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After the wildfire: Indoor smoke residues and cleaning considerations

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Key Messages

Wildfire smoke and indoor environments

- Wildfire smoke pollutants can infiltrate indoor environments and deposit on surfaces, be absorbed into common household materials, or undergo chemical transformation on surfaces.
- Indoor surfaces and materials can act as reservoirs of settled dust and absorbed or adsorbed pollutants that can later be released back into indoor air as resuspended particulate matter or desorbed gases.
- Pollutants can persist for weeks to months after a fire.
- Two months after a wildfire, settled dust in smoke-impacted homes remains highly dominated by wildfire-generated particles if no cleaning or remediation occurs.

Pollutant classes identified indoors after wildfires

- Field and laboratory studies identified changes in the quantity and composition of particulate matter, polycyclic aromatic hydrocarbons (PAHs), volatile organic compounds (VOCs), and trace metals in indoor environments following wildfire smoke episodes.
- VOCs can be absorbed by indoor materials and subsequently be released back into indoor air over time.
- The median sum of 16 priority PAHs was 16% higher in dust from smoke-affected homes than from homes outside smoke-impacted areas.
- Trace metals associated with wildfire smoke have been detected in indoor dust; median heavy metal concentrations in smoke-affected dust were 15% higher six months post-wildfire compared to background samples.

Remediation and cleaning strategies

- Laboratory studies indicate that wiping, laundering, vacuuming, and mopping can reduce concentrations of PAHs and VOCs on surfaces and in indoor air; however, effectiveness varies by pollutant, surface type, and method.
- Cleaning smoke-exposed glass reduced PAHs by 71% (ethanol) and 62% (commercial cleaner); laundering cotton removed 48% of PAHs.
- Combined vacuuming, mopping, and dusting reduced airborne formic acid by 50%, formaldehyde by 32%, furan by 19%, and water-soluble organic carbon by 39%.
- HEPA portable air cleaners and ventilation may reduce VOC pollutant concentrations, but effects may be temporary and influenced by operational conditions.
- Evidence gaps remain regarding the effectiveness of cleaning strategies for settled dust and trace metals following episodes of wildfire smoke.





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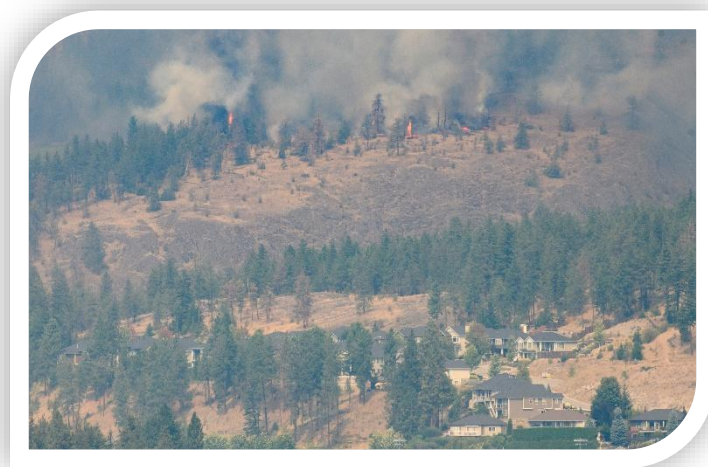


Introduction

As climate change continues to drive more intense and prolonged wildfire seasons, understanding how wildfire smoke impacts nearby communities is needed to help guide recommendations to reduce public health impacts. When wildfire smoke infiltrates indoor environments, pollutants can deposit on surfaces, be absorbed into common household materials, or undergo chemical transformation. These contaminants may persist for weeks to months, with household materials acting as reservoirs for pollutants that can be resuspended or desorbed back into indoor air long after a smoke episode has ended. While the infiltration of smoke during a wildfire event has been studied in some contexts,¹ even less is known about the range of pollutants that remain indoors after a smoke event, the health implications, and effective strategies to reduce ongoing exposures.

The aim of this review is to synthesize evidence available from studies examining the deposition, accumulation, persistence, and resuspension of wildfire smoke pollutants in indoor environments, as well as the effectiveness of cleaning and ventilation strategies. This review examines findings from a combination of field studies conducted after wildfire smoke episodes and experimental studies in a laboratory setting. Lastly, this review identifies key evidence gaps related to remediation effectiveness in real-world residential settings.

In some cases, wildfires can encroach into the built environment causing structural fires (i.e., wildland urban interface fires). The fire damage to homes and impacts of extreme heat in these scenarios is beyond the scope of this review. Damage to structural, electrical, water, and gas systems can cause serious health risks that should be assessed and remediated by qualified professionals.



Methodology

Literature search

A literature search was conducted using variants and Boolean operator combinations of key search terms such as “wildfire,” “smoke,” “combustion,” “prescribed burn,” “biomass burning,” “clean,” “decontaminate,” “mitigation,” “vacuum,” “remove,” and “wash” (a full list of search terms is available upon request). Although the focus of the search strategy was to assemble studies specific to wildfire smoke, the studies also included the impacts of interface fires where combustion of anthropogenic fuels contributed to the overall smoke plumes.

The following databases were searched: MEDLINE, Embase, CINAHL, and Environment Complete. Bibliographies and citations of key articles were used to retrieve additional literature via forward and backward chaining, along with supplemental searches as necessary. English documents were considered that were published over the previous 22 years (2004–2026). The titles and abstracts of results were screened by one reviewer, and the full texts of results meeting inclusion criteria were screened by a single reviewer. Exclusions included sources published ahead of print, before peer review, and mathematical modelling studies that exclusively used estimated data. Forward and backward chaining of key studies identified additional results providing evidence on the composition of post-smoke episode residues indoors and the influence of different cleaning techniques (e.g., wet wiping, mopping, washing, and HEPA vacuuming) on indoor pollutant concentrations.

The quality of included evidence was evaluated using critical appraisal tools, as indicated by the study design below. Completed quality assessments for each included study are available on request.

Study Design	Critical Appraisal Tool
Randomized Controlled Trial	JBIChecklist for Randomized Controlled Trials
Quasi-experimental	JBIChecklist for Quasi-Experimental Studies
Analytical Cross Sectional	JBIChecklist for Analytical Cross Sectional Studies

For each study, all available data relevant to the research question were extracted by one reviewer, including study design, setting, location, population characteristics, interventions, and outcomes. The results were synthesized narratively due to the variation in methodology and outcomes for the included studies. Table 1 provides a summary of the inclusion and exclusion criteria for studies.

Table 1. Inclusion and exclusion criteria

	Inclusion	Exclusion
Setting	Residential environments, clean air spaces, public/institutional buildings (e.g., schools, daycares, malls, libraries, community centres, healthcare centres, long-term care centres, etc.), vehicles	Industrial workplaces
Context	During combustion-derived air pollution episodes that may be caused by fire (wildland, coal mine fires, peat fires, interface fires, landscape fires, agricultural fires, prescribed burns, industrial fires, landfill fires, tire fires, any multi-day structural fires (e.g., 9/11)	During non-combustion derived episodes, air pollution events exclusively due to engine exhaust, traffic pollution, coal combustion, etc. (coal mine fires were included)
Intervention	Cleaning after a combustion-derived air pollution episode as defined above	
Comparator	Any or no comparator; pre- vs. post-cleaning	
Outcomes	Concentrations of airborne pollutants and pollutants in settled dust on surfaces, films, furnace filters	
Study designs	Before-and-after studies, case studies, laboratory studies, grey literature (including technical reports from standard setting bodies, NRC/NIST, etc.)	Modelling studies
Publication date	Published in last 22 years	

There are additional factors to consider when returning to a property that has been evacuated if the building or surrounding land has been directly damaged by fire. Excess heat can damage electrical systems, home appliances, and drinking water systems, potentially causing fire risks, gas leaks, and water damage and the potential for mould growth. In older homes, hazardous materials such as asbestos and leaded paint can become exposed following a fire, posing a risk to residents. In these situations, qualified professionals specializing in remediation of hazardous materials should be consulted. These issues are beyond the scope of this review; however, additional resources are available from professional organizations, such as the American National Standards Institute (ANSI)/ Institute of Inspection, Cleaning and Restoration Certification (IICRC) [Standard for professional fire and smoke damage restoration S700-2025](#) and associated [Technical Guide](#).^{2,3}



Background

Combustion-derived air pollutants (i.e., smoke)

Wildfire smoke is a complex mixture of particulate matter (PM), inorganic gases and organic gases, and trace metals.⁴ The PM component consists of a mixture of organic carbon and trace metals and is often reported based on the particle size fractions. For example, PM₁₀, PM_{2.5}, and PM₁ consists of particles less than or equal to 10, 2.5, and 1 µm in diameter, respectively. For gases, inorganic gases include carbon monoxide (CO), carbon dioxide (CO₂), and nitrogen dioxide (NO₂), and organic gases such as volatile organic compounds (VOCs) and semi-volatile organic compounds (SVOC), which include polycyclic aromatic hydrocarbons (PAHs). The mixture of pollutants in wildfire smoke can also change over time, with PAHs transitioning between being PM-bound and airborne. Though reactions in the atmosphere VOCs can be transformed into gases like ozone (O₃) and secondary organic aerosols that can condense on existing particles or form new particles.

Compared with other types of air pollution, wildfire smoke often has higher levels of VOCs such as benzene, acrolein, and formaldehyde; PAHs such as acenaphthylene, fluorene, and phenanthrene; and trace metals such as magnesium and potassium.⁵⁻⁷ However, pollutant concentrations can change depending on atmospheric conditions and the types of materials burnt. Smoke from biomass burning can be differentiated from other types of air pollution by the presence of chemically stable organic marker compounds such as levoglucosan, which is released in large quantities when wood is combusted. Wildfire smoke also has smaller particles than urban air pollution.⁸ For instance, air sampling during a wildfire period in California revealed that the mean fraction of fine PM (PM₁/PM_{2.5}) was significantly higher during the wildfire period (0.80) compared to the non-wildfire period (0.52).⁴



When wildfire smoke infiltrates indoors, it can contribute to indoor dust as well as absorb onto interior surfaces (Table 2). Indoor dust is a complex mixture of organic and inorganic particles stemming from outdoor and indoor sources. Outdoor sources include soil, pollen, and ambient air pollution. Indoor sources include pet dander, human skin cells, dust mites, and combustion particles from various activities such as cooking, burning candles, and tobacco smoke.⁹ Occupation can also play a role in dust composition. For example, the dust that is resuspended indoors from clothing would be different for an office worker compared with a construction worker. Depending on humidity, particle size, and the type of surface, wildfire smoke PM can contribute to house dust by settling on surfaces (e.g., flooring, walls, worktops, and furniture), contributing to indoor dust. Similarly, gaseous wildfire smoke pollutants can absorb onto surfaces and into household materials (e.g., fabrics) or undergo chemical transformation and condense onto settled particles.^{10,11}

Table 2. Overview of indoor wildfire smoke pollutants and measurement techniques^{12,13}

Pollutant	Definition	Measurement technique
Airborne particles	Particles suspended in air	– Air sampling and analysis or direct-reading monitors – Methods include gravimetric analysis, optical particle counters, beta attenuation monitors (BAM), or tapered element oscillating microbalance (TEOM)
Airborne gases	Inorganic and organic compounds in the gas phase	– Air sampling and laboratory analysis – Direct-reading instruments, for inorganic gases such as CO – Laboratory analytical methods mostly used for organic compounds and include gas chromatography and or spectroscopy (e.g., Time-of-Flight Chemical Ionization Mass Spectrometers)
Settled or deposited pollutants	Particles or pollutants that have agglomerated into larger particles that settle onto surfaces	– Laboratory analysis of sampling wipes – Laboratory analytical methods include extracting pollutants and determining chemical profiles by gas or liquid chromatography and or spectroscopy
Adsorbed pollutants	Pollutants that have settled and adhered onto surfaces	– Indirect measurements via air sampling for organic gases or direct-reading monitors for inorganic gases in a sealed environment into which the pollutants may desorb – Laboratory analytical methods include analysis of air samples and pollutants extracted from wipes by gas or liquid chromatography and or spectroscopy
Absorbed pollutants	Pollutants that have penetrated into materials	– Direct analysis of material or indirect measurements via examining changes in localized airborne/gas concentrations – Laboratory analytical methods for direct measurements include thermal desorption and analysis using gas chromatography-mass spectrometry

After a wildfire smoke episode, contaminated surfaces and materials can act as air pollutant reservoirs, releasing pollutants back into the indoor air by resuspension of deposited particles or desorption of gaseous pollutants. These pollutants can also change over time through reaction with atmospheric oxidants or cleaning chemicals such as bleach, creating different pollutants.^{14,15} Without remediation or cleaning, reservoirs of pollutants can persist in the indoor environment, presenting an ongoing health risk long after the wildfire smoke clears from the outdoor air. For example, residents of homes damaged by smoke or ash have reported experiencing dry cough, itchy or watery eyes, and sneezing for at least one year after a major smoke episode.¹⁶ In these scenarios, uptake of pollutants by the body can occur via inhalation and dermal exposure, as well as ingestion

of dust that settles on food, food preparation surfaces, or skin (e.g., hand-to-mouth behaviours by toddlers). However, the exact concentrations and types of wildfire smoke-related pollutants likely to affect indoor environments is poorly understood, as are the health implications of long-term exposures indoors.

Remediation of indoor smoke contaminants

Remediation of wildfire smoke contaminants after a wildfire can be referred to as restoration or decontamination and includes all activities or strategies used to remove smoke pollutants. Several agencies provide guidance to homeowners and businesses on returning to a premise safely and on cleanup following a smoke episode.^{3,17-26}

Current guidance broadly includes:

- Advice on personal protective equipment (PPE) – matching equipment to the expected hazards. This may include using a respirator, gloves, and protective clothing.
- Advice on cleaning – using wet mopping and wiping with a mild diluted detergent (avoid dry wiping and sweeping because this can resuspend particles), and HEPA-filtered vacuuming.
- Advice on physical removal of airborne pollutants via ventilation and/or filtration.
- Advice on odour management – using activated charcoal or similar odour absorbent media to remove VOCs, and professional use of ozone generators (see ozone generators in Box 1 below).
- Advice on removal of contaminated material – such as replacing furniture and/or clothing.

Box 1. Ozone generators and odour management

The use of ozone generators is not recommended in occupied spaces because ozone and by-products of ozone oxidation can be hazardous to human and animal health. In some cases, ozone can also degrade materials such as textiles and plastics. Health Canada currently recommends against the use of personal ozone generators in homes. In addition, as of 2025, there is no CSA electrical testing standards for these devices, meaning that they may not be compliant with the Canadian Electrical Code, representing a potential safety hazard.



The use of ozone generating machines to remove smoke odours in homes is, however, common practice by some professional restoration or remediation companies, and frequently mentioned in American remediation guidance. International guidance on the use of these machines can be found in a technical document on wildfire smoke restoration from the American Industrial Hygiene Association.²⁷

For some premises, repeated cleaning cycles may be required for pollutants that are more difficult to remove. Specialist smoke damage restoration and cleaning contractors may use different approaches for severely impacted premises.



Results

Overview of studies

This review identified a total of 15 studies of moderate to high quality, including seven quasi-experimental studies, and eight analytical cross-sectional studies (Appendix A). Eight studies investigated wildfire smoke episodes, and six studies used laboratory experimental smoke exposure systems to examine indoor air and/or surface pollutant concentrations post-smoke episode or exposure. The pollutants examined in these studies included settled dust and particulate matter (n = 4), VOCs (n = 8), PAHs (n = 5), and trace metals (n = 5). Experimental laboratory studies examined the effectiveness of cleaning methods on reducing airborne VOCs (n = 4) and PAHs (n = 1) that have been deposited and then released from common indoor surfaces such as glass and cloth. One post-wildfire smoke study examined the use of air cleaning and ventilation on airborne VOCs.

What pollutants are found on indoor surfaces post-wildfire smoke episodes?

In this review, several classes of pollutants were identified in the indoor environment following wildfire-smoke episodes. The deposition, accumulation, and persistence of these pollutants were described in a combination of field community-based and experimental test-chamber or house studies (Table 3). Observations from these studies are described in the following subsections by pollutant type.

Particulate matter (PM)

The PM component of smoke contains VOCs, PAHs, and trace metals along with elemental carbon and organic compounds in solid particles and liquid droplets. Specific sub-classes of pollutants that may be present in PM will be covered in the following sub-sections. Wildfire-related PM may also be referred to as ash, soot, and char. Soot and char are produced by more complete combustion and consist of mostly carbon. Ash consists of residue containing inorganic compounds, minerals, and charred organic components after the combustion of carbon-based compounds.^{28,29}

Wildfire-smoke PM can be airborne or deposit onto surfaces, contributing to settled dust. After settling, dust on indoor surfaces can be resuspended into the air by activities (e.g., vacuuming) or changes in ventilation (e.g., fans). Using polarized light microscopy to look for char, soot, and ash, Spurgeon et al. examined surface samples from 343 residences in northern California after 22 wildfires that occurred from 2017 to 2022.³⁰ In total, surface char was detected in 57% (n = 196) of houses, ash in 8.8% (n = 30) of houses, and soot in 1.2% (n = 4) of houses. The deposition of smoke residue was also found to vary indoors depending on location within the home, with char detected



more often on windowsills versus interior hard surfaces (2.8 times more likely to be detected). In another study, Silberstein et al. examined dust samples collected in smoke-impacted residences in Boulder, Colorado, up to two months after the 2021 Marshall wildland urban interface fire. Prior to cleaning, dust samples contained elevated levels of the biomass combustion marker levoglucosan (median = 4147 ng/g, n = 4) compared to samples from non-smoke-affected homes (median = 256 ng/g, n = 4), indicating that wildfire smoke substantially contributed to dust loading.³¹ Levoglucosan is an important marker of biomass burning because it indicates other common wildfire contaminants of concern are likely to be present.

Silberstein et al. also demonstrated the capacity for resuspension of settled wildfire smoke in dust. When sampled houses were unoccupied, airborne mass concentrations of PM_{2.5} and PM₁₀ were very low (approximately 0–2 µm/m³). However, resuspension of particles was observed during research and cleaning activities, with levels spiking to 8 µm/m³ for PM_{2.5} and 120 µm/m³ for PM₁₀. Spikes in PM_{2.5} and PM₁₀ also occur during normal indoor activity. For example, maximum PM_{2.5} mass concentrations of 28 µm/m³ have been reported for non-smoking homes in Regina, Saskatchewan. Even so, the chemical composition of resuspended dust containing wildfire smoke PM likely differs from that of typical dust,⁴ with higher concentrations of heavy metals and organic compounds (see following subsections), which may result in different health effects on those exposed.

A similar study by Goftari et al. examined PM mass concentrations in 19 homes in Los Angeles County, California, approximately two months after the Palisades and Eaton wildland urban interface fires.³² Indoor PM_{2.5} and PM₁₀ concentrations of 3.45 and 31.66 µg/m³, respectively, were reported. In addition, the average indoor/outdoor (I/O) ratios for PM_{2.5} and PM₁₀ were 1.08 and 2.99, respectively, indicating strong indoor sources of PM. This is notable as only three of the 19 homes were reoccupied at the time of testing, suggesting that the indoor PM is not likely due to occupant activities, but rather infiltrated smoke. Furthermore, the authors asked homeowners about smoke mitigation strategies used during the fire, finding 73% of PM₁₀ and 86% of PM_{2.5} levels could be explained by proximity to the fire, absence of air purifiers, use of non-HEPA vacuums, and open windows. It is also important to note that all the homes in this study suffered at least some level of structural damage from the fire.

Some pollutants in wildfire smoke can undergo physical and chemical transformation after depositing on surfaces. Farmer et al. examined PM generated from combusting Ponderosa pine chips in a smoker. They reported that PM deposited onto glass surfaces in a test house had two distinct morphologies: aggregate-like and smaller rounded. These two morphology types also had differing chemical compositions.³³ Although specific chemical characterization was not performed, atomic force microscopy-infrared spectroscopy showed that the aggregate particles contained more compounds with esters, aliphatic carbon chains, and carboxylic acids. The rounded particles contained more carbon-carbon, aromatic, and aldehyde functional groups common to molecules such as levoglucosan. Farmer et al. did not indicate in this study whether these differences in size

and morphology may cause differences in the toxicity profile and/or the ability to remediate these particles with cleaning strategies. However, aggregate-like particles in general may be more toxic.³⁴

PAHs

Three studies were identified that examined PAH concentrations in house dust post-wildfire episodes. In Silberstein et al., the concentration of 16 priority PAHs (according to the US Environmental Protection Agency, US EPA) were measured in settled dust that was sampled up to two months post-wildfire in smoke-affected homes (n = 4).³¹ Smoke-affected homes were defined as those within the 2021 Marshall fire perimeter set by the fire management agencies. Background homes (n = 4) were selected from both upwind and downwind of the burned area at distances varying from approximately 1 to 15 km. The median summed concentration of the 16 PAHs in smoke-affected homes was higher than in background homes (1859.3 ng/g vs. 1607.5 ng/g, statistical significance not evaluated). However, measurements in both smoke-impacted and background homes were lower than background values reported in other American cities,³⁵ and similar to levels found in urban soil samples.³⁶ Similarly, the authors noted the median sum of the 16 PAHs in Colorado was less than background levels reported for Kingston, Ontario, where the median sum of 12 PAHs in indoor dust was 2050 ng/g.³⁷ All 12 PAHs analyzed in this study were included in the 16 priority PAHs identified by the US EPA.

Smoke-affected dust samples in the Silberstein et al. study had greater proportions of light PAHs (two-, three-, or four-ring structures) such as fluoranthene and pyrene. Among heavy PAHs (more than three rings), the median total concentration of seven PAHs designated as carcinogenic by the US EPA was lower in the smoke-affected homes compared with background homes (380.1 ng/g vs. 451.1 ng/g). When comparing the concentrations of the known toxic and carcinogenic PAH benzo[a]pyrene, median values were lower in both smoke-affected homes (67.8 ng/g) and background homes (94.7 ng/g) compared with dust found in homes in Kingston, ON (221 ng/g).³⁷ However, no human health risk assessment was performed in Silberstein et al., and comparable levels of PAHs across studies does not preclude the potential for adverse health effects.

A parallel study by Dresser et al. also examined a home in the area burned during the 2021 Marshall fire. It reported the mixing ratios of airborne PAHs decreased significantly over the course of serial air sampling 10 to 35 days after the fire.³⁸ Moreover, the authors observed a common trend for the PAHs naphthalene, indene, acenaphthalene, and dimethylnaphthalene. Over 60% of the total reductions observed between days 10 and 35 occurred between days 10 and 15, meaning the rate of decrease slowed between days 15 and 35. Dresser et al. also compared the gas-phase profile of PAHs detected with the PAHs detected in dust samples by Silberstein et al., and found that lower volatility PAH species were less present in gas samples than in dust samples. They hypothesized this was due to lower volatility PAH species being removed from the gas phase through surface interactions.



Kohl et al. reported on PAH concentrations in dust samples collected from residential homes (n = 64) in Fort McMurray, Alberta after the 2016 Horse River wildfire.³⁹ The median sum concentration of all 14 PAHs measured in this study was 915 ng/g (range: 415 to 31,200 ng/g), which is tenfold lower than values reported for house dust samples collected in unimpacted Ottawa homes. Samples in Fort McMurray were collected 14 months after the wildfire, which may have limited applicability for understanding the persistence of PAHs in smoke-affected dust samples. Moreover, the authors note that the areas of the houses that were sampled were areas that are typically cleaned often (e.g., bedrooms), suggesting that cleaning may have removed most of the PAHs. There are currently no guidelines for PAH concentrations in residential house dust in Canada, but the authors noted that naphthalene, phenanthrene, anthracene, benz[a]anthracene, and fluoranthene concentrations exceeded the Alberta Tier 1 Soil and Groundwater Remediation guidelines for residential soils.⁴⁰

In a study of 39 homes distributed along a gradient of smoke exposure from a coal mine fire ignited by a nearby wildfire in the Latrobe Valley, Australia, Wheeler et al. reported elevated concentrations of PAHs in attic dust.⁴¹ The most abundant PAHs, phenanthrene, naphthalene, 2-methylnaphthalene, fluoranthene and pyrene, were all found to have average concentrations above 500 ng/g. The average sum concentration of the 16 priority PAHs (EPA) was 4,010 ng/g (range: 520–48,900 ng/g), with a trend for increasing concentration as a function of distance from the coal mine fire.

Wildfire-derived PAHs can also adsorb or absorb on glass, textiles, and HVAC filters as demonstrated by Laguerre et al. in laboratory chamber experiments using smoke generated by burning Douglas Fir and Western Red Cedar needles.⁴² These materials acted as reservoirs for PAHs, with glass surfaces and bulk materials (cotton and filter samples) having elevated concentrations for up to 40 of the 133 day study period. After 37 days, PAH levels decreased by 88% for glass, 81% for cotton, and 74% for the filter material. PAH decay rates were also calculated for these materials using experiments spiked with standard concentrations. The total mean $\pm$ standard deviation PAH decay rates were 0.025 ± 0.005 , 0.051 ± 0.009 , and $0.010 \pm 0.002 \text{ day}^{-1}$ for glass, cotton, and mechanical air filter media, respectively. The values indicate that PAHs persisted for longer on filters than glass or cotton material, with the most persistent filter PAH being phenanthrene. However, the authors did not specify whether the reduction in PAHs over time was due to desorption back into the air or decomposition.

VOCs

Two studies on smoke-derived VOCs found that they adsorb/absorb and desorb from indoor surfaces. In an experimental test house, Li et al. demonstrated that VOCs generated by burning pinewood chips were either absorbed by surface reservoirs, escaped outdoors, or remained in a gas-phase indoors.⁴³ Following a smoke injection, concentrations of PM_{1.0} peaked at 100 to 300 $\mu\text{g}/\text{m}^3$ and total gas-phase water-soluble organic carbon (WSOCg) peaked at approximately 220 μg carbon/ m^3 . During this period, airborne VOC levels initially declined as they adsorbed/absorbed onto materials but stabilized after several hours when surface material reservoirs desorbed, or

released, VOCs back into the air, with ventilation controlling long-term overall airborne VOC concentrations. For example, surfaces became a net source of acrolein emissions (a known lung irritant) at 4.8 hours.⁴⁴ Most smoke-related VOCs had enough time to penetrate materials in the test house and reach an equilibrium state before being predominantly vented out of the house. Furthermore, organic molecules with higher molecular weights tended to decay faster, but the investigators indicated that it is possible for VOCs to persist for weeks to months in houses repeatedly exposed to wood smoke because VOCs can diffuse deeply into permeable materials (material type not specified). One important caveat to this study was that the major surfaces in the test house were painted walls (42.1%), wooden floors (11.2%), and painted ceilings (9.7%), with no soft flooring such as carpets.

In another test-house study by Link et al., concentrations of VOCs were measured until 100 days after exposing the house to smoke.⁴⁵ Because the aim of this study was to examine the impact of wildland urban interface fires, a mixture of non-combustible gypsum, spruce-pine fir, polyurethane rubber, polyvinyl chloride (PVC), and acrylonitrile-butadiene-styrene plastic was burned to produce smoke that was blown into the house 11 times over 55 days. Air sampling 35 days after the last burn detected significantly decreased concentrations of most VOCs measured compared with two days after the burn period. However, the VOCs benzofuran, naphthalene, benzonitrile, C12 aromatics, C₉H₁₀, C₁₀H₈O, and C₉H₇N were still detected at five times higher concentrations compared with background levels, even though the levels of these VOCs had reduced by over 75% of their maximum detected concentrations on day 2. By 98 days after the burn period, benzofuran, C₁₂H₁₀, C₉H₁₀, and C₉H₇N were still being detected at much higher concentrations than background levels, indicating the presence of significant reservoirs of smoke pollutants in the home.

Persistence of VOCs was also observed by Dresser et al., who examined gas-phase VOC levels in a home from 10 to 35 days after the 2021 Marshall fire in Colorado.³⁸ Relatively high concentrations of VOCs were observed in initial samples taken at day 10, with concentrations decreasing by approximately 80% by day 35. Similar to the PAH data, most of the decrease occurred between days 10 and 15. For example, benzene levels decreased from approximately three parts per billion by volume (ppbv) on day 10 to one ppbv on day 15, and further decreased to 0.6 ppbv by day 35.

In three other studies by Goftari et al., Yao et al., and Pei et al., VOCs were identified in homes following the 2025 Los Angeles Palisades and Eaton wildland urban interface fires in California.^{32,46,47} In Goftari et al., total VOCs were measurable in three of 19 homes, with average values of 42.5, 22.6, and 20.8 ppb.³² Yao et al. examined 22 airborne VOCs in 22 homes across the active burning and post-fire period, reporting increased I/O ratios during the postfire period.⁴⁶ Specifically, significantly higher ratios ($p < 0.05$) were observed for *m,p*-xylene, carbon tetrachloride, and heptane. Benzene was not significant, but an increasing trend of median I/O ratios was noted at 0.94 during the fire, 1.01 during fire containment, and 1.41 postfire.



Pei et al. examined 24 VOCs in 50 homes in the two months following the Los Angeles fires.⁴⁷ Concentrations of airborne benzene, trichloroethylene, ethylbenzene, perchloroethylene, and ethyl acetate in homes were found to exceed EPA cancer risk screening levels in 42, 14, 6, 4, 2% of homes sampled, respectively. However, in a follow up analysis, many of these VOCs were found to be increased in homes that had been cleaned and may be due to other indoor sources of VOCs such as cleaning supplies. Specifically, median concentrations of cleaning related VOCs, acetone, ethyl acetate, ethylbenzene, heptane, *n*-hexane, perchloroethylene, toluene, and xylenes were 51–227% greater in cleaned homes. When considering home distance from the burn area as a proxy for the intensity of smoke impacts, the authors reported a trend for median indoor concentrations of naphthalene increasing with proximity from the burn area. However, naphthalene concentrations did not exceed EPA cancer screening levels. Benzene was found to exceed EPA levels in the greatest proportion of homes in this study (42%), but median indoor concentrations largely increased with distance from the fires, which the authors indicated may be due to other outdoor sources such as vehicle emissions.

Primary VOCs can also undergo chemical transformation into semi-volatile organic compounds (SVOCs) after depositing on surfaces. Deeleepojananan et al. showed that VOCs were likely to undergo chemical transformation on kaolinite (e.g., clay) surfaces but not on glass or rutile (a mineral commonly used as a pigment in paints) surfaces in laboratory chamber experiments.⁴⁸ For instance, the VOC diethyl 3-hydroxyglutarate was detected on kaolinite but not on glass or rutile, indicating that some materials facilitate chemical reactions that may cause secondary pollutants that have the potential to be released back into indoor air. Overall, the most abundant compounds identified on glass samples were levoglucosan and 3-methyl-1,2,3-butanetricarboxylic acid (MBTCA). These are both tracers of biomass smoke, and indicators that smoke VOCs adsorbed to glass surfaces.

Trace metals

Wildfire smoke can also deposit trace metals indoors, mostly as a constituent of PM. Silberstein et al. reported a moderately elevated median sum concentration of 20 heavy metals in house dust from four smoke-affected homes compared with four background homes two months after the 2021 Marshall fire (34.6 mg/g vs. 30.1 mg/g, respectively).³¹ Both smoke-affected and background dust samples contained elevated levels of zinc (Zn), arsenic (As), cobalt (Co), chromium (Cr), and copper (Cu) when compared with ambient levels in Colorado soils; however, the authors indicate the increases in dust samples may be due to sources other than wildfire smoke.

Dingle et al. also reported elevated concentrations of trace metals in indoor dust post-wildfire exposure.⁴⁹ In homes in Fort McMurray (n = 125) that were severely impacted by the 2016 Horse River interface wildfire, aluminium (Al), titanium (Ti), vanadium (V), As, thallium (Tl) concentrations were all significantly higher ($p < 0.05$) compared with less affected neighbourhoods up to two years after the wildfire. The analyses accounted for those who lived in the home and may have had

occupational exposure to metals. In neighbourhoods where buildings burned during the fire, concentrations of As, and V were 62% and 37% higher in settled dust samples, respectively, compared with less affected neighbourhoods. However, all measured concentrations were below Alberta Teir 1 guidelines for soil.⁴⁰ Another study examining house dust samples (n = 64) in Fort McMurry after the same wildfire by Kohl et al. found that all 13 metals analysed remained within Alberta regulatory guidelines with the exceptions of Zn and Cu.^{39,40}

In Goftari et al., surface and airborne concentrations of trace metals were measured in 19 homes following the Los Angeles County wildfires.³² The most abundant airborne metals were Al, iron (Fe), and magnesium (Mg), with reported concentrations (mean $\pm$ SD) of 223.8 ± 153.9 , 61.9 ± 52.3 , and 305.1 ± 168.7 ng/m³, respectively. Analysis of surface wipe samples from a combination of surfaces, including carpets, tables, floors, countertops, and windowsills revealed potassium (K), Mg, and Al to be the most abundant metals, with concentrations (mean $\pm$ SD) of 24061.6 ± 32875.1 , 21475.1 ± 35706 , and 19628.0 ± 26453.4 μ g/m², respectively. Increased levels of Pb were also observed in this study, with levels exceeding EPA exposure guidelines for floors and surfaces of 53.8 μ g/m² in 12 homes. However, the authors noted that most of the homes were built prior to 1978 and may contain lead paint.

When examining attic dust trace metals in 39 homes exposed to smoke from a coal mine fire in the Latrobe Valley, Australia, Wheeler et al. reported concentrations of Al, barium (Ba), calcium (Ca), Co, Fe, Mg, manganese (Mn), and sulfur (S) were all elevated at least ten times relative to background levels.⁴¹ Of these elements, Ba and Mg showed the strongest association with distance to the mine fire. By abundance, Ca, Fe, and Al were the most prominent, with mean concentrations of 30.8, 24.5, and 17 mg/g. Pb was also elevated (0.423 mg/g), but tended to correlate with older homes (i.e., increased exposure to lead paint and gasoline) and was not elevated in the smoke emitted by the mine fire.

Table 3. Summary of post-wildfire studies examining indoor air and surface pollutants in smoke-impacted homes

Reference	Setting	Key findings
Goftari et al. 2026 ³²	Los Angeles County, California (Palisades and Eaton Fires); residential homes (n = 19)	- Indoor PM_{2.5} and PM₁₀ concentrations averaged 3.45 and 31.66 µg/m³, respectively - Al, Fe, and Mg were the most abundant airborne metals with mean ± SD of 223.8 ± 153.9, 61.9 ± 52.3, and 305.1 ± 168.7 ng/m³, respectively - K, Mg, Al were the most abundant metals detected on surfaces with mean ± SD of 24061.6 ± 32875.1, 21475.1 ± 35706, 19628.0 ± 26453.4 µg/m² - Pb exceeded EPA guidelines for floors of surfaces of 53.8 µg/m² in 12 homes
Pei et al. 2026 ⁴⁷	Los Angeles County, California (Palisades and Eaton Fires); residential homes (n = 50, 18 homes within burn area, 25 adjacent, and seven more than three km away)	- Concentrations for five of 24 indoor VOCs measured in homes exceeded EPA cancer risk screening levels; benzene, ethyl acetate, perchloroethylene, trichloroethylene and ethylbenzene - Naphthalene median indoor concentrations increased in homes closer to the burn area (burn area: 0.20 µg/m³; adjacent: 0.14 µg/m³; far: 0.05 µg/m³) - VOCs commonly present in household cleaning products were elevated by 51–227% in cleaned homes compared to uncleaned
Yao et al. 2026 ⁴⁶	Los Angeles County, California (Palisades and Eaton Fires); residential homes (n = 22; eight homes during the fires, 11 homes during fire containment, 11 homes postfire)	VOC I/O ratios significantly higher postfire compared to during the fire for <i>m,p</i>-xylene (p = 0.004), carbon tetrachloride (p = 0.002), and heptane (p = 0.02)
Dresser et al. 2025 ³⁸	Boulder County, Colorado, USA (Marshall Fire); residential home (n = 1)	Airborne VOCs and PAHs elevated compared with typical levels 10 days post-fire, drop significantly by 15 days and slowly return to approximately 20% of day 10 values by 35 days post-fire (e.g., Benzene reduced from approximately 3 ppbv at 10 days to 0.5 ppbv at 35 days)
Silberstein et al. 2023 ³¹	Boulder County, Colorado, USA (Marshall Fire); residential homes (n = 8, four smoke-affected and four background)	- Priority PAHs increased in smoke-affected homes compared with background. Median summed concentration of 16 PAHs 1859.3 ng/g vs. 1607.5 ng/g. Designated carcinogenic PAHs were lower in smoke-affected homes. - Trace metals increased in smoke-affected homes (median 34.6 mg/g vs 30.1 mg/g).

Reference	Setting	Key findings
Spurgeon et al. 2022 ³⁰	Northern California, USA (22 wildfires); residential homes (n = 343)	– Detected char in 57% of houses, ash in 8.8% of houses, and soot in 1.2% of houses. – Char was 2.8 times more likely to be detected on windowsills compared with interior hard surfaces. – Char detected in 67% of homes < one mile from a wildfire compared with 10% of homes one to two miles away.
Dingle et al. 2021 ⁴⁹	Fort McMurray, Alberta, Canada; residential homes (n = 125)	– Al, Ti, V, As, and Tl concentrations were higher in the house dust from fire-damaged neighborhoods (n = 39–49) compared with non-damaged neighbourhoods (n = 44–61, p < 0.05).
Kohl et al. 2019 ³⁹	Fort McMurray, Alberta, Canada; residential homes (n = 64, 32 in most fire-affected neighbourhoods)	– Median total PAH concentration was 915 ng/g (range: 415–31,200 ng/g) in settled dust. – Naphthalene, phenanthrene, anthracene, benz [a] anthracene, and fluoranthene exceeded Alberta tier 1 soil guidelines, but were lower than reported values in Ottawa homes. – As and V were 62% and 37% higher, respectively, in house dust in neighborhoods with burned buildings (t = 2.08, p = 0.041).

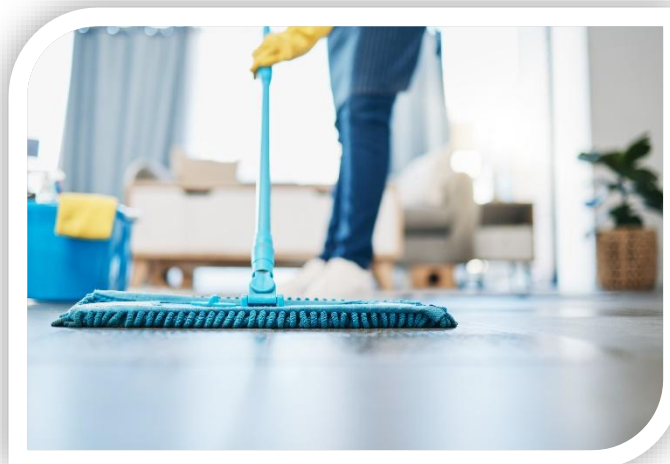
Abbreviations: **PAH:** polycyclic aromatic hydrocarbons, **VOC:** volatile organic compound.



What impact does cleaning have on concentrations of indoor wildfire smoke pollutants?

This section synthesizes the available evidence on the effectiveness of cleaning to remove indoor contaminants post-wildfire smoke episode.

Laguerre et al. tested the effectiveness of several cleaning techniques on glass and cotton that were exposed to smoke generated by burning Douglas Fir and Western Red Cedar needles in a laboratory chamber.⁴² For these experiments, $PM_{2.5}$ reached concentrations of $156 \pm 60 \mu g/m^3$ (mean $\pm$ SD) in the exposure chamber. When comparing the use of ethanol with a commercial cleaner (brand not reported), PAH concentrations on the glass samples were reduced by 71% and 62%, respectively, after cleaning. For the cotton samples, washing in a washing machine and drying the sample removed 48% of PAHs (8.30 vs. $4.33 \mu g/m^2$). However, chrysene and fluoranthene levels remained elevated post cleaning, with a 1.5% increase and a 36% decrease, respectively, compared to exposed samples. From these experiments, Laguerre et al. also estimated the inhalation and dermal exposures for total PAHs to be 43 and 102 ng/day, respectively. These values were reduced to 10 and 50 ng/day, respectively, when cleaning techniques were used.



In Li et al., the effect of HEPA vacuuming, wet mopping, and dry dusting surfaces on airborne VOC concentrations was assessed in a test house exposed to smoke generated from burning pinewood chips.⁴³ The combination of cleaning techniques reduced airborne VOC concentrations of formic acid ($HCOOH$) by 50.3%, formaldehyde ($HCHO$) by 32%, and furan (C_4H_4O) by 19%, as well as WSOCg by 39% for the duration of sampling (12 h). Compared with cleaning, the use of a portable air cleaner (PAC) with an activated charcoal filter only reduced airborne concentrations of formic acid by 3.8%, formaldehyde by 3.1%, and furan by 4.8%. Furthermore, these the concentrations returned to pre-PAC levels 30 minutes after the PAC was turned off. Li et al. also examined the utility of opening windows in removing VOCs, reporting high efficiency values in reducing total VOC concentrations by 74%, but after three hours no further reductions were observed. In summary, vacuuming, mopping, and dusting were more effective at reducing airborne VOC levels than ventilation or use of a PAC.

After the 2021 Marshall fire, the effectiveness of do-it-yourself (DIY) PACs was assessed by Dresser et al. in one home, reporting airborne VOC average concentration reductions of approximately 50%.³⁸ The PACs used in this study were Corsi-Rosenthal boxes with pleated air filters coated with

activated carbon, which may explain the greater effectiveness at reducing VOC levels. Although, after the machines were turned off, VOC concentrations rebounded close to initial values within six hours. Ventilation by opening windows was also assessed in this study, measuring 55–85% reductions in VOC concentrations when five windows upstairs and five windows on the main level of the home were opened. However, approximately two hours after the windows were closed, the VOC concentrations rebounded close to initial levels, indicating the presence of strong indoor reservoirs.

The effectiveness of removing contaminated material was assessed by Link et al. in a test-home study.⁴⁵ The carpets that covered approximately 80% of the floor in the house were removed 100 days after the burn period. After carpet removal, there was a 20% reduction in VOC concentrations compared with air samples taken on day 98. However, the authors indicate that they did not test the carpet itself as a source of VOC emissions prior to the burn period of the study, and some of the observed decrease may have been due to the removal of the carpet material, not smoke pollutants.

The choice of cleaning agent has also been examined when remediating indoor surfaces after a wildfire smoke episode. In Jorga et al., hypochlorous acid mixed with pinewood smoke in a laboratory chamber produced secondary pollutants.¹⁴ Hypochlorous acid is a strong oxidant emitted during cleaning with chlorine-cleaning agents such as bleach. Although the specific secondary pollutants were not reported, the authors noted that the mixture included organochlorine species, which are known to be toxic.

In summary, wet wiping and mopping, and HEPA vacuuming have been demonstrated to be effective methods to reduce wildfire smoke pollutants indoors. However, most studies to date have been in a laboratory setting. Air filtration with PACs and ventilation are also effective at reducing airborne smoke pollutants but do not eliminate the source of the pollutants, namely surfaces where adsorbed and absorbed pollutants may be released over time. There remain evidence gaps on the effectiveness of cleaning strategies for wildfire smoke pollutants, especially settled dust and deposited trace metals after wildfire smoke episodes, including evidence on the effectiveness of different cleaning products and methods for specific types of contaminants (e.g., VOCs, and PAHs). Areas for further study include understanding how wildfire pollutants absorb and desorb from common indoor materials (e.g., furniture, curtains, stuffed animals, clothing, carpets, or hardwood flooring), which could guide the choice of remediation strategy (i.e., disposal vs. cleaning).

Summary

Wildfire smoke introduces a complex mixture of pollutants into indoor environments, where they can deposit on surfaces, persist, and be resuspended or desorbed back into the air. Resuspended wildfire smoke PM is different from normal house dust, and people exposed could experience effects ranging from minor irritation and inflammation of the eyes and respiratory tract to serious exacerbations of chronic respiratory diseases depending on the level of impact and duration of exposure. For this reason, public health and other agencies typically recommend using respiratory protection (e.g., N95 or half-face elastomeric respirator) upon returning home until an initial assessment and/or clean up has been completed.

Evidence from field studies and laboratory experiments shows that PM, polycyclic aromatic hydrocarbons (PAHs), volatile organic compounds (VOCs), and trace metals accumulate indoors following wildfire smoke episodes. Concentrations were generally higher in smoke-impacted homes than in unaffected homes, but multiple studies reported ranges overlapping with typical background levels. These pollutants can persist for weeks to months, with indoor surfaces and materials acting as reservoirs that re-emit contaminants into indoor air, or household activities causing resuspension of pollutants. No human health risk assessment studies on wildfire smoke residues and dust were identified in this review, and expected health risks from the long-term exposure to these pollutants remains to be examined. Cleaning can reduce indoor pollutant levels, but effectiveness varies by pollutant type, surface, and method. Natural ventilation was also effective at reducing airborne concentrations of pollutants, but the effects were transient, with concentrations rising again after windows were closed.

PAHs were elevated in smoke-impacted homes in two studies directly following a wildfire smoke episode, with one study reporting 16% higher median total concentrations of 16 priority PAHs in settled dust. In laboratory experiments, PAHs readily accumulated on test-house indoor materials and persisted for extended periods, up to 40 days. VOCs from wildfire smoke also interacted strongly with indoor surfaces, with surface materials rapidly absorbed VOCs. However, these same materials later acted as emissions sources. VOC persistence varied by chemical properties and surface type, with evidence suggesting that VOCs may remain in indoor environments for weeks to months.

Trace metals associated with wildfire smoke were also detected in indoor dust. For instance, two months after the 2021 Marshall fire, median concentrations of heavy metals in smoke-affected dust were 15% higher than background samples. Similarly, two years after the 2016 Horse River wildfire, concentrations of several metals, including aluminum, arsenic, and vanadium, were significantly higher in homes located in more severely impacted neighborhoods, even after accounting for occupational factors. Elevated levels of lead were also found in homes impacted by the Los Angeles



wildfires and the Latrobe Valley coal mine fire, but in both cases the authors speculated this may be due to the presence of lead paint.

Across field and laboratory studies, smoke PM was consistently detected in settled dust indoors. PM_{2.5} and PM₁₀ concentrations were generally low in unoccupied homes, with spikes noted during cleaning activities two months after a wildfire. Although spikes of similar magnitude can be found in homes not affected by wildfires due to everyday activities, the composition of indoor dust is altered after a smoke episode and may have higher health risks associated with exposure.

Cleaning interventions reduced indoor wildfire smoke pollutants, but the effectiveness varied by method and pollutant. In laboratory experiments, cleaning of smoke-exposed glass reduced PAH concentrations by 62–71%, depending on cleaner type, while laundering cotton reduced PAHs by 48%. In test-house experiments, combined vacuuming, mopping, and dusting reduced airborne VOC concentrations by 19–50%. In contrast, PACs with activated charcoal filters reduced VOC concentrations by less than 5%, and the effects disappeared within 30 minutes after use. However, another study indicated DIY PACs with activated charcoal filters reduced VOC concentrations by approximately 50%, although levels rebounded after the PACs were turned off. Ventilation reduced VOC concentrations by 55–85% across two studies, but this benefit declined within hours. Together, these studies indicate short-term effectiveness of cleaning methods in removing VOCs and PAHs; however, there remain evidence gaps related to long-term effectiveness, material-specific cleaning strategies, and wildfire-related settled dust and trace metals.

Overall, the evidence shows that wildfire-smoke episodes lead to the accumulation and long-term persistence of particulate matter, PAHs, VOCs, and trace metals on indoor surfaces. These pollutants can be resuspended or re-emitted into indoor air during normal household activities, extending potential exposures well beyond the wildfire smoke event itself. While cleaning and ventilation can substantially reduce some contaminants, their short-term effectiveness varies and long-term effectiveness has not been addressed. These limitations underscore the need for careful, evidence-informed guidance to minimize ongoing indoor exposure after wildfire smoke episodes.

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Appendix A: Study details

Reference	Setting	Date	Pollutants	Design	Quality
Goftari et al. 2026	Los Angeles County, California (Palisades and Eaton Fires); residential homes (n = 19)	January 2025 (fires), March 2025 (sampling)	PM _{2.5} , PM ₁₀ , total VOCs, trace metals	Analytical Cross Section	High
Pei et al. 2026	Los Angeles County, California (Palisades and Eaton Fires); residential homes (n = 50)	January 2025 (fires), February 10–April 1, 2025 (sampling)	VOCs	Analytical Cross Section	High
Yao et al. 2026	Los Angeles County, California (Palisades and Eaton Fires); residential homes (n = 22)	January 2025 (fires), January 8–February 18, 2025 (sampling)	VOCs	Analytical Cross Section	High
Silberstein et al. 2023	Boulder County, Colorado, USA (Marshall Fire); residential homes (n = 8)	December 30–31, 2021 (fire), January–February 2022 (sampling)	PM _{2.5} , PM ₁₀ , trace metals, and PAHs	Analytical Cross Section	Moderate
Spurgeon et al. 2022	Northern California, USA (22 wildfires); residential homes (n = 343)	2017–2020	Char, soot, and ash	Analytical Cross Section	Moderate
Dingle et al. 2021	Fort McMurray, Alberta, Canada; residential homes (n = 125)	2016–2018	Trace metals	Analytical Cross Section	High
Wheeler et al. 2020	Latrobe Valley, Victoria, Australia (coal mine fire); residential homes (n = 39)	February 9–March 26, 2014 (fire)	PAHs, trace metals	Analytical Cross Section	High
Kohl et al. 2019	Fort McMurray, Alberta, Canada; residential homes (n = 64)	2016	PAHs, and trace metals	Analytical Cross Section	High
Deeleepojananan et al. 2025	Laboratory chamber experiments using ponderosa pine woodchips	NA	VOCs	Quasi-experimental	High
Farmer et al. 2025	Experimental residential house, ponderosa pine woodchips	NA	PM	Quasi-experimental	High

Reference	Setting	Date	Pollutants	Design	Quality
Studies that include cleaning					
Link et al. 2026	Experimental residential house, pine-spruce wood mixed with gypsum and plastics	NA	Non-Methane VOCs	Quasi-experimental	High
Dresser et al. 2025	Boulder County, Colorado, USA (Marshall Fire); residential homes (n = 1)	December 30-31, 2021 (fire), January–February 2022 (sampling)	PAHs, VOCs	Quasi-experimental	High
Laguerre et al. 2024	Laboratory chamber experiments using Douglas Fir and Western Red Cedar needles	NA	PAHs	Quasi-experimental	High
Jorga et al. 2023	Laboratory chamber experiments using peat tablets and pine wood chips	NA	VOCs	Quasi-experimental	Moderate
Li et al. 2023	Experimental residential house, ponderosa pine woodchips	NA	VOCs	Quasi-experimental	Moderate

Abbreviations: **NA**: not applicable, **PAH**: polycyclic aromatic hydrocarbons, **PM_{x-y}**: particulate matter x-y µm in diameter, **VOC**: volatile organic compound, **SVOC**: semi-volatile organic compound, **WSOCg**: water-soluble organic compounds in the gas phase.

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