Sub-Daily Exposure to Fine Particulate Matter and Ambulance Dispatches during Wildfire Seasons: A Case-Crossover Study in British Columbia, Canada

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BACKGROUND: Exposure to fine particulate matter (PM2.5) during wildfire seasons has been associated with adverse health outcomes. Previous studies have focused on daily exposure, but PM2.5 levels in smoke events can vary considerably within 1 d.

OBJECTIVES: We aimed to assess the immediate and lagged relationship between sub-daily exposure to PM2.5 and acute health outcomes during wildfire seasons in British Columbia.

METHODS: We used a time-stratified case-crossover study design to evaluate the association between modeled hourly PM2.5 and ambulance dispatches during wildfire seasons from 2010 to 2015. Distributed lag nonlinear models were used to estimate the lag-specific and cumulative odds ratios (ORs) at lags from 1 to 48 h. We examined the relationship for all dispatches and dispatches related to respiratory, circulatory, and diabetic conditions, identified by codes for ambulance dispatch (AD), paramedic assessment (PA) or hospital diagnosis (HD).

RESULTS: Increased respiratory health outcomes were observed within 1 h of exposure to a 10-µg/m3 increase in PM2.5. The 48-h cumulative OR [95% confidence interval (CI)] was 1.038 (1.009, 1.067) for the AD code Breathing Problems and 1.098 (1.013, 1.189) for PA code Asthma/COPD. The point estimates were elevated within 1 h for the PA code for Myocardial Infarction and HD codes for Ischemic Heart Disease, which had 24-h cumulative ORs of 1.104 (0.915, 1.331) and 1.069 (0.983, 1.162), respectively. The odds of Diabetic AD and PA codes increased over time to a cumulative 24-h OR of 1.075 (1.001, 1.153) and 1.104 (1.015, 1.202) respectively.

CONCLUSIONS: We found increased PM2.5 during wildfire seasons was associated with some respiratory and cardiovascular outcomes within 1 h following exposure, and its association with diabetic outcomes increased over time. Cumulative effects were consistent with those reported elsewhere in the literature. These results warrant further investigation and may have implications for the appropriate time scale of public health actions. https://doi.org/10.1289/EHP5792

Introduction

Approximately 3% of the global land surface is burned by landscape fires every year, an area equivalent to nearly 20% of North America (Giglio et al. 2013). Over the past few decades, many areas of the world have reported longer wildfire seasons and more severe wildfire activity in terms of fire frequency, size, and intensity (Dennison et al. 2014; Jain et al. 2017; Jolly et al. 2015; Lucas et al. 2007). These trends are partially attributed to the increasing temperatures and more drought as the global climate changes, and projections suggest a continuation of these trends into the future (Aldersley et al. 2011; Barbero et al. 2015; Westerling et al. 2006; Wotton et al. 2017).

Smoke emitted from wildfires can affect large populations, even those distant from the fire, by degrading air quality at the local, regional, and global scales (Dempsey 2013; Dirksen et al. 2009; Jeong et al. 2008; Miller et al. 2011). Although wildfire smoke is a complex mixture of gases and particles, ambient concentrations of fine particulate matter (PM2.5) are the most widely used as a proxy for the mixture. The use of PM2.5 is based on a) concentrations being consistently elevated during wildfire smoke events, at locations both near and far from the fire (Naeher et al. 2007); b) well-established dose–response relationships with a wide range of health outcomes; and c) the availability of continuous measurements in many locations for the purposes of air quality regulation. The most recent estimates suggest that approximately 3.4 million deaths could be attributed to ambient air pollution in 2017 (Stanaway et al. 2018), and fire emissions accounted for up to 8% of these deaths (Lelieveld et al. 2015).

Exposure to wildfire smoke has also been associated with a wide range of acute cardiopulmonary morbidity (DeFlorio-Barker et al. 2019; Dennekamp et al. 2015; Haikerwal et al. 2015; Liu et al. 2015; Reid et al. 2016; Tinling et al. 2016), and evidence is emerging for other health conditions, such as adverse early life outcomes (Black et al. 2017; Holstius et al. 2012) and reduced diabetic control (Johnston et al. 2018). Although diabetes has not been comprehensively studied in response to short-term changes in PM2.5, these recent findings for wildfire smoke are consistent with other findings associating PM2.5 with increased risk of diabetic hospitalization (Zanobetti and Schwartz 2002; Zanobetti et al. 2014). In addition, evidence from studies on long-term exposure to PM2.5 exposure suggests an association with the development of type 2 diabetes (Bowen et al. 2018; Pearson et al. 2010; Rao et al. 2015; Thiering and Heinrich 2015). Finally, people with diabetes were more vulnerable to cardiovascular health effects associated with PM2.5 (Forastiere et al. 2008; Pinault et al. 2018; Zanobetti and Schwartz 2002). These findings warrant further investigation into the impact of wildfire smoke exposure on diabetic conditions.

So far, most epidemiological studies on wildfire smoke exposure have focused on 24-h average PM2.5 concentrations, associating same-day or previous-day exposures with acute health outcomes. Very few studies have examined the relationship between these acute health outcomes and sub-daily exposures, measured in 1-h rather than 24-h periods (Liu et al. 2015; Reid et al. 2016). Given that PM2.5 concentrations can vary considerably within 1 d due to changes in the weather and the intermittent nature of wildfire emissions (Said et al. 2015; Strand et al. 2011), there is uncertainty about the true nature of the exposure–response...
relationship during smoke episodes. First, it is not clear whether smoke exposure can trigger an acute outcome immediately, or whether there is a time lag between the exposure and the effect. Second, it is not clear whether the effects of smoke are driven by peak exposures within the day or by cumulative exposures throughout the day. Although PM$_{2.5}$ from wildfire smoke is the focus of this work, the same uncertainties apply to PM$_{2.5}$ from other sources. These questions are important to address for the development of effective public health response plans with the appropriate time sensitivity.

There are two key challenges in studying the health effects of sub-daily smoke exposures. The first, addressed in our previous study (Yao et al. 2018), is to generate spatially resolved PM$_{2.5}$ estimates at the 1-h time scale. The second is to identify population-based health outcomes recorded on a similar time scale. Because wildfire smoke events are usually sporadic, large populations and long time-series of data are needed to provide enough statistical power to detect their effects. This research is most feasible using routinely collected administrative health data, such as hospital admissions or medical billings. However, many of these databases do not have precise information on the exact time and location of the health events, and this information is necessary for studying sub-daily exposures.

Ambulance dispatches are a promising alternative to other types of administrative health data for assessing the sub-daily impacts of air pollution, including wildfire smoke. These databases typically have records of the exact location (in latitude and longitude) and time (in minutes and seconds) of the dispatch call. Although the spatial and temporal resolutions of ambulance dispatch data are ideal for studying sub-daily exposures, dispatch codes have uncertain diagnostic value in the absence of contextual information from trained medical personnel. As such, recent studies in Australia have found a general association between ambulance dispatch and wildfire smoke exposure (Dennekamp et al. 2015; Johnston et al. 2018; Salimi et al. 2017) but have not been able to examine a wide range of cause-specific dispatches at the sub-daily temporal scale.

We extend this prior work to North America and examine the association between sub-daily exposure to wildfire smoke and acute health outcomes by combining a previously developed exposure model with ambulance dispatch data that have been uniquely linked to subsequent paramedic reports and hospital admissions. This method allows more complete examination of the relationship between wildfire smoke and all ambulances dispatches, as well as those subsets most likely to be due to cardiovascular, respiratory, and diabetic conditions.

**Methods**

**Study Area and Study Period**

British Columbia (BC) is the westernmost province in Canada, with a population of approximately 5 million people in 2018, over half of which reside in the greater Vancouver area located in the southwestern corner. With more than 70% of land covered by forests (Austin et al. 2008), the province is prone to seasonal wildfires, and smoke from these fires is the dominant source of elevated PM$_{2.5}$ during the summer months (McKendry 2006). The health outcomes and exposure data for this study cover the wildfire seasons (1 April to 30 September) from 2010 to 2015, including three severe seasons in 2010, 2014, and 2015 with over 300,000 hectares burned in each of those years.

**Health Outcome Data**

We obtained data for all emergency ambulance dispatches during the study period from BC Emergency Health Services, which is the sole provider of ambulance and emergency health services across the province. The data included the date and time of the call, geographic coordinates of the event, and the reason for the call recorded as one of 33 codes (Table S1) assigned by the dispatcher using the Medical Priority Dispatch System (MPDS) (Clawson et al. 2015). The MPDS is a standardized set of protocols produced by the International Academies of Emergency Dispatch. Calls without a dispatch location or calls from callers who made more than four calls during the study period (5% of all unique callers) were excluded. The latter was done to minimize the occurrence of multiple calls within a short period of time, which may violate the assumptions of the case-crossover design. If there were multiple calls from the same caller within a 24-h period, only the first call was included in the analysis.

Each call in the dispatch database was provided with a linked patient care report as completed by the attending paramedics. Key information retrieved from these reports included the Personal Health Number (PHN, a lifetime unique identifier for health care in the province), the age and sex of the patient, and any assessment of medical conditions by the paramedics, assigned as one of the 18 Paramedic Impression (PI) codes (Table S2) (BC Emergency Health Services 2019). Although each patient had a care report, not all patients had an impression code because it is not a mandatory field for the paramedics to complete.

We also obtained hospital discharge data from the BC Ministry of Health (Canadian Institute for Health Information 2017), which included the date of hospital admission and the primary diagnosis, coded according to the International Classification of Diseases, 10th Revision (ICD-10). The primary diagnosis reflects the primary reason for the total length of the hospital stay and so may or may not reflect the initial reason for admission. Hospital diagnoses were linked to a dispatch call by PHN and included for the study whether a) the admission occurred within a 7-d period of the ambulance dispatch, and b) it was the admission closest to the date of the dispatch for cases where multiple admissions were found within the 7-d period.

Given this chain of data linkage, we could have had up to three measures of health outcome for each dispatch call: a) the MPDS code assigned by the dispatcher at the call center [Ambulance Dispatch (AD)]; b) the PI code assigned by the paramedics at the dispatched location [Paramedic Assessment (PA)]; and c) the primary ICD-10 code associated with the hospital admission record [Hospital Diagnosis (HD)]. These three measures have different advantages and disadvantages for the purposes of our study. The AD code was assigned to every single dispatch (no missing data), and it was assessed at the time closest to the onset of the emergency event. However, it was generally based on information self-reported by a lay caller and recorded as broad categories of health problems. The PA code was assigned by professionals with medical training after a physical examination the patient, but the assessment can be constrained by time, equipment, and demand and is not available on every record. The HD code provided the most robust medical assessment among the three, but it was available only for the most severe cases (i.e., those admitted), and it could have been made hours or even days after the initial call. Considering these different features, we decided to first provide a summary of the relationship among them to assess consistency and then to examine the dispatches related to cardiovascular, respiratory, and diabetic conditions, as identified by each of these three measures (Table 1).

**Exposure Assessment and Assignment**

Hourly exposures to PM$_{2.5}$ during the study period for all subjects were estimated with the 1-h Optimized Statistical Smoke Exposure Model (OSSEM-1h) previously developed and described elsewhere (Yao et al. 2018). Briefly, OSSEM-1h generates hourly
Table 1. Definitions and number of cases for each health outcome measure.

| Case groups | Definition | Number of cases |
|-------------|------------|----------------|
| All         |            | 676,401        |
| Cause-specific cases identified by Ambulance Dispatch (AD) codes based on the Medical Priority Dispatch System (MPDS)⁶ | | |
| Breathing problems | MPDS = 6 | 46,277 |
| Chest pain | MPDS = 10 | 51,996 |
| Arrest | MPDS = 9 | 3,527 |
| Stroke | MPDS = 28 | 21,173 |
| Heart problems | MPDS = 19 | 12,039 |
| Diabetic problems | MPDS = 13 | 5,987 |
| Other Codes | | 535,402 |
| Cause-specific cases identified by Paramedic Assessment (PA) codes based on Paramedic Impressions (PI)⁶ | | |
| Circulatory PI starts with | | 44,122 |
| Stroke PI = 0615 | | 17,495 |
| Myocardial infarction PI = 0860 | | 1,724 |
| Other circulatory | | 24,903 |
| Respiratory PI starts with 09 | | 23,392 |
| Asthma/COPD PI = 0930 | | 5,824 |
| Other respiratory | | 17,568 |
| Diabetic PI = 0305 or 0315 | | 4,722 |
| Hyperglycemia PI = 0305 | | 1,535 |
| Hypoglycemia PI = 0305 | | 3,187 |
| Other codes | | 346,910 |
| Cause-specific cases identified by Hospital Diagnosis (HD) codes based on the International Classification of Diseases, 10th Revision (ICD-10) | | |
| Circulatory ICD-10 = B90 to I99 | | 37,078 |
| Stroke ICD-10 = I60 to I69 | | 10,373 |
| Ischemic heart diseases ICD-10 = I20 to I25 | | 10,653 |
| Other circulatory | | 16,052 |
| Respiratory ICD-10 = J00 to J99 | | 22,038 |
| Asthma/COPD ICD-10 = J40 to J45 | | 9,084 |
| Lower respiratory infection ICD-10 = J13 to J22 | | 7,708 |
| Other respiratory | | 5,246 |
| Diabetic ICD-10 = E10 to E14 | | 2,921 |
| Other codes | | 114,246 |

⁶More about Medical Priority Dispatch System codes can be found in Clawson et al. 2015.
⁷More about PI codes can be found at BC Emergency Health Services 2019.

PM₂.₅ estimates across all populated areas of BC at a 5 km x 5 km spatial resolution using a random forests model, with input data including PM₂.₅ measurements from monitoring stations, fire activity observed by satellites, meteorology assimilated from observations and modeling, and geographic information. Compared with observations, model predictions had a correlation of 0.93 and a root mean squared error of 3.2 μg/m³. The intent was to model fire-related PM₂.₅ by using only data in fire seasons and incorporating fire-related predictors. Model training, however, necessarily relied on total PM₂.₅ measured at air-quality monitors; thus, its predictions do not strictly reflect PM₂.₅ from wildfire smoke alone. Exposure for each dispatch call was assigned based on the date and hour of the call, as well as the dispatch location (latitude and longitude) that was matched to the OSSEM-1h grid.

Statistical Analysis

A time-stratified case-crossover study design (Maclure 1991) was used to assess the association between dispatches and estimated PM₂.₅ exposure during wildfire seasons. Exposure during the case window was compared with exposures during a series of control windows. The case window was defined as the hour immediately before the ambulance was dispatched, and the control windows were defined as the same hour on the same day of the week in the same calendar month of the dispatch to control for day-of-week effects and seasonal trends. Control window exposures were assigned at the same location as the case window exposure. Using conditional logistic regression, individual factors that do not vary over a short time period (e.g., age, smoking status) can be controlled because the exposures during the case and control windows are compared in the same individual.

To examine the lag structure of the association between exposure and outcome, a distributed lag nonlinear model (DLNM) (Gasparrini et al. 2010) was used. This type of model can simultaneously describe complex exposure–response and lag–response relationships by combining the functions for both relationships in the same model. This approach has been applied in studies on the acute health effects of air pollution and ambient temperature (Buteau et al. 2018; Gasparrini et al. 2015; Guo et al. 2011; Guo 2017). We allowed for delayed relationship up to 48 h (lag 1–48 h) because most of previous studies using 24-h average exposures to wildfire smoke found the strongest association or best model fit at lags of 0–2 d (Borchers-Arriagada et al. 2019; Dellino et al. 2009; Elliott et al. 2013; Henderson et al. 2011; Johnston et al. 2007; Morgan et al. 2010; Youssouf et al. 2014). A natural cubic B-spline with 2 or 3 degrees of freedom (df), depending on the health outcome, was used for the lag–response relationship based on exploratory analyses to minimize the Akaike Information Criterion (AIC). Other functions including polynomials and penalized splines, with varying degrees of freedom, were also tested to describe the lag structure in the exploratory analyses, which produced similar results and less desirable model fit compared with cubic splines based on AIC. Both the lag-specific and cumulative odds ratios (ORs) were calculated to evaluate the time course and the overall association, respectively.

A linear exposure–response relationship was assumed in the analyses after preliminary evaluation of linear and nonlinear options found the linear models provided the best fit for most health outcomes (Figures S1–S3). This assumption also simplified the presentation of the results and allowed us to focus on the lag–response relationship. Models were adjusted for the same-day and previous-day maximum apparent temperatures from the nearest weather station maintained by Environment and Climate Change Canada, using a natural cubic B-spline with 3 df. All data preparation and statistical analyses were conducted using R software (version 3.5.1; R Development Core Team). The dlmn package was used to fit DLNM (Gasparrini 2011). Cox regression with Breslow ties was used to fit conditional logistic regression models, adopting the example code provided in a previous publication (Guo 2017). The study was approved by the Behavioural Research Ethics Board at the University of British Columbia (H15-02269).

Results

A total of 676,401 dispatch calls from 500,302 unique individuals were included in the study, among which all calls had an AD code; 444,189 (65.7%) had a PA code; and 244,101 (36.1%) were linked to HD codes (Figure 1). Paramedics arrived at the dispatched location within 1 h of the call in 99% of the cases, regardless of PA code group. Hospital admissions occurred within the same calendar day of the dispatch calls for 73% to 81% of cases, depending on the HD code group.

For each of the six AD code groups, 66%–73% of the calls also had a PA code (Figure 2A). For calls with the AD code Breathing Problems, the most prevalent (45.8%) PA code was related to respiratory conditions (Asthma/COPD and Other Respiratory in Figure 2A). Most of the calls with an AD code for Chest Pain, Heart
Problems, and Arrest (61.0%, 59.2%, and 60.0%, respectively) had PA codes related to circulatory conditions (Myocardial Infarction and Other Circulatory in Figure 2A). On the other hand, only 44.2% of calls with the AD code for Stroke had a PA code for Stroke, and 51.2% of calls with an AD code for Diabetic Problems had the PA code for Diabetic (Figure 2A).

For each of the six AD codes groups assessed, 24%–52% were linked to HD codes (Figure 2B). Compared with the linkage between AD and PA codes, the linkage between AD and HD codes had a larger proportion of cases in the Other Codes category (Figure 2C), indicating a somewhat weaker correspondence. The strongest relationship was observed between PA codes and HD codes (Figure 2C). For example, there were z cases with a PA code for Myocardial Infarction and a subsequent hospital admission, of which 76% also had a HD code for Ischemic Heart Disease, of which Myocardial Infarction was the primary subtype.

The mean (interquartile range) of PM2.5 exposures during the case and control windows in the 1-h window prior to dispatch were 5.5 (3.1, 6.5) μg/m³ and 5.4 (3.0, 6.4) μg/m³, respectively. The means of the maximum apparent temperature on the case days and control days were 17.6°C (13.5°C, 21.7°C) and 17.6°C (13.4°C, 21.7°C), respectively.

The lag–response relationship between PM2.5 exposures and the AD, PA, and HD codes varied by health outcome. There was a small increase in the odds of any AD [at lag 1 h, OR (95% CI) = 1.001 (0.999, 1.002)] within 1 h following increased PM2.5 exposure (Figure 3). An increase in odds was also observed for respiratory conditions identified by the AD code Breathing Problems [at lag 1 h, OR (95% CI) = 1.002 (0.999, 1.004)] and the PA codes Asthma/COPD [at lag 1 h, OR (95% CI) = 1.014 (1.001, 1.027)] and Respiratory [at lag 1 h, OR (95% CI) = 1.002 (0.999, 1.005)]. In all cases, the increase was largest at the 1-h lag interval. On the other hand, the ORs for respiratory outcomes identified by the HD codes Respiratory, Asthma/COPD, and Lower Respiratory Infection increased over time (Figure 4), and the maximum ORs at lag 48 h were 1.003 (0.999, 1.007), 1.006 (1.000, 1.012), and 1.005 (0.998, 1.013), respectively.

Switching to cardiovascular outcomes, the AD codes for Chest Pain and Heart Problems did not show any increase associated with PM2.5 exposure, but elevated odds were observed within 1 h after exposure in the PA code for Myocardial Infarction [at lag 1 h, OR (95% CI) = 1.027 (0.997, 1.057)] and in the HD codes for Ischemic Heart Disease [at lag 1 h, OR (95% CI) = 1.005 (0.999, 1.011)]. The OR for the Ambulance Dispatch code Arrest increased from negative to positive over time, with the largest OR observed at lag 48 h [1.007 (0.997, 1.013)] (Figure 5). The lag–response relationship for Stroke varied by outcome measure: An immediate increase in odds was observed in AD code, whereas the ORs increased from negative to positive over time for the PA and HD codes (Figure 5). Finally, the odds of Diabetic outcomes for the AD and PA codes both increased over time, and the OR became positive at approximately 24 h after the exposure, but the same was not observed for the HD codes (Figure 6).

When we stratified the analysis for the PA codes into Hypoglycemia and Hyperglycemia, the lag–response relationship was different for the two conditions (Figure S4).

Many outcomes reached their maximum cumulative ORs (95% CI) at a 48-h lag, including: 1.038 (1.009, 1.067) for the AD code Breathing Problems; 1.046 (0.995, 1.098) for the HD code Respiratory; 1.034 (0.948, 1.128) for Lower Respiratory Infection; 1.075 (1.001, 1.153) for the AD code Diabetic Problems; and 1.104 (1.015, 1.202) for the PA code Diabetic. On the other hand, the HD code Ischemic Heart Disease reached a cumulative maximum of 1.070 (0.983, 1.165) at a 28-h lag, and the PA codes for Myocardial Infarction and Asthma/COPD reached maximums of 1.189 (0.979, 1.442) at a 13-h lag and 1.106 (1.015, 1.205) at a 15-h lag, respectively (Figures 3–5).

**Discussion**

In this study, we found that a) cause-specific AD codes matched to subsequent PA and HD codes agreed reasonably well, providing more confidence in ambulance dispatches as a measure of health; b) exposure to elevated PM2.5 during wildfire seasons was associated with increased odds of dispatches related to respiratory and cardiovascular conditions, and the largest point estimates were observed in the hour immediately after the exposure; and c) exposure to elevated PM2.5 during wildfire seasons was also associated with dispatches related to diabetic conditions, with positive associations observed after a 24-h lag in exposure.

An important novelty of this study was the focus on the lag–response relationship at the hourly time scale, using a modern statistical method. Previous studies have examined such relationships by including the averaged exposure in specific lag hours as multiple independent variables in one regression model (Bhaskaran et al. 2011), or including exposure averaged over different lag windows in separate models (Enser et al. 2013; Evans et al. 2017; Gardner et al. 2014; Peters et al. 2001; Rosenthal et al. 2008; Sullivan et al. 2005; Wichmann et al. 2013). The biggest limitation of these methods was that they did not account for the correlation between exposures at different lags, whereas the distributed lag nonlinear model (DLNM) framework used in this study accounts for such correlation. As a result, the lag-specific effect estimates from this study may not be directly comparable with estimates from previous studies using conventional methods, and the following discussion will focus on comparison of the lag–response relationship and the cumulative effect estimates.

The associations between respiratory outcomes and PM2.5 estimates were consistent with previous reports. Studies using ambulance data in Australia found an association between daily PM2.5 and breathing problems [relative risk (RR) = 1.04, 95% CI: 1.02, 1.05, per 10 μg/m³ increase in PM2.5] (Salimi et al. 2017), as well as asthma/COPD calls (OR = 1.06; 95% CI: 1.01, 1.11, per 10 μg/m³ increase in PM2.5) (Johnston et al. 2018), similar to the 24-h and 48-h cumulative ORs we report. These cumulative ORs were also consistent with those estimated for respiratory medication dispensations (Elliott et al. 2013; Yao et al. 2016), physician visits (Yao et al. 2016), and hospital admissions (Henderson et al. 2011) from studies using daily PM2.5 measurements during wildfire events in the same region. Increased airway inflammation and decreased lung function have been observed in children with asthma and in the elderly immediately following exposure to ambient PM2.5, with lagged associations lasting from
5 to 12 h (Adamkiewicz et al. 2004; Delfino et al. 2006; Mar et al. 2005; Yamazaki et al. 2011). We also found an immediate increase in ambulance dispatches for respiratory codes following exposure, and the association declined over time.

Odds of myocardial infarction as measured by PA were elevated immediately following exposure, as were odds of ischemic heart disease as measured by HD. Neither result was reported by the previous Australian study (Johnston et al. 2018). However, a few studies using a similar case-crossover design found immediate associations for myocardial infarction following exposure to elevated ambient PM$_{2.5}$ (Evans et al. 2017; Gardner et al. 2014; Peters et al. 2001; Rosenthal et al. 2008). Controlled exposure studies in humans have also suggested there is an increase in subclinical indications of acute cardiovascular responses within hours of elevated exposure to PM$_{2.5}$, including increased cardiac arrhythmia, blood pressure, arterial stiffness, and thrombus
formation (He et al. 2011; Lucking et al. 2008; Lundbäck et al. 2009; Soppa et al. 2017; Urch et al. 2005).

Although previous studies have reported significant associations between out-of-hospital cardiac arrests and wildfire smoke exposure (Dennekamp et al. 2015; Salimi et al. 2017), we did not find the same for the AD code Arrest. Results from studies on ambient PM$_{2.5}$ and out-of-hospital cardiac arrests have also been inconsistent, where some found immediate associations following exposure (Pradeau et al. 2015; Rosenthal et al. 2008, 2013) and others found no association (Ensor et al. 2013; Raza et al. 2014; Wichmann et al. 2013). These inconsistencies could be due to different risk factors for cardiac arrest in different regions. For example, drug overdose was one of the major PA code for cardiac/respiratory arrest calls in this study, but not in the Australian study (Salimi et al. 2017).

Short-term exposure to PM$_{2.5}$ has been associated with insulin resistance (Dang et al. 2018; Haberzettl et al. 2016) and hospitalization for diabetes (Zanobetti and Schwartz 2002; Zanobetti et al. 2014), and long-term exposure has been associated with increased incidence and prevalence of type 2 diabetes (Eze et al. 2015; Pearson et al. 2010; Yang et al. 2018). On the other hand, the association between PM$_{2.5}$ and other chronic conditions, such as heart diseases, may be modified by diabetic comorbidity (Pinault et al. 2018; Zanobetti et al. 2014). The mechanism behind these associations remains inconclusive, but some studies suggest that it may be linked to oxidative stress and systematic inflammation induced by the exposure (Haberzettl et al. 2016; Sun et al. 2009). In this study, we found that the association between diabetic problems and wildfire smoke exposure increased over time and became positive at a 24-h lag. This finding adds to the limited evidence that has been available on the lag–response relationship, especially at the hourly time scale. When we modeled PA codes for hypoglycemia and hyperglycemia separately, the lag–response relationship was different (Figure S4). A recent study in Australia found that daily exposure...
PM$_{2.5}$ exposure was associated with increased same-day and next-day dispatches for hypoglycemia but not hyperglycemia (Johnston et al. 2018). These findings warrant further studies that examine the potential mechanisms behind the association between wild fire smoke and different diabetic conditions.

This study has some unique strengths. The temporal resolution of the exposure and ambulance dispatch data allowed the examination of the exposure–response lag structure on an hourly scale. The linkage among ambulance dispatches, paramedic assessments, and hospital admissions provided the opportunity to evaluate the quality of the data and the internal consistency of the study results. In addition, having a single provider of ambulance services in British Columbia enabled us to conduct a population-based study over a very large geographic area with variable wild fire smoke impacts.

There were also several limitations. First, exposure misclassification was possible due to a) error in the exposure model, b)
uncaptured variability within the 5 km × 5 km prediction grid, c) the assumption that each subject was exposed at the dispatch location during the control windows, and d) the potential lag time between exposure and the dispatch call. Second, some populations may be more likely than others to call ambulance services (Kerr et al. 2006; Rucker et al. 1997). With the case-crossover study design, we were able to control for the confounding effect of this factor, but the results might not be generalizable to the entire population. Third, there is still uncertainty about the temporal relationship between PM$_{2.5}$ exposure, symptom onset, ambulance call, and subsequent care for any given study subject. Although the ambulance dispatch data have finer temporal resolution than many other administrative data sets, there may still be a lag between the onset of symptoms and the action of calling an ambulance. Further, there was also a lag between some dispatches and hospital admissions, the reasons for which are unclear. In some cases, the subject may have been transported to a hospital and held under observation in the emergency room prior to admission, or admission was delayed due to limited availability of hospital beds. Unfortunately, the emergency room data were disparateely collected for each hospital and were not available through an integrated database for the study period. All of these uncertainties may affect the characterization of the lag structure. Finally, with our study design, we are not able to rule out the possibility that some of the increased ambulance calls may be due to the elevated stress level from the perception of fire or smoke, instead of air pollution, during wildfire smoke events.

This study adds to the limited evidence on the acute health effects from sub-daily exposure to PM$_{2.5}$, especially during wildfire seasons. We found associations with some respiratory and cardiovascular outcomes within 1 h of exposure, whereas the association with some diabetic increased over time and became positive at approximately 24 h after exposure. The results varied by whether the outcomes were coded for AD, PA, or HD. These results warrant further investigation into the health effects of sub-daily exposures and may have implications for the appropriate time scale of air quality standards and public health actions during air pollution events.

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