REAL-TIME MEASUREMENT OF OUTDOOR TOBACCO SMOKE PARTICLES

NEIL E. KLEPEIS,* WAYNE R. OTT, AND PAUL SWITZER
Stanford University, Stanford, CA 94305 USA

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ABSTRACT

The current lack of empirical data on outdoor tobacco smoke (OTS) levels impedes OTS exposure and risk assessments. We sought to measure peak and time-averaged OTS concentrations in common outdoor settings near smokers and to explore the determinants of time-varying OTS levels, including the effects of source proximity and wind. Using 5 types of real-time airborne particle monitoring devices, we obtained more than 8,000 min worth of continuous monitoring data, during which there were measurable OTS levels. Measurement intervals ranged from 2 sec to 1 min for the different instruments. We monitored OTS levels during 15 visits to 10 outdoor public places where active cigar and cigarette smokers were present, including parks, sidewalk cafés, and restaurant and pub patios. For 3 of the visits, and during 4 additional days of monitoring outdoors and indoors at a private residence, we controlled smoking activity at precise distances from monitored positions. The overall average OTS respirable particle concentration for the visits to public places during smoking was about 30 µg m⁻³. OTS exhibited sharp spikes in particle mass concentration during smoking that sometimes exceeded 1,000 µg m⁻³ at distances within 0.5 m of the source. Some average concentrations over the duration of a cigarette and within 0.5 m exceeded 200 µg m⁻³, with some average downwind levels exceeding 500 µg m⁻³. OTS levels in a constant upwind direction from an active cigarette source were nearly zero. OTS levels also approached zero at distances greater than about 2 m from a single cigarette. During periods of active smoking, peak and average OTS levels near smokers rivaled indoor tobacco smoke concentrations. However, OTS levels dropped almost instantly after smoking activity ceased. Based on our results, it is possible for OTS to present a nuisance or hazard under certain conditions of wind and smoker proximity.

IMPLICATIONS

This article is the first peer-reviewed publication of systematic measurements of outdoor tobacco smoke (OTS) concentrations.

*Corresponding author. Please visit http://klepeis.net

INTRODUCTION

Secondhand tobacco smoke (SHS), also called environmental tobacco smoke (ETS) or passive smoke, is defined as diluted and dispersed air pollutant emissions generated from the consumption of tobacco products. Emissions may be exhaled by a smoker (mainstream) or leave the burning tip of a cigarette or cigar (sidestream). When occurring outdoors, SHS is called outdoor tobacco smoke (OTS).

Indoor SHS has an established connection to adverse health outcomes in adults and children, such as asthma, respiratory infection, and lung cancer. More recent work has shown an association between SHS exposure and reduced cognitive ability in children, increased respiratory disease in adults from work exposure and increased cancer for people exposed at home as children, increased coronary heart disease in women exposed at home or work, and a general increase in mortality for persons living with smokers. The US Surgeon General’s Report entitled "The Health Consequences of Involuntary Exposure to Tobacco Smoke" concludes that there is no level of exposure to SHS without some associated risk, and the California Air Resources Board recently designated SHS as a "toxic agent", a classification also given to pure compounds such as arsenic or benzene.

The body of evidence demonstrates clear harm from SHS exposure and supports the pursuit of exposure reduction policies. In 1995, California Assembly Bill 13 was passed, which effectively banned smoking inside eating and drinking establishments throughout California. Other state or country-wide initiatives that ban smoking inside bars and restaurants also have
have just started to institute bans on outdoor smoking, such as for parks and beaches. Bans may be supportable due to the drift of OTS inside buildings or from the littering of cigarette butts. Outdoor smoking bans may also serve to discourage smoking behavior in general, by making it more difficult for smokers to find a place to light up or by preventing children from associating smoking with enjoyable outdoor activities. However, the ongoing debate over the appropriateness of outdoor bans from an exposure standpoint suffers from a lack of air monitoring data. To date, no data have been published in the archival literature on the systematic measurement of human exposure to OTS. To meet this need, we performed OTS monitoring surveys and controlled OTS experiments in public outdoor locations and a private residential patio using state-of-the-art, real-time particle sensing instruments. These instruments were anticipated to be useful for pinpointing and understanding transient elevations in OTS pollution. We expect that the results of our study will be helpful to those involved in tobacco-related policy development, as well as to risk assessors and environmental epidemiologists.

METHODS

While there are many potentially toxic compounds in both the gaseous and particle phases of secondhand tobacco smoke (SHS), for the present work we used airborne particle concentrations to characterize SHS levels. The use of particles to indicate the presence of SHS is common practice. Airborne particles comprise a significant portion of the sidestream and mainstream mass emissions from burning cigarettes and other tobacco products, and indoor particle concentrations associated with SHS are substantial. The size range of SHS particles is roughly 0.02 to 2 μm, so that all SHS particles fall within the PM2.5 and respirable suspended particle (RSP; also called PM1.5) size ranges, which consist of particles with diameters under 2.5 and 3.5 μm, respectively. When inhaled, these particles can deposit in the human lung. Other benefits of using particles to characterize SHS are that particle concentrations can be measured using standard techniques, particles have a direct association with adverse health effects, and there are existing health standards for time-averaged particle concentrations.

Since many types of portable continuous monitors for airborne particles are currently available, we decided for the present study to employ a range of different instruments to characterize dynamic outdoor tobacco smoke (OTS) levels in the field and under controlled conditions. The simultaneous use of multiple monitors of the same type and of different types allowed us to achieve a high level of confidence in measured OTS levels, and to perform intensive evaluations and comparisons of the instruments.

We used 5 types of portable real-time airborne particle monitoring instruments to measure OTS concentrations at intervals ranging from 2 sec to 1 min. The monitor types included a piezoelectric microbalance (piezobalance; PZB), a photoelectric aerosol sensor (PAS), and three light-scattering photometers—an integrating nephelometer (NEPH), a laser particle counter (GRIMM), and a laser diode photometer (SIDEPAK). A brief summary of the characteristics of each real-time particle monitoring instrument, along with references to the scientific literature or manufacturers’ guides, is given in Table 1. We selected each instrument because of its sensitivity to tobacco smoke particles, rapid response time, portability, and/or proven reliability in the field. In addition to these instruments, we used a real-time hot wire anemometer to record air flow (0.01 m sec−1 threshold), temperature, and relative humidity every minute (VelociCalc Model 8386, TSI, St. Paul, MN).

The piezobalance (PZB) was designated as the reference particle mass monitor, because it provides direct measurements of respirable suspended particle (RSP) mass concentrations, and it has a long history of use with tobacco smoke. The PZB has been shown to agree well with reference pump- and filter-based RSP measurements. Ott et al. provide a review of prior studies that evaluated the PZB, including one by Sem et al., who report piezobalance mass readings for tobacco smoke to be within 15% of filter-based samples. Based on 9 recent experiments we performed in a 9 m³ chamber using cigarettes and incense as sources, we found that average mass readings of an impactor-equipped PZB were within about 10% of average mass concentrations determined from cyclone mass filter samples ($R^2 = 96\%$).

INSTRUMENT TESTING AND CALIBRATION

The NEPH, SIDEPAK, GRIMM, and PAS continuous monitors can be used to estimate RSP mass concentrations. However, it is essential to first calibrate them with respect to the specific aerosol under study. We tested, calibrated, and compared the monitoring instruments for a tobacco smoke source during a set of 14 side-by-side experiments in a 44 m³ room of a residence. For each experiment, a single cigarette was lit and allowed to burn by itself (smolder-smoke) for 4--10 min. Doors and windows were kept closed, except to clear smoke from the room in between experiments. The room SHS particle concentrations were measured during and after each cigarette burn period. We subtracted background particle levels, which were observed just before smoking began, from all readings. Over the range of relative humidities, which we measured during the experiments (40--70%), we found no influence of relative humidity on measured particle levels.

We calculated one conversion factor for each of 12 experiments where valid PZB readings were available (see Table 2) by taking the ratio of the fresh 5-min average for the PAS,
**TABLE (1): Characteristics of Five Real-Time Airborne Particle Monitors Used in the Present Work**

<table>
<thead>
<tr>
<th>Monitor Type</th>
<th>Abbreviation</th>
<th>Description</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Piezobalance</td>
<td>PZB</td>
<td>The model 3511 (Kanomax, Inc., New York, NY) and model 8510 (TSI, Inc., St. Paul, MN) piezobalances measure respirable suspended particle (RSP) mass in units of $\mu$g m$^{-3}$ by passing an air stream though a 3.5 $\mu$m size-selective particle impactor and onto a vibrating piezoelectric crystal. The frequency change in the crystal is converted to an average particle concentration with a resolution of about 10 $\mu$g m$^{-3}$, which we record automatically in 1-min intervals using a custom-built logging system.</td>
<td>Ott et al.\textsuperscript{16} and Sem et al.\textsuperscript{17}</td>
</tr>
<tr>
<td>Nephelometer</td>
<td>NEPH</td>
<td>The model M903 integrating nephelometer (Radiance, Inc., Seattle, WA) uses a flash lamp and optical filter to measure a light scattering coefficient (&quot;extinction coefficient&quot;) for particles drawn into the instrument at intervals as small as 2 sec. The instrument does not include a size-selective inlet.</td>
<td>Brauer et al.\textsuperscript{18} and Radiance Research\textsuperscript{19}</td>
</tr>
<tr>
<td>Laser photometer</td>
<td>SIDEPAK</td>
<td>The model AM510 Sidepak (TSI, Shoreview, MN) is a 90-degree light scattering system using a 670 nm laser diode that is pre-calibrated by the manufacturer using Arizona road dust to measure aerosol mass in units of mg m$^{-3}$. In the present work, we equipped the monitor with a 2.5 $\mu$m impactor and used the internal logger to record levels at intervals as small as 10 sec. Prior to each monitoring visit, the SIDEPAK inlet flow rate was adjusted to 1.7 L min$^{-1}$ using a Gilibrator primary flow calibrator.</td>
<td>TSI\textsuperscript{20}</td>
</tr>
<tr>
<td>Laser counter</td>
<td>GRIMM</td>
<td>The model 1.108 laser counter (GRIMM, Inc., Douglasville, GA) internally records counts of airborne particles every minute in 14 size ranges from 0.3–20+ $\mu$m with a resolution of 1 particle count per liter. It measures light photons from a semi-conductor laser that have been scattered at an angle of approximately 90 degrees.</td>
<td>Grimm Technologies\textsuperscript{21}</td>
</tr>
<tr>
<td>Photoelectric Aerosol Sensor</td>
<td>PAS</td>
<td>The model PAS 2000CE photoelectric aerosol sensor (EcoChem, Inc., League City, TX) takes advantage of the physics of polycyclic aromatic hydrocarbon (PAH) photoemission on the surface of particles. It uses ultraviolet light to ionize PAH on particles under 1 $\mu$m in diameter and measures the resulting electrical charges. The instrument is pre-calibrated to internally record the mass concentration of PAH in units of ng m$^{-3}$ at intervals as small as 30 s. Because it measures particle-bound PAH, the PAS instrument may respond differently to aerosols that have comparable total mass concentrations but vary in their surface PAH content.</td>
<td>Ott and Siegmann\textsuperscript{22} and EcoChem Analytics\textsuperscript{23}</td>
</tr>
</tbody>
</table>
SIDEPAK, GRIMM, and NEPH readings to the fresh PZB 5-min average RSP mass levels measured during a period starting 5–10 minutes after smoking stopped (at which time concentrations were evenly mixed in the room). Background levels were subtracted prior to taking the ratio. Fresh levels were used to determine conversion factors, because outdoor tobacco smoke (OTS) was expected to consist exclusively of fresh emissions. Except for the GRIMM monitor, we used the raw readings of each instrument to determine the conversion factors. In the case of the GRIMM, we used the sum of all particles from the lowest measured diameter of 0.3 μm up to 3.0 μm, since tobacco smoke particles are expected to be in the 0.02–2 μm range. While linear regressions between 1-min average PZB readings and the other instruments across all well-mixed concentrations (fresh and aged) showed generally good agreement on a per-experiment basis ($R^2 = 80–99\%$), there was evidence of a nonlinear relationship in many cases. Our use of ratios of background-subtracted 5-min average particle concentrations to calculate conversion factors, rather than linear regressions, resulted in lower relative variation for conversion factors – likely because it minimized biases due to deposition, coagulation, or evaporation of tobacco smoke particles occurring over time.

The average conversion factors from NEPH and SIDEPAK native units to RSP units observed in the present study ($4.6 \text{ m}^2 \text{ g}^{-1} \text{ and } 3.3 \times 10^{-3} \text{ mg} \mu \text{g}^{-1}$, respectively) are similar to those determined by other investigators. For example, Brauer et al. found a value of $4.7 \text{ m}^2 \text{ g}^{-1}$ for the NEPH conversion to mass for cigarettes, and both Travers and Lee found values of approximately $3 \times 10^{-3} \text{ mg} \mu \text{g}^{-1}$ for the SIDEPAK conversion. Lee performed 14 laboratory calibration tests of the SIDEPAK using gravimetric PM$_{2.5}$ filter samples and a smoking machine. Previous investigators have also found good agreement between personal nephelometers (e.g., the MIE personal DataRam) and reference gravimetric methods when calibrated for the target aerosols and adjusted properly for high relative humidity. Personal nephelometers, which have been used by the USEPA and others to characterize particle exposures, operate on principles similar to the SIDEPAK and NEPH light-scattering photometers used in the current study. Our SIDEPAK conversion factor corresponds to an internal “custom calibration factor” of approximately 0.3 (dimensionless), which is calculated by multiplying our result by 1000 and taking the reciprocal.

Unlike the other particle instruments, the PAS is expected to exhibit variation in response to RSP based on the PAH content of particle emissions, and it only responds to particles under 1 μm in diameter. However, evidence suggests that the PAS-measured PAH in cigarette smoke consistently tracks RSP mass across a range of cigarette types and smoking styles. Ding et al. report that mainstream smoke for US cigarettes contains 1–1.6 μg of PAH per cigarette. The average PAS-to-RSP conversion factor of 0.83 ng μg$^{-1}$, which we observed in the present study for the PAS monitor’s response to smolder-smoked Marlboro cigarette emissions, implies that 0.083% of the emitted particle mass consists of particulate PAH. Our value for the conversion factor is similar to a value of 0.8 ng μg$^{-1}$ observed by Repace in a casino and values of 1 and 0.8–1.3 ng μg$^{-1}$ observed in two of our prior studies, which used an older version of the PAS monitor (Model PAS 1000i, EcoChem, Inc.). We found that the older PAS 1000i monitor’s response had to be reduced by a factor of 10 relative to the PAS 2000CE, due to the fact that the 1000i uses a krypton bromine UV lamp, whereas the 2000CE uses a mercury vapor UV lamp.

Our use of particles measured by the GRIMM in the 0.3–3 μm range avoided interference from non-tobacco sources of ultrafine particles (< 0.1 μm) and large dust particles (> 3 μm). The empirical GRIMM conversion factor of 6300 counts m$^{-3}$ (μg L)$^{-1}$ agrees well with a theoretical mean value of about 6500 counts m$^{-3}$ (μg L)$^{-1}$ (relative standard deviation of 0.15), which we calculated from the particle counts by assuming spherical particles, a uniform distribution of particle sizes in each size bin, a particle density of 1.1 g cm$^{-3}$, and a lognormal particle size distribution with a mass median diameter of 0.2 μm and a geometric standard deviation of 2.\textsuperscript{14}

We estimated the error associated with readings of a given monitor by computing the ratio of 1-min values for matched instruments of the same type. We also estimated the error associated with conversion of native PAS, GRIMM, NEPH, and SIDEPAK readings to RSP mass units by computing the ratio of the estimated 1-min average RSP mass units for each monitor to the native RSP mass values measured by the PZB. The results of these calculations showed generally good consistency for intra- and inter-instrument comparisons, with the bulk of errors under 10–20%.

**On-Site Monitoring Visits**

To establish typical OTS levels, we conducted 15 on-site field visits to 10 public outdoor locations containing smokers, including restaurant and pub patios, cafés, airport sidewalks, and a public park (see Tables 3 and 4 and the location schematics in Figure 1). These visits were designed so we could measure the average particle exposure attributable to emissions from real smokers that might occur during a meal at an outdoor establishment or while waiting on a sidewalk or in some other public area.

During each on-site visit, we made real-time measurements of airborne particles using the GRIMM and/or the PAS instrument, or the SIDEPAK instrument. We used the PZB as a supplemental instrument during a single visit. We used the GRIMM, PAS, and SIDEPAK for the visits, because they are more portable and unobtrusive than the PZB and NEPH monitors. For each visit, we measured OTS levels during periods with active smoking. To provide background levels, we also measured during times when no tobacco sources were active.

For 9 of the visits (S1–S9), we measured OTS particle lev-
FIGURE (1): Rough schematic diagrams of patios and sidewalks where outdoor tobacco smoke (OTS) particle levels were monitored in proximity to smokers or burning cigarettes. All the patios had at least an open roof and many were open on three sides. Broken lines represent open boundaries, and solid lines indicate a surrounding wall or an adjacent building. Tables and benches are represented by circles or ovals and rectangles indicate doorways to buildings or an opening in the wall or fence surrounding a patio. The approximate positions of active smokers and monitors during one or more visits are indicated by the letters "S" and "M", respectively. See Table 4 for dimensions and other characteristics of each OTS monitoring location.
els using the PAS and/or GRIMM while sitting or standing on each patio or sidewalk and observing the activity of nearby cigarette and cigar smokers, but, because patrons engaged in uncontrolled smoking, we were not able to make precise measurements of the distance between smokers and the monitoring instruments. The monitors were generally positioned at the approximate heights of 3 to 4 ft or table height (~3 ft). The inlets of the GRIMM and PAS monitors were placed within 12 in of each other where possible. The time spent near active smokers ranged from 0.5 to 3.4 hr per visit.

For 3 on-site visits to outdoor patios (OC1–OC3), we smoked or smolder-smoked cigarettes or cigars near the monitoring positions for smoking periods of 0.01 and 0.5 hr. We used the GRIMM and/or PAS during these visits.

Finally, during 3 site visits to sidewalk patios (OP1–OP3), we measured OTS levels using the SIDEPAK at precise distances from active cigarettes, which were either smolder-smoked or human-smoked, for periods ranging from 0.6 to 1.7 hr. We also measured temperature, air speed, and relative humidity continuously during these visits.

### Matched Monitor Experiments

To quantify the relationship between distance from the smoker and OTS concentration, i.e., the proximity effect, and to make direct comparisons between OTS and indoor SHS levels, we performed controlled experiments on four days (E1–E4) at a private residence (Tables 3 and 4, and BP1 in Figure 1) using pairs of matched PAS, NEPH, and GRIMM instruments at different distances from burning cigarettes. We smolder-smoked successive cigarettes both on the outdoor patio and inside the residence. For most experiments, we made continuous measurements of air speed, temperature, and relative humidity.

The E1 experiments consisted of six outdoor patio experiments on a single day in which a cluster of single PAS, NEPH, and GRIMM monitors were surrounded by five burning cigarettes at distances of 2, 4, or 6 ft, and heights of 3–4 feet, for periods of 10 min per experiment. The cigarettes were positioned in concentric pentagonal arrangements so that cigarettes surrounded the monitors at equal distances for each experiment. This arrangement was expected to diminish the impact of wind direction on measured concentrations. In addition to the 6 cigarette experiments, we conducted 2 experiments in which a single cigar was smoked for 20–30 min at a distance of 4 ft from the monitor cluster. For all experiments, a second, identical cluster of particle monitors, which was intended to provide continuous background levels, was positioned about 28 feet (8.5 m) from the first cluster and around the corner of the house.

For experiments E2–E4, we built two mobile particle monitoring assemblies containing PAS, NEPH, and SIDEPAK instruments fastened to wheeled chairs. On each day, we created 7–9 periods of smolder-smoked cigarette activity lasting 30–50 min, using 3 to 5 individual cigarettes burned successively. The monitoring inlets and burning cigarettes were both at an approximate height of 3–4 ft. To provide accurate background levels, we measured particle concentrations during intermediate time periods with no cigarette activity, which were of similar duration as the smoking periods. For each period of smoking activity, the two monitoring assemblies were placed on opposite sides of the source at distances of 0.25, 0.5, 1.0, 2.0, or 4.0 m. On the fourth day, the PZB instrument was added to the suite of monitoring instruments.

Immediately following 5–6 periods of controlled outdoor cigarette combustion on the backyard patio (BP1 location; E2–E4 experiments), we moved the monitoring assemblies indoors and performed several experiments in the bedroom or living room of the residence. The design of the indoor experiments was nearly identical to the outdoor experiments, except that only distances of 0.25 and 0.5 m from the burning cigarette were monitored, and the experiments were performed inside the house where all exterior doors and windows were closed during periods of smoking activity. In addition, for one of the two living room experiments, a small fan was introduced to explore

### Table 2: Native Units and Conversion Factors for Real-Time Particle Monitoring Instrument Readings

| Instrument | Native Units | Conversion Factor from Native Units to RSP Mass Conc. [µg m⁻³] | CI₉₀ | s | s/|X̄|
|------------|--------------|---------------------------------------------------------------|------|---|---|
| PZB        | µg m⁻³       | RSP mass concentration units                                  |      |   |   |
| NEPH       | 10⁻⁶ m⁻¹     | 4.6 m² g⁻¹                                                    | ±0.4 | 0.78 m² g⁻¹ | 0.17 |
| SIDEPAK    | mg m⁻³       | 3.3 x 10⁻³ mg µg⁻¹                                             | ±0.3 | 0.53 x 10⁻³ mg µg⁻¹ | 0.16 |
| GRIMM      | counts L⁻¹   | 6300 counts m⁻³ (µg L⁻¹)                                       | ±800 | 160 counts m⁻³ (µg L⁻¹) | 0.25 |
| PAS        | ng m⁻³       | 0.83 ng µg⁻¹                                                  | ±0.1 | 0.19 ng µg⁻¹ | 0.23 |

*aReal-time airborne particle monitoring instrument abbreviations: PZB = Kanomax or TSI piezobalance; NEPH = Radiance integrating nephelometer; SIDEPAK = TSI Sidepak laser photometer; GRIMM = Grimm laser counter; PAS = Ecochem photoelectric aerosol sensor.

*bThe sample mean of conversion factors from native units to estimated respirable suspended particle (RSP) mass concentration units are given for readings of each real-time airborne particle monitoring instrument. Also given are the 90% confidence intervals for the sample mean (p. 74). Dividing the native units by the conversion factor gives RSP units of µg m⁻³.

### Table 3: Summary of Outdoor Tobacco Smoke (OTS) Surveys and Experiments

<table>
<thead>
<tr>
<th>aName</th>
<th>bLocation(s)</th>
<th>cSources</th>
<th>dDuration</th>
<th>eOverall Avg. OTS RSP Conc. [µg m⁻³]</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>PP1, PP2</td>
<td>H, CG, C</td>
<td>3.3 hr</td>
<td>64 (PAS)</td>
</tr>
<tr>
<td>S2</td>
<td>PP2</td>
<td>H, CG, C</td>
<td>1.0 hr</td>
<td>50 (PAS)</td>
</tr>
<tr>
<td>S3</td>
<td>PP1</td>
<td>H, CG, C</td>
<td>1.3 hr</td>
<td>29 (PAS), 51 (GRIMM)</td>
</tr>
<tr>
<td>S4</td>
<td>RC</td>
<td>H, C</td>
<td>3.0 hr</td>
<td>6 (PAS)</td>
</tr>
<tr>
<td>S5</td>
<td>RC</td>
<td>H, C</td>
<td>3.4 hr</td>
<td>10 (PAS)</td>
</tr>
<tr>
<td>S6</td>
<td>PP3</td>
<td>H, CG, C</td>
<td>1.1 hr</td>
<td>30 (PAS), 42 (GRIMM)</td>
</tr>
<tr>
<td>S7</td>
<td>PP3</td>
<td>H, C</td>
<td>1.4 hr</td>
<td>26 (GRIMM)</td>
</tr>
<tr>
<td>S8</td>
<td>AP</td>
<td>H, C</td>
<td>0.6 hr</td>
<td>31 (PAS), 30 (GRIMM)</td>
</tr>
<tr>
<td>S9</td>
<td>AP</td>
<td>H, C</td>
<td>0.5 hr</td>
<td>56 (PAS), 15 (GRIMM)</td>
</tr>
</tbody>
</table>

### On-site Surveys with Uncontrolled Human Smokers

<table>
<thead>
<tr>
<th>aName</th>
<th>bLocation(s)</th>
<th>cSources</th>
<th>dDuration</th>
<th>eOverall Avg. OTS RSP Conc. [µg m⁻³]</th>
</tr>
</thead>
<tbody>
<tr>
<td>OC1</td>
<td>PP3</td>
<td>H, CG, C</td>
<td>0.5 hr</td>
<td>62 (PAS) 17 (GRIMM)</td>
</tr>
<tr>
<td>OC2</td>
<td>PK</td>
<td>S, C</td>
<td>0.4 hr</td>
<td>67 (PAS), 23 (GRIMM), 60 (PZB)</td>
</tr>
<tr>
<td>OC3</td>
<td>BP2</td>
<td>H, C</td>
<td>0.1 hr</td>
<td>27 (GRIMM)</td>
</tr>
</tbody>
</table>

### On-site Proximity Experiments with Controlled Smolder-Smoked Cigarettes or a Controlled Smoker

<table>
<thead>
<tr>
<th>aName</th>
<th>bLocation(s)</th>
<th>cSources</th>
<th>dDuration</th>
<th>eOverall Avg. OTS RSP Conc. [µg m⁻³]</th>
</tr>
</thead>
<tbody>
<tr>
<td>OP1</td>
<td>SC1, SC3</td>
<td>S, C</td>
<td>1.7 hr</td>
<td>133 (SIDEPAK)</td>
</tr>
<tr>
<td>OP2</td>
<td>RP</td>
<td>S, C</td>
<td>0.6 hr</td>
<td>106 (SIDEPAK)</td>
</tr>
<tr>
<td>OP3</td>
<td>SC2</td>
<td>H, C</td>
<td>1.4 hr</td>
<td>109 (SIDEPAK)</td>
</tr>
</tbody>
</table>

### Private Patio Experiments with Controlled Smolder-Smoked Cigarettes or Machine-Smoked Cigars

<table>
<thead>
<tr>
<th>aName</th>
<th>bLocation(s)</th>
<th>cSources</th>
<th>dDuration</th>
<th>eOverall Avg. OTS RSP Conc. [µg m⁻³]</th>
</tr>
</thead>
<tbody>
<tr>
<td>E1</td>
<td>BP1</td>
<td>S, C</td>
<td>2.0 hr</td>
<td>48 (PAS), 19 (GRIMM), 10 (NEPH)</td>
</tr>
<tr>
<td>E2</td>
<td>BP1</td>
<td>S, C</td>
<td>3.7 hr</td>
<td>47 (PAS), 28 (GRIMM), 10 (NEPH)</td>
</tr>
<tr>
<td>E3</td>
<td>BP1</td>
<td>S, C</td>
<td>3.9 hr</td>
<td>61 (PAS), 29 (GRIMM), 22 (NEPH)</td>
</tr>
<tr>
<td>E4</td>
<td>BP1</td>
<td>S, C</td>
<td>2.5 hr</td>
<td>38 (PAS), 18 (GRIMM), 16 (NEPH)</td>
</tr>
</tbody>
</table>

1. S1–S9 = on-site visits (surveys) of patios and sidewalk areas with human smokers, OC1–OC3 = on-site controlled visits (surveys) for which the investigators controlled the smoking or smolder-smoking of one or more cigarettes or cigars near the monitors, OP1–OP3 = on-site proximity experiments with controlled smolder- or human-smoked cigarettes positioned at precise distances from the monitoring positions, and E1–E4 = controlled experiments performed at a private residence (patio, living room, bedroom) with smolder-smoked cigarettes positioned at precise distances from two separate monitoring positions.

2. Codes refer to one of the outdoor locations listed in Table 4: SC1, SC2, SC3 = sidewalk cafés; PP1, PP2, PP3 = pub patios; RC = resort café; RP = restaurant patio; PK = park plaza; AP = airport sidewalk; BP1, BP2 = private backyard patio.

3. H = human smoked, S = smolder-smoked, CG = cigars, C = cigarettes or cigarillos

4. D = Duration of the monitoring period during which OTS sources were intermittently or continuously active.

5. The estimated average OTS respirable suspended mass (RSP) concentration in µg m⁻³ determined by converted measurements of a PAS, GRIMM, NEPH or SIDEPAK instrument (indicated in parentheses) taken during times when cigarettes or cigars were active. Background levels were subtracted. Abbreviations: PAS = Ekochem photoelectric aerosol sensor, GRIMM = Grimm laser particle counter; PZB = Kanomax or TSI piezobalance; NEPH = Sidepak laser photometer. Results for S1–S9 include time when smokers were intermittently active at a location. Results for OC1–OC3, OP1–OP3, and E1–E4 include times when a cigarette or cigar was smoked or smolder-smoked by the investigators near the monitoring position. While experiments E2–E4 included indoor SHS measurements, they were not included in the calculated average OTS particle concentrations shown in the table.
spikes in airborne particle levels when cigars or cigarettes were active. The structure of the peaks could be observed using the NEPH and SIDEPAK instruments, which provided readings at intervals of 2 sec and 10 sec, respectively (see Figure 2). Some peaks exceeded 1,000 µg m$^{-3}$, which is seen in close proximity to activity sources, have been attributed to “microplumes” by previous investigators, who observed them within 2 m of indoor point sources of pollution. Microplumes are defined as thin concentrated streams of smoke, or some other air pollutant, that follow complex trajectories during periods of release. When the microplumes impinge on a monitor inlet, the monitor momentarily registers a peak in concentration. Over time and at further distance from the source indoors, the microplumes dissipate and pollution becomes well mixed in an interior space, persisting long after the source has been extinguished. In contrast to persistent and mixed indoor levels, which exhibit smooth rises and decays in concentration, OTS consists entirely of periods characterized by microplumes. There is no period where OTS is well mixed, and OTS disappears almost instantly when tobacco sources are extinguished.

We analyzed the OTS data in terms of raw concentration readings, 1-min average concentrations, and averages on a per-experiment, per-site, or overall basis. All results presented are for periods of continuous (experiments) or intermittent (on-site visits) active smoking. Prior to averaging and data analysis, we subtracted background levels for each day’s worth of data from each monitor. We created a consistent and integrated database by calculating 1-min averages for each monitor and by converting the native units of each monitor into units of RSP mass concentration (µg m$^{-3}$) using the mean conversion factors in Table 2. The quantitative discussion of variation in OTS levels during each monitoring episode refers to either peak values over intervals as low as 2 or 10 sec, or to 1-min average levels. During nearly all the outdoor monitoring periods on patios and sidewalks where relative humidity was measured, it was fairly low, averaging about 40% with a range of 20–65%. Therefore, correction of OTS levels due to high relative humidity was deemed unnecessary. Where measured, outdoor temperatures averaged 26 °C with a range of 10–38 °C and outdoor ground-level wind speeds (~1 m above ground) averaged 0.41 m sec$^{-1}$ with a range of approximately 0.0 to 1.2 m sec$^{-1}$.

### RESULTS AND DISCUSSION

Measured concentrations of OTS consistently showed sharp spikes in airborne particle levels when cigars or cigarettes were active. The structure of the peaks could be observed using the NEPH and SIDEPAK instruments, which provided readings at intervals of 2 sec and 10 sec, respectively (see Figure 2). Some peaks exceeded 1,000 µg m$^{-3}$. Transitory peaks of this nature, which are seen in close proximity to activity sources, have been attributed to “microplumes” by previous investigators, who observed them within 2 m of indoor point sources of pollution. Microplumes are defined as thin concentrated streams of smoke, or some other air pollutant, that follow complex trajectories during periods of release. When the microplumes impinge on a monitor inlet, the monitor momentarily registers a high peak in concentration. Over time and at further distance from the source indoors, the microplumes dissipate and pollution becomes well mixed in an interior space, persisting long after the source has been extinguished. In contrast to persistent and mixed indoor levels, which exhibit smooth rises and decays in concentration, OTS consists entirely of periods characterized by microplumes. There is no period where OTS is well mixed, and OTS disappears almost instantly when tobacco sources are extinguished.

We analyzed the OTS data in terms of raw concentration readings, 1-min average concentrations, and averages on a per-experiment, per-site, or overall basis. All results presented are for periods of continuous (experiments) or intermittent (on-site visits) active smoking. Prior to averaging and data analysis, we subtracted background levels for each day’s worth of data from each monitor. We created a consistent and integrated database by calculating 1-min averages for each monitor and by converting the native units of each monitor into units of RSP mass concentration (µg m$^{-3}$) using the mean conversion factors in Table 2. The quantitative discussion of variation in OTS levels during each monitoring episode refers to either peak values over intervals as low as 2 or 10 sec, or to 1-min average levels. During nearly all the outdoor monitoring periods on patios and sidewalks where relative humidity was measured, it was fairly low, averaging about 40% with a range of 20–65%. Therefore, correction of OTS levels due to high relative humidity was deemed unnecessary. Where measured, outdoor temperatures averaged 26 °C with a range of 10–38 °C and outdoor ground-level wind speeds (~1 m above ground) averaged 0.41 m sec$^{-1}$ with a range of approximately 0.0 to 1.2 m sec$^{-1}$.

### TYPICAL OTS LEVELS

Tables 3 and 5 contain overall average OTS particle mass concentrations for periods of smoking during the outdoor on-site monitoring episodes.
A. Real-time outdoor tobacco smoke (OTS) and indoor secondhand smoke (SHS) respirable suspended particle (RSP) mass concentrations determined from raw 2-sec NEPH instrument readings during a suite of patio experiments (E3) performed in the backyard of a residence using smolder-smoked cigarettes. Average RSP mass concentrations are shown for each period when cigarettes were active, indicated by solid horizontal bars, for both northerly and southerly monitoring positions at source-receptor distances of 0.25, 0.5, 1, and 2 m. The southerly average concentrations, shown in larger typeface, were consistently higher than the northerly ones for outdoor measurements, likely because the prevailing winds were in the southerly direction. Outdoor air speed averaged 0.5 m sec\(^{-1}\) on the patio during times that cigarettes were active. The indoor air speed was close to zero.

B. Real-time OTS RSP mass concentrations determined from raw 10-sec SIDEPAK instrument readings during an on-site proximity experiment (OP3) performed on a sidewalk patio with a human smoker. Average mass concentrations during periods of smoking are indicated by solid horizontal bars. The distance of the monitor from the smoker, which ranged over 4 values between 0.5 and 3.7 m, is also given. Air speed averaged 0.16 m sec\(^{-1}\) during times that cigarettes were active.

C. Real-time OTS RSP mass concentrations determined from raw 10-sec SIDEPAK instrument readings during an on-site proximity experiment (OP1) performed on a sidewalk patio where cigarettes were smolder-smoked at 5 different distances from the instrument, ranging from 0.3 m (1 ft) to 2.7 m (9 ft). Average mass concentrations during periods of smoking are indicated by solid horizontal bars. During this set of experiments, wind was consistently blowing in a single direction along the sidewalk. All concentrations were monitored in the downwind direction, except for the second cigarette at 0.6 m, for which concentrations were monitored in the opposite (upwind) direction. Air speed averaged 0.5 m sec\(^{-1}\) during times that cigarettes were active.

Figure (2): A. Real-time outdoor tobacco smoke (OTS) and indoor secondhand smoke (SHS) respirable suspended particle (RSP) mass concentrations determined from raw 2-sec NEPH instrument readings during a suite of patio experiments (E3) performed in the backyard of a residence using smolder-smoked cigarettes. Average RSP mass concentrations are shown for each period when cigarettes were active, indicated by solid horizontal bars, for both northerly and southerly monitoring positions at source-receptor distances of 0.25, 0.5, 1, and 2 m. The southerly average concentrations, shown in larger typeface, were consistently higher than the northerly ones for outdoor measurements, likely because the prevailing winds were in the southerly direction. Outdoor air speed averaged 0.5 m sec\(^{-1}\) on the patio during times that cigarettes were active. The indoor air speed was close to zero. B. Real-time OTS RSP mass concentrations determined from raw 10-sec SIDEPAK instrument readings during an on-site proximity experiment (OP3) performed on a sidewalk patio with a human smoker. Average mass concentrations during periods of smoking are indicated by solid horizontal bars. The distance of the monitor from the smoker, which ranged over 4 values between 0.5 and 3.7 m, is also given. Air speed averaged 0.16 m sec\(^{-1}\) during times that cigarettes were active. C. Real-time OTS RSP mass concentrations determined from raw 10-sec SIDEPAK instrument readings during an on-site proximity experiment (OP1) performed on a sidewalk patio where cigarettes were smolder-smoked at 5 different distances from the instrument, ranging from 0.3 m (1 ft) to 2.7 m (9 ft). Average mass concentrations during periods of smoking are indicated by solid horizontal bars. During this set of experiments, wind was consistently blowing in a single direction along the sidewalk. All concentrations were monitored in the downwind direction, except for the second cigarette at 0.6 m, for which concentrations were monitored in the opposite (upwind) direction. Air speed averaged 0.5 m sec\(^{-1}\) during times that cigarettes were active.
field visits and for breakdowns by various factors. Note that the results presented in Table 5 are not meant to imply direct comparisons of concurrent measurements for the different instruments, since not all the monitors were used during a given visit.

As determined from PAS instrument measurements during the on-site visits with natural and controlled smoking (S1—S6, S8—S9, and OC1—OC2), average estimated RSP mass concentrations of OTS particles on a given day ranged from 6 to 67 µg m\(^{-3}\) with an overall average of 33 µg m\(^{-3}\). The estimated GRIMM RSP levels for similar visits to outdoor patios (S3, S6—S9, and OC1—OC3) ranged from 17 to 51 µg m\(^{-3}\) with an average of 34 µg m\(^{-3}\). The PZB levels from a single visit with controlled smoking near the monitor (OC2) averaged 60 µg m\(^{-3}\) (0.4 hr averaging period).

In general, the variation in 1-min average OTS levels (Table 5) was very high with overall relative standard deviations of 1.7 for the PAS and GRIMM instruments. This variation results from the occurrence of sharp spikes in the OTS concentration time series due to swirling microplumes. Peaks in 1-min average OTS levels during site visits were observed to reach as high as 300–600 µg m\(^{-3}\) as measured by the PAS and GRIMM instruments.

The estimated RSP mass concentrations determined from PAS measurements in the present work may have been influenced by non-tobacco sources or differences in PAH emissions for different types of tobacco products or smoking styles relative to what we used during the calibration experiments. Ott and Siegmann\(^{22}\) report very different PAH concentrations for different combustion sources. In the current study, we found the PAS monitor was more sensitive to some non-OTS particles, such as diesel exhaust and soot from some types of candles, than the other instruments, because these emissions are high in PAH. We minimized bias in the PAS measurements caused by other sources by including only levels for the PAS when no non-OTS sources or unexplained concentrations were observed.

In spite of possible interference from other sources, the general validity of the PAS results – and their applicability to estimating OTS RSP – is supported by their generally good agreement with the estimated RSP levels derived from the GRIMM instrument. Some of the differences we observed between the two instruments may have resulted from microplume effects, in which localized peaks in particle concentration occurred near only one monitor’s inlet at a given instant.

To facilitate direct comparisons to PAS measurements performed in other studies, the estimated RSP values reported here can be converted back to the native ng m\(^{-3}\) units of the PAS instrument by using the conversion factor of 0.83 ng µg\(^{-1}\) presented above. For example, the average per-visit particle-bound PAH concentrations measured during on-site surveys where smoking occurred were 5–56 ng m\(^{-3}\) with an overall average of 27 ng m\(^{-3}\), which is similar to the particulate PAH concentrations reported by Ott and Siegmann\(^{22}\) using the same PAS 2000CE monitor.

As shown in Table 5, overall average OTS concentrations for time periods when both cigarettes and cigars were active (50 and 43 µg m\(^{-3}\) for PAS and GRIMM, respectively) were 40–70% higher than those when only cigarettes were active (16 and 25 µg m\(^{-3}\)). This result may have occurred because cigars are active over a longer period of time than are individual cigarettes. In addition, average OTS concentrations measured by the PAS and GRIMM instruments during visits to outdoor patios that were enclosed by fences or walls (PP1 and PP2 locations) were 50% and 43% higher, respectively, than those observed in more open areas (52 and 51 µg m\(^{-3}\) versus 21 and 29 µg m\(^{-3}\)). In the more open patios (SC1–SC3, BP2, PK, RC, RP, and AP locations), which may have contained tables, chairs, umbrellas, and low fences, air could flow across the patio, perhaps influenced by a “street canyon” effect characterized by air movement in a consistent direction along building boundaries. In contrast, the enclosed patios had walls on four sides that protected patrons from wind and may have contained OTS emissions to a greater degree.

**Outdoor Versus Indoor Concentrations**

The three days of monitoring at a residence (E2—E4), during which parallel measurements were performed indoors and outdoors using the PAS, GRIMM, NEPH, and PZB instruments, provide data for direct comparisons between OTS levels and indoor SHS particle concentrations observed during periods of active smoking for these experiments. Figure 2A shows the complete time series of one set of experiments (E3) for the NEPH instrument.

The effect of accumulation of cigarette emissions indoors and the effect of room volume were plainly evident during the experiments. Whereas OTS concentrations dropped immediately to background levels when the cigarette sources were extinguished, indoor SHS concentrations persisted at relatively high levels and slowly decayed for hours until the doors were opened to ventilate the house.

As expected, the smaller bedroom with a volume of 44 m\(^3\) had a larger average indoor SHS particle concentrations during smoking (105 µg m\(^{-3}\) from PZB) than the living room (35 µg m\(^{-3}\) from PZB), which had a volume of more than 400 m\(^3\) (see Table 6). The average indoor SHS levels observed in this study were similar to those observed by Özkanayak et al.\(^{34}\), who report that secondhand smoke contributes approximately 30 µg m\(^{-3}\) on average to indoor particle levels in homes. In the present study, we observed PZB particle mass peaks in the living room and bedroom of approximately 50 and 200 µg m\(^{-3}\), respectively, which are similar to peak values that we observed in previous real-time monitoring studies of cigar and cigarette smoking in homes.\(^{31,35}\)

The average OTS particle concentrations we observed during each experiment across all distances were 10–22 µg m\(^{-3}\) for the NEPH, 18–29 µg m\(^{-3}\) for the GRIMM, and 38–61
### Table (5): Observed OTS Particle Concentrations During Patio and Sidewalk Café On-Site Surveys in RSP Mass Units [µg m⁻³]

<table>
<thead>
<tr>
<th>Factor</th>
<th>PAS³</th>
<th></th>
<th>GRIMM⁴</th>
<th></th>
<th>PZB⁵</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>µ</td>
<td>s</td>
<td>µ/µ</td>
<td>µ</td>
</tr>
<tr>
<td>Overall³</td>
<td>852</td>
<td>33</td>
<td>55</td>
<td>1.7</td>
<td>339</td>
</tr>
<tr>
<td></td>
<td></td>
<td>µ</td>
<td>s</td>
<td>µ/µ</td>
<td>µ</td>
</tr>
<tr>
<td>Cigarettes³</td>
<td>434</td>
<td>16</td>
<td>32</td>
<td>2.0</td>
<td>178</td>
</tr>
<tr>
<td>Cigars+Cigarettes³</td>
<td>418</td>
<td>50</td>
<td>66</td>
<td>1.3</td>
<td>161</td>
</tr>
<tr>
<td>Closed Area⁶</td>
<td>326</td>
<td>52</td>
<td>69</td>
<td>1.3</td>
<td>75</td>
</tr>
<tr>
<td>Open Area⁸</td>
<td>526</td>
<td>21</td>
<td>40</td>
<td>1.0</td>
<td>264</td>
</tr>
</tbody>
</table>

|                  |      | µ          | s      | µ/µ      | µ    |

³ This table contains grouped descriptive statistics calculated from 1-min average outdoor tobacco smoke (OTS) particle measurements observed during 9 site visits S1–S9, where natural smoking of cigarettes and cigars by smokers occurred (intermittent smoking), and 3 on-site visits OC1–OC3, during which one or more cigarettes or cigars were smoke-stopped by the investigators near the monitor(s) (continuous smoking). The respirable suspended particle (RSP) mass units for the PAS and GRIMM were estimated using conversion factors from native PZB RSP values based on the results of controlled collocation experiments using all monitors and a cigarette source (see text). Background levels were subtracted from all instrument measurements. The monitors are abbreviated as follows: PAS = Ecochem photoelectric aerosol sensor; GRIMM = Grimm laser particle counter; and PZB = Kanomax or TSI piezobalance. The abbreviations for statistics are: n = sample size of 1-min average values; µ = RSP sample mean in [µg m⁻³]; s = RSP sample standard deviation in [µg m⁻³]; and s/µ = RSP relative standard deviation (dimensionless).

⁴ The PAS and GRIMM were used together for 6 out of the 13 visits (see Table 3).

⁵ The PZB was only present at the OC2 visit (see Table 3).

⁶ Results taken over all 13 visits.

⁷ Results for time periods when only cigarettes were observed to be active.

⁸ Results for time periods when both cigars and cigarettes were observed to be active.

⁹ Closed areas are patios located at restaurants or pubs and enclosed with a fence or wall on all sides so that directional air flow was effectively impeded (PP2 and PP3).

Open areas were sidewalks, sidewalk cafés, or parks where, although there may have been trees, umbrellas, and low barriers, there was enough open space that a potential "street canyon" effect could occur whereby air flow was channeled across the patio due to the presence of surrounding buildings (SC1, SC2, SC3, BP2, PK, RC, RP, AP).

### Wind Effect

The experiment in the living room of the residence, where a fan was used to blow the plume of a burning cigarette toward a set of monitors at an air speed of about 0.4 m sec⁻¹, demonstrates how wind can elevate OTS levels in downwind directions (Figure 2A). For this particular experiment, the fan increased average NEPH levels during smoking by approximately 3 times at a downwind monitor relative to an upwind monitor.

This effect is further illustrated by our observation that the two sets of monitors positioned on either side of the active cigarette sources on the outdoor residential patio recorded much different OTS particle levels. The average levels in the northerly direction were about 40–60% lower than those in the southerly direction (Table 6). From the time profiles for one set of measurements (Figure 2A), it is evident that outdoor levels could be higher than corresponding indoor (non-fan) levels in one direction, but near zero in the opposite direction.

The clearest evidence that wind leads to extremely high OTS levels during smoking was provided by the results of the OP1 experiment at the first sidewalk café where 6 cigarettes were smoke-stopped at 5 distances from the SIDEPAK monitor (Figure 2C). For this experiment, the wind was observed to consistently blow the smoke microplumes in a single direction at an average speed of 0.5 m sec⁻¹ when cigarettes were active. Upwind levels were practically zero, whereas the average downwind particle levels during smoking were 582 [µg m⁻³] at 0.3 m and even at 1.2–2.7 m they were still elevated above back-
table (6): observed ots and indoor shs particle concentrations during controlled experiments e1–e4 and op1–op3 in rsp mass units [µg m⁻³]n

<table>
<thead>
<tr>
<th>factor</th>
<th>pas</th>
<th>grimm</th>
<th>neph</th>
<th>pzb</th>
<th>sidepak</th>
</tr>
</thead>
<tbody>
<tr>
<td>outdoor⁴</td>
<td>1029 50 113 2.3</td>
<td>1052 22 50 2.3</td>
<td>1052 15 26 1.7</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>living room⁴</td>
<td>235 33 33 1.0</td>
<td>235 30 35 1.2</td>
<td>235 32 22 0.7</td>
<td>30 35 12 0.3</td>
<td>–</td>
</tr>
<tr>
<td>bedroom⁴</td>
<td>22 46 48 1.0</td>
<td>22 106 105 1.0</td>
<td>22 95 80 0.8</td>
<td>11 105 74 0.7</td>
<td>–</td>
</tr>
<tr>
<td>[0.25, 0.5) m</td>
<td>328 108 175 1.6</td>
<td>332 45 76 1.7</td>
<td>332 35 38 1.1</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>[0.5, 1) m</td>
<td>202 43 72 1.7</td>
<td>202 16 21 1.3</td>
<td>202 11 11 1.0</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>[1, 2) m</td>
<td>301 19 25 1.3</td>
<td>310 12 34 2.9</td>
<td>310 7 8 1.3</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>[2, 4] m</td>
<td>198 8 9 1.1</td>
<td>208 4 5 1.3</td>
<td>208 2 2 0.9</td>
<td>–</td>
<td>33 11 7 0.6</td>
</tr>
<tr>
<td>northerly</td>
<td>465 28 76 2.7</td>
<td>465 17 54 3.3</td>
<td>465 12 26 2.2</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>southerly</td>
<td>451 72 144 2.0</td>
<td>465 27 45 1.6</td>
<td>465 20 28 1.4</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>downwind⁸</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>upwind⁸</td>
<td>10 2.5 1.0</td>
<td>0.4</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

× this table contains grouped descriptive statistics calculated from 1-min average outdoor tobacco smoke (ots) particle measurements observed during 4 controlled day-long experiments at a residence (e1–e4) and 3 on-site proximity experiments (op1–op3) for which distance from continuously active tobacco sources was recorded precisely. the respirable suspended particle (rsp) mass units for pas, grimm, neph, and sidepak instruments are estimated based on conversion factors to pzb rsp mass concentration units that were calculated from the results of controlled cigarette smoking experiments performed using the collocated monitoring instruments. background levels were subtracted. the monitors are abbreviated as follows: pas = ecochem photoelectric aerosol sensor; grimm = grimm laser particle counter; neph = radiance integrating nephelometer; pzb = kanomax piezobalance; and sidepak = tsi laser photometer. the abbreviations for statistics are: n = sample size of 1-min average values; r = rsp sample mean in µg m⁻³; s = rsp sample standard deviation in µg m⁻³; and s/x = relative standard deviation (dimensionless).

⁴ the pas, grimm, and neph were used together for the e1–e4 day-long experiments (see table 3). the pzb was only used during the inside portion of the e4 experiments.

⁵ the sidepak was only used (by itself) during the op1–op3 proximity experiments (see table 3).

⁶ the “outdoor” row contains statistics calculated from outdoor tobacco smoke (ots) levels across all experiments. the “living room” and “bedroom” rows contain indoor secondhand smoke (shs) results for the two indoor locations when the fan was off or monitors were upwind from the fan. indoor shs levels were only measured at distances of 0.25 and 0.5 m from the monitoring positions. apart from the two rows labeled “living room” and “bedroom,” all rows in the table are for ots levels only.

⁷ the distance from the source in four groups for outdoor tobacco smoke (ots) levels only. [“or “] indicates left or right limit is inclusive and “)” indicates right limit is exclusive.

⁸ for three outdoor experiments on the residential patio (e2–e4), groups of monitors were placed in northerly and southerly directions.

ground by 13–41 µg m⁻³. the 10-sec spikes in the downwind ots particle time series sometimes exceeded 1500 µg m⁻³.

proximity effect

we observed a clear reduction in ots levels as the distance from a tobacco source increased. generally, average levels within 0.5 m from a single cigarette source were quite high and comparable to indoor levels, and ots levels at distances greater than 1 or 2 m were much lower. however, during on-site proximity experiments op1 and op3, ots was still detectable by the sidepak at distances of about 3–4 m from a single cigarette on sidewalk patios. a neph instrument also registered slightly elevated particle concentrations at a distance of 8 m from a cluster of burning cigarettes and around the corner of the house during a backyard patio experiment (e1).

to summarize and quantify the proximity effect observed in our study, we fit curves to average ots particle concentrations (y) as a function of the distance from the source (x). figure 3 shows two curves with separate fits for data from the sidewalk cafés (op1–op3: y = 44.4x⁻² + 27x⁻¹ + 4.1) and the backyard patio (e1–e4: y = −0.3x⁻² + 16.8x⁻¹ − 2.8) where distances were measured precisely. every point represents the overall average for a given distance across all smoking periods and instruments at a given type of location. the levels on the private patio were generally lower and dropped off by 1–2 m, whereas the café levels, where winds may have been stronger and/or more directional, started out about 4 times higher and did not entirely drop off by 4 m.

previous ots studies

prior to the current study, few data on ots levels have been available. in an unpublished study, the california air resources board (carb) measured 1-hr and 8-hr average nicotine concentrations, number of active cigarettes, and wind characteristics outside an airport, a college, a government center, an office complex, and an amusement park (chapter 5, pp.v6-v19). average ots nicotine concentrations were strongly affected by counts of the number of smokers and moderately affected by the size of the smoking area and the measured wind speed. the observed 8-hr average ots nicotine levels
in locations with relatively stronger winds or a smaller number of smokers were about 0.1 $\mu g$ m$^{-3}$ or less. In locations with a larger number of smokers, the levels could reach 1 or 3 $\mu g$ m$^{-3}$. These OTS levels are in the middle range of observed indoor SHS nicotine levels, which can average from 0.01 to 10 $\mu g$ m$^{-3}$. Based on the CARB study, Californians spending time close to outdoor smokers could potentially be exposed to OTS levels similar to those associated with indoor SHS concentrations.

The general findings of the CARB study are compatible with the findings of the current work. The CARB results establish the potential for relatively high OTS exposures in places where smokers congregate. The experiments in the current work go further to quantify potential exposures under specific wind and proximity conditions – focusing on single smokers. Extrapolation of our controlled experimental methods and results to multiple smokers is complex, because one must consider the relative positions between each source and the receptor. Generally, we would expect that exposure increases in proportion with the number of smokers. The exact increase depends on the amount of time the receptor spends downwind and at a given distance from each source.

**Incremental Contributions to 24-hr Total Exposure**

It is useful to calculate per-cigarette 24-hr incremental exposure ($IE_{24}$) concentrations for OTS, where $IE_{24}$ is defined as the contribution of a given OTS-related event involving one cigarette to a person’s 24-hr total particle exposure. For example, during on-site experiment OP1 we observed an average OTS particle concentration at a distance of 0.3 m from a cigarette of 582 $\mu g$ m$^{-3}$ in the downwind direction. Since the cigarette lasted about 10 min, we calculate a per-cigarette 24-hr incremental exposure as follows: $IE_{24} = 582 \mu g$ m$^{-3} \times 10$ min / 1440 min = 4 $\mu g$ m$^{-3}$. The calculation amounts to a weighting of the per-cigarette average concentration by the proportion of time the cigarette lasts with respect to the 24-hr (1440-min) day.

The incremental exposure concept allows one to combine exposures for different events and compare the total to health-related standards or other reference levels. For example, if a person experienced 9 cigarette events over the course of their day – with each event similar to the one that occurred at 0.3 m in the OP1 experiments – then their overall 24-hr OTS particle exposure would be $9 \times 4$ $\mu g$ m$^{-3} = 36$ $\mu g$ m$^{-3}$. This exposure would exceed the USEPA 24-hr health-based ambient standard for fine particles, which is currently 35 $\mu g$ m$^{-3}$. Note that the USEPA standard was devised for ambient air pollution, which is likely to have substantially different composition than tobacco smoke pollution. However, since secondhand smoke contains many toxic compounds, including carcinogens, it is likely that, at a given airborne particle concentration, OTS carries the greater risk.

**Summary and Conclusions**

The measurement of OTS is a new area in terms of epidemiologic and human exposure investigations. The present work provides some of the first evidence that OTS levels can be substantial under certain conditions of wind and proximity. The major findings of our research are summarized as follows:

- Real-time particle instruments, especially those based on light scattering, are useful in characterizing the determinants of OTS levels, which fluctuate on a time scale of seconds. The different particle detection instruments provide consistent findings and support the general conclusion that significant OTS levels can occur near smokers.

- Outdoor particle concentrations measured close to a cigar or cigarette exhibit multiple concentration spikes or “microplumes”, which are similar to those that have been observed close to indoor particle sources.

- Average OTS particle levels near active sources over the course of 1 or more cigarettes can be comparable to average well-mixed indoor SHS particle levels observed to occur in living rooms or bedrooms during active smoking. Average OTS particle concentrations can reach 100’s of $\mu g$ m$^{-3}$. Unlike for indoor SHS, OTS levels drop to zero when smoking ends.

- OTS levels are highly dependent on wind conditions. Upwind levels are likely to be very low, whereas downwind
OTS levels during periods of active smoking can be very large with 10-sec peak levels at the closest positions potentially exceeding 1500 µg m⁻³ and average levels over the duration of a single cigarette potentially exceeding 500 µg m⁻³.

- OTS levels are highly dependent on source proximity. Levels at 0.25–0.5 m can drop by half or more as the distance increases to 1–2 m. At distances larger than 2 m, levels near single cigarettes were generally close to background. The concentrations at different distances are influenced by wind conditions. We found that it was possible for there to be detectable OTS levels at downwind positions of 4 m or more from a single active cigarette. Also, as the number of active cigarettes increases, the distance at which OTS is detectable is likely to increase.

- In outdoor restaurant patios and parks, where there may be multiple smokers, between 8 and 20 cigarettes smoked sequentially could cause an incremental 24-hr particle exposure greater than a threshold level of 35 µg m⁻³ for a person who is within 0.5 m of the smokers. This threshold level is the 24-hr USEPA health-based standard for fine particles.

Our results demonstrate that OTS can be high during periods of smoking in locations where persons are near active smokers. Therefore, it is possible for OTS to present a nuisance or hazard under certain conditions. Examples of scenarios where OTS levels might be high include eating dinner with a smoker on an outdoor patio, sitting at a table next to a smoker in a sidewalk café, sitting next to a smoker on a park bench, or standing near a smoker outside a building. Children who accompany a smoking parent or guardian may experience substantial exposure. Outdoor restaurant or pub workers who spend a significant portion of their time within a few feet of active smokers are also likely to receive relatively large total OTS exposures over the course of a day – possibly exceeding the USEPA 24-hr health standard for fine particles. If one is upwind from a smoker, levels most likely will be negligible. However, if the smoker’s position changes or one spends time downwind from a smoker, then moving to a distance of more than 2 m can reduce the likelihood of experiencing elevated particle exposure due to OTS. Future studies should measure OTS levels for dynamic situations with multiple smokers, including continuous measurements of personal OTS concentrations or biomarker levels for workers in outdoor locations.

Support for health-based OTS bans may lie in a potential acute effect on susceptible populations. Short-term OTS exposures might be life threatening for high-risk persons, since the human cardiovascular system is very sensitive to secondhand smoke. A recent before-and-after smoking ban study showed a decreased chance of myocardial infarction when a ban was in place, which suggests that there is an acute risk associated with SHS exposure for persons at increased risk of coronary heart disease or with known coronary artery disease.

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**ABOUT THE AUTHORS**

Neil E. Klepeis is a consulting assistant professor in the Department of Civil and Environmental Engineering at Stanford University. He has studied human exposure to secondhand smoke for over 13 years. Wayne R. Ott is a consulting professor in the Department of Civil and Environmental Engineering and a visiting scholar in the Department of Statistics at Stanford University. Dr. Ott worked for 30 years for the US Environmental Protection Agency, helping to develop exposure science as a recognized field. Paul Switzer is a professor in the Department of Statistics at Stanford University. Professor Switzer has worked on topics in human exposure analysis for over 25 years. Please address correspondence to: Dr. Neil E. Klepeis, URL: http://klepeis.net.