



2004 Particulate Summary

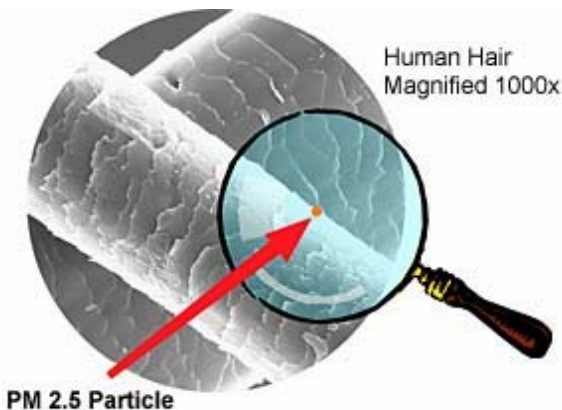
New Jersey Department of Environmental Protection

NATURE AND SOURCES

Particulate air pollution consists of both solid particles and liquid droplets suspended in the atmosphere. Suspended particles can range in size from 70 microns in diameter, approximately the size of a pinhead, to less than 1 micron in diameter. Particles can be directly emitted, or they can form in the atmosphere from gaseous emissions, such as sulfur dioxide (SO₂) and oxides of nitrogen (NO_x). Particles that originate as gases are referred to as secondary particulates.

Particulate matter is generally categorized according to the size of the particles. Coarse particles are defined as particles greater than 2.5 microns in diameter, while particles less than 2.5 microns in diameter are referred to as fine particles (PM_{2.5}) (See Figure 1). Coarse particles are further subdivided into Total Suspended Particulates (TSP), which include all but the largest particles, and PM₁₀, which include particles less than 10 microns in diameter. The human respiratory tract will usually trap particles above about 10 microns in diameter before they reach the lungs. Particles smaller than 10 microns (PM₁₀) are inhalable and are considered to be more harmful to human health than larger particles; fine particles are considered to be even more harmful as they can reach the deep recesses of the lungs.

Figure 1
Size of PM_{2.5} Particle Compared to a Human Hair



Graphics Courtesy of the US Department of Energy

Both fine and coarse particles have anthropogenic, or man-made, as well as natural sources. Anthropogenic sources of coarse particles include industrial processes such as grinding operations, while anthropogenic sources of fine particles include soot from fuel combustion, and secondary particle formation from organic compounds, biomass burning, and emissions of sulfur dioxide (SO₂) and oxides of nitrogen (NO_x). Natural sources of coarse particles include windblown dust, sea salt, and biological debris; and natural sources of fine particles include biogenic gases, which result in the formation of secondary particles.

ENVIRONMENTAL EFFECTS

In addition to health effects, particulate matter is the major cause of reduced visibility in many parts of the United States. Figure 2 provides an example of reduced visibility recorded by our WebCam site in Newark (accessible via the Internet at www.state.nj.us/dep/airmon). Airborne particles can also impact vegetation and aquatic ecosystems, and can cause damage to paints and building materials. More information is provided in the Regional Haze section of this report.

Figure 2
Visibility WebCam



View of New York City Skyline from Newark

HEALTH EFFECTS

Inhalable particles (PM₁₀) and especially fine particles (PM_{2.5}) are a health concern because they easily reach the deepest recesses of the lungs. Various health problems are associated with both long and short-term exposures. When inhaled, these particles can accumulate in the respiratory system and are associated with increased hospital admissions and emergency room visits for heart and lung conditions, such as asthma, bronchitis, cardiac arrhythmias, heart attacks, and even premature death. Groups that appear to be at the greatest risk from particulates include children, the elderly, and individuals with heart and lung diseases, such as asthma (*US EPA, 2001*).

STANDARDS

In 1971, EPA set primary (health based) and secondary (welfare based) standards for total suspended particulate matter (TSP). These standards, known as the National Ambient Air Quality Standards (NAAQS), were based on maximum 24-hour and annual concentrations (*US EPA, 1997*). The annual standards were based on the geometric mean concentrations over a calendar year, and the 24-hour standards were based on the arithmetic average concentration from midnight to midnight. The primary 24-hour average standard for TSP was set at 260 micrograms per cubic meter (µg/m³) and the annual geometric mean health standard was set at 75 µg/m³. The 24-hour secondary standard was set at 150 µg/m³. While EPA did

not establish a secondary annual standard for TSP they did set a guideline of 60 µg/m³ to be used to ensure that the secondary 24-hour standard was being met throughout the year. Although New Jersey still maintains state standards for TSP, the national standards have been replaced with standards for smaller particles as described below. As a result, monitoring for TSP has largely been discontinued, with the exception of one station, where TSP samples are taken to analyze for lead (Pb). See the Lead Summary section for more details.

In 1987, EPA replaced the TSP standards with standards that focused only on inhalable particles. Inhalable particles are defined as particles less than 10 microns in diameter (PM₁₀). The 24-hour PM₁₀ primary and secondary standards were set at 150 µg/m³, and the annual primary and secondary standards were set at 50 µg/m³. The annual standard for PM₁₀ is based on the arithmetic mean, as opposed to the geometric mean that was used for TSP.

In 1997, EPA promulgated new standards for fine particulates, which it defined as particles less than 2.5 microns in diameter (PM_{2.5}). They kept the existing standards for PM₁₀ as well. The PM_{2.5} annual primary and secondary standards were set at 15 µg/m³ and the 24-hour standard was set at 65 µg/m³. Table 1 provides a summary of the Particulate Matter standards.

**Table 1
National and New Jersey
Ambient Air Quality Standards for Particulate Matter**

Micrograms Per Cubic Meter (µg/m³)

Standard	Averaging Period	Type	New Jersey	National
Total Suspended Particulates (TSP)	12-Month [‡]	Primary	75 µg/m ³	---
	24-Hour	Primary	260 µg/m ³	---
	12-Month [†]	Secondary	60 µg/m ³	---
	24-Hour	Secondary	150 µg/m ³	---
Inhalable Particulates (PM ₁₀)	Annual [†]	Primary & Secondary	---	50 µg/m ³
	24-Hour Average	Primary & Secondary	---	150 µg/m ³
Fine Particulates (PM _{2.5})	Annual [†]	Primary & Secondary	----	15 µg/m ³
	24-Hour Average	Primary & Secondary	----	65 µg/m ³

[‡] Annual Geometric Mean

[†] Annual Arithmetic Mean

PARTICULATE MONITORING NETWORK

New Jersey's Particulate Monitoring Network consists of 19 fine particulate monitoring sites, 6 PM₁₀ monitoring sites, 1 TSP monitoring site, and 10 sites where smoke shade is monitored.

At some of these sites, samplers that comply with strict EPA specifications are used for collecting data that are submitted to a national database maintained by the EPA. These filter-based samplers, which are approved by the EPA and known as Federal Reference Method (FRM) samplers, collect particles on a filter over a 24-hour period. The filters are subsequently weighed under controlled environmental conditions. The data from the FRM samplers are used by the NJDEP and EPA to determine whether the state, or portions of the state, meet the federal health and welfare standards for particulate matter. Because the FRM samplers do not provide data in real time, the NJDEP employs additional samplers that continuously measure particulate concentrations. These samplers are used by the NJDEP

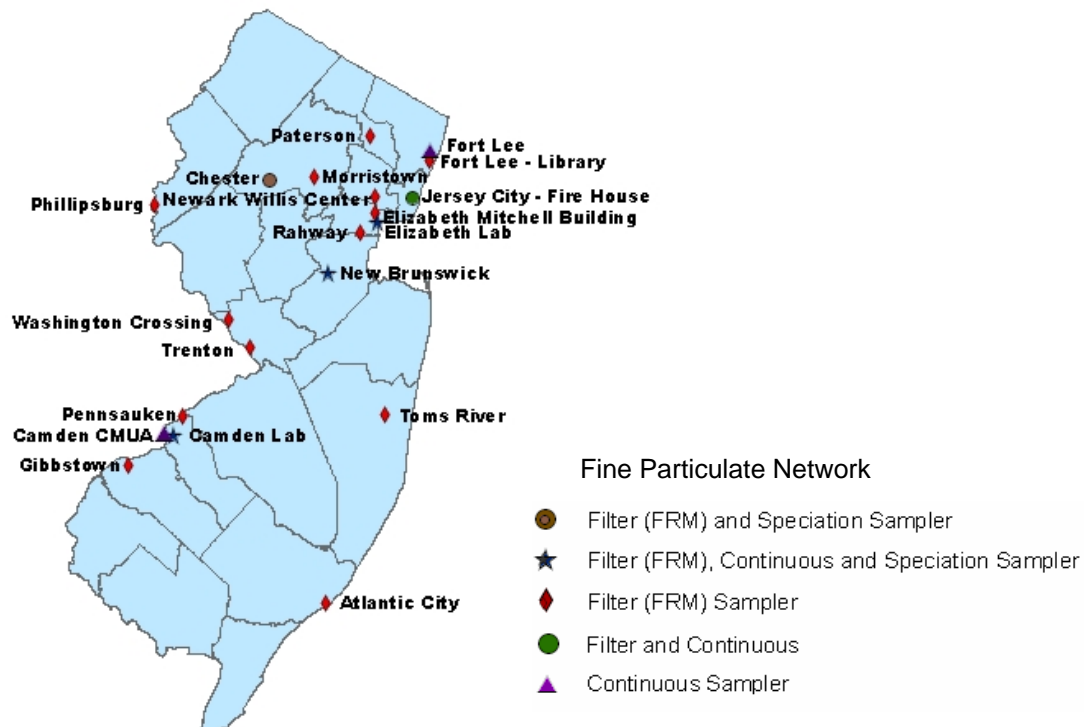
to report current air quality to the public through the Air Quality Index (www.state.nj.us/dep/airmon). The NJDEP uses Tapered Element Oscillating Microbalance (TEOM) analyzers and smoke shade instruments for real-time particle reporting. The TEOM analyzers collect a sample of fine particles on an oscillating filter, and determine the concentration based on the change in the frequency at which the filter oscillates. Smoke shade instruments collect a sample of particles on a paper tape for one hour. At the end of each hour the amount of light that will pass through the spot that has formed on the tape is measured, the tape advanced, and the cycle started over. The amount of light transmittance measured is used as an estimate of actual particle concentrations.

FINE PARTICLE SUMMARY

FINE PARTICLE MONITORING SITES

There are 18 monitoring sites in New Jersey where a filter-based (FRM) sampler routinely collects PM_{2.5} 24-hour samples (see Figure 3). At 6 sites, continuous

**Figure 3
2004 PM_{2.5}
Monitoring Network**



particulate monitors measure the concentration of fine particles every minute and transmit the data to the Bureau of Air Monitoring's central computer, where it is made available on the Bureau's Public Website (www.state.nj.us/dep/airmon). Additionally, at four of these locations a separate sampler collects fine particles on three types of filter media which are subsequently analyzed using ion chromatography (IC), X-ray fluorescence (XRF), and Thermal Optical Analysis (TOA) to determine the concentrations of the chemical analytes that constitute the sample.

FINE PARTICLE CONCENTRATION SUMMARY

The annual mean concentration of PM_{2.5} ranged from 10.2 µg/m³ in Chester to 15.2 µg/m³ at Elizabeth Lab. The maximum 24-hour concentrations ranged from 32.8 µg/m³ at Chester to 56.4 µg/m³ at Elizabeth Lab.

Figure 4 and Table 2 depicts the mean and maximum concentrations at each site.

None of the sites exceeded the 24-hour standard of 65 µg/m³. One site, Elizabeth Lab exceeded the annual standard of 15 µg/m³. **The annual mean concentration at Elizabeth Lab was 15.2 µg/m³.** Three years of data are required to determine compliance with the NAAQS for PM_{2.5}. NJDEP will be evaluating PM_{2.5} data collected to date in making its final determination as to whether the annual NAAQS are being met.

Figure 4
2004 Fine Particulate (PM^{2.5})

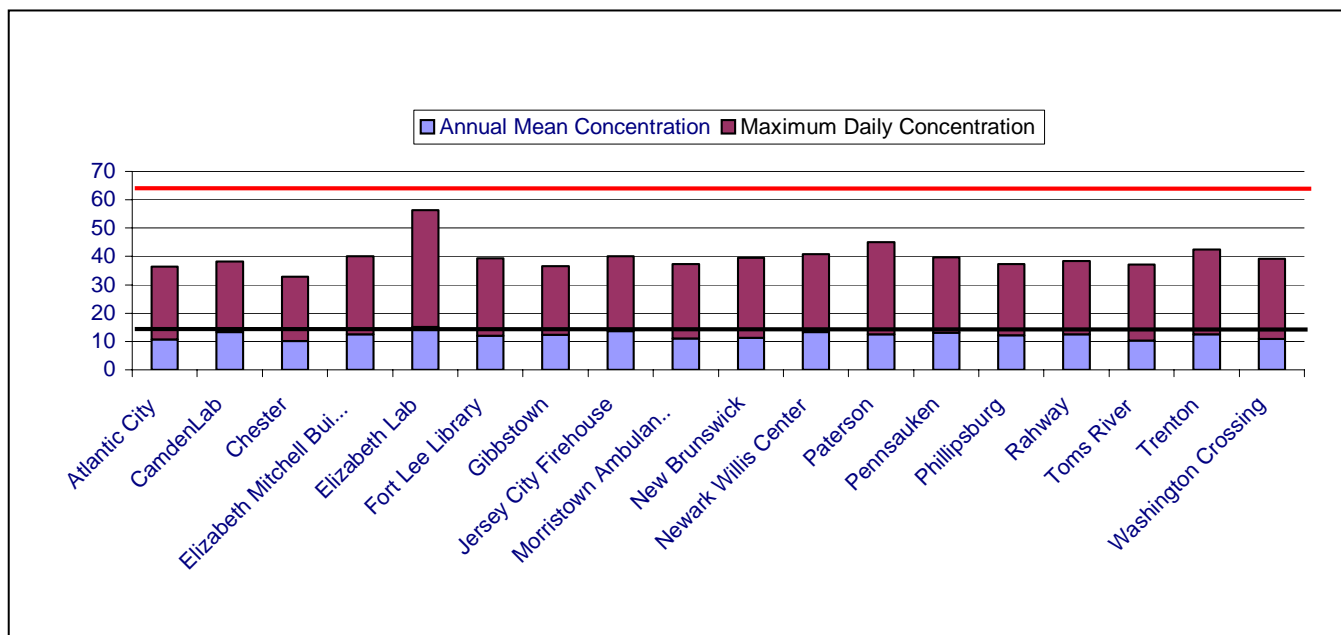


Table 2
PM_{2.5} Summary Data – 2004

Monitoring Site	Number of Samples	24-Hour Maximum µg/m³	Second Highest µg/m³	Annual Mean µg/m³
Atlantic City	89	36.3	29.6	10.6
Camden Lab	110	38.2	36.8	13.3
Chester	113	32.8	30.5	10.2
Elizabeth-Mitchell Building	113	40.1	35.7	12.6
Elizabeth Lab	327	56.4	46.0	15.2
Fort Lee-Library	112	39.3	36.6	12.0
Gibbstown	103	36.6	32.9	12.4
Jersey City-Firehouse	116	40.1	37.6	13.8
Morristown-Ambulance Squad	107	37.3	34.0	11.1
New Brunswick	118	39.5	35.5	11.2
Newark-Willis Center	112	40.9	36.9	13.3
Paterson	104	45.0	39.0	12.6
Pennsauken	109	39.7	38.5	13.2
Phillipsburg	110	37.3	35.6	12.2
Rahway	111	38.5	36.7	12.7
Toms River	106	37.1	35.3	10.4
Trenton	108	42.4	38.4	12.5
Washington Crossing	110	39.2	35.7	11.0

Table 3
2004 Summary of Continuous PM_{2.5} Data

Concentration in Micrograms Per Cubic Meter (µg/m³)

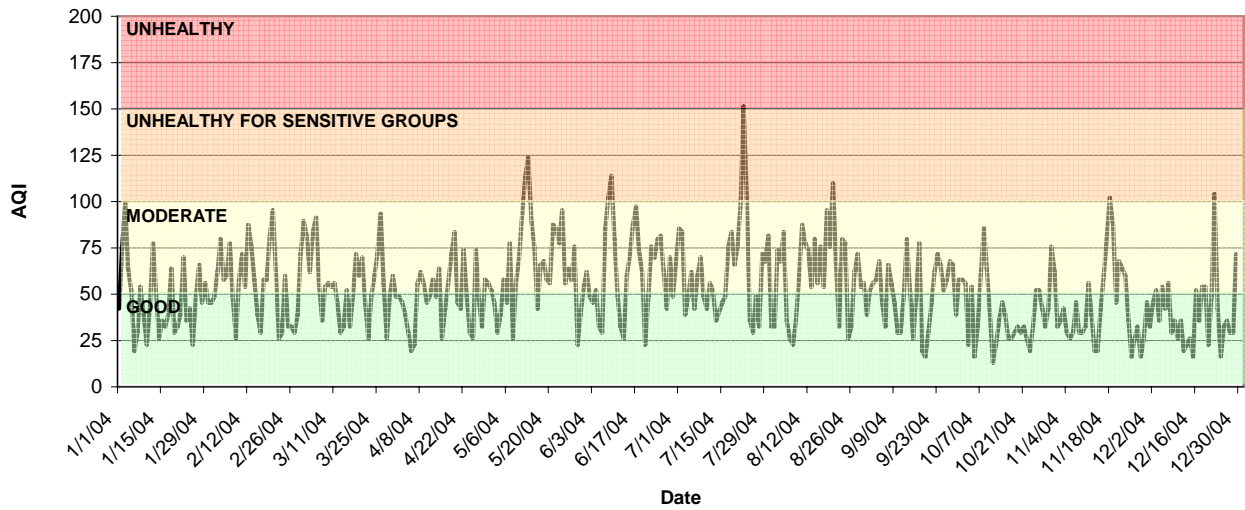
Monitoring Site	Annual Mean	Maximum Daily Concentration	2nd Highest Daily Concentration
Camden Lab	14	51	46
Elizabeth Lab	14	61	43
Fort Lee ^a	17	66	52
Jersey City-Firehouse	14	58	42
New Brunswick	12	47	42
South Camden	15	56	47

^a Data Not Available after October

PM_{2.5} REAL-TIME MONITORING

New Jersey's continuous PM_{2.5} monitoring network consists of 6 sites: Camden Lab, Elizabeth Lab, Fort Lee, Jersey City, New Brunswick, and South Camden. The data is transmitted once a minute to a central computer in Trenton, where it is averaged and automatically updated on the bureau's website every hour. Table 3 provides a summary of the data from these sites, and Figure 5 depicts the health level associated with the maximum daily fine particulate concentration recorded in the state each day for the entire year.

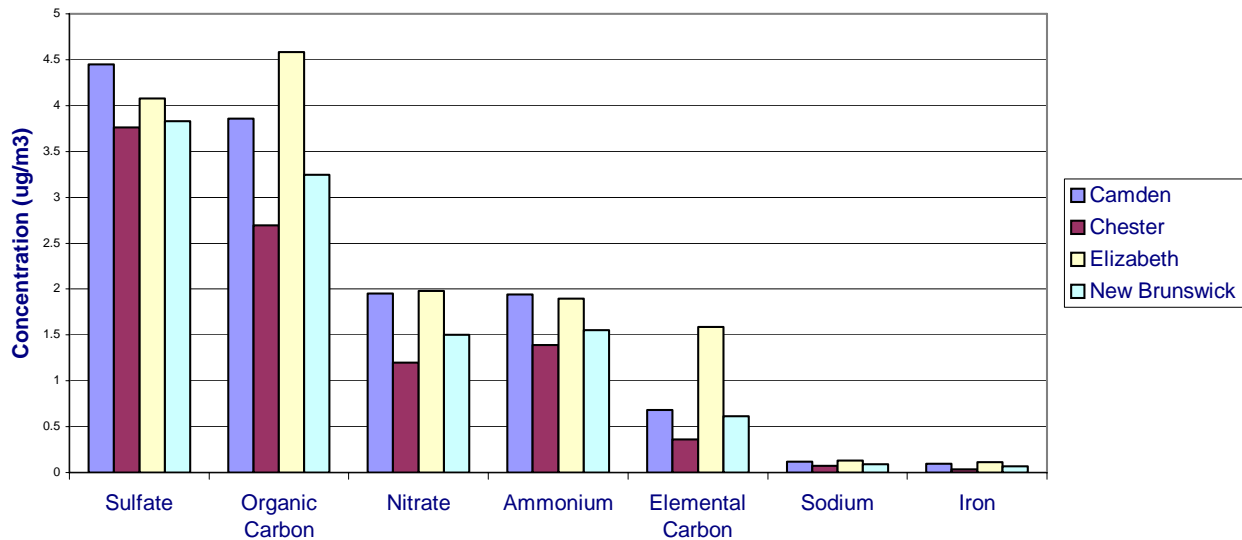
Figure 5
2004 Maximum Daily Fine Particulate Concentration
(Highest site)
Air Quality Index (AQI)



FINE PARTICLE SPECIATION SUMMARY

New Jersey's Fine Particulate Speciation Network consists of 4 monitoring sites: Camden Lab, Elizabeth Lab, New Brunswick, and Chester. Samplers run every third day on a schedule concurrent with the FRM sampling network. Of the 55 measured analytes, organic carbon and sulfate combined make up 59% of the total mass, and nitrate, ammonium, and elemental carbon make up an additional 35% of the particulate mass. Figure 6 depicts the average concentration of each analyte at all the sites, with only the seven most prevalent constituents depicted. Appendix B shows the average, maximum, and 2nd highest concentrations for each compound for 2004.

Figure 6
2004 Fine Particulate Analyte Composition
(Highest 7 Analytes Depicted)



2004 COARSE PARTICLE SUMMARY

COARSE PARTICLE MONITORING SITES

The coarse particulate monitoring network is composed of PM₁₀ sampling sites and TSP sampling sites. Samples are collected on a filter, which is weighed before and after sampling. The amounts of Sulfate and Nitrate are measured on some PM₁₀ samples and Lead is measured on the TSP samples. Figure 7 depicts the PM₁₀ particulate monitoring network in New Jersey.

Figure 7
2004 PM₁₀
Monitoring Network



TSP CONCENTRATION SUMMARY

New Jersey currently operates one site, located in New Brunswick, mainly for the purpose of determining the concentration of lead in the atmosphere. For more information, see the 2004 Lead Summary section. In 2004, the annual geometric mean concentration of TSP in New Brunswick was 25.7 $\mu\text{g}/\text{m}^3$, and the maximum 24-hour concentration recorded was 66 $\mu\text{g}/\text{m}^3$. The site was in attainment for the primary and secondary annual TSP standards of 75 $\mu\text{g}/\text{m}^3$ and 60 $\mu\text{g}/\text{m}^3$ respectively, and the site did not surpass the 24-hour primary standard of 260 $\mu\text{g}/\text{m}^3$ or the 150 $\mu\text{g}/\text{m}^3$ secondary standard.

PM₁₀ CONCENTRATION SUMMARY

In 2004, the annual mean concentration of PM₁₀ ranged from 18.8 $\mu\text{g}/\text{m}^3$ at Trenton to 41.3 $\mu\text{g}/\text{m}^3$ at Camden RRF. Table 4 shows the annual mean and 24-hour maximum PM₁₀ concentrations throughout the state. All areas of the state are in attainment for the annual PM₁₀ standard of 50 $\mu\text{g}/\text{m}^3$. Camden RRF measured one exceedance of the 24-hour standard of 150 $\mu\text{g}/\text{m}^3$.

The concentration of Sulfate and Nitrate were also analyzed on some PM₁₀ filters. The results showed that, on average, about 2% percent of PM₁₀ is nitrate and 15% percent is sulfate; however, these percentages vary across sites and sampling dates. The contributions of sulfate and nitrate to PM₁₀ are significantly less than their contributions to PM_{2.5}. This is because PM₁₀ is predominantly made up of larger particles most of which are directly emitted into the atmosphere. PM_{2.5} is predominantly a secondary pollutant, forming in the atmosphere from gaseous emissions, such as SO₂ and NO_x. For more details on the PM₁₀ sulfate and nitrate results, see the section on atmospheric deposition.

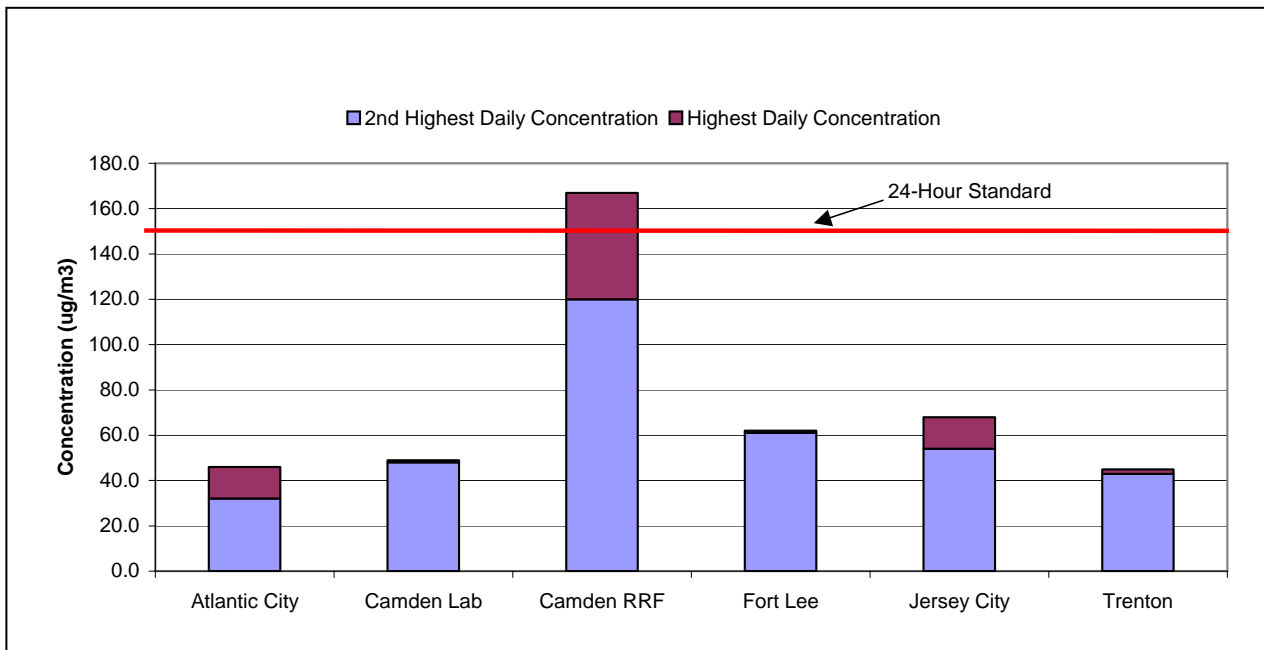
Table 4
PM10 Data - 2004
24-Hour and Annual Averages

Micrograms Per Cubic Meter ($\mu\text{g}/\text{m}^3$)
 24-Hour Standard = $150 \mu\text{g}/\text{m}^3$
 Annual Standard = $50 \mu\text{g}/\text{m}^3$

Monitoring Site	Number of Samples	24-Hour Maximum	Second Highest	Annual Mean
Atlantic City	52	46	32	19.4
Camden Lab	48	49	49	20.8
Camden RRF	55	167	120	41.3
Fort Lee ^a	39	62	61	31.8
Jersey City-Firehouse	59	68	54	27.4
Trenton	57	45	43	18.8

^a Data Not Available after September

Figure 8
Summary of PM₁₀ Concentrations, New Jersey 2004



SMOKE SHADE SUMMARY

SMOKE SHADE MONITORING SITES

In addition to fine and coarse particulate monitoring, smoke shade is also monitored at 10 stations around the state. Smoke shade, which is an indirect measurement of particles in the atmosphere, has been monitored in New Jersey for over 30 years. Smoke shade is primarily used for the daily reporting of particulate levels in the Air Quality Index. The sites monitoring smoke shade are shown in Figure 9.

SMOKE SHADE CONCENTRATION SUMMARY

In 2004, the annual mean concentration of smoke shade ranged from 0.17 Coefficient of Haze units (COH) at Flemington and Camden Lab to 0.55 COH at Elizabeth Lab. COH are units of light transmittance and smoke shade is not a direct measure of particle mass. A 24-hour average level of 2.0 COH is used as a benchmark. Readings above the 2.0 COH benchmark are reported as Unhealthy for Sensitive Groups on the daily Air Quality Index. For more details see the Air Quality Index section of this report. Table 5 lists the maximum and second highest daily average and annual mean smoke shade levels recorded at the monitoring sites in 2004.

**Figure 9
2004 Smoke Shade
Monitoring Network**



**Table 5
Smoke Shade - 2004**

Coefficient of Haze (COHs)
No Standard

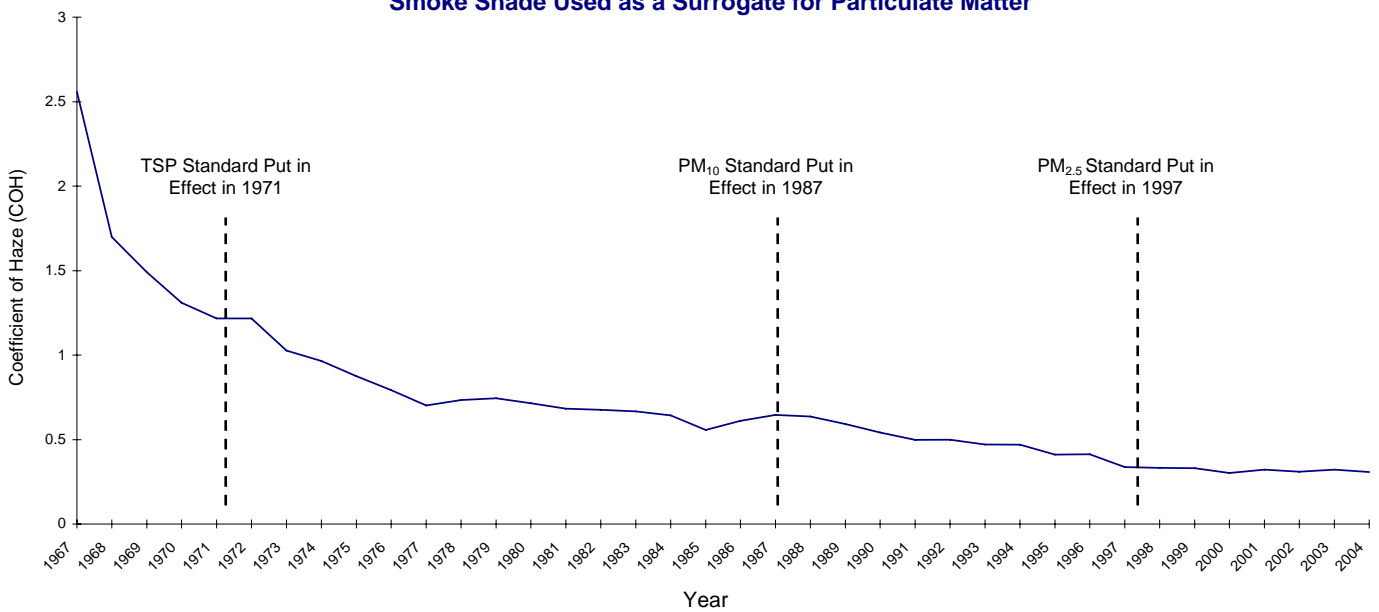
Site	Maximum Daily Average	2nd Highest	Annual Mean
Burlington	0.62	0.58	0.18
Camden Lab	0.84	0.82	0.17
Elizabeth	1.56	1.30	0.48
Elizabeth Lab	1.37	1.37	0.55
Flemington	0.68	0.50	0.17
Freehold	0.62	0.55	0.21
Hackensack	1.13	0.95	0.24
Jersey City	1.65	1.22	0.48
Morristown	0.51	0.42	0.19
Perth Amboy	0.88	0.85	0.27

TRENDS IN PARTICULATE CONCENTRATIONS

The longest continuously operating particle monitoring network in the state that is suitable for looking at trends is the smoke shade network. As noted earlier, this monitoring program has been in effect for over thirty years and still has 10 active sites. The trend graph for smoke shade, shown in Figure 10 indicates that particulate levels have steadily declined over the past thirty years. Smoke shade is not a direct measurement of particle mass, but can be related to TSP, PM₁₀ and PM_{2.5} health standards.

Figure 10
Long Term Trend in Particulate Levels
1967- 2004

Smoke Shade Used as a Surrogate for Particulate Matter



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