ENVIRONMENTAL EXPOSURE

Traffic related pollution and heart rate variability in a panel of elderly subjects

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Background: Particulate air pollution has been associated with increased cardiovascular deaths and hospital admissions. To help understand the mechanisms, the types of particles most involved, and the types of persons most susceptible, the association between exposure to summertime air pollution and heart rate variability (HRV) was examined in a panel study of 28 elderly subjects.

Methods: Subjects were seen once a week for up to 12 weeks and HRV (SDNN, r-MSSD, PNN₅₀, low frequency/high frequency ratio (LFHFR)) was measured for approximately 30 minutes at each session using a defined protocol. Temperature, day of the week, and hour of the day were controlled, and dummy variables for each subject were controlled for subject specific risk factors.

Results: $PM_{2.5}$ was associated with r-MSSD (-10.1% change for an interquartile range (IQR) increase in exposure (95% CI -2.8 to -16.9)) and PNN₅₀, but stronger associations were seen with black carbon, an indicator of traffic particles, which was also associated with SDNN (-4.6% per IQR (95% CI -2.0 to -7.2)) and LFHFR. Secondary particles were more weakly associated with r-MSSD, as was ozone. No associations were seen with SO₂ or NO₂. CO had similar patterns of association to black carbon, which disappeared after controlling for black carbon. Black carbon had a substantially higher effect on SDNN in subjects who had had a previous myocardial infarction (-12.7%, 95% CI -5.7 to -19.25).

Conclusions: Particles, especially from traffic, are associated with disturbances of autonomic control of the heart.

•he air pollution disasters of the 20th century clearly showed that acute changes in air pollution were associated with similarly rapid changes in daily death rates.1-3 Most notable was the London smog episode of December 1952 where 4000 excess deaths were recorded in a week in response to an extreme episode of increased particle and sulphur dioxide (SO₂) concentrations.¹ A key finding was that sudden deaths were increased more than respiratory or cardiovascular deaths in general.¹ In 1994 we analysed death certificate data from Philadelphia contrasting high and low air pollution days.⁴ The percentage increase in sudden deaths associated with particulate air pollution was approximately five times the percentage increase in all deaths. Particulate air pollution has been associated with daily deaths from cardiovascular disease in numerous studies over the last decade.5-7 Out of hospital deaths appeared to account for most of the increase.6 These cardiovascular responses have been shown to occur with little latency after exposure.⁸

One possible mechanism for this association is by altering autonomic control of the heart. Animal studies have shown that exposures to combustion particles can produce reductions in heart rate variability (HRV), a risk factor for sudden death⁹ and, in compromised animals, death from arrhythmia.¹⁰ Our previous study in Boston found a significant decrease in HRV associated with PM_{2.5} and ozone.¹¹ Similar decreases in HRV have been noted in a study of elderly subjects in Baltimore¹² and in a younger occupational cohort in Boston.¹³ The association in the workers persisted when restricted to days when they were off work,¹⁴ indicating that younger subjects are also sensitive to exposures to environmental particles.

The goals of this study were to replicate the results of the 1997 study, to examine the association with particles from different sources (especially traffic), to examine the shape of the concentration-response relation, to look for heterogeneity in response and identify its sources, and to examine other pollutants more fully.

METHODS

Study design and protocol

We recruited a panel of 28 subjects (aged 61-89 years) living at or near an apartment complex located on the same street as the Harvard School of Public Health, a few cross-streets away. A baseline questionnaire regarding medications, pulmonary and cardiac symptoms, and smoking history was administered and a resting 12-lead ECG was performed. Exclusion criteria included unstable angina, atrial flutter, atrial fibrillation, or paced rhythm. Each subject was given a day of the week and a time of day for testing, with the goal of being seen weekly for 12 weeks during the summer of 1999. Each week participants were administered a questionnaire regarding chest pain, medication changes, and whether medications had been taken that morning. Continuous Holter monitoring with electrodes in a modified V5 and AVF position were performed during a protocol similar to that of 1997¹¹ involving (1) 5 minutes of rest; (2) 5 minutes of standing; (3) 5 minutes of exercise outdoors (if the participant felt able, a standard walk was performed involving one climb up a slight incline); (4) 5 minutes of recovery; and (5) 3 minutes and 20 seconds of paced breathing. For 20 respiratory cycles the participant was asked to breathe in for 5 seconds and then out for 5 seconds, with a technician coaching. One subject only completed a single visit and was therefore excluded from the analysis.

Abbreviations: CO, carbon monoxide; HRV, heart rate variability; MI, myocardial infarction; NO₂, nitrogen dioxide; PM, particulate matter; SO_2 , sulphur dioxide

Processing of Holter recordings

A trained engineer reviewed and, when necessary, corrected automatically determined readings of QRS complexes. Regions of noise and artifact were eliminated. After correction, only normal-to-normal (NN) intervals between 150 and 5000 ms with NN ratios between 0.8 and 1.2 were included for analysis of heart rate variability. For the entire protocol duration we computed the standard deviation of normal RR intervals (SDNN), the root mean squared differences between adjacent RR intervals (r-MSSD), and the proportion of adjacent NN intervals differing by more than 50 ms. The low frequency/high frequency ratio (LFHFR), a frequency domain outcome reflecting the balance of the sympathetic versus parasympathetic components of the nervous system,¹⁵ was also computed.

Air pollution measurements

Air pollution measurements (PM2.5, black carbon, carbon monoxide (CO)) were collected at the Harvard School of Public Health within a kilometre of the residences of the subjects. Measurements of SO₂, ozone, and nitrogen dioxide (NO₂) were obtained from the State monitoring sites in Boston. Continuous PM2.5 was measured using the Model 1400A Tapered Element Oscillating Microbalance (TEOM) (Rupprecht and Patashnick, Albany, NY, USA). Since the TEOM sample filter is heated to 50°C, PM_{2.5} concentrations are corrected to compensate for the loss of semi-volatile mass that occurs at this temperature using a season specific calibration factor. The calibration factor was obtained by regressing 24 hour averaged TEOM concentrations on collocated integrated 24 hour Harvard Impactor Teflon filter gravimetric measurements. Black carbon was measured using a model AE-14 Aethalometer (Magee Scientific Inc, Berkeley, CA, USA) which has agreed well with EC in comparisons.16 17 Carbon monoxide (CO) was measured with a ThermoEnvironmental (Franklin, MA) Model 48 gas analyser using a US EPA reference method for CO.

In the summer in Boston, long range transported sulfate particles are a major component of PM_{2.5}. These particles are not black, and the black carbon measurements are surrogates for the primary (that is, directly emitted) carbonaceous particles, most of which derive from traffic. In Boston, hourly sulfate concentrations are not well correlated with hourly black carbon concentrations because they originate from different sources, sulfate particles being regional in nature while traffic particles are primarily emitted from local sources. As a result, sulfate and black carbon tend to have different diurnal profiles, especially in the summer months, with sulfate concentrations have a broad peak in the late morning to early afternoon while the highest levels of black carbon occur during morning and evening rush hours (Oh JA, doctoral dissertation, 2000). Other secondary particles (that is, those formed by reactions in the atmosphere from gaseous emissions), including nitrates and organic carbon also follow different temporal patterns than black carbon. This poor correlation allows us to estimate the concentrations of such secondary particles by regressing PM_{2.5} (which includes particles from all sources) against black carbon concentrations. The residuals represent hourly variations in particle mass concentration that is independent of both fresh traffic particles and transported black carbon, whose primary source is regional traffic. Thus, the residuals represent predominantly secondary particles independent of black carbon, of which sulfate particles are the largest components during the summer in Boston. We have used these residuals as a surrogate for secondary particle concentration.

Analysis of data

The data in our study existed on multiple levels. Data such as HRV were measured for each person, for each observation time. Other data such as medical characteristics of the subjects were fixed and did not change from visit to visit. Our analysis took into account these multiple levels.

HRV measurements were log transformed for normality. We examined HRV measurements for the entire visit (25– 30 minutes). This longer interval has less noise than the 5 minute intervals. Longer wavelength components also contribute a greater fraction of total HRV than in the 5 minute measurements. This allows us to distinguish whether the low frequency to high frequency ratio is disturbed by air pollution.

Each subject served as his or her own control. This was accomplished by including dummy variables for each subject in the regression models. These controlled for all time invariant subject specific risk factors such as baseline medical condition, smoking history, etc, *whether measured or not*. This is a major strength of this study design. The regression also included dummy variables for day of week and hour of day, which controlled for diurnal and weekly variations in autonomic function or air pollution, indicator variables for whether the subjects had taken their medication before the examination, and a regression spline (3 degrees of freedom) for temperature. A time trend variable was also included.

While these models control for subject specific characteristics, they do not account for possible heterogeneity of response to air pollution. To address this question we used hierarchical models. A hierarchical model is simply a model with multiple levels of analysis. Specifically, we assumed:

 $Y_{it} = b_0 + u_i + covariate model + (b_1 + v_i) Pollution_t + e_{it}$

where Y_{it} is the logarithm of HRV in subject *i* at time *t*, b_0 is the overall intercept, and u_i is the separate intercept for subject *i*. For the pollution model b_1 is the slope representing the overall effect of pollution, and v_i captures the variation of the slope for subject *i* about that overall mean. This gives us the opportunity to examine the distribution of subject specific slopes to see, for example, whether any adverse effect is primarily concentrated in a few susceptible individuals or whether pollution has an adverse impact on most or all subjects. In this model we treat the v_i as random effects.

The second level of the model is only at the level of the subject. Here we examine whether the variability in susceptibility of individual subjects may be explained by some factor Z_i which may represent a diagnosis (previous myocardial infarction), medication use (beta blockers), etc. This model takes the form:

$$\mathbf{E}(v_i) = \mathbf{d}_i + \mathbf{h}_i \mathbf{Z}_i$$

where h_i is the coefficient of the susceptibility factor Z_i and d_i represents the remaining subject specific intercept for subject *i* which is not explained by Z_i . This is formally identical to having an interaction term between the susceptibility factor Z and the exposure in the model, but recognises that this may not explain all heterogeneity in slope (reflected in d_i). This multilevel model was estimated using the linear mixed effect function in Splus. The possible effect modifiers were whether the subject had a diagnosis of a myocardial infarction, congestive heart failure or chronic obstructive pulmonary disease, whether their measured systolic blood pressure was greater than 140 mm Hg, and whether they were on β blockers.

Our strategy was first to try to replicate our original finding of an association between $PM_{2.5}$ and ozone with r-MSSD and a weaker association with SDNN. We then turned to a fuller

Variable	25%	50%	75%
Continuous variables			
Age	64	74	81
SDNN (ms)	77	101	132
r-MSSD (ms)	18	29	41
PNN ₅₀	0.9	3.8	11
Heart rate (bpm)	70	77	84
LFHFR	1.38	1.97	2.86
Environmental variables			
Temperature (°C)*	69	72	77
PM _{2.5} (μg/m ³)*	7	10	17
$PM_{2.5} (\mu g/m^3)^{\dagger}$	6	10	19
Black carbon (µg/m³)†	0.9	1.2	1.6
Black carbon (µg/m ³)*	0.8	1.0	1.3
Secondary particles (estimated)	- 4.5	- 1.7	3.5
(µg/m ³)†			
Ozone (ppb)†	20	34	46
NO ₂ (ppb)*	14	18	23
SO ₂ (ppm)*	0.017	0.020	0.54
CO (ppm)*	0.38	0.45	0.54
Proportions			
Female	21/28		
Calcium channel blocker	3/28		
β blocker	5/28		
ACE inhibitor	8/28		
Previous MI	3/28		
Ever CHF	2/28		
COPD	2/28		
Systolic BP >140	11/28		

*24 hour average

†1 hour average

range of HRV measures and measurements of different particle components to test the hypothesis that traffic particles are more important, and finally to inclusion of other air pollutants.

Our previous study found that ozone was associated with HRV using the immediate hour's exposure. The $PM_{2.5}$ association was strongest with the mean concentration in the preceding 4 hours, but also with the preceding 24 hour average. In this study we therefore examined those three time frames: pollution averaged over the preceding hour, the preceding 4 hours, and the preceding 24 hours before each person's HRV measurement.

RESULTS

Table 1 shows the distribution of the personal characteristics of the participants and of the environmental variables in the study. The median age was 74 and participants were predominantly female. Eleven participants completed all 12 visits, 13 completed 11 visits, three completed 10 visits, and one completed five visits. The correlation coefficients among the exposure variables are shown in table 2.

Particle model

As expected, peak periods of $PM_{2.5}$ and black carbon did not occur at the same time. Figure 1 shows a plot of the black carbon versus $PM_{2.5}$ data used in our study. While at low levels there is correlation between the two measures, the peak periods for each pollutant are distinct, enabling us to distinguish between carbonaceous and non-carbonaceous particles. Black carbon explained 30% of the variation in $PM_{2.5}$ concentration, and the residuals from that model were used as a surrogate for non-carbonaceous secondary particles.

Replication

We were able partially to replicate our 1997 study. We found an association between the mean of the previous 24 hour $PM_{2.5}$ (percentage change for an interquartile range (IQR) increase in exposure -10.1% (95% CI -2.8 to -16.9)) and the previous hour's ozone (percentage change for an IQR increase of exposure -8.5% (95% CI 0.3 to -16.6)) with r-MSSD, although the association with ozone was substantially weaker than in the previous study.¹¹ In this study the $PM_{2.5}$ association was stronger with the 24 hour average than

	Black					
Variable	carbon	PM _{2.5}	Ozone	со	SO ₂	NO ₂
Black carbon	1					
PM _{2.5}	0.37	1				
Ozone	- 0.03	0.62	1			
CO	0.83	0.61	0.21	1		
SO ₂	- 0.22	0.10	0.13	- 0.18	1	
NO ₂	0.54	0.29	0.02	0.55	- 0.28	1

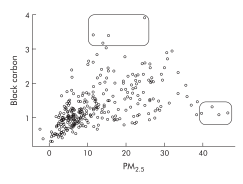


Figure 1 Scatter plot of hourly black carbon versus $PM_{2.5}$ measurements in Boston during the time when HRV measurements were being taken. The boxes highlight the four highest black carbon and $PM_{2.5}$ measurements, respectively, and they each occur at unexceptional levels of other particle measurements.

with the 4 hour average, although the 4 hour average was also associated with decreased r-MSSD (percentage change -8.6% (95% CI -0.1 to -16.3)).

Particle effects

The associations with the three particle measures (fit separately) are shown in table 3. In general we observed a gradient of responses—that is, for $PM_{2.5}$ and black carbon the magnitude of the percentage change in HRV outcome for a given change in exposure was $PNN_{50} > r-MSSD > SDNN$ for both averaging periods. For secondary sulfate particles the magnitude and precision of the effect estimates for effects on HRV were lower than for all fine particles or for the carbonaceous particles, particularly for SDNN. The same pattern of larger effects for PNN₅₀ and lower effects for SDNN was seen.

Only black carbon (percentage change -4.6% (95% CI -2.0 to -7.2)) showed a convincing association with SDNN. Black carbon was also the only particle measure to show a significant effect on the ratio of low frequency to high frequency components of HRV. Figures 2 and 3 further illustrate the pattern of association of particle measures with SDNN. Figure 2 shows the estimated relation between black carbon and SDNN using a loess smooth with about 3 degrees of freedom. The association is quite linear across the range of exposures in the study. In contrast, fig 3 shows the same relation using PM_{2.5} as the exposure metric. For low concentrations, when the correlation between PM_{2.5} and black carbon was good (as in fig 1), there is again a roughly linear association with SDNN. At high doses of PM_{2.5} which are uncoupled from black carbon, the association flattens out. On the other hand, the associations between PM2.5 and r-MSSD and PNN₅₀ were similar in magnitude to the association with black carbon, although somewhat less precise. The magnitude of effect for secondary particles was

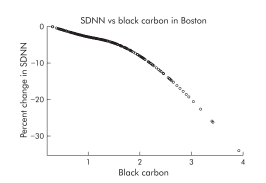


Figure 2 Smoothed plot of the percentage deviation from predicted SDNN (based on the model with all other covariates) versus black carbon concentrations in Boston. The association is nearly linear, despite being fit with about 3 degrees of freedom.

smaller, suggesting that $PM_{2.5}$, which includes fresh traffic particles, is acting in part as a surrogate for the black carbon effect in these relationships.

For all relationships except that between SDNN and black carbon the strongest associations are with the mean of the preceding 24 hours of pollution. In that case, the association with the current hour's exposure was slightly stronger, although not much different from the effect of 24 hour exposure. The most consistent association is therefore seen with the 24 hour exposures.

Other pollutants

Table 4 shows the associations of gaseous air pollutants with HRV. As noted above, a weak association of immediate ozone exposure and r-MSSD was seen, although no associations with other measures of HRV were apparent. This pattern was similar to the pattern for non-traffic secondary particles. No evidence was seen for an association between SO₂, NO₂, and any of the outcomes. CO, in contrast, showed an association, but only with the 24 hour average. The pattern of association was similar to black carbon; there were associations with SDNN and the LFHFR as well as with r-MSSD and PNN₅₀.

Interactions and heterogeneity

For simplicity, we only examined interactions for black carbon, which showed the strongest and most consistent pattern of association. We began by fitting random effects models with random slopes for each subject. The estimated overall effect of black carbon was little changed from the fixed effects models (percentage change for an IQR increase in black carbon -4.0% (95% CI -1.2 to -6.7)). There was evidence of heterogeneity in the effect of black carbon. Figure 4 shows the distribution of the random subject specific effect size estimates relating black carbon to SDNN. While all of the subject specific effects were negative, they had a considerable range and some evidence of skewness in the distribution. This was examined more formally using

Variable	SDNN (ms)	r-MSSD (ms)	PNN ₅₀ (%)	LFHFR
BC*	- 4.6 (-2.0 to -7.2)	- 6.1 (0 to -11.9)	- 9.4 (-0.1 to -17.8)	1.8 (-3.0 to 7.0)
BC†	- 5.1 (-1.5 to -8.6)	- 10.1 (-2.4 to -17.2)	- 16.9 (-6.0 to -26.6)	7.2 (.7 to 14.1)
PM _{2 5} *	- 3.4 (0.6 to -7.3)	- 7.4 (1.6 to -15.5)	- 12.9 (0 to -24.2)	1.4 (-5.5 to 8.8)
PM2.5†	- 2.6 (0.8 to -6.0)	- 10.1 (-2.8 to -16.9)	- 12.7 (-1.6 to -22.5)	0.2 (-4.6 to 5.2)
Secondary PM*	- 0.5 (2.8 to -3.7)	- 3.0 (4.8 to -10.3)	- 6.6 (-16.9 to 5.0)	1.4(-4.1 to 7.2)
Secondary PM ⁺	-0.9(2.5 to -4.1)	- 6.4 (0.6 to -12.9)	- 5.8 (-15.6 to 5.1)	-2.4(-7.6 to 3.1)

All models control for subject, temperature, day of week, hour of day, medication use on that day, and time trend. *1 hour average.

†24 hour average

‡Residuals of PM_{2.5} controlling for black carbon (BC).

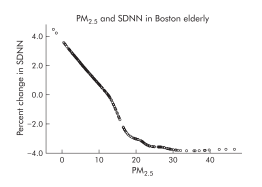


Figure 3 Smoothed plot of the percentage deviation from predicted SDNN (based on the model with all other covariates) versus $PM_{2.5}$ concentrations in Boston. The association flattens out at high concentrations where the correlation between $PM_{2.5}$ and black carbon disappears.

interaction terms for the subject characteristics thought to be potential effect modifiers. We found that a previous MI was associated with a greater decrement in SDNN for an IQR increase in black carbon exposure. For subjects without a MI, an IQR increase in black carbon was associated with a 3.5% decrease in SDNN (95% CI 0.7 to 6.3). For subjects with a prior MI, the effect size was almost four times as great (12.7% decrease, 95% CI 5.7 to 19.25). The other subject characteristics examined (COPD, heart failure, high blood pressure, use of β blockers) did not seem to modify the effect of black carbon.

Two pollutant models

Having found HRV associations with both particles and CO, we examined black carbon and CO in two pollutant models. Black carbon (but not CO) remained significant for two of the HRV measures (SDNN and LFHFR) and neither pollutant remained significant for the other two measures (table 5).

DISCUSSION

Increased particle levels were found to be related to reduced HRV. Moreover, a surrogate for traffic particle exposure (black carbon) had the largest effect on reduced HRV, with secondary particles (primarily sulfates) associated with lower and different effects. The secondary particles were marginally associated with changes in r-MSSD, with effects sizes similar in magnitude to those of black carbon. However, no effect was seen on SDNN or on the LFHFR. The carbonaceous particles, in contrast, increased the LFHFR, suggesting an increase in sympathetic compared with parasympathetic tone. The relation of HRV to cardiac mortality is thought to work through sympathetic predominance and/or diminished parasympathetic tone.

Distribution of effect sizes of black carbon on SDNN

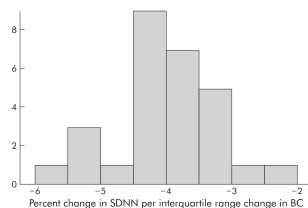


Figure 4 Histogram of the subject specific covariate adjusted effect sizes relating black carbon to SDNN in Boston. All of the slopes are negative, but they vary in magnitude by a factor of 3.

The poorer association of $PM_{2.5}$ with SDNN suggests that the separation between high BC days and high $PM_{2.5}$ days in fig 1 is responsible. The high BC days on average had lower wind speed, ozone, and temperature, but higher CO concentrations than the high $PM_{2.5}$ days. These are all conditions that would be expected to result in lower concentrations of transported photochemical particles.

Laden and co-workers, in a study of six US cities, found that traffic particles (identified in a source apportionment model using elemental composition) were more strongly associated with cardiovascular deaths than particles from coal burning.¹⁸ Janssen and colleagues recently reported that the magnitude of the effect of $10 \ \mu g/m^3 \ PM_{10}$ on hospital admissions for heart disease varied with the proportion of the particles from traffic.¹⁹ Our study and others support the conclusion that the effects of primary traffic particles on HRV may be larger and more significant than the effects of sulfates and other secondary particles.

On the other hand, given the considerably larger degree of measurement error in the estimated secondary particle term, the finding of an association with HRV is noteworthy. While this may reflect the correlation between secondary particle concentrations and ozone, ambient concentrations of sulfates have a much stronger association with personal exposure to sulfates than is true for ambient and personal ozone, suggesting that a real effect of secondary particles is plausible. The greater measurement error may have affected the associations with different outcomes differently (e.g. LFHFR v r-MSSD), raising some caution in conclusions about different mechanisms. Further work with better measures of secondary particles will be necessary to confirm this.

 Table 4
 Percentage change in heart rate variability associated with an interquartile range change in gaseous pollution exposure

Variable	SDNN (ms)	RMSSD (ms)	PNN ₅₀ (%)	LFHFR
O ₃ †	- 3.1 (-7.0 to 0.9)	- 8.5 (-16.6 to 0.3)	- 6.5 (-18.9 to 7.8)	5.7 (-1.5 to -13.4)
O3*	- 1.5 (-5.7 to 2.9)	- 2.3 (-11.6 to 7.9)	0.9 (-13.4 to 17.6)	- 4.8 (-11.8 to 2.7)
SO ₂ †	0.4 (-1.3 to 2.1)	1.4 (-2.6 to 5.5)	3.8 (-12.1 to 22.5)	- 1.2 (-4.2 to 1.8)
SO ₂ *	0.4 (-4.2 to 5.1)	- 0.3 (-1.3 to 0.8)	- 0.2 (20.9 to 17.6)	2.9 (-4.9 to 11.4)
NO_2^{\dagger}	- 0.4 (2.1 to -2.9)	- 0.9 (-6.3 to 4.9)	- 1.2 (-9.5 to 7.9)	1.8 (-2.6 to 6.3)
NO ₂ *	- 0.7 (-3.5 to 2.3)	- 2.3 (-8.7 to 4.4)	- 5.5 (-14.7 to 4.6)	5.0 (-0.2 to 10.5)
COT	- 2.6 (-5.6 to 0.5)	- 3.9 (-10.6 to 3.3)	- 3.5 (-13.7 to 8.0)	4.5 (-1.2 to 10.5)
CO*	- 4.2 (-0.6 to -7.7)	-10.2(-2.4 to -17.4)	- 14.8 (-3.0 to -25.2)	6.2 (-0.6 to 13.4)

All models control for subject, temperature, day of week, hour of day, medication use on that day, and time trend. *24 hour average.

t1 hour average

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 Table 5
 Two pollutant models with black carbon (BC) and carbon monoxide (CO) showing percentage change (95% CI) for interquartile range changes in each pollutant

	Pollutant	%	95% CI
SDNN	BC	- 4.3	(−7.4 to −1.1)
	CO	- 1.3	(-5.5 to 3.0)
r-MSSD	BC	- 0.5	(-17.1 to 19.4)
	CO	- 9.9	(7.1 to -24.3)
PNN ₅₀	BC	- 18.1	(-38.1 to 8.5)
	CO	0.4	(-23.1 to 31)
FHFR	BC	15.6	(0.4 to 33.1)
	CO	- 5.8	(-17.6 to 7.7)

The random slope model is a key feature of this study. It shows that the negative effects of traffic particles are not restricted to a few subjects. Rather, the slope of the association is negative for all participants. It also indicates that there is variation among subjects in those slopes, suggesting that important factors affecting susceptibility remain to be identified. The identification of subjects with MI as being at differential risk from air pollution suggests that ischaemic heart disease is an important predisposing condition for the effect of traffic particles on HRV. This finding, while statistically significant, rests on the difference between three subjects with prior MI and the other 25 subjects, and therefore must be treated with caution. We have previously found that prior diagnosis of conduction disorders or dysrhythmias increased the risk of particle associated hospital admissions for cardiovascular disease.²⁰ It is possible that local traffic exposure may increase the risk of cardiac hospital admissions through altering cardiac autonomic function, as reflected in reduced HRV.

Several studies have now shown that increased ambient particle levels predict reduced HRV.^{11 12 14 21} Between-person reduced HRV as a subject characteristic is a known predictor of poor cardiac outcomes and sudden death.^{22 23} However, a recent study of ischaemic events showed that they were preceded by decreases in high frequency HRV in the hour before the event and in low frequency HRV in the 4 minutes before the event.²⁴ Similarly, a significant decrease in HRV has been reported to precede paroxysmal atrial fibrillation.²⁵ Hence, short term changes in HRV are of interest.

A limitation of the study is the use of a single nearby ambient monitoring station to measure air pollution instead of individual level exposures. Many studies have shown that longitudinal changes in outdoor concentrations of PM_{2.5} are good predictors of longitudinal changes in personal exposure to PM_{2.5} of outdoor origin.^{26–29} This is not the case for some of the gaseous air pollutants. In a Baltimore study of elderly subjects similar to ours, Sarnat et al³⁰ confirmed that outdoor PM_{2.5} was a good surrogate for exposure to PM_{2.5}, but found poor associations between temporal fluctuations in outdoor concentrations of SO₂, ozone, or NO₂ and personal exposure to those pollutants. The outdoor concentrations of all three gases were better correlated with personal exposure to PM_{2.5}. Ozone was a better surrogate for exposure to sulfate particles. A more recent analysis of a Boston panel of elderly subjects found similar results.²⁸ These results are not surprising. SO₂ and ozone are very reactive gases, and their reactivity makes indoor concentrations much lower than outdoor concentrations. The temporal variability in exposure can therefore be dominated by temporal variability in how much time is spent outdoors. Ozone is produced by similar photochemical reactions that produce sulfate particles and is also a long range transported pollutant. It is therefore highly correlated with secondary transported particles in Boston and, indeed, with personal exposure to sulfates. This may be why the

pattern of ozone association with HRV in our study is so similar to the pattern seen with non-traffic particles. Given the observed relations between outdoor concentrations and exposure in Boston, ozone may represent a secondary particle effect and not a true ozone effect in this study. This does not mean that ozone does not affect autonomic function or have other cardiovascular effects, merely that studies that do not use personal exposure measurements are less able to assess that question.

This study suggests the predominant importance of traffic exposure in its influence on cardiac autonomic function. Further investigation is needed regarding the components of traffic particles that may be responsible for cardiac autonomic function and subsequent arrhythmias and acute coronary artery syndromes. The indication of an association between a noisy surrogate for sulfate particles and r-MSSD also deserves further investigation, as those particles may affect autonomic control in a different way from traffic particles.

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REFERENCES

- Her Majesty's Public Health Service. Mortality and morbidity during the London fog of December 1952. London: Her Majesty's Public Health Service, 1954.
- 2 Firket J. Fog along the Meuse Valley. Trans Faraday Soc 1936;32:1192-7.
- 3 Shrenk H, Heimann H, Clayton GD. Air pollution in Donora PA: epidemiology of the unusual smag episode of October 1948. Washington, DC: US Public Health Service, 1949.
- 4 Schwartz J. What are people dying of on high air pollution days? Environ Res 1994;64:26-35.
- 5 Bremner SA, Anderson HR, Atkinson RW, et al. Short-term associations between outdoor air pollution and mortality in London 1992–4. Occup Environ Med 1999;56:237–44.
- 6 Schwartz J. Assessing confounding, effect modification, and thresholds in the association between ambient particles and daily deaths. *Environ Health Perspect* 2000;108:563–8.
- 7 Katsouyanni K, Touloumi G, Samoli E, et al. Confounding and effect modification in the short-term effects of ambient particles on total mortality: results from 29 European cities within the APHEA2 project. Epidemiology 2001;12:521–31.
- 8 Braga AL, Zanobetti A, Schwartz J. The lag structure between particulate air pollution and respiratory and cardiovascular deaths in 10 US cities. J Occup Environ Med 2001;43:927–33.
- 9 Godleski JJ, Verrier RL, Koutrakis P, et al. Mechanisms of morbidity and mortality from exposure to ambient particles. Res Rep Health Eff Inst 2000;91:5–88.
- 10 Watkinson WP, Campen MJ, Costa DL. Cardiac arrhythmia after exposure to residual oil fly ash particles in a rodent model of pulmonary hypertension. *Toxicol Sci* 1998;41:209–16.
- 11 Gold DR, Litonjua A, Schwartz J, *et al.* Ambient pollution and heart rate variability. *Circulation* 2000;101:1267–73.
- 12 Liao D, Creason J, Shy C, et al. Daily variation of particulate air pollution and poor cardiac autonomic control in the elderly. Environ Health Perspect 1999;107:521–5.
- 13 Magari SR, Hauser R, Schwartz J, et al. Association of heart rate variability with occupational and environmental exposure to particulate air pollution. *Circulation* 2001;104:986–91.
- 14 Magari SR, Schwartz J, Williams PL, et al. The association between personal measurements of environmental exposure to particulates and heart rate variability. *Epidemiology* 2002;13:305–10.
- 15 Task Force of the European Society of Cardiology and the North American Society of Pacing and Electrophysiology. Heart rate variability: standards of measurement, physiological interpretation and clinical use. *Circulation* 1996;**93**:1043–65.
- 16 Allen GAKP, Lawrence J. Field validation of a real-time method for aerosol black carbon (Aethalometer) and temporal patterns of summertime hourly black carbon measurements in southwestern Pennsylvania. Atmos Environ 1999;33:817–23.

- 17 Hansen ADA, Rosen H. Vertical distributions of particulate carbon, sulfur, and bromine in the Arctic haze and comparison with ground-level measurements at Barrow, Alaska. Geophys Res Lett 1984;11:381-4.
- 18 Laden F, Neas LM, Dockery DW, et al. Association of fine particulate matter from different sources with daily mortality in six U.S. cities. Environ Health Perspect 2000;108:941–7.
- 19 Janssen NA, Schwartz J, Zanobetti A, et al. Air conditioning and source-specific particles as modifiers of the effect of PM(10) on hospital admissions for heart and lung disease. Environ Health Perspect 2002;110:43–9.
- 20 Zanobetti A, Schwartz J, Gold D. Are there sensitive subgroups for the effects of airborne particles? *Environ Health Perspect* 2000;108:841–5.
- 21 Pope CA 3rd, Verrier RL, Lovett EG, et al. eart rate variability associated with particulate air pollution. Am Heart J 1999; 138:890–9.
- 22 Tsuji H, Larson MG, Venditti FJ Jr, et al. Impact of reduced heart rate variability on risk for cardiac events. The Framingham Heart Study. *Circulation* 1996;**94**:2850–5.
- 23 Odemuyiwa O, Malik M, Farrell T, et al. Comparison of the predictive characteristics of heart rate variability index and left ventricular ejection fraction for all-cause mortality, arrhythmic events and sudden death after acute myocardial infarction. Am J Cardiol 1991;68:434–9.

- 24 Kop W, Verdino R, Gottdiener J, et al. Changes in heart rate and heart rate variability before ambulatory ischemic events. J Am Coll Cardiol 2002;39:744–5.
- 25 Bettoni M, Zimmermann M. Autonomic tone variations before the onset of paroxysmal atrial fibrillation. *Circulation* 2003;105:2753–9.
- 26 Oglesby L, Kunzli N, Roosli M, et al. Validity of ambient levels of fine particles as surrogate for personal exposure to outdoor air pollution: results of the European EXPOLIS-EAS Study (Swiss Center Basel). J Air Waste Manag Assoc 2000;50:1251–61.
- 27 Sarnat JA, Koutrakis P, Suh HH. Assessing the relationship between personal particulate and gaseous exposures of senior citizens living in Baltimore, MD. J Air Waste Manag Assoc 2000;50:1184–98.
- Koutrakis P, Suh HH, Sarnat JA, et al. Characterization of particulate and gaseous exposures of sensitive sub-populations living in eastern US metropolitan areas. Final Report. Boston, MA: Health Effects. Institute, 2002.
 Jansen NA, de Hartog JJ, Hoek G, et al. Personal exposure to fine particulate
- 29 Janssen NA, de Hartog JJ, Hoek G, et al. Personal exposure to fine particulate matter in elderly subjects: relation between personal, indoor, and outdoor concentrations. J Air Waste Manag Assoc 2000;50:1133–43.
- 30 Sarnat JA, Schwartz J, Catalano PJ, et al. Gaseous pollutants in particulate matter epidemiology: confounders or surrogates? Environ Health Perspect 2001;109:1053–61.

LUNG ALERT

Chest wall tenderness does not exclude pulmonary embolism

▲ Le Gal G, Testuz A, Righini M, *et al.* Reproduction of chest pain by palpation: diagnostic accuracy in suspected pulmonary embolism. *BMJ* 2005;**330**:452–3

Pulmonary embolism (PE) is usually associated with pleuritic chest pain, and the presence of chest pain that is reproducible by palpation may lead a clinician to discard this diagnosis. This study assessed whether palpable chest pain predicted the likely absence of a PE in individuals in whom the diagnosis was suspected (acute shortness of breath or chest pain without another obvious aetiology).

Patients (n = 965) were recruited from the database of another study looking at diagnosis of PE. Predictive variables of a PE (the Geneva score) were recorded and whether the pain was reproducible by palpation. A PE was ruled out if the p-dimer concentration was <500 µg/l or if proximal venous ultrasonography and helical computed tomography were both negative. If clinical probability was high, a negative pulmonary angiogram was also required. The overall prevalence of PE was 23%. This was not significantly lower in patients with pain reproducible by palpation (19.9% v 23.8%, p = 0.25). 17% of patients with a PE had palpable chest pain.

In these patients with a suspected PE, chest pain reproducible by palpation was not associated with a lower prevalence of an embolism. Physicians must be cautious not to discount this diagnosis in patients who present with this atypical feature.

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