., Speizer, F.E., and Ferris, B.G. 1991. Associations of indoor nitrogen dioxide with respiratory symptoms and pulmonary function in children. *Am. J. Epidemiol.* 134: 204-209.
24. Melia, R.J., Florey, C.V., Morris, R.W., Goldstein, B.D., John, H.H., Clark, D., Craighead, I.B. and MacKinlay, J.C. 1982. Childhood respiratory illness and the home environment. II. Association between respiratory illness and nitrogen dioxide, temperature and relative humidity. *Int. J. Epidemiol.* 11: 164-169.
25. Shy, C.M., Creason, J.P., Pearlman, M.E., McClain, K.E., Benson, F.B., and Young, B.B. 1970. The Chattanooga School Children Study. Part II. Effects of community exposure to nitrogen dioxide. *J. Air Pollut. Control Assoc.* 20:582-588.
26. Pearlman, M.E., Finklea, J.F., Creason, J.P. Shy, C.M., Young, M.M., Horton, R.J. 1971. Nitrogen dioxide and lower respiratory illness. *Pediatrics.* 47: 391-398.
27. Gardner, D.E., 1984. Oxidant-induced enhanced sensitivity to infection in animal models and their extrapolation to man. *J. Toxicol. Environ. Health.* 30:23-29.
28. Henry, M.C., Findlay, J., Spengler, J., and Ehrlich. 1970. Chronic toxicity of NO₂ in squirrel monkeys. *Arch. Environ. Health.* 20:566-570.
29. Rose, R.M., Fuglestad, J.M., Skornik, W.A., Hammer, S.M., Wolfthal, S.F., Bech, B.D. and Brain, J.D. 1988. The pathophysiology of enhanced susceptibility of murine cytomegalovirus respiratory infection during

- short-term exposure to 5 ppm nitrogen dioxide. *Am. Rev. Resp. Dis.* 137:912-917.
30. Delvin, R., Horstman, D., Becker, S., Gerrity, T., Madden, M., and Koren, H. 1992. Inflammatory response in humans exposed to 2.0 ppm NO₂ (abstract). *Am. Rev. Resp. Dis.* 145: A455.
 31. Frampton, M.W., Smeglin, A.M., Roberts, N.J., Finklestein, J.N., Morrow, P.E., and Utell, M.J. 1989. Nitrogen dioxide exposure in vivo and human alveolar macrophage inactivation of influenza virus in vitro. *Environ. Res.* 48:179-192.
 32. Barnes, P.J. 1986. Neural control of human airways in health and disease. *Am. Rev. Resp. Dis.* 134:1269-1314.
 33. Chatkin, L.W. and Saunders, L.Z. 1974. Experimental chronic bronchitis. *Lab. Invest.* 30: 145-154.
 34. Giddens, W.E., and Fairchild, G.A. 1972. Effects of sulfur dioxide on the nasal mucosa of mice. *Arch. Environ. Health.* 25:166-173.
 35. Carson, J.L., Collier, A.M., Hu, S.C., Smith, C.A., and Stewart, P. 1987. The appearance of compound cilia in the nasal mucosa of normal human subjects following acute in vivo exposure to sulfur dioxide. *Environ. Res.* 42:155-165.
 36. Committee of the Environmental and Occupational Health Assembly of the American Thoracic Society. 1996. Health effects of outdoor air pollution. Part 1. *Am. J. Resp. Crit. Care Med.* 153: 3-50.
 37. Bedi, J.F., Folinsbee, L.J., and Horvath, S.M. 1984. Pulmonary function effects of 1.0 and 2.0 ppm sulfur dioxide exposures in active young male nonsmokers. *J. Air Pollut. Control Assoc.* 34: 1117-1121.
 38. Schacter, E.N., Witek, T.J., Beck, G.J., Hosein, H.R., Colice, G., Leaderer, B.P., and Cain, W. 1984. Airway effects of low concentrations of sulfur dioxide: dose-response characteristics. *Arch. Environ. Health.* 38:34-43.

39. Folinsbee, L.J., Bedi, J.F., and Horvath, S.M. 1985. Pulmonary response to threshold levels of sulfur dioxide (1.0 ppm) and ozone (0.3 ppm). *J. Appl. Physiol.* 58:1783-1787.
40. Sheppard, D., Saisho, A., Nadel, J.A. and Boushey, H.A. 1981. Exercise increases sulfur dioxide-induced bronchoconstriction in asthmatic subjects. *Am. Rev. Resp. Dis.* 123: 486-491.
41. Sheppard, D., Wong, W.S., Uehara, C.D., Nadel, J.A. and Boushey, H.A. 1980. Lower threshold and greater bronchomotor responsiveness of asthmatic subjects to sulfur dioxide. *Am. Rev. Resp. Dis.* 122: 873-878.
42. Kirkpatrick, M.D., Sheppard, D., Nadel, J.A., and Boushey, H.A. 1982. Effects of the oronasal breathing route on sulfur dioxide-induced bronchoconstriction in exercising asthmatic subjects. *Am. Rev. Resp. Dis.* 125: 627-631.
43. Schenker, M.B., Samet, J.M., Speizer, F.E., Gruhl, J., and Batterman, S. 1983. Health effects of air pollution due to coal combustion in the Chestnut Ridge region on Pennsylvania: Results of cross-sectional analysis of adults. *Arch. Environ. Health.* 38:325-330.
44. Euler, G.L., Abbey, D.E., Magis, A.R., Hodgkins, J.E. 1987. Chronic obstructive pulmonary disease symptom effects of long-term cumulative exposure to ambient levels of total suspended particulates and sulfur dioxide in California Seventh-Day Adventist residents. *Arch. Environ. Health.* 32: 279-285.
45. Hatch, G.E., Boykin, E., Graham, J.A., Lewtas, J., Pott, F., Loud, K., and Mumford, J.S. 1985. Inhalable particles and pulmonary host defense: in vitro and in vivo effects of ambient air and combustion particles. *Environ. Res.* 36:67-80.
46. Beck, B.D., Brain, J.D., and Wolfthal, S.F. 1988. Assessment of lung injury produced by particulate emissions of space heaters burning automotive wasteoil. *Ann. Occup. Hyg.* 32:257-265.
47. Dockery, D.W., Ware, J.H., Ferris, B.G., Speizer, F.E., Cook, N.R., Herman, S.M., 1982. Changes in pulmonary function in children

- associated with air pollution episodes. *J. Air Pollut. Control Assoc.* 32: 937-942.
48. Dassen, W., Brunekreef, B., Hoek, G., Hofschreuder, P., Staatsen, B., deGroot, H., Schouten, E., and Biersteker, K. 1986. Decline in children's pulmonary function during an air pollution episode. *J. Air Pollut. Control Assoc.* 36: 1223-1227.
 49. Pope, C.A. & Kanner, R.E. 1993. Acute effects of PM₁₀ pollution on pulmonary function of smokers with mild to moderate COPD. *Am. Rev. Resp. Dis.* 147: 1336-1340.
 50. Levy, D., Gent, M. and Newhouse, M.T. 1977. Relationship between acute respiratory illness and air pollution levels in an industrial city. *Am. Rev. Resp. Dis.* 116: 167-173.
 51. Pope, C.A., Dockery, D.W., Spengler, J.D., and Raizenne, M.E. 1991. Respiratory health and PM₁₀ pollution: A daily time series analysis. *Am. Rev. Resp. Dis.* 144: 668-674.
 52. Ostro, B.D., and Rothschild, S. 1989. Air pollution and acute respiratory morbidity: An observational study of multiple pollutants. *Envir. Res.* 50: 238-247.
 53. Dockery, D.W., and Schwartz, J. 1992. Particulate air pollution and daily mortality in Steubenville, Ohio. *Am. J. Epidemiol.* 135: 12-19.
 54. Samet, J.M., Speizer, F.E., Bishop, Y., Spengler, J.D. and Ferris, B.G. 1981. The relationship between air pollution and emergency room visits in an industrial community. *J. Air Pollut. Control Assoc.* 31: 236-240.
 55. Schwartz, J & Marcus, A. 1990. Mortality and air pollution in London: A time-series analysis. *Am. J. Epidemiol.* 130: 18-194.
 56. Schwartz, J. & Dockery, D.W. 1992. Increased mortality in Philadelphia associated with daily air pollution concentrations. *Am. Rev. Resp. Dis.* 145: 600-604.
 57. Dockery, D.W. & Pope, C.A. 1994. Acute respiratory effects of particulate air pollution. *Annu. Rev. Public Health.* 15: 107-132.

58. Ferris, B.G. 1978. Epidemiology standardization project. *Am. Rev. Resp. Dis.* 118: 1-120.
59. Imbruce R. 1990. Standardized testing of five commercially available peak flow meters. Unpublished data on file, HealthScan Products Inc.
60. U.S. Environmental Protection Agency. 1991. National Air Quality and Emissions Trends Report. Office of Air Quality Planning and Standards. Research Triangle Park, N.C. 450-R-92-001.
61. Knudson, R.J., Lebowitz, M.D., Holberg, C.J., and Burrows, B. 1983. Changes in the normal maximal expiratory flow-volume curve with growth and aging. *Am. Rev. Resp. Dis.* 127: 725-734.
62. Godfrey, S., Kamburoff, P.L., and Narin, J.R. 1970. Spirometry, lung volumes and airway resistance in normal children aged 5 to 18 years. *Brit J. Dis. Chest.* 64:15-24.
63. Colley, J.R., and Holland, W.W. 1967. Social and environmental factors in respiratory disease. A preliminary report. *Arch. Environ. Health.* 14: 157-161.
64. He, Q., Liyo, J.P., Wilson, W.E., and Chapman, R.S. 1993. Effects of air pollution on children's pulmonary function in urban and suburban areas of Wuhan, People's Republic of China. *Arch. Environ. Health.* 48: 382-391.
65. Xu, X.P., Dockery, D.W., and Wang, L.H. 1991. Effects of air pollution on adult pulmonary function. *Arch. Environ. Health.* 46: 198-206.
66. Arrosa, W., Spinaci, S., Bugiani, M., Natale, P., Bucca, C., and de Candussio, G. 1987. Changes in lung function of children after an air pollution decrease. *Arch. Environ. Health.* 42: 170-174.
67. Dockery, D.W., Speizer, F. E., Stram, D.O., Ware, J. H., Spengler, J. D., and Ferris, B. G. 1989. Effects of inhalable particles on respiratory health in children. *Am. Rev. Respir. Dis.* 139: 587-594.

HARVEY CUSHING / JOHN HAY WHITNEY
MEDICAL LIBRARY

MANUSCRIPT THESES

Unpublished theses submitted for the Master's and Doctor's degrees and deposited in the Medical Library are to be used only with due regard to the rights of the authors. Bibliographical references may be noted, but passages must not be copied without permission of the authors, and without proper credit being given in subsequent written or published work.

This thesis by _____ has been
used by the following persons, whose signatures attest their acceptance of the
above restrictions.

NAME AND ADDRESS

DATE

