## Health Effects of Particulate Matter Air Pollution

#### C. Arden Pope III Mary Lou Fulton Professor of Economics



Presented at EPA Wood Smoke Health Effects Webinar July 28, 2011

## What we breath impacts our health

 $\rightarrow$  Pure Air--nitrogen (78%),Oxygen (21%), Argon, CO<sub>2</sub>...

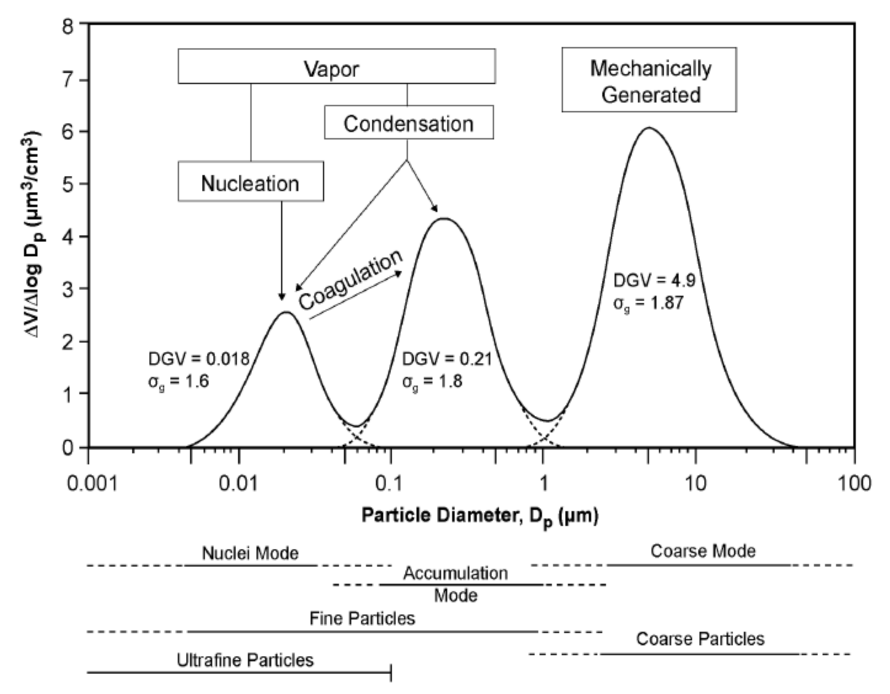
Various gaseous pollutants including: - SO<sub>2</sub>, NO<sub>2</sub>, CO, O<sub>3</sub>...

### Particulate matter:

- Course particles (> 2.5  $\mu$ m in diameter)
- Fine particles (< 2.5 μm in diameter)

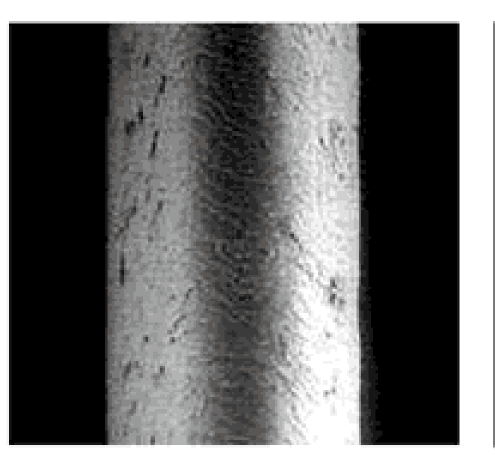


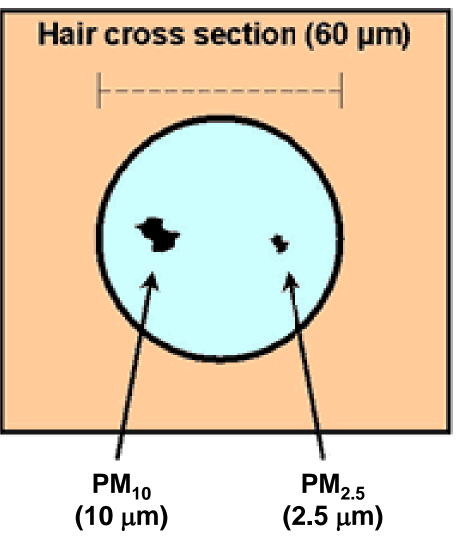
Wood Smoke



EPA PM Criteria Document, 200404

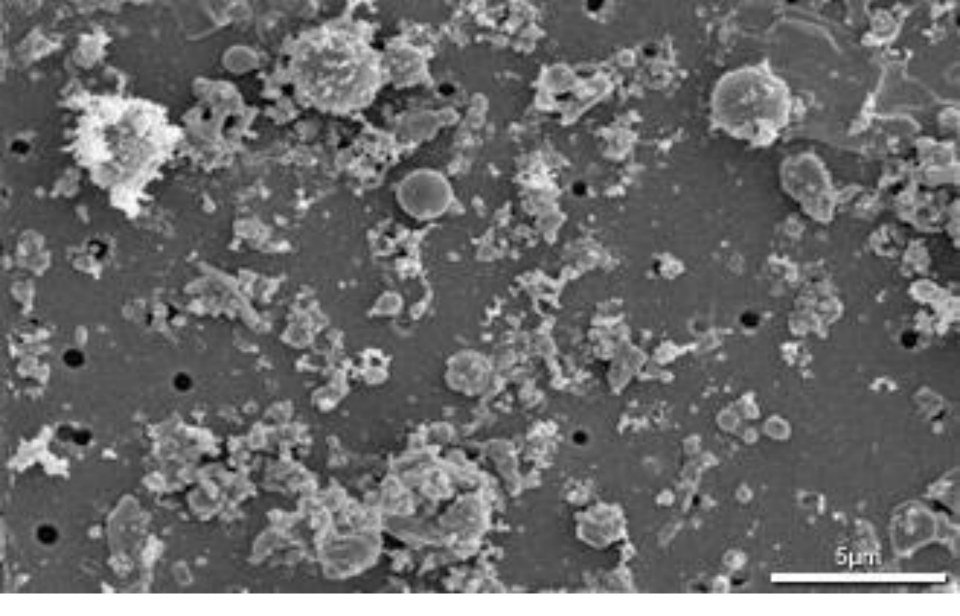
## How small are fine particles?





**Human Hair** 

(60 μm diameter)



Magnified ambient particles (www.nasa.gov/vision/earth/environment)



Studies of short-term exposure (hours-days)

- Episode
- Population-based daily time-series
- Panel-based acute exposure
- Case-crossover

Studies of long-term exposure (years-decades)

- Population-based cross-sectional
- Cohort-based mortality
- Cohort- and panel-based morbidity
- Case-control studies

Intervention/natural experiment (months-years)

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Early "Killer smog" episodes demonstrated that air pollution at extreme levels can contribute to respiratory and cardiovascular disease and death



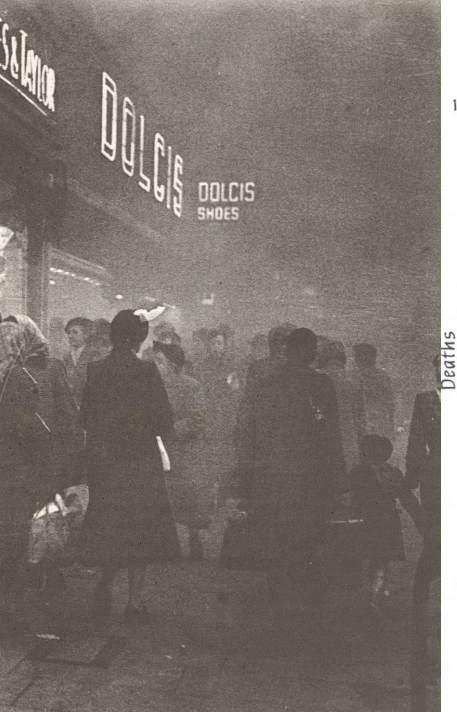
60 deaths (10x expected)



#### Oct. 27-31, 1948: Donora, PA 20 deaths, ½ the town's population fell ill

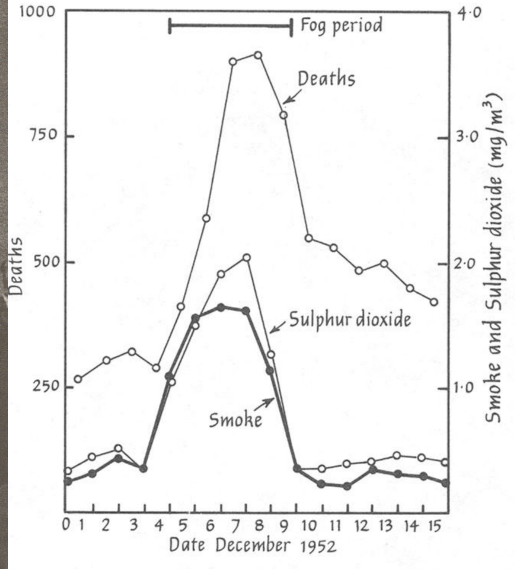






### London Fog Episode, Dec. 1952

THE BIG SMOKE



From: Brimblecombe P. The Big Smoke, Methu

### Utah Valley, 1980s

- Winter inversions trap local pollution
- Natural test chamber



- Local Steel mill contributed ~50% PM<sub>2.5</sub>
- Shut down July 1986-August 1987
- Natural Experiment



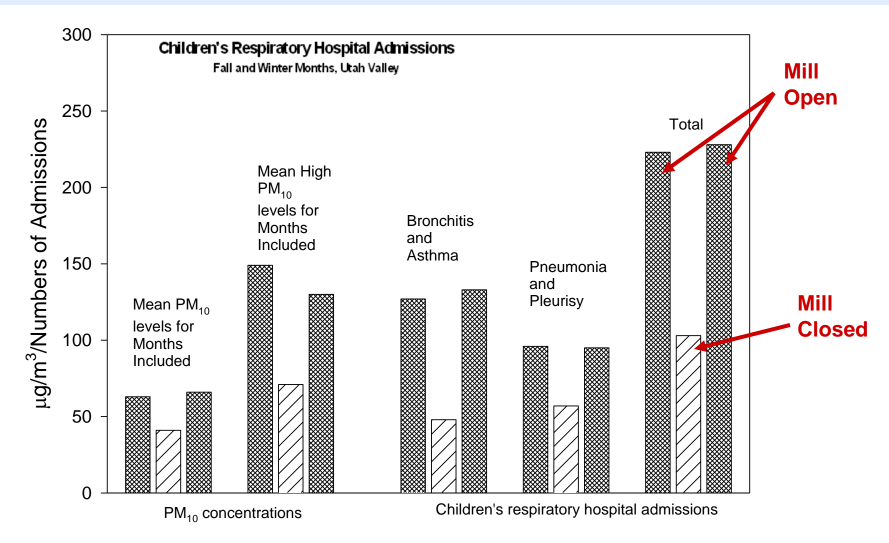


### Large difference in air quality when inversions trap air pollution in valley

**Utah Valley: Clean day** 



## When the steel mill was open, total children's hospital admissions for respiratory conditions **approx. doubled.**



Sources: Pope. Am J Pub Health.1989; Pope. Arch Environ Health. 1991

Studies of short-term exposure (hours-days)

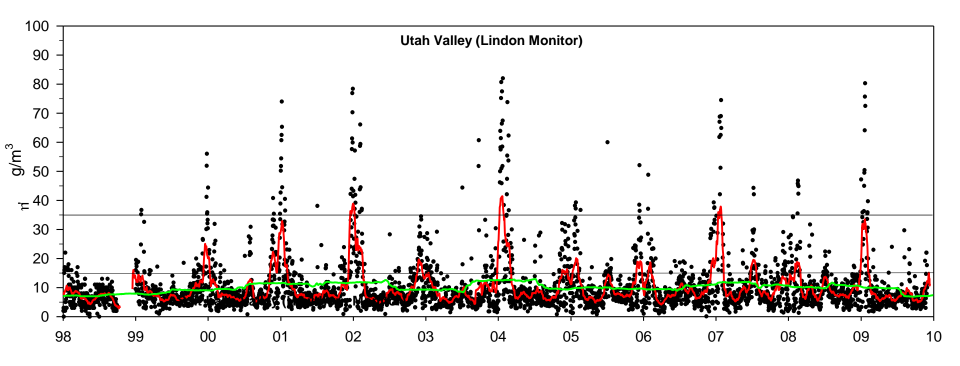
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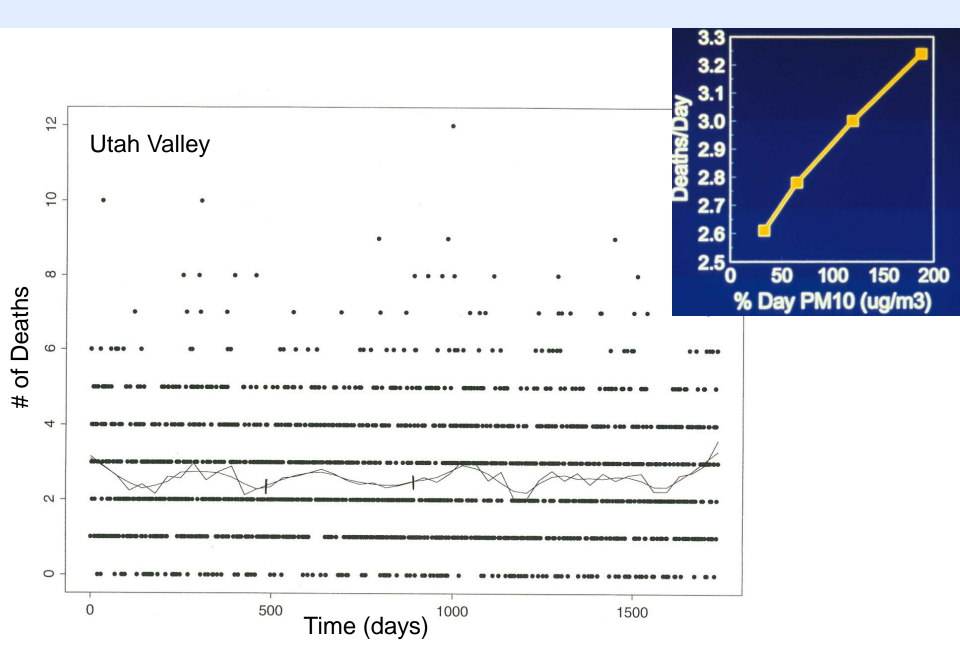
Intervention/natural experiment (months-years)

## Health studies take advantage of **highly variable** air pollution levels that result from inversions.



**PM**<sub>2.5</sub> **concentrations January 1 1998-December 12 2009.** Black dots, 24-hr PM<sub>2.5</sub>; Red line, 30-day moving average PM<sub>2.5</sub>; Green line, 1-yr moving average PM<sub>2.5</sub>.

### Daily changes in air pollution **— daily death counts**

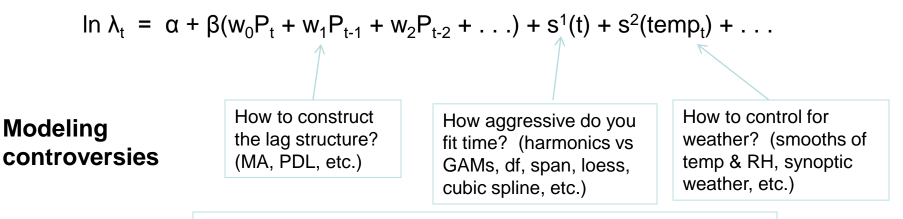


#### **Poisson Regression**

Count data (non-negative integer values). Counts of independent and random occurrences classically modeled as being generated by a Poisson process with a Poisson distribution:

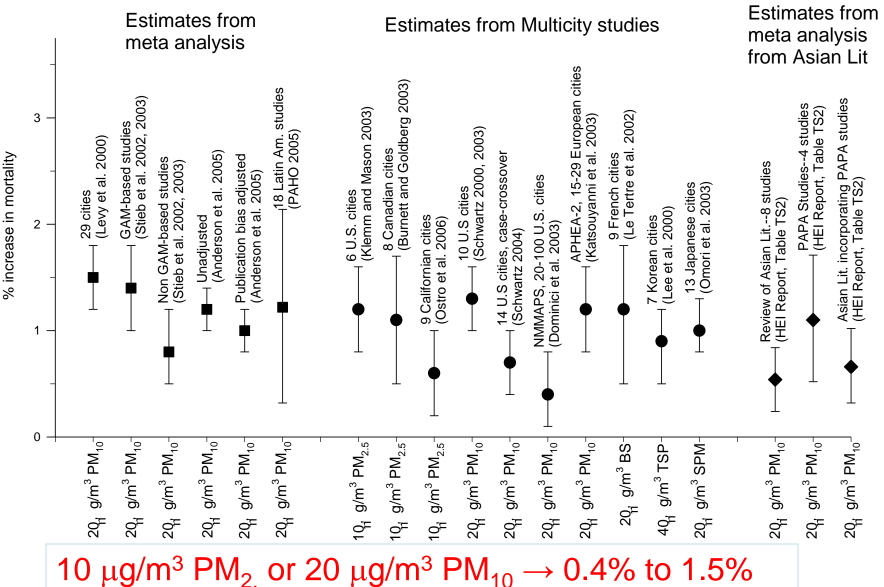
Prob (Y = r) = 
$$e^{(-\lambda)} \frac{\lambda^r}{r!}$$

Note:  $\lambda$  = mean and variance. If  $\lambda$  is constant across time, we have a stationary Poisson process. If  $\lambda$  changes over time due to changes in pollution (P), time trends, temperature, etc., this non-stationary Poisson process can model as:



Also: How to combine or integrate information from multiple cities

#### Daily time-series studies \*\*\*of over 200 cities\*\*\*



10  $\mu$ g/m<sup>3</sup> PIN<sub>2</sub>. or 20  $\mu$ g/m<sup>3</sup> PIN<sub>10</sub>  $\rightarrow$  0.4% to 1.5% increase in relative risk of mortality—Small but remarkably consistent across meta-analyses and multi-city studies.

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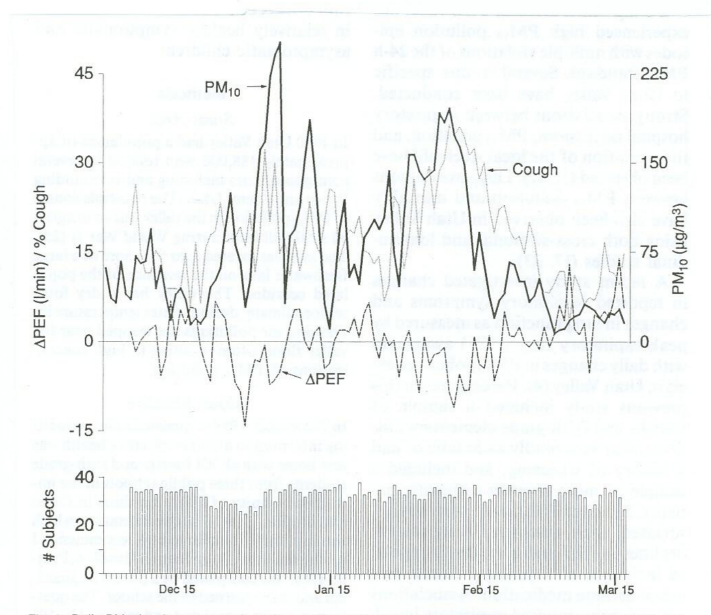
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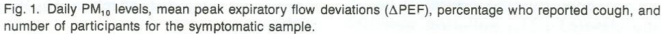
Intervention/natural experiment (months-years)

#### Panel studies of asthmatics and non-asthmatics









### Summary of early Utah Valley epidemiological studies

#### Health effects

- Increased hospital admissions
- Increased respiratory symptoms
- Reduced lung function -
- Increased school absences,
- Increased respiratory and cardiovascular deaths



#### **Study References**

Pope (1989) Am. J. Public Health Pope (1991) Arch. Environ. Health

Pope, Dockery, Spengler, Raizenne (1991) Am. Rev. Resp. Dis. Pope, Dockery (1992) Am. Rev. Resp. Dis. Pope, Kanner (1993) Am. Rev. Resp. Dis.

Ransom, Pope (1992) Environ. Res

Pope, Schwartz, Ransom (1992) Arch. Environ. Health Pope, Kalkstein (1996) Environ. Health Perspect. Pope, Hill, Villegas (1999) Environ. Health Perspect.

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Intervention/natural experiment (months-years)



#### Ischemic Heart Disease Events Triggered by Short-Term Exposure to Fine Particulate Air Pollution

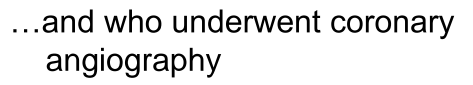
C. Arden Pope III, PhD; Joseph B. Muhlestein, MD; Heidi T. May, MSPH; Dale G. Renlund, MD; Jeffrey L. Anderson, MD; Benjamin D. Horne, PhD, MPH

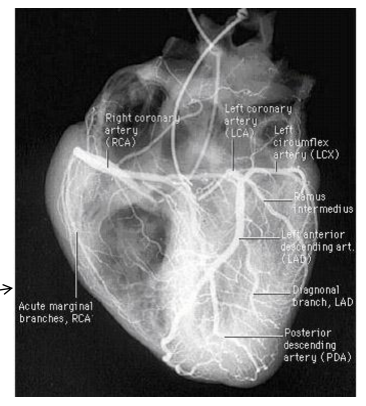


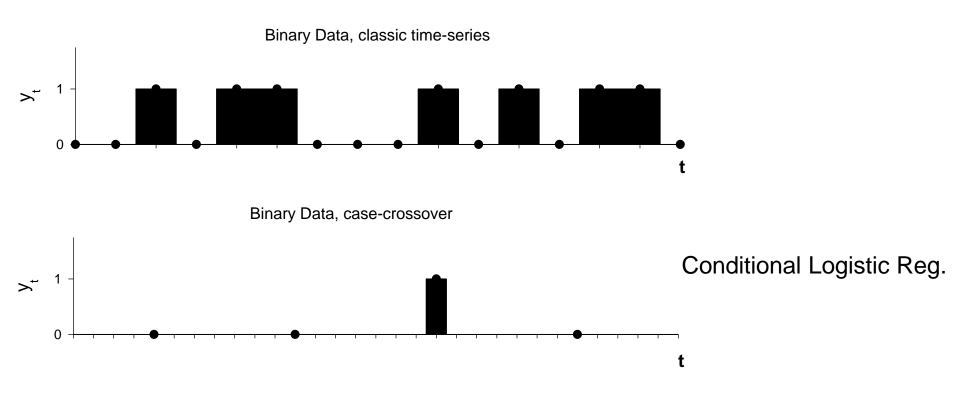
Jeffrey Anderson

### **Methods:**

Case-crossover study of acute ischemic coronary events (heart attacks and unstable angina) in 12,865 well-defined and followed up cardiac patients who lived on Utah's Wasatch Front







Each subject serves as his/her own control.

Control for subject-specific effects, day of week, season, time-trends, etc.—by matching

**Conditional logistic regression:** 

$$\ln \left( \frac{\text{Prob} (Y_t = 1)}{1 - \text{Prob} (Y_t = 1)} \right) =$$

$$\alpha_1 + \alpha_2 + \alpha_3 + \ldots + \alpha_{12,865} + \beta(w_0P_t + w_1P_{t-1} + w_2P_{t-2} + \ldots)$$

Control by matching for: All cross-subject differences (in this case, 12,865 subject-level fixed effects), Season and/or month of year, Time trends, Day of week

**Modeling controversies:** How to select control or referent periods. Time stratified referent selection approach (avoids bias that can occur due to time trends in exposure) (**Holly Janes, Lianne Sheppard, Thomas Lumley** Statistics in Medicine and Epidemiology 2005)

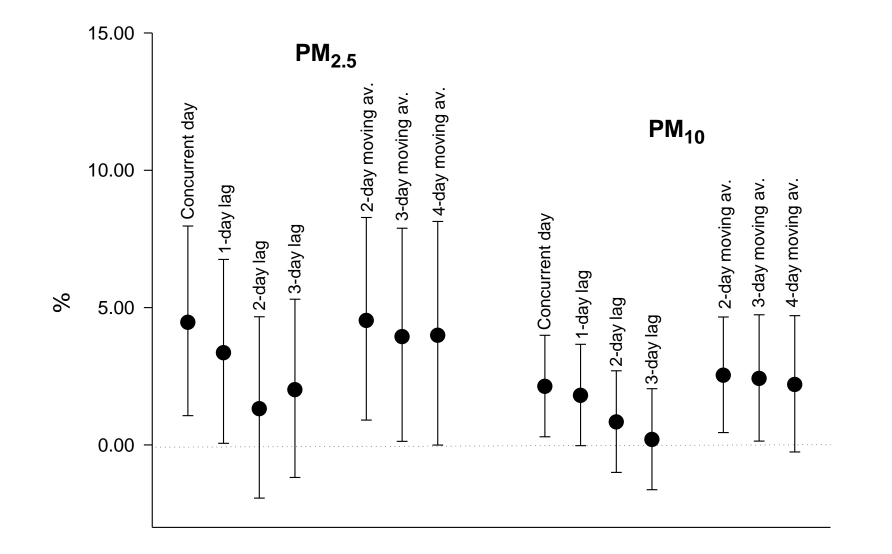
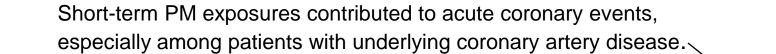


Figure 1. Percent increase in risk (and 95% CI) of acute coronary events associated with 10  $\mu$ g/m<sup>3</sup> of PM<sub>2.5</sub>, or PM<sub>10</sub> for different lag structures.



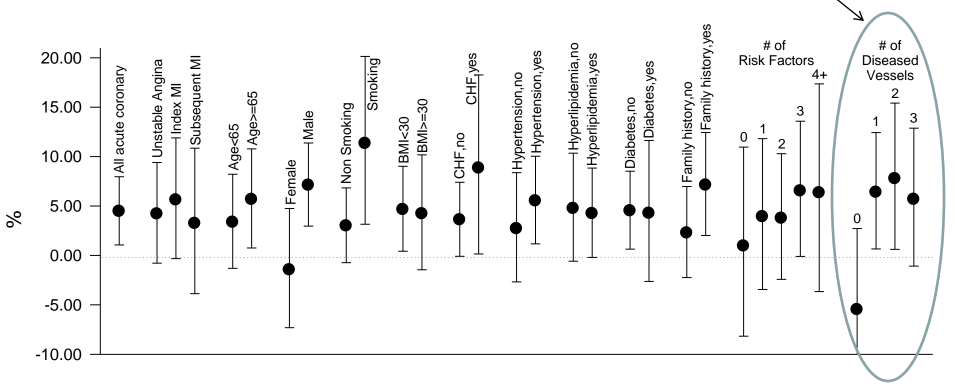


Figure 2. Percent increase in risk (and 95% CI) of acute coronary events associated with  $10 \ \mu g/m^3$  of PM<sub>2.5</sub>, stratified by various characteristics.

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Any Questions?

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Intervention/natural experiment (months-years)

Studies of short-term exposure (hours-days)

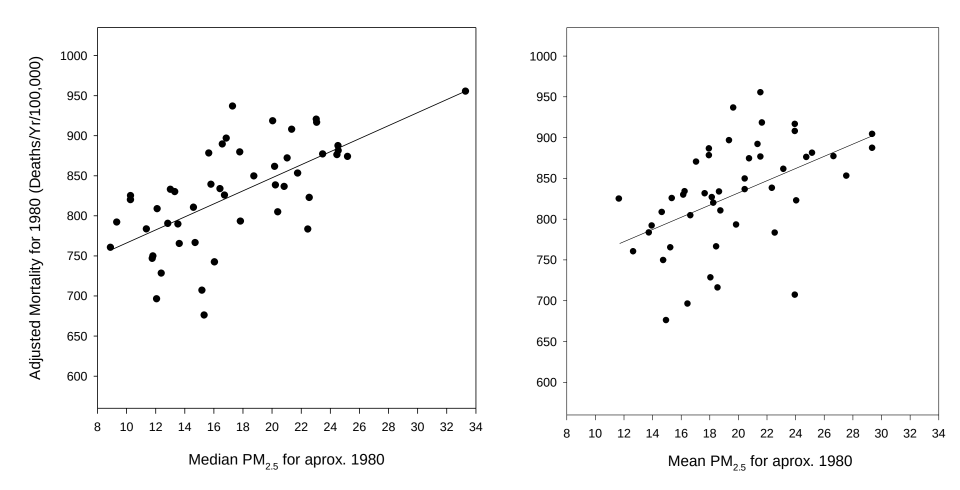
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Age-, sex-, and race- adjusted population-based mortality rates in U.S. cities for 1980 plotted over various indices of particulate air pollution (From Pope 2000).



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### An Association Between Air Pollution and Mortality in Six U.S. Cities



The NEW ENGLAND JOURNAL of MEDICINE 1993

Dockery DW, Pope CA III, Xu X, Spengler JD, Ware JH, Fay ME, Ferris BG Jr, Speizer FE.



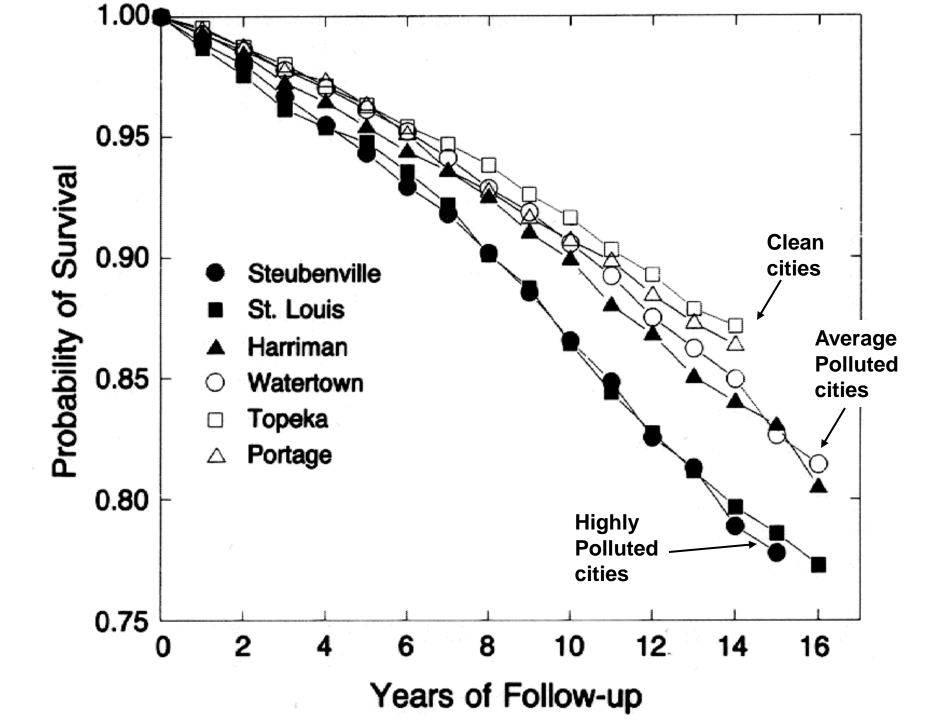
### **Methods:**

14-16 yr prospective follow-up of 8,111 adults living in six U.S. cities.

> Monitoring of TSP  $PM_{10}$ ,  $PM_{2.5}$ ,  $SO_4$ ,  $H^+$ ,  $SO_2$ ,  $NO_2$ ,  $O_3$ .

Data analyzed using survival analysis, including Cox Proportional Hazards Models.

Controlled for individual differences in: age, sex, smoking, BMI, education, occupational exposure.



#### **Cox Proportional Hazards Survival Model**

Cohort studies of outdoor air pollution have commonly used the CPH Model to relate survival experience to exposure while simultaneously controlling for other well known mortality risk factors. The model has the form

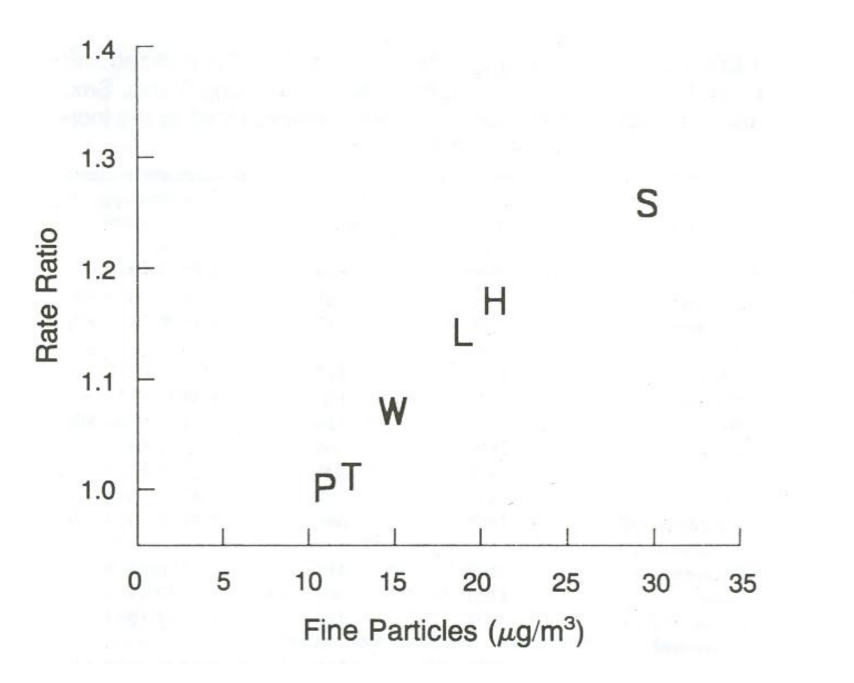
 $\lambda_i^{(l)}(t) = \lambda_0^{(l)}(t) \exp\left(\beta^T x_i^{(l)}(t)\right)$ 

Hazard function or instantaneous probability of death for the *i*<sup>th</sup> subject in the *I*<sup>th</sup> strata.

Baseline hazard function, common to all subjects within a strata. Regression equation that modulates the baseline hazard. The vector  $X_i^{(l)}$ contains the risk factor information related to the hazard function by the regression vector  $\beta$  which can vary in time.

## Adjusted risk ratios (and 95% CIs) for cigarette smoking and $PM_{2.5}$

Cause of	Current Smoker,	Most vs. Least
Death	25 Pack years	Polluted City
All	<b>2.00</b> (1.51-2.65)	<b>1.26</b> (1.08-1.47)
Lung	<b>8.00</b>	<b>1.37</b>
Cancer	(2.97-21.6)	(0.81-2.31)
Cardio-	<b>2.30</b>	<b>1.37</b>
pulmonary	(1.56-3.41)	(1.11-1.68)
All	<b>1.46</b>	<b>1.01</b>
other	(0.89-2.39)	(0.79-1.30)



#### **Particulate Air Pollution as a Predictor of Mortality in a**



Michael Thun

#### **Prospective Study of U.S. Adults**

Pope CA III, Thun MJ, Namboodiri MM, Dockery DW, Evans JS, Speizer FE, Heath CW Jr.

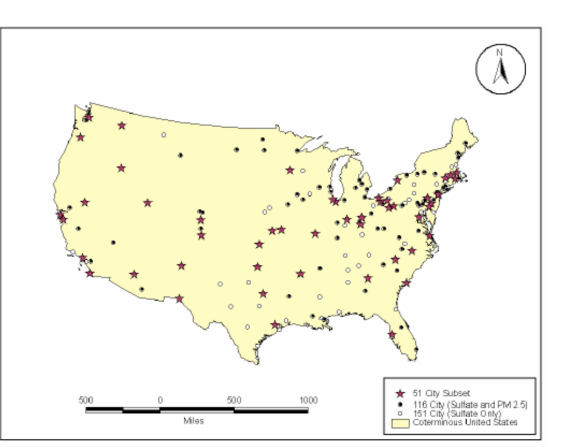


AMERICAN JOURNAL OF Respiratory and Critical Care Medicine®



Clark Heath

**Methods:** Linked and analyzed ambient air pollution data from 51-151 U.S. metro areas with risk factor data for over 500,000 adults enrolled in the ACS-**CPSII** cohort.



1995

Adjusted mortality risk ratios (and 95% CIs) for cigarette smoking the range of sulfates and fine particles

Cause of	Current	Sulfates	Fine
Death	Smoker		Particles
All	<b>2.07</b>	<b>1.15</b>	<b>1.17</b>
	(1.75-2.43)	(1.09-1.22)	(1.09-1.26)
Lung	<b>9.73</b> (5.96-15.9)	<b>1.36</b>	<b>1.03</b>
Cancer		(1.11-1.66)	(0.80-1.33)
Cardio-	<b>2.28</b>	<b>1.26</b>	<b>1.31</b>
Pulmonary	(1.79-2.91)	(1.16-1.37)	(1.17-1.46)
All other	<b>1.54</b>	<b>1.01</b>	<b>1.07</b>
	(1.19-1.99)	(0.92-1.11)	(0.92-1.24)





Dan Krewski Rick Burnett Mark Goldberg and 28 others

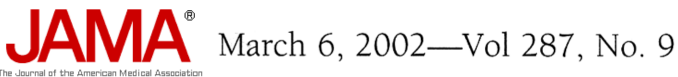
### SPECIAL REPORT

H E A L T H E F F E C T S INSTITUTE

July 2000

Reanalysis of the Harvard Six Cities Study and the American Cancer Society Study of Particulate Air Pollution and Mortality

A Special Report of the Institute's Particle Epidemiology Reanalysis Project



### Lung Cancer, Cardiopulmonary Mortality, and Long-term Exposure to Fine Particulate Air Pollution

C. Arden Pope III, PhD
Richard T. Burnett, PhD
Michael J. Thun, MD
Eugenia E. Calle, PhD
Daniel Krewski, PhD
Kazuhiko Ito, PhD
George D. Thurston, ScD

**Context** Associations have been found between day-to-day particulate ai and increased risk of various adverse health outcomes, including cardiopulmc tality. However, studies of health effects of long-term particulate air pollubeen less conclusive.

**Objective** To assess the relationship between long-term exposure to fir late air pollution and all-cause, lung cancer, and cardiopulmonary mortalit

**Design, Setting, and Participants** Vital status and cause of death data lected by the American Cancer Society as part of the Cancer Prevention II stu going prospective mortality study, which enrolled approximately 1.2 million adu





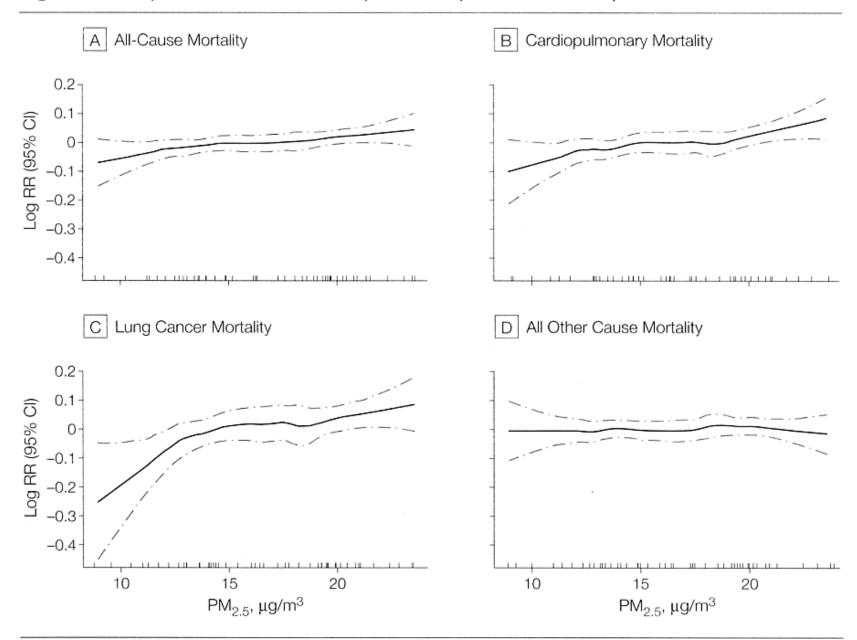








Figure 2. Nonparametric Smoothed Exposure Response Relationship





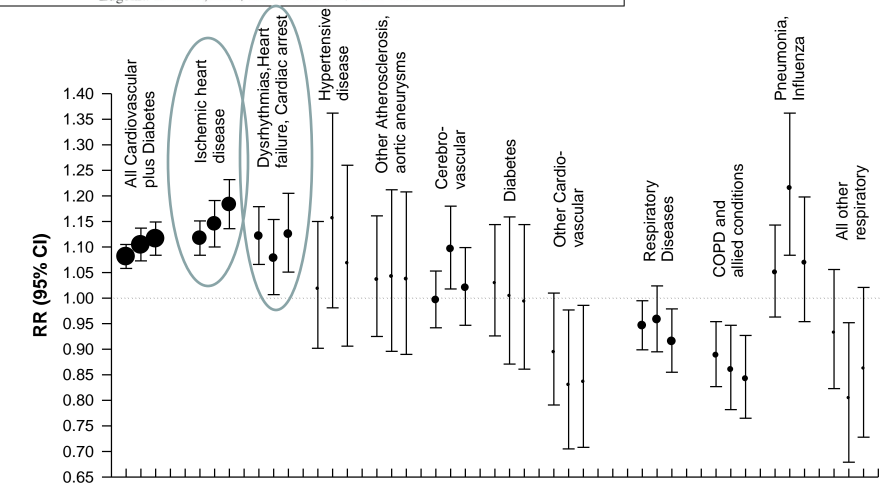
#### Cardiovascular Mortality and Long-Term Exposure to Particulate Air Pollution

Epidemiological Evidence of General Pathophysiological Pathways of Disease

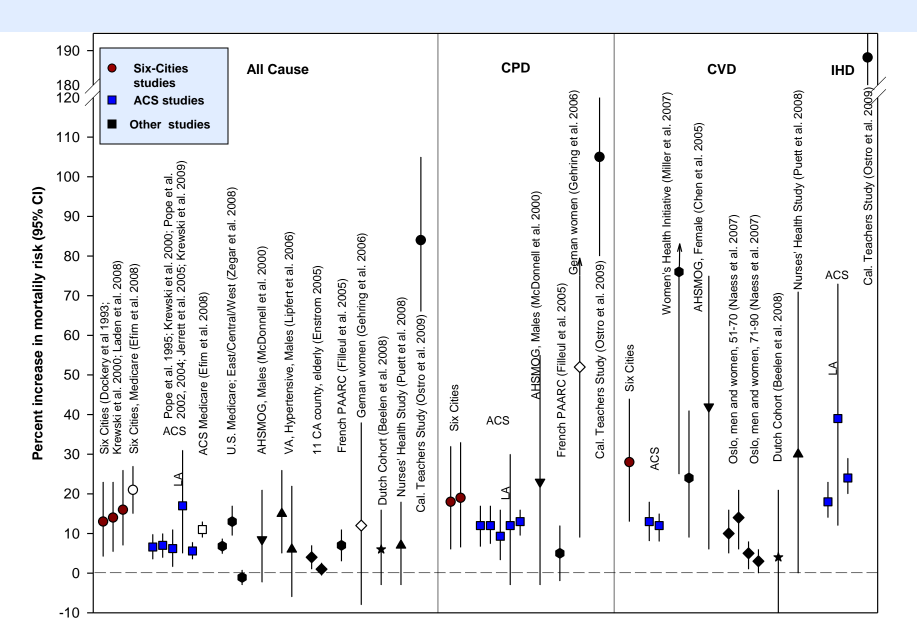
C. Arden Pope III, PhD; Richard T. Burnett, PhD; George D. Thurston, ScD; Michael J. Thun, MD; Eugenia E. Calle, PhD; Daniel Krewski, PhD; John J. Godleski, MD



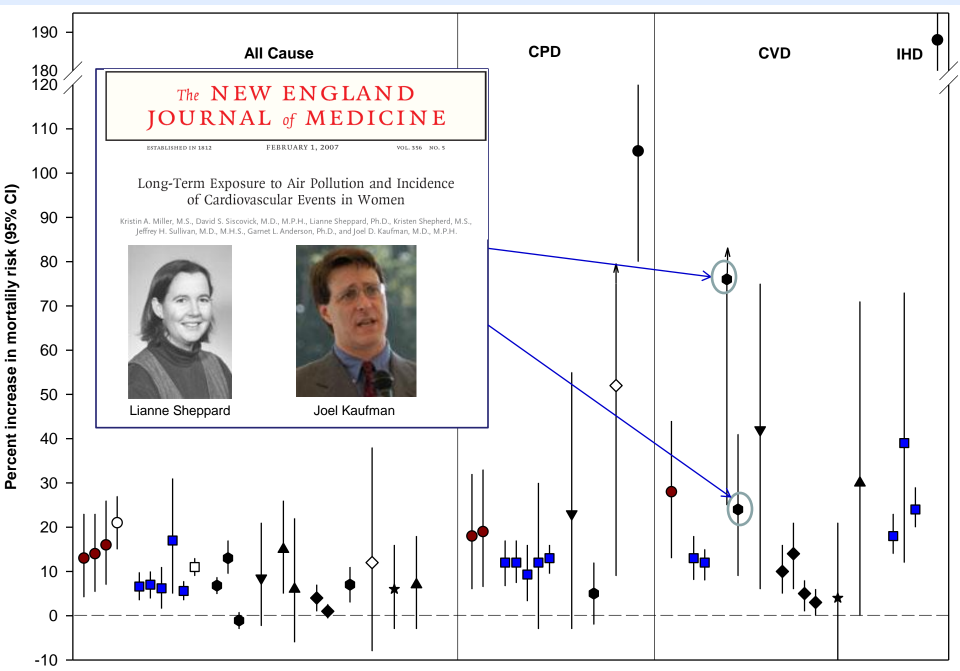
John Godleski

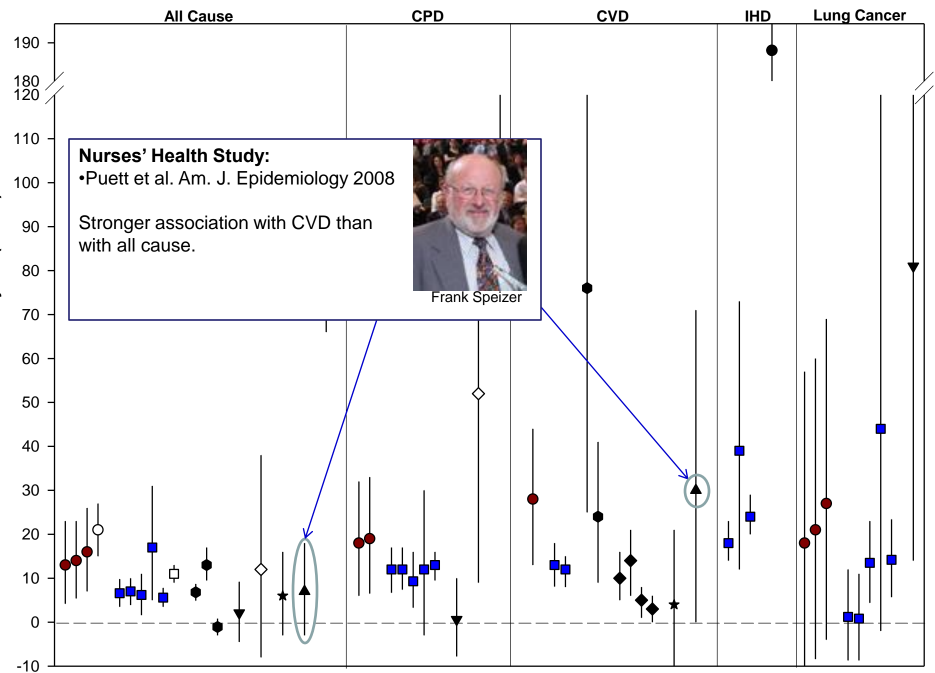


### Other cohort studies have shown associations between exposure to fine PM and increased risk of cardiovascular death.

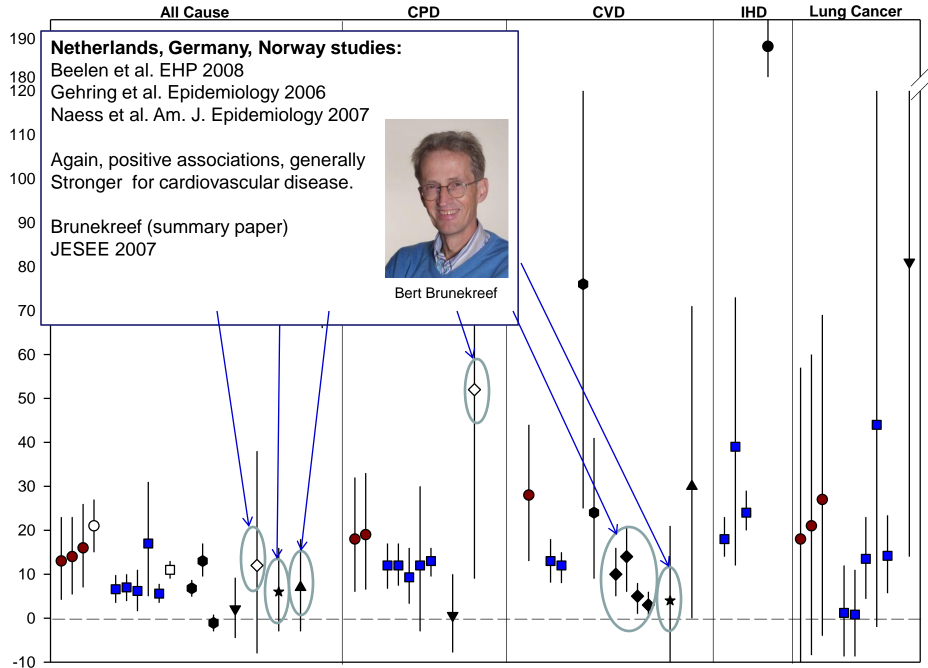


#### Women's Health Initiative Study



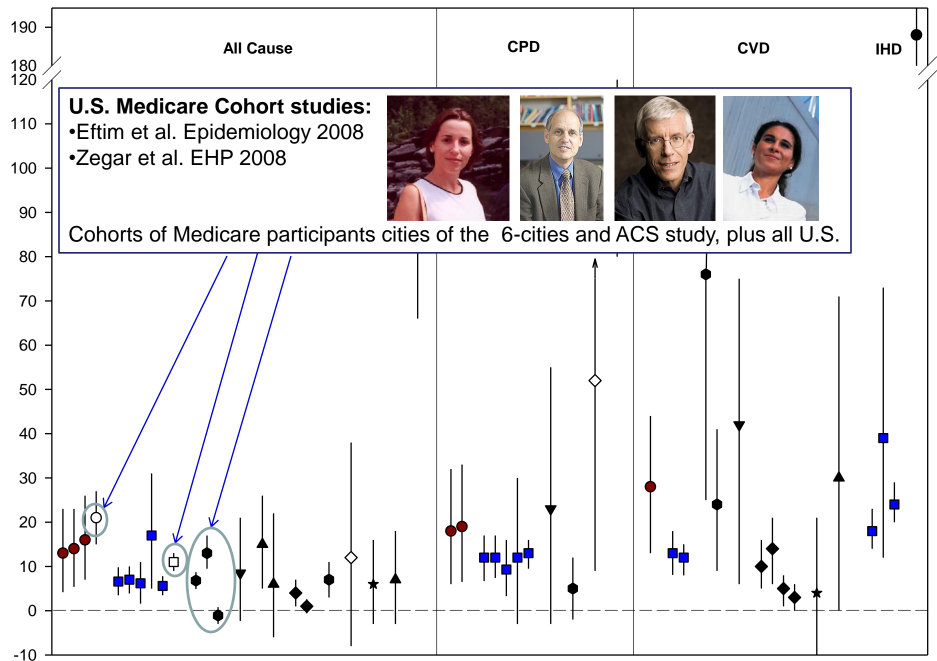


Percent increase in mortalily risk (95% CI)



Percent increase in mortalily risk (95% CI)

#### **U.S. Medicare Cohort Studies**



## This presentation not organized chronologically, but methodologically

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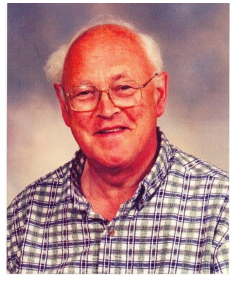
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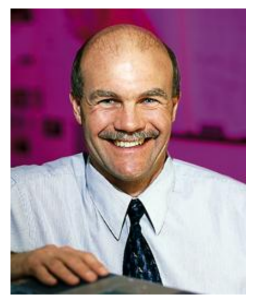
Controlled experimental human and animal

#### Southern California Children's Health Study

# Effects of air pollution on children's health, especially lung function growth.



David Bates, Advisor

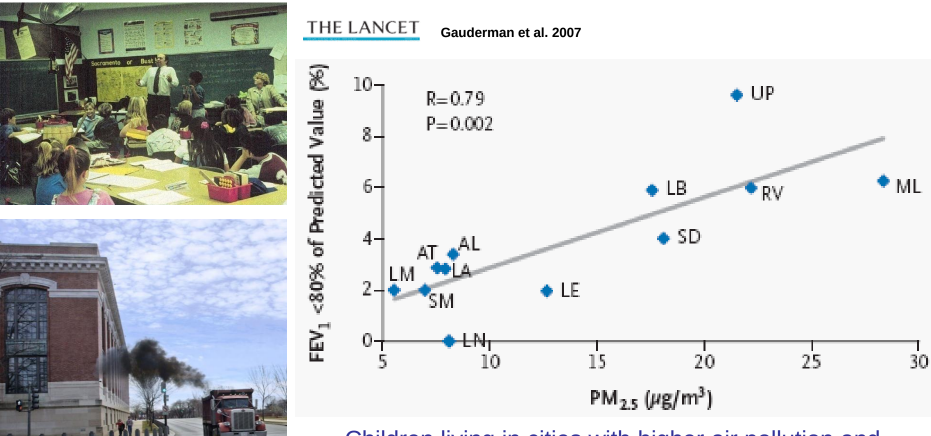


W. James Gauderman



John Peters

### Southern California Children's Health Study, has shown that air pollution impacts lung development in children.



Children living in cities with higher air pollution and living near major traffic sources showed greater deficits in lung function growth.

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#### The NEW ENGLAND JOURNAL of MEDICINE

#### Fine-Particulate Air Pollution and Life Expectancy in the United States

C. Arden Pope, III, Ph.D., Majid Ezzati, Ph.D., and Douglas W. Dockery, Sc.D.

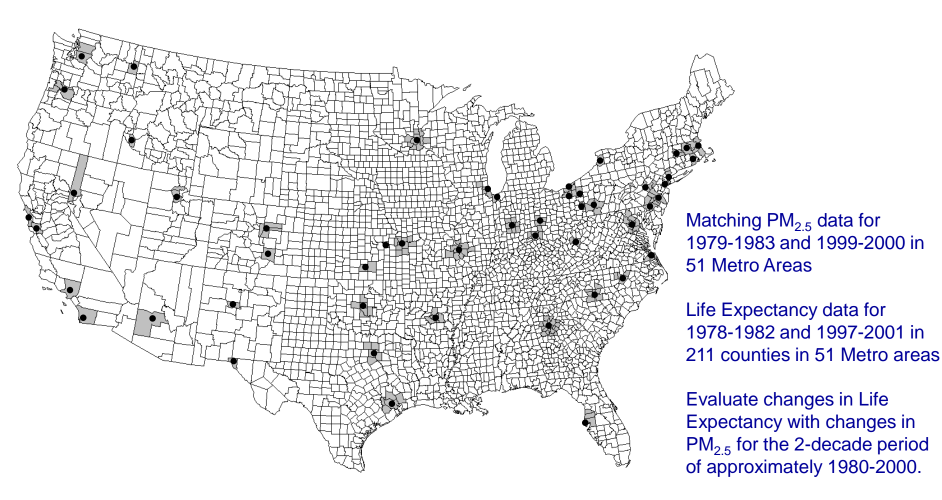
January 22, 2009





Majid Ezzati

Doug Dockery



#### **Covariates included in the regression models**

Changes in socio-economic and demographic variables (from U.S. Census Data):

➢Per capita income

➢Population

≻5-yr in-migration

➢High-school graduates

➤Urban population

➢Black proportion of population

Hispanic proportion of population

Proxy cigarette smoking variables—available for all 211 counties

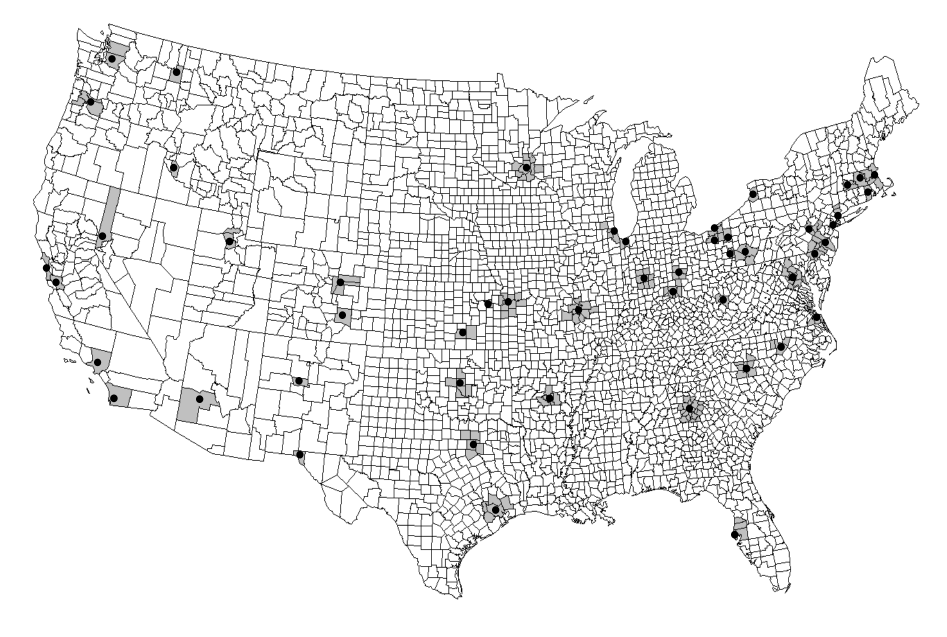
➤COPD mortality rates

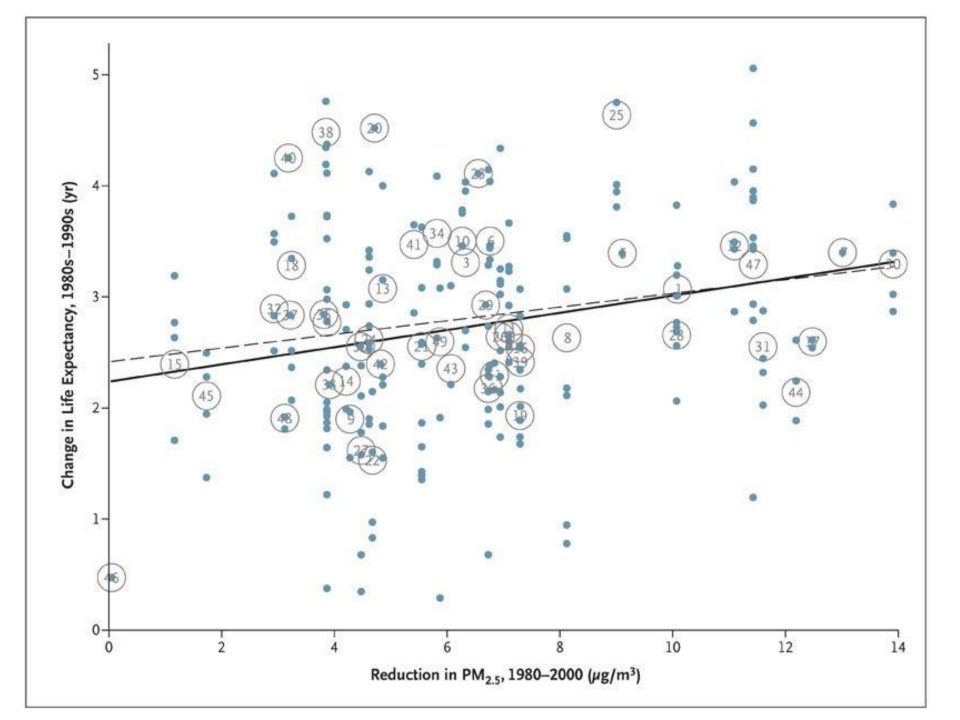
Lung Cancer mortality rates

Survey-based metro-area estimates of smoking prevalence

- National Health Interview Survey (1978-1980)
- Behavioral Risk Factor Surveillance System (1998-2000)
- Matching data available for only 24 of 51 metro areas

Clustered standard errors (clustered by the 51 metro areas) were estimated for all models except for analysis that included only the 51 largest counties in each metro area.



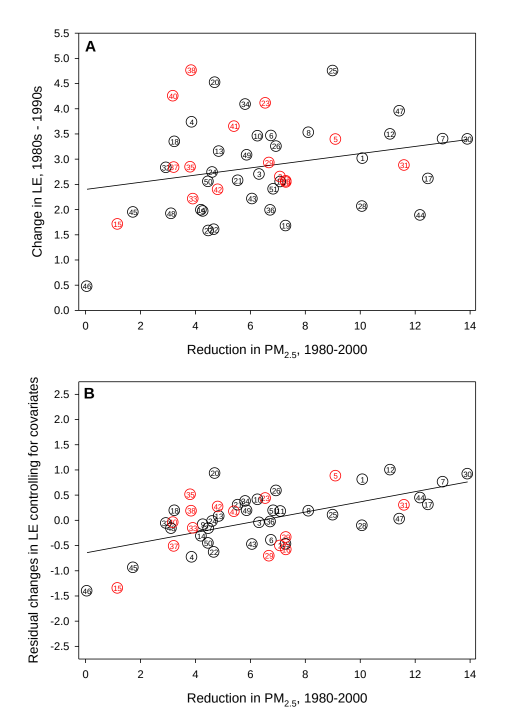


### A 10 $\mu$ g/m<sup>3</sup> decrease in PM<sub>2.5</sub> was associated with a <u>7.3 (± 2.4) month</u> increase in life expectancy.

Table 2. Results of Selected Regression Models, Including Estimates of the Increase in Life Expectancy Associated with a Reduction in PM<sub>2.5</sub> of 10 µg per Cubic Meter, Adjusted for Socioeconomic, Demographic, and Proxy Indicators for Prevalence of Smoking.\*

Variable	Model 1	Model 2	Model 3	Model 4	Model 5†	Model 63	Model 7
				years			
Intercept	2.25±0.21§	0.80±0.19§	1.78±0.27§	1.75±0.27§	2.02±0.34§	1.71±0.51§	2.09±0.36§
Reduction in PM <sub>2.5</sub> (10 µg/m <sup>3</sup> )	0.72±0.29¶	0.83±0.20§	0.60±0.20§	0.61±0.20§	0.55±0.24¶	1.01±0.25§	0.95±0.23§
Change in income (in thousands of \$)	_	0.17±0.02§	0.13±0.02§	0.13±0.01§	0.11±0.02§	0.15±0.04§	0.11±0.02§
Change in population (in hundreds of thousands)	—	0.08±0.02§	0.05±0.02§	0.06±0.02§	0.05±0.02§	0.04±0.02	0.05±0.02¶
Change in 5-yr in-migration (proportion of population) $\ ^{\pm\pm}$	_	0.19±079	1.28±0.80	_	_	-0.02±1.83	_
Change in high-school graduates (proportion of population)	-	0.17-0.56	-0.11±0.53	-	-	-0.90±0.86	—
Change in urban residence (proportion of population)	_	-0/6±0.32¶	-0.40±0.25	-	-	0.03±1.88	_
Change in black population (proportion of population) $ \uparrow\uparrow$	_	-1.94±0.58§	-2.74±0.58§	-2.70±0.64§	-2.95±0.78§	-5.06±2.12§	-5.98±1.99§
Change in Hispanic population (proportion of population)	- /	1.46±1.23	1.33±1.10	-	-	2.44±2.22	_
Change in lung-cancer mortality rate (no./10,000 population)	- /	_	-0.07±0.02§	-0.06±0.02§	-0.07±0.03¶	0.01±0.03	0.02±0.03
Change in COPD mortality rate (no./10,000 population)	_/	_	-0.07±0.02§	-0.08±0.02§	-0.09±0.03§	-0.15±0.06§	-0.19±0.05§
No. of county units	211	211	211	211	127	51	51
R <sup>2</sup> ;;;;	0.05	0.47	0.55	0.53	0.60	0.76	0.74

This increase in life expectancy persisted even after controlling for socio-economic, demographic, or smoking variables



## This presentation not organized chronologically, but methodologically

Studies of short-term exposure (hours-days)

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- Case-crossover

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- Population-based cross-sectional
- Cohort-based mortality
- Cohort- and panel-based morbidity
- Case-control studies

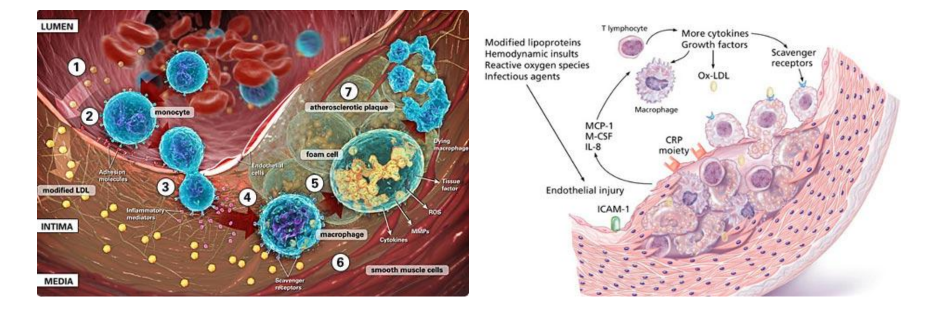
Intervention/natural experiment (months-years)

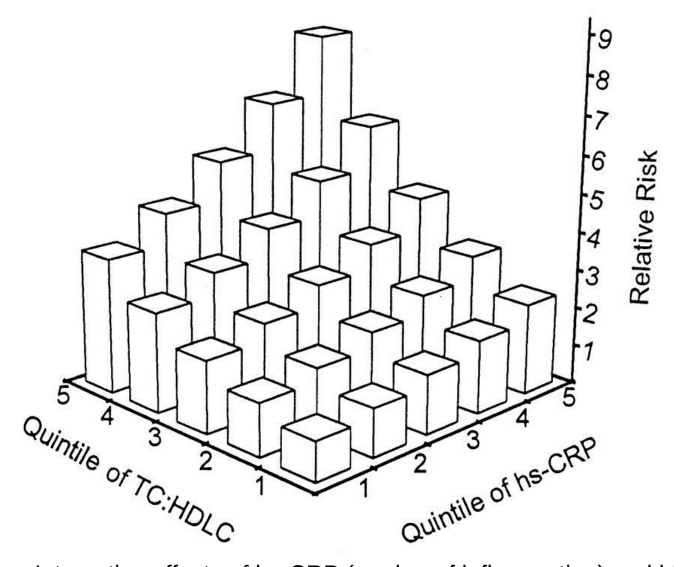
Controlled experimental human and animal

### Cardiovascular disease as part of chronic and acute inflammatory processes.

By the early 2000s, there was increasingly compelling evidence that inflammation is a major accomplice with LDL cholesterol in the initiation and progression of atherosclerosis.

Furthermore, inflammation contributes to acute thrombotic complications of atherosclerosis, increasing the risk of making atherosclerotic plaques more vulnerable to rupture, clotting, and precipitating acute cardiovascular or cerebrovascular events (MI or ischemic stroke).





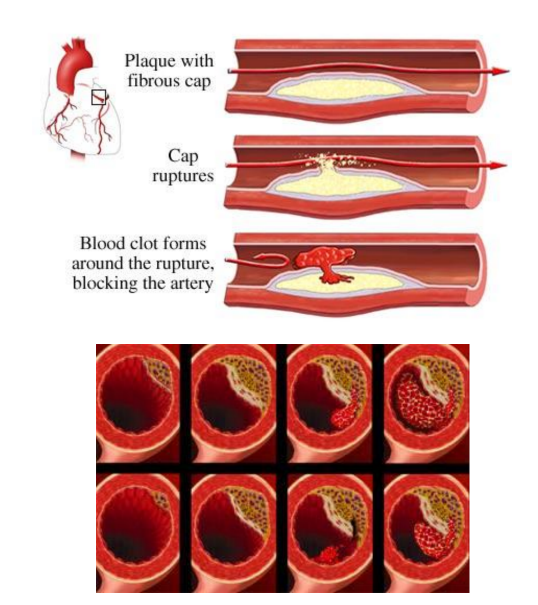


Interactive effects of hs-CRP (marker of inflammation) and blood lipids.



Ridker PM. 2001;103:1813-1818.

**Fine Particulate** exposure **Pulmonary and** systemic inflammation and oxidative stress (along with blood lipids) **Progression and** destabilization of atherosclerotic plaques

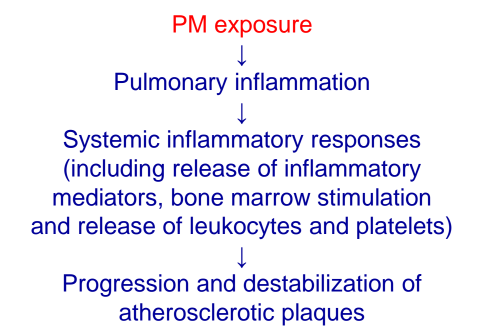




**Experimental evidence of biological effects of PM extracted from filters** (Ghio, Costa, Devlin, Kennedy, Frampton, Dye, et al. 1998-2004)

- Acute airway injury and inflammation in rats and humans
- *In vitro* oxidative stress and release of proinflammatory mediators by cultured respiratory epithelial cells
- Differential toxicities of PM when the mill was operating versus when it was not (metals content and mixtures?)

#### A series of studies by van Eeden, Hogg, Suwa et al. (1997-2002) suggest:



Stephan van Eeden



James Hogg

In rabbits naturally prone to develop atherosclerosis they found that:

#### PM exposure

Accelerated progression of atherosclerotic plaques with greater vulnerability to plaque rupture

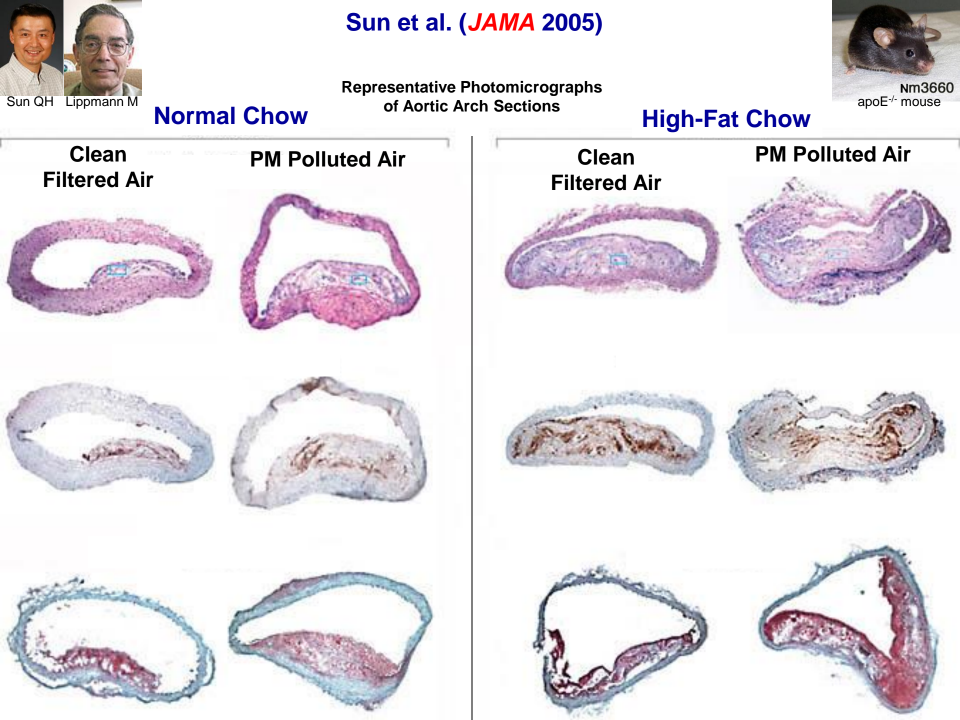


 Table 1. Stylized outline of studies on air pollution and health.

General Study Designs	Examples	Time-scale	Selected Sample of Effects of Elevate	d E <del>xpec</del> ures	Common Statistical Modeling Approaches
Episode	Meuse Valley 1930 <sup>73</sup> Donora 1948 <sup>74</sup> London 1952 <sup>75,76</sup>	Days		e and	Simple Comparative Stats, Graphs
Population-based time-series	Meta analysis <sup>27</sup> Large multi-city mortality <sup>28-31</sup> Large multi-city hospitalizations <sup>77</sup>	Days			Poisson reg., (GAMs, smooths for time , weather etc.)
Panel-based acute exposure	Review resp. effects <sup>19</sup> Review CVD effects <sup>54</sup> HRV/Inflammation <sup>78,79</sup>	Hours-days	<ul> <li>↑ respiratory symptoms</li> <li>↓ pulmonary function</li> <li>↑ markers of inflammation</li> <li>↓ boart rate variability</li> </ul>	,	Linear and Logistic Reg., (fixed effects, temporal autocorr., etc.)
Case-crossover	Multi-c Many stud	dies using	various	ar disease	Conditional Logistic Reg.
Population-based cross-sectional	us.m study des		Linear regression		
Cohort-based mortality	Harvar Americ Wome Medica Europe Swiss c	ality	Survival Analyses, Cox Proportional Hazards models (random effect, spatial autocorr., etc.)		
Cohort- and panel-based morbidity	Harvar so. Cal Athero	nerosclerosis	Various regression modeling strategies (fixed effects, mixed models)		
Case-control studies	Czech Lung C AMI tr Italy DVT <sup>94</sup>	ly coheren	t evidence.		Conditional Logistic Reg.
Intervention/natural experiment	Utah Valley, steel mill <sup>18</sup> Months to years       Various intervention-related improvements in morbidity, mortality and/or life expectancy         Dublin coal ban <sup>65</sup> Forg Kong Sulfur <sup>66</sup> Forg Kong Sulfur <sup>66</sup> Copper smelter strike <sup>64</sup> Forg Kong Sulfur <sup>67</sup> Forg Kong Sulfur <sup>66</sup> U.S. Life Expectancy <sup>67</sup> Forg Kong Sulfur <sup>67</sup> Forg Kong Sulfur <sup>66</sup>				Various comparative stats and regression models
Controlled experimental human studies and animal toxicology	Al Human instillation <sup>96</sup> Variety, usually hours to weeks Human chamber <sup>97,98</sup> Tox, rabbits <sup>99</sup> Tox, hamsters <sup>100,101</sup> Tox, dogs <sup>102</sup> Iox, mice <sup>103-105</sup> Variety, usually hours to weeks Growing complementary evidence of adverse cardiopulmonary health effects of air pollution				Various comparative stats and regression models

#### **AHA Scientific Statement**

#### Brook et al

#### IParticulate Matter Air Pollution and CVDCirculationJune 1, 2010

Table 6.Overall Summary of Epidemiological Evidence of the<br/>Cardiovascular Effects of PM2.5, Traffic-Related, or<br/>Combustion-Related Air Pollution Exposure at Ambient Levels

Health Outcomes	Short-Term Exposure (Days)	Longer-Term Exposure (Months to Years)
Clinical cardiovascular end points from epidemiological studies at ambient pollution concentrations		
Cardiovascular mortality	111	$\uparrow \uparrow \uparrow$
Cardiovascular hospitalizations	111	Ŷ
Ischemic heart disease*	111	$\uparrow \uparrow \uparrow$
Heart failure*	1 1	↑
Ischemic stroke*	1 1	↑
Vascular diseases	1	<b>↑</b> †
Cardiac arrhythmia/cardiac arrest	1	1
Subclinical cardiovascular end points and/or surrogate measures in human studies		
Surrogate markers of atherosclerosis	N/A	Ŷ
Systemic inflammation	1 1	<b>↑</b>
Systemic oxidative stress	1	
Endothelial cell activation/ blood coagulation	↑ ↑	1
Vascular/endothelial dysfunction	↑ ↑	
BP	$\uparrow$ $\uparrow$	
Altered HRV	1 1 1	Ŷ
Cardiac ischemia	1	
Arrhythmias	•1	

Table 7. Summary of Level of Evidence Supporting Global Biological Pathways and Specific Mechanisms Whereby PM<sub>2.5</sub>, Traffic-Related, or Combustion-Related Air Pollution Exposure Can Affect the Cardiovascular System

	Animal Studies	Human Studies
General "intermediary" pathways whereby PM inhalation can instigate extrapulmonary effects on the cardiovascular system		
Pathway 1: Instigation of systemic proinflammatory responses	↑ ↑ ↑	↑↑↑
Pathway 2: Alterations in systemic ANS balance/activity	î	↑ ↑
Pathway 3: PM and/or associated constituents directly reaching the systemic circulation	1	Ŷ
Specific biological mechanisms directly responsible for triggering cardiovascular events		
Vascular dysfunction or vasoconstriction	$\uparrow \uparrow \uparrow$	↑ ↑
Enhanced thrombosis or coagulation potential	↑ ↑	↑ ↑
Elevated arterial BP	$\uparrow$ $\uparrow$	↑ ↑
Enhanced atherosclerosis or plaque vulnerability	an Heart	Ŷ
Arrhythmias	sociation.	1

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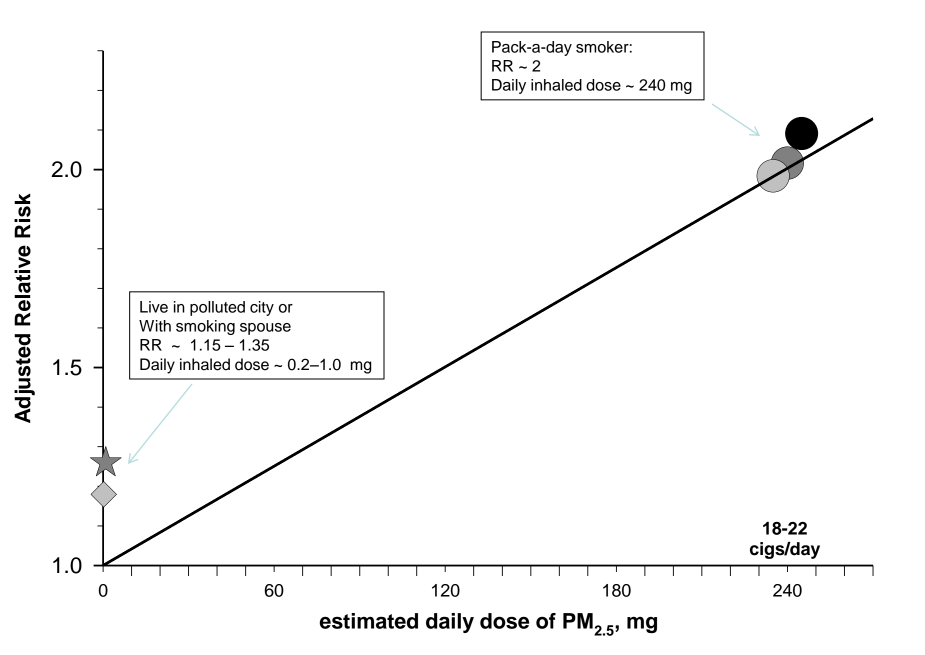
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Controlled experimental human and animal

Any Questions?



Circulation Pope, Burnett, Krewski, et al. 2009.

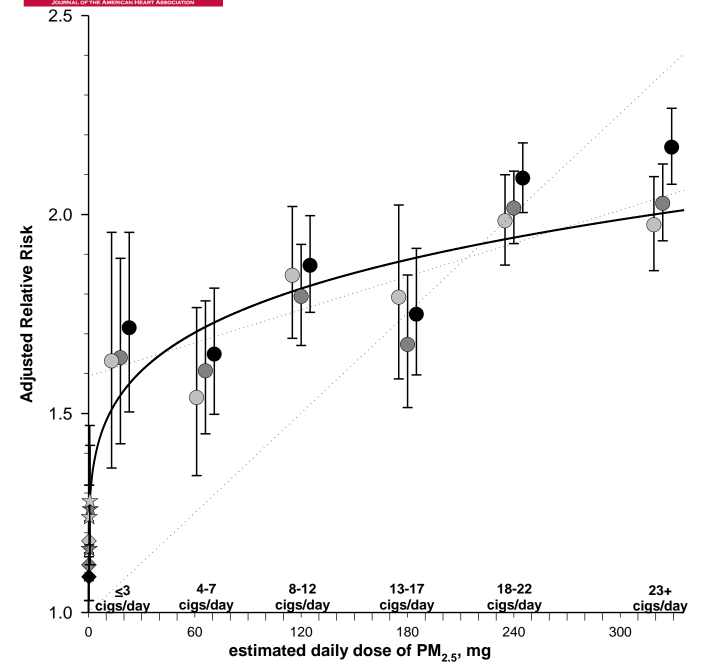


Figure 1. Adjusted relative risks (and 95% CIs) of IHD (light gray), CVD (dark gray), and CPD (black) mortality plotted over estimated daily dose of PM<sub>2.5</sub> from different increments of current cigarette smoking. **Diamonds represent** comparable mortality risk estimates for PM<sub>2.5</sub> from air pollution. Stars represent comparable pooled relative risk estimates associated with SHS exposure from the 2006 Surgeon General's report and from the INTERHEART study.

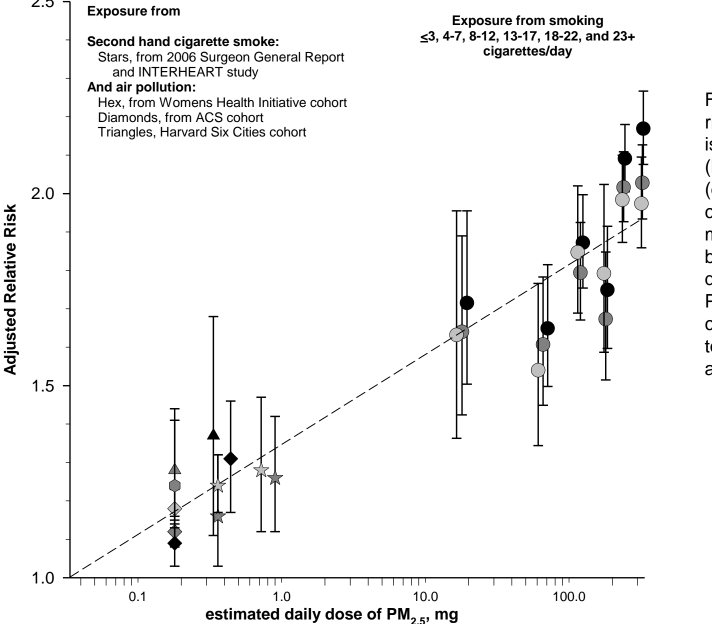


Figure 2. Adjusted relative risks (and 95% CIs) of ischemic heart disease (light gray), cardiovascular (dark gray), and cardiopulmonary (black) mortality plotted over baseline estimated daily dose (using a log scale) of  $PM_{2.5}$  from current cigarette smoking (relative to never smokers), SHS, and air pollution.

#### 2.5