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HTAP3 Fires: towards a multi-model, multi-pollutant study of fire impacts

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Abstract. Open biomass burning has major impacts globally and regionally on atmospheric composition. Fire emissions include particulate matter, tropospheric ozone precursors, and greenhouse gases, as well as persistent organic pollutants, mercury, and other metals. Fire frequency, intensity, duration, and location are changing as the climate warms, and modelling these fires and their impacts is becoming more and more critical to inform climate adaptation and mitigation, as well as land management. Indeed, the air pollution from fires can reverse the progress made by emission controls on industry and transportation. At the same time, nearly all aspects of fire modelling – such as emissions, plume injection height, long-range transport, and plume chemistry – are highly uncertain. This paper outlines a multi-model, multi-pollutant, multi-regional study to improve the understanding of the uncertainties and variability in fire atmospheric science, models, and fires' impacts, in addition to providing quantitative estimates of the air pollution and radiative impacts of biomass burning. Coordinated under the auspices of the Task Force on Hemispheric Transport of Air Pollution, the international atmospheric modelling and fire science communities are working towards the common goal of improving global fire modelling and using this multi-model experiment to provide estimates of fire pollution for impact studies. This paper outlines the research needs, opportunities, and options for the fire-focused multi-model experiments and provides guidance for these modelling experiments, outputs, and analyses that are to be pursued over the next 3 to 5 years. The paper proposes a plan for delivering specific products at key points over this period to meet important milestones relevant to science and policy audiences.

1 Introduction

Open biomass burning (BB), which includes wildland fires and agricultural burning (often called “fires” hereafter), has major impacts on global and regional atmospheric chemistry, climate, air quality, and the health of ecosystems, via emissions of air pollutants and greenhouse gases, their long-range

transport, and their deposition. Fire emissions include particulate matter; tropospheric ozone precursors, such as nitrogen oxides (NO_x), volatile organic compounds (VOCs), and carbon monoxide (CO); long-lived greenhouse gases such as methane, nitrous oxide, and carbon dioxide; persistent organic pollutants; and mercury and other metals. While contributions to poor air quality from industrial and transportation sources are decreasing in many parts of the world due to emission controls, fires are a growing contributor to elevated air pollution episodes. Fire frequency, intensity, duration, and location are changing as the climate warms (UN, 2022; Cunningham et al., 2024), and understanding and modelling these changes to fire regimes and their impacts are becoming more and more critical for climate adaptation and mitigation. At the same time, nearly all aspects of fire modelling – such as emissions, plume injection height, long-range transport, and plume chemistry – are highly uncertain. We propose a multi-model, multi-pollutant, multi-regional study to improve the understanding of the uncertainties and variability in fire atmospheric science and its impacts, in addition to providing quantitative estimates of the air pollution and radiative impacts of biomass burning.

The proposed study (herein referred to as HTAP3 Fires) is being planned under the auspices of the Task Force on Hemispheric Transport of Air Pollution (TF HTAP, <http://htap.org>, last access: 15 May 2025), an expert group organized under the Convention on Long-Range Transboundary Air Pollution (UN, 1979), to improve understanding of the intercontinental flows of air pollutants, including aerosols and their components, ozone and its precursors, mercury and other heavy metals, and persistent organic pollutants. TF HTAP has an interest in understanding the relative contribution of fires as compared to other sources, to air pollution impacts on health, ecosystems, and climate at the regional to global scale. TF HTAP is also well-positioned to bring together the multi-disciplinary, international modelling and fire science communities to work towards the common goal of improving global modelling of air pollutants released from fires. Although initiated under TF HTAP, this paper with the plan presented herein is intended to reflect the interests of this broader community and to facilitate communication and co-

ordination between a variety of related ongoing activities and new activities that may be initiated as part of this community plan.

This paper outlines the research needs, opportunities, and options for improving understanding of the climate, air quality, and toxicological impacts of fires and identifies specific research activities and modelling products that could be pursued over the next 3–5 years. Specifically, Sect. 2 contains the motivational science policy questions; Sect. 3 contains background information and defines the scope of this study; Sect. 4 discusses the available options for the model design, providing consideration and justification for the specific plan. Finally, Sect. 5 provides that specific model design plan, which aims to deliver specific products at key points to meet important milestones relevant for science or policy audiences.

2 Motivation: science policy questions

Several open online meetings were organized by TF HTAP in 2022 and 2023 to identify policy-relevant science questions that could be explored in a study of the transboundary air pollution impacts of fires. The questions identified through those meetings have been subsequently refined into the subsections below. The stated questions are not an exhaustive compilation, but the questions do provide important motivation and direction for the HTAP3 Fires multi-model experiments.

2.1 Transboundary transport of fire-emitted compounds

- What are the impacts of fire emissions on air quality, human health, ecosystems, and climate at different scales, from near to far fields?
- What is the role of transboundary movements of fire plumes in impacting atmospheric composition in different regions? And how will the absolute and relative magnitudes of these contributions change over time?
- How does the location or seasonality of large fire events within regions affect the long-range transport potential? And how might these locations change over time with land use and climate change?
- How do plume dynamics and near-fire chemical transformations (e.g. sequestration of NO_x in peroxyacetyl nitrate (PAN), formation of secondary organic aerosols) affect the long-range transport potential and downwind impacts?
- Do different fire types (e.g. agricultural waste burning and wildland fires) have different extents of long-range transport? What are their relative contributions to regional air pollution?

2.2 Fire variability and uncertainty

- What is the range of variability and uncertainty of the results from multiple models' simulations?
- How do model differences in physical and chemical processes manifest in the varied impacts of climate forcing and health that are due to fire emissions?
- Are there certain fire-related parameterizations that perform particularly well against observations and why?
- What are key model parameters that require improved observational constraints to reduce uncertainty?
- What is the impact of different fire emissions inputs on atmospheric concentrations?
- How sensitive are model results to prescribed fire emissions versus prognostic (interactive fire modules that are coupled to climate) emissions?

2.3 Similarities and differences between different pollutants

- What is the contribution of fires to atmospheric concentrations of different air pollutants?
- How do the footprints of different pollutants differ and what are the principal drivers of those differences?
- How much do source–receptor relationships differ based on model type, which often have a different focus (e.g. air quality versus climate), but provide similar subsets of pollutants?
- How do fire emissions interact chemically with other anthropogenic emissions in the atmosphere?

2.4 Questions identified by the research community but that are beyond the scope of this study

- What are the implications of potential regional changes in prescribed burning, fire suppression policies, and other fire management strategies?
- What is the impact on transboundary smoke from local fire management policies?
- What impact does pyrocumulonimbus have on long-range transport of fire emissions? How often and where does pyrocumulonimbus occur and will they become more frequent with climate change?
- What emissions result when wildfires consume buildings and other infrastructure in the wildland–urban interface? What are the health impacts of built–environment burning?

- How much do fires with small burned areas that are not detected by satellite observations influence the fire emissions amount and composition?

3 Scope and background information

The scope and further motivation for this undertaking are defined in this section, partially informing the multi-model experiment design that will appear in Sect. 5, including the model output table (Sect. 5.4).

3.1 Pollutants of interest

Fires emit all the pollutants that the Convention on Long-Range Transboundary Air Pollution (CLRTAP) is concerned with. This study is an opportunity to address all pollutants with the common emission source of open burning. Below is additional information on these pollutants in the context of fires and this modelling study.

3.1.1 Tropospheric ozone and its precursors

Tropospheric ozone (O_3) is both an air pollutant detrimental to human health and vegetation and a short-lived climate forcer (SLCF) (Monks et al., 2015). O_3 is not emitted directly but rather formed through photochemical processes involving nitrogen oxides ($NO_x = NO + NO_2$); hydrocarbons, such as volatile organic compounds (VOCs); methane (CH_4); and carbon monoxide (CO). This chemistry evolves in fire plumes: freshly emitted plumes, typically containing a lot of particulate matter, may suppress O_3 formation due to low-light conditions, diminishing photolysis rates (Alvarado et al., 2015), or heterogeneous chemistry on smoke particles (e.g. Konovalov et al., 2012), whereas aged fire plumes may produce O_3 more efficiently (e.g. Real et al., 2007). Due to a large quantity of VOC emissions from biomass burning, O_3 formation in wildfire plumes is generally NO_x -limited. However, when VOC-rich smoke plumes are transported into NO_x -rich urban pollution, O_3 formation may be enhanced.

The overall impact of fires on O_3 concentrations remains highly uncertain. While NO_x is short-lived, it can be transported long distances in the form of PAN (a reservoir for sequestering NO_x and HO_x radicals), leading to additional O_3 production in downwind regions for moderate smoke plumes, and production increases with plume age (Jacob, 1999; Lin et al., 2010; Jaffe and Wigder, 2012; Fiore et al., 2018). Recent field measurements show that emissions of NO_x and HONO in wildfire plumes are rapidly converted into more oxidized forms such that O_3 production in wildfire plumes becomes rapidly NO_x -limited (Juncosa Calahorrano et al., 2021; Xu et al., 2021). After a few daylight hours, 86 % of the total reactive oxidized nitrogen species (NO_y) are in the forms of PAN (37 %), particulate nitrate (27 %), and gas-phase nitrates (23 %) (Juncosa Calahorrano et al., 2021). When a VOC-rich smoke plume mixes into a NO_x -rich urban

area, it can also create an environment for enhanced O_3 production (Liu et al., 2016; Baylon et al., 2015). The net impact of fires on regional and extra-regional O_3 therefore depends on the emission of a range of precursor species and their chemical transformation in fresh and aged wildfire smoke plumes. Previous HTAP assessments (HTAP1 and HTAP2) have shown that ground-level O_3 is significantly influenced by long-range transport at the hemispheric scale and have demonstrated the utility of a large ensemble of models for quantifying these effects and their uncertainty (Fiore et al., 2009). While fires contribute only a small amount to annual average ground-level O_3 in the major Northern Hemisphere receptor regions, they can be important episodically and may become more important with global warming and reduction of traditional anthropogenic emissions.

The 1999 CLRTAP Gothenburg Protocol (GP; EMEP, 1999) as amended in 2012 regulates the emissions of O_3 precursors in member states. In a recent review, it was concluded that current air quality legislation in the United Nations Economic Commission for Europe (UNECE) region is not sufficient to meet the long-term clean-air objectives of CLRTAP. In support of the CLRTAP response to the recent GP review, TF HTAP is currently organizing a new set of multi-model experiments (HTAP3) aimed at quantifying the contribution of long-range transport to ground-level O_3 in all world regions from remotely emitted O_3 precursors, including from fire emissions (the “Ozone, Particles, and the deposition of Nitrogen and Sulfur”, or HTAP3-OPNS project). To avoid duplication of effort, the model runs contributing to both exercises will be harmonized as much as possible (e.g. using common emission datasets and simulation years).

3.1.2 Methane

CH_4 is the second most important greenhouse gas after CO_2 and modulates the chemistry of many other air pollutants via its impact on atmospheric concentrations of the hydroxy radical (OH). It is also involved in tropospheric O_3 photochemistry (Sect. 3.1.1). In addition to CH_4 being directly emitted from biomass burning, NO_x , CO, and non-methane VOCs (NMVOCs) emitted by fires have the potential to alter regional and global OH concentrations, thus influencing the atmospheric lifetime of CH_4 (e.g. Naus et al., 2022). Modelling studies suggest significant suppression of global OH concentration following enhanced CO emissions from extensive wildfires in Southeast Asia during El Niño events (Duncan et al., 2003; Manning et al., 2005; Rowlinson et al., 2019). Butler et al. (2005) and Bousquet et al. (2006) both found that this change in global OH significantly contributed to the observed increase in global CH_4 concentration during the 1997 El Niño fires. The influence of fires on global OH appears to depend on the location of the fires. Leung et al. (2007) showed that the CO emissions from extensive boreal fires in 1998 did not significantly lower global OH and thus did not significantly contribute to enhanced CH_4

growth. Rowlinson et al. (2019) showed that the increase in CH₄ lifetime induced by El Niño-related fires in the tropics offsets an El Niño-driven reduction in CH₄ lifetime caused by changes in humidity and in atmospheric transport.

Extreme fires and fire seasons may lead to increased CH₄ emissions from wildland fires. For example, the 2020 extreme fire year in California accounted for approximately 14 % of the state's total CH₄ budget, including all anthropogenic CH₄ sources (Frausto-Vicencio et al., 2023). Fires in Arctic tundra will also lead to more CH₄ emissions in the future, as recent observations in Alaska revealed that previously burned tundra (within 50 years) emits more CH₄ than the surrounding landscapes (Yoseph et al., 2023).

3.1.3 Particulate matter

Particulate matter (PM) is emitted in great quantities from fires and is usually the main cause of air quality exceedances during fire episodes. In addition, it has consequences for cloud interactions and radiative forcing. It is comprised of a range of species including black carbon (BC, also known as elemental carbon or soot), primary organic carbon (OC, related to organic aerosol, OA), sulfate (SO₄), nitrate (NO₃), ammonium (NH₄), and crustal material (CM, or dust). Particulate matter may be emitted directly or can be formed as secondary aerosols through gas-to-particle conversion. Secondary organic aerosol (SOA) is particularly important in the context of long-range transport (see Sect. 3.2.5). If smoke is transported through a cloudy boundary layer, aqueous-phase processing can also facilitate the transformation of SO₂ gas into sulfate, with consequences for cloud interactions (e.g. Dobracki et al., 2025). The chemical and radiative properties, as well as cloud interactions, are all dependent on the chemical composition, size, and vertical distributions of the particulate matter (e.g. Huang et al., 2012). BC accounts for about 10 % of smoke plume mass and is the largest contributor to aerosol radiative forcing (RF) (Veira et al., 2016). In contrast to other aerosol components, BC introduces a radiative warming into the Earth's climate system (Sect. 3.2.2). Compared to BC from fossil fuel combustion, BC from biomass burning consists of more, generally larger, particles that are more thickly coated with more absorption per unit mass (Schwarz et al., 2008).

3.1.4 Mercury

Mercury (Hg) is a potent neurotoxin that bioaccumulates in the environment, endangering human health, wildlife, and ecosystems. Wildfires release mercury from plants and soils into the atmosphere, where it may be carried and deposited over great distances, contaminating water bodies and terrestrial ecosystems (Obrist et al., 2018; Chen and Evers, 2023). The Minamata Convention on Mercury (UN, 2013), a worldwide convention enacted in 2013, seeks to safeguard human health and the environment against mercury's negative ef-

fects. It examines the complete life cycle of mercury, including extraction, trading, use, and emissions, emphasizing the need of reducing mercury pollution internationally. A third set of multi-model experiments being organized under HTAP3, known as the Multi-Compartmental Mercury Modelling and Analysis Project (HTAP3-MCHgMAP), is aimed at attributing trends in environmental mercury concentrations to changes in primary mercury emissions and releases or to changes in other drivers or processes (Dastoor et al., 2024). All three sets of HTAP3 experiments (Fires, OPNS, and MCHgMAP) will aim to harmonize inputs and experimental designs as much as possible and avoid duplication of effort.

3.1.5 Persistent organic pollutants

Persistent organic pollutants (POPs) are synthetic chemicals that are also bioaccumulative, toxic, and subject to long-range transport. POPs that have been trapped through wet and dry deposition by trees and shrubs (Su and Wania, 2005; Daly et al., 2007) can be re-released during a wildland fire. The high temperature and vertical winds of wildland fires can remobilize POPs from fuels such as leaves and needles and the forest soil, which otherwise act as a sink for POPs. Eckhardt et al. (2007) reported record high concentrations of polychlorinated biphenyls (PCBs) at the Arctic station of Zeppelin (Svalbard) in a forest fire plume after a transport time of 3–4 weeks. Many atmospheric models do not simulate POPs; however, several POPs models exist, with some listed in Table A2.

The UNEP Stockholm Convention on POPs has provided the framework for global regulation and monitoring of POPs since 2004. However, many POPs (e.g. polychlorinated biphenyls, dichlorodiphenyltrichloroethane, and its degradation products (DDTs); other organochlorine pesticides, polybrominated diphenyl ethers (PBDEs), and per- and polyfluoroalkyl substances (PFASs)) were in use for decades before being regulated. While most legacy POPs in air are declining globally (Wong et al., 2021; Shunthirasingham et al., 2018; Kalina et al., 2019), increasing trends are observed for chemicals of emerging concern, e.g. PFASs (Wong et al., 2018; Saini et al., 2023).

Dioxins are one class of POPs that are formed during incomplete combustion processes. Dioxins are emitted from waste incineration, industrial and residential combustion of fossil fuels, and biomass burning. Global gridded emission inventories are now available for dioxins (EDGAR at <http://edgar.jrc.ec.europa.eu>, last access: 15 May 2025; and Song et al., 2023b). Compared to the early 2000s, global dioxin emission reduced by 26 % in the late 2010s, attributable to emission mitigation in upper- and lower-middle income countries. However, the declining trend of dioxin emissions over the past decades terminated from the early 2010s due to increasing significance of wildfire-induced emissions in the total emission. The highest levels of dioxin emissions (ex-

pressed as polychlorinated dibenzodioxins/dibenzofurans – PCDD/Fs) were identified in East and South Asia, Southeast Asia, and part of sub-Saharan Africa. In East and South Asia, growing dioxin emissions are attributed to industrialization, whereas wildfire is a major contributor to high dioxin emissions in Southeast Asia and sub-Saharan Africa.

3.1.6 Polycyclic aromatic hydrocarbons

Polycyclic aromatic hydrocarbons (PAHs) are organic pollutants primarily generated by incomplete combustion. PAHs are of concern because their concentrations have remained stable despite global emission reductions. PAHs exist in both gas and particulate phase in the atmosphere, allowing them to undergo long-range transport to remote locations (Muir and Galarneau, 2021; Zhou et al., 2012). PAHs are regulated under the UNECE Aarhus Protocol on POPs in the CLRTAP (Yu et al., 2019), yet they are still observed in pristine, remote areas, such as the Arctic and Antarctic regions. The long-range atmospheric transport of PAHs has been extensively investigated and partly attributed to sources in global emission inventories (e.g. PEK-FUEL at <http://inventory.pku.edu.cn/>, last access: 15 May 2025 and EDGAR at <http://edgar.jrc.ec.europa.eu>, last access: 15 May 2025). Further efforts to update global monthly PAH emissions from wildland fire sources from 2001 to 2020 use carbon stock data up to 2020 based on satellite remote sensing (Luo et al., 2020; Song et al., 2023a). The new inventories improve modelling of wildfire-induced PAH levels and trends particularly in the Arctic, sub-Saharan Africa, Southeast Asia, and South America. In the Arctic, source-tagging methods have identified local wildfire emissions as the largest sources of benzo[a]pyrene (BaP), a PAH with high carcinogenicity, accounting for 65.7 % of its concentration in the Arctic, followed by wildfire emissions of North Asia. Wildland fires account for 94.2 % and 50.8 % of BaP levels in the Asian Arctic during boreal summer and autumn, respectively, and 74.2 % and 14.5 % in the North American Arctic for the same seasons (Song et al., 2023a). In the Arctic remote regions, the highly variable, non-changing long-term time trends of PAHs are inconsistent with the global PAH emission reduction and have significantly increased during summers with more frequent wildland fire events in Nordic countries (Yu et al., 2019). Retene (a PAH) was often used as a tracer for wildland fire activities. However, volcanic eruption (Overmeiren et al., 2024) and volatilization from soil and ocean due to warming can also elevate PAHs' air concentrations in remote locations. Models together with observations can better link BB and long-range transport of fire-related substances to remote sites.

3.1.7 Other metals and trace elements

Biomass-burning aerosols also contain a large variety of metals and other trace elements (Perron et al., 2022). The source

can be the vegetation consumed and/or surrounding soils entrained into plumes by strong pyroconvective updrafts (Wagner et al., 2018; Hamilton et al., 2022) or mixing of BB aerosol emissions into advecting dust plumes, as happens in sub-Saharan Africa (Quinn et al., 2022). Entrained soil dust is estimated to be the major (two-thirds) source component for the iron contained in smoke plumes (Hamilton et al., 2022), with other elements needing further investigation. Many of these elements are important components for biogeochemical cycles, human health impacts, and/or aerosol RF.

The mass of iron emitted by fires is particularly important to quantify because iron is a limiting nutrient in many open-ocean regions, playing an important role in CO₂ sequestration, particularly in the southern oceans through increasing phytoplankton primary productivity (Tang et al., 2021; Hamilton et al., 2020).

Other nutrients (e.g. phosphorus) are also emitted from fires in sufficient quantities to warrant deeper understanding of their fluxes and related impact assessment on terrestrial and marine biogeochemical cycles. For example, African fires have been identified as an equal source to African dust in terms of the intercontinental transport of phosphorus to the Amazon rainforest (Barkley et al., 2019). There is also growing evidence that increasing United States (US) fire activity is impacting downwind freshwater ecosystems through depositing phosphorous (Olson et al., 2023).

One practical issue in determining the impact of changes in fire activity on metal aerosol emission and deposition fluxes is quantifying the contribution of fire to the atmospheric loading of a given metal. There are many other sources of metals to the atmosphere, including mineral and anthropogenic dust, fossil fuels and vehicular transport, metal smelting and mining, and volcanoes to name a few (Mahowald et al., 2018; Hamilton et al., 2022). Once sources become well-mixed in the atmosphere, it becomes much more difficult to trace their individual source contributions. One potential avenue in “fingerprinting” the fire source contribution is the use of metal isotopes. In general, different metal sources have different isotopic fractionations (Fitzsimmons and Conway, 2023), and this difference in aerosol characteristic has been used successfully to differentiate iron aerosol between dust and anthropogenic sources (Conway et al., 2019). However, there are currently no data on the iron isotopic signature of fire, so that aspect is beyond the scope of this study.

3.2 Impacts from fires

3.2.1 Human health

Densely populated areas like Southeast Asia, North America, and the Mediterranean experience episodes of intense air pollution from wildfires exceeding the ambient air quality standards that last multiple days or weeks on a regu-

lar basis (Liu et al., 2015; Jaffe et al., 2020; Dupuy et al., 2020; and see Supplement, Sect. S1 for further regional discussions and Sect. S2 for acute exposure health impacts). An estimated 339 000 premature deaths per year (interquartile range: 260 000–600 000) can be attributed to exposure to wildfire smoke worldwide (Johnston et al., 2012). Xu et al. (2023) estimated that each person in the world had an average of 9.9 d of smoke exposure from 2010–2019, a 2.1 % increase compared to the previous decade. The impacts are projected to increase under future climate change (Xie et al., 2021). In many regions of the world, farmers commonly burn crop residue to clear land for crop cultivation. However, these agricultural fires have health implications as air pollution increases (Jones and Berrens, 2021). During peak fire periods, these agricultural fires can contribute more than half of the particulate matter (PM) pollution, even in urban settings (Cusworth et al., 2018; Liu et al., 2018).

Health risk assessment models and air quality health indices are often based on surface level concentrations of $\text{PM}_{2.5}$, CO , O_3 , and NO_x . Emissions of $\text{PM}_{2.5}$ from fires are of particular health concern, with no known safe $\text{PM}_{2.5}$ concentration in air, as noted by the World Health Organization (WHO, 2006). Fine particles impact lung function, encouraging respiratory and cardiovascular mortality and morbidity, including asthma and emphysema (Davidson et al., 2005; Lampe et al., 2008; Jain et al., 2014; Reid et al., 2016; Cascio, 2018; Ghosh et al., 2019; Chen et al., 2021; Aguilera et al., 2021a, b; Sonwani et al., 2022; Gao et al., 2023; Bauer et al., 2023). There is also evidence that wildfire smoke affects mental health (Eisenman and Galway, 2022; To et al., 2021), such as due to displacement and smoke exposure following wildfires, which can lead to increased cases of anxiety and post-traumatic stress disorder (e.g. Humphreys et al., 2022).

An additional consideration is how smoke influences the structure of the boundary layer and thus the concentration of pollutants that people are exposed to. Fire aerosols, by cooling the surface and reducing boundary layer turbulence (Sect. 3.2.2), can suppress mixing of air in the boundary layer, effectively increasing pollution exposure at the surface (Bernstein et al., 2021). This effect has been studied extensively in polluted urban environments, but its importance for fires, where the composition of aerosol may be substantially different, remains unclear.

Finally, the chemical composition of the PM influences its health impacts. For example, benzo(a)pyrene, the most toxic congener of 16 parent PAHs, has been linked to high lifetime cancer risk from inhalation. Knowledge of PM size distribution (e.g. Sparks and Wagner, 2021) and chemical composition is essential for understanding health impacts, thus motivating our multi-pollutant approach to these model simulations.

3.2.2 Climate and radiative forcing (RF)

While wildland fires have long been considered a natural and relatively carbon-neutral component in the Earth system (CO_2 emitted during burning is reabsorbed as the forest regrows), land use change and anthropogenic climate change have caused the frequency and intensity of fires to rapidly change, potentially altering the global carbon budget. As radiative forcing is typically expressed as a change relative to the preindustrial era, and the magnitude of preindustrial fires is highly uncertain, there is a factor of 4 uncertainty in RF from fires (Hamilton et al., 2018; Wan et al., 2021; Mahowald et al., 2024).

Fire emissions have diverse effects on the climate. In addition to the direct effects from released greenhouse gases and aerosols, additional indirect effects arise from the formation of tropospheric O_3 , reduction in lifetime of CH_4 by enhancing tropospheric oxidation capacity, and changes in stratospheric water vapour caused by responses of the atmospheric chemistry. Co-emitted SO_2 can also become converted to SO_4^{2-} , an effective cloud nucleator, thereby affecting cloud lifetime (Dobracki et al., 2025). The aerosols have indirect (microphysical) and semi-direct (radiative) impacts on cloud fields and large-scale circulation (Adebiyi and Zuidema, 2018; Diamond and Wood, 2020; Ding et al., 2021). Short-term radiative effects of smoke on surface wind, temperature, moisture, and precipitation can also substantially enhance fire emissions and weaken smoke dispersion (Grell et al., 2011; Huang et al., 2025). Snow and ice albedos also change dramatically when fire-emitted black and brown carbon are deposited. Additionally, indirect effects on biogeochemistry result from wildfire emissions (Sects. 3.1.6 and 3.2.3).

The RFs from fire plume components are summarized in Table 1. Though most studies focus on specific components or regions for wildfire RF (e.g. Mao et al., 2013; Chang et al., 2021; Moubarak et al., 2023), Ward et al. (2012) conducted a comprehensive global analysis of wildfire emission's RF, encompassing all components.

The large range in aerosol indirect effect heavily depends on the background conditions. Aged smoke is an excellent source of cloud condensation nuclei (e.g. Kacarab et al., 2020), but increasing cloud condensation nuclei from other emission sources can reduce the RF from wildfire emissions (e.g. Ward et al., 2012; Hamilton et al., 2018), which is a general feature for natural emissions (Spracklen and Rap, 2013). A reduction of anthropogenic emissions in the future could increase the effects of natural emissions (see example for tropospheric ozone by Mertens et al., 2021). The estimate of the aerosol albedo effect also varies in sign, but the magnitude is in general rather small compared to the indirect aerosol effect (Tian et al., 2022). The height of the fire plume influences its RF, and recent studies suggest a large climate impact of fire emissions that rise into the stratosphere (Stocker et al., 2021; Damany-Pearce et al., 2022). Moreover, new measure-

Table 1. Summary of present-day RFs from specific fire plume components. Please note that the different studies use very different model simulations and approaches to estimate the RF (see original papers). This table only serves as a general overview of the order of magnitude of the effects found in previous studies.

Fire emission component	RF (W m^{-2})	Comments
Tropospheric O_3	0.03 to 0.05	Dahlmann et al. (2011), Ward et al. (2012). Depends heavily on the emissions from other sources, emission location, and plume height (e.g. Naik et al., 2007; Paugam et al., 2016).
Aerosol direct effect	−0.20 to 0.25	Rap et al. (2013), Tian et al. (2022). Depends on uncertainties in BC absorption and height of the smoke plume.
Aerosol indirect effect	−1.11 to −0.09	Tian et al. (2022), Rap et al. (2013). Depends on background conditions.

ment data indicate a larger warming potential of the aerosol emissions from grassland fires, in part because of low single-scattering albedos resulting from a high fraction of refractive black-carbon (rBC)-containing particles and relatively low OA : rBC mass ratios (Dobracki et al., 2023). New data on long-range transported aerosol might help to reduce these discrepancies between the models (Zhong et al., 2022).

3.2.3 Ecosystems

Fires impact land cover, runoff/infiltration, soil erosion, and water quality, via reducing water use by plants and increasing soil hydrophobicity. The impact depends on the surface (topography, vegetation type, soil type) and fire properties as well as the quantity and intensity of precipitation following the fires. For example, high forest fire counts in India can decrease the soil moisture content, evapotranspiration, and normalized difference vegetation index (Jain et al., 2021). Further regional discussions can be found in Sect. S1. Note that human intervention/management practices to reduce these fire impacts vary by region, but those activities may or may not be accounted for or represented well in atmospheric and Earth system models.

Fires can also positively or negatively impact aquatic and land ecosystems nearby and afar via deposition. Specifically, fires can impact downwind marine ecosystems if deposition is sufficient to alleviate nutrient limitation in the surrounding waters (Hamilton et al., 2022). For example, Siberian fires were recently linked to anomalously high phytoplankton growth in the Arctic Ocean through the additional atmospheric supply of nitrogen (Ardyna et al., 2022). Ozone produced from fire and other emissions can reduce the productivity of O_3 sensitive ecosystems, perturbing biogenic emissions.

The estimated deposition fluxes depend highly on the models' deposition schemes and vary by chemical species and surface types (e.g. Tan et al., 2018; Huang et al., 2022). Through radiative impacts which are only accounted for in some models, fires can perturb numerous variables relevant to the calculation of deposition velocity/coefficient and sec-

ondary pollutant formation (e.g. see Huang et al., 2025, for a Canadian wildfire event in 2023 that enhanced O_3 and nitrogen deposition in the eastern USA).

3.2.4 Socioeconomics and fire management decisions

In cases of forest fires that encroach on the wildland–urban interface, people are forced to evacuate or permanently relocate their homes. High fatalities of residents (e.g. Molina-Terrén et al., 2019), firefighters, and fauna; severe air pollution ranging over a few to thousands of kilometres; and huge economic losses from property damages, national park closures, tourism and recreational activity curbs, highway blocks, air travel diversions, and forest-based livelihood losses (e.g. Psaropoulos, 2021) result from large scale, recurrent forest fires (Bowman et al., 2011).

Catastrophic wildfires around the world are increasingly more frequent and hazardous. For example, in the United States, fire-loss events increased from an average of 1.5 events per decade from 1980–1999 to 7 per decade from 2000–2019, costing the nation a cumulative USD 10 billion and USD 75 billion, respectively (Smith et al., 2020). Few studies have reported on the increasing socioeconomic impacts and diversity of people and communities being affected (Moritz et al., 2014; Bowman et al., 2018). Further studies denoting the dollar cost of fire events include Masters (2021) for the 2019–2020 Australia fires and Wang et al. (2021) for the 2018 California fires. Additional regional discussions can be found in Sect. S1.

The wildland–urban interface (WUI) is the area where human development meets or intermixes with wildlands (Stewart et al., 2007; Platt, 2010). Increased human availability in the WUI leads to more human caused ignitions, while simultaneously wildfires in this area pose a greater risk to structures and lives. Thus, WUI fires are harder to manage yet must be suppressed (Choi-Schagrin, 2021). The demographics of the WUI are regionally dependent (e.g. Wigtill et al., 2016; Davies et al., 2018; Tang et al., 2024) and are changing with time, as housing costs (Greenberg, 2021) and immigration (Shaw et al., 2020) evolve over time.

Moreover, some studies focused on environmental justice describe various impacts to, and the social vulnerability of, different communities. Wildfires preferentially impact US regions with lower populations of minorities and higher populations of elderly (Masri et al., 2021). Elderly populations are particularly vulnerable to the effects of fire (Masri et al., 2021; Liu et al., 2015; Murphy and Allard, 2015). Indigenous communities also have high vulnerability, because they are disproportionately located in areas of high fire risk (Davies et al., 2018).

Land management decisions have an important role in determining ecological and socioeconomic pathways. Prescribed or controlled burning is an important tool within holistic land management plans for enhancing ecosystem resilience, biodiversity conservation, plant response, air quality, and carbon sequestration. Each of these benefits are expanded upon in Sect. S3, with the general conclusion that collaboration with local communities, incorporation of traditional ecological knowledge, and adaptive management techniques guarantee that land management decisions are consistent with sustainable practices. Further research beyond the scope of this study is needed to incorporate these kinds of land management decisions into fire emissions scenario inputs for atmospheric models.

3.2.5 The role of atmospheric long-range transport

Long-range transport of fire-related pollutants makes open biomass burning relevant for regions that are not typically impacted by widespread, frequent, or intense fires. For example, recent Canadian 2018 and 2023 fires were reported to cause high PM and O₃ pollution episodes in the USA (e.g. Xie et al., 2020; Lin et al., 2024b; Huang et al., 2025), and these plumes can reach Europe through long-range cross-Atlantic transport (Real et al., 2007; Alvarado et al., 2020; CAMS, 2023). In tropical regions, prevailing easterlies and the African Easterly Jet South (Adebiyi and Zuidema, 2018) can readily transport biomass-burning aerosol from Africa to South America (Holanda et al., 2020). The biomass-burning aerosol interactions with a large subtropical low cloud deck vary microphysically and radiatively with the vertical collocation of aerosol and cloud (Kacarab et al., 2020; Zhang and Zuidema, 2019, 2021). Smoke is also an annual occurrence in northern Thailand, and northern part of Southeast Asia, transported regularly to southern China and Taiwan. At even larger scales, global teleconnections such as the El Niño–Southern Oscillation allow Indonesian peat fires to impact atmospheric loadings as far away as equatorial Africa (Doherty et al., 2006; Lin et al., 2014). Smoke also impacts the Southeast Asian monsoon through increasing the low cloud coverage (Ding et al., 2021).

Long-range transport depends on many factors including but not limited to source proximity, plume height, synoptic weather conditions, large-scale general circulation, atmospheric chemistry, and deposition rates. Long-lived primary

pollutants such as CO may be transported on a hemispheric scale, while short-lived species such as PM and NO_x typically affect a much smaller region. However, the formation of secondary pollutants within the plume introduces substantial uncertainty into the broader atmospheric impacts of fires. In particular, the formation of longer-lived pollutants such as O₃, PAN, and secondary fine particles can substantially impact atmospheric composition over intercontinental distances, documented in both observational and modelling studies (Real et al., 2007; Lin et al., 2024b). The timing and magnitude of secondary pollutant formation in transported plumes strongly influence the health and ecosystem impacts of distant downwind regions and introduce much uncertainty in our assessment of these impacts. Sensitivity experiments with atmospheric chemistry transport models, constrained with estimates of formaldehyde which can be detected from space (Zhong et al., 2022; Alvarado et al., 2020), are important for understanding the long-range impact of fire-related primary and secondary pollutants on receptor regions.

As nations implement more stringent air quality targets, long-range transport will start to play an increasingly important role in determining if these targets are met. The multi-model study proposed here will include regional emissions perturbation experiments (Sects. 4.5 and 5.3) to quantify the long-range impacts on local atmospheric composition.

3.3 Leveraging recent and ongoing efforts

Several distinct scientific communities are addressing fire research and applications in line with their specific objectives. Table A1 lists the recent and ongoing efforts in the community that are complementary but not duplicating the research outlined in this paper. For example, the IGAC BBURNED activity hosted a workshop in November 2023 to assess current global biomass-burning emissions datasets and recommend one as the baseline fire emissions dataset for this work (Sect. 5.2 and 5.3). The Arctic Monitoring and Assessment Programme (AMAP) SLCF expert group may utilize the model output from this work for a future Arctic-focused biomass-burning report. A further example is the Climate Model Intercomparison Projects CMIP6 and CMIP7 activities: CMIP6 and FireMIP included simulations from dynamic vegetation models with interactive fire modules. These provide future fire emissions for different climate scenarios as input for this work (Sects. 4.2.3 and 5.2). AerChemMIP2, planned for CMIP7, will include fire-focused simulations for their aerosols and gas chemistry climate impacts.

4 Discussion of modelling options

In this section, we establish the range of model types expected to participate and then discuss different options for model inputs, such as emissions and driving meteorology. We also discuss what kinds of simulations could be carried out to

answer the science policy questions of Sect. 2. Final guiding decisions on all of these topics are provided in Sect. 5.

4.1 Model types and scope

Models suitable for exploring the local, regional, and global impacts of fires have a wide range of different geographic and temporal scales and resolutions. Models of atmospheric processes have widely differing treatments of chemical complexity and differ in their vertical and horizontal extent. Some models incorporate physical processes to simulate their own meteorology, which may be nudged to match meteorological reanalyses, while others are driven with reanalysis data, either directly or following downscaling with a meteorological model. More complex models may incorporate other Earth system components including the land surface and vegetation (which may or may not be interactive), ocean exchange (and sometimes biogeochemistry), and the cryosphere. In some models, fire ignition, spread, and pollutant emission are explicitly represented, governed by vegetative fuel loading and meteorology, while in others they are a specified input. This diversity in model types and scope presents a technical challenge in comparing the simulated impacts of fires between models (e.g. Shinozuka et al., 2020; Doherty et al., 2022), but the different approaches and levels of complexity present a valuable opportunity to provide fresh insight into our understanding of fire processes and how they are best represented for specific goals. The models participating may fall into these categories:

- Earth system models (ESMs) or coupled chemistry–climate models (CCMs)
- regional or global chemical transport models (CTMs)
- multi-media POP models
- Lagrangian transport models
- reduced-form, surrogate models (e.g. emulators)
- inverse models (see Sect. S4 for more information).

Of note, in hindcast or historical simulations, regional prognostic meteorological models can ingest (or downscale) reanalysis data in two different ways, i.e. with or without nudging. The former deals only with initial and boundary conditions. The latter dynamically nudges model output towards selected reanalysis fields, which helps preserve or maintain the underlying meteorological conditions generally at meso- and synoptic scales. Modellers in the HTAP3 Fires can weigh which way is more justifiable to their purposes. However, nudging in online coupled modelling may not be encouraged for some applications since it potentially obscures or affects interactions between meteorology and chemistry. The modelling centres in Table A2 have indicated interest in participating in this study. The characteristics of the models in Table A2 are taken into consideration for the experimental design.

4.2 Available emissions inputs for historical and future simulations

Almost all atmospheric models will require some information about anthropogenic and natural emissions as inputs. In this section, we discuss available datasets for both historical and future anthropogenic and natural emissions relevant for a global multi-model study. Extricating truly natural from anthropogenic biomass burning is a tricky endeavour that is beyond the scope of this study. For example, while agricultural and deforestation fires are considered uncontroversially as anthropogenic, would accidental human ignition of a wild-fire be considered natural or anthropogenic biomass burning? Similarly, would wildland fires that are more frequent and intense due to anthropogenic climate change be considered natural or anthropogenic? For the model design and interpretation of results, we simplify the total fire emissions into those with and without agricultural burning and classify traditional fossil fuel emissions as anthropogenic. Agricultural burning appears in both kinds of emissions datasets, so guidance is provided in Sect. 5.2 on which to use to not double-count those emissions.

4.2.1 Historical and future anthropogenic emissions

The HTAPv3 global anthropogenic emissions mosaic (Crippa et al., 2023) covers the time period 2000–2018 at $0.1 \times 0.1^\circ$ spatial resolution and monthly temporal resolution. This mosaic inventory is based on the EDGAR 6.1 global inventory and incorporates detailed emissions (for 16 sectors) for SO_2 , NO_x , CO, NMVOC, NH_3 , PM_{10} , $\text{PM}_{2.5}$, BC, OC, and four POPs species from several national and regional inventories using the original spatial distributions wherever possible. Speciation profiles are supplied for PM and NMVOC emissions. The REASv3.2.1 regional inventory is used for Asia (South Asia, East Asia, and Southeast Asia); the CAMS-REGv5.1 regional inventory is used for Europe; the CAPS S-KU national inventory is used for South Korea; and emission data from the respective national authorities of Japan, Canada, and the United States of America are used for the respective geographical zones. Wherever the respective regional or national inventories did not include specific emission sectors, or wherever these sectors did not include the full set of species provided by EDGAR 6.1, these emissions were gap-filled using EDGAR 6.1. Conversely, the minor sources that regional and national inventories had that were *not* present in EDGAR 6.1 (e.g. CO, NO_x , and SO_2 from the solvent sector) were included in the HTAPv3 mosaic. HTAPv3 is thus a complete and model-ready dataset representing the best-available emissions for global and regional model simulations aimed at informing air quality policy. By January 2025, HTAPv3.1 global anthropogenic emissions are expected to be released, which are as above, except covering the years 2000–2020, based on EDGARv8, and including updated emissions from the regional inventories.

Future scenarios of anthropogenic air pollutant and CH₄ emissions are available from the IIASA GAINS integrated assessment model for the period 1990–2050. The scenarios are based on those originally produced in 2021 by IIASA to support the review of the amended Gothenburg Protocol carried out under the Convention on Long-Range Transboundary Air Pollution, as well as the AMAP SLCF (AMAP, 2021) assessment report, and shown to be more realistic than those used in CMIP6 in Ikeda et al. (2022). The next version of these scenarios, called GAINS LRTAP, is available as of January 2025 (Klimont et al., 2025) and will be used to support HTAP3 activities aimed at modelling future air quality to inform the CLRTAP policy response to the Gothenburg Protocol review. Three scenarios are provided: CLE (current legislation) is based on realistic implementation of existing air quality plans; MTFR (maximum technically feasible reduction) is based on the same underlying activity data as CLE but with full implementation of all proven technical measures to abate CH₄ and air pollutant emissions regardless of cost effectiveness; and LOW, which builds on MTFR, adding additional structural measures representing climate policies consistent with Paris Agreement goals and dietary changes aimed at reducing emissions from the agriculture sector.

The HTAPv3.1 historical emissions and LRTAP future scenarios will be used in other concurrent HTAP3 projects (MCHgMAP and OPNS). Use of these emissions datasets would provide consistency across the HTAP3 experiments and would maximize policy relevance of the experiment results. While the historical emissions from HTAPv3.1 and the future scenarios from LRTAP do overlap in time (2010–2020), they have not been harmonized with each other, so they do not provide a seamless time series of anthropogenic emissions from 2000 to 2050. Therefore, results from the historical simulations and future simulations should be analyzed separately. For example historical trends from the early 2000s to 2020 can be assessed, and future trends from 2010 to 2050 can be assessed, each with a consistent source for anthropogenic emissions. In addition, in Sect. 4.2.3, we will see that the future BB emissions are harmonized to the historical BB emissions; thus an absolute change for BB emissions from early 2000 to 2050 is also possible. Note that the HTAPv3.1 emissions do not contain POPs, PAHs, Hg, and other toxic emissions. The anthropogenic emissions for those species are available from EDGARv8.1 (Hg), PKU-GEMS (PAHs), and PKU-LZU (Song et al., 2023a, b) for PCDD/Fs.

Each modelling centre will need to pre-process the selected emissions datasets to account for vertical profiles and diurnal variations of these emissions. As these processes may differ across models and it may not be possible to harmonize these characteristics, these processes will introduce a source of variability in emissions inputs across models. However, if models use their own default assumptions for vertical and temporal allocation, their methods and assumptions should

be reported with their output and taken into consideration in the analysis of outputs.

4.2.2 Historical biomass-burning emissions

The latest available major global fire emissions datasets are GFEDv4s (van der Werf et al., 2017), GFASv1.2 (Kaiser et al., 2012), FEERv1.0-G1.2 (Ichoku and Ellison, 2014), FINNv2.5 (Wiedinmyer et al., 2023), FLAMBE (Reid et al., 2009), QFEDv2.5 (Darmenov and da Silva, 2013), GBBEXv4, and IS4FIRES (Sofiev et al., 2009). Developers of each of these datasets attended and presented their methods at the Fire Emissions Workshop (FEW2023 at <https://www2.acom.ucar.edu/bburned/fire-emission-workshop-virtual-2023>, last access: 16 May 2025; co-hosted by BBURNED and TF HTAP) in November 2023. Intercomparison studies such as Griffin et al. (2023), Pan et al. (2020), Wiedinmyer et al. (2023), and Liu et al. (2020) were also presented there, and the workshop attendees discussed options for which dataset to recommend for consistent baseline fire emissions. An intercomparison tool called FIRECAM (<https://globalfires.earthengine.app/view/firecam>, last access: 16 May 2025) was useful for intercomparison. The different methodologies used to estimate fire emissions (e.g. Table 2) account for how and why the emissions results are so different from one another (Fig. 1). The intercomparison studies demonstrated that no one fire emissions dataset performed best for all locations and all pollutants. The Parrington et al. (2025) report summarizes all of these results.

Fire emissions from peat

Satellite data assimilation studies have shown that emission inventory underestimations may often be due to lack of peat fires. For example, Nechita-Banda et al. (2018) found that incorporating satellite measurements of CO increased CO emissions (compared to GFAS and GFED) from peat fires in Indonesia during the 2015 El Niño event. The ability to account for emissions from peat fires is a key issue in several regions. Recent work to improve peat fires for Indonesia was done in Kiely et al. (2019). Out of several global fire emissions datasets, only GFASv1.2 and GFEDv4 have tropical peat fires, and only GFASv1.2 contains high-latitude Siberian peatland fires. GFEDv5 emissions will have high-latitude peat fires, but as of this writing, it has not yet been released, nor evaluated. Similarly, a newer version of GFAS (v1.4) has not been published or documented yet, though it could have improvements for long-term trends in fire emissions.

Regardless of their inclusion, peat fire emissions are highly uncertain (McCarty et al., 2021). There are different EFs for high-latitude and low-latitude peat fires, given the different vegetation that grows on top, and global maps of

Table 2. Summary of characteristics of major global fire emissions, adapted from Liu et al. (2020). Dashes indicate missing values. Note: FRP – fire radiative power, BA – burned area.

	Horizontal resolution	Temporal resolution	Near-real time availability	Input satellite fire product	Peatlands included	Cloud gap adjustment	References
GFEDv4s	0.25°	monthly	–	MODIS BA + active fire geolocations	✓ (tropical)	–	van der Werf et al. (2017)
GFASv1.2	0.1°	daily	✓	MODIS FRP	✓ (Siberian and tropical)	✓	Kaiser et al. (2012)
FEERv1.0	0.1°	monthly	✓	MODIS FRP	–	✓	Ichoku and Ellison (2014)
FINNv2.5	1 km	daily	✓	active fire geolocations from MODIS & VIIRS	–	–	Wiedinmyer et al. (2011, 2023)
FLAMBE	1–3 km	hourly	✓	MODIS thermal anomalies	–	–	Reid et al. (2009)
QFEDv2.5	0.1°	daily	✓	FRP	–	✓	Darmenov and da Silva (2013)
GBBEXPv4	0.1°	hourly	✓	VIIRS FRP	–	–	Parrington et al. (2025)
IS4FIRES	0.1°	daily	✓	MODIS FRP	–	–	Sofiev et al. (2009)

peatland are out of date (McCarty et al., 2021). It is also very difficult to detect smouldering (low-intensity) peat fires from satellite measurements. That said, the consensus recommendation from FEW 2023 was to use GFASv1.2 based on its inclusion of high-latitude peat fires, ease of adjusting EFs, possibly somewhat better sensitivity to small fires in fire radiative power (FRP) than in-burnt area, and availability of information on the diurnal cycle (more on timing of emissions below). Unlike other types of wildland fire emissions, tropical and mid-latitude peat fires generally have a flat diurnal cycle, apparent in the FRP observations during daytime and night-time (e.g. Fig. 10 in Kaiser et al., 2012). Diurnal information is directly available in the separately assimilated daytime and night-time FRP in GFASv1.4, but this database has not yet been used to adapt the emission factors.

Magnitude of emissions

Substantial uncertainty arises from estimates of the magnitude and location of emissions. This can be explored through short case study simulations investigating the use of alter-

native emission datasets, along with comparison of these with observations and baseline model studies. Such sensitivity studies implicitly include differences in resolution and species fractionation (and possibly injection height and timing), as well as fire magnitude and location but nevertheless can provide a useful estimate of uncertainty to fire emissions across the models (Pan et al., 2020).

Timing of emissions

Most long-term model studies, such as those performed for CMIP intercomparisons, apply monthly mean fire emissions rather than considering more temporally resolved emissions that capture the largely episodic nature of fires. The implications of this, either for comparison with surface observations or for regional and global budgets, remain unclear. In addition, there are substantial diurnal cycles in fire intensity, local meteorology, and boundary layer dynamics that suggest that the impacts of fires are likely sensitive to the timing of emissions throughout the day. Observational evidence indicates emerging overnight fires due to increasing drought

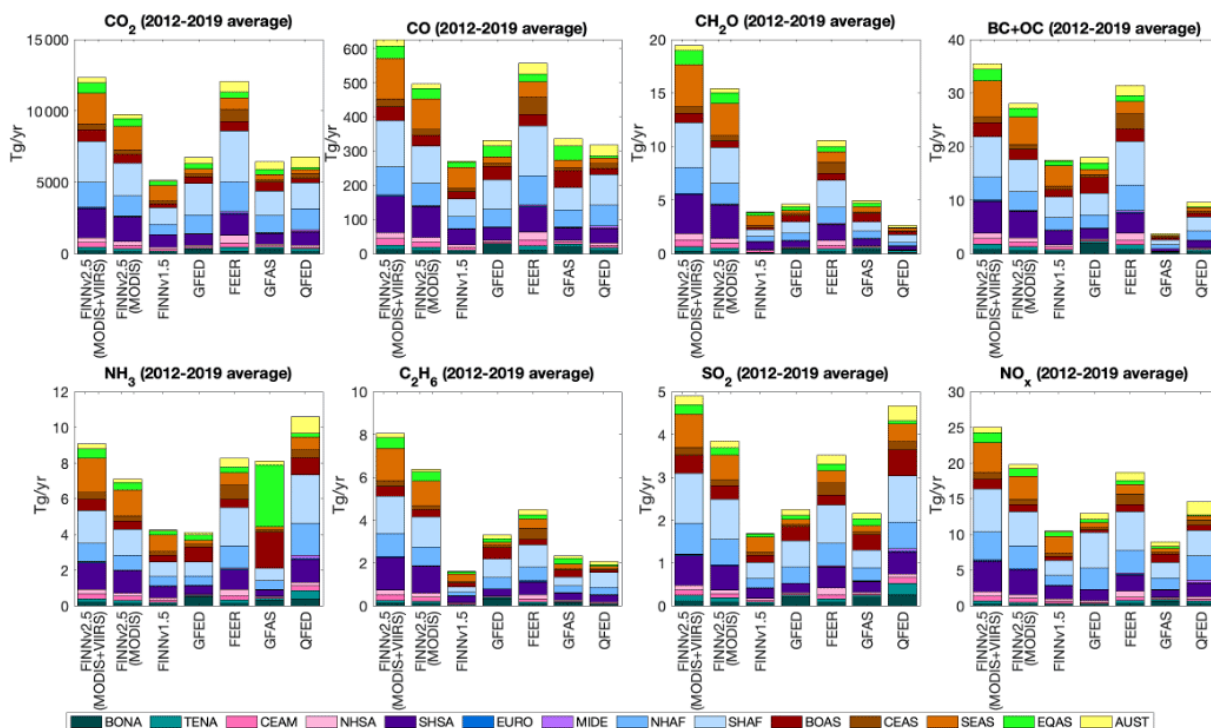


Figure 1. Multi-species, multi-regional intercomparison of fire emissions datasets, from Wiedinmyer et al. (2023), where the regions (colours) are shown on a map in our Fig. 3c.

conditions that challenge the traditional diurnal cycle characterized by “active day, quiet night” (Luo et al., 2024). These uncertainties can be explored through short studies (1 year/several years) that consider (1) monthly mean fire emissions, based on the same set of emissions used at higher temporal resolution in the baseline run, and (2) emissions provided without a diurnal variation in magnitude or injection height.

Fire emissions of other species: Hg, POPs, and PAHs

Mercury fire emissions are not included by default in most global fire emissions datasets, like GFASv1.2. For the HTAP3 MCHgMAP project, Hg fire emissions are based on FINNv2.5 global fire emissions, using emission factors from Andreae (2019) but replacing EFs for certain biomes with mean EFs from Friedli et al. (2003a, b) and McLagan et al. (2021). The MCHgMAP team also apply those EFs to GFED4 fire emissions for sensitivity simulations. Those biome-specific EFs could be applied to the chosen fire emissions datasets (GFAS4HTAP and Hamilton and Kasoar) for this project to generate consistent Hg fire emissions.

Similarly, EFs used for POPs and PAHs could be added and applied to the base fire (and anthropogenic) emissions for this study. For PAHs, there is a recently updated Peking University (PKU-FUEL) “global PAH emission inventory” spanning from 1961 to 2020 at <http://inventory.pku.edu.cn/> (last access: 15 May 2025), which takes wildfire emission into account and used measured PAH emission factors. That

group also developed global OPFR, SCCP, and PCDD/Fs emissions, often using experimentally derived emission factors from the USEPA and UNEP as well as the literature (He et al., 2004; Jiang et al., 2017; Song et al., 2023b; Li et al., 2023).

Post-fire dust emissions

The removal of vegetation creates a more exposed soil surface from which dust can be emitted (Dukes et al., 2018; Jeanneau et al., 2019; Whicker et al., 2006). The emission of dust from a post-burn landscape will continue until the vegetation sufficiently recovers, spanning a period of days to potentially years. Approximately one in two large fires is estimated to be followed by increased dust emissions, with savanna ecosystems the most susceptible (Yu and Ginoux, 2022). Emission estimates are highly uncertain with the only global estimate to date of 100 Tg yr^{-1} of additional soil dust emissions with an order of magnitude uncertainty (Hamilton et al., 2022). As there are no existing emissions datasets for this process, further research beyond the scope of this study would be needed to address this impact of fires.

Wildland–urban interface fires

Wildland–urban interface (WUI) fires account for $\sim 4\%$ of total fires globally. WUI fires can involve built-structure burning, and hence their emissions may be more harmful.

They are also closer to humans and properties causing expensive damages. Studies have been conducted for specific WUI fires and regions (Holder et al., 2023), and a future version of FINN will include WUI fire emissions. However, currently there is no global BB emissions dataset that explicitly addresses WUI fire emissions, and future research beyond the scope of this study would be needed to address this aspect of fire emissions.

Summary of the recommended historical fire emissions dataset

The discussions at FEW 2023 suggested that several characteristics are important in selecting a fire emissions dataset for the multi-model experiments: (1) high temporal resolution, given the high variability of fires; (2) the inclusion of boreal peatland fires, particularly for those interested in boreal and Arctic locations; and (3) the inclusion of fire plume height for atmospheric modelling. For these reasons, GFASv1.2 became the recommended fire emissions dataset for the historical period (starting in 2003), but it was modified with an updated fire type map and emission factors (hereafter called GFAS4HTAP; Kaiser et al., 2024b).

4.2.3 Future biomass-burning emissions

Land models, such as those that participated in FireMIP (Li et al., 2019), can provide fire emission projections that are reflective, not only of future land use changes, but also of the changing climate under different future climate scenarios. However, as of this writing, the FireMIP future simulations have yet to be conducted.

Current CMIP6 SSP future fire datasets only account for human impacts on future fire activity, whereby fire activity is assumed to decrease and includes no impact from the changing climate conditions on those future fire emissions (Fig. 2, left bars). An alternative set of future fire emission projections does however exist in six fire–climate coupled models from CMIP6 (Xie et al., 2021). The models have future fire results that take into account the changing climate under different SSP scenarios: “a climate-consistent future fire emissions estimate” (Fig. 2, right bars). Emissions for three SSP scenarios (SSP1-2.6, SSP3.70, SSP5-8.5) have been produced by Hamilton et al. (2025), and other SSP scenarios using the same methodology can be generated (i.e. SSP2-4.5 for this study from 2020 to 2050). In each emission projection “natural” fire emissions are defined as boreal and temperate forest and all grassland fires and calculated as a product of the CMIP6 multi-model mean, accounting for similarities in land models. “Human”-controlled agricultural and deforestation fires are then added to natural fires. Agricultural fires are included using the GAINS anthropogenic emission estimates (Sects. 4.4.1 and 5.2 for this recommendation). Tropical forest fires are assumed to be primarily due to deforestation practices and were added from the SSP

dataset in place of CMIP6 model estimates in that forest biome. Peat fires are held at present-day levels throughout the century, also following SSP dataset protocols, very likely underestimating their contribution to future emission fluxes, but this is because most interactive ESM fire modules did not contain these uncertain types of fires. Finally, each emission dataset is bias-corrected regionally to dry matter consumed in the present day (2015–2020 average). In Hamilton et al. (2025), the present-day bias correction used GFED4s emission data, but for this project, the dry-matter-consumed bias correction is being redone with GFAS4HTAP so that the absolute changes in BB emissions for the historical and future time period would be consistent. The bias correction was changed from emissions to dry matter to also allow the same GFAS4HTAP fire emission factors to be applied.

Note that in Tang et al. (2023) and Romanello et al. (2023), the Community Earth System Model 2 (CESM2) was used to project future burned area and total fire carbon emissions under different climate scenarios; however, it is only based on the one ESM, so it is not recommended for this project. As mentioned in Sects. 2.4 and 3.2.4, applying fire management policies to future scenarios is beyond the scope of this study. However, further information on that topic is included in the Sect. S3.3.

4.2.4 Other emissions

Aside from the emissions mentioned above, models typically include biogenic and geological emissions from natural sources. These can include isoprene and other VOC emissions from vegetation, NO_x emissions from soil microbes and lightning, and sulfur emissions from volcanos. Most models rely on the same interactive biogenic emission database, namely MEGAN (available at <https://bai.ess.uci.edu/megan>, last access: 16 May 2025; Guenther et al., 2012) (Table A2) or a derivative thereof.

4.2.5 Methane emissions and concentrations

While a few models are able to simulate CH₄ from emissions, it is common in many participating models to prescribe CH₄ concentrations instead. For emissions-driven models, BB emissions of CH₄ are available (Sect. 5.2) for the historic and future time periods, and anthropogenic CH₄ emissions are included in the IIASA GAINS emissions. However, CH₄ is not included in the historical HTAPv3.1 anthropogenic emissions. For concentration-driven CH₄ models, the future surface concentrations of CH₄ will be created by MSC-W (Met Norway) based off of the IIASA GAINS emissions and a climate model and provided to participants. The historic CH₄ concentrations are provided as annual average surface mixing ratios from the NOAA Global Monitoring Laboratory (Thoning et al., 2022), which are based on measurements from the NOAA Global Monitoring Laboratory,

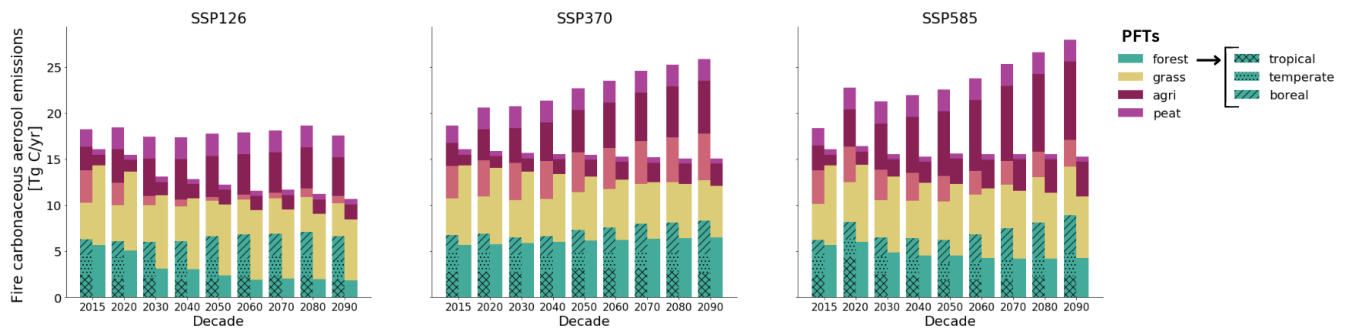


Figure 2. Decadal average time series of future fire aerosol emissions from Hamilton et al. (2025). Right-hand bars are fire emissions from interactive ESMs, and left-hand bars are fire emissions from CMIP6, based on land use change only.

which operates a network of background monitoring stations in remote locations around the world.

4.3 Available meteorological inputs

The height reached by a smoke plume, its horizontal transport, vertical mixing, and subsequent impact on a region are greatly determined by the prevailing weather conditions. These effects occur across a wide range of scales, from turbulent mixing of pollutants in the boundary layer, to lifting into the free troposphere, and subsequent transport by the prevailing winds. Observation-based reanalysis datasets provide an important source of meteorological information needed to drive some of the models (included in Table A2), but differences between these products, as well as between reanalyses and model-generated meteorology, provide an additional source of uncertainty (e.g. Adebisi et al., 2023).

4.3.1 Historical meteorological datasets

Currently, several meteorological reanalysis datasets are available and could be utilized, such as MERRA2 (Modern-Era Retrospective Analysis for Research and Applications, version 2), ERA5, NCEP-NCAR (National Centers for Environmental Prediction – National Center for Atmospheric Research), or JRA-55 (Japanese 55-year Reanalysis). They are summarized in Table S2.

4.3.2 Future meteorological input

To assess the alterations in meteorological conditions across the 21st century and their potential implications on fires (frequency, intensity, transport), the CMIP6 multi-model ensemble is the best available source for future meteorology that would occur in the changing future climate. The IPCC (Intergovernmental Panel on Climate Change) defined the Shared Socioeconomic Pathway (SSP) future scenarios, which illustrate different potential pathways for societal development throughout the 21st century and analyze their potential impacts on greenhouse gas emissions. The SSPs are classified into five trajectories: SSP1 represents a sustainable world,

SSP2 outlines a moderate pathway, SSP3 depicts a fragmented world with considerable challenges, SSP4 illustrates a world emphasizing equality and sustainability, and SSP5 envisions a world driven by rapid economic growth and dependence on fossil fuels. These five categories define different SSP emissions and concentration pathways, providing unprecedented detail of input data for climate model simulations: SSP1 (1.9 and 2.6), SSP2-4.5, SSP3-7.0, SSP4 (3.4 and 6.0), and SSP5 (3.4-OS and 8.5). The SSPX-Y scenarios refer to the estimated RF levels at the end of the 21st century; for instance, the “1.9” in the SSP1-1.9 scenario signifies an estimated RF level of 1.9 W m^{-2} in 2100.

The SSP emissions and concentration pathways are used as input for freely running ESMs, which then simulate future meteorological conditions for the Coupled Model Intercomparison Project Phase 6 (CMIP6) (Eyring et al., 2016). Access to the meteorological fields generated by these ESMs under each of the specified SSP scenarios is facilitated through platforms such as those provided by CMIP6 (<https://pcmdi.llnl.gov/CMIP6/>, last access: 16 May 2025), the IPCC Data Distribution Centre (DDC) (<https://www.ipcc-data.org/>, last access: 16 May 2025), and the Climate Data Store (CDS) by Copernicus (<https://cds.climate.copernicus.eu/>, last access: 16 May 2025). It is generally believed at the present that SSP2-4.5 is the most likely future scenario (Gidden et al., 2019; Meinshausen et al., 2020), and this one was chosen as the basis for the HTAP3 OPNS project.

4.4 Observational data available for model evaluation

The comparison of model results to observations is valuable for assessing how well models represent the real world and is critical for identifying gaps in our current understanding or weaknesses in how key processes are represented in models. Given the known uncertainties in fire and other model processes, observational comparisons provide a valuable opportunity to critically assess current parameterizations and identify which are most appropriate under particular conditions.

Comparisons with satellite-derived atmospheric composition will enable large-scale simulations to be evaluated consistently over the historical period under consideration. Observations of trace gases such as CO and aerosol properties can be used to evaluate long-range transport simulation (transport pathways, altitude, plume vertical extent and dilution). All surface monitoring measurements of the pollutants of Sect. 3.1 could be used for model evaluation, but we focus the rest of this section on highly relevant fire-specific observational datasets and field campaigns. See also Table S1 in the Supplement for a non-exhaustive list of relevant observational datasets. Note that the simulation time periods are chosen based on the prevalence of fire-relevant observational data.

As shown in Sect. 3.1, no single tracer is emitted by wildfires only, and domestic wood burning has the same signature as wildfires. Enhanced Hg and POP concentrations are also often observed within biomass-burning emissions, and come from the burned matter itself as well as being reemitted from soil. If those concentrations are enhanced simultaneously with those of other primary pollutants like CO, BC, and SOA, it is a strong indication that wildfire emissions are observed (e.g. Eckhardt et al., 2007).

For detecting wildfire plumes in observations, statistical methods use a combination of different trace species. For example, SO₄, BC, CO, and NO₂ have been combined with a positive matrix factorization to identify biomass-burning plumes (Karl et al., 2019). Yttri et al. (2024) used aerosol absorption coefficients recorded at different wavelengths by an aethalometer to distinguish BC emitted by fossil fuel or by biomass burning. Those observations are available for several stations in Europe.

Evaluation of modelled fluxes (e.g. deposition) is more challenging. These, as well as Nr impacts, may also be dynamically modelled in some systems. Cross-disciplinary satellite and in situ data (atmospheric, land surface, water quality, etc.) can be used to evaluate the modelled deposition results, helping identify weakness in individual models and reduce uncertainty in impact assessments (e.g. Fu et al., 2022; Huang et al., 2025).

4.5 Experiment design and sensitivity analyses

This section outlines different model experiments to help answer the science policy questions of Sect. 2. These fall into several distinct sets, targeting different aspects of our understanding, and some include a range of subexperiments to explore specific aspects in greater depth. Model groups may contribute to any number of experiments but are not required to complete them all. Where applicable, we indicate in Sect. 5.4 which experiments are higher priority for HTAP and which experiments may be dependent upon completion of other experiments.

4.5.1 How well do models perform? Baseline and case study simulations

Models should conduct baseline simulations of recent historical conditions, with a common set of anthropogenic and fire emissions, as both a basis of comparison for perturbation and sensitivity experiments and for general model inter-comparisons and evaluation with observations. The results can then be used to quantify the uncertainties and variability in atmospheric modelling. As the type of models participating is highly variable, with a range of computational costs, both short and long time periods are suggested for the baseline simulations. These time periods are selected based on the availability of reliable emission assessments and periods with abundant observations. Very computationally expensive models (e.g. very high resolution, inclusion of complex atmosphere chemistry) may only be able to simulate 1 year or less. Given how highly regional and interannually variable fires are, we can identify short-term fire case studies for evaluation of those models and explore particular fire events in detail. Fire event case studies can include the particularly large Australian fires of 2019–2020 (Filkov et al., 2020; Johnston et al., 2021; Collins et al., 2021; van der Velde et al., 2021; Anema et al., 2024), the fires in the USA that coincided with the 2018 WE-CAN and 2019 FIREX-AQ measurement campaigns (Juncosa Calahorrano et al., 2021; Warneke et al., 2023), and the significant fire season in Indonesia in 2015 due to a strong El Niño (Chen et al., 2016; Nechita-Banda et al., 2018).

4.5.2 What is the magnitude of pollution that comes from fires? Source–receptor/emissions perturbation experiments

To determine the magnitude of pollution from fires, species concentrations from baseline simulations can be compared to simulations with fire emissions removed. For additional detail, fire emissions from different geographical regions and from different types of burning can be perturbed for separate species, locations, and seasons to quantify source–receptor relationships and their uncertainties. However, the number of perturbation experiments can increase rapidly, so care is needed to prioritize and not define regions and sectors too finely.

Geographical regions

Coarsely defined regions help reduce the number of perturbation simulations. Figure 3 shows several options for the geographical source regions that were considered for these experiments, including those used within the HTAP2 multi-model experiments (Fig. 3a). While those distinguished boreal fires in higher latitudes from the low-latitude fires were associated with agricultural, temperate, and grassland regions, further refinement is needed for HTAP3 Fires. Re-

regions used for anthropogenic emissions perturbation experiments in the HTAP3 OPNS project are shown in Fig. 3b. We note that Southern Hemisphere Africa has been a focus of recent field campaigns (Zuidema et al., 2016) as the region emanates a third of the world's carbon from biomass-burning aerosol (van der Werf et al., 2010). South America also emits a significant fraction of the world's total BB aerosols. Therefore, for global modelling completeness, the scientific modelling community would benefit from including those southern hemispheric regions in HTAP3 Fires perturbation experiments as well. Figure 3c shows the GFED BB emissions regions used in many analyses that balance political regions and fire-relevant biomes. However, there are 14 GFED regions, which would be costly to run. Therefore, we merge the 14 GFED regions into 8 larger regions to make perturbation experiments (experiment 5 in Table 5) more feasible in Fig. 3d. These merged regions are broadly consistent with the HTAP2 regions, but with improved coverage, and are loosely aligned with the regions used for anthropogenic emissions in HTAP3 OPNS. Regional models may have geographical domains that differ from these, and where possible, these should simulate a subset of the regional perturbation experiments.

Fire sectors

Management decisions and policies are best informed by perturbing biomass-burning sectors separately. The two main categories are agricultural burning and wildland fires. Agricultural biomass burning is the deliberate burning of agricultural waste products, such as crop waste products, stubble, and other organic matter left in fields after harvest, as a method of waste disposal or as a practice in land management. The burning of grasslands towards coaxing new growth is also included. Deliberate burning is frequently applied in agricultural areas, especially where traditional practices are still widely practised. The United National Economic Commission for Europe (UNECE) adopted a guidance document on how to define and build policies around reducing open agricultural burning (UNECE, 2023).

Perturbing emissions from these 2 sectors separately over the 9 regions (8 regions + all) implies that global models participating in perturbation experiments would have 18 simulations to run. Figure 4 shows the distribution of the dominant fire types, which was developed for the application of fire type-specific FRP-to-dry matter burnt conversion and smoke constituent emission factors. It is based on the original GFAS fire type classification and spatial maps of ESA CCI Land cover (ESA, 2017) and PEATMAP (Xu et al., 2018). GFAS4HTAP will use a classification that is closer to the one of ESA CCI Land cover.

4.5.3 What is the impact of different fire processes? Process perturbation experiments

The following sensitivity studies perturbing key processes one at a time are meant to better understand and potentially reduce these key uncertainties.

Fire plume height

Most of the pollutants emitted from wildfires are released directly into the atmospheric boundary layer. However, depending on the meteorological conditions and the strength of the fire, material can be lifted well into the free troposphere or, in extreme cases, the stratosphere. This can have a substantial influence on the downwind impacts of the fire, as horizontal transport is typically faster in the free troposphere, the chemical processing of oxidants such as O_3 is typically more efficient, and the removal of pollutants by deposition processes is less efficient than in the boundary layer. Previous model studies have quantified the importance of injection height for key pollutants (e.g. Leung et al., 2007; Feng et al., 2024), but this has not been explored in a rigorous manner across a range of models. On longer timescales, the presence of high levels of BC in plumes can lead to local heating, which causes further lofting of the plume (e.g. Ohneiser et al., 2023). The altitude of tropospheric O_3 also influences the magnitude of its warming potential. Therefore, fire plume height introduces substantial uncertainty into assessment of impacts.

Some fire emission datasets (such as GFAS and GBBEPx) are based on FRP, whereas others, like GFED and FINN emissions, are based on burned area (BA). Both FRP and BA are (mainly) based on MODIS satellite observations. Daily information on wildfire injection heights, and/or FRP (fire radiative power), in combination with meteorological parameters, can be used in the calculation of the vertical distribution of fire plumes. Daily fire emissions based on FRP and BA differ substantially on a daily basis. Some fire emissions datasets, such as GFAS, provide injection height parameters based on satellite-observed FRP and available meteorological parameters (Rémy et al., 2017).

Some models represent plume rise in their simulations, while other models do not. Among the models that address plume rise, some include online parameterization of fire plume rise. For example, the Freitas scheme (Freitas et al., 2007, 2010) calculates plume rise by solving a set of 1D differential equations vertically, while the Sofiev scheme considers the conservation of the heat energy (Sofiev et al., 2012, 2013). The Canadian Forest Fire Emission Prediction System (CFFEPS) contains a thermodynamically based fire plume height parameterization based on fire energy and neutral buoyancy (Chen et al., 2019). Other models use a simpler approach of a constant plume injection height climatology (e.g. Dentener et al., 2006; Val Martin et al., 2010, 2012), which usually depends on region, season, and vege-

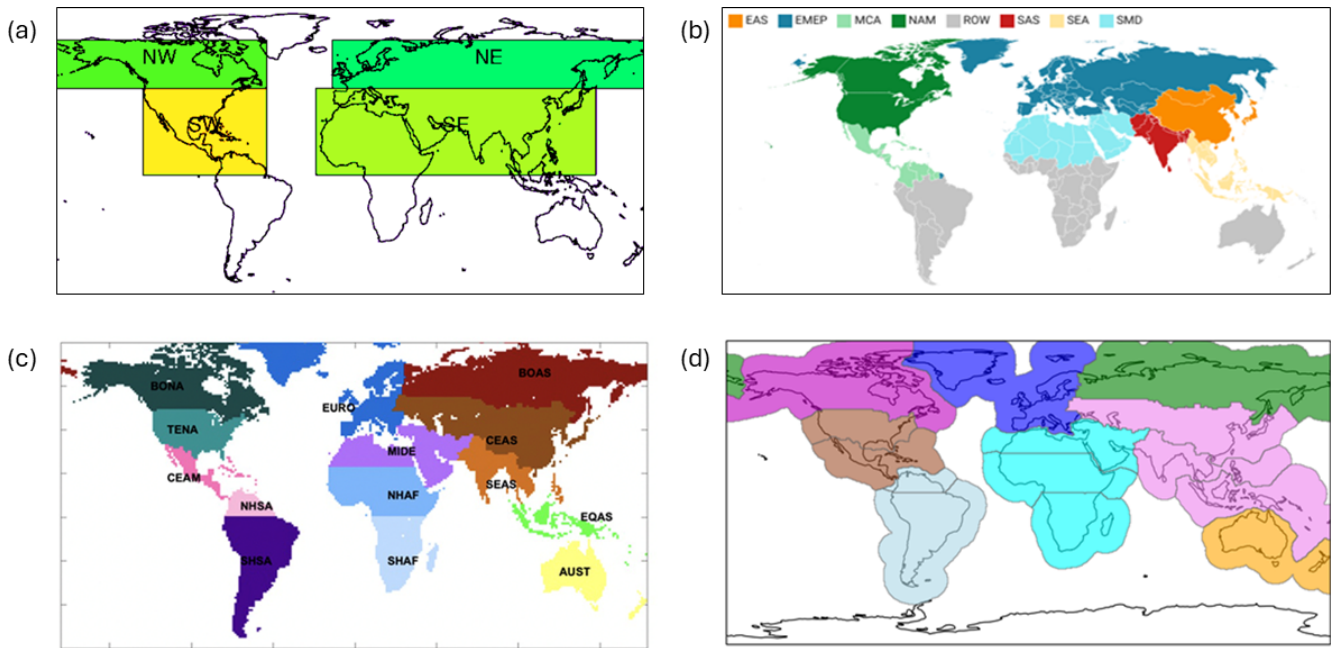


Figure 3. Regions for perturbation experiments: (a) BB source regions used in HTAP2 experiments, (b) Regions used for anthropogenic emissions in the HTAP3 O3PNS project, (c) GFED regions often used for fire emissions datasets, and (d) GFED regions (grey lines) and proposed merged regions (coloured areas). Panels (a)–(c) are those used in other comparable studies, and (d) is the choice for this study.

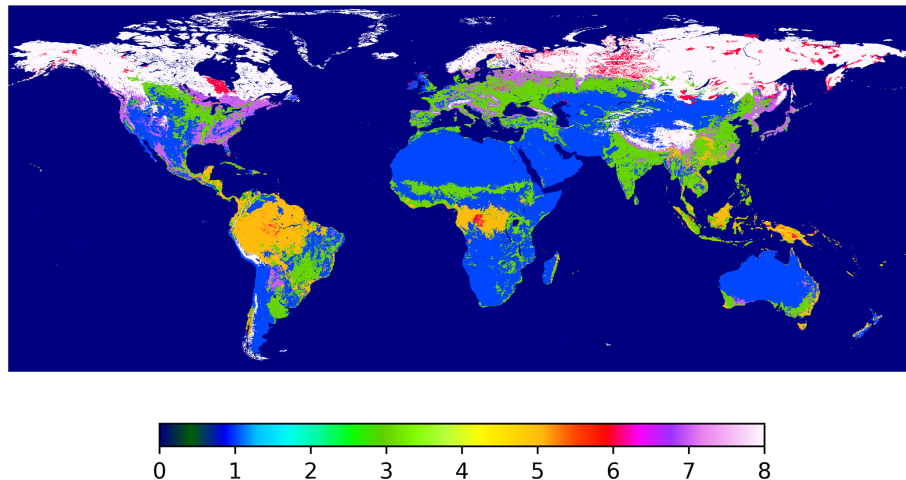


Figure 4. Land cover map for the GFAS4HTAP BB emissions for CAMS: 0 is water and ice, 1 is savanna, 2 is savanna organic soil, 3 is agriculture, 4 is agriculture organic soil, 5 is tropical forest, 6 is peat, 7 is extratropical forest, and 8 is extratropical forest organic soil.

tation type and does not consider FRP or fire size for specific fires. It is important to understand the impacts of different plume rise treatments on the model results, exploring the impacts of fuel type, moisture, and heat flux assumptions across the plume rise schemes used. Fire plume heights derived from CALIPSO, MISR, and TROP-OMI (Griffin et al., 2020) global satellite data and regional airborne instruments (Shinozuka et al., 2020; Doherty et al., 2022) could be used for quantitative model evaluation of fire plume height and vertical distribution.

Impacts of fire plume height were found to be different when looking at regional simulations versus at global climatological scales. In Field et al. (2024), using GFAS injection heights improved model performance at regional scales, whereas long-range transport patterns, influenced by the winds in the driving meteorology, mattered more than individual fire events at climatological timescales.

Fire plume chemistry

Biomass burning emits particles along with NO_x , nitrous acid (HONO), ammonia (NH_3), CO and CH_4 , and hundreds of VOCs, including a large number of oxygenated VOCs (OVOCs) (Jaffe et al., 2020). Representing this chemical complexity is a key challenge for modelling fire impacts on air quality, especially for secondary pollutants such as O_3 (Sect. 3.1.1) and SOA.

State-of-the-art atmospheric chemistry models typically overpredict O_3 close to fires but have difficulty simulating the influence of aged wildfire smoke plumes on downwind O_3 (e.g. Pfister et al., 2008; Singh et al., 2012; Zhang et al., 2014, 2020; Fiore et al., 2014; Lin et al., 2017; Baker et al., 2016; Jaffe et al., 2020). This may reflect (1) inaccurate fire emissions, especially underestimates of oxygenated VOC emissions from wildland fires (Arnold et al., 2015; Jin et al., 2023; Permar et al., 2023; Lin et al., 2024b); (2) lack of sufficient resolution or parameterization of smoke plume rise dynamics (Paugam et al., 2016); and (3) shortcomings in model representation of rapid photochemical processes in a concentrated smoke plume (Singh et al., 2012). Several modelling studies have shown strong sensitivity of O_3 production to differences in VOC chemistry, fire plume vertical transport, and NO_y partitioning (Zhang et al., 2014; Arnold et al., 2015; Lin et al., 2024b). Rapid conversion of NO_x to more oxidized forms typically reduces excessive ozone production simulated in near-fire smoke plumes. A recent study by Lin et al. (2024b) shows that sequestration of wildfire NO_x emissions in Canada as PAN enhances ozone production during smoke transport and thereby increases the impacts of Canadian wildfires on ozone air quality in US cities.

Additionally, large uncertainties in carbonaceous aerosol emissions from biomass burning (Pan et al., 2020; Carter et al., 2020; Xie et al., 2020) can also influence simulations through the impacts of aerosols on heterogeneous chemistry and photolysis rates. Further suggestions for model experiments to assess O_3 chemistry uncertainties appear in Sect. 5.3.1.

An additional challenge is the rate of SOA formation (Sect. 3.1.3). While SOA formation increases near-source, measurements taken after long-range free-tropospheric transport suggest SOA loss (Sedlacek et al., 2022; Dobracki et al., 2023), hypothesized to occur through heterogeneous oxidation primarily. Estimates of the reaction rates with OH vary, and measurements focused on constraining these rates would improve model depictions. OA loss is not included in many models (e.g. Lou et al., 2020, only consider photolysis, although their modelling construction could be using photolysis as a proxy for heterogeneous oxidation OA loss as well) but could be encouraged in the model output for this project.

Dry and wet deposition

Modelled dry and wet deposition fluxes are highly variable, uncertain, and a possibly significant cause for inter-model differences in pollutant concentrations. Models can test out different wet and dry deposition schemes and/or turn deposition on and off to quantify its impact. Deposition is also important for evaluating ecosystem impacts. Wet and dry deposition fluxes should be diagnosed from all model simulations.

4.5.4 How will fires and their impacts change in the future? Future scenario experiments

The frequency and severity of wildland fires are likely to increase within a warming climate, particularly in the Northern Hemisphere (van Wees et al., 2021). Quantifying the influence of these changes, given different future emission scenarios, is an important application of models (e.g. Xie et al., 2021). Future modelling experiments can be performed with chemical transport models that use provided future emissions and meteorology (see Sect. 5.2). Experimentation can also be performed by ESMs with and without interactive fire modules. ESMs can typically simulate future climate/meteorological conditions in a free-running state out to 2100. Experiments for future fires with both interactive ESMs and other atmospheric models driven offline will help determine the range of uncertainty on future fire projections and their impacts. While fire emissions are likely to change under the effects of changes in human management practices and policies, those aspects that are not already included in the CMIP SSP scenarios (Xie et al., 2021) are beyond the scope of this study due to a lack of scenario emissions datasets.

5 Recommended plan

5.1 Simulation periods

Given the combination of emissions dataset availability (Sect. 4.4) and existing observational datasets to compare against (Sect. 4.4), we suggest the following time frames for simulation years (Table 3). The short historical option for the HTAP3 OPNS and Hg projects was selected to be 2015. However, 2015 had a strong El Niño and was an extreme fire year in Indonesia as a result. Fires are so greatly variable on interannual scales that it would be unwise to base policy decisions on the analysis of a single year. We therefore encourage use of the medium historical option, which includes field campaigns of 2016–2018 that were offshore from African fires (Redemann et al., 2021; Haywood et al., 2021; Zuidema et al., 2018) and 2019, which had a field campaign in the USA. The medium option stops by the end of 2019 to avoid incorporating the complexity in anthropogenic emissions that arose with the COVID-19 pandemic in 2020. The medium future option includes 5 years on either side of the

2015 start and 2050 end dates of the GAINS future emissions to enable 10-year averages to be created around these start and end dates, thus accounting for interannual variability, consistent with the HTAP3 OPNS project. The 2015 emissions may be used for 2010–2014 and the 2050 emissions for 2051–2055. Finally, while the climate community routinely does simulations out to 2100, given that the GAINS anthropogenic emissions end in 2050, and the AerChemMIP2/CMIP7 community (see Table A1 and Sect. 3.3) will focus on future simulations, including climate impacts from fires, we have elected not to include a long future option within this study.

5.2 Inputs: emissions and meteorology

Based on discussions in Sect. 4.2, the following emissions datasets are recommended and summarized in Table 4 below. For methane, the emissions and concentrations described in Sect. 4.2.5 are recommended.

Historical fire emissions

The historical fire emissions datasets were carefully considered during and following a 4 d online workshop hosted by IGAC BBURNED in November 2023. The methodology, advantages, and disadvantages of each major global fire emissions dataset were discussed. It was agreed to recommend use of the fire emissions based on GFASv1.2 because (a) they provide daily emissions (providing improved temporal variability over monthly emissions); (b) they include peatland fires, including in the boreal region, the latter particularly important for the AMAP scientific community; and (c) they provide fire plume heights as well as speciated emissions. The GFASv1.2 emissions have since been updated with the most recent emission factors and land categories and are called GFAS4HTAPv1.2.1. We furthermore note that peat fire emissions remain highly uncertain and that these fire emissions do not include special treatment of WUI fires. We also note that modellers may apply these emissions at their preferred temporal resolution for baseline simulations.

Future fire emissions

The future fire emissions dataset that is derived from a multi-model ensemble that includes the influence of the changing climate on fires is that from Hamilton and Kasoar (Douglas Hamilton, personal communication, 2024, 2025). We recommend use of this future fire emissions dataset, but note that it does not include scenarios for future changes to fire management policy and practice, as these quantitative emissions adjustments are not available yet.

Historical anthropogenic emissions

These are chosen to be consistent with the other concurrent HTAP3 projects. They are the HTAPv3.1 anthropogenic

emissions, which were delivered in early 2025, for the years 2000 to 2020, inclusive. They include all relevant species except Hg, PAHs, and PCDD/Fs. The anthropogenic emissions for those species should be EDGARv8.1 (Hg), PKU-GEMS (PAHs), and PKU-LZU (PCDD/Fs), as discussed in Sect. 4.2.1.

Future anthropogenic emissions

For consistency with other HTAP3 projects, the CLE (current legislation) future emissions from IIASA GAINS (LR-TAP) will be used in future simulations. Climate modellers may wish to simulate out to 2100, and while the SSP2-4.5 anthropogenic emissions for 2015 to 2100 are available and are roughly equivalent to the GAINS CLE emissions scenario for CO₂ and energy, they are not necessarily similar for other pollutant emissions. We therefore recommend for this project ending the future simulation in 2050 and participating in CMIP7/AerChemMIP2 for longer future simulations.

Biogenic and other natural emissions

While it is useful to have consistent emissions across models, this can be difficult to achieve due to the dependence of natural emissions on structural aspects of models, including vegetation, soil, and land use. Therefore, we suggest that each modelling centre use their preferred emissions from biogenic and other natural sources. These should be documented and taken into consideration in the analysis.

Driving meteorology

As discussed in Sect. 4.3, there are several data reanalysis collections that could potentially be employed. Although the ERA5 collection offers greater spatial, temporal, and vertical resolution overall, any of the mentioned datasets would be suitable for use. It is recommended that modellers use ERA5 if possible but otherwise use their preferred meteorology for historical simulations and ensure that they document this clearly. For future simulations, as discussed in Sect. 4.3.2, we suggest using interannually varying, monthly mean sea surface temperatures and sea ice distributions from the SSP2-4.5 multi-model CMIP6 ensemble.

5.3 Model experiments

The following model experiments in Table 5 are proposed based on the discussions in Sect. 4.5, and further details on selected experiments are described below.

5.3.1 Details on fire process perturbation experiments (experiment 6)

While a short time range for perturbation experiments can help keep model simulations manageable, they may not provide generalizable results, given the high interannual vari-

Table 3. Simulation time periods, with options for different types of models.

	Short option	Medium option	Long option
Historical	See case studies (Sect. 4.5.1 and Table 5)	2015–2019	2003–2020
Future	2045	2010–2020 and 2040–2050	2010–2050

Table 4. Emissions inputs for model experiments. See “Data availability” for more information.

Emission type	Recommendation	Notes
<i>Historical simulations (2003–2020)</i>		
Fire	GFAS4HTAP for BB, including agricultural burning	Daily gridded global 0.1° resolution. Including its agricultural burning emissions.
Anthropogenic	HTAPv3.1	Minus its agricultural burning.
<i>Future simulations (2010–2050)</i>		
Fire	GFAS4HTAP for 2010–2020 and Hamilton et al. (2025) for 2020–2050	SSP2-4.5-scenario-based climate-influenced future fire emissions, calibrated to GFAS4HTAP historical fire emissions (both <i>not</i> including agricultural burning).
Anthropogenic	IIASA GAINS CLE	Including agricultural burning
<i>Biogenic and other natural emissions: each modelling centre use their default</i>		

ability of fires. Therefore, the time ranges of Table 3 should be followed for perturbation experiments as well.

Injection height

Repeat experiment 1, but with alternative fire plume height schemes. We suggest the following options, where modellers can opt into any number of these when possible:

- model’s default fire plume height system, whatever it may be;
- FRP-based plume heights provided by GFAS (Sect. 4.5);
- climatological plume heights from AeroCom (Dentener et al., 2006), assuming standard vertical profiles; and
- no plume rise, assuming all pollutants are released into the lower part of the planetary boundary layer.

Chemistry

To assess the impacts and uncertainties around fire plume chemistry, a few sensitivity runs are recommended:

- partitioning total NO_y emissions from biomass burning into PAN (37 %), HNO_3 (27 %), and NO_x (36 %), rather than emitting only NO in the baseline simulation (experiment 1), as recommended by Lin et al. (2024a, b)

based on recent aircraft measurements (WE-CAN 2018 and FIREX-2019);

- doubling BB emissions of all NMVOCs, including formaldehyde and acetaldehyde producing acetyl peroxy radical (CH_3CO_3) for PAN formation;
- for models with suitable capability, exploration of the effects of different levels of complexity in VOC chemistry or differences in volatility or reactivity of VOC;
- increasing BB emissions of OC and BC aerosols by 50 % to explore their impacts on oxidative chemistry through heterogeneous chemistry or photolysis.

Emissions temporal resolution

As a repeat of experiment 1 with hourly, daily, and monthly versions of the fire emissions to quantify the importance of temporal resolution, many previous major studies, such as CMIP6, have used monthly fire emissions, and this sensitivity study will allow these results to be put into context.

Meteorology

Using repeating annual meteorology for 2018 with interannually changing emissions will determine how much of the interannual variability in impacts seen in experiment 1 is due to meteorology and not emissions.

Table 5. Model experiment types.

Experiment name	Description	Purpose	Priority
1. Baseline simulation	Historical time period(s) given in Table 3. Common set of emissions given in Table 4.	Model evaluation; baseline for subsequent sensitivity and perturbation experiments.	High.
2. Case study/studies	More detailed, specific fire events (Indonesia 2015; North America 2018, 2019; and Australia 2019–2020) at higher spatial and temporal resolution.	Model evaluation.	High for regional models. Low for global models.
3. Fire emissions sensitivity	Same as experiment 1, but driven by different sets of fire emissions (GFED, FINN, etc.).	Model/emissions evaluation; to gauge differences between fire emission datasets across models.	Low.
4. Prescribed future fires	Future time period(s) given in Table 3. Future emissions given in Table 4.	To determine how wildland fires and their impacts will change in the future.	High.
5. Regional and sectoral emissions perturbations	Turn off all BB emissions for all species everywhere. Turn BB emissions off in each region of Fig. 3d, and each of the 2 sectors: agricultural burning and wildland fires, over the historical time periods in Table 3.	To quantify regional source–receptor relationships and uncertainties.	High for both fire sectors combined. Low for separate sectors.
6. Fire process perturbations	Parameter/process perturbations, for fire plume height, chemistry, emissions, and meteorology (see Sect. 5.3.1). Short-to-middle time periods of Table 3.	To determine importance of different processes and impacts of different model fire parameterizations.	Medium.
7. Interactive fire modules	Historical and future simulations (Table 3) with coupled land–atmosphere models.	To determine how wildland fires will change in the future with an interactive climate and compare to experiment 4 results.	Medium.
8. Data assimilation	Inverse modelling to combine CTMs with observed atmospheric volume mixing ratios of CO (MOPITTv9), O ₃ (OMI), NO ₂ (OMI). See Table 6.	Infer surface–atmosphere emissions/fluxes.	Low.

5.3.2 Details on future experiments (experiment 4)

The SSP2–4.5 future climate scenario will be the driver for the future time period, which includes those future fire emissions from Hamilton et al. (2025) and the GAINS CLE anthropogenic emissions. For future agricultural burning emissions, which appear in both the BB and the anthropogenic emissions datasets, we recommend that the GAINS future agricultural burning emissions be used and those removed from the BB emissions so as not to double-count them.

5.3.3 Details on data assimilation experiments (experiment 8)

Models that can do data assimilation of observations are recommended to assimilate measurements depending on the period of interest (see Table 6), in order to constrain CO, NO₂, or O₃ fire prior emissions used in the baseline experiments.

Satellite observations to be considered are the Measurement of Pollution in the Troposphere (MOPITT; Deeter et al., 2003; Edwards et al., 2006) CO measurements. Since MOPITTv9 has overpasses only every 16 d with a resolution of 25 km, we recommend assimilating the Tropospheric

Table 6. Measurements to be assimilated depending on experiment and period of simulation.

Gases		Long option (2003–2020)	Medium option (2015–2019)	2015 Indonesian fires	2019–2020 Australian fires
CO	Sensor, satellite		MOPITT, TERRA		TROPOMI, Sentinel 5P
	Version		Level 2 version 9		Level 2 v2.4.0
	Reference		Deeter et al. (2022)		Apituley et al. (2022)
NO ₂	Sensor, satellite		OMI, AURA		TROPOMI, Sentinel 5P
	Version		Level 2 v3		Level 2 v2.4.0
	Reference		Lamsal et al. (2021)		Eskes et al. (2022)
O ₃	Sensor, satellite		OMI, AURA		TROPOMI, Sentinel 5P
	Version		Level 2 v3		Level 2 v2.4.0
	Reference		Veefkind (2012)		Romahn et al. (2022)

Monitoring Instrument (TROPOMI; Veefkind et al., 2012) Level 2 measurements, launched in fall 2017, which has daily global coverage and fine resolution ($5 \times 7 \text{ km}^2$ for CO at nadir) for the case study simulation of Australia. For some CTMs at coarse spatial resolution (even at $1^\circ \times 1^\circ$ spatial resolution), and because of the fine resolution of TROPOMI, large computational costs can arise. It is then possible to use TROPOMI super-observations (area-weighted average of the pixel at coarser resolution) following the approach of Miyazaki et al. (2012). For NO₂ and O₃ long- and medium-term inversions, observations such as from the Ozone Monitoring Instrument (OMI; Levelt et al., 2006) with data available since 2005 can be used. Surface measurements can be considered to constrain background concentrations, particularly for regional emission estimates (such as the nested zoom model TM5 and GEOS-Chem). The assimilation here will help to determine how the GFAS4HTAPv1.2.1 emissions would be constrained. Assimilation algorithms (3Dvar, 4Dvar, or ensemble Kalman filter), as well as prior and observation uncertainty, and OH fields do not have specific recommendations, and each user is free to choose these parameters.

5.4 Model outputs

To maximize accessibility of the results, the model output data request for this project is based on the AerChem-MIP tables from CMIP6, as adopted by the HTAP OPNS project, with some additions for the extra species and impacts. The table of model outputs is located online at <https://nextcloud.gfz-potsdam.de/s/sp8XmMY2rQizjA4> (last access: 16 May 2025) surface NO₂, PM_{2.5}, and O₃, as well as hourly O₃ deposition parameters needed for air quality, health, and ecosystem impact analysis, in addition to monthly radiative flux output for climate impacts. When measurements and impacts are only related to surface concentrations (e.g. POPs), we have suggested only surface-level 2D model output be provided to save storage space.

Data workspace

The model output can be uploaded to METNO's AeroCom database and infrastructure as part of the HTAP3 component of the AeroCom database. Instructions for obtaining access to the AeroCom user server, formatting, uploading, downloading are found here: https://aerocom.met.no/FAQ/data_access (last access: 16 May 2025). The AeroCom database infrastructure is available to host HTAP model data on a read-only permanent database, which can be accessed by authorized users with an account on the AeroCom user server. A scratch area on the AeroCom-user server can be used to upload data. Uploaded data can be transferred on demand by METNO to the read-only permanent database section for HTAP, under the directory HTAP-PHASE-III.

5.5 Post-processing and analysis

This multi-pollutant, multi-model experiment will generate a large amount of data that will be analyzed to answer the science policy questions of Sect. 2.

5.5.1 Model evaluation: comparison of experiments 1, 2, 3, and 6 to observations

By comparing the results of experiments 1, 2, 3, and 6 to the observations discussed in Sect. 4.4 (and listed in Table S1), specific model inputs and processes can be evaluated. Note that Table S1 is not an exhaustive list of measurements that may be used in model evaluation, and it also includes some measurements that may not overlap in time with the simulations. The aim of the evaluation would be to improve our understanding of fire processes such as plume rise and plume chemistry and improve their parameterizations in models. We may also be able to determine which inputs (emissions, meteorology) and parameterizations are best, as well as identify gaps that require further research. One example would be to analyse the impacts of injection heights on PAN concentrations in the free troposphere and downwind O₃ formation as fire plumes subside into the boundary layer, by compar-

ing the model simulations of PAN and related tracers to recent aircraft measurements. We suggest that, when possible, community tools like MELODIES (Model Evaluation using Observations, Diagnostics and Experiments Software) and ESMValTool be used for inter-model comparisons and evaluation against observations. Regardless, the evaluation will require a large effort by the community.

5.5.2 Assessing health impacts of fires

The most cited and widely used approaches of risk analysis are the following: all causes of deaths, mortality and morbidity impacts, emergency hospitalization, reduced life expectancy, premature mortality, incremental lifetime cancer risk, and health-related cost of air pollution (Goel et al., 2021; Sonwani et al., 2022; Nagpure et al., 2014; Gidhagen et al., 2009; Guttikunda and Kopakka, 2014; Ghozikali et al., 2014; Farzaneh, 2019). Human health risk assessment is the mathematical estimation and modelling of several processes, including population estimates, population exposure to pollutants, and adverse health impact assessment through specific concentration–response functions (WHO, 2021). Widely used quantitative health risk assessment tools of different agencies are listed in Table S3, while Table S4 represents the comparison between the air pollution health risk assessment tools (methodologies, scopes, input parameters, and predicted health impacts). The surface-level model outputs of atmospheric composition at high spatial (e.g., 10 km for global, 1 km for regional) and temporal (monthly down to daily) resolution will be invaluable for new health risk assessments, especially when fused with other modelling (e.g. land use regression) and observational (e.g. remote sensing) techniques (Johnson et al., 2020).

5.5.3 Assessing climate impacts of fires

Climate impacts can be assessed through the RF from fire-emitted pollutants by comparing the differences of the radiative fluxes of the simulations with and without fire emissions (i.e. the baseline simulation and the regional and sectoral emissions perturbations). To assess the component-specific RFs, more detailed simulations with source attribution techniques, such as those for O₃ (Grewe et al., 2017; Butler et al., 2018) or for aerosols (Righi et al., 2021), are helpful. Models capable of such possibilities therefore perform additional pollutant-specific perturbations, including source attribution techniques. Moreover, the model's composition fields can be applied in offline radiative transfer models or via the kernel method to calculate the component-specific RF. These should be included in the regional perturbation experiments in order to have the required data to assess the RF impacts of biomass burning. For estimation of future fire RFs, it is key to quantify the effects caused by the nonlinearities on the O₃ RF and for the aerosol–cloud interactions.

5.5.4 Assessing future vs. historical changes

While the future BB emissions are calibrated to the historical emissions, the future anthropogenic emissions are not calibrated to the historical anthropogenic emissions. Therefore, analysis of the full historical to future time period (2003–2050) should be done with care: the absolute, but not relative, changes to BB results can be assessed for 2003–2050. However, the future versus present changes (absolute and relative of 2040–2050 vs. 2010–2020 for example) should be assessed from simulations using only the future anthropogenic emissions input for consistency (see Table 3).

6 Conclusions

In this paper we have described the need for a multi-model, multi-pollutant study focused on fires and highlighted a range of important science policy questions arising from discussions with the scientific and policy communities that this study is intended to answer. The study will address gaps in our current scientific understanding of fire processes and provide a more robust quantification of fire pollution and its impacts to inform decision-making. We have thoroughly discussed the scope of this study (Sect. 3), based on extended consultation with the science, impact, and policy communities, and have outlined a number of model design options that were considered (Sect. 4), with the ultimate choices justified. We then provide the recommended specifications for the modelling study (Sect. 5) to be carried out over the next ~ 3 years that will provide maximal benefit for the scientific community and for key policy-adjacent communities, including HTAP and AMAP. HTAP3 Fires is aimed at providing fresh understanding of the atmospheric and environmental impacts of fires and providing the foundation for sound policy decisions

Appendix A

Table A1. Complementary fire-related research activities.

Name	Objective	Website	Notes
Biomass Burning Uncertainty: Emissions, ReactionNs, and Dynamics (BBURNED)	To coordinate fire research community towards better understanding fire variability and uncertainty, particularly as it relates to atmospheric chemistry	https://www2.acom.ucar.edu/bburned (last access: 16 May 2025)	An International Global Atmospheric Chemistry (IGAC) activity
Arctic Monitoring and Assessment Programme (AMAP)	Inform the Arctic Council through science-based, policy-relevant assessments regarding pollution and climate change issues	https://www.amap.no/about (last access: 16 May 2025)	Expert groups on SLCFs, POPs, Hg, local vs. long range. SLCF EG may use these HTAP experiments for a future AMAP report.
Arctic Black Carbon impacting on Climate and Air Pollution (ABC-iCAP)	Creation of fire management scenarios for Arctic Council countries/states	https://abc-icap.amap.no/ (last access: 16 May 2025)	
WMO Vegetation Fire Smoke Pollution Warning Advisory and Assessment System (VFSP-WAS)	To enhance the ability of countries to deliver timely and quality vegetation fire and smoke pollution forecasts, observations, information, and knowledge to users through an international partnership of research and operational communities	https://community.wmo.int/en/activity-areas/gaw/science/modelling-applications/vfsp-was (last access: 16 May 2025)	
International Association of Wildland Fire (IAWF)	Organizes large-scale conferences around wildfire	https://www.iawfonline.org/ (last access: 16 May 2025)	
Integrated Land Ecosystem-Atmosphere processes Study (iLEAPS)	Recently conducted a meeting on fires in South Asia, focusing on the prescribed fires and their modelling and planning the next workshop in March. Carry out the conversion on the prescribed fires and their impact on air quality and health and modelling, including fire emission estimate.	https://ileaps.org/future-earth (last access: 16 May 2025) and https://www.tropmet.res.in/204-event_details (last access: 16 May 2025)	

Table A1. Continued.

Name	Objective	Website	Notes
Arctic Community Resilience to Boreal Environmental change: Assessing Risks from fire and disease (ACRoBEAR)	To predict and understand health risks from wildfire air pollution and natural–focal disease at high latitudes, under rapid Arctic climate change, and resilience and adaptability of communities across the region to these risks.	https://www.acrobear.net/ (last access: 16 May 2025)	Integrating health data and knowledge, community knowledge and stakeholder dialogue, with satellite and in situ observations, and numerical modelling.
Air Pollution in the Arctic: Climate Environment and Societies (PACES)	Review existing knowledge and foster new research on the sources and fate of Arctic air pollution and its impacts on climate, health, and ecosystems	https://igacproject.org/activities/PACES (last access: 16 May 2025)	IGAC/IASC initiative: Improving knowledge of high-latitude forcing from fire emissions. Key questions around ageing of fire plumes, mixing with anthropogenic pollution following export (e.g. POLARCAT cases). NO _y speciation, BC ageing, how these vary between models. Need for improved observational constraint on these processes
AeroCom		https://aerocom.met.no (last access: 16 May 2025), and more specifically e.g. https://aerocom.met.no/node/110 (last access: 16 May 2025) and https://aerocom.met.no/node/115 (last access: 16 May 2025)	
The Fire Model Intercomparisons Project (FireMIP)	Systematic examination of global fire models, which have been linked to different vegetation models. Relevant for ESM/coupled fire simulations.	https://gmd.copernicus.org/articles/10/1175/2017/ (last access: 16 May 2025)	ISIMIP3 (the Intersectoral Impacts MIP phase 3), which FireMIP is now merging with – ISIMIP3b (simulations currently in progress) will produce multi-model projected future fire emissions for different SSP scenarios
Support for National Air Pollution Control Strategies (SNAPCS)	Part of this project involves investigating the impact of local and long-range transport of fire-related pollutants on the UK. There is particular interest in implications for health/air quality and model development		Project involvement from the UKCEH, Imperial College London, EMRC, and DEFRA

Table A1. Continued.

Name	Objective	Website	Notes
European Network on Extreme fire behaviOr (NERO)	Bringing together wildfire researchers and practitioners to advance the current state of the science, thus making a crucial step in improving fire management, firefighter training and safety, and public safety planning. Science-based wildfire management.	https://www.cost.eu/actions/CA22164/ (last access: 16 May 2025)	European Cooperation in Science and Technology (COST) action
FLARE: Fire science Learning AcROSS the Earth system (workshop)	The goal is to develop a roadmap for coordinated wildfire research for the next 5–10 years.	https://futureearth.org/initiatives/funding-initiatives/esa-partnership/ (last access: 16 May 2025)	Held 18–21 September 2023. Article: https://futureearth.org/2023/12/13/ (last access: 16 May 2025)
AerChemMIP2 in CMIP7	Historical and future climate change simulations focused on aerosols and trace gas chemistry for CMIP7.		Will include a focus on wildland fires and biomass burning. Simulation design in 2024.

Table A2. Model characteristics of potential participants. PI – principal investigator(s), BC/IC – boundary/initial conditions. Model types (see Sect. 4.1 for abbreviations).

Organization	PI	Model(s)	Type	Species	Domain	Resolution (spatial and temporal)	BC/IC
(a)							
University of Hertfordshire, Centre for Climate Change Research	Ranjeet S Sokhi	WRF/CMAQ, WRF-Chem	CTM	PM _{2.5} (components), O ₃ , NO ₂	Europe, CORDEX	5–10 km over Europe, hourly, daily, monthly (for future projections)	CAM-chem, ECMWF, GFS
Environment and Climate Change Canada Climate Research Division	Cynthia Whaley, Knut von Salzen, David Plummer, Vivek Arora	CanAM-PAM, CMAM, CanESM, CLASSIC (global & regional)	ESM and CCM	PM _{2.5} , O ₃ , NO _x , CH ₄	Global, Arctic, North America	T64, typically 3-hourly to monthly output	CanESM provides BC/IC for CanRCM
NSF National Center for Atmospheric Research Atmospheric Chemistry Observations and Modelling	Louisa Emmons, Rebecca Buchholz, Douglas Hamilton	MUSICAv0 (CESM/CAM-Chem)		O ₃ , NO _x , CO, VOCs, PM _{2.5} and speciated aerosols, metals	Global, USA	12 km over USA (and other regions), 1° global	
NASA Goddard/University of Maryland	Min Huang	WRF-Chem (NASA version, LIS)	ESM	O ₃ , PM, CO, NO _x , VOCs	Eastern USA	10 km or finer	CAMS, CAM-Chem/WACCM
NASA Goddard Institute for Space Studies	Keren Mezuman, Kostas Tsigaridis	NASA GISS ModelE	ESM	CO ₂ , CO, HCHO, CH ₄ , acetone, alkenes, paraffin, SO ₂ , NO _x , NH ₃ , aerosols/PM _{2.5} (organic, black carbon)	Global	Cubed sphere 1 × 1 effective resolution. 30 min to monthly	
Cyprus Institute Climate and Atmosphere Research Centre	Theo Christoudias	EMAC, WRF-Chem	CTM	PM (including black carbon), CO, O ₃	Global, Middle East, North Africa	20 km over Middle East (2 min), 1° global (15 min)	GFS/WACCM

Table A2. Continued.

Organization	PI	Model(s)	Type	Species	Domain	Resolution (spatial and temporal)	BC/IC
University of Bremen, Institute of Environmental Physics	Nikos Daskalakis, Sarah-Lena Meyer, Mihalis Vrekousis	TM5-MP		PM _{2.5} , O ₃ , NO _x , NO ₂ , CO, CH ₄ , VOCs, OA, speciated aerosols	Global	1° × 1°; 3-hourly to monthly	
UKCEH/Edinburgh University, Atmospheric Chemistry and Effects/School of Chemistry	Damaris Tan, Stefan Reis, Mathew Heal, Massimo Vieno, Eiko Nemitz	EMEP4UK, EMEP MSC-W WRF	CTM	PM _{2.5} and components	Global, Europe, UK	1 or 3 km over UK, 27 km over Europe, 1° global resolution. Hourly output.	WRF with GFS/ERA5 reanalysis
UK Met Office, Hadley Centre	Steven Turnock, Gerd Folbert, Joao Teixeira	UKESM1+ INFERNO	ESM	PM _{2.5} , O ₃ , CH ₄	Global	Global grid at 1.25 × 1.75° resolution (~ 140 km)	
Lancaster University, Lancaster Environment Centre	Oliver Wild	FRSGC/UCI CTM	CTM	O ₃ , NO _x , CO, VOC, CH ₄ , gas-phase oxidants	Global	Usually T42 (2.8 × 2.8°) but T106 (1.1 × 1.1°) feasible; output hourly/monthly	
Thailand Team (currently, King Mongkut's University of Technology Thonburi, Thammasat University)	Kasemsan Manomaiphiboon, Vanisa Surapipith	WRF-Chem	CTM	PM _{2.5} (primary/secondary), O ₃	Upper Southeast Asia (with focus on Lower Mekong Basin)	4–12 km; output hourly	
IITM Pune, India, AQEWS Urban Air Modelling	Rupal Ambulkar, Sachin D, Gaurav Govardhan	WRF-Chem	CTM	PM _{2.5} , PM ₁₀ , CO	India	10 km over India run with daily output	GFS
CICERO Center for International Climate Research, Oslo, Norway	Marianne Tronstad Lund	OlsoCTM3	CTM	O ₃ , NO _x , CO, VOCs, PM _{2.5} , and speciated aerosols	Global	2.25 × 2.25° (possibly 1 × 1° depending on no. of simulations/scope), 60 vertical layers. Monthly output, with option of 3-hourly	
DLR Institute of Atmospheric Physics, Earth system modelling	Mariano Mertens	EMAC, MECO(n)		O ₃ , NO _x , CO, VOCs, PM _{2.5} , and speciated aerosols	Global, Europe, West Africa		
NOAA Geophysical Fluid Dynamics Laboratory (GFDL)	Meiyun Lin	GFDL AM4VR		O ₃ , PAN, NO _y , CO, VOCs, PM _{2.5} , and speciated aerosols	Global, North America	~ 13 km over North America, 25–50 km over Europe, and 50–100 km over Asia. Monthly, daily	
MIT, Earth, Atmospheric and Planetary Sciences	Noelle Selin, Lexia Cicone, Eric Roy	GEOS-Chem	CTM	PAHs, Hg	Global	2 × 2.5, 47 vertical layers, monthly outputs	
Meteorological Research Institute, Japan Met Agency (MRI-JMA), Department Of Atmosphere, Ocean, and Earth System Modelling Research	Naga Oshima	MRI-ESM2, TL159	ESM	PM _{2.5} , speciated aerosols, O ₃ , NO _x , NO ₂ , CO, CH ₄ , VOCs	Global	AGCM: TL159; aerosol: TL95; ozone: T42	
Institut National Polytechnique Félix Houphouët-Boigny (INP-HB) Department of Forestry and Environment	Jean-Luc Kouassi			PM _{2.5} , O ₃ , NO ₂ , CO, CH ₄	Global, West Africa		
Space Research of the Netherlands, Earth Department	Helene Peiro, Ilse Aben, Ivar van der Velde	TM5-4DVar zoom	Inverse model	CO	Global, North America, Europe, Africa	Global 3 × 2, regional 3 × 2 and 1 × 1; daily to monthly	
Norwegian Meteorological Institute, Research Department	Jan Eiof Jonson	EMEP		O ₃ , NO _x , CO, NMVOCs, PM	Global, Europe	Daily to annual output	
NILU, Atmospheric Chemistry	Sabine Eckhardt, Nikolaos Evangelinou	FLEXPART	Lagrangian transport model	CO, BC	Global	0.5°, 3 h resolution	

Table A2. Continued.

Organization	PI	Model(s)	Type	Species	Domain	Resolution (spatial and temporal)	BC/IC
Finnish Meteorological Institute, Atmospheric Composition Research	Mikhail Sofiev, Rostislav Kouznetsov, Risto Hanninen, Andreas Uppstu, Evgeny Kadantsev	IS4FIRES-SILAM		O ₃ , NO _x , CO, VOCs, PM _{2.5} , and speciated aerosols	Global to local	Several options, e.g. global 5 d forecast 10 km for fire PM, 20 km for full AQ tropo + strato, 10 km Europe. Multi-annual reanalysis up to 50 km global	Global nested, CAMS
Tsinghua University, School of Environment	Shuxiao Wang, Bin Zhao, Yicong He, Lyuyin Huang	CESM, WRF-Chem	ESM and CTM	PM _{2.5} , O ₃ , NO _x , NO ₂ , CO, CH ₄ , VOC, OA	Global, southern China, and SE Asia	0.9 × 1.25 hourly, daily, monthly. 27 km overall domain with 9 km nested domain; hourly	
University of Augsburg, Faculty of Medicine	Christophe Knote, Bin Zhou	WRF-Chem	CTM	PM _{2.5} , OA	Europe	20 and 2 km nest, hourly	GFS, CAM-Chem
Jožef Stefan Institute/MSCE, Department of Environmental Sciences	Oleg Travnikov	GLEMGLEOS	Multi-media POPs (?)	POPs, Hg, metals	Global, Europe	1 × 1°, monthly or daily output	
Peking University, Lanzhou University	Jianmin Ma, Tao Huang	CanMETOP, CMAQ	Long-range atmospheric physical transport and CTM	POPs, heavy metals	Global, China, North America	from 10 km to 1° × 1°. Hourly, daily, and yearly	
Sorbonne Université, LATMOS/IPSL	Solène Turquety	CHIMERE	CTM	O ₃ , CO, VOCs, PAN, NH ₃ , aerosols	Northern Hemisphere, Europe	1° × 1° hemispheric, 10 km Europe	CAMS
Stockholm University	Matthew MacLeod	BETR Global	Multi-media POPs	POPs, PAHs	Global	3.75° × 3.75°, weekly or monthly output	
(b)							
Organization	Meteorology	Anthro emissions	Fire emissions	Natural emissions	Simulation length	Fire plume height	References
University of Hertfordshire, Centre for Climate Change Research		CAMS regional	CAMS GFAS, NCAR FINN	MEGAN			
Environment and Climate Change Canada Climate Research Division	Free-running or nudged to ERA-Interim	CMIP6, ECLIPSEv6b	CMIP6/GFED4 or use online interactive fire module from CLAS-SIC		Season to multi-decade	Climatological distribution based on AeroCom, or online with CFFEPS plume height scheme	
NSF National Center for Atmospheric Research Atmospheric Chemistry Observations and Modelling	Nudged to MERRA2 reanalysis	CAMS	FINN, QFED, or other	MEGAN online in CLM, prognostic sea spray, and dust	Season to multi-year		https://wiki.ucar.edu/display/camchem/Home (last access: 16 May 2025)
NASA Goddard/University of Maryland	WRF, initial/boundary conditions from NARR	CAMS (multi-year), HTAPv3 for 2018	QFED, plume rise	Online biogenic and lightning	Case studies and multi-year warm seasons		Huang et al. (2022, 2025)
NASA Goddard Institute for Space Studies		CMIP6	GFED4s, pyrE (interactive fire model)	MEGAN, wind-driven sea salt and dust, lightning			
Cyprus Institute Climate and Atmosphere Research Centre		CAMS, EDGAR	FINN or GFED or other	MEGAN, lightning, dust/sea salt	2010–present		
University of Bremen, Institute of Environmental Physics		CMIP6	GFEDv3, GFEDv4, CMIP6	MEGAN-MACC	Multi-year		

Table A2. Continued.

Organization	Meteorology	Anthro emissions	Fire emissions	Natural emissions	Simulation length	Fire plume height	References
UKCEH/Edinburgh University, Atmospheric Chemistry and Effects/School of Chemistry	WRF, with reanalysis from GFS/ERA5 depending on version/setup. Nudged every 6 h, 1° res.	NAEI for UK, CEIP for Europe, HTAP (2010) for global	NCAR FINN (v1.5 and transitioning to v2.5)	Online biogenic VOCs (BVOCs) soil NO _x , volcano, sea salt, dust	Month to multi-year	Emissions evenly distributed over 8 lowest vertical layers (Simpson, 2014)	Simpson et al. (2012), Vieno et al. (2016)
UK Met Office, Hadley Centre		Mainly CMIP6, but flexible	Can run with pre-scribed emission datasets or use online interactive fire model IN-FERNO	Various and depends on the configuration setup but mostly interactive for dust, BVOCs, sea salt, DMS	Typically years, multi-year, multi-decade		
Lancaster University, Lancaster Environment Centre	Driven by ECMWF-IFS cy38 met at TL159L60 3 h resolution	Flexible	Flexible	MEGAN	Single-year/multi-year	Surface/planetary boundary layer (PBL) emissions only	Wild (2007)
Thailand Team, KMUTT		HTAP/CAMS, locally adjusted	FINN and others	MEGAN	Sub-seasons, selected events		
IITM Pune, India, AQEWS Urban Air Modelling		EDGAR-HTAP	FINNv1.5	MEGAN	Air quality forecast for 10 d		
CICERO Center for International Climate Research, Oslo, Norway		Flexible, but currently CEDS and ECLIPSE most used.	GFED4	MEGAN (online or offline)	Single-year/multi-year, likely time slice for selected years on longer timescales.		
DLR Institute of Atmospheric Physics, Earth system modelling	Free-running or nudged (ERA5)	Flexible	Flexible	Lightning NO _x , air–sea exchange, dust, biogenic and soil NO _x			Jöckel et al. (2010, 2016)
NOAA Geophysical Fluid Dynamics Laboratory (GFDL)	Driven by observed sea surface temperatures or nudged to reanalysis winds	CEDS-v2021-04-21	GFED4 (daily or monthly), but flexible	Interactive MEGAN BVOCs; Interactive dust coupled to vegetation cover; Lightning NO coupled to subgrid convection	Multi-year, multi-decade	Distributed vertically up to 6 km, based on an injection height climatology from MISR	Lin et al. (2024a, b)
MIT, Earth, Atmospheric and Planetary Sciences		PKU	PKU	PKU			
Meteorological Research Institute, Japan Met Agency (MRI-JMA), Department of Atmosphere, Ocean, and Earth System Modelling Research	Free-running or nudged to JRA55	CMIP6, ECLIPSEv6b	CMIP6/GFED, GFED				
Institut National Polytechnique Félix Houphouët-Boigny (INP-HB) Department of Forestry and Environment	ERA5	CMIP6	GFED4				
Space Research of the Netherlands, Earth Department	Mainly ECMWF	CAMS	GFED4.1s, or GFED5, or GFAS			IS4FIRES	
Norwegian Meteorological Institute, Research Department	ECMWF	Variable, mainly EMEP	FINN, GFAS				
NILU, Atmospheric Chemistry	ECMWF, ERA5, CESM GCM, WRF, etc.	ECLIPSEv6b	GFED, GFAS		Years	From GFAS emissions dataset	Pisso et al. (2019)

Table A2. Continued.

Organization	Meteorology	Anthro emissions	Fire emissions	Natural emissions	Simulation length	Fire plume height	References
Finnish Meteorological Institute, Atmospheric Composition Research	Nudged to MERRA2 reanalysis data	CAMS, CMIP, ECLISE, EDGAR	IS4FIRES	MEGAN or own model (Europe)	From 5 d forecasts up to multi-decades in climate mode	Sofiev et al. (2012)	
Tsinghua University, School of Environment	NCEP FNL reanalysis data	Huang et al. (2023) for full-volatility organic, CMIP6 for other pollutants. Chang et al. (2022) for full-volatility organic ABaCAS-EI for other pollutants	VOC emissions based on the burning area of GFEDv4; Other pollutants from GFEDv4, FINN	Dust/biogenic/sea salt: calculated online dust/biogenic/sea salt: calculated online	Multi-year (2015–2020), 2018	Daily fire info from FRP and met data. Freitas plume rise scheme	
University of Augsburg, Faculty of Medicine	Nudged to GFS during spinup above PBL	EDGARv5 + national German inventory	FINN	Online	Season to multi-year		
Jožef Stefan Institute/MS-C, Department of Environmental Sciences	ECMWF, WRF	EMEP, EDGAR, PKU	FINN, MCHgMAP, PKU	MCHgMAP, PKU	Multi-year		
Peking University, Lanzhou University	ECMWF, NCEP FNL	PKU, LZU, EDGAR	PKU, LZU		Month to multi-year	Gaussian plume model to distribute to 3 km height	Luo et al. (2020); Song et al. (2023b)
Sorbonne Université, LATMOS/IPSL	ERA5, WRF	CAMS	CAMS GFAS, API-FLAME	MEGAN, dust, sea salt, lightning	Seasonal	Satellite observations, GFAS plume height	Menut et al. (2021), Turquety et al. (2020)
Stockholm University	Driven by ECHAM 5 model outputs	Flexible	Flexible	Flexible	Multi-year, multi-decade	Emission to boundary layer or free troposphere	MacLeod et al. (2011)

Data availability. The regional land mask (Fig. 3d) is available here: <https://doi.org/10.5281/zenodo.15270649> (Whaley, 2025).

The HTAPv3.1 anthropogenic emissions files for the historical period are available here: https://edgar.jrc.ec.europa.eu/dataset_htap_v31 (European Commission, 2025) and <https://doi.org/10.5281/zenodo.1449944> (Crippa, 2024).

The IIASA GAINS anthropogenic emissions files for the future period are available here: <https://doi.org/10.5281/zenodo.14748815> (Klimont et al., 2025).

The GFAS4HTAPv1.2.1 biomass-burning emissions for the historical period are available here: <https://doi.org/10.5281/zenodo.13753452> (Kaiser and Holmendal, 2024b).

The Hamilton et al. (2025) biomass-burning emissions for the future period are freely available from the Coupled Model Inter-comparison Project at <https://aims2.llnl.gov/search> (ESGF, 2025; Hamilton et al., 2025), and the post-processed emissions will be provided to participants.

The ERA5 reanalysis recommended for meteorology is available here: <https://doi.org/10.24381/cds.adbb2d47> (Hersbach et al., 2023).

The observational datasets for the assimilation experiments are available here:

- MOPITTv9 CO Level 2 data are available through the NASA EarthData portal at <https://doi.org/10.5067/TERRA/MOPITT/MOP02J.009> (MOPITT Team, 2022).
- TROPOMI NO₂, CO, and O₃ Level 2 datasets are made available operationally through the ESA Sentinel-5P data hub (<https://dataspace.copernicus.eu/>, Copernicus, 2025).
- OMI NO₂ v4 data are available at https://search.earthdata.nasa.gov/search?q=OMNO2G_003 (NASA, 2025a).
- OMI O₃ v3 data are available at https://search.earthdata.nasa.gov/search?q=OMTO3_003 (NASA, 2025b).

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