Pilot study of the vertical variations in outdoor pollutant concentrations and environmental conditions along the height of a tall building

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ABSTRACT

It is generally assumed that vertical pollutant dispersion can reduce exposures to ambient pollutants in tall buildings, as concentrations of some ground-source pollutants are diluted at higher floors. However, we are aware of very few measurements of airborne pollutant concentrations that have been made specifically along the height of tall buildings. Therefore, we conducted a pilot study to measure the vertical variation in the concentrations of several outdoor pollutants and environmental parameters along the height of a 60-story (∼300 m) building in downtown Chicago, IL during a one-week period in the summer of 2017. Simultaneous measurements of concentrations of size-resolved particulate matter 0.3–10 μm (which were also used to estimate PM\textsubscript{1}, PM\textsubscript{2.5}, and PM\textsubscript{10} mass concentrations), ozone (O\textsubscript{3}), nitrogen dioxide (NO\textsubscript{2}), carbon dioxide (CO\textsubscript{2}), and carbon monoxide (CO), as well as temperature and relative humidity, were made using multiple sets of instrumentation installed in the outdoor air intakes of the mechanical systems upstream of any filtration or mixing processes on the 2nd, 16th, 29th, and 44th floors and in an open-air area on the 61st floor. The average PM\textsubscript{1}, PM\textsubscript{2.5}, and PM\textsubscript{10} concentrations estimated on the top two floors were more than 30% lower than on the 2nd floor. Temperature, humidity ratio, and CO\textsubscript{2} concentrations decreased with height, O\textsubscript{3} concentrations increased with height, and NO\textsubscript{2} concentrations were less consistent. Most of the differences between floors were statistically significant. Floor height was more strongly correlated with PM\textsubscript{1}, PM\textsubscript{2.5}, PM\textsubscript{10}, CO\textsubscript{2}, and O\textsubscript{3} concentrations than with local wind speed and direction.

1. Introduction

Elevated outdoor concentrations of airborne pollutants such as particulate matter, ozone, and oxides of nitrogen have been consistently associated with increased risks of respiratory symptoms, mortality, and lung cancer [1–7]. Concentrations of many of these pollutants have increased in many urban environments globally in recent years [8–10]. Associations between outdoor pollutant concentrations and adverse health effects are typically made in large epidemiological studies using stationary ambient measurements with inlet heights of ∼2–15 m [11]. However, because outdoor pollutants can infiltrate and persist indoors where Americans spend the majority of their time [12], much of their exposure to pollutants of outdoor origin often occurs inside buildings [13–20]. Indoor exposures to outdoor pollutants are a function of several key factors including outdoor air ventilation rates, envelope pollutant penetration efficiency, HVAC filtration efficiency, indoor pollutant deposition rates, and, importantly, outdoor pollutant concentrations at the source of ventilation air [21]. While previous research has assessed many of these parameters in smaller residential and commercial buildings [22,23], very few measurements have ever been made in tall buildings where inlet heights for outdoor air can be hundreds of meters above ground level.

Most previous studies on vertical pollutant dispersion or the vertical distribution of other environmental parameters in urban street canyons have relied on computational fluid dynamics (CFD) simulations [24–26] or wind tunnel experiments [27–29]. There have been very few field measurements of the vertical dispersion of outdoor pollutants specifically along the height of tall buildings. As an example, one recent study of a mid-rise (i.e., ∼22 stories, or ∼55 m tall) building in Chile showed that outdoor ozone concentrations were found to increase with height [30]. Measured outdoor ozone concentrations were approximately 10–15% higher on the 21st story (53 m above ground level) than on the 3rd story (6 m above ground level). These measurements suggest that occupants of the higher floors in this building may be exposed to...
higher indoor concentrations of outdoor ozone, depending on detailed ventilation system characteristics such as ventilation rates and the location of the outdoor air intakes.

Other limited previous experimental research, while not necessarily sampling in and around tall buildings, has shown that outdoor pollutant concentrations can vary greatly with elevation within the range of height of many tall buildings [31]. Perhaps most relevant to tall (i.e., < 300 m) and super-tall (i.e., > 300 m) buildings [32], aircraft measurements have shown that the vertical variation in outdoor ozone concentrations may be even greater at higher elevations. For example, one study showed that the highest outdoor ozone concentrations during nighttime periods were observed above 200 m (note that 200 m roughly corresponds to ~ 55 stories with typical floor height) [33]. This variation was more scattered during mornings and afternoons, but still suggested an overall similar pattern. These data suggest that occupants on the highest floors of tall or super-tall buildings may be subjected to more than twice the outdoor ozone concentrations than someone in the bottom third floors of the same building, depending on a number of detailed HVAC system characteristics. Vertical variations in outdoor ozone concentrations tend to vary with the height of the atmospheric boundary layer [34], which can vary highly between rural and urban environments and can vary diurnally [35,36]. There is also strong experimental evidence from ambient monitoring that outdoor particulate matter concentrations often decrease with building height [37,38], potentially offering a protective effect at higher floors. However, very few measurements of the vertical variation in outdoor pollutant concentrations exist, particularly along the height of tall buildings in urban environments.

Despite the lack of measurements to date, a study in Switzerland recently suggested that differences in environmental exposures may have contributed to reductions in all-cause mortality that are associated with increased residential floor height in buildings [39]. Similarly, a study of office buildings in the U.S. found significantly higher building-related symptoms reported by occupants working on the floors of buildings that had outdoor air intakes less than 60 m above ground level, which may have been due to greater levels of pollutants from vehicles at air intakes nearer the ground level [40]. The need to better understand pollutant exposures in tall buildings is growing, as there are now a total of over 1300 buildings taller than 200 m in the world, with 144 (11% of the total) being completed in 2017 alone [41]. To begin to fill this knowledge gap, here we report results from a pilot study in which we measured the vertical variation of several outdoor pollutants and environmental parameters along the height of a single tall building in downtown Chicago, IL, USA. The aim is to quantify the dispersion of ambient pollutant concentrations and environmental parameters measured along the height of the test building and to determine the importance of building height and local meteorological factors in influencing the observed variability in the resulting data.

2. Material and methods

A single tall building in Chicago, IL, USA, was recruited for measurements. The building, which will remain unnamed and whose ownership will not be identified, was approximately 60 stories (~ 300 m) tall. Time-resolved measurements were conducted over one weeklong period from June 22, 2017 to June 29, 2017 to monitor concentrations of size-resolved particulate matter (PM), ozone (O3), nitrogen dioxide (NO2), carbon dioxide (CO2), carbon monoxide (CO), and temperature and relative humidity along the height of the tall building. To best represent outdoor air coming into the building, simultaneous measurements were made using multiple sets of instruments placed in the outdoor air intakes on the mechanical systems located on four different floors (i.e., the 2nd, 16th, 29th, and 44th floors), as well as in an open-air area on the 61st floor located underneath a ~2 m high metal rack on which a cooling tower was located. The location of measurements within the outdoor air intakes was upstream of any filtration or mixing processes. Measurements were made within approximately 0.2 m downstream of a coarse metallic grate located on the exterior facade of the building through which outdoor air flowed, and approximately 3 m upstream from adjustable louvers that were located downstream of the exterior grate. The louvers controlled mixing between outdoor air and return air, and were located 2–3 m upstream of a downstream filter bank. A photo of the measurement location in one outdoor air intake is shown in the SI (Fig. S1).

2.1. Instrumentation

The following sections describe the instruments that were used to measure pollutant concentrations and environmental conditions in the five sampling locations along the vertical height of the test building.

2.1.1. Particulate matter (PM)

MetOne GT-526S optical particle counters (OPCs) were used to measure size-resolved optical particle concentrations for particles from 0.3 to 10 + μm in optical diameter in 6 size bins: 0.3–0.5 μm, 0.5–1 μm, 1–2 μm, 2–5 μm, 5–10 μm, and 10+ μm [42]. We primarily used estimates of PM mass concentrations rather than number concentrations for the analyses herein because of their greater, or at least better known, implications for human health. However, we also show the measured size-resolved particle number concentrations in the results section and in the SI.

To estimate integral PM mass concentrations, the mass concentration of particles in each size bin smaller than 10 μm was estimated by assuming spherical particles with diameter equal to the geometric mean diameter of each size bin and uniform density of 1.5 g/cm3 for all particle sizes [43,44]. The mass concentration of PM1, PM2.5, and PM10 was then estimated by adding the mass of particles in the size bins associated with each fraction, as shown in Equations (1)–(3) [45–47]. The assumption for uniform particle density is taken from existing literature sources and may not be accurate for the Chicago area [48–50]. Further, this approach does not account for any mass below 0.3 μm, which will greatly underestimate total number concentrations and may also underestimate PM mass concentrations [51]. However, for the purposes of this study (i.e., to explore the pattern of pollutant concentrations measured along the height of the test building), only repeatable, not absolutely accurate, PM measurements are required on each floor.

\[
C_{PM1} = \left( \frac{1}{6} \pi d^{3}_{0.3-0.5} \rho N_{0.3-0.5} + \frac{1}{6} \pi d^{3}_{0.5-1.0} \rho N_{0.5-1.0} \right) \times 10^{-6}
\]

\[
C_{PM2.5} = \left( \frac{1}{6} \pi d^{3}_{0.3-0.5} \rho N_{0.3-0.5} + \frac{1}{6} \pi d^{3}_{0.5-1.0} \rho N_{0.5-1.0} + \frac{1}{6} \pi d^{3}_{1.0-2.0} \rho N_{1.0-2.0} \right. + \left. \frac{1}{6} \pi d^{3}_{2.0-5.0} \rho N_{2.0-5.0} \times \log \frac{2.5 - \log 2.0}{\log 5.0 - \log 2.0} \right) \times 10^{-6}
\]

\[
C_{PM10} = \left( \frac{1}{6} \pi d^{3}_{0.3-0.5} \rho N_{0.3-0.5} + \frac{1}{6} \pi d^{3}_{0.5-1.0} \rho N_{0.5-1.0} + \frac{1}{6} \pi d^{3}_{1.0-2.0} \rho N_{1.0-2.0} + \frac{1}{6} \pi d^{3}_{2.0-5.0} \rho N_{2.0-5.0} + \frac{1}{6} \pi d^{3}_{5.0-10} \rho N_{5.0-10} \right) \times 10^{-6}
\]

where \( C_{PM1}, C_{PM2.5}, \) and \( C_{PM10} \) are the mass concentrations of PM1, PM2.5, and PM10, respectively (μg/m3); \( d_{j} \) is the geometric mean of particle diameter sizes from i to j (μm); \( \rho \) is the assumed particle density (1.5 g/cm3); and \( N_{i} \) is the particle number concentration measured in size range i (#/m3). To estimate PM2.5 mass concentrations, we used a log-basis differential method to estimate the mass concentration in the 2–2.5 μm size range based on measurements in the 2–5 μm size bin. The number concentration in a virtual size bin of 2–2.5 μm was estimated by multiplying the number concentration of the default 2–5 μm size bin by the ratio of the logarithmic difference of the virtual 2–2.5 μm and actual 2–5 μm size bins (i.e., \((\log 2.5 – \log 2)/\log 5 – \log 2))\).
2.1.2. Gaseous pollutants: \(O_3\), \(NO_3\), \(CO_2\), and \(CO\)

Aeroqual SM50 OEM gas-sensitive semiconductor (GSS) ozone monitors (0–0.5 ppm) were used to measure ozone concentrations with a manufacturer-reported accuracy of ± 15% and a lower limit of detection (LOD) of 1 ppb. These monitors have recently been shown to have accuracy comparable to research and regulatory grade equipment [52], particularly at ambient concentrations well above the reported LOD. Aeroqual S500 monitors with gas-sensitive electrochemical (GSE) NO\(_2\) sensor heads were used to measure NO\(_2\) concentrations with a manufacturer-reported accuracy of ± 20 ppb, a resolution of 1 ppb, and a limit of detection of 5 ppb. These instruments have been used successfully in a few recent studies of indoor and outdoor microenvironments of which we are aware [53–55], and have been shown to be reasonably accurate after calibration with higher-grade equipment.

Lascar EL-USB-CO300 CO monitors were used to measure CO with a manufacturer-reported accuracy of ± 6% of reading and a resolution of 0.5 ppm. These inexpensive instruments are typically most useful when there are large CO sources from combustion (e.g., environmental tobacco smoke or water pipe smoking) [56,57], but other recent residential indoor investigations have used them with success as well [58]. Extech SD8000 dual wavelength non-dispersive infrared (NDIR) CO\(_2\) monitors were used to measure CO\(_2\) concentrations with an accuracy of ± 40 ppm for concentrations less than 1000 ppm or ± 5% of the reading for concentrations greater than 1000 ppm. They have been used in prior successful demonstrations by other field research teams [58]. However, we do not report any of the data from the CO data loggers because the outdoor CO concentration was consistently below the LOD of the CO monitors; thus, the monitors recorded zero values for the vast majority of the measurement intervals.

2.1.3. Environmental conditions

Onset HOBO U12-013 2-Channel Temperature/Relative Humidity (RH) data loggers were used for logging data from the Aeroqual SM50 OEM ozone monitors as well as measuring the temperature and relative humidity of the ambient air. The humidity ratio at each recorded interval was calculated based on the temperature and relative humidity readings following procedures described in the ASHRAE Handbook of Fundamentals [59]. Moreover, we obtained data for wind speed and wind direction from the same time period as the field measurements from a nearby weather station via the Weather Underground Personal Weather Station Network [60]. These data were reported typically at 5-min intervals (albeit with some variation throughout the week), and were summarized as hourly averages for subsequent analyses.

2.1.4. Instrument calibrations

Because multiple versions of each instrument were used to measure each parameter on each of the five floors simultaneously, instrument calibration was conducted using a combination of approaches either before or after the field measurements occurred. At least one of two types of calibrations was performed for each instrument: (1) co-localizations against factory-calibrated research-grade instruments and/or (2) co-localizations against other instruments of the same type, assuming one serves as an arbitrary reference. The former approach allows raw data from each instrument to be adjusted to provide a reasonably accurate measure of the absolute value of the parameter in question. The latter approach allows raw data from each instrument to be adjusted to provide a reasonably accurate measure of the relative value of the parameter compared to an arbitrary reference instrument. Calibration procedures and resulting data are reported in full for each instrument in the SI.

A summary of the calibration results is as follows. The Onset temperature and relative humidity loggers and the Lascar CO loggers were not calibrated based on past experience demonstrating little need to do so. The Aeroqual SM50 \(O_3\) monitors were successfully calibrated using a research grade instrument (2B Technologies Model 211) as a reference, although the instruments were shown to have a fairly high limit of detection (LOD). The highest observable LOD was ∼30 ppb; thus, all data below 30 ppb were excluded from analysis to allow for a fair comparison between instruments on each floor. The MetOne OPCs, Extech CO\(_2\) monitors, and the Aeroqual S500 NO\(_2\) monitors were successfully calibrated using one of the same instruments as an arbitrary reference. The arbitrary reference in all cases was chosen to be the instrument that was deployed on the lowest floor (i.e., the 2nd floor) in the field-testing.

2.2. Field measurement methods

All five sets of instruments described in Section 2.1 were placed in the top drawer in five identical rolling tool carts with uninterruptible power supplies installed in the bottom drawer. The top drawer of each rolling tool cart was modified to include a small exhaust fan on one side and 12 small holes for air intake drilled on the opposite side to continuously draw in sample air flow (Fig. S1). A team of researchers distributed the monitoring instruments to be installed on the 2nd, 16th, 29th, 44th, and 61st floors with the help of the building’s facilities and engineering personnel. In the mechanical rooms, the rolling tool carts were placed as close as possible to the exterior grates on the outdoor air intakes and a box fan was operated continuously to ensure that outdoor air was flowing into the plenum area even if/when the HVAC outdoor air intake happened to shut off for periods of time. Unfortunately we do not have data on the operation of the outdoor air dampers or the operation of HVAC systems. For the 61st floor installation, the rolling tool cart was placed underneath a cooling tower stand that was approximately 2 m tall and located in an otherwise open area that provided for substantial outdoor air flow to the instrument cart.

All of the instruments were synchronized to collect data at approximately the same time. First, we set the internal clock of all instruments to be consistent immediately before the field deployment. The Onset HOBO data loggers recorded the outdoor temperature and relative humidity and data from the SM50 ozone monitors at 1-min intervals using the setting “at interval” to synchronize. Therefore, no additional synchronization was needed for those measurements. The same was true for the CO data loggers, which synchronized with the CPU clock that was used to launch the devices. In order to launch the other monitors simultaneously, individual researchers were deployed to each floor and communicated via two-way radios to manually initiate data logging on each instrument at the same interval at approximately the same time. The result is a set of data that includes synchronized time-stamped data for which each instrument for each measurement type is synchronized to the other instruments with the same measurement type, while all measurement types are synchronized to within approximately 30 s of each other (or closer).

The monitors were then left to record data for approximately one week. The SM50 ozone monitors, the SD800 CO\(_2\) monitors, and the HOBO T/RH data loggers successfully collected data for the entire period while synchronized at 1-min intervals, while the GT-526S OPcs successfully collected data for one week at 2-min intervals (there was not enough storage space on the OPCs to collect data for the entire week at 1-min intervals). The S500 NO\(_2\) monitors recorded data at 1-min intervals for only the last ∼5.5 days of the weeklong measurements because their internal memory cards were filled and the earlier data points were automatically overwritten.

2.3. Data analysis

Upon data collection, calibration factors were applied to the raw data collected from each instrument following procedures described in the SI. Reported measurements are not corrected for air temperature or density, as volume corrections for air density are estimated to be less than 3% at maximum along the ∼300 m height of the test building. We explored the resulting data set in the following ways: (1) time-series plots; (2) box plots and summary statistics for each floor and building.
3. Results and discussion

In each subsection below, floor-by-floor comparisons of synchronized parameter measurements are shown using box plots that show the interquartile range (IQR) as the 25th and 75th percentile values (boxes), median values (line within the boxes), the upper and lower adjacent values (i.e., the upper or lower quartile plus/minus 1.5 times the IQR), and any outside values as dots (unless excluded for clarity) (i.e., Figs. 1–4). Time-series data from the same measurements are also shown in the SI, and average (± standard deviation) values for all parameters are plotted versus approximate building height in Fig. 5. Table S1 provides a full statistical summary of the calibrated results and also calculates the relative differences in the arithmetic means of each measured parameter from each floor compared to the 2nd floor as the closest-to-ground-level reference. Results from statistical significance testing of simultaneous measurements on each floor are shown in Table 1.

3.1. Temperature and humidity ratio

Fig. 1 shows box plots of the temperature and humidity ratio data measured on the five floors. Full weeklong time-series data are shown in Figs. S18 and S19. Humidity ratio is shown instead of relative humidity because it is an absolute measure of humidity and it is not a function of temperature. The average temperature across all floors ranged from \(\sim 21.5^\circ\text{C}\) to \(\sim 24.0^\circ\text{C}\) throughout the week, with minimum and maximum values ranging from \(\sim 14^\circ\text{C}\) to \(\sim 32^\circ\text{C}\). The average temperature was \(\sim 2.8\%\) higher on the 16th floor compared to the 2nd floor, but was \(\sim 1.7\%\) (i.e., \(\sim 0.4^\circ\text{C}\)), \(\sim 2.3\%\) (i.e., \(\sim 0.5^\circ\text{C}\)), and \(\sim 7.6\%\) (i.e., \(\sim 1.7^\circ\text{C}\)) lower on the 29th, 44th, and 61st floors compared to the 2nd floor, respectively. All measured temperature differences between floors were statistically significant (Table 1).

The average temperature difference of \(\sim 1.7^\circ\text{C}\) between the 61st floor (height of \(\sim 300\text{ m}\)) and the 2nd floor (height of \(\sim 5\text{ m}\)) yields an average temperature lapse rate of about \(\sim 0.58^\circ\text{C}\) per 100 m along the height of the building, which is within \(\sim 10\%\) of the commonly used Standard Lapse Rate of \(\sim 6.5^\circ\text{C}\) per 1000 m (i.e., \(\sim 0.65^\circ\text{C}\) per 100 m) \([61,62]\). However, the temperature lapse was not constant across each floor comparison, which suggests that the temperature lapse rate assumption for a building of this size in this urban context may not be linear and may be influenced by other factors such as surrounding buildings or highly localized meteorological conditions \([63]\).

The average relative humidity across all floors ranged from \(\sim 45\%\) to \(\sim 71\%\) throughout the week, with minimum and maximum values ranging from \(\sim 26\%\) to \(\sim 77\%\). The average relative humidity was \(\sim 9.1\%\), \(\sim 5.5\%\), and \(\sim 4.8\%\) lower on the 16th, 29th, and 44th floors compared to the 2nd floor, respectively, but was \(\sim 4.8\%\) higher on the 61st floor compared to the 2nd floor. Differences in RH were not as consistent as differences in humidity ratio because of the dependence of RH on temperature. The average humidity ratio across all floors was \(\sim 0.0084 \text{ kg}_{\text{H}_2\text{O}}/\text{kg}_{\text{dry}}\) throughout the week, with minimum and maximum values ranging from \(\sim 0.005 \text{ kg}_{\text{H}_2\text{O}}/\text{kg}_{\text{dry}}\) to \(\sim 0.016 \text{ kg}_{\text{H}_2\text{O}}/\text{kg}_{\text{dry}}\). The average absolute humidity ratio was \(\sim 5.2\%\), \(\sim 7.9\%\), \(\sim 8.0\%\), and \(\sim 5.1\%\) lower on the 16th, 29th, 44th, and 61st floors compared to the 2nd floor, respectively. There was no clear linear trend observed between humidity ratio and building height, but humidity ratio was lower on all floors above ground level. All differences in relative and absolute humidity measured between floors were statistically significant except the comparison of humidity ratio between the 16th and 61st floors (Table 1).

3.2. Particulate matter (PM)

3.2.1. Size-resolved particle number concentrations

Fig. 2 shows box plots of size-resolved particle number concentrations measured on each floor. Outside values (i.e., those above/below the upper/lower adjacent values) are not shown for visual clarity. Fig. S20 shows full weeklong time-series particle number concentration data. Raw number concentrations were adjusted to 2nd floor monitor equivalent values as an arbitrary reference following procedures.
described in the SI. Number concentrations for each particle size bin measured on the 16th floor and above were consistently lower than those measured on the 2nd floor, with reasonably similar trends for most particle size bins (albeit with some deviations, particularly in the 61st floor measurements). The lowest number concentrations for most particle sizes along the height of the building were observed on the 44th floor, as median concentrations in the 0.5–1 μm and 5–10 μm size ranges were both ∼51% lower than the median concentrations on the 2nd floor. The smallest decreases in number concentrations along the height of the building were observed for particles in the 2–5 μm size range, with median decreases between measurements on the 16th floor and above and the 2nd floor ranging from ∼15% to ∼21%.

3.2.2. Estimates of PM1, PM2.5, and PM10 mass concentrations

Fig. 3 shows box plots of estimates of PM1, PM2.5, and PM10 mass concentrations made using number concentrations measured on each of the five floors, with outliers excluded for clarity. Figs. S21, S22, and S23 show full weeklong time-series data. Again, raw concentrations were adjusted to 2nd floor monitor equivalent values as an arbitrary reference. Estimates of PM mass concentrations were strongly correlated with number concentrations from the closest corresponding size bins (i.e., Spearman rank correlation coefficients were greater than 0.9 between PM1 and the 0.3–1 μm bins; greater than 0.8 between PM2.5 and the 0.3–2 μm bins; and greater than 0.85 between PM10 and the 2–10 μm bins).

The median (and mean) PM1, PM2.5, and PM10 concentrations estimated from number measurements in the 2nd floor outdoor air intake as a near-ground reference were ∼1.3 (∼1.5) μg/m3, ∼2.2 (∼2.3) μg/m3, and ∼9.3 (∼10.6) μg/m3, respectively, throughout the week, which are surprisingly low for an urban environment such as Chicago. However, the average daily PM2.5 concentration measured at the nearest ambient regulatory monitor (#17-031-0057, 1745 N Springfield Ave, Chicago, IL, ∼9 km away) was only 2.8 μg/m3 during the measurement campaign [64]. For comparison, the average daily PM2.5 concentration for the year 2017 measured at the same regulatory monitor was ∼8.6 μg/m3. Although this presents only a limited...
comparison, it demonstrates that the field campaign happened to occur during a period of relatively low ambient PM concentrations.

The average PM$_1$ concentration was estimated to be $\sim$18.4%, $\sim$24.8%, $\sim$34.5%, and $\sim$23.7% lower on the 16th, 29th, 44th, and 61st floors compared to the 2nd floor, respectively, suggesting a fairly consistent trend of PM$_1$ concentrations decreasing with building height. Similarly, the average PM$_{2.5}$ concentration was estimated to be $\sim$10.4%, $\sim$18.0%, $\sim$30.3%, and $\sim$31.7% lower on the 16th, 29th, 44th, and 61st floors compared to the 2nd floor, respectively. All PM$_1$ and PM$_{2.5}$ comparisons were statistically significant except for the comparison between PM$_{2.5}$ concentrations measured on the 44th and 61st floors. The trend for both PM$_1$ and PM$_{2.5}$ was nearly linear from floors 2 through 44, with a deviation in the open-air 61st floor location. The PM$_1$ and PM$_{2.5}$ concentration dispersion data are reasonably consistent with prior ambient measurements [37,38].

The average PM$_{10}$ concentration was estimated to be $\sim$12.9%, $\sim$32.4%, and $\sim$31.5% lower on the 29th, 44th, and 61st floors compared to the 2nd floor, respectively, but actually $\sim$15.8% higher on the 16th floor compared to the 2nd floor (although the median PM$_{10}$ concentration on the 16th floor is still lower than the 2nd floor as shown in Fig. 3c). All PM$_{10}$ comparisons were statistically significant (Table 1). This inconsistent trend at the lower levels may be suggestive of local ground sources with greater dilution occurring at higher elevations. Interestingly, the standard deviation of PM$_{10}$ concentrations was largest

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**Fig. 4.** Box plots of (a) ozone (O$_3$), (b) nitrogen dioxide (NO$_2$), and (c) carbon dioxide (CO$_2$) concentration data measured on each of the five floors.

Fig. 5. Average (± standard deviation) of the CO$_2$, O$_3$, NO$_2$, PM$_1$, PM$_{2.5}$, and PM$_{10}$ concentrations and the air temperature and humidity ratios measured (or estimated) during the weeklong field campaign plotted against the approximate floor height of the test building.
on the 16th floor, which means that there were periodically very high PM10 concentrations measured on the 16th floor and suggests an influence from nearby transient PM10 sources around this height.

### 3.3. Gaseous pollutants: O3, NO2, and CO2

Fig. 4 shows box plots of the O3, NO2, and CO2 concentrations measured on the five floors and Figs. S24, S25, and S26 show the full weeklong time-series data. For O3, each data point was adjusted to the federal equivalent method (FEM) reference instrument (2B Technologies Model 211) following calibration procedures described in the SI. Only data above the highest measured LOD for the SM50 instruments (∼30 ppb) are shown, as varying LODs make it impossible to compare null values with actual values recorded at concentrations lower than ∼30 ppb. For NO2 and CO2, each data point was adjusted to 2nd floor monitor equivalent values as an arbitrary reference. All observed differences in O3, NO2, and CO2 concentrations between floors were statistically significant (Table 1).

### Table 1

Significance testing on paired data sets of each parameter compared between each floor using nonparametric Wilcoxon signed-rank tests.

<table>
<thead>
<tr>
<th>Sample size, n</th>
<th>Temp.</th>
<th>RH</th>
<th>Humidity Ratio</th>
<th>CO2</th>
<th>O3</th>
<th>PM1</th>
<th>PM2.5</th>
<th>PM10</th>
<th>NO2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>9928</td>
<td>9928</td>
<td>9928</td>
<td>10089</td>
<td>9901</td>
<td>4959</td>
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</tr>
<tr>
<td>Adj. p-value threshold</td>
<td>5.17 × 10⁻⁶</td>
<td>5.17 × 10⁻⁶</td>
<td>5.17 × 10⁻⁶</td>
<td>5.08 × 10⁻⁶</td>
<td>5.18 × 10⁻⁶</td>
<td>1.03 × 10⁻⁵</td>
<td>1.03 × 10⁻⁵</td>
<td>1.03 × 10⁻⁵</td>
<td>6.31 × 10⁻⁴</td>
</tr>
</tbody>
</table>

**p < 10⁻²⁵⁰**
**10⁻²⁵⁰ < p < 10⁻¹⁰⁰**
* 10⁻¹⁰⁰ < p < threshold.

n.s. = not significant.

* Comparisons are made using 1-min or 2-min interval data as dictated by each instrument. Only those cells with bold text failed to reach statistical significance based on a threshold p-value that is adjusted for sample size.

The average O3 concentration across all floors (when limiting to data above an LOD of ∼30 ppb) was ~48 ppb throughout the week, with a maximum of ~122 ppb. The measured O3 concentrations were similar to the average daily O3 concentration of ~46 ppb measured at the three nearest ambient regulatory monitors (#17-031-0003 6545 W Hurlbut St; #17-031-0032, 3300 E Cheltenham Pl; and #17-031-0076, 7801 Lawndale; each ~15–20 km away) during the field campaign [64]. Further, historical records of O3 concentrations measured by these same regulatory monitors suggest that hourly O3 concentrations are typically below 30 ppb during periods in late June approximately 20–30% of the time [64]; a similar fraction of data points were excluded for being below the 30 ppb LOD on most floors in our data set as well.

The average O3 concentration above this LOD was ~11.9% and ~11.3% lower on the 16th and 29th floors compared to the 2nd floor, respectively, but ~16.0% and ~18.0% higher on the 44th and 61st floors compared to the 2nd floor, respectively. This inconsistent vertical trend in O3 concentrations is not unlike the limited data from aircraft measurements reported in the literature reviewed in Section 1, in which concentrations first decrease and then increase with elevation [32]. This may be due in part to titration of O3 by NO from ground-level tailpipe emission sources, which might not reach the higher elevations or might be diluted and/or reacted by the time air masses reached higher elevations. We should also note that we have somewhat lower confidence in these ozone concentration measurements based on the potential for calibration factors to vary with temperature, relative humidity, and interference by other compounds that have not been captured in our calibration procedures or field measurements [52].

The average NO2 concentration across all floors (when reporting data using the 2nd floor monitor as an arbitrary reference) was ~50 ppb throughout the week, with a maximum of ~129 ppb encountered on the 29th floor. Conversely, the average daily NO2 concentration measured by the nearest regulatory monitor (#17-031-0063, 321 S Franklin, Chicago, IL) was only 21 ppb. Because of the close proximity of the test building to the nearest NO2 regulatory monitor, reasons for this discrepancy are most likely attributed to artifacts in the monitoring devices and issues that prevented a true calibration against research grade equipment. For example, Section 1.2.4 in the SI (including Fig. S15) clearly shows a positive offset of ~30–50 ppb for the Aeroqual S500 monitors compared to the 28 Technologies Model 405 monitor that unfortunately could not be systematically accounted for in data processing. Despite this offset issue, data from the S500 NO2 monitors still yield meaningful comparisons on a relative basis (i.e., when calibrated against each other). Using these data, the average NO2 concentration was ~25.3% lower on the 16th floor, ~47.0% higher on the 29th floor, ~15.1% lower on the 44th floor, and ~5.3% lower on the 61st floor, each compared to the 2nd floor. Reasons for this inconsistent trend are not clear, but it is worth noting that, similar to the O3 measurements, we also have somewhat lower confidence in these NO2 concentration measurements based on the potential for calibration factors to vary with temperature, relative humidity, and interference by other compounds that have not been captured in our calibration procedures or field measurements [52].

Finally, the average CO2 concentration across all floors was ~387 ppm throughout the week, with a maximum of ~700 ppm observed on the 2nd floor. The average CO2 concentration was ~7.6%, ~1.5%, ~4.9%, and ~6.9% lower on the 16th, 29th, 44th, and 61st floors compared to the 2nd floor, respectively. These relative differences correspond to average absolute differences of ~30 ppm, ~6 ppm, ~20 ppm, and ~28 ppm, respectively. There was no consistent linear trend in average CO2 concentrations across all elevations, although once again, concentrations were consistently lower on all floors above the 2nd floor, suggesting dilution or dispersion of ground-level sources at higher floors.

### 3.4. Diurnal patterns between floors

Fig. 6 shows box plots (excluding outliers) of the diurnal variations...
in relative differences in PM$_{1}$, PM$_{2.5}$, PM$_{10}$, CO$_2$, O$_3$, and NO$_2$ concentrations measured on the 16th, 29th, 44th, and 61st floors compared to the 2nd floor. Relative differences are calculated as the simultaneous nth floor values minus the simultaneous 2nd floor values, divided by the simultaneous 2nd floor values. Values are reported for each of the 24 h of each day, aggregated across all days of the measurement campaign. The maximum differences in estimates of PM$_{1}$ and PM$_{2.5}$ concentrations tended to occur in the morning and midday from $\sim$7 a.m. to $\sim$2 p.m., while smaller and relatively consistent differences were observed during other time periods. Some large peaks observed around 5 a.m.–8 a.m. drove most of the differences in PM$_{10}$ concentrations, particularly on the 16th floor. A few other transient peaks in estimates of PM$_{10}$ concentrations were also seen at other hours on the 29th and 44th floors, suggesting some highly localized sources or perhaps even resuspension of settled dust by the presence of maintenance personnel within the outdoor intake areas (although this could not be confirmed).

The maximum differences in CO$_2$ concentrations were observed in the late afternoons from $\sim$2 p.m. to $\sim$5 p.m., suggesting likely contributions from vehicle sources as traffic starts to increase in the area. There were no clear diurnal patterns in the differences in observed O$_3$ and NO$_2$ concentrations other than a mild increase in differences in O$_3$ concentrations at higher elevations in the late afternoons. Combined, these data generally suggest that variations along the building’s height are reasonably consistent throughout the day, albeit with some periodic pollutant-specific deviations.

Similarly, Fig. 7 shows box plots (excluding outliers) of the diurnal variations in temperature and relative humidity measured on the 16th, 29th, 44th, and 61st floors compared to the 2nd floor. The largest differences in temperature and, to a greater extent, humidity ratio, tended to occur in the late afternoons and early evenings (e.g., $\sim$2–$\sim$5 p.m.), with some periodically large deviations among certain floors at other times of day. For example, there were
large increases in temperatures on the 44th floor in the early morning hours (≈6–8 a.m.) and large decreases in humidity ratios on the 29th floor at night (≈8–10 p.m.), likely caused by temporary anthropogenic sources, perhaps including operation of HVAC systems in nearby buildings, changes in vehicle traffic patterns [65], or other common street canyon effects [24–26].

### 3.5. Potential drivers of variations in the measured data

The obtained wind speed and wind direction data from the Weather Underground Personal Weather Station Network [44] were combined with building height to investigate the potential drivers of the observed variations in measured (or estimated) parameters. The average (± standard deviation) hourly wind speed and wind direction near the measurement site were 2.5 (±1.3) m/s and 193° (±47°) (i.e., from the south-southwest), respectively (Fig. 8). The most prevalent wind direction was from the southwest (supporting the hypothesized effects [24–26]). The comparison between NO2 concentrations and building height was not significant; however, wind direction was positively correlated with measured NO2 concentrations, with building height to investigate the potential drivers of the observed variations in measured (or estimated) parameters. The average (± standard deviation) hourly wind speed and wind direction near the measurement site were 2.5 (±1.3) m/s and 193° (±47°) (i.e., from the south-southwest), respectively (Fig. 8). The most prevalent wind direction was from the southwest (supporting the hypothesized effects [24–26]).

Spearman rank correlation coefficients were calculated using hourly averages of each measured parameter as the dependent variable and building height, hourly average wind speed, and hourly average wind direction as independent variables (Table 2). Results show that the variable that was most strongly correlated with most of the measured (or estimated) pollutant concentrations was floor height, with the highest Spearman rank correlation coefficients calculated for hourly average PM10, PM2.5, PM1, O3, and CO2 concentrations in particular. Spearman rank correlation coefficients were negative for all of these pollutants, suggesting a decreasing trend in concentration with building height. Moreover, each of these comparisons with building height was statistically significant (p < 0.0001), but relatively weak (i.e., Spearman rank correlation coefficients with building height ranging from –0.18 to –0.36). The comparison between NO2 concentrations and building height was not significant; however, wind direction was positively correlated with measured NO2 concentrations, which suggests NO2 concentrations were higher when the prevailing wind direction was from the southwest (supporting the hypothesized transport of vehicular NO2 emissions).

Building height was also significantly correlated with temperature and humidity ratio, but wind direction was more strongly correlated with both parameters. Wind speed showed the strongest association with temperature, but was weakest for humidity ratio. These data demonstrate that the majority of floor-by-floor comparisons shown...
Corporation for their generous acknowledgment at scales that are relevant to occupants of these building super-tall buildings in other climate zones and geographic regions to ventilation and particle dispersion and local environmental conditions in urban environments within the context of tall buildings. These pilot data also suggest the following implications for the design and operation of tall buildings: (1) the dry bulb temperature lapse rate of a building can deviate from the linear Standard Lapse Rate assumption during some periods, which may need to be accounted for in HVAC design and energy simulation; (2) concentrations of some ambient pollutants or constituents, especially measurements of PM number concentrations and estimates of PM mass concentrations, and, to a lesser extent, CO2, showed strong signatures of ground-level emissions that become dispersed or diluted at higher floors, which may need to be accounted for in designing and operating ventilation and particle filtration systems; and (3) concentrations of O3 were clearly highest at the lowest elevations of the building, which may also need to be considered in the design and operation of ventilation and gas-phase filtration systems.

4. Conclusions

To our knowledge, this data set represents the first known measurements of the vertical variation in concentrations of several health relevant outdoor pollutants and climatically relevant outdoor environmental parameters in outdoor air intakes along the height of a tall building in a major metropolitan area. In general, the arithmetic mean values of most measured parameters tended to decrease with building height, albeit with some exceptions. The magnitude of measured differences among floors was statistically significant but typically small for most parameters (i.e., less than 10% for temperature, relative humidity, and CO2) but large for others (i.e., up to a maximum decrease of ~32%, with averages consistently decreasing with floor height, for estimates of PM1 and PM2.5 concentrations). Variations in other parameters such as PM10, O3, and NO2 concentrations were less consistent and varied in magnitude. Given some of the relatively large magnitudes of differences in measured values observed herein, we recommend that additional measurements be made in other tall and super-tall buildings in other climate zones and geographic regions to better understand how and why pollutant concentrations vary with elevation at scales that are relevant to occupants of these building types.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.buildenv.2018.04.031.

References


