

AerChemMIP2 - Unraveling the role of reactive gases, aerosol particles, and land use for air quality and climate change in CMIP7

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Abstract. Phase 2 of the Aerosol and Chemistry Model Intercomparison Project (AerChemMIP2) is a registered model intercomparison project (MIP) of the Coupled Model Intercomparison Project phase 7 (CMIP7). The focus of AerChemMIP2 is the quantification of the atmospheric composition, biogeochemical feedbacks, air quality and climate responses to changes in emissions of chemically reactive gases, aerosol particles, and land use. AerChemMIP2 aims to facilitate a better understanding of their relative contributions to changes in atmospheric composition, radiative forcing, and the climate response and feedbacks from the pre-industrial period to the present day and for projected future emission pathways. Some experiments from the first phase of AerChemMIP are requested in the second phase to track changes in the results of CMIP7 compared to phase six of CMIP. New experiments in AerChemMIP2 open scientific opportunities to address knowledge gaps and persistent uncertainties. Specifically, AerChemMIP2 requests experiments (1) to assess the dependence of effective radiative forcing for aerosols on the fidelity of resolved processes and the simulated base climate, (2) to provide first estimates of forcing for hydrogen and individual volatile organic compounds in the context of CMIP, (3) to enable studies on non-linearity in the Earth system response, (4) to understand the response of wild fires to historical forcings, and (5) to quantify the influence of desert dust increases on climate change. AerChemMIP2 further requests variants of the ScenarioMIP-CMIP7 high-end and

overshoot scenarios to quantify future responses to policy implementations for air quality management. Diagnostic requests
15 of AerChemMIP2 are made from CMIP7 core experiments to facilitate offline experiments for chemistry and aerosols. The
experimental protocol of AerChemMIP2 presented here closely aligns with the CMIP7 core experimental design, and its
other registered MIPs. Selected AerChemMIP2 experiments are performed in the Assessment Fast Track (AFT) of CMIP7.
Participation of modelling centres in AerChemMIP2 would help to gain new insights for atmospheric composition and implications
for air quality in a warming world with rapidly changing emissions.

20 **1 Introduction**

Atmospheric composition is key to understanding climate change and air quality. Atmospheric constituents that typically stay
in the atmosphere for weeks to two decades are referred to as short-lived climate forcers (SLCFs, Szopa et al., 2021), and
affect both climate change and air quality, schematically depicted in Figure 1. Emissions and concentrations of SLCFs are
spatially and temporally variable. SLCFs may therefore influence climate differently in time and space. Due to their short
25 atmospheric residence time, emission reductions to improve air quality may lead to benefits or adverse influences on global
and regional climate on timescales much shorter than for long-lived greenhouse gases (GHGs). Despite their significant role in
regional climate change, the magnitude of radiative forcing, climate responses, and feedbacks associated with SLCFs remain
insufficiently understood (e.g., Bellouin et al., 2020; Schaeffer et al., 2025). Addressing these critical knowledge gaps is the
primary goal of the Aerosol and Chemistry Model Intercomparison Project (AerChemMIP).

30 AerChemMIP phase one (Collins et al., 2017) targeted the role of SLCFs including methane, and was designed in support
of the Coupled Model Intercomparison Project phase six (CMIP6, Eyring et al., 2016). Of interest was the role of methane
(CH₄), tropospheric ozone (O₃) and its non-CH₄ precursor gases, stratospheric O₃, nitrous oxide (N₂O) and ozone-depleting
substances (ODSs), and aerosol particles and their precursors. The MIP facilitated the quantification of climate and air quality
impacts of aerosols and chemically reactive gases (Collins et al., 2017), recently reviewed by Griffiths et al. (2025). It
35 included estimates of forcings and responses from changes in aerosols, tropospheric O₃ and their precursor emissions, and
halocarbons. Feedbacks from climate-driven changes in natural emissions of individual species were also quantified. There
were detailed studies on the role of SLCFs in future climate and air quality with a focus on tradeoffs and co-benefits between air
quality and climate policies, highlighting potential win-win scenarios. AerChemMIP experiments highlighted how SLCFs pose
attractive options for near-term climate mitigation, and prompted new research on their regional impact and the uncertainties
40 and challenges in assessing the impact of these short-lived climate forcers (Griffiths et al., 2025, and references therein).
AerChemMIP focused on the process level within the complex and diverse range of Earth System and climate models, aiming
to build confidence in our understanding of the role of these processes in climate forcing and Earth system responses and
feedbacks. The experiments also allowed the attribution of historical surface temperature changes to composition changes,
and the calculation of radiative forcing of individual components. This newly gained understanding was included in the sixth
45 assessment report (AR6) of Working Group I (WGI) of the Intergovernmental Panel on Climate Change (IPCC), e.g., in the
dedicated chapter on SLCFs (Szopa et al., 2021).

AerChemMIP phase two (AerChemMIP2) builds on the experimental protocol of AerChemMIP (Collins et al., 2017) in two ways. First, we largely adopt the same strategy for the experimental designs by again requesting time-slice experiments with prescribed annually repeating conditions for computing radiative forcing and feedbacks, as well as fully coupled and atmosphere-only experiments for the historical and future time periods as in CMIP to study climate forcing and the response of air quality and climate. Second, we partly request that some of the experiments as in AerChemMIP are repeated with CMIP7 models and climate forcings data sets in order to track changes in forcings, responses, and feedback estimates since AR6. Nitrate aerosols were, for instance, simulated by only a few models in CMIP6 (Turnock et al., 2020; Allen et al., 2020, 2021). As such, there is currently a large uncertainty in future changes and the associated implication of nitrate aerosols for climate and air quality. AerChemMIP2 experiments can help to address these uncertainties based on the larger number of models with interactive nitrate aerosol capability.

New experiments in AerChemMIP2 are opportunities to address knowledge gaps, some of which have already been known for some time. One of these gaps relates to the historical and future changes of natural dust aerosols, which have important implications for climate and air quality. Desert dust aerosol constitutes the largest fraction of the total aerosol mass in the Earth's atmosphere (e.g., Adebisi et al., 2023) and affects climate through its direct radiative effect and via its influence on cloud microphysics (e.g., Kok et al., 2023). Moreover, dust influences biogeochemical processes with links to the carbon cycle, air quality and human health, as well as other socio-economic impacts including the transportation, agricultural, and renewable energy sectors. Despite the importance of natural dust, CMIP6 models did not capture observed historical trends for dust aerosols (Kok et al., 2023), although large uncertainties in global dust simulations have been known for some time (Huneus et al., 2011; Evan et al., 2014). AerChemMIP2 therefore includes new experiments with prescribed monthly dust emission data to induce observationally informed trends for dust aerosols in CMIP7 model simulations. The aim is a first assessment of the spatio-temporal changes in climate forcing and response associated with dust aerosol trends in CMIP7.

Another opportunity in AerChemMIP2 concerns the past evolution of wildfire activities and associated emissions of SLCFs. Fires have devastating impacts on humans and the environment (e.g., Martin et al., 2016) and their emissions in the pre-industrial era have implications for the quantification of present-day anthropogenic aerosol forcing (Hamilton et al., 2018, 2024). Moreover, humans are directly involved in altering the environment in which fires can occur, e.g., through changes in land use, fire weather, and fire management policies, altering the magnitude and extent of past and future wildfires. More models are anticipated to have the capability to perform experiments with co-evolving fire and climate conditions, which AerChemMIP2, in collaboration with FireMIP (Li et al., 2025), will exploit for new climate forcing and response experiments targeting fire emissions.

AerChemMIP2 experiments further enable the quantification and understanding of the role of SLCFs aligned with future scenarios in CMIP7 (Dunne et al., 2025) that complement the set of scenarios expected to be provided by ScenarioMIP-CMIP7 (van Vuuren et al., 2025). AerChemMIP explored the air quality and climate responses to strong air pollution controls with and without CH₄ mitigation in a world with increasing long-lived GHGs by requesting variants of the Shared Socioeconomic Pathway SSP3-7.0 (e.g., described by Allen et al., 2021). AerChemMIP2 requests experiments building on two ScenarioMIP-CMIP7 scenarios that have alternative SLCF emission trajectories. SLCFs are of particular importance for regional climate

change (Wilcox et al., 2023; Persad et al., 2023), especially in the next few decades, due to both air quality abatement measures and new emerging technologies for sustainability. Reaching climate neutrality implies, for instance, a large-scale increase in renewable energy sources, which may include a growing hydrogen economy, in which hydrogen (H₂) is produced from
85 renewables and/or from fossil fuels combined with carbon capture and storage (CCS). Regardless of the production method, H₂ is prone to leakage during the whole value chain, and will indirectly contribute to warming by changing methane (CH₄), ozone (O₃) and stratospheric water vapor levels (Sand et al., 2023; Warwick et al., 2023). The potential implication of a hydrogen economy can be studied with Chemical Transport Models (CTMs) driven by output from selected CMIP7 model experiments that have been included in the full CMIP data request for atmospheric variables (Dingley et al., 2025).

90 Finally, AerChemMIP2 interacts synergistically with other MIPs for creating multi-purpose experiments, i.e., requesting identical experiments across MIPs, where possible. We do so to avoid duplication of effort and to reduce the overall computational burden for modelling centres. We further identified research areas where novel opportunities arise from the use of consistent parallel experiments across MIP boundaries and other modelling initiatives.

2 Scientific Aims

95 The experiment protocol of AerChemMIP2 is guided by our scientific aims, which are embedded in the scientific goals of the Coupled Model Intercomparison Project phase seven (CMIP7, Dunne et al., 2025). The scientific aims of AerChemMIP2 were defined through community consultation and discussions via online meetings and workshop of the Composition Air quality Climate inTeractions Initiative (CACTI, Fiedler et al., 2024), resulting in four scientific questions for AerChemMIP2.

- 100 1. *Process Understanding*: How have our process understanding and associated impact assessments advanced for global and regional atmospheric composition changes, radiative forcing, and climate responses?
2. *Feedbacks*: How important are climate feedbacks on natural SLCF emissions, atmospheric composition, and radiative effects?
3. *Air quality*: What is the relative importance of climate change and emissions of SLCFs for atmospheric composition and air quality over the historical period and into the future?
- 105 4. *Sustainability*: What future climate benefits and/or penalties are expected from improving air quality and what are the climate benefits/trade-offs arising from policies for improved sustainability?

Through addressing these questions, AerChemMIP2 aims to contribute to quantifying and advancing the scientific understanding of the role of SLCFs for (1) global climate forcing and patterns of radiative effects, (2) the spatio-temporal response of atmospheric composition, air quality, and climate to SLCF changes, and (3) climate and biogeochemical feedback mechanisms
110 induced by emission and atmospheric composition changes. The aims of AerChemMIP2 contribute to the four guiding science questions of CMIP7 outlined by Dunne et al. (2025) as follows.

Patterns of sea surface change: SLCFs are much more spatially heterogeneously distributed than long-lived, well-mixed GHGs. Through the focus on SLCFs like aerosol particles, AerChemMIP2 supports the CMIP7 science goal concerning a better understanding of historical changes in spatial patterns of radiative effects (Section 3.1), climate responses, feedback processes, and their interactions with natural variability (Section 3.2). Past research has for instance shown northern hemisphere responses of sea-surface temperature patterns to the aerosol increase until the 1980s and their later West to East shift due to changes in atmosphere and ocean dynamics (Fiedler and Putrasahan, 2021; Kang et al., 2021), including a response of the Atlantic Meridional Overturning Circulation (Booth et al., 2012) which is seen as a tipping element in the climate system (McKay et al., 2022). AerChemMIP2 experiments will help to improve the understanding of the role of SLCF emissions in shaping the historical and future spatial patterns of air quality and climate change, and their temporal evolution (Sections 3.2 and 3.3) paired with precise model-based estimates of patterns of their radiative effects (Section 3.1). AerChemMIP2 experiments will therefore be useful in assessing how SLCF emissions contributed to observed changes and whether changes attributed to individual emission perturbations will be different in future emission trajectories.

Changing weather: AerChemMIP2 experiments enable the impact of SLCFs on weather patterns and their connection to extremes, including impacts on air quality and associated extremes, to be assessed. Emissions of SLCFs, such as aerosol, have marked regional patterns and are known for different effects on atmospheric radiative transfer and associated weather responses, e.g., ranging from differences in atmospheric stability, cloud microphysical processes, and precipitation formation, to larger scale responses of the atmosphere and ocean circulation (e.g., Allen and Sherwood, 2011; Bellouin et al., 2020; Wilcox et al., 2020; Fiedler and Putrasahan, 2021; Xie et al., 2022; Myhre et al., 2024). AerChemMIP2 experiments may help to better understand both the historical and potential future changes in weather patterns influenced by SLCFs in a warming world, e.g., monsoon precipitation, including extremes. Additionally, extreme weather events, such as heatwaves, can in turn create meteorological conditions favourable for poor air quality which may lead to compound high-impact events (e.g., Fiore et al., 2015; Schnell and Prather, 2017; Guo et al., 2021; Jain et al., 2022). Such co-occurrences of extreme weather and pollution events can be assessed with output from the AerChemMIP2 ensemble of experiments (e.g., Sections 3.2.1, 3.2.2, and 3.3.1).

Water-carbon-climate nexus: Advancing the understanding of biogeochemical feedbacks for climate change is again a component of AerChemMIP2. AerChemMIP2 requests experiments with Earth system models allowing the community to assess chemical cycles in the context of air quality and climate change. New to AerChemMIP2, and hence CMIP7, are experiments with prescribed desert dust aerosols for which CMIP models have been facing uncertainty across multiple CMIP phases such that the past trends were missed by models. As carriers of nutrients for both terrestrial and marine eco-systems, desert dust aerosols influence the carbon cycle and AerChemMIP2 experiments will allow the regional-to-global implication of dust changes in the context of past climate change and future trajectories of SLCF emissions to be assessed (Section 3.2.3). Moreover, AerChemMIP2 addresses the role of wild fires through fully coupled simulations with land use, the carbon cycle and SLCFs (Section 3.2.4). It also targets the influence of future land-use change on climate and atmospheric composition through experiments with future afforestation and reforestation with feedbacks on emissions (Section 3.3.2).

Points of no return/ratcheting: Scenarios in AerChemMIP2 include both an overshoot and a high-end emission scenario to quantify the role of SLCF emissions, including CH₄, in the future. AerChemMIP2 requests experiments from Earth system

models that can simulate feedback mechanisms that other CMIP7 models do not account for. As such, AerChemMIP2 experiment output can address to what extent changes in atmospheric composition from, e.g., increasing CH₄ emissions from wetlands due to warming (e.g., O'Connor et al., 2010) and decreasing aerosol load due to mitigation policies (e.g., Wood et al., 2024) can both exacerbate near-term warming. Earth system models capable of answering these kinds of questions are increasingly becoming available (e.g., Folberth et al., 2022). As such, AerChemMIP2 will help to highlight potential surprises regarding atmospheric composition changes by providing a basis for analyzing CH₄ feedbacks which are unique in the MIP family listed by CMIP7 (Section 3.3.1).

3 Experimental Design

The AerChemMIP2 experiments are designed to allow the community to address the scientific questions in Section 2 while keeping model capabilities in mind. Models contributing to AerChemMIP2 are primarily those with a capability to interactively simulate climate processes with spatio-temporal changes in aerosols (AER) or in trace gas chemistry *and* aerosols (CHEM). Specifically, AER means that the model should at least have a time-evolving treatment of aerosols, either through time-dependent prescribed input fields or a fully interactive aerosol scheme whereas CHEM means that the model is also required to have an interactive chemistry scheme.

Experiments should preferably be performed with as much capability as possible with respect to the atmosphere, atmospheric composition, land, ocean, and climate components. AerChemMIP2, therefore, expects more complexity in simulated processes in the participating models than is the case for most models in CMIP7. However, model contributions with prescribed concentrations or optical properties, like in many CMIP7 models, are equally welcome (e.g., for aerosols, ozone and their precursors) to facilitate broad intercomparison studies building on model output with different complexities. This diversity in model complexity is deliberately chosen, since too few model contributions in AerChemMIP was challenging for interpreting results and models with intermediate complexity are useful for transient experiments (Griffiths et al., 2025). In the CMIP7 experimental design, Sanderson et al. (2024) argue that emission-driven simulations for carbon dioxide (CO₂) should be prioritized. Emission-driven capability for other climate forcers, like CH₄ and other SLCFs, is not explicitly expected in CMIP7 (Sanderson et al., 2024) but is encouraged in AerChemMIP2. Therefore, it is anticipated that the complexity of atmospheric composition process representation (e.g., AER or CHEM) will vary across the AerChemMIP2 ensemble, and the capability required to perform a particular simulation is incorporated into the experimental designs described in the following sections. Irrespective of the level of model complexity chosen for participation in AerChemMIP2, we request that the model complexity is consistent with that used in the DECK and ScenarioMIP-CMIP7, whose experiments in some instances act as control experiments for AerChemMIP2. Modelling groups are specifically requested to document the model complexity in the metadata of their output, e.g., by providing a reference for the simulated chemical and aerosol processes.

Motivated by the wide usage of AerChemMIP experiments for research studies (Griffiths et al., 2025, and references therein), we ask modelling centres to replicate a selection of the AerChemMIP phase one experiments (Collins et al., 2017) in phase two. Note that we adjust some of the experiment names for improved clarity, e.g., in AerChemMIP2, we have experiments

180 on air quality (AQ), such as *piClim-AQ*, to better reflect that these experiments account for perturbations of tropospheric
O₃, aerosols and their precursor emissions instead of all near-term climate forcers (NTCF). We have, therefore, changed the
name of such experiments compared to AerChemMIP phase one, e.g., from *piClim-NTCF* to *piClim-AQ*. For improved clarity,
we have also changed the name of the experiments involving stratospheric ODSs to *piClim-ODS*, rather than using HC for
halocarbons from the first phase (*piClim-HC*). Repeating a selection of experiments in AerChemMIP2 allows the community
185 to document the degree to which results change between CMIP6 and CMIP7. Although the experimental designs are similar, the
model output will deliver new information, e.g., additional diagnostics and updating the present-day effective radiative forcing
for individual anthropogenic perturbations up to the year 2021 based on the latest model versions and the CMIP7 climate
forcing data sets. All AerChemMIP2 experiments are expected to utilize the same version of forcing datasets that are used
for the CMIP7 Diagnostic, Evaluation and Characterization of Klima (DECK) suite of experiments (Dunne et al., 2025, also
190 <https://input4mips-cvs.readthedocs.io/en/latest/dataset-overviews/>), unless stated otherwise. This will ensure that the specific
responses are due to the experimental protocol and not to any potential forcing differences.

In addition to the request to repeat selected experiments of AerChemMIP, newly designed experiments in AerChemMIP2
address current challenges in composition-climate modelling, where the community perceives major gaps in our understanding
of climate change with potential implications for future developments. These challenges include for instance the poor representation
195 of changes in desert dust aerosols and the lack of inter-model diversity in fire emissions via fully coupled fire schemes in
CMIP6 models. AerChemMIP2 identifies eight topics for advancing our understanding with the help of new experiments and
five topics where updates based on previously existing experiment designs are perceived useful. These topics are distributed
over three families of AerChemMIP2 experiments: time slice experiments with prescribed annually repeating climatologies
for sea-surface temperature and sea-ice conditions (Section 3.1), atmosphere-only and fully coupled historical experiments
200 with transient changes in climate forcings for 1850–2021 inclusive (Section 3.2), as well as atmosphere-only and fully coupled
future scenarios for 2022–2125 inclusive (Section 3.3). Some of the AerChemMIP2 experiments are included in the Assessment
Fast Track (AFT) of CMIP7, schematically depicted in Fig. 2. The reasons for their inclusion and the mix of experiments in
AerChemMIP2 from which the scientific opportunities arise are described in the following sections.

3.1 Time slice experiments

205 We request atmosphere-only experiments for time slices with a prescribed annually repeating monthly climatology of sea-
surface temperatures and sea ice. The climatology is based on the last thirty years from the model's own PI fully coupled control
experiment for *piClim-X* and the model's own fully-coupled historical experiment for *pdClim-X* experiments in AerChemMIP2,
where *X* refers to the emission of an individual or a combination of SLCFs. Models that interactively simulate emissions that
depend on the ocean surface state, e.g., surface water concentrations of dimethyl sulfide (DMS) or chlorophyll, are asked to
210 also prescribe them as monthly climatology, which are to be diagnosed from the same reference experiments.

The categorization of biomass-burning emissions in such experiments is a challenge. AerChemMIP2 encourages models
to use the fullest complexity of interactive processes available, and therefore request that models determine and document
the PI fire-related emissions as a reference to compute the ERF, and which (if any) are treated as anthropogenic. Not all

biomass burning emissions have an anthropogenic origin. For models which apply prescribed agricultural and/or deforestation biomass-burning emissions, we recommend that post-1850 SLCF emissions from agricultural and deforestation fires should, for instance, be attributed to anthropogenic emissions and included in the respective experiments. However, the exact distinction will depend on the complexity of the fire representation in the models and should be documented. For instance, in models which interactively simulate agricultural burning alongside unmanaged wildfire emissions, a separation between natural and anthropogenic emissions might not be possible.

The number of *piClim-X* and *pdClim-X* experiments defined below is relatively large but they come with a comparably small computational burden. These are single-forcing atmosphere-only experiments of 30 years in length post model spinup. This is a deviation from phase 1 where the simulation length was 30 years in total, with no recommendation for spinup. However, some models found that up to 15 years was required for spinup, particularly for perturbations to longer-lived GHGs such as ODSs (e.g., O'Connor et al., 2021). Therefore, the recommendation is allow sufficient time for spinup and then run the experiments for 30 years. The necessary time for the spinup might depend on the model configuration and could differ substantially between models. We recommend to run the spinup until the simulated radiation balance at the top-of-the atmosphere and the chemistry state reach an equilibrium, i.e., trends across annual values (if any) become small. Some simulations with CMIP6 models required for instance 10 years for ozone depletion to reach an equilibrium after the concentrations of ODSs were changed from PI to PD levels (Figure S2 in Collins et al., 2026). Modellers with interactive chemistry schemes may choose to accelerate the spinup by initialising concentrations of CH₄ or ODSs throughout the atmosphere to the new present-day values, e.g., take them from the historical experiment rather than starting from PI levels. Experience from the community during CMIP6 showed that these experiments are relatively easy to perform once the model setup for the *piClim-control* experiment has been completed; this latter experiment was common to AerChemMIP and the CMIP6-endorsed Radiative Forcing Model Intercomparison Project (RFMIP, Pincus et al., 2016), and it is now a core experiment performed for the AFT of CMIP7 (Dunne et al., 2025). The computational burden is justified by their potential for scientific exploitation and their direct relevance for climate change assessments. AerChemMIP2 and RFMIP2 (Kramer et al., 2025) will again use the same PI base state for their experimental setups to ensure the comparability of the results and to keep the use of computing resources small. It allows modelling centres to easily contribute to both MIPs.

3.1.1 Partitioning of ERF

The *piClim-X* experiments of AerChemMIP and RFMIP were essential in IPCC AR6 for quantifying individual SLCF effective radiative forcings (ERFs) at the present day (PD) relative to the pre-industrial (PI) period. These estimates allow a detailed assessment of the relative contributions from changes in individual emissions to the radiation imbalance, with simulations accounting for internal natural variability in the radiation budget. Specifically, *piClim-X* provided the basis for assessing the contribution of PI to PD anthropogenic SLCF emission changes to effective radiative forcings and historical global mean surface temperature change (Szopa et al., 2021). AerChemMIP2 will again enable an analysis of the contributions from individual changes in emissions of individual aerosol species and reactive gases to present-day effective radiative forcing. In doing so, AerChemMIP2 enables attributing shares of radiative forcing to individual composition changes, e.g., for SO₂,

which gives additional information that complements estimates of effective radiative forcing for all aerosols taken together as in *piClim-aer*, also requested by RFMIP (Pincus et al., 2016) and CMIP7 (Dunne et al., 2025). The experiment *piClim-aer* in AerChemMIP2 uses aerosols and aerosol precursor emissions of BC, organic carbon (OC), ammonia (NH₃), and sulphur dioxide (SO₂), and omits changes in anthropogenic NO_x and VOC emissions. Although these latter species may act as aerosol precursors and influence oxidising capacity and secondary aerosol formation, they are omitted here to be consistent with the same experiment in RFMIP2 (Kramer et al., 2025) and to maximise potential contributions from models with AER capabilities.

AerChemMIP2 request partly similar *piClim-X* experiments compared to AerChemMIP to track changes since CMIP6 and gain new insights, not only because of updated climate forcings data sets and a more recent year for PD values, but also because of new model developments. AerChemMIP had, for instance, few results for nitrate aerosol, due to few models having interactive nitrate aerosol schemes, which limited the understanding of their role in climate and air quality in CMIP6 (Turnock et al., 2020; Allen et al., 2020, 2021) - an aspect that will potentially change in CMIP7 due to additional models with the capability of interactively simulating nitrate aerosols (e.g., Jones et al., 2021), e.g., [GFDL-ESM4.5, which is an updated configuration of ESM4.1 \(Dunne et al., 2020; Horowitz et al., 2020\), and other models \(Kelley et al., 2020; Jones et al., 2021; Tilmes et al., 2021\)](#).

The *piClim-X* experiments listed in Table 1 contribute to science question 1 and are part of the AFT of CMIP7 (Fig. 2). We request 12 *piClim-X* experiments (Table 1) for calculating ERFs of anthropogenic SLCFs with experiment *piClim-control* serving as a reference. The PD conditions are representative of the year 2021, consistent with CMIP7 (Dunne et al., 2025). The conditions of the specified year are to be used in all model components, e.g., PD CH₄ levels (either concentrations or emissions) in *piClim-CH4* are used in both the chemistry and radiation schemes to allow for both direct radiative effects and chemical adjustments via CH₄-driven influences on O₃, stratospheric water vapor and CH₄ lifetime via changes in the hydroxyl (OH) radical. An exception is the experiment *piClim-O3*, in which the CH₄ levels are for 1850 in the radiative transfer calculation, but for 2021 in the chemistry scheme to account for chemical adjustments (i.e., via CH₄ oxidation) influencing the radiative forcing of O₃ (Table 1).

3.1.2 State-dependence of ERF

New in AerChemMIP2 are parallel calculations of the effective radiative forcing (ERF) based on two reference states, namely the PI as outlined above and the PD base state. The chemical and physical base states of the atmosphere in the PI and PD differ and are thought to influence the magnitude of ERF, e.g., for anthropogenic aerosols (Carslaw et al., 2013). Models without interactive chemistry and aerosols, as often used for CMIP experiments, however, might only show a weak dependency on the base state, e.g., for anthropogenic aerosol ERF (Fiedler et al., 2025; White et al., 2025). AerChemMIP2 experiments can be used to systematically address the influence of the PI to PD differences in the base state on ERF magnitudes based on the most comprehensive Earth system models that are currently available. In addition to studying the base-state dependence, performing *pdClim-X* experiments is beneficial for an additional evaluation of the climate model results over the past few decades, for which a rich collection of observational data exists.

To explore the role of the base state for estimates of forcing by SLCFs, AerChemMIP2 adds a new family of *pdClim-X* experiments. In the *pdClim-X* experiments, we ask to prescribe the model's own PD climatology for sea surface temperatures and sea ice, and PD SLCFs instead of using their PI equivalents. The PD climatology can be created from the last thirty years of a fully coupled historical experiment. These *pdClim-X* experiments, listed in Table 2, can be evaluated against observations, which is an advantage over *piClim-X* to the extent that the PI state is inherently uncertain. The *pdClim-X* experiments can also be more relevant for understanding the climate impacts of future changes in SLCFs starting from present conditions. Moreover, the comparison of the new *pdClim-X* paired experiments against their *piClim-X* counterparts allows for an assessment of the state-dependence of ERF for O₃, aerosols, and black carbon (BC) separately, with a substantially reduced internal variability than for atmosphere-only experiments with transient changes. We separate BC and all aerosols from the estimate for all SLCFs taken together, since aerosol effects showed large differences in ERF in previous assessments (Bellouin et al., 2020).

3.1.3 Step change for aerosol ERF

Aerosol-cloud interactions are a major uncertainty in understanding and quantifying the ERF of anthropogenic aerosols (Bellouin et al., 2020) - a problem tightly linked to the representation of meteorological processes influencing clouds (Stevens and Bony, 2013; Bony et al., 2015). For instance, model biases in precipitation patterns are known and have not been resolved in CMIP models (Respati et al., 2024), although progress in modelling some precipitation metrics is noticeable across the CMIP phases (Fiedler et al., 2020). Moreover, aerosol effects on clouds co-develop with meteorological conditions, which requires an adequate analysis of the effects that consider different cloud regimes embedded in meso-scale dynamical processes of the atmosphere across world regions. Storm-resolving models have been proposed to overcome some of the longstanding challenges in simulating cloud coupling to atmospheric dynamics and associated precipitation (Guendelman et al., 2024), although accurately modelling cloud microphysical processes remains a challenge (Naumann et al., 2025).

We encourage modelling centres that develop models for storm-resolving simulations with spatial resolutions of a few kilometers (1–20 km) to perform *pdClim-control* and *pdClim-aer*. Output from kilometer-scale simulations from models with any implemented aerosol treatment would be welcome, including both models using interactive aerosol schemes and models with prescribed data, e.g., for aerosol concentrations or optical properties. We ask the modelling centres to document their aerosol treatment in the metadata of the model output, e.g., with information on their aerosol treatment along with the reference for the prescribed aerosol data or the implemented aerosol parameterization schemes. Having these experiments from storm-resolving models would create a new line of evidence for aerosol ERF. The consistent experimental setup warrants an unbiased comparison against CMIP7 model results not only for aerosol ERF, but also for the simulated PD atmosphere for which the rich observational data can be leveraged.

3.1.4 Hydrogen

Using hydrogen (H₂) is a potential alternative to fossil fuels and is considered a mitigation option for some economic sectors, yet H₂ also has a climate warming potential (Goita et al., 2025). One reason is the intended and unintended leakage of H₂ during its production, transport, storage and end-use, which affects the atmospheric concentration of CH₄ which has a far

greater warming potential than CO₂. As green hydrogen from renewable energy and blue hydrogen with carbon capture are developed to support the transition to low-carbon energy systems, concerns exist that H₂ can contribute to warming (Sand et al., 2023; Warwick et al., 2023). By reacting with OH, H₂ reduces the atmosphere's ability to remove CH₄, leading to longer CH₄ lifetimes. H₂ can also contribute to the formation of tropospheric ozone and stratospheric water vapor, further enhancing its indirect warming effect. This warming could potentially be large enough to offset positive accomplishments for mitigating the effects of long-lived GHGs in the next few decades (Ocko and Hamburg, 2022), but the magnitude of these leakages is highly uncertain (Esquivel-Elizondo et al., 2023). Although, policies for H₂ are currently not planned, implications of H₂ leakage for climate change are beginning to be quantified (Paulot et al., 2021; Brown et al., 2025; Chua et al., 2025).

We request a new *pdClim-H2* experiment to estimate the potential forcing from a future hydrogen economy (*pdClim-H2*; Table 2) from models that include H₂ emissions. The experiment will enable the quantification of ERF of H₂. The Community Emissions Data System (CEDS) team will develop an H₂ emission data set for the experiment (pers. comm. Steve Smith, PNNL). Models could take the soil uptake into account following Paulot et al. (2024). Interested modelling centres are asked to contact the authors for coordinating the experimental setup in more detail, e.g., for using emissions or concentrations of H₂ for the experiment. Being aware of few models having the capability for H₂ experiments at the time of writing, namely UKESM1.0 (Brown et al., 2025) and GFDL-AM4.1 (Paulot et al., 2024), we also request output from CMIP7 experiments to setup simulations with chemical transport models (CTMs, Section 4.2).

3.1.5 Volatile organic compounds

A set of new *pdClim-X* experiments in AerChemMIP2 addresses the role of volatile organic compounds (VOCs), listed in Table 3. We choose here *pdClim-X* experiments for enabling direct comparison to observational data sets for present-day conditions. Non-methane volatile organic compounds (NMVOCs) contribute to air pollution as precursors of carbon monoxide (CO) and non-methane tropospheric O₃ and secondary organic aerosols, and influence the Earth's radiation budget via their effects on tropospheric O₃, aerosols, CH₄ lifetime and carbon dioxide (CO₂) concentrations (Stevenson et al., 2013). They may also cause aerosol-mediated cloud adjustments via changes in oxidising capacity and secondary aerosol formation (O'Connor et al., 2021; O'Connor et al., 2022). In the first phase of AerChemMIP, the *piClim-VOC* experiment included emission perturbations of both NMVOCs as well as carbon monoxide (CO) (Collins et al., 2017), complicating the attribution of ERFs to individual NMVOC and CO emission changes. In AerChemMIP2, we add two tier 2 experiments to explicitly quantify the ERFs due to PI to PD changes in anthropogenic emissions of CO (*pdClim-CO*) and NMVOCs (*pdClim-NMVOC*).

Additionally, we request 6 *pdClim-XVOC* simulations, where *XVOC* = ethane (C₂H₆), propane (C₃H₈), ethene (C₂H₄), propene (C₃H₆), butane (C₄H₁₀), and alcohols, to attribute air pollution and ERFs to changes in anthropogenic emissions of these individual NMVOCs. These experiments are assigned as tier 3 and are not part of the AFT of CMIP7. We expect the ERFs diagnosed from these perturbations to be small. However, these experiments will provide first estimates of the contributions of individual NMVOCs to air pollution and atmospheric composition in broader terms, e.g., particulate matter (PM) and O₃, CH₄ lifetime, and aerosol burden. The contributions of speciated NMVOCs to air pollution and climate have not been studied before in a consistent manner, highlighting a knowledge gap and an opportunity within AerChemMIP2.

3.1.6 Quantifying Biogeochemical Feedbacks

The *piClim-2X* experiments of AerChemMIP were essential for quantifying the magnitude of biogeochemical climate feedbacks on individual natural emissions of SLCFs (Thornhill et al., 2021a) and were used in IPCC AR6 (IPCC, 2021). In these climatology experiments, the emissions of the component of interest are doubled relative to PI levels. AerChemMIP2 again requests such experiments and, to a large extent, uses an identical experimental design as in AerChemMIP. In so doing, the *piClim-2X* experiments in AerChemMIP2 can be used to quantify changes in the feedback estimates since AerChemMIP arising from model improvements and forcing data updates since CMIP6. The *piClim-2X* experiments fall under science questions 2 and 3.

We request the 8 *piClim-2X* experiments listed in Table 4; they allow us to estimate the strength of climate feedbacks on sea spray, desert dust, fires, biogenic volatile organic compounds (BVOCs), wetlands, and primary marine organic aerosols (PMOA) plus dimethyl sulfide (DMS), following the approach in AerChemMIP (Collins et al., 2017) with changes as follows. New in AerChemMIP2 is the *piClim-2xWet* experiment, in which the CH₄ emissions from wetlands are doubled. Output from that experiment allows for a first multi-model assessment of the role of wetlands. The state dependence of the interactive emissions (if any) is not yet well understood. To what extent interactive emissions change due to a simulated warmer climate state compared to the preindustrial can be addressed with the newly added *piClim-p4K* experiment, where the sea surface temperatures (SSTs) of ice-free ocean grid cells are artificially increased from the model's PI control climatology by +4 Kelvin. Output of this experiment serves to aid feedback analysis by isolating the effect of warming on natural emissions, an approach inspired by experiments in the Cloud Feedback Model Intercomparison Project (CFMIP) protocol (Webb et al., 2017) and future scenario experiments with fixed PI sea-surface conditions (Zanis et al., 2022). Output from *piClim-p4K* can be used to isolate the response of emission changes to a temperature change and to eliminate any conflation between the biophysical and radiative effects of CO₂. The requested experiment *piClim-p4K* is the same as in RFMIP phase two (RFMIP2.0 Kramer et al., 2025) to help reduce the overall computational burden of modelling centres. Furthermore, we request a new *piClim-2xflash* experiment, which builds on the *piClim-2xNOx* experiment from AerChemMIP. In AerChemMIP, *piClim-2xNOx* doubled the amount of NO_x produced per lightning flash with respect to *piClim-control*. For AerChemMIP2, we instead request that the lightning flash rate itself is doubled. This approach will still increase the production of NO_x by lightning, but will also have further impacts on atmospheric composition and top-of-atmosphere radiative fluxes in models with interactive parameterizations for wildfire emissions that account for natural fire ignitions from model-derived lightning flashes.

3.2 Historical Experiments

In AerChemMIP, attribution experiments were performed in which an all-forcings historical experiment is compared with experiments in which selected species are held at PI levels, which became known as the all-but-one experimental design. Both fully coupled experiments (*hist-piX*) and experiments with prescribed sea-surface temperatures and sea ice (*histSST-piX*) were requested for CMIP6 covering the period 1850–2014. These experiments targeted aerosol and aerosol precursors (*hist-piAer* and *histSST-piAer*), non-CH₄ tropospheric O₃ precursors (*histSST-piO3*), CH₄ (*histSST-piCH4*), nitrous oxide

(*histSST-piN2O*), and halocarbons (*hist-1950HC* and *histSST-1950HC*). The *histSST-piX* experiments enabled the attribution of transient changes in radiative forcing and climate responses in some cases, as well as quantifying drivers of composition and air quality changes over the historical period (Stevenson et al., 2020). Building on that all-but-one attribution approach from AerChemMIP, AerChemMIP2 requests historical experiments for 1850–2021 inclusive, using CMIP7 forcing data sets as provided by the CMIP climate forcings task team (Durack et al., 2025). All experiments *histSST-piX* are requested to be consistent with historical experiments of CMIP7.

3.2.1 Attribution of climate and air quality responses

AerChemMIP2 experiments for the historical period aim to advance the scientific understanding of the contribution of anthropogenic SLCF emissions to atmospheric composition, air quality, and human-induced climate change, leveraging models that have new capabilities in process representation and use updated forcing data provided for CMIP7. Differences in aerosol radiative forcing arising from different emission inventories could be of the order of 0.1 Wm^{-2} (Lund et al., 2023). The *hist-piX* experiments help to improve the physical science basis of climate change. Specifically, *hist-piX* experiments are relevant for obtaining information on model performance through the evaluation against observational data, for better understanding drivers of observed climate and air quality changes associated with spatially varying anthropogenic SLCF emissions, particularly on regional scales. Such information is useful to explain, anticipate, and possibly mitigate changes we expect in the future. For instance, the role of anthropogenic aerosol emissions for regional climate and air quality responses remains an outstanding source of uncertainty in our understanding of human-influenced climate change (e.g., Bellouin et al., 2020; Persad et al., 2023). The experiments can help to quantify the uncertainty in the role of heterogeneous SLCFs in global and regional water cycle changes observed over the historical period. Results from the Precipitation Driver and Response MIP (PDRMIP, Myhre et al., 2017, 2018), AerChemMIP (Allen et al., 2020, 2021), DAMIP (Monerie et al., 2022) and RAMIP (Wilcox et al., 2020) pointed to the role of atmospheric composition in modulating regional precipitation changes, but large uncertainties remain. Reasons for the large range of model results may include model diversity in radiative forcing, model state biases and structural differences, and ensemble size that causes difficulty in obtaining a highly precise estimate of forced responses under variable weather conditions at regional scales.

The AerChemMIP2 historical experiments include the fully coupled Earth system model simulations listed in Table 5, namely, *hist-piAQ*, where anthropogenic SLCF emissions contributing to air pollution are set at PI levels, and *hist-piX*, where a choice of individual or group of SLCF emissions are set at PI levels. Experiment *hist-piAer* is identical to that in AerChemMIP phase 1, with aerosols and aerosol precursor emissions of BC, organic carbon (OC), ammonia (NH_3), and sulphur dioxide (SO_2) set to PI levels. The experiment *hist-piAQ* is the same as *hist-piNTCF* from AerChemMIP phase one (Collins et al., 2017) except the new name accurately reflects the experimental protocol wherein air pollutant emissions, including non- CH_4 tropospheric O_3 precursors, aerosols and their precursor emissions (BC, OC, NH_3 and SO_2) are set to PI levels. The experiment *hist-piAQ* can be used to diagnose the climate and air quality responses to the regionally heterogeneous evolution of anthropogenic non- CH_4 SLCF emissions, which falls under science questions 3 and 4. A single-forcing experiment for O_3 (*hist-piO3*) allows responses to O_3 from all tropospheric O_3 precursors (CH_4 , NMVOCs, CO, and NO_x) to be quantified which was not possible

415 in AerChemMIP phase one. We request that CH₄ concentrations evolve in the radiation scheme but concentrations or emissions
are fixed at PI levels in the chemistry treatment of *hist-piO3* to suppress CH₄-driven changes on surface-level and tropospheric
O₃ concentrations, documented elsewhere (e.g., Fiore et al., 2002). Another single-forcing experiment for CH₄ (*hist-piCH4*)
allows the net effect of CH₄ changes on atmospheric composition and associated climate responses to be cleanly assessed. The
420 fully coupled CMIP7 DECK historical experiment (Dunne et al., 2025) is needed as a reference for computing differences in
responses due to PI to PD aerosol and tropospheric O₃ precursor emissions changes.

AerChemMIP2 atmosphere-only historical experiments will help to provide policy-relevant information by quantitatively
relating a mass change in emissions to radiative forcing and temperature changes. Some interactions are indirect, e.g., perturbations
in non-CH₄ tropospheric O₃ precursor (NMVOCs, CO, and NO_x) emissions do not directly affect radiative forcing but do so
indirectly via their effects on O₃, CH₄ lifetime, aerosols, and cloud adjustments. Surface O₃ is a critical pollutant and its
425 precursor emissions are controlled to meet air quality standards. However, tropospheric O₃ is also a GHG so a policy-relevant
question is – what is the climate response and air quality impact from changes in O₃ precursor emissions? Similarly, what is the
implication of policy-induced emission reductions of aerosols and their precursors for air quality and unmasking warming due
to GHGs? The set of atmosphere-only historical experiments *histSST-X* listed in Table 6 help to address such policy-relevant
questions based on observed changes in emissions of SLCFs and CH₄. Since composition changes and air quality respond to
430 both climate and emission changes, we request complementary historical experiments with fixed PI sea-surface conditions and
transient emission changes (*piClim-histall*). It means *piClim-histall* can be used, for example, to establish the extent to which
changes in CH₄ lifetime over the historical period are driven by climate (e.g., Stevenson et al., 2020) and to determine the
associated climate penalty on air quality (Fu and Tian, 2019; Zanis et al., 2022; Murray et al., 2024; Akritidis et al., 2024).

3.2.2 Disentangling forced response from variability

435 Three-member ensembles are included in the AFT of CMIP7 for *hist-piAQ* and *hist-piAer*, but larger ensemble sizes would
be beneficial for separating forced responses from internal variability if modelling centres can afford to provide them. Models
which can simulate SLCFs other than aerosols (i.e., CHEM) should perform *hist-piAQ* while models without such capability
(i.e., AER) are asked to perform *hist-piAer*. The review of AerChemMIP (Griffiths et al., 2025) noted that a relatively small
ensemble size prohibited some analyses, particularly with regard to climate responses to anthropogenic emissions at the
440 regional scale. At least three ensemble members for all requested *hist-piX* experiments as this is seen as the minimum required
to disentangle responses from internal variability at large spatial scales, and has been identified as being important when
investigating the scale and uncertainty of future air pollution episodes (Fiore et al., 2022; Doherty et al., 2022), but larger
ensembles are necessary for estimates of precipitation changes (Monerie et al., 2022). Three-member ensembles for these
experiments is also desirable for comparability of the results to the *hist-aer* experiments from the Detection and Attribution
445 Model Intercomparison Project v2.0 (DAMIP v2.0, Gillett et al., 2025).

We encourage the creation of a larger ensemble size to study pattern effects where modelling centres have the computational
capacity, e.g., as was done for the Community Earth System Model (CESM, Deser et al., 2020a; Simpson et al., 2023) and
the Seamless System for Prediction and EArth System Research (SPEAR, Delworth et al., 2020) models. Based on experience

in the Regional Aerosol Model Intercomparison Project (RAMIP, Wilcox et al., 2023), we suggest ten-member ensembles
450 of *hist-piAQ* (CHEM) or *hist-piAer* (AER) experiments per model. Creating ensembles of simulations implies a comparably
large request for computational resources but it is justified by the need to explain past climate change and the large interest
in the scientific exploitation of AerChemMIP’s coupled experiments (Griffiths et al., 2025; Fiedler et al., 2024). Since similar
experiments have also been performed for AerChemMIP phase one, changes in the understanding of CMIP7 results against
CMIP6 can be documented through a comparison of results from the *hist-piX* experiments for the overlapping period 1850–
455 2014.

For all fully coupled historical experiments, we request corresponding atmosphere-only experiments (*histSST-piAQ*, *histSST-
piAer*, *histSST-piO3*, and *histSST-piCH4* listed in Table 6), with prescribed time-varying boundary conditions (e.g., sea surface
temperatures and sea ice) taken from the CMIP7 historical experiment of the model to compute the historical changes in
effective radiative forcing. We request an additional single-forcing experiment for NO_x (*histSST-piNOx*) to assess the role
460 of anthropogenic NO_x as a precursor of tropospheric O₃ (Nguyen et al., 2022). A corresponding atmosphere-only experiment
including all emission changes and climate change (*histSST*) is needed as a reference for calculating differences in the chemical
and radiation budgets induced by the individual SLCF emission perturbations. The additional experiment *piClim-histall*, as in
RFMIP, is needed to separate the influence of emissions and climate change on atmospheric composition and air quality.

3.2.3 Non-linearity in the climate response

465 A cross-MIP collaboration between AerChemMIP2 and DAMIP v2.0 (Gillett et al., 2025) is enabled through the parallel
request for experiments for single forcing historical simulations for aerosols, following the “all-but-one” approach in AerChemMIP2
and the “only” approach in DAMIP, respectively. The parallel experiments allow non-linearities in climate responses to aerosols
to be systematically studied for the first time.

AerChemMIP2 follows again the all-but-one experimental design like in AerChemMIP (Collins et al., 2017). One reason
470 is to retain the direct comparability to AerChemMIP results. Another reason is that the all-but-one design is better suited for
studying responses of chemical interactions because the chemical composition of the atmosphere itself influences the response
of aerosols and air quality to emission changes, although neither of the two approaches is necessarily superior to the other for
studying climate response to atmospheric composition changes.

Having both experimental designs for aerosols in the framework of CMIP7 will facilitate the study of nonlinearities more
475 systematically than was possible in the past. For example, it might be found that simple emulators are insufficient to accurately
model the physical climate response to realistically co-varying chemical and aerosol forcings, for which machine learning
emulators might be better suited (e.g., Watson-Parris et al., 2022). Model results can differ for the two different experimental
setups, e.g., seen in CESM (Simpson et al., 2023). The ensemble of DAMIP and AerChemMIP available from CMIP6 is, to
that end, not conclusive since different models contributed to the two MIPs. For CMIP7, we therefore encourage modelling
480 centres to perform both the AerChemMIP2 *hist-piAer* experiment following the all-but-one approach of AerChemMIP2 and
the *hist-aer* experiment of DAMIP (Gillett et al., 2025).

3.2.4 Desert dust particles

Dust is known to have varied substantially over the historical period, showing both large decadal variability (Prospero and Lamb, 2003; Mahowald et al., 2010; Shao et al., 2013) and a long-term increase of 55 ± 30 % since PI times (Kok et al., 2023).
485 These large changes in dust have important implications for various Earth system processes, including for radiative forcing and biogeochemical feedbacks that fall under the science questions 1 and 2 of AerChemMIP2. Reproducing the past variability and trend of desert dust aerosols has been a longstanding challenge for CMIP-class models (Evan et al., 2014; Kok et al., 2018; Zhao et al., 2022; Kok et al., 2023). CMIP6 historical experiments failed to reproduce the past increase of desert dust aerosols, e.g., illustrated by the percentage change in dust aerosol optical depth in Fig. 3, which hinders a better understanding of the
490 implications of increased dust amounts for the climate response and biogeochemical feedbacks.

To address this knowledge gap, we request a new coupled historical experiment *hist-Dust* (Table 5), where the historical development of desert dust aerosols is prescribed while all other SLCFs are treated as in an historical experiment. Participating models are asked to use the dust emission data from Leung et al. (2025), which was obtained by combining an inversion of dust deposition fluxes from ice cores and other sedimentary records (Hooper and Marx, 2018) with constraints on the modern
495 day dust cycle (Kok et al., 2021a, b). Moreover, we ask for a parallel atmosphere-only historical and future experiments - *histSST-Dust* (Table 6) and *esm-scen7-vl-SST-Dust* (Table 7) - to diagnose the radiative effects of desert dust aerosols from the PI to the future. In addition to the historical experiments for dust, we request a future extension with a linear increase of dust aerosols in an overshoot scenario (*esm-scen7-vl-Dust*; see Section 3.3.1). Note that this future increase in dust is hypothetical in that there is currently no consensus on how dust will change in the future (Pu and Ginoux, 2017; Kok et al., 2023), although
500 that gap in our understanding is known for at least two decades (Mahowald and Luo, 2003; Tegen et al., 2004).

A pilot study using the dust dataset in AeroCom models is ongoing (<https://aerocom.met.no/experiments/DURF>) and successful tests of the dataset in CESM2 are complete. Specifically, the standard configuration of CESM2 failed to reproduce past dust changes like CMIP6 models (blue line in Fig. 3). Prescribing the dust dataset from Leung et al. (2025) in CESM2 yields global variability and trends in dust aerosol optical depth similar to the observational reconstruction from Kok et al. (2023) for the
505 entire historical period (black and red lines in Fig. 3). Community support for adjusting the dust data from Leung et al. (2025) to model-specific requirements will be available, e.g., adjustments of the emission data for different aerosol size bins and scaling the data over time to mimic the observed dust trends in the participating models. Such experience already exists, e.g., from the dust radiative forcing experiment in the AeroCom community. A flexible parameterization for prescribing changes of dust aerosol optical properties is currently developed for use in *hist-Dust* and *esm-scen7-vl-SST-Dust* in CMIP models without
510 interactive dust parameterization schemes.

3.2.5 Advancement in Fires

In AerChemMIP2, we expect contributions from models with and without biomass burning feedbacks. The mix of representations of biomass burning across contributing models has the potential to better link to assessments of fires based on experiments in AFT. Emissions from biomass burning are an important contributor to atmospheric composition and climate change (Ward

515 et al., 2012; Zhong et al., 2024), with regionally-varying trends (Earl and Simmonds, 2018; Jones et al., 2022). Extra-tropical
fire emissions have increased with warming to a degree that is comparable to the reduction of fires in the tropics for 2001–2023
(Zheng et al., 2021; Jones et al., 2024). Biomass burning is also notable as one of the main drivers of interannual variability
in atmospheric composition (Voulgarakis et al., 2015). Simulating the variability in biomass burning may result in a less
520 negative anthropogenic aerosol radiative forcing than prescribing time-averaged biomass-burning emissions, e.g., due to non-
linear interactions (Heyblom et al., 2023), as well as surprising coupled responses (Fasullo et al., 2022; Heyblom et al., 2022).
Biomass burning is also a major contributor to poor air quality with concomitant effects on human health (Xue et al., 2021;
Xu et al., 2024), which similarly require simulating the variability in fire emissions for a full characterization due to non-linear
relationships between pollutant concentrations and health impacts.

Historical fire datasets produced for use in CMIP (BB4CMIP, van Marle et al., 2017) show minimal interannual variability
525 in CO₂ emissions from fires before the availability of satellite remotely-sensed burned area, in contrast with the historical fire
emissions that were diagnosed by ESMs with interactive fire capabilities in CMIP6 (Figure 4). Moreover, the earlier start in use
of recorded visibility data from weather stations in 1960s influenced the representation of interannual variability in emissions
data (not shown). Interactively simulated biomass burning emissions across CMIP6 models varied by up to a factor of five
(Figure 5). Most individual historical experiments show much more biomass burning emissions than the CMIP6 and CMIP7
530 climate forcings datasets. The biomass burning prescribed emissions datasets from the Shared Socioeconomic Pathways (SSPs)
in CMIP6 lack future climate-driven changes in biomass burning, whereas evidence for a substantial increase of regional fire
activity with warming exists (Dowdy et al., 2019; Abatzoglou et al., 2019; Jones et al., 2022; Galizia et al., 2023).

AerChemMIP2 encourages modelling centres with the capability to perform experiments with interactive fires, to perform
piControl and *hist* experiments with fire feedbacks switched on. The additional *hist-piFire* (Table 5) experiment, with SLCF
535 emissions from fires held fixed at the PI level but all other forcings evolve as in *hist* allows an assessment of the role of fire
emissions on climate change and air quality, which is a contribution to science questions 1 and 3. Models with the capacity
to perform experiments with interactive fires that use it in their model configuration for other experiments of CMIP7 and
AerChemMIP2, would only need to additionally perform *hist-piFire*. As noted above, temporal smoothing of models' biomass-
burning emissions may introduce a change in radiative forcing, and therefore for *hist-piFire*, we recommend that models with
540 interactive fires prescribe monthly fire emissions taken from multiple decades of a PI control simulation in order to maintain
both the same seasonal and interannual variability in emissions, rather than prescribing a monthly climatology which is repeated
each year. Only SLCF emissions from fires should be held at PI levels, with everything else free to evolve as in *hist*.

We similarly encourage modelling centres to perform future scenario experiments with interactive fires to address concerns
that fire activities increase with warming (Jones et al., 2022) and anthropogenic aerosol reductions (Allen et al., 2024). In
545 addition to the direct impact of fires on lives, livelihoods, and property, fire also accounts for about half of the carbonaceous
aerosol emissions globally (Jones et al., 2024), and paired with the projected reduction in anthropogenic SLCF emissions in
ScenarioMIP-CMIP7 (van Vuuren et al., 2025) could become a more significant source of SLCFs in the future compared to the
past. Despite the consequences, the implications of fires for future atmospheric composition, radiative forcing, and air quality
are not currently fully explored or well-quantified. To address this gap in knowledge, we ask those models that use interactive

550 fires in their default configuration to perform the experiments *esm-scen7-h* and *esm-scen7-vl* of ScenarioMIP-CMIP7 with fire feedbacks switched on and to submit associated emissions as diagnostic output. More specific model experiments with a focus on fires are part of FireMIP (Li et al., 2025).

3.3 Future Scenario Experiments

Potential future changes in air quality and climate change are tightly connected. SLCFs are co-emitted with CO₂, such that
555 CO₂ emission reduction targets have a co-benefit of improving air quality (Turnock et al., 2019). Moreover, future climate change can have a regional influence on surface concentrations of pollutants, e.g., by enhancing or decreasing O₃ or aerosol particle concentrations over populated areas (e.g., Fu and Tian, 2019; Zanis et al., 2022; Murray et al., 2024; Akritidis et al., 2024). Likewise, Earth system feedbacks in a warming world could have consequences for future air quality (Gomez et al., 2023). The degree to which different SLCFs are reduced, either individually or in combination, can have potentially large
560 impacts on both near-term climate forcing and air quality. Figure 6 (adapted from results in Turnock et al., 2022) highlights that co-benefits and penalties to air quality and climate can occur depending on the emission pathway of each scenario, with the ideal aim of maximizing future co-benefits. Aggressive aerosol mitigation measures are, for instance, thought to be strong enough to threaten the 2-degree warming goal of the Paris Climate Agreement (Wood et al., 2024), by unmasking the warming associated with GHGs (Allen et al., 2021; Turnock et al., 2022). As such, strong mitigation of SLCF emissions that imply
565 improved air quality in some scenarios of ScenarioMIP-CMIP7 ensures that future warming from GHGs will be clearly visible due to reduced aerosol effects (van Vuuren et al., 2025).

AerChemMIP2 requests a set of variants of ScenarioMIP-CMIP7 scenarios to systematically assess the influence of individual climate forcers, such as CH₄ in two differently warming worlds. These experiments can provide information on the effectiveness of net-zero policies, and the implication of their potential failure for air quality and climate change mitigation. As such,
570 AerChemMIP2 addresses the implication of a near-term reduction in SLCF emissions for air quality and climate change to allow investigations that fall under scientific questions 3 and 4. To that end AerChemMIP2 chooses two tier 1 scenarios of ScenarioMIP-CMIP7 (van Vuuren et al., 2025) as baselines, which has been coordinated with CMIP7 and its registered MIPs to reduce the overall computational burden for modelling centres. Specifically, AerChemMIP2 uses the scenario Very Low Low Overshoot (VL) that follows an SSP1 trajectory assuming low air pollution and the high-end (H) scenario that follows an SSP3
575 development with emissions that adversely affect air quality. The relatively clean VL scenario (*esm-scen7-vl*) assumes limiting global warming to 1.5°C in the 21st century with a temporally limited small overshoot of the temperature limit, whereas the polluted H scenario (*esm-scen7-h*) assumes a lack of ambitious mitigation of climate change which also deteriorates air quality (van Vuuren et al., 2025).

While these scenarios have not yet been finalized, their description in van Vuuren et al. (2025) indicates reductions of
580 emissions across the board for VL, for air pollutants and greenhouse gases alike. Such changes can be driven by a decrease in the use of coal, gas, and oil in the energy system, which are replaced mainly by non-biomass renewables (Figure 7a). Conversely, the H scenario would see stable or increasing trends in emissions, with no fossil fuel phase out or phase down. For instance for SO₂ emissions, the differentiated pathways for coal use, from phase-out to continued or increased use, lead to

strong future differences between the two scenarios (Figure 7b). Depending both on how much vegetation is modelled in the
585 future in these scenarios and how much fire management is assumed, emissions from open burning fires may be different as
well. While for SO₂, emissions from fires are a small part of global total emissions, this share is larger for some other species
such as OC.

The spatial differences in future energy sources between the two scenarios lead also to diverging regional patterns of SLCF
emissions in the two future projections (Figure 8). In the middle of the 21st century (2050), VL assumes a level of emissions
590 consistent with a substantial reduction of the use of conventional energy sources in stark contrast to H. Several regions such as
sub-Saharan Africa and India could see increases in the use of coal in H, while China could keep high levels of coal use for
electricity as well (not shown). That development is in contrast to VL which could see a projected (near) phase-out of the use
of these conventional energy sources consistent with low emissions of SLCFs and CH₄ (Figure 7b). Similarly, oil and gas use
could continue to increase in H, while growth in renewables slows down.

595 3.3.1 SLCF emission trajectories

The AerChemMIP2 future scenarios *esm-scen7-h-X* and *esm-scen7-vl-AQ* are fully coupled experiments and variants of the
two high-priority ScenarioMIP-CMIP7 experiments VL and H. They are fully consistent with the underlying socio-economic
developments of ScenarioMIP-CMIP7 with variants of the future development of SLCFs (Tables 5 and 7). Specifically, these
experiments allow an assessment of the implication of SLCFs for atmospheric composition, air quality, and climate response
600 by isolating the influence of SLCF mitigation compared to the baseline scenarios defined by ScenarioMIP-CMIP7 (van Vuuren
et al., 2025).

The AerChemMIP2 variants of the H scenario are to quantify the role of SLCF emissions in a future with high GHG
emissions driven by little ambition for climate change mitigation. To that end, we request experiment *esm-scen7-h-AQ* (Table
5) in which aerosols, O₃ and their precursor emissions are set to PD level while all other emissions follow the h scenario.
605 The setup allows *esm-scen7-h-AQ* to be a consistent extension of the historical experiment with a future trajectory of strong
warming (AerChemMIP2 tier 1). Through comparison of *esm-scen7-h-AQ* against *esm-scen7-h* the potential influence of a
future implementation of clean air policies for minimizing air pollution despite a lack of climate change mitigation can be
assessed (upper left hand quadrant in Fig. 6). All emissions in *esm-scen7-h-AQ* will follow the prescribed spatio-temporally
evolving emissions as in *esm-scen7-h*, except for aerosols, O₃ and their precursors. Aerosol influences can be separately
610 assessed through the additionally requested experiment *esm-scen7-h-Aer* in models with interactive chemistry. Models without
interactive chemistry are asked to prioritize *esm-scen7-h-Aer* over *esm-scen7-h-AQ*.

The variants of the VL scenario explore the role of individual SLCFs for climate change in an overshoot of warming before
the world follows a trajectory towards climate-neutral conditions (*esm-scen7-vl-AQ*). Experiments *esm-scen7-vl-X* assumes
higher SLCF emissions, e.g., taken from *esm-scen7-h*, to mimic a theoretical failure of policies for cleaning the air. In contrast to
615 the different long-term pathways of *esm-scen7-vl* and *esm-scen7-h*, their expected near-term global mean temperature increase
until 2050 might be similar (van Vuuren et al., 2025). AerChemMIP2 could exploit this near-term similarity between those
scenarios to explore the impact of different spatial patterns of SLCF emissions for the sensitivity scenario *esm-scen7-vl-AQ*

compared to *esm-scen7-h-AQ*, and *esm-scen7-vl-Aer* compared to *esm-scen7-h-Aer*. The different regional patterns in SLCF emissions could, for instance, mimic a theoretical SLCF emission increase in regions with delayed action in controlling air pollution during economic growth (right hand side of Fig 6).
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AerChemMIP2 also addresses the future air quality and climate response that might arise from more desert dust aerosols through the sensitivity scenario *esm-scen7-vl-Dust* (Table 5) in which a future extension of the desert dust aerosol increase from the dataset by Leung et al. (2025) is prescribed (compare Section 3.2.4). The experiment assumes a hypothetical increase in dust-aerosols over time with a rate similar to what has been seen for past decades. The experiment applies a linear increase to the dust emissions on the PD spatial pattern of sources. The future extension is part of the dust dataset listed for AerChemMIP2 and CMIP6plus via input datasets for Model Intercomparison Projects (ESGF input4MIPs, 2025). In so doing, *esm-scen7-vl-Dust* is a seamless future extension of *hist-Dust*. In comparison to output from *esm-scen7-vl-Aer*, *esm-scen7-vl-Dust* enables to study the potential air quality and climate responses to a future with a continuous desert dust increase to be assessed along with SLCF policies targeting anthropogenic sources for improving air quality. The idealized design of experiment *esm-scen7-vl-Dust* allows us to investigate to what extent the climate response to dust forcing might be linear in a future scenario targeting net zero.
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We request three-member ensembles for each of the future scenario experiments in AerChemMIP2 as part of CMIP7-AFT. Models without interactive chemistry schemes (i.e., AER) are encouraged to contribute *esm-scen7-h-Aer*, *esm-scen7-vl-Aer* and *esm-scen7-vl-Dust* simulations, with a lower priority to create *esm-scen7-vl-AQ* and *esm-scen7-h-AQ* also. The AerChemMIP2 future experiments will allow the community to quantify the role of future mitigation actions for SLCFs on climate and air quality responses (i.e. which part of Fig 6 do these actions lead to), which can otherwise not be diagnosed from other CMIP7 output. For instance, *esm-scen7-h-Aer* and *esm-scen7-vl-Aer* in comparison to *esm-scen7-h* and *esm-scen7-vl* can be used to estimate how much regional warming might be masked by aerosol particles. The experiments can therefore inform policymakers on near- and long-term impacts arising from future SLCF emission changes targeting air quality and climate priorities. The parallel experiments, using ScenarioMIP-CMIP7 experiments as baselines, keep the computational burden for AerChemMIP2 smaller and allow a direct link to the scenarios used for CMIP7.
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Analogous AerChemMIP future atmosphere-only scenario experiments *esm-scen7-vl-SST-X* *esm-scen7-h-SST* are requested and listed in Table 7, e.g., to diagnose the time-evolving effective radiative forcing of SLCFs in the scenarios described above. Prescribed monthly sea-surface temperatures and sea-ice should be from the model's own fully coupled simulation of *esm-scen7-vl* and *esm-scen7-h*, respectively. Models with emission fluxes that depend on the sea-surface state should also prescribe these as time-evolving fields taken from their own fully-coupled simulations. The perturbation experiments *esm-scen7-vl-SST-X* and *esm-scen7-h-SST-X* use the same forcings as for their fully coupled future experiment counterparts. Moreover, we request two additional future sensitivity scenarios for the individual perturbations of CH₄ and O₃, with *esm-scen7-vl-SST* as the reference. These additional experiments can be used to address the potential future implication of different CH₄ (*esm-scen7-vl-SST-CH4*) and O₃ precursor (*esm-scen7-vl-SST-O3*) emissions compared to the development in the VL scenario. Specifically, we ask to prescribe the emissions of the polluted pathway in H (not VL) for CH₄ in *esm-scen7-vl-SST-CH4* and
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for O₃ precursors in *esm-scen7-vl-SST-O3*. The *esm-scen7-vl-SST-X* experiments are deliberately counterfactual scenarios that mimic a theoretical failure of air pollution mitigation in a world that limits warming to 1.5°C.

3.3.2 Disentangling influences of re/afforestation

655 AerChemMIP2 aims to address the potential influence of future large-scale forest expansions on atmospheric composition and climate change. Afforestation and reforestation (A/R) efforts are among the most widely-suggested climate change mitigation strategies with a high likelihood of deployment in one form or another (Verdone and Seidl, 2017). A/R will affect the Earth's radiative budget by changing surface albedo (Betts, 2000) and atmospheric composition that can offset a substantial part of the cooling associated with the CO₂ uptake by forests (Weber et al., 2024). In addition to sequestering CO₂, forests emit large
660 quantities of BVOCs (Guenther et al., 2012) which react chemically, affecting CH₄ lifetime, O₃ and aerosol abundances, as well as water vapour and cloud properties (Weber et al., 2022). Forest fires are also important sources of reactive gases and aerosols (Ward et al., 2012) while changes in land cover will also affect dust emissions, which influence the Earth's radiative budget (Thornhill et al., 2021a). Influences of A/R on fire and dust emissions have not been assessed in this context. Thus, the net impact of A/R on climate, and so its efficacy as a mitigation strategy, requires a systematic assessment of affected processes
665 alongside the benefit of CO₂ sequestration.

The response of atmospheric composition to A/R is also dependent on climate. BVOC emissions as well as fire frequency and severity, and thus associated emissions depend on temperature (Zheng et al., 2021; Burton et al., 2024). Moreover, mineral-dust emissions depend on meteorological and surface conditions (Shao et al., 2011). The impact on composition and climate from a change in BVOC and fire emissions further depends on the contemporaneous anthropogenic emissions of reactive gases
670 and aerosols. For example, a reduction in anthropogenic aerosol emissions amplifies the climate impact of aerosols from A/R (Carslaw et al., 2013) while anthropogenic NO_x emissions influence the ozone-forming potential of BVOCs (Seinfeld and Pandis, 1998). Trees themselves can also be damaged by elevated levels of ozone (Cheesman et al., 2024), leading to reduced productivity and carbon sequestration. Therefore, an assessment of the net climate impact of A/R must include changes to non-CO₂ processes and be embedded in multiple possible future transient climate scenarios which span the range of possible
675 future surface temperatures (Weber et al., 2024).

To simulate A/R, AerChemMIP2 will use the fully coupled *esm-scen7-vl* simulation of ScenarioMIP-CMIP7 (van Vuuren et al., 2025), specifically its land use which will feature some level of A/R. The VL scenario assumes a large reliance on negative emission technologies and so is the best option for A/R for AerChemMIP2. In some ESMs participating in ScenarioMIP-CMIP7, the land cover is preferably simulated interactively, e.g., through coupled vegetation schemes (Cox, 2001; Sellar et al.,
680 2019). Output from ESMs with prescribed evolving land cover consistent with the climate forcing data for the VL scenario are equally welcome and should include information on the land-use treatment in the metadata information. Modelling centres should use the same land cover treatment as for their *esm-scen7-vl* experiment for consistency with the AerChemMIP2 land use (LU) experiments and ScenarioMIP-CMIP7. We encourage modelling centres to run all experiments with interactive chemistry and aerosol (CHEM), e.g., to account for the influence via BVOC and fire emissions. Some of the experiments can also be

685 performed by models without such capability (AER), since these can nevertheless simulate the influence of land-use changes on surface albedo.

We request six atmosphere-only experiments with prescribed changes in sea-surface temperature and sea-ice with details listed in Table 8. These simulations allow the calculation of the net impact on the Earth's radiation budget from the expansion of tree cover simulated in the coupled *esm-scen7-vl* simulation and, separately, allow the radiative impact of non-CO₂ composition and surface albedo to be isolated. These experiments are consistent with the *esm-scen7-vl-SST* and *esm-scen7-h-SST* experiments (Table 7) concerning climate forcings such as GHGs and anthropogenic SLCF emissions. The two scenarios *esm-scen7-vl* and *esm-scen7-h* are again chosen as references since they span different anthropogenic emission pathways and different future surface temperature evolutions. Thus, these simulations will provide information as to the likely impact of A/R in a future where, after some delay, action is taken to mitigate climate change (VL) and a future where A/R represents the only mitigation method deployed (H).
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The anticipated information from the set of future land-use experiments is summarized in Fig 9. For the *esm-scen7-vl* (VL) background, the impact of A/R on the radiative budget comes from the comparison of *esm-scen7-vl-SST-pdLU-BFD*, which uses a fixed PD LU climatology and transient SSTs from VL that can influence BVOC, fire and dust emissions, and *esm-scen7-vl-SST*, which uses transient LU from VL capturing A/R. This can be considered as the difference between a scenario where A/R is pursued (*esm-scen7-vl-SST*) and a scenario where SSTs continue to evolve but LU is kept fixed at PD levels. Likewise, for the *esm-scen7-h* background, comparison of *esm-scen7-h-SST-pdLU-BFD* and *esm-scen7-h-SST-vl-LU* will yield the full impact of A/R on the radiation budget. Note that while SSTs follow the *esm-scen7-h* scenario, the LU in *esm-scen7-h-SST-vl-LU* follows that in VL since that scenario has A/R. The experiments *esm-scen7-vl-SST-pdLU* and *esm-scen7-h-SST-vl-LU* use the model's own prescribed PD climatology for land-use (based on that recommended for CMIP7), but BVOC, fire and dust emissions prescribed from *esm-scen7-vl-SST* and *esm-scen7-h-SST* respectively.
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Comparison of *esm-scen7-vl-SST-pdLU-BFD* against *esm-scen7-vl-SST-pdLU* and *esm-scen7-h-SST-pdLU-BFD* against *esm-scen7-h-SST-pdLU* isolates the influence of changes to BVOC, fire, and dust emissions from LU change. Likewise, comparison of *esm-scen7-vl-SST-pdLU* against *esm-scen7-vl-SST* and *esm-scen7-h-SST-pdLU* against *esm-scen7-h-SST-vl-LU* isolates the radiative impact of changes to surface albedo due to LU change. Thus the removal of atmospheric CO₂ by the biosphere due to A/R is not captured by the proposed experiments. Estimating the carbon removal for the A/R sink can be explored via other methods using output from *esm-scen7-vl*, e.g. as in Weber et al. (2024)). It would allow, for instance, for the reporting of an instantaneous radiative forcing due to A/R's CO₂ removal.
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The A/R experiments have different tiers. Simulation *esm-scen7-vl-SST*, also used as a reference for other future scenarios in AerChemMIP2 (Table 7) is Tier 1 while the two additional simulations based on VL are Tier 2. All simulations using the H scenario as a reference are Tier 3 and will simulate multiple influences, including fire and BVOC emission feedbacks and changes to vegetation productivity in a substantially warmer world to be explored. Combining A/R from VL with the H scenario can be interpreted as a future pathway where mitigation via A/R is chosen after climate change impacts have been experienced for longer than projected in the VL scenario. The AerChemMIP2 experiments *esm-scen7-h-SST-X* therefore help to build an understanding of the effectiveness of A/R as a mitigation policy in an even warmer world.
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720 Technical support will be available for modelling centres to perform the A/R experiments and we encourage interested
centres to contact the authors. We also note that a subset of the simulations can be done by models which don't have interactive
aerosol and/or chemistry since useful information about the forcing from surface albedo and water vapour changes can still be
extracted. Specifically, *esm-scen7-vl-SST* and *esm-scen7-vl-SST-pdLU*, which differ only in the land use, can be done by models
with a minimum capability. Such contributions from CMIP7 models that would typically not participate in AerChemMIP2, for
725 example, would be informative for the multi-model differences in surface-albedo changes due to A/R.

4 Diagnostic request

To provide further consistency with AerChemMIP and minimize model development overhead, the diagnostic request for
AerChemMIP2 builds on the AerChemMIP and RAMIP requests. Recent analysis has highlighted the most commonly used
CMIP variables (Juckes et al., 2025), but there remains a long tail of rarely used valuable outputs that support key science
730 goals of AerChemMIP2. Following the new data request protocol of CMIP7, a review of the CMIP6 AerChemMIP request
was made, a minor number of deletions were made, and new variables were added through a series of community consultation
meetings. Together, these variables were proposed as a scientific “opportunity” to CMIP7 and included in version 1.2 of the
CMIP7 harmonized data request for simulations in the AFT (Anstey and Ellis, 2025) so that they can be included in key
baseline DECK and ScenarioMIP-CMIP7 simulations. For new additions, this involved proposing new names to be included
735 in the Climate and Forecast (CF) standard (e.g., for the tendency of atmosphere mole concentration of ozone due to net
chemical production). Moreover, it required proposing new physical parameters (e.g., do3chm), proposing new variable names
(a combination of both the physical parameter and its temporal and spatial sampling; e.g., AERmon.do3chm), and organising
all variables into groups (e.g., aerchemmip_3d_monthly).

For AerChemMIP2, the data request is organised into 10 variable groups. Additional lower-priority variables will be added
740 to later versions of the data request in order to provide driving fields for offline chemical transport models (CTMs). Further
details on how the v1.2 data request for the Atmosphere, including AerChemMIP2, was developed can be found in Dingley
et al. (2025).

4.1 Aerosol properties for forcing uncertainty

We take this opportunity to highlight the value of modelling centres providing key diagnostics. For example, while requested in
745 CMIP6, aerosol optical depth (AOD) was not routinely output by models even though it can provide a crucial constraint on the
aerosol direct radiative forcing (ERFari) (Watson-Parris et al., 2020). In turn, further details, including the absorption AOD or
the single scattering albedo for natural and anthropogenic aerosol provides valuable context for their relative contributions to
absorption. CMIP models show considerable uncertainty in the forcing magnitude (Smith et al., 2020; Thornhill et al., 2021b;
Fiedler et al., 2023) and the aerosol optical properties are measurable variables to better understand and constrain the simulated
750 aerosol ERFari. In a possible synergy with the Geoengineering Model Intercomparison Project (GeoMIP Visioni et al., 2025),
we also request that models output the three-dimensional fields of aerosol extinction on native model levels for comparison

with observational products (such as from CALIOP) and the diagnosis of stratospheric AOD for separating historical volcanic eruptions in total AOD.

755 A large component of the aerosol forcing uncertainty stems from aerosol-cloud interactions (ERFari; Bellouin et al., 2020). We therefore request key diagnostics such as the cloud droplet number and optical depth. In combination with diagnostics requested by the next phase of the Cloud Feedback MIP (CFMIP, Webb et al., 2017), the output can help to constrain these variables influencing ERFari against historical observations from remote sensing instruments. Synergies with the Clouds and the Earth's Radiant Energy System MIP (CERESMIP, Schmidt et al., 2023) exist to constrain the radiation budget. To further understand the impact on air quality, and health, it is requested that as many modelling centres as possible provide a direct
760 output of surface concentrations of fine particulate matter ($PM_{2.5}$), in addition to the request for individual aerosol components. This will avoid any underestimation from an approximate calculation of this metric after AerChemMIP2. The model-dependent species included in $PM_{2.5}$ diagnostics should be documented. Consistent calculations of $PM_{2.5}$ across multiple-models are additionally possible (Allen et al., 2020).

4.2 Driving fields for Chemistry Transport Models

765 Studies of future impacts of climate change on surface air quality and stratospheric ozone have traditionally fallen under the realm of free-running climate models that include atmospheric chemistry, due to the necessity of simulating future meteorology. However, there are well-established international modelling initiatives independent of CMIP that study air quality in the present day using CTMs (e.g., Galmarini et al., 2017). CTMs use prescribed meteorology, usually from historical meteorological reanalyses, and therefore have better signal-to-noise ratios for atmospheric composition changes from emission perturbations.
770 They are also able to spend more computational power on more elaborate chemistry mechanisms, as well as perform more sensitivity simulations. One example is H_2 for which CTM simulations would enable a better understanding of its role for atmospheric composition.

For AerChemMIP2, we request that modelling centres provide the PD and future archived meteorology that would be necessary for driving CTMs, which typically require hourly temporal resolutions for 2-D fields and 3-hourly resolution for
775 3-D fields (Dingley et al., 2025). In later versions of the data request than v1.2, we will specifically request the necessary meteorological fields for driving the widely used GEOS-Chem CTM (<https://geos-chem.org>), which is normally driven by reanalysis products from the NASA Global modelling and Assimilation Office. By using different meteorologies from different models to drive a CTM with a consistent chemical mechanism, we can explicitly isolate the impact of different meteorological processes alone on atmospheric composition and surface air pollution. Nevertheless, we realize that this is a large data request,
780 so it is lower in priority than other diagnostics and is only requested from one to two transient coupled simulations, e.g., from a single realization of both a *hist* and a future scenario experiment.

5 Summary

AerChemMIP2 offers an experimental protocol that facilitates an Earth system model intercomparison, including benchmarking experiments, extended model validation via comparison with observations and other state-of-the-science models. The experiments
785 enable new scientific studies aimed at the role of SLCFs in the Earth system based on state-of-the-science models and forcing data sets, with assessment and quantification of radiative forcing, Earth system feedbacks, carbon budgets and climate sensitivity. The focus is on short-lived climate forcers, methane and land use, targeting the most complex Earth system models currently available, with implications for atmospheric composition, air quality and associated policy development.

While some experiments from AerChemMIP phase one are repeated to update and track changes in the scientific understanding
790 since CMIP6, new experiments explore aspects that would otherwise not be addressed in the context of CMIP7. Specifically, AerChemMIP2 highlights eight scientific opportunities that will enable research studies (1) to advance the understanding of state-dependence of ERFs through new present-day climatology experiments with CMIP7, (2) to globally upscale the aerosol radiative effects for a new line of evidence using storm-resolving models, (3) to provide first multi-model estimates of ERF for hydrogen emissions consistent with CMIP7 forcings, (4) to assess the contributions of non-methane volatile organic compounds
795 to the present-day anthropogenic ERF, (5) to address non-linearity in climate responses through parallel experimental designs across MIP boundaries, (6) to fill knowledge gaps by inducing previously missing desert dust trends in CMIP7 models, (7) to exploit new model capabilities for interactively simulating wild fires, and (8) to disentangle influences of re- and afforestation for future atmospheric composition and climate responses. Updates are enabled through repeating selected AerChemMIP experiments that will allow computations of individual ERFs and biogeochemical feedbacks, attribution studies of climate
800 and air quality responses accounting for internal variability, and future oriented assessments based on emission trajectories consistent with ScenarioMIP-CMIP7.

Contributing to AerChemMIP2 provides the essential basis for multi-model assessments of radiative forcing, response and feedbacks from interactive simulations of atmospheric composition and climate changes, which justifies the computational needs. Contributing experiments to AerChemMIP2 is rewarding for modelling centres. We encourage participation in AerChemMIP2
805 with models of different complexity in terms of number and fidelity of represented processes as well as different spatial resolutions ranging from a hundred to a few kilometers. We encourage modelling centres to contribute to AerChemMIP2 experiments listed under all Tiers, which are meant as a help for temporally planning resources for completing experiments rather than misinterpreting tier 1 as being more important than tier 3 experiments. Indeed, tier 3 experiments from AerChemMIP phase one have for instance been cited similarly often as experiments in other tiers and informed IPCC AR6 thanks to
810 contributions from many (Griffiths et al., 2025). Output of AerChemMIP2 experiments will provide avenues to enhance understanding of air quality and climate interactions, to better quantify Earth system feedbacks involving short-lived climate forcers and methane, and to improve traceability and evolution of model performance in comparison to other contemporary models. Moreover, the participation of modelling centres will allow the community to update climate assessments and projections and to bear down on key uncertainties in our understanding of the role of short-lived climate forcers and methane on air quality
815 and climate change.

Data availability

News from AerChemMIP2 will be distributed via a mailing list (subscription by sending an empty email to: AERCHEMMIP2-subscribe-request@listserv.uni-heidelberg.de). Technical questions on experimental setups and diagnostic output for AerChemMIP2 can be directed to the CACTI committee: CACTI-committee@listserv.uni-heidelberg.de.

820 All output from AerChemMIP2 will be published following the same workflows as for CMIP7. Climate forcings data will be available via input4MIPs with details documented by the CMIP Climate Forcings Task Team. The AerChemMIP2 data request for medium and high priority variables from CMIP7 is available on AirTable v1.2 with details in Dingley et al. (2025). Lower priority variables will be added to later versions with updated information provided on the webpage for the CMIP7 data request.

825 Biomass burning emissions are taken from historical experiments of CMIP6 models available from ESGF (<https://aims2.llnl.gov/search/cmip6>, last access: Nov 2025), the BB4CMIP data set (CMIP6 - version 1.1, CMIP7 - version 2.0, van Marle et al., 2017) available via input4MIPs (<https://aims2.llnl.gov/search/input4mips/>, last access: Nov 2025), the Global Fire Assimilation System (GFAS version 1.0, Kaiser et al., 2012) from the Climate Data Store (<https://doi.org/10.24381/a05253c7>), the Fire Inventory from NCAR (FINN version 2.5, Wiedinmyer et al., 2023) downloaded from NCAR's Research Data Archive (<https://doi.org/10.5065/XNPA-AF09>), from Zhang et al. (2024) accessible on Harvard Dataverse (<https://doi.org/10.7910/DVN/KB0ESS>), and a beta data set from the Global Fire Emissions Database (<https://www.globalfiredata.org/data.html>, last access: Nov 2025). Dust data are taken from CMIP6 historical experiments from ESGF (<https://aims2.llnl.gov/search/cmip6>) and from UCLA 1.0.2: DustCOMM, made available via input4MIPs (<https://aims2.llnl.gov/search/input4mips/>, last access: Nov 2025). Data display from ScenarioMIP-CMIP7 has been curated by Elmar Krieger and Nico Bauer from REMIND-MAGPIE for the
830 vl scenario and by Christoph Bertram from GCAM for the h scenario.

Author contribution

The following authors had a primary role in developing the new experiments in AerChemMIP2: WJC, SF, MK, JFK, VN, FMO'C, DW-P, FP, ER, MS, JK, and JW. The AerChemMIP2 diagnostic request was developed via community engagement, with input from WJC, SF, PTG, MK, LTM, VN, STT, DW-P, and LJW, and co-ordinated and implemented in Airtable by
840 FMO'C. Figures were provided by JFK, JW, MK, STT, and SF. SF led the writing and the coordination for AerChemMIP2. All authors have reviewed and agreed on the content of the manuscript.

Competing interests

Some authors are members of the editorial board of Geoscientific Model Development. There are no competing interests.

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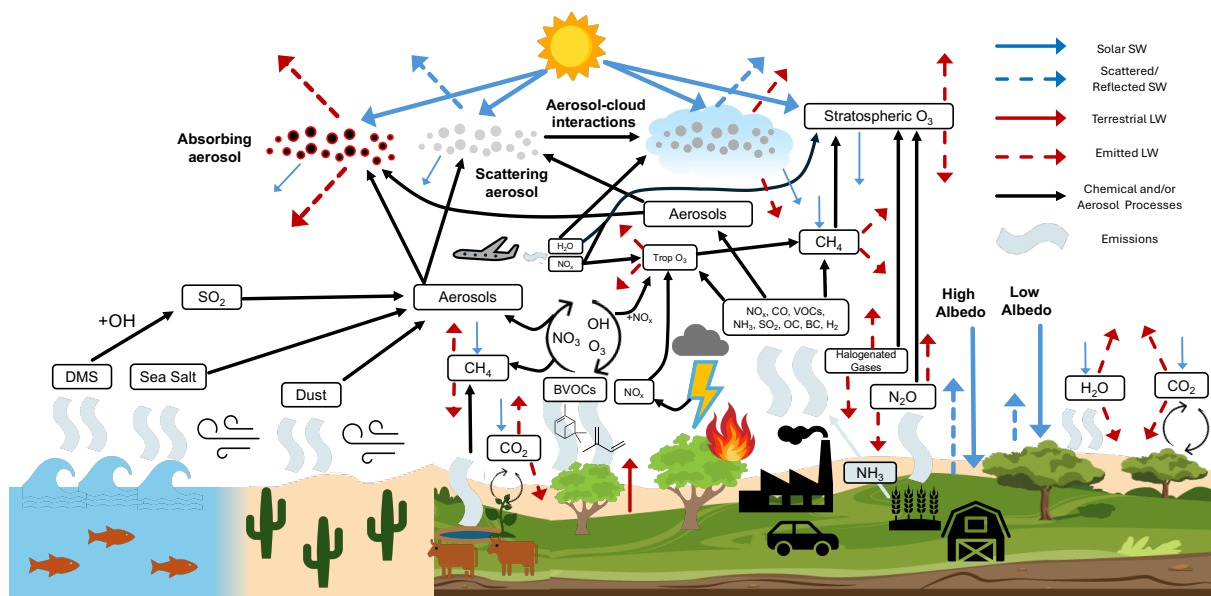


Figure 1. Schematic depicting the interaction of natural and anthropogenic trace gas and aerosol emissions with atmospheric composition, air quality, and climate response (Schematic inspired by Bonan, 2016; Fiore et al., 2015).

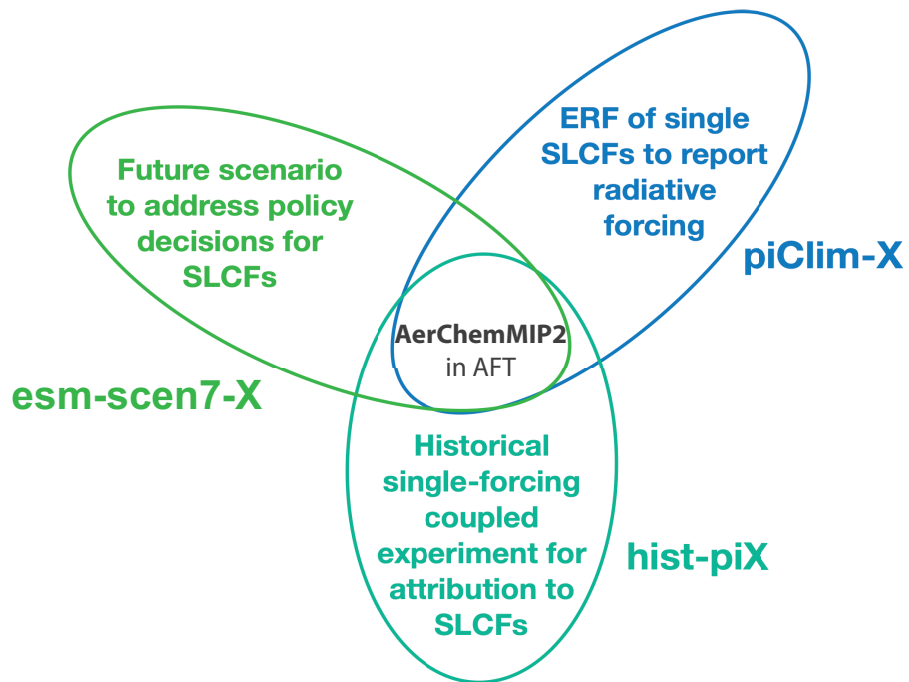


Figure 2. Overview of AerChemMIP2 experiments in the Assessment Fast Track (AFT) of CMIP7

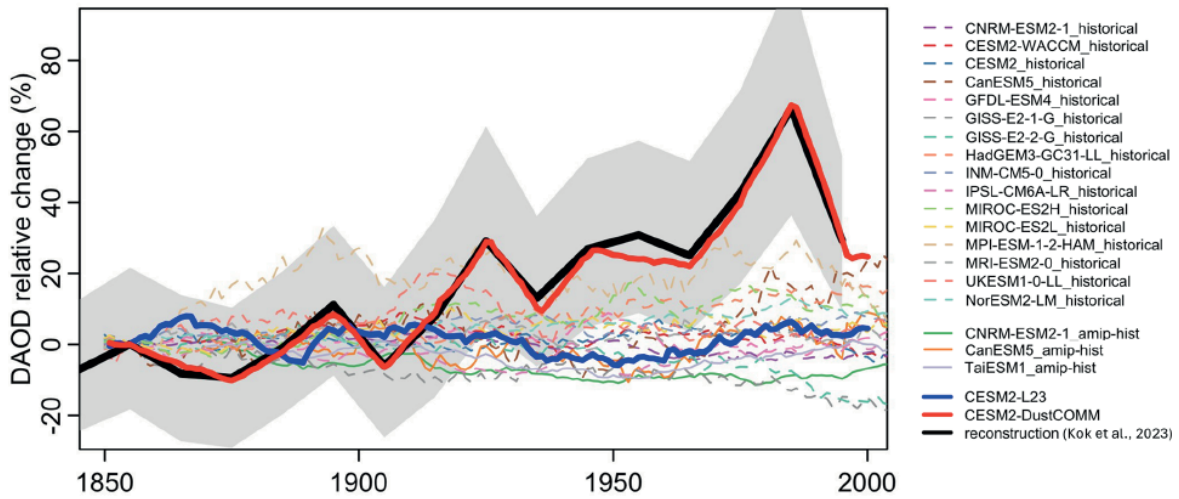


Figure 3. Historical development of the percentage change in dust aerosol optical depth. Shown are the ten-year running means of the dust aerosol optical depth (DAOD) from color-coded historical experiments of CMIP6 models from (dashed lines) fully coupled experiments and (solid lines) atmosphere-only experiments, (black) the historical reconstruction from Kok et al. (2023), and the proof-of-concept use of (red) the dust data set from Leung et al. (2025) in CESM2 for the proposed *hist-Dust* and *histSST-Dust* experiments of AerChemMIP2 against (blue) the results from the standard setup of CESM2. From Leung et al. (2025).

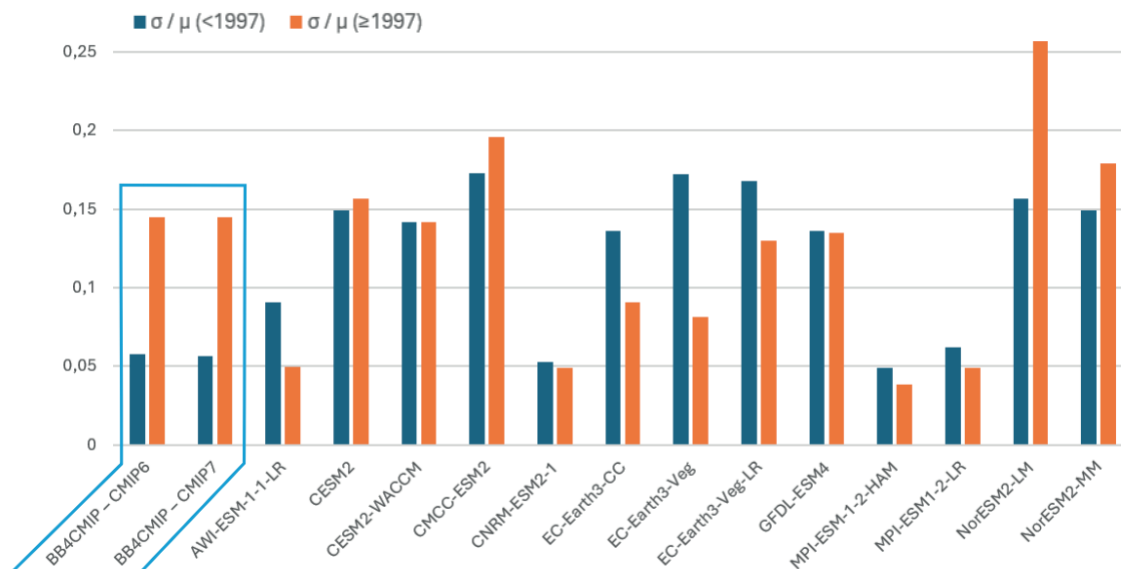


Figure 4. Relative interannual variability in fire emissions before and during the satellite era. Shown are the ratios of the year-to-year standard deviation normalized by the mean of annual total fire emissions of CO₂, for years (blue, 1850–1996 inclusive) before and (orange, 1997–2014 inclusive) during the availability of satellite data in the Global Fire Emissions Database (GFED) from the CMIP climate forcings data sets for prescribing biomass burning emissions in CMIP6 and CMIP7 (BB4CMIP CMIP6 version 1.1 and CMIP7 version 2.0, van Marle et al., 2017) visually highlighted by the blue frame, and individual historical experiments of CMIP6 models which interactively diagnosed fire emissions.

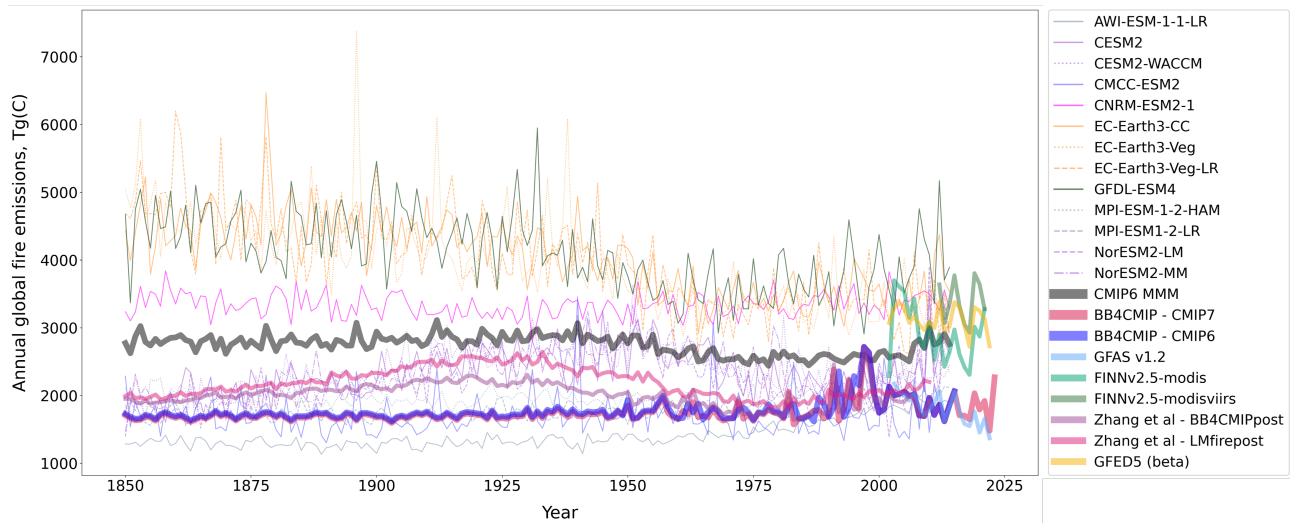


Figure 5. Historical development of CO₂ emissions associated with fires. Shown are annual global totals of CO₂ emission fluxes in Tg(C) from color-coded historical experiments of CMIP6 models of (thin lines) individual fully coupled experiments with interactive fire emission schemes switched on and (thick grey line) the multi-model mean of CMIP6, as well as (thick colored lines) reference data sets including the CMIP climate forcings data sets for CMIP6 and CMIP7 (BB4CMIP CMIP6 version 1.1 and CMIP7 version 2.0, van Marle et al., 2017), observational estimates for the satellite era from the Global Fire Emissions Database (GFED version 5), the Global Fire Assimilation System (GFAS version 1.2, Kaiser et al., 2012), the Fire Inventory from NCAR (FINN version 2.5, Wiedinmyer et al., 2023) and post-corrected data by Zhang et al. (2024). All data are shown for 1850–2014, except the retrieval products from satellite measurements.

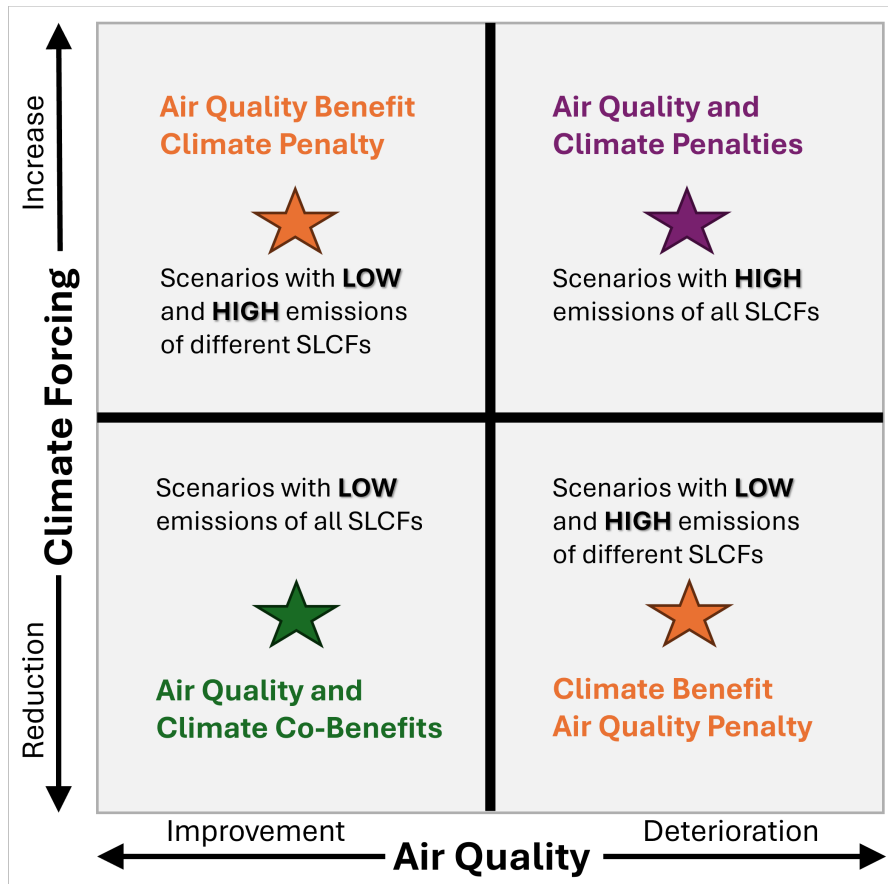
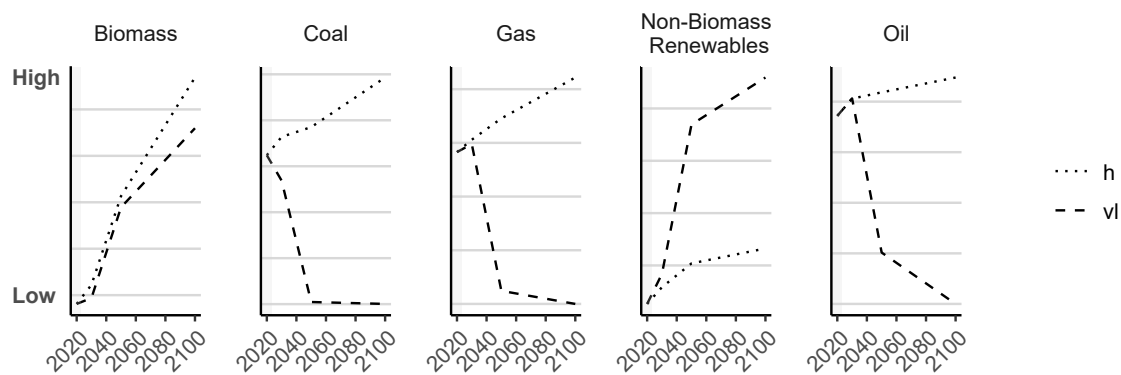


Figure 6. Schematic showing how the choice of SLCFs in future scenarios could impact both air quality and climate forcing, adapted from results in Turnock et al. (2022). While ScenarioMIP-CMIP7 will focus on the left quadrants, AerChemMIP2 will request experiments in the right quadrants to address the effectiveness of air quality policies in synergy with climate change developments.

A

Illustration of possible marker pathways

Global energy projections



B

Global emissions projections of CH₄ and SLCFs

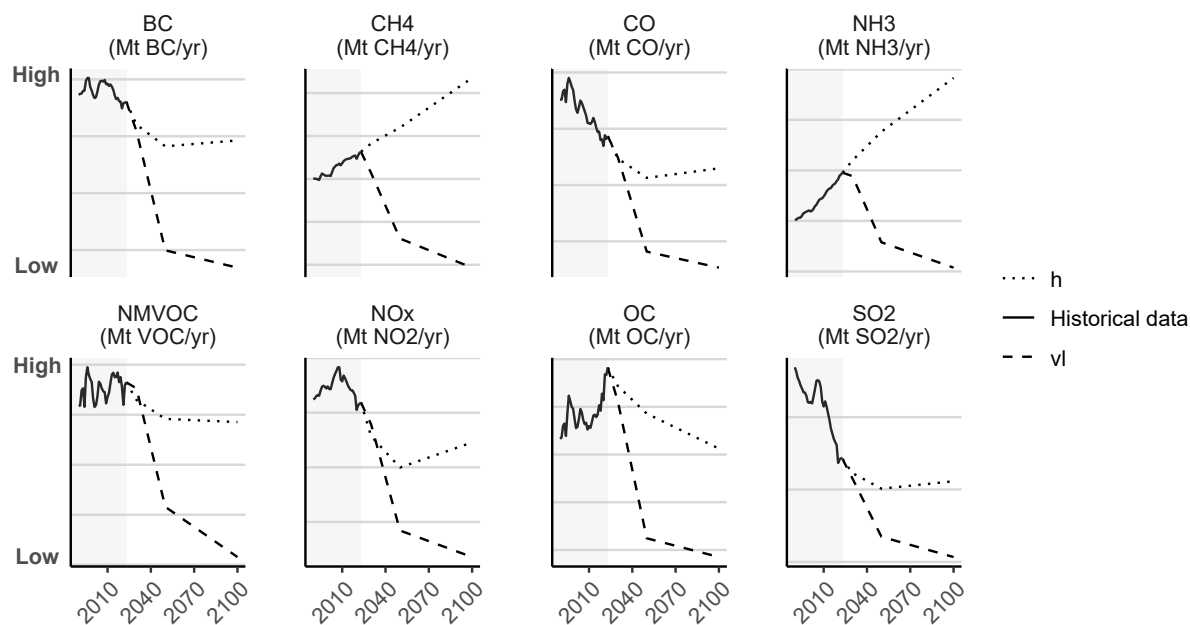


Figure 7. An indicative illustration of possible global trajectories for *esm-scen7-vl* (VL) and *esm-scen7-h* (H) marker pathways in CMIP7 scenarios for 2023–2100. Shown are (a) primary energy split by different energy carriers, and (b) CH₄ and a set of selected SLCFs, harmonized to historical emissions for CMIP7 until 2023 inclusive.

Illustrative Gridded SO₂ Emissions

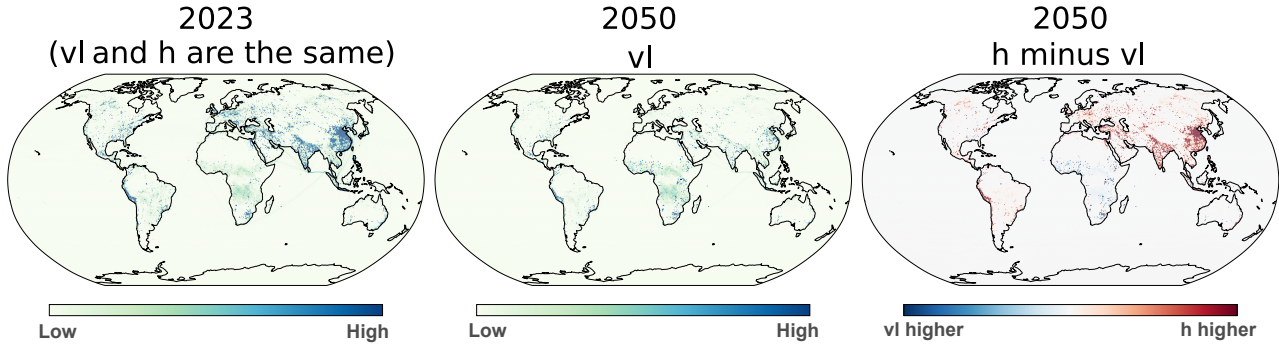


Figure 8. An indicative illustration of possible gridded SO₂ emissions for *esm-scen7-vl* (VL) and *esm-scen7-h* (H) marker pathways. Shown are the total emissions for (left to right) the harmonization year 2023 (end of historical data and start of scenarios), *esm-scen7-vl* in 2050, and their difference between *esm-scen7-vl* and *esm-scen7-h* for 2050. The emission totals include anthropogenic emissions at the surface, vertically integrated aircraft emissions, and open burning emissions.

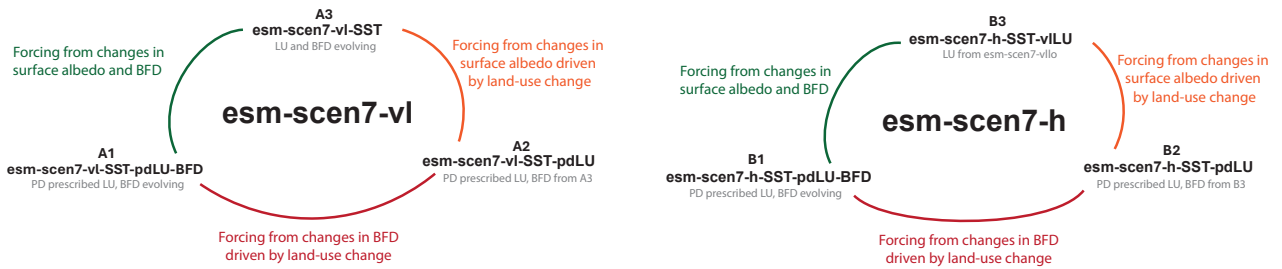


Figure 9. Schematic illustrating the relationship between the 6 simulations proposed for the A/R experiment. BFD refers to BVOC, fire and dust emissions. Anthropogenic aerosols and O₃ precursor emissions, SSTs and CO₂/CH₄/N₂O/CFC/HCFCs as in *esm-scen7-vl-SST* and *esm-scen7-h-SST*, respectively. Further details are in Tab. 8.

Table 1. Experiments for computing present-day anthropogenic ERFs relative to the pre-industrial period. *piClim-X* experiments are atmosphere-only, with a prescribed climatology of the model’s own pre-industrial (PI) sea-surface temperature and sea ice concentration, determined from the model’s respective *piControl* experiment from the CMIP7 DECK. *X* denotes different present-day (Year 2021) anthropogenic perturbations to individual SLCFs. AER means that the model should, at least, have a time-evolving treatment of aerosols, either through time-dependent prescribed input fields or fully interactive aerosol treatments. CHEM means that the models are required to have interactive chemistry. Experiments marked with * follow the setup from AerChemMIP phase 1 (Collins et al., 2017) although *piClim-CH4* may be concentration- or emission-driven in AerChemMIP2.

Experiment ID	Minimum model capability	CH ₄	N ₂ O	Aerosol precursors	O ₃ precursors	CFC/HCFC	Tier	Synergy
<i>piClim-control*</i>	AGCM AER	1850	1850	1850	1850	1850	1	DECK, RFMIP, AerChemMIP, AFT
<i>piClim-AQ</i>	AGCM CHEM	1850	1850	2021	2021	1850	1	RFMIP, AerChemMIP, AFT
<i>piClim-aer*</i>	AGCM AER	1850	1850	2021	1850	1850	1	RFMIP, AerChemMIP, AFT
<i>piClim-BC*</i>	AGCM AER	1850	1850	2021 for BC, other 1850	1850	1850	1	RFMIP, AerChemMIP, AFT
<i>piClim-O3</i>	AGCM CHEM	1850 (2021 for chemistry)	1850	1850	2021	1850	1	RFMIP, AerChemMIP, AFT
<i>piClim-CH4*</i>	AGCM CHEM	2021	1850	1850	1850	1850	1	RFMIP, AerChemMIP, AFT
<i>piClim-N2O*</i>	AGCM CHEM	1850	2021	1850	1850	1850	1	RFMIP, AerChemMIP, AFT
<i>piClim-ODS*</i>	AGCM CHEM	1850	1850	1850	1850	2021	1	RFMIP, AerChemMIP, AFT
<i>piClim-NOx*</i>	AGCM CHEM	1850	1850	1850	2021 for NO _x , 1850 for non-NO _x	1850	1	RFMIP, AerChemMIP, AFT
<i>piClim-SO2*</i>	AGCM AER	1850	1850	2021 for SO ₂ , other 1850	1850	1850	1	RFMIP, AerChemMIP, AFT
<i>piClim-OC*</i>	AGCM AER	1850	1850	2021 for OC, other 1850	1850	1850	1	RFMIP, AerChemMIP, AFT
<i>piClim-NH3*</i>	AGCM AER	1850	1850	2021 for NH ₃ , other 1850	1850	1850	1	RFMIP, AerChemMIP, AFT

Table 2. As Table 1, but for new *pdClim-X* experiments for the assessment of a potential state dependence of ERF. *pdClim-X* experiments use present-day sea surface and sea ice conditions calculated from the last 30 years of a single realization of the model’s own coupled historical (*hist*) experiment.

Experiment ID	Minimum model capability	CH ₄	N ₂ O	Aerosol precursors	O ₃ precursors	CFC/HCFC	Tier
<i>pdClim-control</i>	AGCM AER	2021	2021	2021	2021	2021	3
<i>pdClim-AQ</i>	AGCM CHEM	2021	2021	1850	1850	2021	3
<i>pdClim-aer</i>	AGCM AER	2021	2021	1850	2021	2021	3
<i>pdClim-BC</i>	AGCM AER	2021	2021	1850 for BC, other 2021	2021	2021	3
<i>pdClim-O3</i>	AGCM CHEM	2021 (1850 for chemistry)	2021	2021	1850	2021	3
<i>pdClim-H2</i>	AGCM CHEM	2021 (1850 for H ₂ effects)	2021	2021	2021 (1850 for H ₂ effects)	2021	3

Table 3. As Table 1, but for new *pdClim-X* experiments that quantify the influence of carbon monoxide (CO) and non-methane volatile organic compounds (NMVOCs) on effective radiative forcing.

Experiment ID	Minimum model capability	CH ₄	N ₂ O	Aerosol precursors	O ₃ precursors	CFC/HCFC	Tier
<i>pdClim-CO</i>	AGCM CHEM	2021	2021	2021	1850 CO, 2021 others	2021	2
<i>pdClim-NMVOC</i>	AGCM CHEM	2021	2021	1850 NMVOCs, 2021 others	1850 NMVOCs, 2021 others	2021	2
<i>pdClim-C2H6</i>	AGCM CHEM	2021	2021	2021	1850 C2H6, 2021 others	2021	3
<i>pdClim-C3H8</i>	AGCM CHEM	2021	2021	2021	1850 C3H8, 2021 others	2021	3
<i>pdClim-C2H4</i>	AGCM CHEM	2021	2021	2021	1850 C2H4, 2021 others	2021	3
<i>pdClim-C3H6</i>	AGCM CHEM	2021	2021	2021	1850 C3H6, 2021 others	2021	3
<i>pdClim-C4H10</i>	AGCM CHEM	2021	2021	2021	1850 C4H10, 2021 others	2021	3
<i>pdClim-alcohol</i>	AGCM CHEM	2021	2021	2021	1850 alcohols, 2021 others	2021	3

Table 4. Experiments for quantifying the radiative effect of natural emission changes. These are *piClim-X* experiments like in Table 1, but here for doubled emission fluxes. The experiment *piClim-p4K* is identical to *piClim-control* except that a uniform temperature perturbation of 4 K is applied to sea surface temperatures in ice-free conditions. Experiments marked with * follow the setup from AerChemMIP phase 1 (Collins et al., 2017).

Experiment ID	Minimum model capability	Variable to be increased	Tier	Synergy
<i>piClim-2xdust*</i>	AGCM AER	2 x Mineral dust emissions	1	AerChemMIP, AFT
<i>piClim-2xss*</i>	AGCM AER	2 x Sea salt emissions	1	AerChemMIP, AFT
<i>piClim-2xfire*</i>	AGCM AER	2 x Fire emissions (NO _x , BC, OC, CO, VOCs, ...)	1	AerChemMIP, AFT
<i>piClim-2xBVOC*</i>	AGCM CHEM	2 x Biogenic VOC emissions	1	AerChemMIP, AFT
<i>piClim-2xWet</i>	AGCM CHEM	2 x Wetland emissions (CH ₄)	1	AFT
<i>piClim-2xPOApDMS</i>	AGCM AER	2 x primary marine organic compounds and 2 x dimethyl sulphate	1	AFT
<i>piClim-2xflash</i>	AGCM CHEM	2 x flash rate (NO _x , natural fire ignitions (if applicable))	1	AFT
<i>piClim-p4K</i>	AGCM AER	Sea-surface temperature + 4 K	1	AFT

Table 5. Experiments with a coupled ocean model for the historical period (1850–2021) and future scenarios with baselines *esm-scen7-h* and *esm-scen7-vl* from ScenarioMIP-CMIP7 (2022–2125). The experiments *piControl* and *hist* are part of CMIP7 DECK and are used here as reference experiments in AerChemMIP2. Ref and Hist are the component’s developments as in the baseline scenario and historical experiments. Experiments marked with * follow the setup from AerChemMIP phase 1 (Collins et al., 2017). Experiments marked with ^ address feedbacks via natural emissions that were not fully represented in past AOGCMs. Experiments marked with ° will allow an assessment of policy impacts on atmospheric composition and climate change.

Experiment ID	Minimum model capability	CH ₄	N ₂ O	Aerosol precursors	O ₃ precursors	CFC/HCFC	Tier	Minimum Ensemble Size	Synergy
<i>hist-piAQ*</i>	AOGCM CHEM	Hist	Hist	1850	1850	Hist	1	3	AFT, DAMIP
<i>hist-piAer*</i>	AOGCM AER	Hist	Hist	1850	Hist	Hist	1	3	AFT, DAMIP
<i>hist-Dust ^</i>	AOGCM AER	Hist	Hist	Hist with prescribed dust	Hist	Hist	2	3	AeroCom
<i>hist-piFire ^</i>	AOGCM CHEM	Hist	Hist	Hist but 1850 fire emissions	Hist but 1850 fire emissions	Hist	2	3	FireMIP
<i>hist-piCH4*</i>	AOGCM CHEM	1850	Hist	Hist	Hist	Hist	2	3	DAMIP
<i>hist-piO3</i>	AOGCM CHEM	Hist (1850 for chemistry)	Hist	Hist	1850	Hist	2	3	DAMIP
<i>esm-scen7-h-AQ °</i>	AOGCM CHEM	H transient	H transient	2021	2021	H transient	1	3	AFT
<i>esm-scen7-h-Aer °</i>	AOGCM AER	H transient	H transient	2021	H transient	H transient	1	3	AFT
<i>esm-scen7-vl-AQ °</i>	AOGCM CHEM	VL transient	VL transient	H transient	H transient	VL transient	1	3	AFT
<i>esm-scen7-vl-Aer °</i>	AOGCM AER	VL transient	VL transient	H transient	VL transient	VL transient	1	3	AFT
<i>esm-scen7-vl-Dust ^</i>	AOGCM AER	VL transient	VL transient	dust increase, other VL transient	VL transient	VL transient	2	3	AeroCom

Table 6. Historical experiments with prescribed transient changes in sea-surface conditions. These experiments follow the all-but-one approach of AerChemMIP, where the climate forcing of interest is held fixed at the pre-industrial level whereas all other forcings evolve as in a historical experiment. *piClim-histall* uses a prescribed annually repeating climatology of the pre-industrial sea-surface conditions (as used in *piClim-control*) but time-varying climate forcings.

Experiment ID	Minimum model capability	CH ₄	N ₂ O	Aerosol precursors	O ₃ precursors	CFC/HCFC	Tier	Synergy
<i>piClim-histall</i>	AGCM AER	Hist	Hist	Hist	Hist	Hist	2	RFMIP
<i>histSST</i>	AGCM AER	Hist	Hist	Hist	Hist	Hist	1	AerChemMIP
<i>histSST-piAQ</i>	AGCM CHEM	Hist	Hist	1850	1850	Hist	1	AerChemMIP, DAMIP
<i>histSST-piAer</i>	AGCM AER	Hist	Hist	1850	Hist	Hist	1	DAMIP
<i>histSST-Dust</i>	AGCM AER	Hist	Hist	Hist with prescribed dust	Hist	Hist	2	AeroCom
<i>histSST-piFire</i>	AGCM CHEM	Hist	Hist	Hist but 1850 fire emissions	Hist but 1850 fire emissions	Hist	2	FireMIP
<i>histSST-piCH4</i>	AGCM CHEM	1850	Hist	Hist	Hist	Hist	2	AerChemMIP
<i>histSST-piO3</i>	AGCM CHEM	Hist (1850 for chemistry)	Hist	Hist	1850	Hist	2	AerChemMIP
<i>histSST-piNOx</i>	AGCM CHEM	Hist	1850	Hist	Hist	Hist	3	AerChemMIP

Table 7. Experiments for future development with prescribed sea-surface conditions. We cover the period of the future scenarios of CMIP7 (2022–2125). Ref means developments identical to the reference scenario H and VL as in ScenarioMIP-CMIP7. Ref refers to conditions as in the corresponding ScenarioMIP-CMIP7 experiments, e.g., *esm-scen7-vl-SST* with a relatively cleaner atmosphere compared to *esm-scen7-h-SST*.

Experiment ID	Minimum model capability	CH ₄	N ₂ O	Aerosol precursors	O ₃ precursors	CFC/HCFC	Tier
<i>esm-scen7-h-SST</i>	AGCM AER	H transient	H transient	H transient	H transient	H transient	1
<i>esm-scen7-h-SST-AQ</i>	AGCM CHEM	H transient	H transient	2021	2021	H transient	1
<i>esm-scen7-h-SST-Aer</i>	AGCM AER	H transient	H transient	2021	H transient	H transient	1
<i>esm-scen7-vl-SST</i>	AGCM AER	VL transient	VL transient	VL transient	VL transient	VL transient	1
<i>esm-scen7-vl-SST-AQ</i>	AGCM CHEM	VL transient	VL transient	H transient	H transient	VL transient	1
<i>esm-scen7-vl-SST-Aer</i>	AGCM AER	VL transient	VL transient	H transient	VL transient	VL transient	1
<i>esm-scen7-vl-SST-Dust</i>	AGCM AER	VL transient	VL transient	dust increase, other VL transient	VL transient	VL transient	2
<i>esm-scen7-vl-SST-CH4</i>	AGCM CHEM	H transient	VL transient	VL transient	VL transient	VL transient	2
<i>esm-scen7-vl-SST-O3</i>	AGCM CHEM	VL transient (H transient for chemistry)	VL transient	VL transient	H transient	VL transient	3

Table 8. Experiments with prescribed sea-surface conditions examining afforestation and reforestation (A/R). Int(eractive) with PD LU means BVOC, fire and dust emissions can evolve with climate (SST change) but land use is fixed at a PD (2021) climatology so BVOC and fire emissions are not influenced by LU changes. Int(eractive) with VL LU means BVOC, fire and dust emissions can evolve with climate and land use changes. Simulations A1, A3, B1 and B3 require models which can calculate BVOC, fire and dust emissions interactively based on climate and land cover (emissions of aerosol and O₃ precursor emissions generated interactively in A3/B3 are then prescribed in A2/B2). If a modelling centre cannot do this but would like to participate, we recommend prescribing emissions diagnosed by a model which has such interactive capability. The authors actively encourage modelling centres in this position to contact them so connections can be made with groups who can generate interactive emissions. Index relates each simulation to those in Fig 9. Int = Interactive, B = BVOC, F = Biomass burning, D = Dust. LU should be prescribed as a monthly climatology from 2021 for pdLU or monthly transient timeseries following VL (VL transient). Further notation as in table 7.

Index*	Experiment ID	Min model capability	CO ₂ , CH ₄ , N ₂ O, CFC/HCFC, SSTs	Aerosol & O ₃ precursors	LU	Tier
A1	esm-scen7-vl-SST-pdLU-BFD	AGCM AER	VL transient	VL transient (Anthro), Int with PD LU (BFD)	2021 climatology	2
A2	esm-scen7-vl-SST-pdLU**	AGCM	VL transient	VL transient (Anthro), BFD from A3	2021 climatology	2
A3	esm-scen7-vl-SST***	AGCM	VL transient	VL transient (Anthro), Int with VL LU (BFD)	VL transient	1
B1	esm-scen7-h-SST-pdLU-BFD	AGCM AER	H transient	H transient (Anthro), Int with PD LU (BFD)	2021 climatology	3
B2	esm-scen7-h-SST-pdLU**	AGCM	H transient	H transient (Anthro), BFD from B3	2021 climatology	3
B3	esm-scen7-h-SST-vl-LU**	AGCM	H transient	H transient (Anthro), Int with VL LU (BFD)	VL transient	3

* A1 vs. A3 / B1 vs. B3 gives total RF; A1 vs. A2 / B1 vs. B2 gives RF from chemistry and aerosols; A2 vs. A3 / B2 vs. B3 gives surface albedo RF

**Min model capability is AGCM since A2 vs. A3 / B2 vs. B3 gives surface albedo RF

***Identical to esm-scen7-vl-SST in Table 7