Final Report of the Public Health Investigation to Assess Potential Exposures to Airborne and Settled Surface Dust in Residential Areas of Lower Manhattan



New York City Department of Health and Mental Hygiene and U.S. Department of Health and Human Services Public Health Service Agency for Toxic Substances and Disease Registry

> as part of the World Trade Center Environmental Assessment Working Group





Final Technical Report of the Public Health Investigation to Assess Potential Exposures to Airborne and Settled Surface Dust in Residential Areas of Lower Manhattan

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New York City Department of Health and Mental Hygiene and Agency for Toxic Substances and Disease Registry U.S. Department of Health and Human Services

As a part of the

World Trade Center Environmental Assessment Working Group

September 11, 2001

In remembrance of those who were lost on September 11, 2001, and to the families and friends who will love them forever and keep their memories alive.

In grateful recognition of the countless responders and the many who supported their efforts.

New York City Department of Health and Mental Hygiene

U.S. Agency for Toxic Substances and Disease Registry

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EXECUTIVE SUMMARY

From November 4 through December 11, 2001, environmental samples were collected in and around 30 residential buildings in lower Manhattan. In addition, four buildings above 59th Street were sampled and used as a comparison area for this investigation. The New York City Department of Health and Mental Hygiene (NYC DOHMH) and the U.S. Agency for Toxic Substances and Disease Registry (ATSDR) conducted this limited investigation with support and collaboration from the U.S. Public Health Service Commissioned Corps Readiness Force and the World Trade Center Environmental Assessment Working Group. The purpose of the sampling was to assess the composition of both outdoor and indoor settled surface and airborne dust within residential areas around the World Trade Center. This information was used to help determine whether additional public health actions are needed to address any remaining World Trade Center-related dust inside residential areas. The information collected could also be used to compare the findings from the locations that were known or were likely to have received dust directly from the collapse of the World Trade Center towers, to findings from areas that were unlikely to have received dust directly from the disaster (comparison areas).

Attention was given to those materials reasonably expected to be in the original dust cloud and in dust generated by ongoing activities at the World Trade Center. The focus was on building materials that have been shown to have irritant properties (e.g., synthetic vitreous fibers [SVF] and gypsum) and be associated with long-term health concerns (i.e., crystalline silica and asbestos). The samples collected during this investigation were analyzed for the following materials: asbestos, SVF, mineral components of concrete (crystalline silica, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite). Efforts were made to get as much information as possible with the sampling that could be conducted, given accessibility and equipment limitations.

Results from this investigation do not necessarily reflect conditions that would be found in other buildings, at other times just following the collapse, or after the sampling period. The measurements reflect conditions present at the time of the sampling (November 4– December 12, 2001) in the buildings and areas sampled. The limited number of results obtained from the comparison areas above 59th Street was an attempt to determine the New York City–specific background levels of asbestos, SVF, mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite).

Sampling Overview

U.S. Public Health Service Commissioned Corps Readiness Force Officers were detailed to the New York City Department of Health and Mental Hygiene and composed the sampling teams for this investigation. The teams completed a survey form for each sampling location. The information collected by the survey was obtained from direct observations and from owners or representatives of building management when available. The survey form and photographs were reviewed along with the analytical results to put the sampling results into the context of where and how the samples were obtained and to determine the general conditions of the sampling locations.

Air and settled surface dust samples were collected and analyzed for the following materials used in WTC construction components: asbestos, SVF, crystalline silica, calcite, portlandite, gypsum, mica, and halite. SVF (e.g., fiberglass) is used in thousands of products because of its chemical resistance, strength, and ability to insulate against heat and sound. All of the crystalline minerals analyzed in this project are commonly used in building construction materials. *Quartz*, a form of crystalline *silica*, is a naturally occurring mineral and is a component of cement. *Cristobalite* and *tridymite* are different crystalline forms of silica. *Calcite and portlandite* are naturally occurring mineral used in plaster, wallboard, and in some cements. *Mica* is a group of naturally occurring minerals that are used in paint, joint cements, plastics, roofing, and rubber. *Halite* (also known as rock salt or sodium chloride) is used in ceramic glazes, fire extinguishing, metallurgy, and highway de-icing and table salt.

In addition to analyzing the samples for the fibers and minerals found in common building materials, when a sufficient quantity of settled surface dust was present, it was mixed with water and the pH of the resulting mixture was measured.

Air Samples

Air samples were collected using vacuum pumps to draw air through a filter positioned at an adult's breathing level. In addition to a filter for fiber analysis, multiple particulate samplers were placed at each sampling location to collect different particle sizes including respirable (PM4 and PM2.5), thoracic (PM10), and total inhalable (PM100). This resulted in the collection of overlapping, or nested, particulate size fractions.

The particle size determines how deeply a particle can travel into the respiratory tract. Respirable particles with diameters of 4 microns or less (PM4) are very small particles that can be breathed in and can travel deeply into the air sacs of the lung. At some locations another sampler was used that collected a smaller set of respirable particles, those with diameters of 2½ microns (PM2.5) or less. Another sampler collected particles with diameters of 10 microns (PM10) or less. This sampler would collect not only respirable particles, but also those slightly larger particles that tend to land in the upper regions of the respiratory tract, including the throat. A fourth sampler collected a larger set of particles, those 100 microns (PM100) or less in diameter. The larger particles in this fraction can land in the nose and throat. Where a particle lands determines what types of health effects might be experienced. Particles too large to travel into the narrower passages of the lower respiratory tract might deposit in the upper airways (e.g., the nose and throat) and cause irritation. Similarly, larger particles might cause eye irritation. If the settled surface dust contains sufficient amounts of particles of a very small size, then effects consistent with particles reaching the lower respiratory tract might occur (coughing, shortness of breath).

Each of the particulate air sample fractions was analyzed for crystalline minerals (quartz, cristobalite, tridymite, calcite, portlandite, gypsum, mica, and halite). X-ray diffraction analysis (XRD) identifies a mineral by the unique diffraction pattern given off when the x-ray strikes its surface. The diffraction pattern serves as a "fingerprint" to identify the mineral present.

Other air samples were screened for fibers using phase contrast microscopy (PCM) to count the total number of all fibers present. If the concentration of total fibers was higher than the maximum concentration of fibers found in the comparison homes (0.003 fibers per cubic centimeter of air) the sample was re-analyzed for asbestos by transmission electron microscopy (TEM). Additionally, scanning electron microscopy (SEM) was used to look for SVF if the PCM fiber counts was higher than 0.003 fibers per cubic centimeter (f/cc) and if the settled surface dust sample from that area contained SVF.

Settled Surface Dust Samples

The composition of settled surface dust reflects particles and fibers that have been tracked into an area or that were present in the air and have with time, deposited on surfaces. Settled surface dust was collected indoors using a specialized vacuum cleaner. Outside settled surface dust was collected by scooping any visible dust-type material into a container. As with the air samples, the settled surface dust samples were analyzed to assess the presence of fibers and minerals. The analysis performed does not allow the determination of what size particles are present in the dust. The composition of the settled surface dust was evaluated to determine if, on the basis of its composition, it could be an irritant if it became airborne or came in contact with skin or eyes during cleaning activities.

The dust samples were analyzed for the presence of asbestos and SVF using polarized light microscopy (PLM). PLM can distinguish between fiber types by their unique appearance and color when viewed under different wavelengths of light. In addition, TEM analysis was conducted on all dust samples that showed asbestos content by PLM as less than 1%. TEM analysis is very specific, can detect smaller fibers of asbestos, and is subject to less interference than PLM.

The dust samples were also analyzed for mineral content using XRD as described previously for the air samples. For the dust samples, the mineral and the fiber results are expressed as a percent, indicating the weight percent each component represents in the portion of the dust sample analyzed. The different analytical techniques, XRD and microscopy (PLM, TEM, SEM), help to characterize what was in the settled surface dust; however the results from the two methods each performed on a portion of the dust collected, cannot simply be added together to "sum up" what was in the dust. XRD results are based upon the weight of the dust sample analyzed. While, PLM, TEM, and SEM results are based on the surface area viewed under the microscope.

Limitations

While the goal of the sampling was to evaluate the air and dust samples for a range of contaminants, there is no one sample collection method or analysis that can provide all of the determinations that were desired. The analysis strategy was to identify what components were present in order to provide the public with information quickly if any of the analyses revealed a significant concern. In order to strive for both of those goals, some modifications to conventional protocols were made which limited some analyses from determining the precise level of the materials in the sample (e.g., XRD conducted on air and dust samples). Whenever an analysis is expanded beyond its specific intent, there is the potential loss of sensitivity and an increase in the variability in the data. Those particular analytical results are reported as estimated values and marked with a "J."

Results for the airborne particulate matter were rejected due to issues found during the quality assurance analysis. However, the XRD analysis for airborne minerals that used the same filter samples as the particulate matter analysis are considered valid because the analysis method does not depend on the filter sample weighing process used to determine the particulate matter concentration. The XRD mineral air concentrations are based on individual mineral weight standard curves—not the weight of the dust sample collected on the filters. These standard curves are not impacted by pre- and post-sampling filter weight variability.

Air Sampling Results

Total fiber counts of air samples taken in lower Manhattan were similar to the comparison areas above 59th Street sampled during this investigation. The six lower Manhattan areas that had elevated total fiber counts were re-examined by TEM and SEM to determine the types of fibers. The TEM and SEM results indicated that neither asbestos nor synthetic vitreous fibers (e.g., fiberglass) contributed to the elevated total fiber counts.

Air sampling results for minerals detected quartz (a form of crystalline silica) and other building-related materials in lower Manhattan. The other forms of crystalline silica were not detected in any air samples except for a one-time detection of cristobalite. The estimated concentrations of these minerals in air were low. In some locations, mineral components of concrete (quartz, calcite, and portlandite) and mineral components of building wallboard (gypsum, mica, and halite) were detected in air samples at higher estimated levels in lower Manhattan residential areas than in samples taken at comparison residential areas above 59th Street. The levels of minerals seen in airborne dust do not pose potential health hazards even for a continuous year of exposure at the highest levels detected.

Settled Surface Dust Results

Results of the settled surface dust analysis indicate the composition of settled surface dust in lower Manhattan is different than the dust analyzed from the comparison areas above 59th Street. Although the materials found are consistent with materials expected from the World Trade Center collapse, these results cannot determine the actual source of the materials present because these materials are common building components and can come from other sources in a busy urban area.

In lower Manhattan, asbestos was found in indoor dust in 15 of 83 (18%) samples from residential units and common areas at levels ranging from less than 1% (<1%) to 1.5%. Asbestos was detected in 6 of 14 (43%) outdoor samples at levels ranging from <1% to 3.4%. Indoor settled surface dust contained SVF in 40 of 83 (48%) locations ranging from 2% to 35% of the dust content. SVF was detected in 11 of 14 (79%) outdoor locations at levels ranging from 1% to 72%. No asbestos or SVF was detected in dust in the comparison areas above 59th Street.

The XRD analysis for crystalline minerals in settled surface dust is semiquantitative (estimated values, indicated by "J"). However, quartz, calcite, portlandite and gypsum appear to make up a higher percentage of dust in some buildings in lower Manhattan when compared to settled surface dust samples from buildings above 59th Street. Quartz was detected up to an estimated 31%J versus up to 2%J found in the comparison areas above 59th Street. Neither cristobalite nor tridymite was detected in any of the settled surface dust samples. Similarly gypsum was found at a maximum estimated concentration of 30%J in settled surface dust, higher than the 4%J estimated in the comparison areas above 59th Street. Calcite and portlandite had maximum concentrations of 21%J and 8%J respectively. At lower Manhattan locations sampled, quartz was detected in 81% of common areas and 53% of residences. Gypsum was seen in 88% of common areas and 79% of residences. Minerals were found in all lower Manhattan outdoor settled surface dust samples at estimated values ranging as high as 27%J quartz, 19%J calcite, 6%J portlandite, and 27%J gypsum. No visible settled outdoor dust was available in the comparison areas above 59th Street.

Several of the minerals detected in the settled surface dust samples, such as portlandite and calcite, can make the dust more alkaline, or raise the pH. Only two dust samples provided enough material for the determination of pH. The samples, collected from two outdoor locations in lower Manhattan, had pH levels of 8.6 and 9.8. On the pH scale, values less than 7 are considered acidic, a value of 7 is neutral, and values above 7 are alkaline or basic. Based on the results of the pH analyses, these dust samples are slightly alkaline. This is consistent with the detection of portlandite and calcite (alkaline minerals present in concrete) in the mineral analysis of the dust sample from the same location.

Conclusions

Based upon the analytical results of samples taken between November 4 through December 11, 2001, and information collected during the sampling effort, the New York City Department of Health and Mental Hygiene and the U.S. Agency for Toxic Substances and Disease Registry make the following conclusions (not in order of importance).

• Total fiber counts of air samples taken in lower Manhattan were similar to the comparison areas above 59th Street sampled during this investigation. The six lower Manhattan areas that had elevated total fiber counts were re-examined by transmission electron microscopy (TEM) and scanning electron microscopy (SEM). The TEM and SEM results indicated that neither asbestos nor SVF contributed to the elevated fiber counts.

• Low levels of asbestos were found in some settled surface dust, primarily below Chambers Street. Many of the lower Manhattan locations sampled had been previously cleaned prior to this investigation. No asbestos was detected in the comparison indoor dust samples taken north of 59th Street. The City of New York has conducted follow-up activities at the locations where asbestos was detected in settled surface dust. Only two follow-up locations, outdoor areas, required professional asbestos abatement. Following-up activity at the other locations did not find any asbestos containing materials.

• When compared with the results obtained from the comparison sampling locations, the lower Manhattan residential areas sampled by this investigation tended to have a greater percentage of SVF (primarily fiberglass), mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite) in settled surface dust. However, the frequency of detections and prevalence patterns of these minerals are similar in both residential areas.

• Exposure to significant amounts of synthetic vitreous fibers (SVF), mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum) may cause skin rashes, eye irritation, and upper respiratory irritation, all of which have been voiced as concerns by citizens and first responders. If the reported irritant effects are associated with World Trade Center related materials, these effects will subside once exposure to SVF, mineral components of concrete, and mineral components of building wallboard end. Some people with pre-existing heart or lung disease (e.g., asthma) or a previous history of very high levels of exposures (occupational) to SVF, mineral components of SVF, mineral components of concrete, and mineral components of building wallboard end. Some people with a previous fiber (components may be more sensitive to the irritant effects of SVF, mineral components of concrete, and mineral components of building wallboard.

• Sometimes mineral components of concrete (calcite and portlandite) and mineral components of building wallboard (gypsum, mica, and halite) were detected in air samples at higher estimated levels in lower Manhattan residential areas than in samples taken at comparison residential areas. These detected mineral levels are orders of magnitude below occupational standards. Although the occupational standards do not account for sensitive individuals or extended periods of exposure, they provide a comparison to an established health guidance value. The levels of minerals seen in airborne dust do not pose potential health hazards even for a continuous year of exposure at the highest levels detected.

• Some settled surface dust could become airborne if disturbed. Therefore, people could potentially inhale the asbestos, SVF, mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite) found in settled surface dust of some lower Manhattan residences. Because we did not determine the weight of dust present in the areas sampled, it is not possible to determine whether any particular residence had an elevated dust loading. Appropriate continued frequent cleaning should minimize exposures.

• Several worst-case assumptions were made in order to assess the potential longterm public health risks of airborne asbestos and quartz. Some of the assumptions were that no cleaning of indoor spaces has occurred or will occur, all fibers found in air were asbestos fibers, and the highest levels detected last fall in air represent long-term air levels. Using these worst-case assumptions, prolonged exposure (decades) to airborne asbestos and quartz *may* increase the long-term, theoretical risk of people developing lung cancer and other adverse lung health effects (more than 1 additional case in 10,000 people exposed). For individuals who conduct frequent cleaning of their residences, as recommended in this report, or participate in the U.S. Environmental Protection Agency cleaning/sampling program, it is unlikely that their exposure would resemble these worst-case conditions.

• A review of the building sampling results from this investigation indicates that there is not a consistent spatial distribution pattern of asbestos, SVF, mineral components of concrete, and mineral components of wallboard in air and settled surface dust. This indicates that the materials are heterogeneously distributed. There are many factors that may contribute to the heterogeneous distribution, including whether the area was cleaned (indoors and outdoors), cleaning method, date since last cleaning, and how much dust was initially in the area. It is not clear which factors contributed to this pattern.

• Results from this investigation do not necessarily reflect conditions that would be found in other buildings, at other times just following the collapse, or after the sampling period. The measurements reflect conditions present at the time of the sampling (November 4–December 12, 2001) in the buildings and areas sampled. The limited number of results obtained from the comparison areas above 59th Street may or may not reflect the New York City-specific background levels of asbestos, SVF, mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite).

Recommendations

Based upon the conclusions of this investigation, the New York City Department of Health and Mental Hygiene and the U.S. Agency for Toxic Substances and Disease Registry make the following recommendations.

• Because more asbestos, synthetic vitreous fibers (e.g., fiberglass), mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite) were found in settled surface dust in lower Manhattan residential areas when compared to comparison residential areas, the New York City Department of Health and Mental Hygiene and the U.S. Agency for Toxic Substances and Disease Registry are recommending that people continue to conduct frequent cleaning with HEPA vacuums and damp cloths/mops to reduce the potential for exposure.

• To ensure that the recommended frequent cleaning is effective and to ensure that the health of the people of New York City is protected, the New York City Department of Health and Mental Hygiene and the U.S. Agency for Toxic Substances and Disease Registry are recommending additional monitoring of residential areas in lower Manhattan. In addition, an investigation should be conducted to better define background levels specific to the city of New York for asbestos, synthetic vitreous fibers, mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite).

• Lower Manhattan residents concerned about possible World Trade Centerrelated dust in their residential areas can request cleaning and/or testing from the U.S. Environmental Protection Agency (EPA) by logging on to the agency's World Trade Center Web page at www.epa.gov/wtc or by calling the EPA hotline at 1-877-796-5471.

ABSTRACT

The New York City Department of Health and Mental Hygiene and the U.S. Agency for Toxic Substances and Disease Registry in collaboration with the U.S. Public Health Service Commissioned Corps Readiness Force and the World Trade Center Environmental Assessment Working Group conducted a limited investigation of residential areas near and around the World Trade Center (WTC). The purpose of the investigation was to assess potential exposures to airborne and settled surface dust that entered residential areas following the collapse of the WTC buildings.

Sampling of the residential areas occurred from November 4 through December 11, 2001, and was conducted by U.S. Public Health Service Commissioned officers detailed to the New York City Department of Health and Mental Hygiene with assistance from the New York City Department of Health and Mental Hygiene.

The results of this investigation do not necessarily reflect levels that would be found in other lower Manhattan buildings or residential areas; the measurements reflect levels that were present at the time of sampling in the specific residential areas sampled. The limited number of results obtained from the comparison areas above 59th Street may or may not reflect the New York City-specific background levels of asbestos, synthetic vitreous fibers (SVF), mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite).

Low levels of asbestos were found in some indoor and outdoor settled surface dust, primarily below Chambers Street. Many of the lower Manhattan locations sampled had been cleaned prior to this investigation. No asbestos was detected in the comparison indoor dust samples taken north of 59th Street. Not enough visible settled dust was available to collect outdoor samples in the comparison area. When compared with the results obtained from the sampling locations above 59 Street, the lower Manhattan residential area settled surface dust sampling results tended to have higher percentages of SVF, mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite) in settled surface dust.

Because asbestos, SVF, mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite) were found at higher levels in settled surface dust in lower Manhattan residential areas when compared to comparison residential areas above 59th Street, the New York City Department of Health and Mental Hygiene and the U.S. Agency for Toxic Substances and Disease Registry are recommending that people continue to conduct frequent cleaning with HEPA vacuums and damp cloths/mops to reduce the potential for exposure and/or participate in the U.S. Environmental Protection Agency cleaning/sampling program.

Total fiber counts of air samples taken in lower Manhattan were similar to the comparison areas above 59th Street sampled during this investigation. The six lower Manhattan areas that had elevated total fiber counts were re-examined by transmission electron microscopy (TEM) and scanning electron microscopy (SEM). The TEM and SEM results indicated that

neither asbestos nor SVF contributed to the elevated fiber counts. Sometimes mineral components of concrete (calcite and portlandite) and mineral components of building wallboard (gypsum, mica, and halite) were detected in air samples at higher estimated levels in lower Manhattan residential areas than in samples taken at comparison residential areas. These detected mineral levels are orders of magnitude below occupational standards. Although the occupational standards do not account for sensitive individuals or extended periods of exposure, they provide a comparison to an established health guidance value. The levels of minerals seen in airborne dust do not pose potential health hazards even for a continuous year of exposure at the highest levels detected.

INTRODUCTION

At 8:45 AM eastern standard time on September 11, 2001, a commercial jetliner struck the north tower of the World Trade Center (One World Trade Center). At 9:03 AM, the south tower (Two World Trade Center) was also struck by a commercial jetliner. The south tower began to collapse at 10:05 AM. At 10:30 AM, the north tower began to collapse. (1,2,3,4)

The collapse of these structures released massive amounts of dust and debris that covered lower Manhattan. Residents were evacuated from many areas south of Canal Street. On September 19, pedestrian and vehicular traffic restrictions for below Canal Street were modified (5). As residents moved back into the areas, they requested information on the safety of the dust and air in the area and the types of cleaning methods that should be used. The New York City Department of Health and Mental Hygiene provided guidance on cleaning for area residents and requested federal assistance in assessing potential exposures to airborne and settled surface dust that entered residential areas following the collapse of the World Trade Center (WTC) buildings.

BACKGROUND

A. World Trade Center Towers

The World Trade Center was a complex of 7 buildings on 16 acres surrounding a 5-acre plaza in lower Manhattan. The complex was bounded by Vesey Street on the north, Church Street on the east, Liberty Street on the south, and West Street on the west. The twin towers were the center of the complex. Each tower had 110 floors; the north tower was 1,368 feet tall and the south tower was 1,362 feet tall. Each tower had approximately 43,200 square feet on each floor. There were seven underground levels that contained services, shopping areas, and a subway station. There were two plaza buildings (WTC 4 and WTC 5) on Church Street that had nine stories. WTC 7 was a 47-story office building. Design and structure details of the towers are shown in Table 1 (1,2,3,4,6). Basic location and U.S. Census population information about lower Manhattan is provided in Table 2 and Figures 1 through 4.

B. Potential Contaminants

The dust cloud released at the collapse of the buildings was comprised of materials that were used in the construction of the buildings (such as concrete and insulation materials). In addition to the dust cloud from the building collapse, contaminants were released in fires fed by jet fuel and the combustible materials in the buildings.

The dust contained the constituents of concrete, wallboard, and insulation. Concrete is a mixture of cement, aggregate (sand and gravel), and water. Cement contains limestone, clay, gypsum, and metals. Sometimes other minerals (such as fly ash or crystalline silica) are added to concrete to change its properties. Concrete tends to have a high pH, and if dust from the concrete was mixed with moisture, the resulting solution would tend to have a high pH. Asbestos was used as insulation in a portion of the towers. Fiberglass, a synthetic vitreous fiber, was also used as insulation in the buildings and is likely to have been used in building furnishings. Surface dust and debris containing asbestos have been found in lower Manhattan. Asbestos was detected in the personal air samplers of rescue and recovery workers. In addition, low levels of asbestos, fibrous glass, and the constituents of concrete and wallboard have been detected in air samples collected at the WTC site perimeter. These materials are common building components, and there are other sources of these materials in any large urban areas.

C. Monitoring in Lower Manhattan After the Collapse of the World Trade Center

The U.S. Environmental Protection Agency (EPA) and other federal, state, local agencies, and private firms have been conducting environmental monitoring in a variety of media since September 11 (some of the ambient air sampling locations are depicted on Figure 5). In November 2001, EPA tasked the IT Corporation Quality Assurance Technical Support Program contractor to evaluate the environmental monitoring data that had been collected. Through April 24, 2002, there were approximately 263,000 sample results analyzed (7). There were 605 analytics that were included in the database such as asbestos, fibers, particulates, metals, volatile organic compounds (VOCs), polycyclic aromatic hydrocarbons (PAHs), and polychlorinated biphenyls (PCBs). The greatest number of analytical samples was collected for analysis of total fiber concentration, asbestos fiber concentration, or the percentage of asbestos. The overall largest number of samples in the monitoring was for particulate matter. The monitoring stations for particulate matter (roof-top locations) are a part of the New York State Department of Environmental Conservation air quality-monitoring program. Additional stations were added after the WTC towers collapsed.

The analytical results of air samples showed decreasing concentrations that appeared to have reached pre-existing background levels by the end of April 2002 (7). Bulk dust samples were tested for asbestos, dioxins/furans, PCBs, PAHs, and metals. The levels for some of the analytics (e.g., asbestos) were relatively high; however, only a few samples obtained after mid-October showed elevated levels (7).

In general, air monitoring near the WTC has not detected any pollutants from the fire and building collapses at levels that are of public health concern. Within the restricted zone (within one block of the WTC), low levels of asbestos have been found in the airborne dust from the building collapse (7). Bulk dust and air samples (mostly on or near the work site), as well as surface runoff water, river sediment, and river water have been analyzed for other contaminants, including metals, VOCs, PAHs, dioxins and PCBs. These contaminants were found at low levels that did not indicate a need to conduct further sampling for them at the time of the residential sampling (8).

D. Community Exposures

The most likely routes of community exposures to contaminants would be through breathing in airborne dust and through direct contact with surface dust and debris. The sampling plan was developed to determine the potential for exposure to building material-related contaminants in air and dust.

E. Human Health Effects of Concern

The primary health effects of immediate concern are respiratory effects and irritant effects from inhalation of or direct contact with building-related materials. Many of the materials expected to be in airborne dust can cause eye, nose, and throat irritation. In addition, those materials that can be inhaled or respired from airborne dust can cause respiratory irritation and exacerbation of pre-existing problems such as asthma, emphysema, and cardiopulmonary disease. Materials in the surface dust can cause skin irritation on contact. There have been reports of burning throat and eyes from residents returning to the area.

OBJECTIVES

The **overall objective** of this sampling was to provide results upon which public health agencies could further determine the potential for environmental exposures to WTC collapse-related materials and possible health implications of the exposures. Sampling was intended to better characterize ambient and indoor airborne and potentially airborne particles (surface dust) in residential areas of lower Manhattan from the collapse of WTC.

Specifically, the objective was to characterize the makeup and size distribution of airborne breathing zone dust in residential areas—both indoors and outside—near the WTC. This information was used to further determine whether materials were present in air at levels of public health concern and whether settled surface dust contained materials that could potentially be re-entrained and result in continued exposures.

Note: The sampling approach was focused on contaminants that emanated from the collapse of the WTC towers and that might cause health problems. However, this investigation could not conclude if contaminants or materials found actually came from the disaster. The contaminants that were selected for testing could be present in some areas from other sources. Some typical examples of this include cooking on a stove top, which can increase indoor airborne particulate matter; vehicle exhaust,

which can increase outdoor airborne particulate matter; and recent home renovation activities, which can result in an increase in levels of gypsum, fiberglass, and other materials in air and surface dust.

METHODS

Details on the methods used in the residential sampling investigations are included in the residential sampling plan that is on the New York City Department of Health and Mental Hygiene's Internet page (<u>http://www.nyc.gov/html/doh/pdf/alerts/indoor1.pdf</u>). Printed copies of the plan can be requested from the New York City Department of Health and Mental Hygiene.

A. Sampling Approach

A series of three concentric circles was drawn around the WTC location. The circles were then divided by direction or zone from the WTC (north, south, and west). Attempts were made to obtain access to the same number of residential buildings per zone for sampling. Four residential buildings above 59th Street were selected for testing to determine what levels of fibers and minerals might be routinely found in the city.

Once a building was selected, air and settled surface dust sampling occurred in up to four different locations: an area outdoors near a main entryway to the building; an area inside of the main entry (or other common area); and two residential units. Informed consent for sampling was obtained and questions were asked about details that could aid in interpreting the sampling results (such as whether the windows were open on the morning of September 11, 2001).

Both air and settled surface dust samples were collected in high traffic or routinely used areas of the space. For example, settled surface dust was collected from areas where individuals frequently walked, not from behind furniture or underneath kitchen appliances. Air sample inlets were at approximately 4 feet in height above the ground/floor surface to provide results more reflective of what might be inhaled.

Attempts were made during air sampling to obtain data that would reflect what might be in the air during periods of high activity. If the selected area was not occupied at the time of sampling, and if there was no central heating or air conditioning system that could result in contaminant migration into another occupied area, aggressive air sampling was conducted. If the area was occupied, the heating or air conditioning system fan was turned on during the period of sampling. This would not represent a period of high activity, but would present data that might be found during typical or normal activity.

B. Samples Taken

Sample collection was conducted from November 4, 2001, to December 11, 2001, for the lower Manhattan area and from December 6–10, 2001, for the comparison buildings above 59th Street. Because of the limited and time critical essence of this investigation, thirty

buildings in lower Manhattan were sampled, and four comparison buildings were sampled from the area north of 59th Street. In the 30 lower Manhattan buildings, the following locations were sampled: 59 residential units, 29 common areas, and 29 outdoor areas.

There were 127 air samples taken from 117 different areas of the lower Manhattan buildings. Ten of the areas had co-located sampling equipment. In the 59 different residential units, there were 64 air samples taken. Thirty air samples were taken from 29 different common areas.

Samples taken from the four comparison buildings above 59th Street included three outside air samples, three of common areas with both indoor settled surface dust and air, and five residential units with both indoor settled surface dust and air. No outside settled surface dust samples were collected because there was no visible dust to collect.

Tables 3, 4, and 5 show the number of buildings and areas within the buildings that were sampled for asbestos, SVF, and minerals in lower Manhattan and the comparison units above 59th Street. The tables also show whether the samples were settled surface dust and air samples.

C. Field and Laboratory Changes From the Original Sampling Protocol

Once the sampling teams went to the various sampling locations, it was determined that some changes needed to be made to the sampling protocol. The sections below indicate the changes that were made to the original sampling protocol.

1. Field Changes

The sampling protocol indicated that a total of 60 residential units would be tested in lower Manhattan. A total of 59 residential units were sampled.

Indoor settled surface dust was collected using EPA's Environmental Response Team Standard Operating Procedure (ERT SOP) for household dust, as stated in the sampling protocol. However, due to cost and equipment availability constraints, Omega Ultivac vacuums with high efficiency particulate air (HEPA) filters were used instead of NilfiskTM vacuums. To avoid the need to decontaminate the vacuum hose between each sampling event, a sample collection filter (Alsock, by Midwest Filtration) was inserted into the air-intake end of the vacuum hose. The Alsock has a 97% capture of particles with diameters of 1.1 microns and larger. The ERT SOP-specified vacuum bag filter has a 4- to 5-micron pore size.

Instead of one settled surface dust sample per unit, two co-located indoor settled surface samples were collected. This change was made as two different laboratory locations were involved in the analysis. The EMSL Analytical laboratory in New York City analyzed samples for fibers (using PCM, PLM, and TEM); the EMSL Analytical laboratory in Westmont, New Jersey, analyzed samples for the remaining materials and conducted the SEM analysis for fibers.

Delays in obtaining equipment resulted in only 10 buildings being sampled for the PM2.5 air fraction.

There was no visible settled surface dust around the comparison homes selected above 59th Street; therefore, no outdoor settled surface dust samples were collected from the comparison areas above 59th Street.

2. Sample Analysis Changes

Airborne and settled surface dust samples were analyzed for crystalline silica using EMSL Analytical's Material Science Division Operating Procedure MSD 0700, January 2000. XRD analysis was conducted for the nonsilica crystalline minerals in air and settled surface dust samples by EMSL using Material Science Division Operating Procedure MSD 9300, January 2002. The samples were not ashed prior to XRD analysis for the following reasons: ashing samples could have resulted in loss or breakdown of analytes of interest including gypsum; and there was not enough organic material present to result in loss of method sensitivity for crystalline minerals of interest.

SEM analysis was conducted on air samples for fibrous glass and other synthetic vitreous fibers by EMSL using Material Science Division Operating Procedure MSD 0300. This analysis was conducted on all samples collected from the comparison areas above 59th Street and on only those air samples from lower Manhattan with PCM fiber concentrations higher than the comparison PCM fiber concentrations.

Settled surface dust samples were analyzed for fibrous glass by EMSL using New York State Department of Health Environmental Laboratory Approval Program Method 198.1 polarized-light microscopy methods for identifying and quantifying asbestos in bulk samples.

Settled surface dust samples were not weighed, so surface loading of materials could not be calculated. Although settled surface dust samples were sieved, sieving was not conducted as specified in the ERT SOP. Rather, sieving was conducted as specified in MSD 0700 and 9300.

D. Data Management and Quality Assurance/Quality Control Procedures

EPA provided the laboratory used for this investigation. The laboratory is part of EPA's "certified" Contract Laboratory Program. It used all of the EPA quality assurance/quality control laboratory methodologies when analyzing the samples collected for this investigation. EPA conducts audits of all the EPA-certified laboratories to assure that appropriate quality assurance/quality control laboratory methodologies are being used.

Chain-of-custody procedures were established and maintained throughout sample collection, transportation, and analysis of samples. These procedures helped to assure that analytical results could be matched to the sampling locations.

Following the sampling effort, more than 5,400 analytical results were individually returned from the laboratory. All results were entered into a Microsoft EXCEL spreadsheet along with the other raw data describing where and how the sample was taken. Asbestos-in-air laboratory results were submitted calculated in concentration and were used as received. The laboratory typically submitted particulates-in-air results as the initial and final filter weights (pre- and post-sampling weights). The total amount of material deposited on the filter and resulting concentrations for the sampling location were estimated from the laboratory results. Results for asbestos, fiberglass, and mineral components in settled surface dust were used as provided by the laboratory: percentage of material analyzed.

All of the results were reviewed as a part of normal data quality analysis. The general purpose of this quality assurance/quality control (QA/QC) review is not to pronounce the data "accurate," but to determine whether the results are obviously affected by sampling or analytical errors. For this investigation, the reported analytical results were accepted, rejected, or used conditionally. Decisions on how to treat a data set were made primarily by comparing the measured results to information gathered from the blanks using graphs and statistical analyses.

Most of the results were used as received from the laboratory. The laboratory reported the asbestos in air results as a concentration of asbestos in the sampled location. The asbestos and fibrous glass in airborne and settled surface dust were used as received.

Results for the airborne particulate matter were rejected due to issues found during the quality assurance analysis. Variability of field blank sample weights was beyond the acceptable limits. A more thorough discussion of why the airborne particulate matter results were rejected is located in Appendix A of this report.

The XRD analysis was conducted on the air samples collected for particulate matter (PM100, PM10, PM4, and PM2.5). The PM results were rejected, but the XRD results were accepted as semiquantitative. The XRD results are considered valid because the analysis method does not depend on the filter-weighing process used to determine PM concentration. The XRD mineral concentrations are based on individual mineral weight standard curves—not the weight of the dust sample collected. These standard curves are not impacted by pre- and post-sampling filter weight variability.

Minerals were not detected on the blank air samples. This indicates that sample contamination or cross-contamination did not occur during air sampling, shipping, or sample analysis. The XRD information for the air samples provided semiquantitative values (estimated values, indicated by "J"). The PM100 and PM10 results may be potential overestimates, as the crystalline mineral size used for the standard curve is close to PM4 in size. This will result in the XRD overestimating the concentration of minerals present in the larger fractions. Similarly, the PM2.5 results may be potential underestimates of mineral concentrations.

The XRD settled surface dust results are also considered semiquantitative ("J") due to particle sizes used for standard curve/calibration (as described for the air samples) and due to the lack of standard analytical error (SAE) information from the laboratory.

E. Data Analysis

The data were entered into an Excel spreadsheet, and descriptive statistics were developed for the sample results. Basic trend analysis on the analytical data was conducted using the SAS statistical program (version 8.01). The SAS System is an integrated system of software providing complete control over data access, management, analysis, and presentations. Examples of some of the SAS procedures ("Proc") used were the "Proc Mean" and "Proc t-Test."

F. Limitations and Uncertainties

The intent of the sampling was not to characterize the total extent of contamination in lower Manhattan or determine the extent of exposures for all people in lower Manhattan that may have resulted from the collapse of the WTC towers. Rather, it was a preliminary investigation to determine if building collapse-related materials could be present at levels of possible health concern in lower Manhattan. Results from this investigation do not necessarily reflect conditions that would be found in other buildings, at other times just following the collapse, or after the sampling period. The measurements reflect conditions present at the time of the sampling (November 4–December 12, 2001) in the buildings and areas sampled.

Some uncertainties tend to underestimate the risk of exposure while other uncertainties tend to overestimate the risk of exposure.

• The locations selected for sampling within each building may not be representative of the rest of the building; the building-specific results may either *overestimate or underestimate* the levels of WTC-related materials in other areas of the building.

• Sampling was conducted in November and December of 2001. Levels of particulate matter and airborne irritants were likely to have been higher in the days and weeks immediately after the WTC collapse. Levels of settled surface dust were likely higher as well. Outdoor dust contamination would have been reduced by wind, rain, and cleaning (HEPA trucks to vacuum the streets and sidewalks). Indoor settled surface dust may have been reduced if appropriate cleaning had been conducted prior to this sampling event. Therefore, these results serve as estimates, and are not necessarily reflective of conditions at earlier time points or of current conditions.

• Likewise, if continued cleaning has occurred since the November–December 2001 sampling event, levels of contaminants should be lower today. Therefore, the results of this sampling effort would tend to *overestimate* the current levels of WTC-related materials in the buildings.

Uncertainty also exists in determining at what level and duration of exposure people might experience irritation or other symptoms from exposure to fibers and crystalline minerals.

• The likelihood of experiencing irritation from individual contaminants or contaminant mixtures is uncertain. A short exposure to a small amount of an irritant may be tolerated, but exposures that continue for a longer period of time or are to a larger amount of an irritant, may overwhelm the body's ability to remove the irritant and increase the likelihood of irritation. Individual differences also play a significant role in whether or not irritation occurs. Using a risk estimate from an occupational situation may *underestimate* risks for more sensitive or susceptible individuals in the general population. Some of the information needed to estimate the risk of irritant symptoms, is not available from this study. When assumptions are made in the absence information (e.g., the duration and intensity of the total exposure) additional uncertainty is introduced.

• Persons with pre-existing respiratory (e.g., asthma, emphysema) or cardiopulmonary problems are more likely than healthy individuals to experience adverse health symptoms at lower levels of airborne particles and contaminants. Airborne dust and crystalline mineral levels based on studies of healthy individuals are levels that *underestimate* the risk for a susceptible individual.

• Individuals who were in the lower Manhattan area when the WTC collapse occurred may have received a bolus inhalation dose of dust, fibers, crystalline minerals, and other materials. Health effects of a bolus dose are not known. Additional exposures would add to the potential for health effects. Therefore, the short and long-term health effects, if any, of such an exposure cannot be determined from the measurements made in this study.

RESULTS

This section of the report will provide an overview of what was determined by this investigation. The overall analytical findings will be presented according to what environmental media was sampled (air or settled surface dust, Figures 7-17).

A. Survey

The primary duty of the sampling team was to coordinate with the building management to identify sampling locations that would yield results representative of the area sampled. These duties included identifying unoccupied residences that could have aggressive air sampling conducted. In addition the sampling teams photographed equipment setup and operation, sampling locations, distribution of surface dust, and general conditions around the building.

The sampling team completed building surveys, in part, to gather the information needed to cite sample locations and to provide information to be used in interpreting the results. These surveys were completed on the basis of their observations and on discussions with building management when representatives were available. However, information was not always available to completely answer each survey question. This was especially true when evaluating specifics of the heating and air conditioning systems. The validity of the information is dependent on the memory and experience of the building representatives and, in the case of residential units, may not reflect the condition or treatment of the entire building.

The survey form and photographs were reviewed along with the analytical results to put the sampling results into the context of where and how the samples were obtained and to determine the general conditions of the sampling locations. Given the purpose of this investigation, the limited amount of locations sampled, and the uncertainty in some of the survey results, no attempt was made to correlate survey information with measured concentrations of any of the materials. The table in Appendix B provides a summary of results from the survey form questions that were most consistently answered.

B. Settled Surface Dust

Settled surface dust samples were analyzed for materials consistent with debris from a building collapse. Materials of interest include insulation material (asbestos, synthetic vitreous fibers, components of concrete (portlandite, calcite, and crystalline silica) and components of other building materials (gypsum, mica, and halite). A detailed summary of results is given in Appendix C. To protect the privacy of individuals who allowed sampling in their residences and buildings, specific detailed locations are not shown. Results of the settled surface dust analysis indicate the composition of settled surface dust in lower Manhattan is different from that of the dust analyzed from the comparison areas above 59th Street. Although the materials found are consistent with materials expected from the World Trade Center collapse, these results cannot determine the actual source of the materials present because these materials are common building components and can come from other sources in a busy urban area.

1. Fiber Analysis of Settled Surface Dust

All settled surface dust samples were analyzed by polarized light microscopy (PLM) for fiber type and content. This technique distinguishes between asbestos fibers, textile fibers, and synthetic vitreous fibers (SVF), which include glass wool and mineral wool. The results are reported as a

Asbestos and SVF, although not found in homes in the comparison areas above 59th Street, were present in the settled surface dust in some areas of lower Manhattan.

percent. Because PLM is an optical method, the amount reported is a visual estimation of the percent of area viewed. Therefore it is neither strictly a weight or a volume percent.

These results do provide an indication of the presence of asbestos or SVF in the dust, and some idea of relative amount.

In lower Manhattan, asbestos was only detected in settled surface dust in one indoor and two outdoor locations by PLM (Figure 8). If asbestos was not seen in the sample or reported as <1% by PLM, a second test was done which is more sensitive, transmission electron microscopy (TEM). An additional 4 outdoor and 15 indoor locations contained asbestos when examined by TEM (Figure 9). Chrysotile asbestos was the only form of asbestos identified, either by PLM or TEM, in all settled surface dust samples. In this study TEM is primarily used as an indicator of the presence or absence of asbestos. For the PLM analysis, values <1% are an indicator of the presence of asbestos because this is the detection limit for PLM. For TEM a detection of asbestos is confirmation of its presence. Therefore, values <1% and values greater indicate the presence of asbestos, by either the PLM or TEM methods (Figure 10).

Asbestos was found in indoor settled surface dust in 10 of 57 residences sampled in lower Manhattan (18%) (Table 6). The settled surface dust in common areas contained asbestos in 5 of 26 areas sampled (19%), and asbestos was found in outdoor dust in 6 of 14 areas sampled (43%) (Table 6). Asbestos levels ranged from none detected up to 3.4% (the highest level was found in an outside sample). However, without surface-loading information, the percentage of asbestos present does not allow for characterization of potential exposures. No asbestos was found in the comparison homes north of 59th Street. Figure 10 depicts the results of analysis of the asbestos content of settled surface dust in samples from residences in lower Manhattan.

The SVF results, seen in Table 6, show a broad range of SVF content in the settled surface dust from none detected up to 35% in indoor dust samples and as high as 72% outdoors. The location and range of SVF found in settled surface dust is shown in Figure 11 for all areas sampled in lower Manhattan. SVF was found in indoor dust in 26 of 57 residences sampled as analyzed by PLM (46%). The settled surface dust in common areas contained SVF in 14 of 26 areas sampled (54%), and SVF was found in outdoor dust 11 of 14 times sampled (79%). The majority of SVF detected was glass wool. Only four buildings contained mineral wool in settled surface dust, ranging from 1%–7%. Because no surface-loading information is known, the exact percentage is less informative than its presence. No SVF was found in the comparison homes north of 59th Street by PLM analysis. Therefore the SVF content of settled surface dust in some areas of lower Manhattan is higher than that of the selected comparison areas above 59th Street (Figure 11).

2. Mineral Analysis of Settled Surface Dust

Settled surface dust samples were analyzed for crystalline minerals potentially associated with the material from the collapsed World Trade Centers. The results reported here are expressed as a mass percent of the dust (weight of minerals per weight of dust). Quantification of the minerals in dust is not exact due to variations in particle size and

other methodological interference. Therefore, these data will only be considered semiquantitatively and should be considered estimated values, noted by the "J" qualifier.

Quartz, calcite, portlandite, and gypsum were the most abundant minerals detected (see Table 7). Mica was detected with less frequency than the other minerals, and generally estimated at approximately 0.1%J of the dust. Halite (salt) was also detected at trace levels.

Quartz was detected in all 14 outdoor dust samples, 21 of the 26 common area samples, and only 30 of the 57 samples from residences in lower Manhattan. Neither cristobalite nor tridymite, other forms of crystalline silica, were found in any of the settled surface dust samples. For this discussion quartz will refer to the alphaquartz form of crystalline silica. Levels of quartz were estimated as from 0.05%J to 31%J in residences, and from 0.03%J to 25%J in common areas. Estimated quartz content in outdoor samples ranged from 1%J

Mineral components of concrete (quartz calcite, and portlandite) and other building materials (gypsum, mica, and halite) in the indoor settled surface dust of some areas of lower Manhattan were estimated at higher percentages than for the comparison areas above 59th Street.

to 27%J. Because quartz is a common material (i.e., silica sand), finding this mineral in the city where there is a lot of concrete building material is not unusual. However, quartz in dust from the comparison areas above 59th Street was estimated from nondetect to 1%J in the common areas and from nondetect to 2%J in the residences. Figure 16 shows the locations where the estimated maximum quartz content in settled surface dust was greater than or similar to estimated quartz content in the dust samples from the comparison areas above 59th Street. Fifteen residences, six common areas, and 12 outdoor areas had estimated quartz levels higher than the associated comparison areas above 59th Street. Therefore, quartz was elevated in some indoor areas of lower Manhattan relative to the comparison areas, due to insufficient amounts of dust. Figures 13-16 compare the indoor and outdoor locations in lower Manhattan to the indoor samples above 59th to show the relative distribution of the minerals.

Calcite (calcium carbonate) and portlandite (calcium hydroxide) are also components of concrete. They occurred with similar frequency in the dust samples and were often co-located with the quartz. Calcite ranged from an estimated 0.8%J to 19%J in outdoor areas, and from 0.02%J to 21%J in indoor areas. Portlandite ranged from an estimated 0.07%J to 6%J in outdoor areas and from 0.04%J to 8%J in indoor areas. In contrast, the maximum levels found in indoor comparison areas above 59th Street were 0.9%J calcite and 0.08%J portlandite. Figures 13 and 15 show the locations where the estimated calcite and portlandite content in settled surface dust was higher than to that estimated in dust samples from the comparison areas above 59th Street. For calcite, 13 residence and 9 common area samples had estimated levels higher than those found in the comparison areas above 59th Street. Similarly for portlandite, 17 residence and 9 common area samples were estimated

as higher than values in comparison areas above 59th Street. So although these materials are found in urban areas, perhaps due to erosion of concrete, building construction and maintenance, the estimated percentages of calcite and portlandite were larger in some areas of lower Manhattan relative to the comparison areas above 59th Street (Figures 13 and 15.)

Gypsum (hydrated calcium sulfate) is a major component of dry wall. Gypsum was found in 11 of 14 outdoor settled dust samples at levels estimated as 0.03%J to 27%J. Indoor levels of gypsum ranged from an estimated 0.07%J to 20%J in 23 of 26 common areas and from 0.05%J to 30%J in 45 of 57 residences. These estimated percentages are higher than the estimated maximum of 4%J seen in the comparison areas above 59th Street. Figure 14 shows the locations where the estimated gypsum content in settled surface dust was higher than or similar to the levels in the dust samples from the comparison areas above 59th Street. Nine residence and six common area samples had estimated levels higher than those found in the comparison areas above 59th Street. Gypsum was elevated in settled indoor dust in some areas of lower Manhattan relative to the comparison areas above 59th Street (Figure 14).

The data can also be evaluated in a nonquantitative manner by just inspecting the frequency of detects of mineral in dust for lower Manhattan and the comparison areas. As shown in Table 7, the pattern of prevalence for the minerals appears to be similar in both study areas. In other words, the minerals were detected in both locations with about the same frequency. The data imply that building materials were the source for both areas. This source was likely building materials; however, the magnitude of the impact was greater in areas closer to the World Trade Center.

3. Alkalinity of Settled Surface Dust

Several of the minerals detected in the settled surface dust samples, such as portlandite, can make the dust more alkaline, or raise the pH. Originally, the pH of all of the settled surface dust samples was to be determined. However, enough dust for this analysis was only available at two locations. Both dust samples were alkaline (pH of 8.6 and 9.8). On the pH scale of 0 (acidic) to 14 (alkaline or basic), these values are slightly alkaline. Mineral analysis of these two settled surface dust samples estimate levels of calcite (15%J, 19%J) and portlandite (6%J, 3%J) were present, respectively. These crystalline minerals along with other components of concrete would have contributed to the measured alkalinity.

C. Air

The settled surface dust results discussed previously indicate the materials present in each area sampled. Fibers and small particles within the dust may become airborne during daily activity or during periods of high activity such as cleaning. Therefore, the settled surface dust sample results are an indicator for areas with a potential for airborne contaminants.

Air sampling was conducted to determine the actual levels of fibers, dust, and crystalline minerals in the air at the time of settled surface dust sampling. Where feasible, these results are correlated to the settled surface dust results.

1. Fibers in Air

Samples for airborne fibers were first analyzed by phase contrast microscopy (PCM) to count all fibers present according to the National Institute for Occupational Safety and Health (NIOSH) Method 7400 counting rules. Phase contrast microscopy identifies fibers based on size and shape, but not the specific type of fiber. Therefore, the fibers seen by PCM include asbestos fibers as well as other fibers, such as textiles, fiberglass, and cellulose that might be present. Any sample with airborne fibers above the highest PCM levels found in the comparison locations was re-analyzed by transmission electron microscopy (TEM) and scanning electron microscopy (SEM). The TEM methodology can distinguish what types of fibers are in a sample, particularly asbestos fibers. The SEM method identifies SVF. Some fibers present in the settled dust of the area may become suspended in the air and be present in the air sample.

a. Indoor Air

Fiber levels in air are expressed as the number of fibers in each cubic centimeter (cc) of air or fibers per cc (f/cc). Low levels of fibers were found in the comparison areas above 59th Street homes, where fiber levels ranged from less than 0.001 f/cc (<0.001 f/cc) to 0.003 f/cc. Because no asbestos was detected in the settled surface dust of the comparison areas above 59th Street, these fibers are not likely to have been asbestos. SEM analysis showed they were not SVF. Figure 7 shows all airborne fiber levels determined by PCM in the lower Manhattan area with respect to the fiber levels found in the comparison areas above 59th Street. These data represent all fibers found according to the analytical method used (NIOSH 7400) and are not specific to asbestos or SVF.

The majority of the lower Manhattan air samples from indoor areas without asbestos in the dust were within the range of fibers found in the comparison areas above 59th Street (<0.001 to 0.003 f/cc). Six of these air samples (from 3 common areas and 3 residences) had higher levels of fibers in air than the comparison areas above 59th Street (Table 8). Four of these areas had measurable levels of SVF in the settled surface dust. All six-air samples were re-analyzed by TEM to determine if the fibers present included asbestos fibers. Asbestos fibers were not found in these six air samples (Table 8). A third analysis was done by SEM to determine if SVF fibers were present in those air samples from areas with SVF in the settled surface dust. SVF fibers were found in three of these air samples at concentrations from 0.000037 f/cc to 0.000255 f/cc. Although SEM is a more sensitive method, detecting thinner fibers than PCM, reported SVF levels were well below the PCM method detection limit of 0.001f/cc. SVF fiber concentrations in the comparison areas above 59th Street ranged from none detected to 0.000087 f/cc. Although air samples from comparison areas above 59th Street were not re-analyzed by TEM, no asbestos fibers were noted on the subsequent SEM analysis for SVF. Therefore the six air samples that had total

fiber counts greater than comparison areas above 59th Street were not elevated due to either asbestos or SVF fibers.

There were 15 out of the 83 (18%) indoor areas in lower Manhattan with measurable asbestos in the settled surface dust, 10 residences and 5 common areas (Table 9). Nine of these residences were sampled aggressively (the air exhaust of the vacuum used for settled dust sampling was used to stir up all dust into the air before an air sample was taken). Four of these aggressively sampled residences had measurable fiber levels in air ranging from 0.001 f/cc to 0.003 f/cc. One residence was occupied and could not be aggressively sampled. Air sampling at that location indicated <0.001 f/cc by PCM analysis. Because the levels found were within the range measured in the comparison areas above 59th Street, no TEM or SEM re-analysis was performed on these air samples. Therefore the composition of these fibers is unknown (i.e., asbestos, SVF, cellulose).

None of the common areas were aggressively sampled because all of these areas were open to the public. Of the five common areas with some asbestos in the dust, three had measurable levels of fiber in the air ranging from 0.001 f/cc to 0.002 f/cc. One area in building 5 where there was not enough settled surface dust to sample had an airborne fiber count of 0.002 f/cc. No TEM or SEM re-analysis was performed on these air samples, and the composition of these fibers is unknown (i.e., asbestos, SVF, cellulose).

b. Outdoor Air

Six of the 14 lower Manhattan outdoor locations sampled had detectable levels of asbestos in the settled surface dust, indicating a potential in these locations for asbestos fibers to become airborne. Air sampling indicates three of the locations did not have detectable fibers in the air (Table 9). The air samples from buildings 1, 2, and 5 did have fibers detected at levels from 0.001 f/cc to 0.003 f/cc. Of the 15 locations where no settled surface dust samples were taken, the majority of the results showed <0.001 f/cc in the air, with only 4 samples reported at the detection limit of 0.001 f/cc. Because the levels found were within the range measured in the comparison areas above 59th Street, no TEM or SEM re-analysis was performed on these air samples, and the composition of these fibers is unknown (i.e., asbestos, SVF, cellulose).

2. Minerals in Air

Air samples of different particulate sizes, representing inhalable (PM100), thoracic (PM10), and respirable (PM4 and PM2.5) fractions were collected in selected building locations. Not all size fractions were collected at every location. The samples were analyzed by XRD for the same crystalline minerals seen in the settled surface dust samples. Each of the four size fractions was collected with a separate sampling device. The PM100 fraction collected particles with an aerodynamic diameter of 100 microns and lower. The PM10 fraction collected particles with an aerodynamic diameter of 10 microns and lower and so on. The results reported here are expressed as the mass of mineral per unit volume of air or micrograms per cubic meter of air (μ g/m³). Quantification of the minerals is not exact due to variations in particle size as compared to the particle size used in the standard

curve and other methodological interference. Therefore, these data will only be considered semiquantitative and should be treated as estimated values, noted by the "J" qualifier.

Due to changes in sample volume, the detection limit of minerals in each fraction varied, allowing lower concentrations to be recorded in the PM10 and PM100 fractions compared to the PM4 fraction. Care must be taken when comparing results between fractions for this reason. Additionally, although the PM4 fraction should be a subset of the PM10 and PM100 fractions, PM4 results are sometimes higher than PM10 or PM100 from the same location. Therefore, due to uncertainties in the quantification and general quality assurance/quality control concerns, data will not be associated to a specific building sampling location or correlated between size fractions at the same location.

The comparison air samples, taken in areas above 59th Street, contained no quartz, cristobalite, tridymite, calcite, portlandite, mica, or halite above method detection limits (Table 11). Gypsum was found at least once in all size fractions collected in the comparison areas above 59th Street. Gypsum concentrations were estimated at $3 \mu g/m^3 J$ (PM2.5), $5 \mu g/m^3 J$ (PM4), $3 \mu g/m^3 J$ (PM10), and $3 \mu g/m^3 J$ (PM100) (Table 11).

All of the minerals found in the settled surface dust were seen in at least one air sample from lower Manhattan (Table 11). Quartz, calcite, portlandite, and gypsum were detected at similar frequencies in each of the larger size fractions, PM100, PM10, and PM 4. Table 11 shows the frequency of detection and the range of estimated values for each size fraction sampled. Quartz and gypsum were each detected once in the PM 2.5 fraction. The PM 2.5 size fraction was only collected toward the end of the project, and there are fewer samples for this size fraction.

Approximate quartz levels in air ranged from $4 \mu g/m^3 - 19 \mu g/m^3 J$ in the respirable fraction (PM4), $3 \mu g/m^3 - 12 \mu g/m^3 J$ in the thoracic fraction (PM10) and $3 \mu g/m^3 - 13 \mu g/m^3 J$ in the inhalable fraction (PM100). Detection frequencies were similar in these fractions at 11%, 10%, and 14% respectively. However, quartz was only seen in one of the PM2.5 samples (3%) at an estimated $3 \mu g/m^3 J$. Cristobalite, a different crystalline form of crystalline silica, was found in one lower Manhattan air sample at

Estimated levels of minerals in air, including quartz, calcite, gypsum, and portlandite, appear higher in some areas of lower Manhattan than in the comparison areas above 59th Street.

 $15 \ \mu g/m^3 J$ in the PM100 fraction. Cristobalite was not seen in any other sample, either air or settled surface dust. Across the size fractions, quartz was detected in air samples taken in lower Manhattan; it was not detected in the comparison areas above 59th Street. Therefore, the estimated airborne levels of quartz appear to be higher in some air samples from lower Manhattan than the chosen comparison areas above 59th Street.

Similar observations can be seen in the results for calcite and portlandite. Calcite and portlandite occur at a similar frequency to quartz (Table 11) and tend to be present in the same air samples (Appendix C.) Both minerals are present in PM4, PM10, and PM100 size fractions. Calcite was estimated at $3 \mu g/m^3 J - 14 \mu g/m^3 J$ and portlandite at $14 \mu g/m^3 J - 95$

 μ g/m³J in inhalable dust. Mica and halite are not seen with the same frequency, but are detected in several air samples from lower Manhattan. Calcite, portlandite, mica, and halite are all detected in air samples taken in lower Manhattan, but not in air samples from the comparison areas above 59th Street (Table 11). Therefore, the airborne levels of calcite, portlandite, mica and halite appear to be higher in some air samples from lower Manhattan than in the samples from the comparison areas above 59th Street.

Gypsum was the most common mineral detected in air samples from lower Manhattan for all size fractions and was detected in 40 out of 114 respirable fraction (PM4) air samples at levels ranging from approximately $4 \mu g/m^3 J-15 \mu g/m^3 J$. Gypsum was estimated at $3 \mu g/m^3 J-14 \mu g/m^3 J$ and detected less frequently in the PM10 (33 of 105) and PM100 (24 of 101) size fractions, even though the detection limits for these fractions were lower. Gypsum was found in one of the PM2.5 samples, at approximately $3 \mu g/m^3 J$. Across the size fractions, gypsum was detected in air samples taken in lower Manhattan at higher estimated levels than air samples taken from the comparison areas above 59th Street (up to $5 \mu g/m^3 J$). Therefore, the airborne levels of gypsum seem to be higher in some air samples from lower Manhattan than the chosen comparison areas above 59th Street.

DISCUSSION

This section of the report will provide a more in-depth review of where the various materials were detected. In addition, the public health concerns potentially associated with these materials is also presented.

A. Discussion of Results

Settled surface dust samples obtained from lower Manhattan as part of this investigation contained materials consistent with debris from a building collapse. Both indoor and outdoor settled surface dust samples contained some amount of insulation material (asbestos, glass wool, and mineral wool), components of concrete (portlandite, calcite, and quartz), and components of other building materials (gypsum, mica, and halite). These materials were not present at the same levels in the four buildings above 59th Street sampled for comparison.

Asbestos levels in dust in the lower Manhattan area, ranging from <1% to 3.4%, were consistent with samples taken of outdoor material by EPA after the WTC collapse. SVF levels in the settled surface dust in lower Manhattan were higher than in the comparison areas above 59th Street, ranging from 1% to 72%. No data were available for SVF in lower Manhattan prior to this investigation. Analysis of archived dust samples collected after the WTC collapse by NIOSH, confirms the presence of SVF in bulk samples taken in lower Manhattan.

The minerals found in settled surface dust from residential buildings in lower Manhattan were consistent with minerals reported by the United States Geological Survey (USGS) for outdoor settled surface dust across lower Manhattan after the WTC collapse (9). Thirty-three dust, two concrete, and two steel insulation samples were analyzed by USGS,

including two from an indoor area. Quartz, calcite, and gypsum were reported for all of these samples. Each of these was occasionally reported as a major constituent of the material, i.e., estimated at 20% or higher. Twenty-six of these samples contained mica, usually estimated as trace or less than 5% of the material. Portlandite was only noted in two samples at trace levels, including the one indoor sample analyzed.

1. Association of Asbestos and Synthetic Vitreous Fibers

Based upon the sampling results, it appears that SVF is likely to be found in settled surface dust whenever asbestos is detected in the settled surface dust. All of the six outdoor and five common areas that had asbestos in settled surface dust also had SVF in the settled surface dust (Table 9). Of the ten residences that contained asbestos in settled surface dust, a little more than half also contained SVF. However, this does not hold true in areas without asbestos, where the absence of asbestos does not indicate an absence of SVF. Of the eight outdoor areas without asbestos in the settled surface dust, three contained SVF. Similarly, SVF was found in approximately 25% of the common areas and in approximately 36% of the residences without asbestos in the dust. Although the presence of asbestos in the dust seems to correspond to SVF, the absence of asbestos does not predict or correspond to a presence or absence of SVF in settled surface dust, in either indoor or outdoor areas of lower Manhattan.

2. Association of Asbestos and Synthetic Vitreous Fiber Levels Outside of and Within a Building

Outdoor settled surface dust was collected at 14 buildings in lower Manhattan. One purpose for collecting outdoor dust was to determine if the contents of the outdoor dust at a location would reflect the type of indoor dust. In the case of fibers, there is little correlation in the indoor and outdoor dust composition. High levels of outdoor SVF are associated with both high and low indoor levels. The presence or absence of asbestos in settled surface dust shows a similar lack of correlation. The outdoor settled surface dust samples do not correspond with indoor dust composition for either group of fibers. This lack of association could, in part, be due to whether and how frequent interiors and exteriors had been cleaned.

In contrast, indoor levels of fibers in settled surface dust did seem to correspond within a building. Buildings with elevated SVF levels in the common area tended to have elevated levels in the residences. A similar pattern was seen between the two residences sampled in the same building. For example, SVF was often found in both residences sampled within the same building, or was absent in both residences. Within this limited data set, fiber levels in one area of a building seem to weakly correspond with fiber levels in other areas of the same building. However, none of these data represent total dust loading, only the relative amount of fibers within the dust.

3. Occurrence of Crystalline Minerals

Indoor areas that have elevated levels of quartz in settled surface dust appear to also have elevated levels of calcite, portlandite, and gypsum (see Figure 10). Additionally, these four minerals appear together in all outdoor samples with only minor exceptions. Although only 14 of the 30 buildings had samples taken of outdoor dust, there does seem to be a correlation between elevated outdoor and indoor levels of quartz, gypsum, and the other minerals. It is unknown if this would be seen in outdoor areas above 59th Street for comparison, as not enough outdoor dust was available for analysis.

Most elevated quartz and gypsum levels indoors were found in the first ten buildings sampled during the first 5 days of the project (with the exception of 14% quartz in the common area of building 22). Although many of these buildings were closer to the WTC site, location may not be the only factor contributing to higher mineral levels in the dust. Other buildings sampled in these areas later (e.g., 13, 14, and 28) did not have elevated levels of minerals. Many factors may contribute to this finding including whether the area was cleaned (indoors, outdoors), the cleaning method used, the time since the last cleaning, and the amount of dust that was initially in the area. It is not clear which factors contributed to variation in mineral levels by building.

Within a building, levels of quartz, calcite, and portlandite tended to be similar, and in some cases showed a similar profile in areas with elevated quartz (see Figures 13, 15, and 16). In general, where quartz was greater than 5%, calcite and portlandite were also present, but as smaller proportions of the dust. Gypsum, although often present with the other minerals, varied from being the most abundant to the least abundant in the dust.

4. Buildings Impacted by Asbestos, Synthetic Vitreous Fibers, and Crystalline Minerals

There was no clear pattern of which buildings were impacted by asbestos, SVF, and crystalline minerals. Figure 17 shows the approximate locations of the buildings and the level of contaminants in settled surface dust that were above or below the levels found in the comparison areas. The data are summarized and presented to indicate the indoor areas within a building with higher levels of a material in settled surface dust than was found in the comparison areas above 59th Street (Table 12). In this table, quartz was considered representative of the two other minerals associated with concrete, portlandite and calcite.

In 6 of the 29 buildings (approximately 21%) sampled in lower Manhattan, concentrations of these materials were not detected at levels higher than the levels in the comparison buildings above 59th Street. These six buildings tend to be among the buildings farthest from the World Trade Center and not located along what appear to be major streets or intersections. Map 17 shows the approximate locations of the buildings and the level of contaminants in settled surface dust that were above or below the levels found in the comparison areas.

Twenty-three buildings (approximately 79%) had levels of at least one of these materials in the dust at levels higher than the maximum found in the comparison buildings. Four of these buildings (buildings 4, 5, 6, and 9) contained all four materials above levels seen in the comparison areas above 59th Street in at least one indoor sample (Table 7). Seven of the 23 buildings were only impacted by SVF in indoor areas (buildings 13, 14, 17, 18, 25, 26, and 28). One building was only impacted by asbestos (building 11). The remaining 11 buildings were impacted by some combination of fibers and minerals. No clear pattern emerges in terms of which fibers or minerals occurred in the indoor dust of a building. Three of these building had only the two fiber types present (buildings 15, 24, and 27). One of these building had only the two minerals present (building 8).

The limited number of buildings sampled and the limited number of indoor locations that were sampled make it difficult to draw conclusions about the relationship of geographical location and the number of different materials found in the dust above the levels found in the comparison locations. However, buildings close to the World Trade Center tend to have a greater number of these materials found.

5. Significance of Minerals and Fibers in the Settled Surface Dust

The results previously discussed indicate minerals, asbestos, and SVF are all found at higher percentages in the samples from lower Manhattan when compared to the sampled areas above 59th Street. These findings are based on comparing the percent of each mineral and fiber in the settled surface dust. However, to determine the total mineral and fiber content in a room, the amount of dust present would also need to be known. Because surface loading is not known, we cannot demonstrate that the total amount of mineral and fiber is increased in any given area. However, these data indicate the components and quality of the dust in lower Manhattan is different from that of the comparison areas above 59th Street.

The increased crystalline minerals and fibers in the settled surface dust indicate a potential for contact with these materials. The magnitude of this contact will vary with dust loading (which is unknown) and activity in the area. Residents may come into contact with the dust directly (i.e., skin, eyes). Additionally, the dust may also become re-suspended in the air through activities such as cleaning. In this way the settled surface dust may be a reservoir for material in the air. Therefore settled surface dust results are important both to evaluate direct contact, as well as to evaluate the potential for airborne/inhalation exposures.

6. Significance of Minerals and Fibers in Air

The settled surface dust samples suggest the potential for airborne particles and fibers. The air samples taken indicate that some of these materials are present in the air at measurable levels. Because the air samples were taken over several hours, they provide an understanding of average air levels over the course of the day. Some of these air samples were taken during normal daily activity outdoors in common areas and in residences. They represent the impact of vehicular traffic, walking, and normal activity in the area. As such they are a good measure of average exposure at that time and place. No samples were taken

to assess high levels of activity, such as cleaning, which might generate short-term periods of higher levels of materials in the air. However, about 40% of the residences were aggressively sampled, which gives some indication of the impact of increased activity on airborne levels.

Although semiquantitative, the minerals detected in air provide important information. Quartz and mica, which have potential health effects on the lung, were found in the respirable fraction of the dust (PM4). This indicates that some portion of the quartz and mica found in the settled surface dust is of respirable size. Minerals found in the larger size air fractions, thoracic (PM10) and inhalable (PM100), indicate that larger particles may also be present that could irritate the upper areas of the respiratory tract. It should be noted that these larger fractions also include thoracic and respirable material.

The fiber analysis by PCM provides an upper bound of airborne fibers. Although fiber type is not identified in this analysis, there can be no more asbestos fibers than total fibers detected. Re-analysis of some of these samples shows that very few fibers of public health interest (SVF or asbestos) were present. However, care must be taken when interpreting these results because there were samples taken which showed fibers by PCM, which were not re-analyzed. Any of these air samples, especially those taken in areas with asbestos or SVF in the dust, may have some of these fibers as part of the PCM fiber count. Generally, the levels of dust and air did not indicate a consistent pattern at different locations. It was not possible to conduct a more accurate correlation analysis between asbestos and SVF material in settled surface dust and fibers in air because the majority of air samples were not analyzed specifically for asbestos or SVF (i.e., TEM or SEM analysis). The limited data gathered during this investigation does not support nor refute the assumption that all the fibers in the air samples will be asbestos or SVF even when asbestos was found in dust.

B. Public Health Evaluation

The potential for health effects is possible after exposures to certain materials. An exposure to a material occurs when it comes into contact with or is taken into our bodies. The route of exposure is the method by which a material comes into contact with the body (e.g., dermal contact, inhalation, or ingestion). The nature of the health effect will depend on the material involved, how it comes into contact with the body, the amount of material (exposure level), the duration of the exposure, resultant dose, and the susceptibility of the individual (e.g., pre-existing health conditions may make an individual more sensitive to a contaminant).

Initially, screening values were established and listed in the project-sampling plan. These screening values were established to provide a guideline for evaluating the data as it was collected for any indication of a hazard requiring immediate attention, and to ensure the proper analytical detection limit to support a chronic health evaluation. These screening values were not used in this report because this public health evaluation is specific to the levels of materials found in the study area.

Because levels of asbestos, SVF, and some crystalline minerals are elevated in the settled surface dust of some buildings in lower Manhattan, individuals may be exposed to those materials in their homes, out of doors, or at work either by direct contact or by inhalation when the dust is disturbed and suspended in the air. Depending on the environment, one or all of these materials may be present in the dust of an area.

1. Sensitive Populations

Sensitive populations are individuals who may experience greater responses when exposed to a material than normally predicted. This sensitivity may be due to life stage, pre-existing medical or exposure conditions, or genetic susceptibility.

Children are considered a sensitive population for several reasons. The first consideration is the potential for greater exposure than adults. With a lower body weight, and higher relative ventilation rate, children have higher doses than adults given exposure to the same levels of materials in air, food, or drinking water. For biopersistent materials, such as some of the minerals discussed previously, childhood exposures present a greater lifetime risk due to longer residence time in the lung. This is demonstrated for asbestos where an exposure with onset at 1-year old presents a greater risk than an exposure at age 18 (10). Finally, because they are shorter than adults, children breathe air that is closer to the ground, and they may play in contaminated areas, creating situations where they are impacted more by materials in soil or dust.

Children and adults with pre-existing medical conditions may be more susceptible. Many of the materials found in the areas sampled have irritant effects on skin, mucous membranes, and the respiratory tract. Individuals with pre-existing allergic and cardiopulmonary diseases may be more sensitive to the effects of exposure to this dust. One of the most clinically important and frequent responses of the airways to dust and irritant exposures is bronchoconstriction. Therefore, individuals with pre-existing asthma are much more likely to develop bronchoconstriction in response to exposure to dust particles, fibers, and irritants.

Obstructive lung disease and other respiratory and cardiopulmonary disorders are factors for increased sensitivity. Any individuals in lower Manhattan who received a large dose of dust during or immediately following the WTC collapse may also be more sensitive. Anyone who has been experiencing respiratory symptoms since the collapse should consult their physician for proper treatment.

Individuals who smoke are at greater risk. Smoking can reduce lung function and may cause pulmonary disease that would make individuals more sensitive. Additionally, smoking has been shown to synergistically increase the risk of developing lung cancer from asbestos exposures. Smoking also increases the risk of developing lung cancer in workers exposed to crystalline silica (e.g., quartz).

2. Soluble Minerals: Gypsum, Portlandite, and Calcite

The potential health effects of gypsum (hydrated calcium sulfate), portlandite (calcium hydroxide), and calcite (calcium carbonate) are similar as these minerals are similar chemically and physically. All three minerals are crystalline in form and soluble (11,12). Because these minerals are soluble in

body fluid, they are easily cleared from the respiratory tract and

Although possible skin, eye, and respiratory tract irritants, the soluble minerals gypsum, portlandite, and calcite do not pose any hazard for long-term health effects.

lungs when inhaled (13,14,15). Therefore, no long-term respiratory effects have been observed, even at the very high exposures that occur during the mining and processing of these minerals (16,17).

Calcite, and portlandite increase the alkalinity of a fluid. Therefore, when these materials come into contact with tissue, they can cause irritation of skin and mucous membranes due to increased alkalinity. Any anhydrous gypsum in settled dust would be hygroscopic, which would be drying and irritating to the skin and eyes. Similarly, inhaled material may irritate or burn the respiratory tract (nose, throat, and lungs). The signs of an inflammatory response include redness, swelling, heat, and pain. Reaction to a skin irritant usually occurs at the point of contact between the skin and the substance. Eye irritants can cause redness, tearing, and itching of the eyes. Respiratory irritants can affect the nose, throat, and lungs. They may cause an itchy, watery nose, a scratchy or sore throat, coughing, chest tightness, wheezing, or difficulty breathing. All of these irritant effects are generally temporary and will begin to resolve once exposures cease.

Occupational exposure guidelines (NIOSH) and standards (OSHA) for these minerals were developed to prevent these irritant symptoms in most workers for a 40-hour workweek exposure (13,14,15). Although no guidelines or standards are available for skin contact or residential exposures, air exposure levels are established for both total and respirable minerals in air (see the following table). The airborne mineral levels measured in lower Manhattan are orders of magnitude below these occupational values. Although the occupational standards do not account for sensitive individuals or for extended periods of exposure, they provide a point of comparison. The levels of minerals seen in airborne dust of lower Manhattan are not at the levels known to be irritating to healthy adult workers. However, it is possible that sensitive individuals may feel some irritation. Additionally, direct contact with the dust—or shorter periods of higher air concentrations caused by disturbing the dust—may result in more irritation than would be predicted by the reported airborne mineral levels.

Mineral	NIOSH REL (µg/m ³)	OSHA PEL (µg/m ³)	Maximum Estimated Value in Lower Manhattan (μ g/m ³)
Gungum	10,000 (total)	15,000 (total)	14J (PM100)
Gypsum	5,000 (resp)	5,000 (resp)	15J (PM4)
Portlandite	5,000	15,000 (total)	95J (PM100)
Portialidite	3,000	5,000 (resp)	84J (PM4)
Calcite	10,000 (total)	15,000 (total)	14J (PM100)
Calche	5,000 (resp)	5,000 (resp)	10J (PM 4)
NIOSH = Na	tional Institute for Occu	pational Safety and Hea	alth, Centers for Disease
Control and	Prevention		
REL = recon	nmended exposure level	/limit	
OSHA = Occ	cupational Safety and He	ealth Administration	
	ssible exposure limit.		
$\mu g/m^3 = micr$	ogram of mineral per cu	bic meter of air	
total = all air	borne particles (could in	nclude particles >100 n	nicrometers)
resp = respira	able fraction (PM4)		

3. Mica

Mica is a group of water-insoluble silicate minerals (18). The mica detected in these dust and air samples of lower Manhattan was muscovite. Mica has been shown to cause pneumoconiosis in workers exposed to high levels during mining and processing operations (19,20). Although workers may have also been exposed to quartz in some of these studies, pneumoconiosis in workers with only mica exposures has been documented. (21,22)

Respirable mica particles that deposit in the deep lung may be cleared slowly due to the relative insolubility of the mineral. Over time, the presence of the mineral will irritate the lung tissue, generate immune response, and result in scarring or fibrosis (23,24). The magnitude of this response is believed to be a function of both the dose and duration of exposure. The current worker standard set at $3,000 \ \mu g/m^3$ is a level at which mild pneumoconiosis was seen in some workers only after extended exposure (18 years).

The airborne levels of mica seen in lower Manhattan with an estimated maximum of $43 \ \mu g/m^3 J$ in respirable dust are well below levels of mica that are associated with the development of pneumoconiosis in workers (25). Although disturbing settled surface dust might generate short-term exposures above the measured values, it is the cumulative exposure that is believed to predict health effects (23). Given the low airborne levels of mica seen in lower Manhattan and the low frequency of detection in both air and settled surface dust samples, no health effects are expected from mica exposure alone.

Mica and halite, although elevated in lower Manhattan relative to the comparison areas above 59th Street, are not likely to contribute to any adverse health effects.

4. Halite

Halite is simply sodium chloride, or rock salt (18). This crystalline form of the salt will readily dissolve in the body and is not expected to have any adverse health effects. Although large changes in salinity can disrupt tissues, the trace amounts of halite and the low frequency of detection do not indicate any potential health concern.

5. Crystalline Silica (Quartz and Cristobalite)

Silica is chemically inert for the most part, so it is the physical form of silica that imparts its toxicity. Small crystals of silica may be deposited in lung tissue. Once deposited, the crystals remain and fibrogenic lesions may appear over time, damaging the lung tissue and reducing lung capacity (26). Silica has been demonstrated to be a pulmonary toxin linked both to silicosis (a fibrotic lung disease) and lung cancer (27,28,29,30). These findings are supported by both human and animal studies. Chronic silica exposure may contribute to lung cancer; emphysema; obstructive airway, etiology, immune, or connective tissue disease; and lymph node fibrosis (27,30,31,32,33).

a. Silicosis

Generally silicosis is seen after years of high-level occupational exposures, although it can result from short-term acute exposure (26,30). It is not known if chronic ambient environmental exposure contributes to silicosis (26). Because the lung damage is proportional to the total amount of silica deposited in the lung, current evaluations are based on a measure of cumulative exposure of air concentration over work-years of exposure (28,34,35).

Short-term exposures to quartz, even for a continuous year of exposure at the highest estimated air concentration, is not expected to result in any adverse health effects.

Assuming worst-case theoretical assumptions, the estimated quartz levels measured cannot rule out adverse health effects from chronic exposures (i.e., 30 years). For individuals who conduct frequent cleaning of their residences, as recommended in this report, or participate in the U.S. Environmental Protection Agency cleaning/sampling program, it is unlikely that their exposure would resemble these worst-case conditions.

Exposures of less than 1 milligram per cubic meter 1 (mg/m³) of air work-year^{*} of exposure are believed to not result in significant incidence of silicosis in workers (28,34,35). The peak respirable quartz level estimated in lower Manhattan of 19 μ g/m³J would be

^{*} Cumulative exposure is expressed as a constant air concentration over a specified period of time. If a worker is exposed to a constant concentration of 1 mg/m^3 of quartz throughout the workday, for an entire year, that is equivalent to $1 \text{ (mg/m}^3)$ work-year. (One milligram per cubic meter of air is 1 mg/m^3).

equivalent to 0.08 (mg/m³)-work year.[†] Therefore, at this measurement even an entire year of exposure 24 hours a day 7 days a week would not be expected to result in silicosis when compared to work environments known to result in silicosis.[‡]

b. Lung Cancer

The International Agency for Research on Cancer (IARC) has determined that there is sufficient evidence that crystalline silica in the form of quartz or cristobalite is carcinogenic in humans (28). The IARC cites only limited evidence for the carcinogenic potential of tridymite from animal studies and inadequate evidence in either humans or animals to classify amorphous silica. Respirable crystalline silica is listed as a known human carcinogen in the National Toxicology Program's ninth report to Congress (27).

Although the exact mechanism for silica-induced lung cancer is not known, it is believed cancer incidence is linked to the development of other silica-induced lung disease (34). It is possible the mechanisms involved in the development of silicosis are linked to the carcinogenic process through the production of reactive oxygen species and tissue responses to immune signals and fibrotic changes (36,37). Some researchers have developed dose cancer risk estimates from epidemiologic studies by building off risk estimates for silicosis (34,38,39). A risk assessment produced for both silicosis and lung cancer on the basis of crystalline silica exposures reviews major studies and proposes that lung cancer incidence may be expected to increase by 14% for each additional 1 (mg/m³) work-year exposure above the baseline of 1 (mg/m³) work-year exposure (36).^{*}

The maximum estimated detection of quartz (crystalline silica) in air in lower Manhattan was 19 μ g/m³J. Using the above risk estimates, the cumulative exposure for 1-year continuous exposure 0.08 (mg/m³) work-year poses negligible increased cancer risk. Although there are uncertainties in this evaluation, current scientific opinion would indicate short-term exposures to the measured quartz levels do not pose a health hazard.[†]

If the highest measured airborne levels are representative of long-term conditions, the cumulative exposure for a continuous 30-year residence would exceed the $1 \text{ (mg/m}^3)$ work-year cumulative exposure which is believed to be the departure point for adverse effects

[†] In order to compare a continuous residential exposure to a workday exposure, the estimated air level was adjusted to be comparable to a shorter workday exposure across the entire year.

[‡] It should be noted that children might be at slightly higher risk than adults in sensitive populations. However, the estimated 0.08 (mg/m³) work-year exposure is significantly below the departure threshold of 1 mg/m³ per work-year exposure, and therefore no adverse effects would be expected.

^{*} This estimate was derived by looking at cancer incidence studies and by relating the incidence of lung cancer to silicosis risk estimates. Both methods provided similar risk estimates. However it should be noted other authors caution about directly linking silicosis and lung cancer risk estimates (39).

[†] There are many uncertainties about whether this risk would be realized. The first is that even though the risk of cancer is not linear, the risk estimates treat it as linear (36). Therefore there may be an overestimate of risk at low exposures compared with the studied populations. The air levels of silica we are examining here are well below those studied in worker populations; therefore even this small increased risk may be overestimated. Conversely, the studied populations were adult workers. So risk estimates based on these studies may underestimate risks to children and to adults with respiratory disorders.

for both silicosis and lung cancer. Therefore these data cannot rule out the potential for long-term risk. However, the highest airborne level measured during this study is not necessarily representative of long-term exposure. In addition, the XRD analysis for quartz in air is semiquantitative (i.e., an estimated value). This adds additional uncertainty to the risk estimation. As presented in the discussion section of this report, it is expected that the ongoing cleaning of lower Manhattan and natural dispersion should reduce these materials over time, thus reducing long-term risk. Therefore cleaning should be continued and levels of respirable crystalline silica monitored, until the data demonstrate that long-term risk of adverse health effects is minimal.

6. Synthetic Vitreous Fibers

The term synthetic vitreous fibers (SVF) refers to several types of manufactured vitreous fibers: glass wool, mineral wool, and refractory ceramic fibers (RCF) (41,42). Because RCF was not found during this investigation, its health effects are not discussed. The types of SVF found in settled surface dust, glass wool and mineral wool, are primarily skin and upper respiratory tract irritants. Although some data do indicate a potential for pulmonary effects, no adverse effects on the lung are expected from the levels measured in this investigation.

a. Irritant Effects

The primary short-term and acute health effects of SVF are skin, upper respiratory tract, and eye irritation (42). It is the physical shape and size of the SVF fibers that cause irritation, generally fibers more than 5 microns in diameter cause skin irritation (41,43,44). The affected area may itch and a rash may form. These symptoms subside as exposure is ended, as the fibers slowly work themselves out of the affected tissue. Upper respiratory tract irritation due to inhaled fibers has been reported when fiber concentrations were greater than 1 f/cc in air (41,42,45).

No long-term health effects are expected from the skin and eye irritation that may have occurred because of exposure to synthetic vitreous fibers.

No adverse pulmonary health effects are expected from the levels of airborne synthetic vitreous fibers measured in this investigation.

Generally this type of irritation has been observed in individuals who are manufacturing and installing SVF insulation materials. Although the settled surface dust results indicate SVF is present in many areas of lower Manhattan, total dust loading is not known. Because these irritant health effects would be due to the total amount of SVF contact exposure, dust loading as well as activity levels would influence whether a particular environment was irritating. Extremely low levels were detected in the air samples analyzed via SEM. In the PCM analysis, even if all of the fibers counted are assumed to be SVF, the detected maximum value of 0.003 f/cc is well below the documented level of irritation. These data indicate that SVF is present at elevated levels in settled surface dust and may be an irritant under certain conditions, or to sensitive individuals. No long-term health effects are expected from the skin, eye, or upper respiratory tract irritation related to the SVF exposure.

b. Pulmonary Effects

SVF less than 3 microns in diameter are respirable and available to enter and deposit in the pulmonary regions of the lung (41). Clearance of these fibers from the lung will be determined by fiber solubility and length (41,42). Fibers cleared from the lung have less potential to create long-term health effects.^{*} Although some animal studies have demonstrated both fibrotic and carcinogenic potential for fibrous glass and mineral wools (41,42,46), more recent studies do not fully support this finding.[†] Epidemiologic studies on workers exposed to fibrous glass do not provide consistent evidence of pulmonary effects, although some effects were noted (42,47). Similarly, when assessing deaths due to lung cancer in workers exposed to glass wool, studies do not provide strong evidence for increased risk of cancer deaths attributable to glass wool fiber exposures.

The carcinogenic potential of fiberglass has been reviewed by several agencies. The IARC originally classified both glass and mineral wool fibers as Group 2B carcinogens, possibly carcinogenic to humans, based on animal studies (50). Similarly these materials were classified as carcinogens by the National Toxicology program and the American Conference of Governmental Industrial Hygienists (41,49). However a review of the carcinogenic potential of these fibers by IARC in 2001, which takes into account updated human studies, animal inhalation studies, and mechanistic studies, recommends a change in this classification. The IARC has announced that the recent monograph designates both glass and mineral wool as Group 3, unclassifiable as to carcinogenicity in humans, because of inadequate evidence of carcinogenicity in humans and the relatively low biopersistence of the materials (50).

Although both glass wool and mineral wool have been found at elevated levels in the settled surface dust in lower Manhattan, airborne levels are extremely low, and well below a level associated with the worker exposures reviewed in this report. No adverse pulmonary health effects are expected from the airborne SVF levels measured in this investigation.

^{*} Less soluble materials have a longer residence time in the lung and therefore have a greater potential to contribute to tissue damage or malignant disease. Within SVF types, glass fibers and slag wool are considered the most soluble, and therefore least toxic. Mineral wool is less soluble than glass wool. Fibers longer than approximately 15 microns, the diameter of a macrophage, are also cleared less efficiently, leading to a greater potential for adverse effects (26).

[†] Early studies often relied upon injection and implantation studies, which may not accurately predict a pulmonary response from inhalation exposures. A review of inhalation studies indicates that glass wool did not cause pulmonary fibrosis or lung cancer in these animal studies (47). A recent study by Hesterberg indicates no increase in pulmonary fibrosis or lung cancer even at doses of 222 f/cc, although cancer incidence in control animals was considered high (51).

7. Asbestos

Asbestos refers to a group of natural fibrous silicate minerals of various structures (18). Used primarily for its fire resistant properties and as an insulation material during the 1900s, asbestos is now known to cause pulmonary damage, lung cancer, and mesothelioma. Asbestos fibers less than 2 microns in diameter and less than 5 microns in length with a 3:1 aspect ratio may enter and deposit in the alveolar and pleural regions of the lung (10). Material that is not cleared by the normal processes of the lung will remain, resulting in possible damaged tissue, fibrotic lesions, and cancer (10,52–54). Fibers may also migrate from the lung to the pleural lining, causing mesothelioma. This discussion will focus on the most studied health effects, asbestosis and cancer. It should be noted that the chrysotile asbestos found in settled surface dust during this investigation is believed to be cleared from lungs to a greater degree than other forms of asbestos (10). Therefore the following review, which considers all forms of asbestos, would be conservative for chrysotile asbestos exposures.

a. Asbestosis

Asbestosis is similar to other mineral pneumoconiosis of the lung. Fibers are deposited in the lung and remain due to their insoluble nature and the inability of lung macrophages to engulf and remove longer fibers (>5 microns in length) (10,54). These fibers can create localized tissue damage, stimulate the immune system, and create a scarring of the lung.

Asbestosis has been observed in workers exposed over several years to levels much higher than seen environmentally. It is believed the cumulative exposure to asbestos correlates to the development of asbestosis, and some cases of asbestosis have been reported with only a brief high exposure documented (55). Because cumulative asbestos exposure is believed to be the appropriate metric for assessing asbestos health effects, it is possible long-term, continuous low-level exposures could contribute to asbestosis. Recent studies of a population exposed to tremolite asbestos in Libby, Montana, do indicate increased pleural anomalies, consistent with asbestosis, for individuals who were not exposed occupationally. Individuals who lived with plant workers or those who had contact with vermiculite piles containing tremolite had increased odds ratios for these anomalies (56).

Although asbestos is present in some settled surface dust in lower Manhattan, aggressive sampling produced a maximum level of 0.003 f/cc (PCM). This level is not high enough to result in asbestosis after short exposures. Although it is theoretically possible that lower level environmental exposures could contribute to disease over extended exposures, the data from this pilot study are too limited to support that finding.

Short-term exposures to asbestos, even for a continuous year of exposure at the highest estimated air concentration, are not expected to result in any adverse health effects.

Assuming worst-case theoretical assumptions, the highest fiber levels measured cannot rule out adverse health effects from chronic exposures (i.e., 30 years), but it is unlikely levels will remain elevated for that long. For individuals who conduct frequent cleaning of their residences, as recommended in this report, or participate in the U.S. Environmental Protection Agency cleaning/sampling program, it is unlikely that their exposure would resemble these worst-case conditions.

b. Lung Cancer and Mesothelioma

Asbestos exposure is known to increase lung cancer incidence, particularly for high-level occupational exposures over an extended period of time (e.g., decades) (10,48,52). Additionally, some studies have indicated increased lung cancer and mesothelioma due to high naturally occurring levels of asbestos or low dose environmental exposures (57–60). Longer asbestos fibers can penetrate into the deeper portions of the lung. Because asbestos is relatively insoluble, longer fibers cannot be physically removed by the body's immune systems and may remain in the lung and cause damage. Although the exact mechanism for lung cancer is not known, it is believed the localized immune response to the asbestos may contribute to cancer. Fibers may also migrate out of the lung and cause mesothelioma, a cancer specific to the lining around the lungs. In general, shorter fibers (less than 5 microns) are more efficiently cleared from the lung and therefore are less potent (10). Additionally, there is some indication that chrysotile asbestos is less potent than other forms of asbestos due to better clearance from the lungs.

EPA has established a cancer slope factor for asbestos to estimate risk in the general population due to chronic environmental exposures to asbestos (48).^{*} Factors that influence excess cancer estimates are age at time of exposure, sex, and cigarette smoking (52).[†] EPA's Integrated Risk Information System (IRIS) is used to develop risk estimates for comparison, with the understanding that smokers will be at higher risk.

Air samples from several indoor areas with asbestos fibers detected in the settled surface dust did contain fibers as described by PCM analysis (0.001 f/cc to 0.003 f/cc).[‡] Because

^{*} This slope factor is based on epidemiologic studies and assumes a linear dose response for cancer, even at very low environmental doses (0.23 excess cancers per million people exposed for each f/cc of asbestos.) This type of cancer risk estimate may over predict risk and does not take into account any lower threshold for effect, which may exist. Additionally, the EPA slope factor was designed to apply equally to all types of asbestos, and all fibers greater than 5 microns in length (IRIS).

[†] Variability in these factors is not accounted for in the EPA slope factor because it does not include synergistic effects with the effects of smoking (48).

[‡] However, because asbestos as a percentage of settled surface dust was elevated, an area with significant dust may be capable of generating higher airborne levels of asbestos, especially during high activity.

none of these air samples were analyzed by TEM, it is not known if they were asbestos fibers. For this analysis of a worst-case exposure, it will be assumed they are asbestos. Four of these samples were taken in common areas during normal daily activity. All other air samples, including the highest, 0.003 f/cc, were taken in residences that were aggressively sampled. Using the IRIS slope factor, no appreciable increase in cancer risk is seen when considering a 1-year continuous exposure at the highest fiber level cited $(1 \times 10^{-5} \text{ excess cancer risk})$.

If the highest measured PCM levels are assumed to be all asbestos fibers (a worst-case assumption) and were representative of long-term conditions (another worst-case assumption), the cumulative exposure for a continuous 30-year residence would result in a greater risk estimate for lung cancer and mesothelioma. Therefore these data cannot rule out the potential for long-term risk. However, the highest airborne level measured during this study is not necessarily representative of long-term exposure nor is the highest level necessarily all asbestos fibers. As presented in the discussion section of this report, it is expected that the ongoing cleaning of lower Manhattan should reduce levels of these materials over time, thus reducing long-term risk. Therefore cleaning should be continued and levels of asbestos monitored, until the data demonstrate long-term risk of lung cancer and mesothelioma are minimal.

8. Multiple Material Exposures

Each of the materials discussed previously may exist singly or in combination with other materials found in lower Manhattan. The combinations of materials found in this investigation are described in the discussion of results. When exposures to multiple materials occur together, there is some potential that the agents may interact with respect to their adverse effects. Agents may be additive if they impact an organ system similarly. Synergistic and antagonistic effects may also be seen if the agents either potentiate or disrupt the effects of other agents.

a. Irritant Effects

The effects of the alkaline minerals are most likely additive, although it should be noted that not all of the materials are equally caustic. Therefore, cumulative effects on pH would be the best measure of irritant effects based on the caustic nature of the dust. There may also be potentially additive or synergistic effects with the irritant effects of the SVF. The ability of either caustic action or the mechanical action of the fibers or minerals to degrade the protective properties of skin and mucus membranes may make tissue more susceptible to irritant effects. Additionally, because both types of actions may trigger localized immune responses, these irritants may potentiate each other.

b. Noncancer Pulmonary Effects

Several of the minerals detected contribute to fibrotic lung damage and may result in several diseases that reduce lung function including silicosis, asbestosis, other mineral fibroses, emphysema, and other respiratory disorders. The mechanisms and pathology of

these conditions are similar, and the effects of these agents may be considered at least additive. However, each material has a different toxicity based on the mineral properties of that material. Less biopersistent materials, such as SVF, have the least long-term toxicity, and more biopersistent material, such as asbestos, have greater long-term toxicity. In general, the order of toxicity may be SVF < mica < silica < asbestos, based on solubility and availability for clearance. No factors currently exist to combine risk estimates for these different materials.

CONCLUSIONS

On the basis of the analytical results of samples taken between November 4 through December 11, 2001, and information collected during the sampling effort, the New York City Department of Health and Mental Hygiene and the U.S. Agency for Toxic Substances and Disease Registry make the following conclusions (not in order of importance).

• Total fiber counts of air samples taken in lower Manhattan were similar to the comparison areas above 59th Street sampled during this investigation. The six lower Manhattan areas that had elevated total fiber counts were re-examined by transmission electron microscopy (TEM) and scanning electron microscopy (SEM). The TEM and SEM results indicated that neither asbestos nor SVF contributed to the elevated fiber counts.

• Low levels of asbestos were found in some settled surface dust, primarily below Chambers Street. Many of the lower Manhattan locations sampled had been previously cleaned prior to this investigation. No asbestos was detected in the comparison indoor dust samples taken north of 59th Street. The City of New York has conducted follow-up activities at the locations where asbestos was detected in settled surface dust. Only two follow-up locations, outdoor areas, required professional asbestos abatement. Following-up activity at the other locations did not find any asbestos containing materials.

• When compared with the results obtained from the comparison sampling locations, the lower Manhattan residential areas sampled by this investigation tended to have a greater percentage of SVF (primarily fiberglass), mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite) in settled surface dust. However, the frequency of detections and prevalence patterns of these minerals are similar in both residential areas.

• Exposure to significant amounts of synthetic vitreous fibers (SVF), mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum) may cause skin rashes, eye irritation, and upper respiratory irritation, all of which have been voiced as concerns by citizens and first responders. If the reported irritant effects are associated with World Trade Center related materials, these effects will subside once exposure to SVF, mineral components of concrete, and mineral components of building wallboard end. Some

people with pre-existing heart or lung disease (e.g., asthma) or a previous history of very high levels of exposures (occupational) to SVF, mineral components of concrete, and mineral components may be more sensitive to the irritant effects of SVF, mineral components of concrete, and mineral components of building wallboard.

• Sometimes mineral components of concrete (calcite and portlandite) and mineral components of building wallboard (gypsum, mica, and halite) were detected in air samples at higher estimated levels in lower Manhattan residential areas than in samples taken at comparison residential areas above 59th Street. These detected mineral levels are orders of magnitude below occupational standards. Although the occupational standards do not account for sensitive individuals or extended periods of exposure, they provide a comparison to an established health guidance value. The levels of minerals seen in airborne dust do not pose potential health hazards even for a continuous year of exposure at the highest levels detected.

• Some settled surface dust could become airborne if disturbed. Therefore, people could potentially inhale the asbestos, SVF, mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite) found in settled surface dust of some lower Manhattan residences. Because we did not determine the weight of dust present in the areas sampled, it is not possible to determine whether any particular residence had an elevated dust loading. Appropriate continued frequent cleaning should minimize exposures.

• Several worst-case assumptions were made in order to assess the potential longterm public health risks of airborne asbestos and quartz. Some of the assumptions were that no cleaning of indoor spaces has occurred or will occur, all fibers found in air were asbestos fibers, and the highest levels detected last fall in air represent long-term air levels. Using these worst-case assumptions, prolonged exposure (decades) to airborne asbestos and quartz *may* increase the long-term, theoretical risk of people developing lung cancer and other adverse lung health effects (more than 1 additional case in 10,000 people exposed). For individuals who conduct frequent cleaning of their residences, as recommended in this report, or participate in the U.S. Environmental Protection Agency cleaning/sampling program, it is unlikely that their exposure would resemble these worst-case conditions.

• A review of the building sampling results from this investigation indicates that there is not a consistent spatial distribution pattern of asbestos, SVF, mineral components of concrete, and mineral components of wallboard in air and settled surface dust. This indicates that the materials are heterogeneously distributed. There are many factors that may contribute to the heterogeneous distribution, including whether the area was cleaned (indoors and outdoors), cleaning method, date since last cleaning, and how much dust was initially in the area. It is not clear which factors contributed to this trend. • Results from this investigation do not necessarily reflect conditions that would be found in other buildings, at other times just following the collapse, or after the sampling period. The measurements reflect conditions present at the time of the sampling (November 4–December 12, 2001) in the buildings and areas sampled. The limited number of results obtained from the comparison areas above 59th Street may or may not reflect the New York City-specific background levels of asbestos, SVF, mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite).

RECOMMENDATIONS

On the basis of the conclusions of this investigation, the New York City Department of Health and Mental Hygiene and the U.S. Agency for Toxic Substances and Disease Registry make the following recommendations.

• Because more asbestos, synthetic vitreous fibers (e.g., fiberglass), mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite) were found in settled surface dust in lower Manhattan residential areas when compared to comparison residential areas above 59th Street, the New York City Department of Health and Mental Hygiene and the U.S. Agency for Toxic Substances and Disease Registry are recommending that people continue to conduct frequent cleaning with HEPA vacuums and damp cloths/mops to reduce the potential for exposure.

• To ensure that the recommended frequent cleaning is effective and to ensure that the health of people of New York City is protected, the New York City Department of Health and Mental Hygiene and the U.S. Agency for Toxic Substances and Disease Registry are recommending additional monitoring of residential areas in lower Manhattan. In addition, an investigation should be conducted to better define background levels specific to the city of New York for asbestos, synthetic vitreous fibers, mineral components of concrete (quartz, calcite, and portlandite), and mineral components of building wallboard (gypsum, mica, and halite).

• Lower Manhattan residents concerned about possible World Trade Centerrelated dust in their residential areas can request cleaning and/or testing from the U.S. Environmental Protection Agency (EPA) by logging on to the agency's World Trade Center Web page at www.epa.gov/wtc or by calling the EPA hotline at 1-877-796-5471.

PUBLIC HEALTH ACTION PLAN

The Public Health Action Plan (PHAP) for the World Trade Center response contains a description of actions to be taken by the New York City Department of Health and Mental Hygiene (NYC DOHMH), the U.S. Agency for Toxic Substances and Disease Registry (ATSDR), and/or other government agencies. The purpose of the PHAP is to ensure that this investigation not only identifies potential public health hazards, but also provides a plan of action designed to mitigate and prevent possible adverse human health effects resulting from exposure to hazardous substances in the environment.

A. Public Health Actions Taken

Shortly after the collapse of the World Trade Center towers, NYC DOHMH staff, along with other governmental and private organizations, provided respirators and fit checking for people participating in the recovery operations.

Beginning September 13, 2001, NYC DOHMH staff coordinated daily conference calls with the various governmental and private organizations that were conducting environmental sampling at and around the World Trade Center. These coordination calls ended in January 2002.

On September 15, 2001, the U.S. Department of Health and Human Services, the U.S. Environmental Protection Agency, and the U.S. Department of Labor formed the World Trade Center Environmental Assessment Working Group. The purpose of the group is to coordinate public health and occupational sampling and data review among the three federal agencies in support of the New York City Department of Health and Mental Hygiene and the New York State Department of Health. Other governmental (e.g., New York State Department of Health, New York State Department of Environmental Conservation, New York City Department of Health and Mental Hygiene, and the New York City Department of Environmental Protection) and private organizations (unions, research institutions, companies) participated on the Working Group.

Shortly after September 11, 2001, and through June of 2002, the U.S. Environmental Protection Agency (EPA), the Occupational Safety and Health Administration, the New York State Department of Environmental Conservation, the New York State Department of Health, and the New York City Department of Environmental Protection conducted frequent ambient environmental and occupational monitoring of many different types specifically for the World Trade Center response.

From September 18 through October 4, 2001, the National Institute for Occupational Safety and Health, Centers for Disease Control and Prevention, conducted occupational sampling and evaluation of the conditions at the World Trade Center.

From November 4 through December 11, 2001, NYC DOHMH and ATSDR, with assistance from the U.S. Public Health Service Commissioned Corps, conducted limited sampling of residential areas in lower Manhattan.

On February 4, 2002, NYC DOHMH and ATSDR provided the public with the preliminary findings (asbestos air and dust and synthetic vitreous fibers dust sampling results) of the limited residential sampling investigation conducted in November and December of 2001. Various public meetings were conducted to provide the public with the most relevant information.

On February 13, 2002, EPA formed the Indoor Air Task Force to help address the remaining indoor air quality issues in lower Manhattan.

On March 8, 2002, New York City created the lower Manhattan Air Quality Task Force.

On May 8, 2002, EPA and its federal, state, and city partners announced a comprehensive plan to ensure that residences impacted by the collapse of the World Trade Center have been properly cleaned.

On June 1, 2002, lower Manhattan residents concerned about possible World Trade Center-related dust in their residential areas could begin to request cleaning and/or testing from EPA by logging on to the agency's World Trade Center Web page at www.epa.gov/wtc or by calling the EPA hotline at 1-877-796-5471.

B. Public Health Actions Planned

During the summer of 2002, EPA and its federal, state, and city partners will conduct additional sampling of settled surface dust and air of residential areas not impacted by the collapse of the World Trade Center towers. The purpose of this investigation is to better define the New York City-specific background levels of asbestos, synthetic vitreous fibers, and quartz. Mineral components of concrete (calcite and portlandite) and mineral components of building wallboard (gypsum, mica, and halite) will also be investigated.

During the summer of 2002, EPA and its federal, state, and city partners will conduct an investigation to determine the effectiveness of various cleaning methods to remove World Trade Center-related material.

In late summer/early fall 2002, EPA will begin to conduct cleaning and/or testing of residential areas requested by lower Manhattan residents.

The New York City Department of Health and Mental Hygiene and the U.S. Agency for Toxic Substances and Disease Registry will establish a unified World Trade Center Exposure Registry of people who may have been exposed to harmful substances resulting from the collapse of the World Trade Center towers.

The New York City Department of Health and Mental Hygiene and the U.S. Agency for Toxic Substances and Disease Registry will continue to work with other federal, state, and local agencies to provide information on the sampling results to the citizens of New York and to help interpret the possible health effects of contaminants found.

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World Trade Center Environmental Assessment Working Group

Formed on September 15, 2001, the World Trade Center Environmental Assessment Working Group is made up of representatives from the U.S. Department of Health and Human Services, the U.S. Environmental Protection Agency, and the U.S. Department of Labor. Other governmental (e.g., New York State Department of Health, New York State Department of Environmental Conservation, New York City Department of Health and Mental Hygiene and New York City Department of Environmental Protection) and private organizations (unions, research institutions, companies) participated on the Working Group. The purpose of the group is to coordinate public health and occupational sampling and data review among the three federal agencies in support of the New York City Department of Health and Mental Hygiene and the New York State Department of Health. TABLES

General Info	Table 1 General Information on the World Trade Center Towers										
Height	Tower One: 1,368 feet	Tower Two: 1,362 feet									
Weight	1.5 million tons										
Statistics	 110 floors 6 basements (with 2 subway stations and PATH train station) 1 underground mall 43,600 windows 97 passenger elevators and 6 freight elevators 300 computer mainframes Air conditioning had 60,000 tons of cooling capacity 										
Rentable Space	43,200 square feet per floor	10,000,000 total square feet									
Owners and Operators	Port Authority of New York and	d New Jersey									
Groundbreaking	August 5, 1966										
Opening Ceremony	April 4, 1973 (Parts were opened between 1970–1973)										
Construction Materials	Steel: 200,000 tons										
	Concrete: 425,000 cubic yards										
	Glass: 600,000 square feet	uare feet									
	Electric cables: 12,000 miles										

Table 2 2000 Census Block Level Population									
Distance From WTC	Total Population	Housing Units							
One-Quarter Mile	6,965	4,617							
One-Half Mile	17,596	10,768							
Three-Quarter Mile	29,667	16,482							
The 16-acre site of the World T	Trade Center was buffered and a	a serial proportion technique							

The 16-acre site of the World Trade Center was buffered, and an aerial proportion technique was used to estimate the number of persons living in the area.

Table 3

Number of Buildings and Areas Within Buildings Sampled for Settled Surface Dust: Analyzed for Asbestos by PLM^a and TEM^b, SVF^c by PLM, and Minerals by XRD^d Due to available sample size and analytical objectives, some samples were not analyzed by each method.

	Lower Manhattan	Above 59th Street
Buildings	29	4
Residential Units	57	5
Common Areas	26	3
Outdoors	14	0

Table 4

Number of Buildings and Areas Within Buildings Sampled for Fibers in Air: Analyzed for Total Fibers by PCM^e, Asbestos by TEM, and SVF by SEM^f Lower Manhattan

	L	ower Manhatta	an	Above 59th Street			
	Total Fibers (PCM)	Asbestos (TEM)	SVF (SEM)	Total Fibers (PCM)	Asbestos (TEM)	SVF (SEM)	
Buildings	30	5	4	4	0	4	
Residential Units	59	3	3	5	0	5	
Common Areas	29	3	2	3	0	1	
Outdoors	29	0	0	3	0	3	

	Table 5Number of Locations Sampled for Minerals in Airborne Dust, Analyzed by XRD for Minerals										
		Lower N	Ianhattan			Above 5	9th Street				
	PM100	PM10	PM4	PM2.5	PM100	PM10	PM4	PM2.5			
Buildings	26	28	30	10	3	4	4	3			
Residential Units	47	52	57	18	3	5	5	2			
Common Areas	26	27	29	8	2	2	3	2			
Outdoors	26	26	29	8	2	3	3	2			

Notes:

a. PLM = polarized light microscopy

b. TEM = transmission electron microscopy

c. SVF = synthetic vitreous fibers

d. XRD = x-ray diffraction

e. PCM = phase contrast microscopy

f. SEM = scanning electron microscopy

Settled surface dust and air samples could not always be obtained together from the same sampling location.

		Table 6			
	nary of Locations With Asbe tions that had either asbestos in				
	Ranges of asbestos and SVF				
Sampling Locations: Lower Manhattan	Total Number of Locations Sampled ^a	Number of Locations with Asbestos in Settled Surface Dust (% of locations detected) ^b	Range of Asbestos Found ^e	Number of Locations with SVF in Settled Surface Dust (% of locations detected) ^d	Range of SVF Found ^e
Outdoor	Asbestos (PLM) = 14Asbestos (TEM) = 12SVF (PLM) = 14	6 (43%)	<1%-3.4 %	11 (79%)	1%-72%
Common	Asbestos (PLM) = 26Asbestos (TEM) = 25SVF (PLM) = 26	5 (19%)	<1%-1.5%	14 (54%)	5%-27%
Residential	Asbestos (PLM) = 57Asbestos (TEM) = 52SVF (PLM) = 57	10 (18%)	<1%-1.5%	26 (46%)	2%-35% ^f
Whole Building	Asbestos (PLM) = 29 Asbestos (TEM) = 29 SVF (PLM) = 29	12 (41%)	<1%-3.4%	21 (72%)	2%-72%
Sampling Locations: Comparison Buildings	Total Number of Locations Sampled ^a	Number of Locations With Asbestos in Settled Surface Dust (% of locations detected) ^b	Range of Asbestos Found ^e	Number of Locations With SVF in Settled Surface Dust (% of locations detected) ^d	Range of SVF Found ^e
Common	Asbestos (PLM) = 3 Asbestos (TEM) = 3 SVF (PLM) = 3	0 (0%)	None Detected by PLM or TEM	0 (0%)	None Detected by PLM
Residential	Asbestos (PLM) = 5 Asbestos (TEM) = 4 SVF (PLM) = 5	0 (0%)	None Detected by PLM or TEM	0 (0%)	None Detected by PLM

a. All dust samples were initially analyzed by polarized light microscopy (PLM). Transmission electron microscopy (TEM) analysis was performed to confirm the absence of asbestos if not detected by PLM analysis.

b. The number of locations containing asbestos in dust reflects locations with asbestos detected by either PLM or TEM (reported as <1% or more). The percentage of locations with asbestos detected is calculated from the total number of dust samples.</p>

c. The range represents the highest value detected at a location (PLM or TEM).

d. The percentage of locations with synthetic vitreous fibers (SVF) detected is calculated from the number of SVF PLM samples.

e. Some locations had an additional co-located sample obtained, the range shown considers the highest measured value for each sampling location.

f. One residential unit had an extra settled surface dust sample (taken from a window sill), which contained 40% SVF by PLM.

					Table 7						
			mary of Minera Surface Dust of Lo	*	Settled Surfac	e Dust by Sampl	ing Location				
	Number of Occurrences, 14 samples totaf ^a	Minimum Detected (% by weight)	Maximum Detected (% by weight)	Average of Detections (% by weight)	Number Above Comparison ^b	Outdoor samples were not taken from comparison areas above					
Quartz	14 (100%)	$1 J^d$	27 J	12 J	NA ^c	59th St	reet because no set	tled surface dust wo	as visible.		
Calcite	13 (93%)	0.8 J	19 J	6 J	NA						
Portlandite	12 (86%)	0.07 J	6 J	2 J	NA						
Gypsum	11 (79%)	0.03 J	27 J	6 J	NA						
Mica	9 (64%)	0.05 J	0.3 J	0.1 J	NA						
Halite	7 (50%)	<0.03 J	0.1 J	0.05 J	NA						
	Co	mmon Area Settle	ed Surface Dust of	^r Lower Manhatta	in	Commo	n Areas of Compar	ison Areas Above 5	9th Street		
	Number of Occurrences, 26 samples total ^a	Minimum Detected (% by weight)	Maximum Detected (% by weight)	Average of Detections (% by weight)	Number Above Comparison ^b	Number of Occurrences, 3 samples total a	Minimum Detected (% by weight)	Maximum Detected (% by weight)	Average of Detected Values (% by weight)		
Ouartz	21 (81%)	0.03 J	25 J	5 J	6 (23%)	2 (67%)	1 J	1 J	NA		
Calcite	15 (58%)	0.02 J	10 J	3 J	9 (35%)	2 (67%)	0.03 J	0.4 J	NA		
Portlandite	13 (50%)	0.04 J	4 J	2 J	9 (35%)	1 (33%)	0.05 J	0.05 J	NA		
Gypsum	23 (88%)	0.07 J	20 J	5 J	6 (23%)	2 (67%)	2 J	3 J	NA		
Mica	5 (19%)	0.06 J	0.6 J	0.2 J	5 (19%)	0 (0%)	ND	ND	NA		
Halite	4 (15%)	0.04 J	0.06 J	0.05 J	1 (4%)	1 (33%)	0.04 J	0.04 J	NA		
	Res	idential Units Sett	led Surface Dust o	of Lower Manhat	tan	Residen	tial Units of Compa	rison Areas Above	59th Street		
	Number of Occurrences, 57 samples total ^a	Minimum Detected (% by weight)	Maximum Detected (% by weight)	Average of Detections (% by weight)	Number Above Comparison ^b	Number of Occurrences, 5 samples total ^a	Minimum Detected (% by weight)	Maximum Detected (% by weight)	Average of Detected Values (% by weight)		
Quartz	30 (53%)	0.05 J	31 J	9 J	15 (26%)	2 (40%)	1 J	2 J	NA		
Calcite	20 (35%)	0.02 J	21 J	8 J	13 (23%)	1 (20%)	0.9 J	0.9 J	NA		
Portlandite	21 (37%)	0.05 J	8 J	2 J	17 (30%)	2 (40%)	0.08 J	0.08 J	NA		
Gypsum	45 (79%)	0.05 J	30 J	4 J	9 (16%)	4 (80%)	2 J	4 J	3 J		
Mica	5 (9%)	0.03 J	0.3 J	0.1 J	1 (2%)	1 (20%)	0.08 J	0.08 J	NA		
Halite	6 (11%)	0.03 J	0.1 J	0.06 J	0 (0%)	1 (20%)	0.4 J	0.4 J	NA		

a. Shows the number of samples where the mineral was detected and the percentage based on the number of samples obtained from this area.

b. Shows the number of results, and the percentage of samples obtained from this area, that had estimated values greater than the maximum levels found at locations above 59th Street.

c. NA = not applicable.

d. J = estimated.

Table 8Summary of Results for Locations With Airborne Fiber LevelsHigher Than Levels in Comparison Areas Above 59th Street.

Air samples were collected and analyzed by phase contrast microscopy for total fibers (NIOSH 7400). All samples with fiber counts higher than the comparison areas above 59th Street were re-analyzed to determine if those fibers were asbestos, synthetic vitreous fibers (SVF), or other material.

Building	Aggressive	Fibers in Air by	Asbestos in	Asbestos in Air	SVF in Dust	SVF in Air by
(Area)	Sampling ^a	PCM	Dust by PLM	by TEM	by PLM	SEM
		(f/cc) ^b	or TEM ^c	$(f/cc)^{b}$	-	$(f/cc)^{b}$
1						
(Residence)	No	0.006	ND^{g}	<0.001 ^d	20%	0.000162
2						
(Common)	No	0.005	ND^{g}	< 0.001 ^d	27%	0.000255
19						
(Common)	No	Overloaded ^e	ND^{g}	< 0.006 ^f	ND	Not analyzed
24						
(Residence)	No	0.005	ND^{g}	< 0.001 ^d	10%	0.000037
26						
(Common)	No	0.004	ND^{g}	< 0.001 ^d	5%	< 0.00004
26						
(Residence)	No	0.012	ND^{g}	< 0.001 ^d	ND	< 0.00004

a. Aggressive sampling refers to a technique used in some residential units where the vacuum exhaust (used for settled dust sample collection) was used to stir up the settled surface dust before the air sampling began.

b. f/cc = fibers in each cubic centimeter (cc) of air as determined using phase contrast microscopy (PCM), transmission electron microscopy (TEM), or scanning electron microscopy (SEM) methods. This is calculated from the number of fibers seen on the air filter and the volume of air pulled through the filter measured in cubic centimeters of air.

c. Value shown represents the highest of polarized light microscopy (PLM) or TEM results for this area.

d. The TEM method employed here, National Institute of Occupational Safety and Health (NIOSH) 7402, counts fibers of the same size as those detected by PCM analysis. Fibers reported here are greater than or equal to 5 microns in length and 0.25 microns in width.

e. A building renovation project was occurring near the area of this sampling equipment. Construction dust and building insulation material may have influenced this sample.

f. Sample processing of the overloaded filter involved transferring the material to a new filter; this process provides less analytical sensitivity, resulting in a higher detection limit.

g. ND = not detected.

		Outdoor A	Areas in Lower Manhatta	n		
Building	Area	Asbestos in Dust (PLM or TEM) ^a	SVF in Dust (PLM) ^b	Aggressive Sampling ^c	Fibers in Air (PCM) ^d	
1	Outdoor	<1%	22%	No	0.001 f/cc	
2 ^e	Outdoor	1.3%	28%	No	0.003 f/cc	
5	Outdoor	3.4%	25%	No	0.002 f/cc	
7	Outdoor	1.7%	35%	No	<0.001 f/cc	
15	Outdoor	1.9%	72%	No	<0.001 f/cc	
27	Outdoor	<1%	15%	No	<0.001 f/cc	
		Common	Areas in Lower Manhatta	n		
Building	Area	Asbestos in Dust (PLM or TEM) ^a	SVF in Dust (PLM) ^b	Aggressive Sampling ^c	Fibers in Air (PCM) ^d	
4	Common	<1%	15%	No	0.001 f/cc	
6	Common <1%		10%	No	<0.001 f/cc	
10	Common	ommon 1.5% 20%		No	0.002 f/cc	
24	Common	mon <1% 5%		No	0.001 f/cc	
27	Common	<1%	10%	No	<0.001 f/cc	
		Residential	Units in Lower Manhatt	an		
Building	Area	Asbestos in Dust (PLM or TEM) ^a	SVF in Dust (PLM) ^b	Aggressive Sampling ^c	Fibers in Air (PCM) ^d	
4	Residence 1	<1%	2%	Yes	<0.001 f/cc	
4	Residence 2	<1%	5%	Yes	0.001 f/cc	
5	Residence 1	<1%	10%	Yes	0.002 f/cc	
5	Residence 2	<1%	20%	Yes	<0.001 f/cc	
9	Residence 1	<1%	2%	Yes	0.001 f/cc	
9	Residence 2	<1%	5%	Yes	0.003 f/cc	
11	Residence 1	<1%	ND	Yes	<0.001 f/cc	
11	Residence 2	1.5%	ND	Yes	<0.001 f/cc	
15	Residence 1	<1%	ND	Yes	<0.001 f/cc	
27	Residence 1	<1%	10%	No	<0.001 f/cc	

b. SVF = synthetic vitreous fibers. Measured by PLM analysis.

c. Aggressive sampling refers to a technique used in some residential units where the vacuum exhaust (used for settled dust sample collection) was used to stir up the settled surface dust before the air sampling began.

d. f/cc = the fibers in each cubic centimeter (cc) of air. This is calculated from the number of fibers seen on the air filter and the volume of air pulled through the filter measured in cubic centimeters.

e. Building 2 also had a co-located dust sample; values represent the highest measured result.

Table 10 Summary of Airborne Fibers at Locations Where Only Synthetic Vitreous Fibers (SVF) Were Detected in Settled Surface Dust

Building	Area	Asbestos in Dust (PLM or TEM) ^a	SVF in Dust (PLM) ^b	Aggressive Sampling ^c	Fibers in Air (PCM) ^d
8	Outdoor	ND^{f}	30%	No	<0.001 f/cc
24	Outdoor	ND^{f}	55%	No	0.002 f/cc
28	Outdoor	ND^{f}	15%	No	<0.001 f/cc
		Common	Areas in Lower Manhatta	an	
Building	Area	Asbestos in Dust (PLM or TEM) ^a	SVF in Dust (PLM) ^b	Aggressive Sampling ^c	Fibers in Air (PCM) ^d
2	Common	ND^{f}	27%	No	0.005 f/cc ^e
7	Common	ND^{f}	5%	No	0.001 f/cc
14	Common	ND ^f	5%	No	0.003 f/cc
25	Common	ND^{f}	5%	No	<0.001 f/cc
26	Common	ND^{f}	5%	No	0.004 f/cc ^e
28	Common	ND^{f}	10%	No	0.001 f/cc
		Residential	Units in Lower Manhatt	an	
Building	Area	Asbestos in Dust (PLM or TEM) ^a	SVF in Dust (PLM) ^b	Aggressive Sampling ^c	Fibers in Air (PCM) ^d
1	Residence 1	ND^{f}	20%	No	0.006 f/cc ^e
1	Residence 2	ND^{f}	20%	Yes	<0.001 f/cc
2	Residence 1	ND^{f}	25%	No	<0.001 f/cc
2	Residence 2	ND^{f}	20%	Yes	0.002 f/cc
6	Residence 1	ND^{f}	15%	Yes	<0.001 f/cc
6	Residence 2	ND^{f}	15%	Yes	<0.001 f/cc
7	Residence 2	ND^{f}	5%	No	<0.001 f/cc
10	Residence 1	ND ^f	15%	Yes	0.001 f/cc
10	Residence 2	ND^{f}	10%	Yes	0.001 f/cc
12	Residence 1	ND ^f	5%	No	0.001 f/cc
12	Residence 2	ND ^f	5%	Yes	<0.001 f/cc
13	Residence 1	ND ^f	10%	Yes	0.003 f/cc
15	Residence 2	ND ^f	5%	Yes	<0.001 f/cc
18	Residence 1	ND ^f	30%	No	0.002 f/cc
	Residence 2	ND ^f	35%	No	0.002 f/cc
18	1	ND^{f}	10%	No	0.005 f/cc ^e
18 24	Residence 2	r			.0.001.0/
18 24 25 27	Residence 2 Residence 1 Residence 2	ND ^f	5% 10%	No No	<0.001 f/cc <0.001 f/cc

c. Aggressive sampling refers to a technique used in some residential units where the vacuum exhaust (used for settled dust sample collection) was used to stir up the settled surface dust before the air sampling began.

d. f/cc = the fibers in each cubic centimeter (cc) of air. This is calculated from the number of fibers seen on the air filter and the volume of air pulled through the filter measured in cubic centimeters.

e. Scanning electron microscopy (SEM) and TEM results for these air samples are available and shown on Table 8.

f. ND = not detected.

Sum	mary of]	Minerals i	in Indoor	and Outdo	or Air Fr	Tabl om Lower		tan and (Compari	son Build	lings Above	e 59th Str	eet ^{a, b, c}	
				s from Lower							From Compa			
	Number of Samples	Quartz	Calcite	Portlandite	Gypsum	Mica	Halite	Number of Samples	Quartz	Calcite	Portlandite	Gypsum	Mica	Halite
PM 100														
Number (Frequency detected %)	101	14 (14%) ^d	9 (9%)	8 (8%)	24 (24%)	2 (2%)	4 (4%)	7	ND^{f}	ND	ND	1 (14%)	ND	ND
Air Concentration $(\mu g/m^3)$		3–13 J ^e	3–14 J	16–95 J	3–14 J	9–13 J	4–14 J					3 J		
PM 10					r			•	r					
Number (Frequency detected %)	105	11 (10%)	10 (10%)	10(10%)	33 (31%)	1 (1%)	5 (5%)	10	ND	ND	ND	3 (30%)	ND	ND
Air Concentration $(\mu g/m^3)$		3–12 J ^e	3–5 J	14–25 J	3–14 J	8 J	4–5 J					3 J		
PM 4					I				1		1			
Number (Frequency detected %)	114	13 (11%)	13 (11%)	12 (11%)	40 (35%)	4 (4%)	3 (3%)	11	ND	ND	ND	3 (27%)	ND	ND
Air Concentration (µg/m ³)		4–19 J	4–10 J	21–84 J	4–15 J	14–43 J	7–19 J					5 J		
PM 2.5					I				1		1			
Number (Frequency detected %)	34	1 (3%)	ND	ND	1 (3%)	ND	ND	6	ND	ND	ND	1 (17%)	ND	ND
Air Concentration $(\mu g/m^3)$		3 J			3 J							3 J		
Range of Detection	Limits (µg/m							1			1			
PM 100 PM 10 PM 4		2–5 J 2–5 J 4–6 J	2–5 J 2–5 J 4–6 J	12–26 J 12–26 J 20–30 J	2–5 J 2–5 J 4–6 J	7–15 J 7–15 J 11–17 J	4–8 J 4–8 J 6–9 J		2–5 J 2–5 J 4–6 J	2–5 J 2–5 J 4–6 J	12–26 J 12–26 J 20–30 J	2–5 J 2–5 J 4–6 J	7–15 J 7–15 J 11–17 J	4-8 J 4-8 J 6-9 J
PM 2.5		2–4 J	2–4 J	13–20 J	2–4 J	7–11 J	4–6 J		2–4 J	2–4 J	13–20 J	2–4 J	7–11 J	4-6.

a. Where two samples were co-located—the greatest value of the two was included in this summary. Each location is only represented once in this table.

b. Concentrations shown are estimated values, indicated by "J."

c. Air samples from all locations, indoor and outdoor, are pooled in this table due to uncertainties in the data.

d. Cristobalite, a different crystalline form of silica, was found in one Lower Manhattan air sample at 15 micrograms per cubic meter of air (µg/m³) J in the PM 100 fraction.

e. J = estimated.

f. ND = not detected.

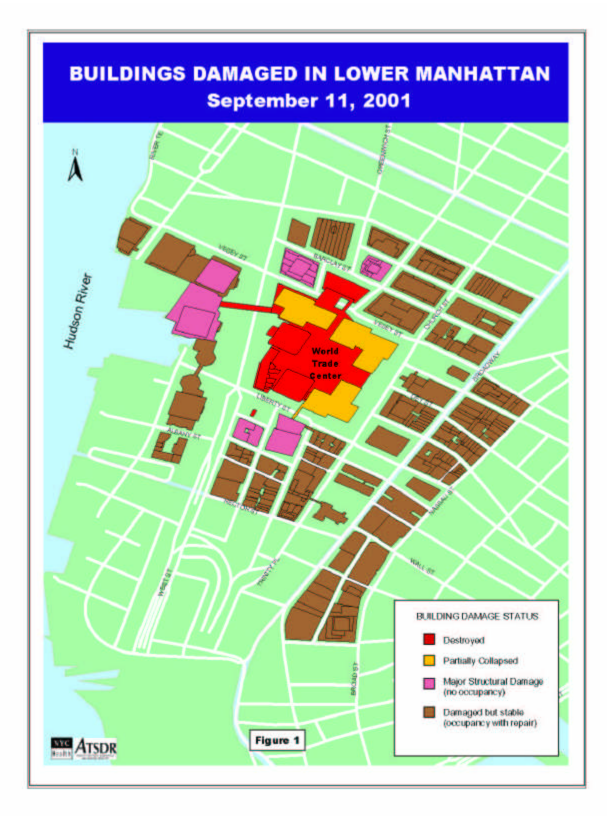
Building Number	Quartz (%) ^{a, d, e}	Gypsum (%) ^{a, d, e}	Asbestos (%) ^{b, f}	SVF (%) ^{c, f}
Comparison				
Locations 31-34	2 J	4 J	Non-Detect	Non-Detect
1	31 J	30 J	Non-Detect	20
2	23 J	14 J	Non-Detect	27
3	Not Sampled	Not Sampled	Not Sampled	Not Sampled
4	14 J	9 J	<1	15
5	11 J	9 J	<1	20
6	27 J	20 J	<1	15
7	21 J	15 J	Non-Detect	5
8	28 J	17 J	Non-Detect	Non-Detect
9	25 J	16 J	<1	7
10	3 J	0.8J	1.5	20
11	2 J	2 J	1.5	Non-Detect
12	4 J	1 J	Non-Detect	5
13	0.05 J	1 J	Non-Detect	10
14	0.03 J	2 J	Non-Detect	5
15	0.4 J	1 J	<1	5
16	Non-Detect	0.9 J	Non-Detect	Non-Detect
17	2 J	2 J	Non-Detect	2
18	0.9 J	1 J	Non-Detect	35
19	1 J	2 J	Non-Detect	Non-Detect
20	0.9 J	1 J	Non-Detect	Non-Detect
21	0.9 J	2 J	Non-Detect	Non-Detect
22	14 J	2 J	Non-Detect	3
23	2 J	1 J	Non-Detect	Non-Detect
24	0.03 J	Non-Detect	<1	10
25	Non-Detect	2 J	Non-Detect	5
26	0.7 J	2 J	Non-Detect	5
27	0.04 J	1 J	<1	10
28	Non-Detect	1 J	Non-Detect	10
29	1 J	2 J	Non-Detect	Non-Detect
30	1 J	4 J	Non-Detect	Non-Detect

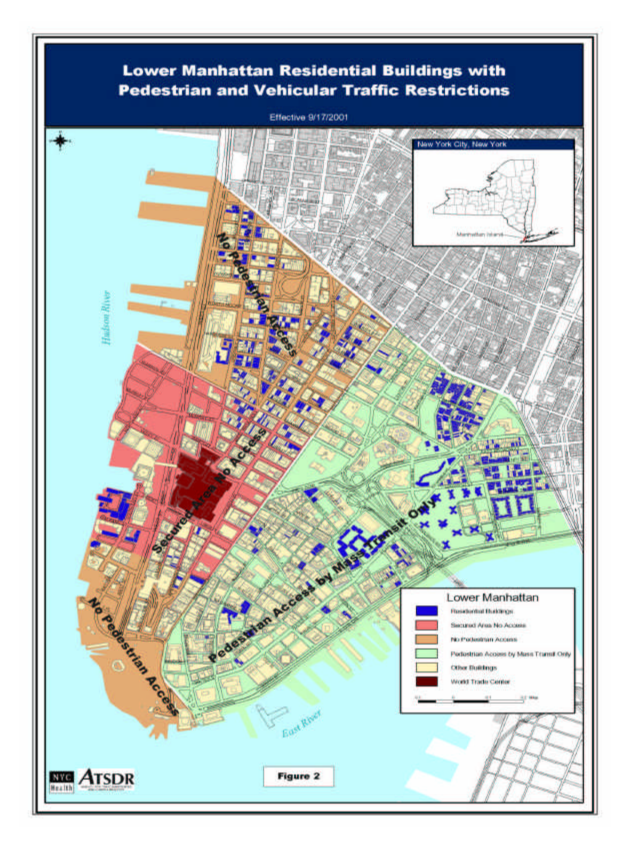
b. Asbestos value represents the highest of the PLM and TEM result for each location.c. SVF is synthetic vitreous fiber and was measured by PLM.

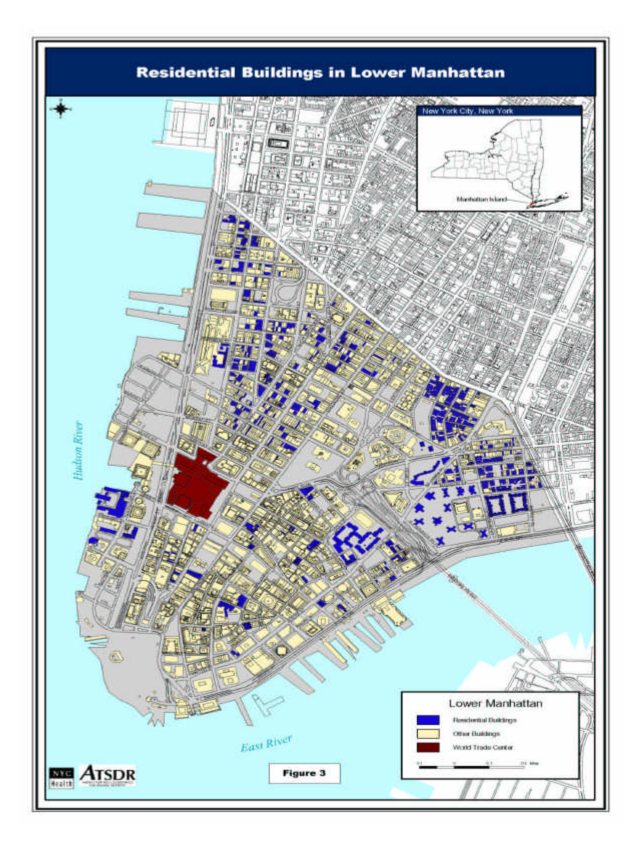
d. Results shown are estimated values, indicated by "J."

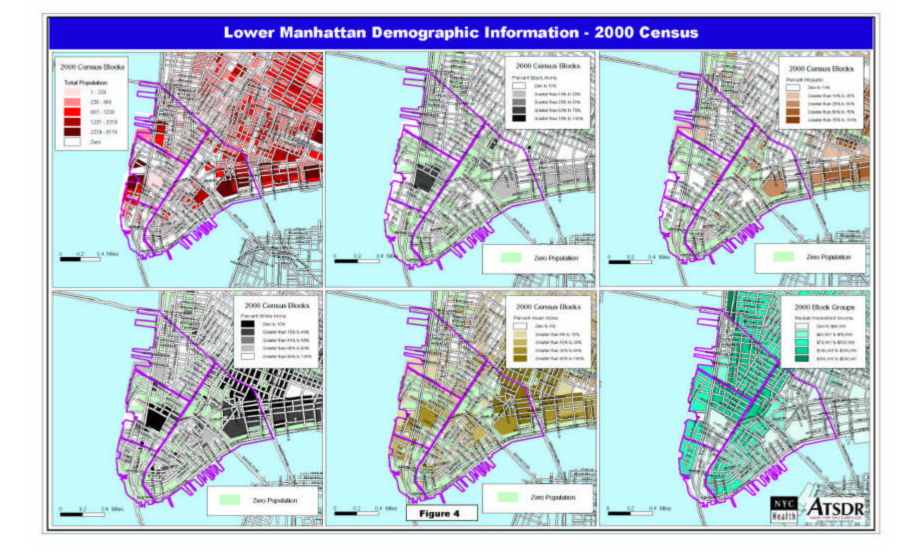
e. % = weight of mineral per weight of dust.

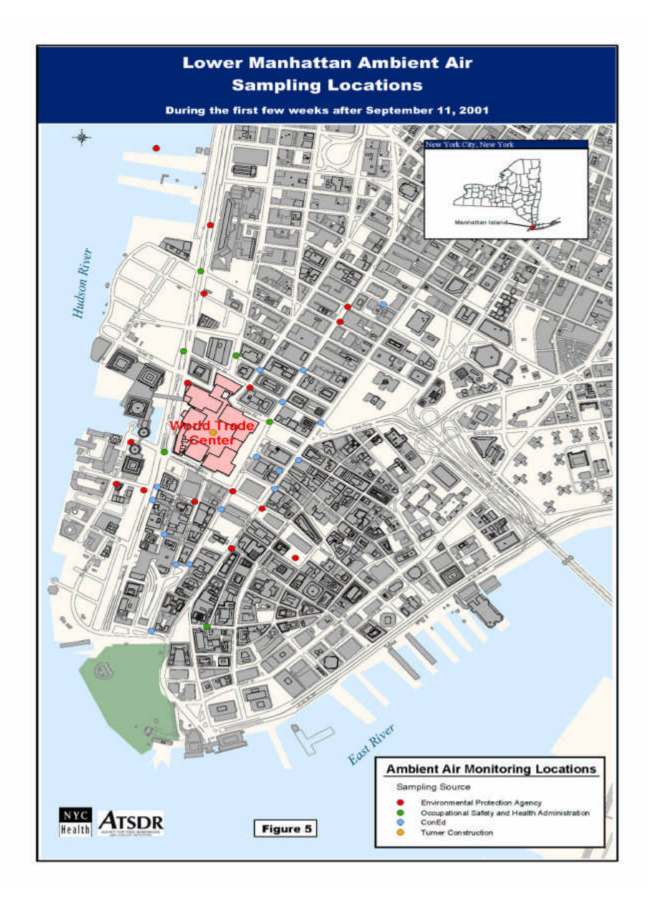
f. % = roughly area of fibers per area of dust. FIGURES

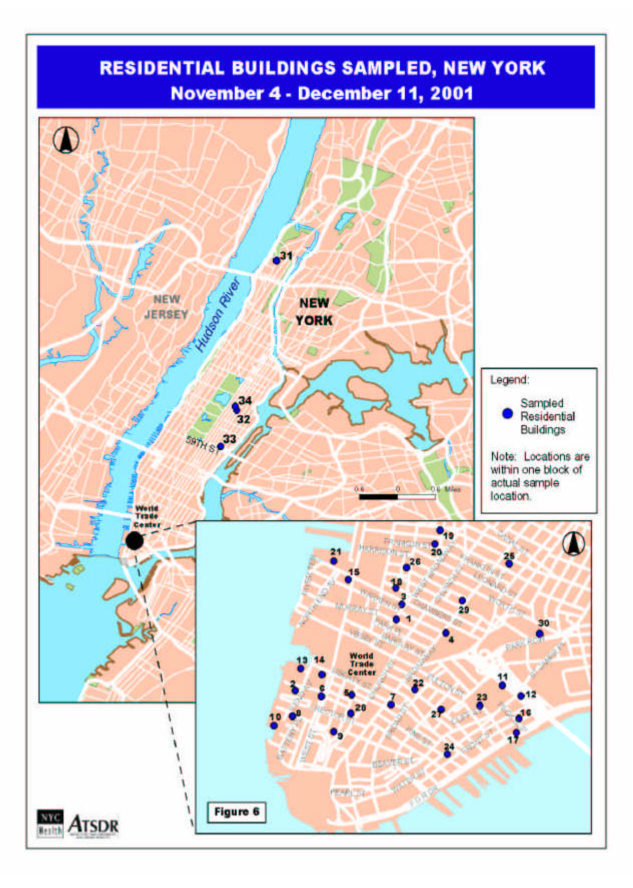


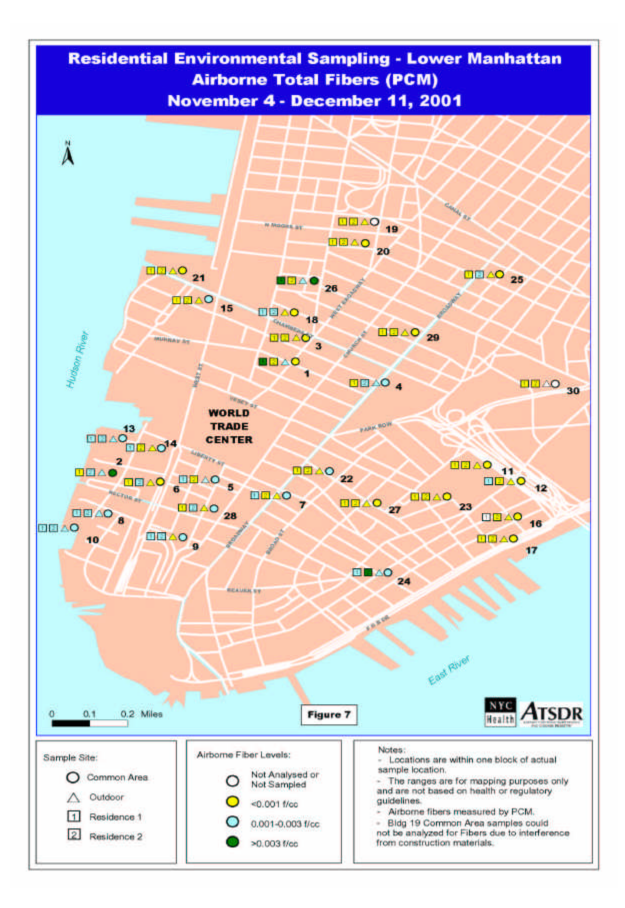




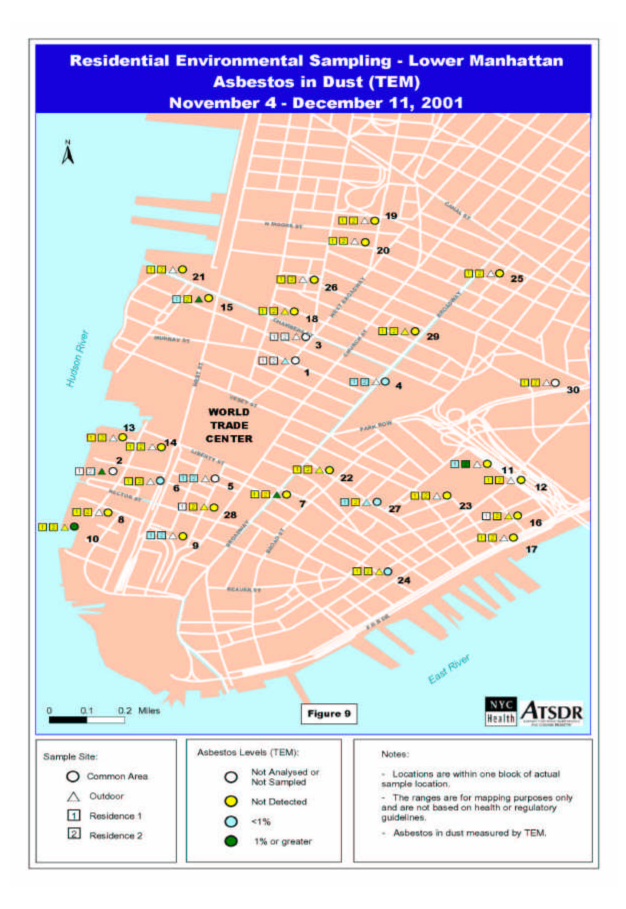






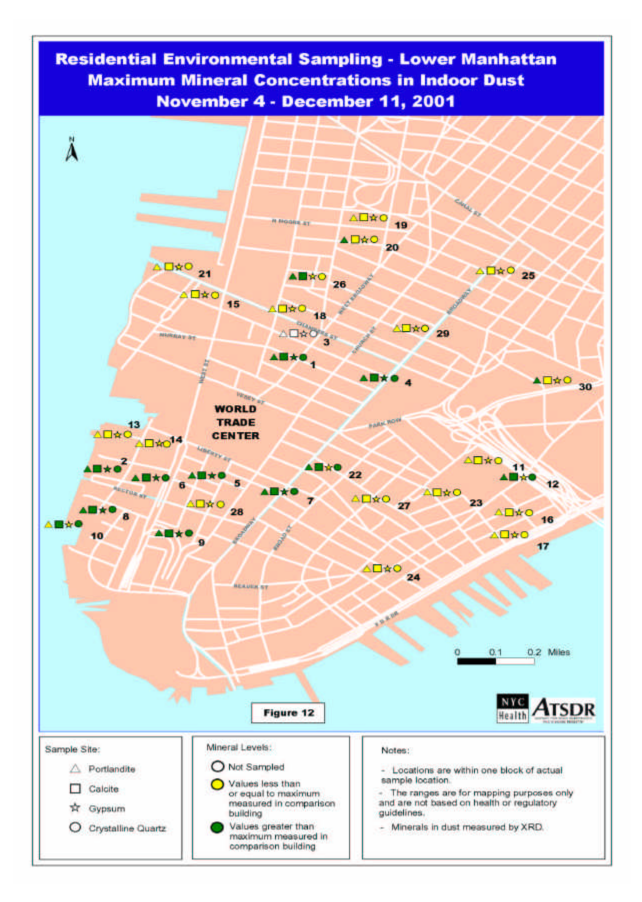






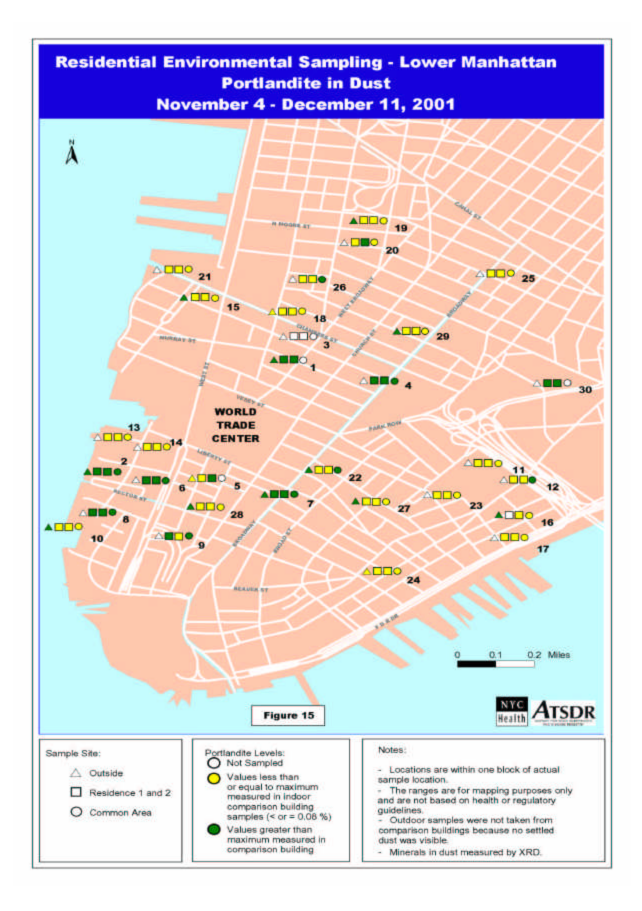


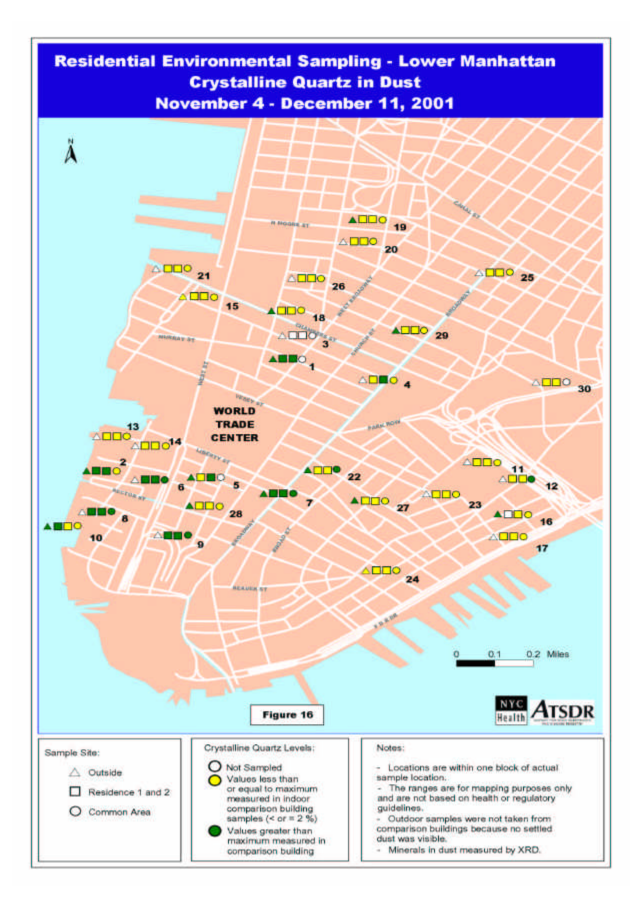


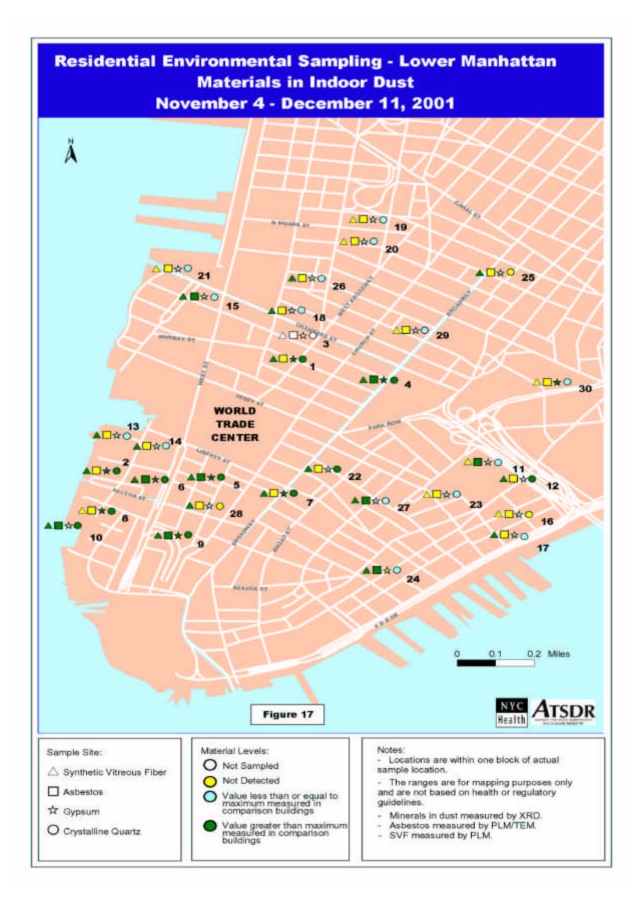












SAMPLING EVENT PHOTOGRAPHS



Photo 1. Residential Vacuum Sample



Photo 2. Residential Air Sampling



Photo 3. Residential Air Sampling



Photo 4. Residential Air Sampling



Photo 5. Residential Air Sampling



Photo 6. Residential Air Sampling



Photo 7. Outside Bulk Sampling



Photo 8. Sampling Head

APPENDICES

Appendix A. Particulate Matter Quality Assurance/Quality Control Discussion

General Notes on Quality Assurance/Quality Control Procedures

Quality assurance/quality control (QA/QC) procedures are used to ensure the precision, accuracy, completeness, representativeness, comparability, and method detection limit of the results. The colocated samples and blanks are a primary means of assessing each of the data quality indicators. Comparing the results of two co-located samples provides information on the precision of the results as a whole. Comparing sample results to their associated blanks can help identify some potential errors in accuracy. Comparing the results received to those expected and necessary to draw conclusions about the data can help in understanding the completeness and reliability of the results. Comparability can be assessed by looking at the results taken from different sampling locations of the same building—or from different buildings for the same type of location. The reviews are generally qualitative and provide a qualitative assessment of how well the data actually represents the sampled location.

The method detection limit is a more quantitative review. It is accomplished by calculating the lowest result the analytical method can accurately identify. It is based on an analysis of the blank samples and is specific for the sampling/analysis method. Reliable information can only be obtained from sample results with method detection limits significantly lower than the average value of the results and significantly lower than any comparison values to which the sample results will be compared.

The analytical results of the concentration of airborne particulate matter were provided by the laboratory as the initial weight of the filter prior to the sampling event and the final weight of the filter after the sampling event. The concentration of the particulate matter was calculated by dividing the weight of the material collected on the filter by the volume of air drawn through the filter by the pump using the following formulas.

Weight Gain of Filter = Final Weight – Initial Weight

Sample Volume = Average Pump Flow Rate × Sample Collection Time

Average Air Sampling Pump Flow Rate=(Presampling Flow Rate + Postsampling Flow Rate) ÷ 2

Concentration = Weight Change ÷ Sample Volume

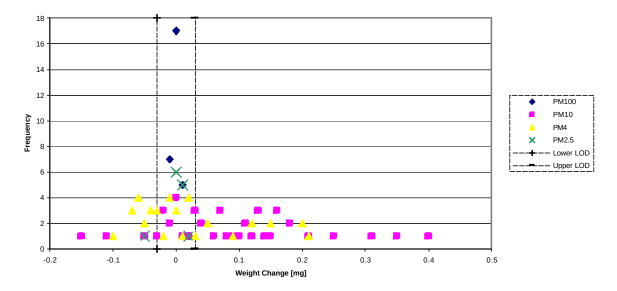
Airborne Particulate Matter Sample Results

Each area that was sampled for airborne asbestos was also sampled for airborne particulate matter (PM). The sample results from the PM measurements did not meet the data QA/QC objectives. A variety of statistical and graphical analyses were performed on, and with, the measured results in an attempt to identify the cause and extent of inconsistencies of the measured results. A subset of the measured results that are not potentially affected by these inconsistencies were not identified. Therefore, the entire data set describing airborne particulate matter concentrations was rejected. In addition, the specific cause of the inconsistencies was not identified. The following information explains why the airborne particulate matter results were rejected.

Ideally, the particulate matter data would have provided information about the airborne concentration with an aerodynamic diameter of 100 microns and less (PM100), of 10 microns and less (PM10), of 4 microns and less (PM4), and of 2.5 microns and less (PM2.5). These different size fractions (PM100, PM10, PM4, and PM2.5) are collected using slightly different equipment, but following the same basic procedures. A pump draws air into a sampling head, and size specific particulate matter is deposited onto a filter. The difference in filter weight, before and after the sampling, represents the mass of the particulate matter that was captured by the filter. Knowing the presampling and postsampling air flow rates of the pump and the time duration for sample collection allows us to calculate the average volume of air that was drawn through the filter. The concentration of the size fraction in the sampled air is then calculated by dividing the PM mass by the air volume sampled.

In addition to the filters that were used to measure the concentration of airborne particulate matter, the laboratory also sent filters to be used as field blanks. Two blanks were sent for each PM fraction for each of the two sampling teams for each building. The blanks traveled with the sample filters, went to the sampled building or area, and were treated just like the sample filters—except that they were not used. Ideally, there should be very little difference in the weight of the blanks before the sampling and after the sampling because they were not used. A weight gain in field blanks may indicate improper sample handling in the field or problems in filter weighing in the lab. For the latter a decrease in post field blank filter weight is also an indication of lab weighing error.

The graph shown with this discussion shows the frequency distribution of the air sample filter weight change in all but seven of the blanks used. Seven blanks were not included because their weight change (from a negative -90 milligrams (mg) to 5.99 mg) was greater than the limits shown in this graph (-0.2 mg to 0.5 mg). The vertical axis represents the number of blanks that had a weight change within 0.01 mg of the weight change shown on the horizontal axis.



Frequency Distribution of Weight Change in Selected Blanks

The dashed vertical lines show the upper and lower limits of detection expected for this method (+/-0.03 mg). The distance between limits of detection represents the maximum weight gain expected for the blanks if the entire sampling procedure was followed as required by standard sampling methodologies. All of the size fractions had at least two blanks outside the limit of detection boundaries. (The two out of range blanks for PM100 are not shown on the graph because they are also beyond the boundaries of the graph). Approximately 6% of the PM100, 70% of the PM10, 57% of the PM4, and 8% of the PM2.5 blanks had weight changes greater than the limit of detection.

In general, blanks are used (1) to assess the ability of the sampling and laboratory analysis methodology to accurately estimate the concentration of particulate matter in the air at the sampled location and (2) to validate that there is no filter contamination problem from the time the blanks are initially weighed in the laboratory until the time the blanks are weighed in the laboratory after sampling. Errors could occur at any step in the process.

The large weight change of the blanks indicates errors in either the laboratory weighing process or the air sampling process. The graph illustrates that the results of the airborne particulate matter sampling cannot be used to reliably estimate the actual concentration of any of the particulate matter size fractions; therefore, the entire data set was rejected from further consideration.

Appendix B. Survey Result

	Summary of Residential Sampling Survey Form Results and Review of Photographs (Abbreviations and terms used in the table are defined on the last page in this table.)											
Event Number	Location	Number of Broken Windows	Dust Visible Initially	Dust Visible Currently	Residence Occupied			Level of Dust in Photo	Photo Comments			
1	Outside	28						None Visible				
1	Common	0	SI	Ν		С		None Visible	Prepared for asbestos abatement			
1	Residence 1	2	LI	Y	U	None	N	Large Amount	Large amount of WTC dust; more than other locations sampled			
1	Residence 2	0	NI	Ν	U	None	Y	None Visible				
2	Outside	1		Y				None Visible				
2	Common	0	NI	Ν		С		Slight Amount				
2	Residence 1	0	SI	N	Y	С	N	None Visible	Residence window faces WTC site			
2	Residence 2	0	SI	Ν	U	С	Y	None Visible				
3	Outside	0		N				None Visible	Sampled from courtyard several floors up			
3	Common	0	LI	N		AA		None Visible				
3	Residence 1	0	LI	N	Y	AA	N	None Visible				
3	Residence 2	0	МІ	N	Y	С	N	None Visible				
4	Outside	0		N				Slight Amount				
4	Common	0	NI	Y		0		None Visible				
4	Residence 1	0	SI	Y	N	None	Y	None Visible				
4	Residence 2	0	SI	Y	N	None	Y	None Visible				
5	Outside	50		Y				Large Amount				
5	Common	0	SI	N		AA		None Visible				
5	Residence 1	6	LI	LI	Y	AA	Y	None Visible				
5	Residence 2	6	LI	LI	Y	AA	Y	None Visible	Room looks clean but lots of WTC dust outside one window sill			
6	Outside	200		N				None Visible				
6	Common	0	SI	N		С			Couldn't identify photos for common area			
6	Residence 1	0	LI	N	U	С	Y	None Visible				
6	Residence 2	0	SI	N	U	С	Y	None Visible				
7	Outside	2		N				None Visible				
7	Common	0	MI	N		С		None Visible	Many window sills (not sure where) have significant amount of WTC			
7	Residence 1	0	MI	N	Y	С	N	None Visible	dust on outside ledge, not noticeable on inside of window.			
7	Residence 2	0	MI	N	Y	С	N	None Visible	Resident has window view of WTC site			

	Summary of Residential Sampling Survey Form Results and Review of Photographs (Abbreviations and terms used in the table are defined on the last page in this table.)											
Event Number	Location	Number of Broken Windows	Dust Visible Initially	Dust Visible Currently	Residence Occupied		Aggressive Sampling	Level of Dust in Photo	Photo Comments			
8	Outside	0		N				None Visible				
8	Common	0	MI	N		С		None Visible				
8	Residence 1	1	SI	Ν	Y	0	Ν	None Visible				
8	Residence 2	1	MI	N	Y	С	N	None Visible	Resident has window view of WTC site			
9	Outside	2		N				None Visible in Picture	Utility/road construction occurring near building and across the			
9	Common	0	LI	N		С		None Visible	street from the front door.			
9	Residence 1	0	MI	Y	Y	С	Y	None Visible				
9	Residence 2	0	MI	Y	Y	С	Y	Slight Amount				
10	Outside	0		Y				Slight Amount				
10	Common	0	SI	N		AA		None Visible				
10	Residence 1	0	NI	N	Y	AA	Y	None Visible	View of Statue of Liberty			
10	Residence 2	1	SI	N	Y	AA	Y	None Visible				
11	Outside	0	LI	N				None Visible	Looks like many dried, fallen leaves on sidewalk			
11	Common	0	SI	Ν		MS		None Visible				
11	Residence 1	0	SI	Ν	V	None	Y	None Visible				
11	Residence 2	0	UNK	N	V	UNK	Y	None Visible	Wall AC unit was removed, it was just a hole to the outside			
12	Outside	0		Ν				None Visible	Looks like many dried, fallen leaves on sidewalk			
12	Common	0	SI	Ν		MS		None Visible				
12	Residence 1	0	MI	Y	V	None	N	Large Amount	Very messy, doesn't look like "WTC dust" (has "'post-move-out" look)			
12	Residence 2	0	SI	N	V	None	Y	None Visible				
13	Outside	0		N				Slight Amount	A little messy, but does not look like WTC dust			
13	Common	0	МІ	N		MS		None Visible				
13	Residence 1	0	МІ	N	V	С	Y	None Visible				
13	Residence 2	0	МІ	Y	V	С	Y	Large Amount	Very messy, but not with WTC dust (post-move-out look)			
14	Outside	0		N				None Visible	Air sample from garden area; no bulk sample taken; WTC dust			
14	Common	0	SI	Ν		С		None Visible	visible on neighboring building.			
14	Residence 1	0	MI	Y	U	None	Y	Large Amount	Light dust on floors; one pile of "material," could be from remodeling activity			

	Summary of Residential Sampling Survey Form Results and Review of Photographs (Abbreviations and terms used in the table are defined on the last page in this table.)												
Event Number	Location	Number of Broken Windows	Dust Visible Initially	Dust Visible Currently	Residence Occupied		Aggressive Sampling	Level of Dust in Photo	Photo Comments				
14	Residence 2	0	MI	Y	U	None	Y	None Visible	Pile of trash swept into center of room; no WTC dust				
15	Outside	0		N				None Visible	Bulk sample from roof top courtyard,				
15	Common	0	SI	Y		MS		None Visible	Dust visible in (and sampled from) cracks between tiles.				
15	Residence 1	0	SI	Y	U	0	Y	None Visible					
15	Residence 2	0	SI	N	U	0	Y	None Visible					
16	Outside	0		N				None Visible					
16	Common	0	MI	N		С		None Visible					
16	Residence 1	0				MS		None Visible					
16	Residence 2	0	LI	N	Y	0	N		No picture				
17	Outside	0		N				Slight Amount	Dirty/messy, but doesn't look like WTC dust				
17	Common	0	SI	Y		С		None Visible					
17	Residence 1	0	SI	N	Y	С	N	None Visible					
17	Residence 2	0	МІ	Y	U	С	Y	Slight Amount					
18	Outside	0		N				М	Sample looks granular and sandy, not gray like WTC dust				
18	Common	0	LI	N		MS		None Visible	Leaf track-in visible				
18	Residence 1	0	МІ	N	Y	С	N	None Visible	Window view of WTC site				
18	Residence 2	0	LI	N	Y	С	N	None Visible					
19	Outside	0		N				Moderate	Visible in, and sampled from, sidewalk joint with building; sidewalk				
19	Common	0	SI	N		С		None Visible	looked cleaner.				
19	Residence 1	0	LI	N	Y	0	N	None Visible					
19	Residence 2	0	NI	N	Y	0	N		No picture				
20	Outside	0		N				Unk	Too distant to see surface dust				
20	Common	0	SI	N		MS		None Visible					
20	Residence 1	0	NI	N	Y	0	N	None Visible					
20	Residence 2	0	SI	N	Y	0	N		No picture				
21	Outside	0		N				None Visible					
21	Common	0	SI	N		с		None Visible					
21	Residence 1	0	SI	N	Y	с	N	None Visible					
21	Residence 2	0	NI	N	Y	С	N	None Visible					

	Summary of Residential Sampling Survey Form Results and Review of Photographs (Abbreviations and terms used in the table are defined on the last page in this table.)												
Event Number	Location	Number of Broken Windows	Dust Visible Initially	Dust Visible Currently	Residence Occupied			Level of Dust in Photo	Photo Comments				
22	Outside	8		Y				None Visible	Sidewalk/entrance to building and roof courtyard look clean, but				
22	Common	1	LI	N		С		None Visible	there is a pile of material to sample on roof; location is unknown.				
22	Residence 1	0	МІ	Y	Y	MS	N	None Visible					
22	Residence 2	0	LI	N	Y	С	N	None Visible					
23	Outside	0		N				None Visible					
23	Common	0	SI	N		С		None Visible					
23	Residence 1	0	МІ	Y	Y	0	N	None Visible					
23	Residence 2	0	МІ	N	Y	0	N	None Visible					
24	Outside	0		N				None Visible					
24	Common	0	LI	N		MS		None Visible					
24	Residence 1	0	SI	N	Y	0	N	None Visible					
24	Residence 2	0	SI	N	Y	С	N	None Visible					
25	Outside	0		N				None Visible					
25	Common	0	NI	N		MS		None Visible					
25	Residence 1	0	NI	N	Y	0	N	None Visible					
25	Residence 2	0	SI	N	Y	С	N	None Visible					
26	Outside	0		N				None Visible					
26	Common	0	NI	N		MS		None Visible	Floor area sampled contained some broken tiles.				
26	Residence 1	0	NI	N	Y	0	N	None Visible					
26	Residence 2	0	SI	N	Y	0	N	None Visible					
27	Outside	0		N				None Visible					
27	Common	0	МІ	N		MS		None Visible					
27	Residence 1	0	LI	N	Y	С	N	None Visible					
27	Residence 2	0	SI	N	Y	С	N	None Visible					
28	Outside	0						None Visible	Bulk sample from 4-inch wide strip between two different tiled areas:				
28	Common	0	LI	Y		С		None Visible	thick with granular/dusty/other material, not sure if any is WTC.				
28	Residence 1	3	LI	N	U	С	N	None Visible					
28	Residence 2	0	МІ	N	U	С	N	None Visible					
29	Outside	0		N				None Visible	Bulk sample from small pile at an inside corner of the building wall				

	Summary of Residential Sampling Survey Form Results and Review of Photographs (Abbreviations and terms used in the table are defined on the last page in this table.)												
Event Number	Location	Number of Broken Windows	Dust Visible Initially	Dust Visible Currently	Residence Occupied			Level of Dust in Photo	Photo Comments				
29	Common	0	NI	N		С		None Visible					
29	Residence 1	0	SI	Ν	Y	0	N	None Visible					
29	Residence 2	0	SI	Y	Y	0	N	None Visible					
30	Outside	0							No picture				
30	Common	0							No picture				
30	Residence 1	0	SI	Ν	Y	0	N	None Visible					
30	Residence 2	0	MI	Ν	Y	С	N	None Visible					
31	Outside	0						None Visible					
31	Common	0	NI	N		MS		None Visible					
31	Residence 1	0	NI	Ν	N	MS	Y	None Visible					
31	Residence 2	0	NI	UNK	Y	0	N	None Visible					
32	Outside	0						None Visible	Trash visible on sidewalk				
32	Common	0	NI	Ν		MS		None Visible					
32	Residence 1	0	NI	N	Y	0	N	None Visible					
32	Residence 2	0											
33	Outside	0						None Visible					
33	Common	0											
33	Residence 1	0											
33	Residence 2	0	NI	Y	Y	0	N	None Visible					
34	Outside	0		N				None Visible					
34	Common	0	NI	N		MS		None Visible					
34	Residence 1	0											
34	Residence 2	0	UNK	Ν	Y	0	Ν	None Visible					

	Summary of Residential Sampling Survey Form Results and Review of Photographs (Abbreviations and terms used in the table are defined on the last page in this table.)									
Event Number	Location	Number of Broken Windows	Dust Visible Initially	Dust Visible Currently		-	Aggressive Sampling	Level of Dust in Photo	Photo Comments	
AA = asbes C = contrac	AA = asbestos abatement professionals									
LI = large in										
M = modera	ate increase									
MS = buildir	ng managem	ent staff								
N = no										
NI = no Incr	ease									
O = owner/t	O = owner/tenant									
SI = slight ir	SI = slight increase over normal									
UNK = unkr	nown									
Y = yes										

Appendix C. Detailed Analytic Results

Results of Fiber Analyses

Results of fiber analyses in air and dust samples from 30 residential buildings (1–30) in lower Manhattan and 4 comparison buildings (31–34) above 59th Street. The range of values measured in the comparison buildings is shown in the summary of comparison areas above 59th Street.

		Resu	ts From Air Sa	amples	Res	ults From Dust San	nples
Building Number	Sample Location	Fibers in Air PCM (f/cc)	Asbestos in Air TEM (f/cc)	SVF in Air SEM (f/cc)	Asbestos in Dust PLM (%)	Asbestos in Dust TEM (%)	SVF in Dust PLM (%)
Summary o	f Comparison Areas A	bove 59th Stre	eet			1	
	Outside	<.001-0.001	NA	ND-<.000043	Not Sampled	Not Sampled	Not Sampled
	Common	<.001–0.002	NA	ND-0.000043	ND	ND	ND
	Residences	<.001-0.003	NA	ND-0.000087	ND	ND	ND
Results for	Individual Buildings	Sampled in Lov	ver Manhattan	(Buildings 1–3	0)		
1	Outside	0.001			ND	<1	22
1	Common	<.001			Not Sampled	Not Sampled	Not Sampled
1	Residence 1	0.006	<.001	0.000162	ND	NA	20
1	Residence 2	<.001			ND	NA	20
2	Outside	0.003			<1	1.2	28
2	Outside co-located	Not Sampled	Not Sampled	Not Sampled	1.3	NA	25
2	Common	0.005	<.001	0.000255	ND	NA	27
2	Residence 1	<.001			ND	NA	25
2	Residence 2	0.002			ND	NA	20
2	Window sill	Not Sampled	Not Sampled	Not Sampled	<1	<1	30
3	Outside	<.001			Not Sampled	Not Sampled	Not Sampled
3	Res 2 co-located	<.001			Not Sampled	Not Sampled	Not Sampled
3	Common	<.001			Not Sampled	Not Sampled	Not Sampled
3	Residence 1	<.001			Not Sampled	Not Sampled	Not Sampled
3	Residence 2	<.001			Not Sampled	Not Sampled	Not Sampled
4	Outside	0.001			Not Sampled	Not Sampled	Not Sampled
4	Outside co-located	<.001			Not Sampled	Not Sampled	Not Sampled
4	Common	0.001			ND	<1	15
4	Common co-located	0.001			Not Sampled	Not Sampled	Not Sampled
4	Residence 1	<.001			ND	<1	2
4	Residence 2	0.001			ND	<1	5
5	Outside	0.002			3.4	NA	25

5 5 5 6 6 6 6 6 6 6 7	Sample Location Outside co-located Common Residence 1 Residence 2 Outside Common Res 2 co-located Residence 1 Residence 2 Outside Common Res 2 co-located Residence 1 Residence 1 Residence 2 Outside Common Residence 1 Residence 1 Residence 1 Residence 1	Fibers in Air PCM (f/cc) 0.003 0.002 0.002 <.001 <.001 0.002 <.001 <.001 <.001 0.001 0.001	Asbestos in Air TEM (f/cc)	SVF in Air SEM (f/cc)	Asbestos in Dust PLM (%) Not Sampled <1 ND Not Sampled ND Not Sampled ND ND ND	Asbestos in Dust TEM (%) Not Sampled <1 <1 <1 Not Sampled <1 Not Sampled ND ND	SVF in Dust PLM (%) Not Sampled 10 20 Not Sampled 10 Not Sampled 15 15
5 5 5 6 6 6 6 6 6 7	Common Residence 1 Residence 2 Outside Common Res 2 co-located Residence 1 Residence 2 Outside Common Residence 1	0.002 0.002 <.001 <.001 0.002 <.001 <.001 <.001 0.001			Not Sampled <1 ND Not Sampled ND Not Sampled ND	Not Sampled <1 <1 Not Sampled <1 Not Sampled ND	Not Sampled 10 20 Not Sampled 10 Not Sampled 15
5 5 6 6 6 6 6 6 7	Residence 1 Residence 2 Outside Common Res 2 co-located Residence 1 Residence 2 Outside Common Residence 1	0.002 <.001 <.001 <.001 0.002 <.001 <.001 <.001			<1 ND Not Sampled ND Not Sampled ND	<1 <1 Not Sampled <1 Not Sampled ND	10 20 Not Sampled 10 Not Sampled 15
5 6 6 6 6 6 7	Residence 2 Outside Common Res 2 co-located Residence 1 Residence 2 Outside Common Residence 1	<.001 <.001 <.001 0.002 <.001 <.001 <.001			ND Not Sampled ND Not Sampled ND	<1 Not Sampled <1 Not Sampled ND	20 Not Sampled 10 Not Sampled 15
6 6 6 6 6 7	Outside Common Res 2 co-located Residence 1 Residence 2 Outside Common Residence 1	<.001 <.001 0.002 <.001 <.001 <.001			Not Sampled ND Not Sampled ND	Not Sampled <1 Not Sampled ND	Not Sampled 10 Not Sampled 15
6 6 6 6 7	Common Res 2 co-located Residence 1 Residence 2 Outside Common Residence 1	<.001 0.002 <.001 <.001 <.001 0.001			ND Not Sampled ND	<1 Not Sampled ND	10 Not Sampled 15
6 6 6 7	Res 2 co-located Residence 1 Residence 2 Outside Common Residence 1	0.002 <.001 <.001 <.001 0.001			Not Sampled ND	Not Sampled ND	Not Sampled 15
6 6 7	Residence 1 Residence 2 Outside Common Residence 1	<.001 <.001 <.001 0.001			ND	ND	15
6 7	Residence 2 Outside Common Residence 1	<.001 <.001 0.001					15
7	Outside Common Residence 1	<.001 0.001			ND	ND	15
	Common Residence 1	0.001					
	Common Residence 1	0.001			ND	1.7	35
•	Residence 1				ND	ND	5
7					ND	ND	ND
		<.001			ND	ND	5
	Window sill (R2)	Not Sampled	Not Sampled	Not Sampled	ND	ND	40
8	Outside	0.001			Not Sampled	Not Sampled	Not Sampled
	Common	0.001			ND	ND	ND
	Residence 1	0.002			ND	ND	ND
	Residence 2	0.003			ND	ND	ND
9	Outside	<.001			Not Sampled	Not Sampled	Not Sampled
9	Common	0.001			ND	ND	7
9	Common co-located	0.001			Not Sampled	Not Sampled	Not Sampled
9	Residence 1	0.001			ND	<1	2
9	Residence 2	0.003			ND	<1	5
10	Outside	0.001			ND	ND	ND
10	Common	0.002			ND	1.5	20
10	Common, TEM re-ana	lysis				<1	
10	Residence 1	0.001			ND	ND	15
10	Residence 2	0.001			ND	ND	10
11	Outside	<.001			Not Sampled	Not Sampled	Not Sampled
	Common	<.001			ND	ND	ND
	Common co-located	<.001			Not Sampled	Not Sampled	Not Sampled
	Residence 1	<.001			ND	<1	ND
	Residence 2	<.001			ND	1.5	ND
12	Outside	<.001			Not Sampled	Not Sampled	Not Sampled

		Resu	Its From Air Sa	amples	Res	ults From Dust San	nples
Building Number	Sample Location	Fibers in Air PCM (f/cc)	Asbestos in Air TEM (f/cc)	SVF in Air SEM (f/cc)	Asbestos in Dust PLM (%)	Asbestos in Dust TEM (%)	SVF in Dust PLM (%)
12	Common	<.001			ND	ND	ND
12	Residence 1	0.001			ND	ND	5
12	Residence 2	<.001			ND	ND	5
13	Outside	0.001			Not Sampled	Not Sampled	Not Sampled
13	Common	0.001			ND	ND	ND
13	Residence 1	0.003			ND	ND	10
13	Residence 2	0.002			ND	ND	ND
14	Outside	<.001			Not Sampled	Not Sampled	Not Sampled
14	Common	0.0013			ND	ND	5
14	Residence 1	0.001			ND	ND	ND
14	Res 1 co-located	<.001			Not Sampled	Not Sampled	Not Sampled
14	Residence 2	<.001			ND	ND	ND
15	Outside	<.001			ND	1.9	72
15	Common	0.001			ND	ND	ND
15	Residence 1	<.001			ND	<1	ND
15	Residence 2	<.001			ND	ND	5
16	Outside	<.001			ND	ND	1
16	Common	<.001			ND	ND	ND
16	Residence 1	Not Sampled	Not Sampled	Not Sampled	Not Sampled	Not Sampled	Not Sampled
16	Residence 2	<.001		- tot Campion	ND	ND	ND
16	Res 2 co-located	Not Sampled	Not Sampled	Not Sampled	ND	NA	ND
16	Res 2 filter piece	Not Sampled	Not Sampled	Not Sampled	ND	NA	ND
17	Outside	<.001			Not Sampled	Not Sampled	Not Sampled
17	Common	<.001			ND	ND	2
17	Residence 1	<.001			ND	ND	ND
17	Residence 2	<.001			ND	ND	ND
18	Outside	<.001			ND	ND	30
18	Common	<.001			ND	ND	ND
18	Residence 1	0.002			ND	ND	30
18	Residence 2	0.002			ND	ND	35

		Resu	Its From Air Sa	mples	Res	ults From Dust San	nples
Building Number	Sample Location	Fibers in Air PCM (f/cc)	Asbestos in Air TEM (f/cc)	SVF in Air SEM (f/cc)	Asbestos in Dust PLM (%)	Asbestos in Dust TEM (%)	SVF in Dust PLM (%)
19	Outside	<.001			ND	NA	ND
19	Common	overloaded	<.006		ND	ND	ND
19	Residence 1	<.001			ND	ND	ND
19	Residence 2	<.001			ND	ND	ND
20	Outside	<.001			Not Sampled	Not Sampled	Not Sampled
20	Outside co-located	<.001			Not Sampled	Not Sampled	Not Sampled
20	Common	<.001			ND	ND	ND
20	Residence 1	<.001			ND	ND	ND
20	Residence 2	<.001			ND	ND	ND
21	Outside	<.001			Not Sampled	Not Sampled	Not Sampled
21	Common	<.001			ND	ND	ND
21	Residence 1	<.001			ND	ND	ND
21	Residence 2	<.001			ND	ND	ND
22	Outside	<.001			ND	ND	7
22	Common	0.001			ND	ND	3
22	Residence 1	<.001			ND	ND	2
22	Residence 2	<.001			ND	ND	ND
22	Res 2 co-located	<.001			Not Sampled	Not Sampled	Not Sampled
23	Outside	<.001			Not Sampled	Not Sampled	Not Sampled
23	Common	<.001			ND	ND	ND
23	Residence 1	<.001			ND	ND	ND
23	Residence 2	<.001			ND	ND	ND
24	Outside	0.002			ND	ND	55
24	Common	0.001			ND	<1	5
24	Residence 1	0.001			ND	ND	ND
24	Residence 2	0.005	<.001	0.000037	ND	ND	10
24	Res 2 co-located	0.005	<.001		Not Sampled	Not Sampled	Not Sampled
25	Outside	<.001			Not Sampled	Not Sampled	Not Sampled
25	Common	<.001			ND	ND	5
25	Residence 1	<.001			ND	ND	5
25	Residence 2	0.001			ND	ND	ND
26	Outside	0.001			Not Sampled	Not Sampled	Not Sampled
26	Common	0.004	<.001	<0.00004	ND	ND	5
26	Residence 1	0.012	<.001	<0.00004	ND	ND	ND

		Resu	Its From Air Sa	amples	Res	ults From Dust San	nples
Building Number	Sample Location	Fibers in Air PCM (f/cc)	Asbestos in Air TEM (f/cc)	SVF in Air SEM (f/cc)	Asbestos in Dust PLM (%)	Asbestos in Dust TEM (%)	SVF in Dust PLM (%)
26	Residence 2	<.001			ND	ND	ND
27	Outside	<.001			ND	<1	15
27	Common	<.001			ND	<1	10
27	Residence 1	<.001			ND	<1	10
27	Residence 2	<.001			ND	ND	10
28	Outside	<.001			ND	ND	15
28	Common	0.001			ND	ND	10
28	Residence 1	<.001			ND	Insufficient Material	ND
28	Residence 2	0.002			ND	ND	ND
29	Outside	<.001			ND	ND	ND
29	Common	<.001			ND	ND	ND
29	Residence 1	<.001			ND	ND	ND
29	Residence 2	<.001			ND	ND	ND
30	Outside	Not Sampled	Not Sampled	Not Sampled	Not Sampled	Not Sampled	Not Sampled
30	Common	Not Sampled	Not Sampled	Not Sampled	Not Sampled	Not Sampled	Not Sampled
30	Residence 1	<.001			ND	ND	ND
30	Residence 2	<.001			ND	ND	ND
Deculto for	^r Comparison Building	commind Abo	EOth Street	(Duildings 21	24)		
31	Outside	<.001		(Buildings SI-	Not Sampled	Not Sampled	Not Sampled
31	Outside co-located	0.001		0.000039	Not Sampled	Not Sampled	Not Sampled
31	Common	<.001		0.000039	ND	ND	ND
31	Residence 1	<.001		<0.000039	ND	ND	ND
31	Residence 2	<.001		<0.000039	ND	ND	ND
51		<.001		<0.000038	ND	ND	ND
32	Outside	<.001		<0.000039	Not Sampled	Not Sampled	Not Sampled
32	Common	0.002		<0.00004	ND	ND	ND
32	Residence 1	<.001			ND	ND	ND
32	Res 1 co-located	0.001		0.000087	Not Sampled	Not Sampled	Not Sampled
32	Residence 2	Not Sampled	Not Sampled	Not Sampled	Not Sampled	Not Sampled	Not Sampled
00	Outoido	Net Complet			Net Complet	Not Commission	Not Complete
33	Outside	Not Sampled			Not Sampled	Not Sampled	Not Sampled
33	Common	Not Sampled			Not Sampled	Not Sampled	Not Sampled
<u>33</u> 33	Residence 1 Residence 2	Not Sampled <.001		<0.000041	Not Sampled ND	Not Sampled ND	Not Sampled ND
ు		<.001		<0.000041	עא	שאו	שא
34	Outside	<.001		<0.000043	Not Sampled	Not Sampled	Not Sampled

		Resu	ts From Air Sa	amples	Results From Dust Samples				
Building Number	Sample Location	Fibers in Air PCM (f/cc)	Asbestos in Air TEM (f/cc)	SVF in Air SEM (f/cc)	Asbestos in Dust PLM (%)	Asbestos in Dust TEM (%)	SVF in Dust PLM (%)		
34	Common	<.001		0.000043	ND	ND	ND		
34	Residence 1	Not Sampled	Not Sampled	Not Sampled	Not Sampled	Not Sampled	Not Sampled		
34	Residence 2	0.002			ND	Insuf. Material	ND		
34	Res 2 co-located	0.003		<0.000047	Not Sampled	Not Sampled	Not Sampled		

f/cc: fibers per cubic centimeter of air

NA: not analyzed

ND: not detected

PCM: phase contrast microscopy

PLM: polarized light microscopy

SEM: scanning electron microscopy

SVF: synthetic vitreous fibers

TEM: transmission electron microscopy

%: percent

Results of Mineral Analyses in Dust and Air

The following table shows the results of analyses of the minerals in settled dust and air samples for 30 residential buildings in lower Manhattan (1-30) and 4 comparison buildings (31-34) above 59th Street. For each building, the table shows the highest estimated air concentration of mineral for each size fraction. In addition, estimated levels of each mineral in the settled surface dust are shown for each sampled location within a building.

Building Number	Sample Type	Method or Location	Quartz	Cristobalite	Tridymite	Calcite	Portlandite	Gypsum	Mica	Halite
Summary	/ of Comparison Areas	Above 59th Street								
	Air (µg/m³)	PM100	ND	ND	ASN	ND	ND	ND–3 J	ND	ND
	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	ND–3 J	ND	ND
	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	ND–5 J	ND	ND
	Air (µg/m³)	PM2.5	ND	ND	ASN	ND	ND	ND–3 J	ND	ND
	Settled surface dust (%)	Common Area	ND-1 J	ND	ND	ND-0.4 J	ND-0.05 J	ND–3 J	ND	ND
	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
	Settled surface dust (%)	Residence 1&2	ND–2 J	ND	ND	ND-0.9 J	ND-0.08 J	2–4 J	ND-0.08 J	ND-0.4 J
				•	•	•				
Results f	or Individual Buildings	Sampled in Lower	Manhattan (B	uildinas 1–30)						
1	Air (µg/m³)	PM100	ND	ND	ASN	ND	ND	6 J	ND	ND
1	Air (μg/m³)	PM10	ND	ND	ASN	ND	ND	14 J	ND	ND
1	Air (µg/m³)	PM4	ND	ND	ASN	6 J	84 J	12 J	ND	ND
1	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
1	Settled surface dust (%)	Common area	NS	NS	NS	NS	NS	NS	NS	NS
1	Settled surface dust (%)	Outdoor	27 J	ND	ND	15 J	6 J	14 J	ND	ND
1	Settled surface dust (%)	Residence 1	19 J	ND	ND	18 J	2 J	30 J	0.3 J	ND
1	Settled surface dust (%)	Residence 2	31 J	ND	ND	21 J	6 J	4 J	0.05 J	ND
2	Air (µg/m³)	PM100	ND	ND	ASN	ND	ND	ND	ND	ND
2	Air (μg/m³)	PM10	ND	ND	ASN	ND	ND	ND	ND	ND
2	Air (μg/m³)	PM4	ND	ND	ASN	ND	ND	7J	ND	ND
2	Air (µg/m³) Settled surface dust	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
2	(%)	Common area	1 J	ND	ND	ND	2 J	14 J	ND	ND
2	Settled surface dust (%)	Outdoor	21 J	ND	ND	19 J	3 J	27 J	ND	<0.03 J
2	Settled surface dust (%)	Residence 1	3 J	ND	ND	2 J	0.9 J	6 J	ND	ND
2	Settled surface dust (%)	Residence 2	23 J	ND	ND	15 J	4 J	9 J	ND	ND
3	Air (µg/m³)	PM100	6 J	15 J	ASN	5 J	24 J	5 J	13 J	ND
3	Air (µg/m³)	PM10	5 J	ND	ASN	5 J	25 J	6 J	ND	ND
3	Air (µg/m³)	PM4	6 J	ND	ASN	6 J	26 J	7 J	ND	ND
3	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS

Building Number	Sample Type	Method or Location	Quartz	Cristobalite	Tridymite	Calcite	Portlandite	Gypsum	Mica	Halite
3	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
	Settled surface dust									
3	(%) Settled surface dust	Common	NS	NS	NS	NS	NS	NS	NS	NS
3	(%)	Residence 1	NS	NS	NS	NS	NS	NS	NS	NS
3	Settled surface dust (%)	Residence 2	NS	NS	NS	NS	NS	NS	NS	NS
4	Air (µg/m³)	PM100	4 J	ND	ASN	4 J	16 J	3 J	9 J	5 J
4	Air (µg/m³)	PM10	4 J	ND	ASN	3 J	16 J	3 J	ND	4 J
4	Air (µg/m³)	PM4	6 J	ND	ASN	6 J	28 J	5 J	14 J	8 J
4	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
4	Settled surface dust (%)	Common area	ND	ND	ND	1 J	0.4 J	8 J	ND	ND
	Settled surface dust									
4	(%) Settled surface dust	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
4	(%)	Residence 1	0.2 J	ND	ND	0.06 J	0.7 J	1 J	ND	ND
4	Settled surface dust (%)	Residence 2	14 J	ND	ND	5 J	2 J	9 J	ND	0.1 J
5	Air (µg/m³)	PM100	ND	ND	ASN	14 J	95 J	14 J	ND	14 J
5	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	14 J	ND	ND
5	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	15 J	ND	ND
5	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
5	Settled surface dust (%)	Common are a	NS	NS	NS	NS	NS	NS	NS	NS
	Settled surface dust									
5	(%) Settled surface dust	Outdoor	23 J	ND	ND	8 J	ND	ND	ND	ND
5	(%) Settled surface dust	Residence 1	2 J	ND	ND	ND	ND	9 J	ND	ND
5	(%)	Residence 2	11 J	ND	ND	4 J	0.5 J	2 J	ND	ND
6	Air (µg/m³)	PM100	ND	ND	ASN	ND	ND	ND	ND	ND
6	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	ND	ND	ND
6	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	ND	ND	ND
6	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
6	Settled surface dust (%)	Common area	17 J	ND	ND	10 J	3 J	20 J	0.6 J	ND
6	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
0	Settled surface dust		113	NO	ING	113		ING	ING	113
6	(%) Settled surface dust	Residence 1	27 J	ND	ND	17 J	8 J	13 J	ND	ND
6	(%)	Residence 2	24 J	ND	ND	15 J	4 J	4 J	0.03 J	ND
7	Air (µg/m³)	PM100	ND	ND	ASN	8 J	54 J	10 J	ND	ND
7	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	11 J	ND	ND
7	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	11 J	ND	ND
7	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
7	Settled surface dust (%)	Common area	11 J	ND	ND	8 J	3 J	15 J	0.09 J	0.04 J
	Settled surface dust	Outdoor								
7	(%) Settled surface dust		14 J	ND	ND	7 J	1 J	12 J	ND	0.1 J
7	(%)	Residence 1	14 J	ND	ND	13 J	2 J	2 J	ND	ND

Building Number	Sample Type	Method or Location	Quartz	Cristobalite	Tridymite	Calcite	Portlandite	Gypsum	Mica	Halite
7	Settled surface dust (%)	Residence 2	21 J	ND	ND	16 J	3 J	4 J	ND	0.09 J
7	Settled surface dust (%)	Residence 2-co	18 J	ND	ND	9 J	1 J	2 J	ND	ND
1	(76)	Residence 2-00	10.5	ND	ND	30	15	2 J	ND	ND
8	Air (μg/m³)	PM100	ND	ND	ASN	7 J	ND	10 J	ND	ND
8	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	9 J	ND	ND
8	Air (µg/m³)	PM4	ND	ND	ASN	6 J	ND	12 J	43 J	19 J
8	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
8	Settled surface dust (%)	Common area	19 J	ND	ND	9 J	2 J	15 J	ND	ND
8	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
8	Settled surface dust (%)	Residence 1	15 J	ND	ND	10 J	3 J	17 J	ND	ND
8	Settled surface dust (%)	Residence 2	28 J	ND	ND	21 J	4 J	6 J	ND	ND
9	Air (µg/m³)	PM100	ND	ND	ASN	ND	ND	3 J	ND	ND
9	Air (µg/m³)	PM10	ND	ND	ASN	4 J	ND	6 J	ND	ND
9	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	ND	ND	ND
9	Air (µg/m³) Settled surface dust	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
9	(%) Settled surface dust	Common area	25 J	ND	ND	9 J	4 J	16 J	0.09 J	0.06 J
9	(%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
9	Settled surface dust (%)	Residence 1	10 J	ND	ND	2 J	0.9 J	5 J	ND	ND
9	Settled surface dust (%)	Residence 2	4 J	ND	ND	ND	ND	2 J	ND	ND
10	Air (µg/m³)	PM100	ND	ND	ASN	ND	ND	ND	ND	ND
10	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	ND	ND	ND
10	Air (μg/m³)	PM4	ND	ND	ASN	8 J	ND	ND	ND	ND
10	Air (µg/m³) Settled surface dust	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
10	(%)	Common area	2 J	ND	ND	3 J	ND	1 J	ND	ND
10	Settled surface dust (%)	Common area-co	NS	NS	NS	NS	NS	NS	NS	NS
10	Settled surface dust (%)	Outdoor	21 J	ND	ND	14 J	3 J	5 J	0.09 J	0.05 J
10	Settled surface dust (%)	Residence 1	3 J	ND	ND	ND	0.06 J	ND	ND	ND
10	Settled surface dust (%)	Residence 2	2 J	ND	ND	ND	ND	0.8 J	ND	0.04 J
11	Air (μg/m³)	PM100	NS	NS	NS	NS	NS	NS	NS	NS
	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	ND	ND	ND
	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	ND	ND	ND
	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
11	Settled surface dust (%)	Common area	2 J	ND	ND	ND	ND	1 J	ND	ND
11	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
11	Settled surface dust (%)	Residence 1	0.2 J	ND	ND	0.02 J	ND	2 J	ND	ND
11	Settled surface dust (%)	Residence 2	0.07 J	ND	ND	ND	ND	ND	ND	ND

Building Number	Sample Type	Method or Location	Quartz	Cristobalite	Tridymite	Calcite	Portlandite	Gypsum	Mica	Halite
12	Air (µg/m³)	PM100	NS	NS	NS	NS	NS	NS	NS	NS
12	Air (µg/m³)	PM10	3 J	ND	ASN	5 J	18 J	4 J	8 J	5 J
12	Air (µg/m³)	PM4	5 J	ND	ASN	7 J	28 J	6 J	15 J	7 J
12	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
12	Settled surface dust (%)	Common area	4 J	ND	ND	2 J	3 J	1 J	ND	ND
12	Settled surface dust	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
12	(%) Settled surface dust (%)	Residence 1	ND	ND	ND	ND	ND	0.3 J	ND	ND
12	Settled surface dust (%)	Residence 2	0.4 J	ND	ND	0.09 J	0.06 J	0.8 J	ND	ND
	(70)		0.10			0.000	0.000	0.00	110	
13	Air (µg/m³)	PM100	5 J	ND	ASN	5 J	16 J	3 J	ND	ND
13	Air (µg/m³)	PM10	4 J	ND	ASN	5 J	16 J	3 J	ND	ND
13	Air (µg/m³)	PM4	7 J	ND	ASN	10 J	26 J	5 J	ND	ND
13	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
13	Settled surface dust (%)	Common area	0.05 J	ND	ND	0.3 J	0.04 J	1 J	ND	ND
13	Settled surface dust	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
13	Settled surface dust	Residence 1	ND	ND	ND	ND	ND	1 J	ND	ND
13	Settled surface dust									
13	(%)	Residence 2	ND	ND	ND	ND	ND	0.8 J	ND	ND
14	Air (μg/m³)	PM100	ND	ND	ASN	ND	ND	ND	ND	ND
14	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	ND	ND	ND
14	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	ND	ND	ND
14	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
14	Settled surface dust (%)	Common area	0.03 J	ND	ND	ND	ND	2 J	ND	0.04 J
14	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
14	Settled surface dust (%)	Residence 1	ND	ND	ND	ND	ND	ND	ND	ND
14	Settled surface dust (%)	Residence 2	ND	ND	ND	ND	ND	0.08 J	ND	ND
	(70)							0.000	110	
15	Air (μg/m³)	PM100	8 J	ND	ASN	3 J	18 J	ND	ND	ND
15	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	3 J	ND	ND
15	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	5 J	ND	ND
15	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
15	Settled surface dust (%)	Common area	0.4 J	ND	ND	0.02 J	ND	1 J	ND	ND
15	Settled surface dust (%)	Outdoor	1 J	ND	ND	ND	0.3 J	2 J	ND	ND
15	Settled surface dust (%)	Residence 1	ND	ND	ND	ND	ND	1 J	ND	ND
15	Settled surface dust (%)	Residence 2	ND	ND	ND	ND	ND	ND	ND	ND
16	Air (μg/m³)	PM100	4 J	ND	ASN	3 J	17 J	3 J	ND	ND
16	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	4 J	ND	ND
16	Air (µg/m³)	PM4	18 J	ND	ASN	5 J	27 J	7 J	ND	ND

Building Number	Sample Type	Method or Location	Quartz	Cristobalite	Tridymite	Calcite	Portlandite	Gypsum	Mica	Halite
16	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
16	Settled surface dust (%)	Common are a	ND	ND	ND	ND	ND	0.09 J	ND	ND
16	Settled surface dust (%)	Outdoor	3 J	ND	ND	1 J	0.6 J	0.04 J	0.06 J	0.04 J
16	Settled surface dust (%)	Residence 1	NS	NS	NS	NS	NS	NS	NS	NS
16	Settled surface dust (%)	Residence 2	ND	ND	ND	ND	ND	0.9 J	ND	ND
16	Settled surface dust (%)	R2 filter piece	2 J	ND	ND	0.3 J	0.4 J	1 J	ND	ND
17	Air (µg/m³)	PM100	NS	NS	NS	NS	NS	NS	NS	NS
17	Air (µg/m³)	PM10	12 J	ND	ASN	4 J	17 J	6 J	ND	5 J
17	Air (µg/m³)	PM4	5 J	ND	ASN	5 J	27 J	6 J	15 J	ND
17	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
17	Settled surface dust (%)	Common area	2 J	ND	ND	ND	ND	1 J	0.06 J	ND
17	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
17	Settled surface dust (%)	Residence 1	0.9 J	ND	ND	ND	0.05 J	2 J	0.05 J	0.03 J
17	Settled surface dust (%)	Residence 2	1 J	ND	ND	0.8 J	ND	1 J	ND	ND
18	Air (µg/m³)	PM100	3 J	ND	ASN	ND	ND	ND	ND	ND
18	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	3 J	ND	ND
18	Air (µg/m³)	PM4	5 J	ND	ASN	ND	ND	5 J	ND	ND
18	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
18	Settled surface dust (%)	Common area	0.9 J	ND	ND	ND	ND	ND	ND	ND
18	Settled surface dust (%)	Outdoor	7 J	ND	ND	1 J	ND	ND	0.2 J	ND
18	Settled surface dust (%)	Residence 1	ND	ND	ND	ND	ND	0.3 J	ND	ND
18	Settled surface dust (%)	Residence 2	ND	ND	ND	ND	ND	1 J	ND	ND
19	Air (µg/m³)	PM100	ND	ND	ASN	ND	ND	ND	ND	ND
19	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	ND	ND	ND
19	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	ND	ND	ND
19	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
19	Settled surface dust (%) Settled surface dust	Common area	1 J	ND	ND	0.08 J	0.05 J	1 J	ND	ND
19	(%)	Outdoor	5 J	ND	ND	1 J	0.6 J	1 J	0.09 J	0.04 J
19	Settled surface dust (%)	Residence 1	ND	ND	ND	ND	ND	1 J	ND	ND
19	Settled surface dust (%)	Residence 2	ND	ND	ND	ND	ND	2 J	ND	ND
20	Air (µg/m³)	PM100	3 J	ND	ASN	3 J	16 J	3 J	ND	ND
20	Air (μg/m ³)	PM100	3 J	ND	ASN	3 J	16 J	4 J	ND	ND
20	Air (µg/m³)	PM4	5 J	ND	ASN	5 J	29 J	6J	ND	ND
20	Air (μg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
20	Settled surface dust (%)	Common area	ND	ND	ND	ND	ND	0.07 J	ND	0.04 J

Building Number	Sample Type	Method or Location	Quartz	Cristobalite	Tridymite	Calcite	Portlandite	Gypsum	Mica	Halite
20	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
20	Settled surface dust (%)	Residence 1	0.05 J	ND	ND	ND	ND	1 J	ND	ND
	Settled surface dust									
20	(%)	Residence 2	0.9 J	ND	ND	0.6 J	0.09 J	1 J	ND	0.05 J
- 24	A:= (DM100	ND	ND	ACN	ND		ND	ND	ND
21 21	Air (μg/m³) Air (μg/m³)	PM100 PM10	ND R	ND R	ASN R	ND R	ND R	ND R	ND R	ND R
21	Air (μg/m³)	PM10	ND	ND	ASN	ND	ND	5 J	ND	ND
21	Air (µg/m³)	PM2.5	ND	ND	ASN	ND	ND	ND	ND	ND
21	Settled surface dust	1 11/2.5	ND	ND	Adiv	ND		ND		ND
21	(%) Settled surface dust	Common area	0.9 J	ND	ND	ND	ND	2 J	ND	ND
21	(%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
21	Settled surface dust (%)	Residence 1	ND	ND	ND	ND	ND	ND	ND	ND
21	Settled surface dust (%)	Residence 2	ND	ND	ND	ND	ND	1 J	ND	ND
22	Air (µg/m³)	PM100	13 J	ND	ASN	ND	ND	ND	ND	ND
22	Air (µg/m³)	PM10	3 J	ND	ASN	ND	ND	ND	ND	ND
22	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	ND	ND	ND
22	Air (µg/m³)	PM2.5	ND	ND	ASN	ND	ND	ND	ND	ND
22	Settled surface dust (%)	Common area	14 J	ND	ND	1 J	0.4 J	ND	ND	ND
22	Settled surface dust (%)	Outdoor	26 J	ND	ND	2 J	0.8 J	0.03 J	0.07 J	ND
	Settled surface dust									
22	(%) Settled surface dust	Residence 1	2 J	ND	ND	0.07 J	0.08 J	1 J	ND	ND
22	(%)	Residence 2	0.6 J	ND	ND	ND	ND	2 J	ND	ND
23	Air (µg/m³)	PM100	ND	ND	ASN	ND	ND	ND	ND	ND
23	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	ND	ND	ND
23	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	ND	ND	ND
23	Air (µg/m³) Settled surface dust	PM2.5	ND	ND	ASN	ND	ND	ND	ND	ND
23	(%) Settled surface dust	Common area	2 J	ND	ND	0.04 J	0.05 J	0.9 J	ND	ND
23	(%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
23	Settled surface dust (%)	Residence 1	ND	ND	ND	ND	ND	ND	ND	ND
23	Settled surface dust (%)	Residence 2	ND	ND	ND	ND	ND	1 J	ND	ND
20	(10)									
24	Air (µg/m³)	PM100	ND	ND	ASN	ND	ND	ND	ND	ND
24	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	ND	ND	ND
24	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	ND	ND	ND
24	Air (µg/m³)	PM2.5	3 J	ND	ASN	ND	ND	ND	ND	ND
24	Settled surface dust (%)	Common area	0.03 J	ND	ND	ND	ND	ND	ND	ND
	Settled surface dust									
24	(%) Settled surface dust	Outdoor	2 J	ND	ND	0.8 J	0.07 J	ND	0.05 J	0.04 J
24	(%) Settled surface dust	Residence 1	ND	ND	ND	ND	ND	ND	ND	ND
24	(%)	Residence 2	ND	ND	ND	ND	ND	ND	ND	ND

Building Number	Sample Type	Method or Location	Quartz	Cristobalite	Tridymite	Calcite	Portlandite	Gypsum	Mica	Halite
25	Air (µg/m³)	PM100	ND	ND	ASN	ND	ND	3 J	ND	ND
25	Air (µg/m³)	PM10	R	R	R	R	R	R	R	R
25	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	4 J	ND	ND
25	Air (µg/m³)	PM2.5	ND	ND	ASN	ND	ND	3 J	ND	ND
25	Settled surface dust (%)	Common area	ND	ND	ND	ND	ND	2 J	ND	ND
25	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
	Settled surface dust									
25	(%) Settled surface dust	Residence 1	ND	ND	ND	ND	ND	ND	ND	ND
25	(%)	Residence 2	ND	ND	ND	ND	ND	2 J	ND	ND
26	Air (µg/m³)	PM100	8 J	ND	ASN	ND	ND	ND	ND	ND
26	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	ND	ND	ND
26	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	ND	ND	ND
26	Air (µg/m³) Settled surface dust	PM2.5	ND	ND	ASN	ND	ND	ND	ND	ND
26	(%)	Common area	0.7 J	ND	ND	2 J	0.6 J	2 J	0.07 J	ND
26	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
26	Settled surface dust (%)	Residence 1	ND	ND	ND	ND	ND	ND	ND	ND
26	Settled surface dust (%)	Residence 2	ND	ND	ND	ND	ND	0.05 J	ND	ND
27	Air (µg/m³)	PM100	3 J	ND	ASN	ND	ND	ND	ND	ND
27	Air (μg/m³)	PM10	ND	ND	ASN	ND	ND	ND	ND	ND
27	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	ND	ND	ND
27	Air (µg/m³)	PM2.5	ND	ND	ASN	ND	ND	ND	ND	ND
27	Settled surface dust (%)	Common area	0.04 J	ND	ND	0.02 J	ND	0.1 J	ND	ND
27	Settled surface dust (%)	Outdoor	4 J	ND	ND	1 J	0.8 J	0.04 J	0.07 J	ND
27	Settled surface dust (%)	Residence 1	ND	ND	ND	ND	ND	0.6 J	ND	ND
27	Settled surface dust	Residence 2	ND	ND	ND	ND	ND	1 J	ND	ND
21	(%)		THE	NB				10		
28	Air (µg/m³)	PM100	3 J	ND	ASN	ND	ND	3 J	ND	4 J
28	Air (μg/m³)	PM10	ND	ND	ASN	ND	ND	3 J	ND	ND
28	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	5 J	ND	ND
28	Air (µg/m³)	PM2.5	ND	ND	ASN	ND	ND	ND	ND	ND
	Settled surface dust									
28	(%) Settled surface dust	Common area	ND	ND	ND	ND	ND	1 J	ND	ND
28	(%) Settled surface dust	Outdoor	8 J	ND	ND	3 J	2 J	0.3 J	0.09 J	ND
28	(%)	Residence 1	ND	ND	ND	ND	ND	ND	ND	ND
28	Settled surface dust (%)	Residence 2	ND	ND	ND	ND	ND	1 J	ND	ND
29	Air (µg/m³)	PM100	4 J	ND	ASN	ND	ND	ND	ND	ND
29	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	3 J	ND	ND
29	Air (µg/m³)	PM4	19 J	ND	ASN	ND	ND	5 J	ND	ND

Building Number	Sample Type	Method or Location	Quartz	Cristobalite	Tridymite	Calcite	Portlandite	Gypsum	Mica	Halite
29	Air (µg/m³)	PM2.5	ND	ND	ASN	ND	ND	ND	ND	ND
29	Settled surface dust (%)	Common area	1 J	ND	ND	0.09 J	0.06 J	2 J	ND	ND
29	Settled surface dust (%)	Outdoor	8 J	ND	ND	3 J	2 J	2 J	0.3 J	0.07 J
29	Settled surface dust (%)	Residence 1	0.2 J	ND	ND	ND	ND	1 J	ND	ND
29	Settled surface dust (%)	Residence 2	ND	ND	ND	ND	ND	ND	ND	ND
30	Air (µg/m³)	PM100	NS	NS	NS	NS	NS	NS	NS	NS
30	Air (µg/m³)	PM10	3 J	ND	ASN	ND	ND	3 J	ND	ND
30	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	5 J	ND	ND
30	Air (µg/m³)	PM2.5	ND	ND	ASN	ND	ND	ND	ND	ND
30	Settled surface dust (%)	Common area	NS	NS	NS	NS	NS	NS	NS	NS
30	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
30	Settled surface dust (%)	Residence 1	1 J	ND	ND	ND	0.9 J	4 J	0.06 J	0.05 J
30	Settled surface dust (%)	Residence 2	0.6 J	ND	ND	0.4 J	0.4 J	4 J	ND	ND
Results	for Individual Co	mparison Build	ings Samp	led Above 5	9th Stree	t (Buildin	igs 31–34)	<u> </u>		
		•	<u> </u>			•	• /			
31	Air (µg/m³)	PM100	NS	NS	NS	NS	NS	NS	NS	NS
31	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	3 J	ND	ND
31	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	5 J	ND	ND
31	Air (µg/m³)	PM2.5	NS	NS	NS	NS	NS	NS	NS	NS
31	Settled surface dust (%)	Common area	1 J	ND	ND	0.03 J	0.05 J	2 J	ND	0.04 J
31	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
31	Settled surface dust (%)	Residence 1	ND	ND	ND	ND	ND	2 J	ND	ND
31	Settled surface dust (%)	Residence 2	ND	ND	ND	ND	ND	ND	ND	ND
32	Air (µg/m³)	PM100	ND	ND	ASN	ND	ND	3 J	ND	ND
32	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	3 J	ND	ND
32	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	5 J	ND	ND
32	Air (µg/m³)	PM2.5	ND	ND	ASN	ND	ND	3 J	ND	ND
32	Settled surface dust (%)	Common area	1 J	ND	ND	0.4 J	ND	ЗJ	ND	ND
32	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
32	Settled surface dust (%)	Residence 1	2 J	ND	ND	0.9 J	0.08 J	4 J	0.08 J	0.4 J
32	Settled surface dust (%)	Residence 2	NS	NS	NS	NS	NS	NS	NS	NS
33	Air (μg/m³)	PM100	ND	ND	ASN	ND	ND	ND	ND	ND
33	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	ND	ND	ND
33	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	ND	ND	ND
33	Air (µg/m³)	PM2.5	ND	ND	ASN	ND	ND	ND	ND	ND
33	Settled surface dust (%)	Common area	NS	NS	NS	NS	NS	NS	NS	NS

Building Number	Sample Type	Method or Location	Quartz	Cristobalite	Tridymite	Calcite	Portlandite	Gypsum	Mica	Halite
33	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
33	Settled surface dust (%)	Residence 1	NS	NS	NS	NS	NS	NS	NS	NS
33	Settled surface dust (%)	Residence 2	1 J	ND	ND	ND	0.08 J	2 J	ND	ND
34	Air (µg/m³)	PM100	ND	ND	ASN	ND	ND	ND	ND	ND
34	Air (µg/m³)	PM10	ND	ND	ASN	ND	ND	ND	ND	ND
34	Air (µg/m³)	PM4	ND	ND	ASN	ND	ND	ND	ND	ND
34	Air (µg/m³)	PM2.5	ND	ND	ASN	ND	ND	ND	ND	ND
34	Settled surface dust (%)	Common area	ND	ND	ND	ND	ND	ND	ND	ND
34	Settled surface dust (%)	Outdoor	NS	NS	NS	NS	NS	NS	NS	NS
34	Settled surface dust (%)	Residence 1	NS	NS	NS	NS	NS	NS	NS	NS
34	Settled surface dust (%)	Residence 2	ND	ND	ND	ND	ND	3 J	ND	ND

ASN: Analytical sensitivity not available. Mineral was not detected at any measured location using the associated method, analytical sensitivity is not known.

Detect: Mineral was detected above the analytical sensitivity in at least one of the measured locations of the building using the associated method.

J: Estimated value. Due to inconsistencies during sample collection/analysis, the result is only an estimate of the actual value. The actual value could be higher or lower than the value shown.

ND: Not detected. Mineral was not detected at any measured location using the associated method.

NS: Not sampled. The associated method was not used at any of the measured locations in the building. R: Result was rejected. Due to inconsistencies during sample collection/analysis, the result values for the sample were rejected.

 $\mu g/m^3$: microgram per cubic meter of air.

Mineral Analyses in Air

Estimated levels of minerals in air for samples from 30 residential buildings (1–30) in lower Manhattan and 4 comparison residential buildings (31–34) above 59th Street are shown in the following table. The estimated air concentration of each mineral by size fraction is shown for each area sampled. Typically, samples were obtained from an outdoor area, an indoor common area, and two residences. In addition, co-located samples were obtained from some locations. However, due to inconsistencies during sample collection or analysis, the specific sampling location within the building is not noted.

Building Number	Particle Size Fraction	Sample Volume (L)	Quartz (µg/m³)	Cristobalite (µg/m³)	Tridymite (μg/m³)	Calcite µg/m³)	Portlandite (µg/m³)	Gypsum (µg/m³)	Mica µg/m³)	Halite (µg/m³))
Summary of	Comparison A	reas Above	59th Street		-					•
	PM100							ND-3 J		
	PM10							ND-3 J		
	PM4							ND-5 J		
	PM2.5							ND-3 J		
Results for I	ndividual Buil	dings Sam	oled in Lower N	lanhattan (Buildin	gs 1–30)					
1	PM100	882						6 J		
1	PM100	830								
1	PM100	818								
1	PM100	892								
1	PM10	861								
1	PM10	835						7 J		
1	PM10	838						14 J		
1	PM10	899						8 J		
1	PM4	809					84 J	10 J		
1	PM4	813						12 J		
1	PM4	820				6 J	68 J	10 J		
1	PM4	837								
2	PM100	1229								
2	PM100	1162								
2	PM100	766								
2	PM100	1240								
2	PM10	1194								
2	PM10	1168								
2	PM10	1390								
2	PM10	1216								
2	PM4	757						7J		
2	PM4	717								
2	PM4	849								

Building Number	Particle Size Fraction	Sample Volume (L)	Quartz (µg/m³)	Cristobalite (μg/m³)	Tridymite (µg/m³)	Calcite µg/m³)	Portlandite (µg/m³)	Gypsum (µg/m³)	Mica µg/m³)	Halite (µg/m³))
2	PM4	762								
3	PM100	868						5 J		
3	PM100	821	6 J	15 J		5 J	24 J	5 J	13 J	
3	PM100	756						5 J		
3	PM100	761								
3	PM10	858						6 J		
3	PM10	815	5 J			5J	25 J	5 J		
3	PM10	765						5 J		
3	PM10	783								
3	PM4	814	5 J					6 J		
3	PM4	824	6 J			6 J	26 J	7 J		
3	PM4	841						5 J		
3	PM4	823								
4	PM100	1230						3 J		
4	PM100	1289	4 J			4 J	16 J	3 J	9 J	5J
4	PM100	1278								
4	PM100	1221						3 J		5J
4	PM10	1218					16 J	3 J		
4	PM10	1415	4 J			3 J	16 J	3 J		4J
4	PM10	1294						3 J		
4	PM10	1201								
4	PM4	751					27J	5J		
4	PM4	793	6 J			6J	28J	5J	14 J	8 J
4	PM4	804						5 J		
4	PM4	757								
5	PM100	783				14 J	95 J	14 J		14 J
5	PM100	786								
5	PM100	804								
5	PM100	800								
5	PM10	780								
5	PM10	797								
5	PM10	846						14 J		
5	PM10	789								
5	PM4	817								
5	PM4	664								
5	PM4	813						9 J		
5	PM4	813						15 J		
5		2.0								
6	PM100	1241								

e	Fraction	Volume (L)	Quartz (µg/m³)	Cristobalite (µg/m³)	Tridymite (µg/m³)	Calcite µg/m³)	Portlandite (µg/m³)	Gypsum (µg/m³)	Mica µg/m³)	Halite (µg/m³))
6	PM100	1212								
6	PM100	1238								
6	PM100	NS	NS	NS	NS	NS	NS	NS	NS	NS
6	PM10	1226								
6	PM10	1197								
6	PM10	1252								
6	PM10	1254								
6	PM4	775								
6	PM4	762								
6	PM4	775								
6	PM4	772								
7	PM100	1277								
7	PM100	1260				8 J	54 J	10 J		
	PM100	1243								
7	PM100	1213								
7	PM10	1227								
7	PM10	1225						11 J		
7	PM10	1240								
7	PM10	1231								
7	PM4	787								
7	PM4	784						10 J		
7	PM4	765						11 J		
7	PM4	748								
										+
8	PM100	1213				7 J		10 J		
8	PM100	1254						6 J		
8	PM100	1223						7 J		
8	PM100	1206								
8	PM100	1175								
8	PM100	1259								
8	PM10	1188						9 J		
8	PM10	1223						6 J		
8	PM10	1172								
8	PM10	1256								
8	PM4	757						12 J		
8	PM4	784				6 J		10 J		
8	PM4	738								
8	PM4	777						10J	43J	19 J
9	PM100	1210								

Building Number	Particle Size Fraction	Sample Volume (L)	Quartz (µg/m³)	Cristobalite (μg/m³)	Tridymite (µg/m³)	Calcite µg/m³)	Portlandite (µg/m³)	Gypsum (µg/m³)	Mica µg/m³)	Halite (µg/m³))
9	PM100	NS	NS	NS	NS	NS	NS	NS	NS	NS
9	PM100	NS	NS	NS	NS	NS	NS	NS	NS	NS
9	PM10	1203								
9	PM10	1217				4 J		6 J		
9	PM10	1231								
9	PM10	1131								
9	PM4	776								
9	PM4	782								
9	PM4	763								
9	PM4	781								
10	PM100	1201								
10	PM100	1225								
10	PM100	1220								
10	PM100	1226								
10	PM10	1181								
10	PM10	1202								
10	PM10	1227								
10	PM10	1224								
10	PM4	757								
10	PM4	766								
10	PM4	762				8 J				
10	PM4	767								
11	PM100	NS	NS	NS	NS	NS	NS	NS	NS	NS
11	PM100	NS	NS	NS	NS	NS	NS	NS	NS	NS
11	PM100	NS	NS	NS	NS	NS	NS	NS	NS	NS
11	PM100	NS	NS	NS	NS	NS	NS	NS	NS	NS
11	PM10	1305								
11	PM10	1301								
11	PM10	1337								
11	PM10	1237								
11	PM4	832								
11	PM4	806								
11	PM4	834								
11	PM4	779								
					1					1
12	PM100	NS	NS	NS	NS	NS	NS	NS	NS	NS
12	PM100	NS	NS	NS	NS	NS	NS	NS	NS	NS
12	PM100	NS	NS	NS	NS	NS	NS	NS	NS	NS
12	PM100	NS	NS	NS	NS	NS	NS	NS	NS	NS
12	PM10	1412	3 J			3 J	14 J	3 J		

Building Number	Particle Size Fraction	Sample Volume (L)	Quartz (µg/m³)	Cristobalite (μg/m³)	Tridymite (µg/m³)	Calcite µg/m³)	Portlandite (µg/m³)	Gypsum (µg/m³)	Mica µg/m³)	Halite (µg/m³))
12	PM10	1315	3 J			5 J	18 J	4 J	8 J	5 J
12	PM10	1337	3 J			3 J	17 J	3 J		5 J
12	PM10	1553	R	R	R	R	R	R	R	R
12	PM10	1479	R	R	R	R	R	R	R	R
12	PM4	925	4 J			4 J	22 J	4 J		
12	PM4	828	5 J			7 J	28 J	6 J	15 J	7 J
12	PM4	948	R	R	R	R	R	R	R	R
12	PM4	932	R	R	R	R	R	R	R	R
13	PM100	1281						3 J		
13	PM100	1300								
13	PM100	1298								
13	PM100	1284	5 J			5 J	16 J	3 J		
13	PM100	1211								
13	PM100	1203								
13	PM100	1184								
13	PM10	1263						3 J		
13	PM10	1325	4 J			5 J	16 J	3 J		
13	PM10	1223								
13	PM10	1205								
13	PM4	802				5 J		5 J		
13	PM4	818	7 J			10 J	26 J	5 J		
13	PM4	762								
13	PM4	761								
14	PM100	1151								
14	PM100	1169								
14	PM100	1166								
14	PM100	1142								
14	PM100	1171								
14	PM100	1149								
14	PM10	1134								
14	PM10	1138								
14	PM10	1155								
14	PM10	1179								
14	PM10	1161								
14	PM4	719								
14	PM4	731								
14	PM4	732								
14	PM4	730								
15	PM100	1219								

Building Number	Particle Size Fraction	Sample Volume (L)	Quartz (µg/m³)	Cristobalite (μg/m³)	Tridymite (µg/m³)	Calcite µg/m³)	Portlandite (µg/m³)	Gypsum (µg/m³)	Mica µg/m³)	Halite (µg/m³))
15	PM100	1199	8 J			3 J	18 J			
15	PM100	1211								
15	PM100	1156								
15	PM10	1218						3 J		
15	PM10	1115	R	R	R	R	R	R	R	R
15	PM10	1212								
15	PM10	1225								
15	PM10	1171								
15	PM4	783						5 J		
15	PM4	778								
15	PM4	766								
15	PM4	751								
16	PM100	1254						3 J		
16	PM100	1225	4 J			3 J	17 J	3 J		
16	PM100	1217								
16	PM10	1269						4 J		
16	PM10	1228						4 J		
16	PM10	1210								
16	PM4	808						7 J		
16	PM4	774						5 J		
16	PM4	769	18 J			5 J	27 J	7 J		
16	PM4	769								
17	PM10	1306						4 J		
17	PM10	1469	3 J			3 J	16 J	3 J		4 J
17	PM10	1250								
17	PM10	1259						3 J		
17	PM10	1293	12 J			4 J	17 J	6 J		5 J
17	PM4	844						5 J		
17	PM4	816	5 J			5 J	27 J	6 J	15 J	
17	PM4	786								
17	PM4	786						5 J		
17	PM4	803					26 J	6 J		
18	PM100	1277								
18	PM100	1247	3 J							
18	PM100	1241								
18	PM100	1265								
18	PM100	1177								
18	PM10	1234								
18	PM10	1269								

Building Number	Particle Size Fraction	Sample Volume (L)	Quartz (µg/m³)	Cristobalite (μg/m³)	Tridymite (μg/m³)	Calcite µg/m³)	Portlandite (µg/m³)	Gypsum (µg/m³)	Mica µg/m³)	Halite (µg/m³))
18	PM10	1258								
18	PM10	1268								
18	PM10	1196						3 J		
18	PM4	791								
18	PM4	780								
18	PM4	785	5 J							
18	PM4	778								
18	PM4	735						5 J		
19	PM100	1275								
19	PM100	1288								
19	PM100	1291								
19	PM100	1215								
19	PM10	1270								
19	PM10	1296								
19	PM10	1286								
19	PM10	1218	R	R	R	R	R	R	R	R
19	PM4	796								
19	PM4	819								
19	PM4	822								
19	PM4	761	R	R	R	R	R	R	R	R
20	PM100	1278						3 J		
20	PM100	1226				3 J	16 J	3 J		
20	PM100	1412						3 J		
20	PM100	1417						3 J		
20	PM100	1366	3 J					3 J		
20	PM10	1282						3 J		
20	PM10	1226				3 J	16 J			
20	PM10	1407								
20	PM10	1357	3 J			3 J		4 J		
20	PM4	792						6 J		
20	PM4	753	5 J			5 J	29 J	5 J		
20	PM4	891								
20	PM4	842	5 J			5 J		5 J		
21	PM100	1227								
21	PM100	1238								
21	PM100	1203								
21	PM100	1218								
21	PM10	1248	R	R	R	R	R	R	R	R
21	PM10	1256	R	R	R	R	R	R	R	R

Building Number	Particle Size Fraction	Sample Volume (L)	Quartz (µg/m³)	Cristobalite (μg/m³)	Tridymite (µg/m³)	Calcite µg/m³)	Portlandite (µg/m³)	Gypsum (µg/m³)	Mica µg/m³)	Halite (µg/m³))
21	PM10	1212	R	R	R	R	R	R	R	R
21	PM10	1241	R	R	R	R	R	R	R	R
21	PM4	776								
21	PM4	780								
21	PM4	780								
21	PM4	750								
21	PM4	771						5 J		
21	PM2.5	1236								
21	PM2.5	1249								
21	PM2.5	1211								
21	PM2.5	1238								
22	PM100	1336	5 J							
22	PM100	1385	13 J							
22	PM100	1365	7 J							
22	PM100	1364								
22	PM100	1261								
22	PM10	1346	3 J							
22	PM10	1389								
22	PM10	1361								
22	PM10	1383								
22	PM10	1271								
22	PM4	870								
22	PM4	872								
22	PM4	863								
22	PM4	787								
22	PM2.5	1343								
22	PM2.5	1334								
22	PM2.5	1354								
22	PM2.5	1252								
23	PM100	1546								
23	PM100	1626								
23	PM100	1554								
23	PM100	1550								
23	PM10	1578								
23	PM10	1596								
23	PM10	1568								
23	PM10	1592								
23	PM10	1612								
23	PM4	991								
23	PM4	1012								

Building Number	Particle Size Fraction	Sample Volume (L)	Quartz (µg/m³)	Cristobalite (μg/m³)	Tridymite (µg/m³)	Calcite µg/m³)	Portlandite (µg/m³)	Gypsum (µg/m³)	Mica µg/m³)	Halite (µg/m³))
23	PM4	978								
23	PM4	1014								
23	PM2.5	1548								
23	PM2.5	1009								
23	PM2.5	1544								
23	PM2.5	1584								
24	PM100	1467								
24	PM100	1471								
24	PM100	1424								
24	PM100	1490								
24	PM100	1485								
24	PM10	1475								
24	PM10	1427								
24	PM10	1482								
24	PM10	1481								
24	PM4	920								
24	PM4	891								
24	PM4	880								
24	PM4	929								
24	PM4	920								
24	PM4	935								
	PM4 PM2.5									
24		1467								
24	PM2.5	1411	3 J							
24	PM2.5	1486								
24	PM2.5	1479								
25	PM100	1454						3 J		
25	PM100	1437								
25	PM100	1457								
25	PM100	1451								
25	PM10	1456	R	R	R	R	R	R	R	R
25	PM10	1451	R	R	R	R	R	R	R	R
25	PM10	1453	R	R	R	R	R	R	R	R
25	PM10	1458	R	R	R	R	R	R	R	R
25	PM10	1447	R	R	R	R	R	R	R	R
25	PM10	1454	R	R	R	R	R	R	R	R
25	PM4	913						4 J	N N	
	PM4	909						4 J 		
25 25	PIVH PM4	909 917								
	PM4 PM4									
25	PM4 PM2.5	967 1431						 3 J		

Building Number	Particle Size Fraction	Sample Volume (L)	Quartz (µg/m³)	Cristobalite (μg/m³)	Tridymite (µg/m³)	Calcite µg/m³)	Portlandite (µg/m³)	Gypsum (µg/m³)	Mica µg/m³)	Halite (µg/m³))
25	PM2.5	1414								
25	PM2.5	1433								
25	PM2.5	1444								
26	PM100	1447								
26	PM100	1445	7 J							
26	PM100	1424								
26	PM100	1432								
26	PM10	1453								
26	PM10	1462	R	R	R	R	R	R	R	R
26	PM10	1469								
26	PM10	1445								
26	PM10	1429	R	R	R	R	R	R	R	R
26	PM10	1424	R	R	R	R	R	R	R	R
26	PM4	894								
26	PM4	909								
26	PM4	889								
26	PM4	886								
26	PM2.5	1449								
26	PM2.5	1444								
26	PM2.5	1436								
26	PM2.5	1429								
27	PM100	1473								
27	PM100	1450	3 J							
27	PM100	1469								
27	PM100	1454								
27	PM10	1467								
27	PM10	1457								
27	PM10	1434								
27	PM10	1435								
27	PM4	923								
27	PM4	902								
27	PM4	917								
27	PM4	906								
27	PM4	902								
27	PM2.5	1462								
27	PM2.5	1457								
27	PM2.5	1458								
27	PM2.5	1451								
28	PM100	1427						3 J		

Building Number	Particle Size Fraction	Sample Volume (L)	Quartz (µg/m³)	Cristobalite (µg/m³)	Tridymite (μg/m³)	Calcite µg/m³)	Portlandite (µg/m³)	Gypsum (µg/m³)	Mica µg/m³)	Halite (µg/m³))
28	PM100	1458								
28	PM100	1468	3 J							4 J
28	PM100	1380								
28	PM100	1346								
28	PM100	1337								
28	PM10	1400						3 J		
28	PM10	1450								
28	PM10	1366						3 J		
28	PM10	1343								
28	PM4	898						5 J		
28	PM4	891								
28	PM4	859						5 J		
28	PM4	810								
28	PM2.5	1396								
28	PM2.5	1371								
28	PM2.5	1319								
28	PM2.5	1451								
29	PM100	1519								
29	PM100	1468	4 J							
29	PM100	1447								
29	PM10	1527								
29	PM10	1481								
29	PM10	1447								
29	PM10	1212						3 J		
29	PM4	927	12 J							
29	PM4	897	19 J							
29	PM4	887						5 J		
29	PM4	746						5 J		
29	PM2.5	1451								
30	PM10	1307	3 J					3 J		
30	PM10	1297								
30	PM4	822						5 J		
30	PM4	781								
30	PM2.5	1307								
Results for l	ndividual Com	parison Bu	ildings Sample	d Above 59th Stree	et (Buildings 31	-34)				
31	PM10	1346								
31	PM10	1328						3 J		
31	PM10	1347								
31	PM10	1324								

Building Number	Particle Size Fraction	Sample Volume (L)	Quartz (µg/m³)	Cristobalite (µg/m³)	Tridymite (µg/m³)	Calcite µg/m³)	Portlandite (µg/m³)	Gypsum (µg/m³)	Mica µg/m³)	Halite (µg/m³))
31	PM10	1371								
31	PM10	1401						3 J		
31	PM4	828								
31	PM4	819								
31	PM4	852								
31	PM4	867						5 J		
32	PM100	1318						3 J		
32	PM100	1336								
32	PM100	1351								
32	PM100	1316								
32	PM10	1312						3 J		
32	PM10	1337								
32	PM10	1317								
32	PM10	1312								
32	PM4	808						5 J		
32	PM4	805						5 J		
32	PM4	776								
32	PM4	828								
32		826								
32	PM4 PM2.5							 3 J		
		1311								
32	PM2.5	1418								
32	PM2.5	1315								
33	PM100	1338								
33	PM10	1338								
33	PM4	831								
33	PM4	838								
33	PM2.5	1338								
34	PM100	1307								
34	PM100	1315								
34	PM100	1309								
34	PM100	1232								
34	PM10	1318	R	R	R	R	R	R	R	R
34	PM10	1327								
34	PM10	1238								
34	PM4	816								
34	PM4	848								
34	PM4	830								
34	PM4	782								
34	PM4 PM4	782								

Building Number	Particle Size Fraction	Sample Volume (L)	Quartz (µg/m³)	Cristobalite (µg/m³)	Tridymite (µg/m³)	Calcite µg/m³)	Portlandite (µg/m³)	Gypsum (µg/m³)	Mica µg/m³)	Halite (µg/m³))
34	PM2.5	1308								
34	PM2.5	1348								
34	PM2.5	1223								

--: Not detected. Mineral was not detected above the analytical sensitivity in the associated sample.

Detect: Mineral was detected above the analytical sensitivity for the associated sample.

J: Result presented is an estimate. Due to inconsistencies during sample collection/analysis, the result is only an estimate of the actual value. The actual value could be higher or lower than the value shown.

ND: Not detected. Mineral was not detected above the analytical sensitivity in the associated sample.

NS: Not sampled. The associated method was not used at any of the measured locations in the building. R: Result was rejected. Due to inconsistencies during sample collection/analysis, the result values for the sample were rejected.

 $\mu g/m^3$: microgram per cubic meter of air.