

Anonymous Referee #2

We would like to thank the reviewer for the dedicated time he/she spent on reading the manuscript and providing valuable feedback to us. A point-by-point response is given below.

This study explores the dependency of emissions-normalized radiative forcing and regional temperature response to the location and magnitude of black carbon emissions, using numerous simulations with the NorESM model. Given the struggles of past efforts to reliably quantify climate impacts from small emissions of short-lived species, this study provides very useful information that can be applied to inform future efforts, such as those of the AMAP expert group on short-lived climate forcers. I recommend publication of this article in ACP after the following minor issue are addressed.

Minor issues:

Abstract: I suggest including a statement on your findings of the ARTP approach, or namely that this approach (when used with DRF) rather accurately reproduces the fully-coupled temperature response of NorESM.

Thank you for pointing this out. We have added such a statement to the abstract.

It is a bit surprising to me that the ITCZ can shift in fixed-SST simulations. If you are aware of other studies that have shown this behavior (especially any of the studies cited on lines 104-105), I suggest noting this.

Yes, this is part of the fast response resulting from atmospheric heating and land cooling before sea-surface temperature (SST) has time to respond and this causes a northward ITCZ shift. This has been shown in Voigt et al. (2017) for seven atmosphere models with large aerosol forcing. We have added this reference.

Voigt, A., Pincus, R., Stevens, B., Bony, S., Boucher, O., Bellouin, N., Lewinschal, A., Medeiros, B., Wang, Z., and Zhang, H. (2017), Fast and slow shifts of the zonal-mean intertropical convergence zone in response to an idealized anthropogenic aerosol, J. Adv. Model. Earth Syst., 9, 870–892, doi:10.1002/2016MS000902.

It is mentioned that the BC-on-snow effect is included in NorESM, but is it included in the estimates of DRF? Please clarify and comment on any associated implications.

We have clarified that the DRF only include the atmospheric forcing, and not BC-on-snow. The BC-on-snow effect is included when calculating the surface temperature response/albedo changes etc.

Section 2.2: More details are needed on the experimental set-up. Specifically: Were annually-repeating or annually-changing boundary conditions (GHGs, SSTs, etc) used in the runs, and from which years do the conditions represent? How long were the ERF fixed-SST runs and what time frame was used for the analysis? (line 93: "after the atmosphere is allowed to respond to the forcing" needs precision). The text indicates that the final 80 years of 100-year fully-coupled runs were analyzed. Is 20 years sufficient for equilibration to occur? Perhaps for BC it is, but a brief discussion on this should be added.

Please describe the baseline emissions that are used, e.g., which inventory are they from and what are the global BC emissions? I also suggest adding the annual emissions for each experiment/region to Table 1.

The have added the following sentences in Methods to clarify:

'We have calculated the ERF as the annual mean change in the net radiation at TOA after the atmosphere is allowed to respond to the forcing (i.e. rapid adjustments in temperature, relative humidity and clouds), over the last 10 years of total 15 years simulations '

'The baseline is run with year 2000 annually repeating emissions of aerosols and precursors, land use conditions and greenhouse gas concentrations. The aerosol emissions (and their precursors) are the historical emissions of CMIP5 described in Lamarque et al. (2010). The global mean BC emissions for the baseline simulation are 7.7 Tg/yr.'

'20 year spin-up was sufficient for equilibrium to occur for BC perturbations (this is also shown in Sand et al, (2015))'

We have also added a column with global annual BC emissions in Table 1.

line 189: Sentence beginning "For South Asia..." - Does this South Asia response occur only with emissions from South Asia, or is it also seen with emissions from elsewhere that transport to South Asia?

Yes, this could easily be misunderstood. We have changed this to 'For BC emitted in South Asia'.

Technical issues:

line 14: "a rate of" -> "a factor of"

This has been changed.

line 17: "BC emitted in South Asia shows a different geographical pattern by changing..." - This sentence needs clarification. Geographical pattern of which variable(s)?

We have added: 'geographical pattern in surface temperatures ..'

line 75: "according different" -> "according to different"

line *This has been changed.*

84: "perturbed by 10 and 20" -> "perturbed by factors of 10 and 20"

This has been changed.

line 128: "global BC burden": This is the <emissions-normalized> global BC burden, correct?

No, this is the total global BC burden change, as stated in the manuscript, but we have changed the text to be clearer. When we double the emissions, we more than double the burden (117% increase (typo 217 % has been changed, thanks for noting this!). We have also changed the description further down: 'For the simulations with the high emission rate, the relative concentration increase normalized to global emissions is largest at high altitudes above 200 hPa (25-55% increase shown in zonal mean plots of concentrations normalized to emissions per altitude in Fig S3-S6).

line 132: Please clarify the meaning of "emission-driven vs. concentration-driven BC"

We agree that this was confusing; we have changed this to: 'BC perturbed by increasing the BC emission rate compared to increasing the BC concentrations.'

line 144-145: Sentence beginning "For the Arctic..." is unclear. Please rework this.

We have change this to: 'The Arctic region shows a higher climate sensitivity for the lower emission rate compared to the higher emission rate (the difference is not significant for North American emissions).'

Review of “Surface temperature response to regional black carbon emissions: Do location and magnitude matter?” by Sand et al., submitted to Atmospheric Chemistry and Physics.

We would like to thank the reviewer for the dedicated time he/she spent on reading the manuscript and providing valuable feedback to us. A point-by-point response is given below.

This paper discusses the impacts of regional emissions of black carbon on global and regional temperatures. Scenarios are examined with different levels of emissions. One conclusion is that a higher emission rate results in lower temperature change per unit emissions, except in the Arctic. Impacts of emissions from North America and Europe are found to be similar to each other but different from emissions from East Asia. Overall, the paper contributes to our understanding of the impacts of black carbon. However, the paper can be improved if the model is described more completely and if previous understanding of black carbon climate effects were discussed more completely. Also, the authors do not discuss the physical processes they are not including that could affect results. Finally, there is no evaluation of the model. Below are more specific comments.

With respect to the model, there should be some validation of baseline cloud fields and aerosol optical depth and aerosol absorption optical depth against satellite and/or in situ data. Some validation is required in virtually all papers.

A comprehensive validation of the same version of the model has already been performed in Bentsen et al (2013) and Kirkevåg et al (2013) and this is the main reason why we chose this model version. We have added the following: ‘A comprehensive assessment of the mean model state is available in Bentsen et al. (2013). Here it is shown that the global mean energy fluxes and associated cloud forcing are close to or within the observations range, but that the cloud cover is underestimated. The aerosols fields are evaluated in Kirkevåg et al. (2013). Here it is shown that global BC concentrations is underestimated by -18 % compared to observations. The global anthropogenic aerosol DRF is -0.08 Wm^{-2} and the indirect radiative forcing is -1.2 Wm^{-2} . ‘

Abstract. “BC emissions are increase by a rate of 10 and 20. . .” 10 and 20 what? A factor of 10 and 20? 10 and 20%? Please clarify.

We have changed ‘rate’ to ‘factor’.

Introduction. “There has been a growing interest for reducing black carbon emissions to slow global warming emissions and improve air quality.” This was proposed at least as far back as 2002 in Jacobson (2002), which states in the abstract, “Reducing BC + OM will not only slow global warming but also improve human health.” Please include this information.

We have added this reference.

Introduction. “BC absorbs solar radiation and therefore rapidly influence heating rates, humidity and clouds in the atmosphere.” That feedback, along with 11 other feedbacks of black carbon to climate, is discussed Jacobson (2002), Sections 3.1-3.12. Please discuss.

We agree that BC climate effects were not described in detail, and we have expanded the Introduction to include more references and examples of responses and feedbacks.

NorESM. Please state explicitly how, in your model, “aerosols can act as CCN based on their size and composition.” Is it an empirical parameterization? Do cloud drops of different size and composition grow physically in time, or do you assume an equilibrium cloud drop size distribution.

There is not an empirical parameterization of cloud droplets in NorESM. CDNC and LWC are prognostically calculated, with CCN activation estimated based on supersaturations calculated from Köhler theory (Abdul-Razzak and Ghan, 2000) using a sub-grid scale vertical velocity. The rate of activation depends on the size distribution and the hygroscopicity. The cloud droplet size distribution is affected by microphysical processes and varies between the time steps. We agree that the model description was too brief. We have added the information above to Methods.

Please state explicitly in the text whether you include Cloud Absorption Effects I and II of black carbon (Jacobson, 2012). CAE I is cloud burnoff due to absorption by black carbon inclusions within individual cloud drops and CAE II is cloud burnoff due to the hydration of BC-containing aerosol particles between cloud drops at the ~100% relative humidity of the cloud, and the resulting enhanced absorption and heating of the cloud due to optical focusing. If you do not include it, please discuss briefly the potential impacts of not including this treatment.

The CAE I effect is, as in most global climate models, not included in NorESM. We have included in the Methods: ‘Any cloud ‘burn-off’-effects due to absorption by BC within individual cloud droplets are not included’.

Studies show that the lensing effect might enhance BC absorption, but the magnitude of this enhancement remains controversial (Lack et al., 2009; Cappa et al., 2012) due to the various assumptions (coating thickness/morphology/mixing geometry) (e.g. Adachi et al., 2010, Scarnato et al., 2013). Coating of BC will also reduce the BC atmospheric lifetime (Boucher et al. 2016). It is therefore difficult to discuss this in a profound way, unfortunately.

*Adachi, K., Chung, S. H., and Buseck, P. R.: Shapes of soot aerosol particles and implications for their effects on climate, *J. Geophys. Res.-Atmos.*, 115, D15206, <https://doi.org/10.1029/2009JD012868>, 2010*

*Boucher, O., Balkanski, Y., Hodnebrog, Ø., Myhre, C. L., Myhre, G., Quaas, J., Samset, B. H., Schutgens, N., Stier, P., and Wang, R.: Jury is still out on the radiative forcing by black carbon, *Proceedings of the National Academy of Sciences*, 113, E5092-E5093, [10.1073/pnas.1607005113](https://doi.org/10.1073/pnas.1607005113), 2016.*

*Scarnato, B. V., Vahidinia, S., Richard, D. T., and Kirchstetter, T. W.: Effects of internal mixing and aggregate morphology on optical properties of black carbon using a discrete dipole approximation model, *Atmos. Chem. Phys.*, 13, 5089–5101, <https://doi.org/10.5194/acp-13-5089-2013>, 2013.*

*Lack, D. A., Cappa, C. D., Cross, E. S., Massoli, P., Ahern, A. T., Davidovits, P., and Onasch, T. B.: Absorption Enhancement of Coated Absorbing Aerosols: Validation of the Photo-Acoustic Technique for Measuring the Enhancement, *Aerosol Sci. Tech.*, 43, 1006–1012, <https://doi.org/10.1080/02786820903117932>, 2009.*

*Cappa, C. D., Onasch, T. B., Massoli, P., Worsnop, D. R., Bates, T. S., Cross, E. S., Davidovits, P., Hakala, J., Hayden, K. L., Jobson, B. T., Kolesar, K. R., Lack, D. A., Lerner, B. M., Li, S.-M., Mellon, D., Nuaaman, I., Olfert, J. S., Petäjä, T., Quinn, P. K., Song, C., Subramanian, R., Williams, E. J., and Zaveri, R. A.: Radiative Absorption Enhancements Due to the Mixing State of Atmospheric Black Carbon, *Science*, 337, 1078–1081, <https://doi.org/10.1126/science.1223447>, 2012.*

Please state explicitly how “The BC on snow effect is included in the model.” Do you model the size-resolved deposition of BC (through wet and dry deposition) to the surface. How is radiative transfer solved through the snow if it is? If it is not, please state so explicitly.

We have changed this to: Deposited aerosols on snow-cover and bare sea-ice are calculated prognostically and include hydrophobic and hydrophilic black carbon and dust. While the effects of deposition of these light-absorbing aerosols are taken into account, we have not calculated the radiative transfer of BC in snow explicitly.

What is the mixing state of BC aerosol components in the model? Externally mixed, well-internally mixed, or something in between?

We have added the following: 'BC primary particles are assumed externally-mixed. BC is internally-mixed with organic matter and SO₄, the latter through either condensation or coagulation. The mass of internally mixed coating species is used in the calculations of the optical properties and cloud droplet number concentration, but it does not alter wet deposition rates'

Abstract. "For these regional BC emissions perturbations, we find that the effective radiative forcing is not a good measure of the climate response." Although it is not clear if your model includes this, one explanation may be that "when absorbing aerosols exist in clouds, instantaneous direct radiative forcing (DRF) and surface temperature change are anticorrelated because when absorbing aerosol burns off a cloud, the aerosol DRF decreases due to a decrease in optical focusing, yet surface temperature escalates rapidly due to the pouring in of sunlight to the surface." (Jacobson, 2014). Please discuss.

Again, as this effect is not included in NorESM unfortunately, is it difficult to discuss this.

Figure 5. Change figure captions to "Delta TOA" rather than just "TOA" Are these "net downward minus upward" irradiances with versus without black carbon emissions? If so, please clarify.

We have changed the caption to: 'Change in delta TOA net SW and LW radiative fluxes (sum of downward and upward) for the 10×BC in East Asia perturbation minus the baseline.'

The text mentions that the Indian Monsoon was displaced. How was it displaced? In what direction and what was the magnitude in wind speed change? Were Monsoon wind speeds decreased due to black carbon? Were globally or regionally averaged wind speeds decreased as might be expected (Jacobson and Kaufman, 2006).

We have expanded this section, and added more references:

'This cooling is linked to a strong decrease in downwelling SW radiation over India caused by the absorption and scattering by BC aerosols, less upward motion and an increase in low level clouds. Northern hemisphere BC emissions cause a northward displacement of the ITCZ that is most pronounced for emissions in South Asia and a corresponding shift in the Indian monsoon. The tropospheric jet is moving northward to adjust for a northward shift in the meridional temperature gradient (not shown). This type of displacement caused by a hemispherical asymmetric heating is a robust feature found in several climate model studies (Chung and Seinfeld, 2005; Jones et al., 2007; Meehl et al., 2008; Allen et al., 2012, Voigt et al. 2017; Kovilakam and Mahajan, 2015)'

References

Jacobson, M. Z., Control of fossil-fuel particulate black carbon plus organic matter, possibly the most effective method of slowing global warming, J. Geophys. Res., 107 (D19), 4410, doi:10.1029/2001JD001376, 2002

Jacobson, M.Z., and Y.J. Kaufmann, Wind reduction by aerosol particles, Geophys. Res. Lett., 33, L24814, doi:10.1029/2006GL027838, 2006.

Jacobson, M.Z., Investigating cloud absorption effects: Global absorption properties of black carbon, tar balls, and soil dust in clouds and aerosols, *J. Geophys. Res.*, 117, D06205, doi:10.1029/2011JD017218, 2012.

Jacobson, M.Z., Effects of biomass burning on climate, accounting for heat and moisture fluxes, black and brown carbon, and cloud absorption effects, *J. Geophys. Res.*, 119, 8980-9002, doi:10.1002/2014JD021861, 2014.

Surface temperature response to regional Black Carbon emissions: Do location and magnitude matter?

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10 **Abstract.** Aerosol radiative forcing can influence climate both locally and far outside the emission region. Here we investigate
Black Carbon (BC) aerosols emitted in four major emissions areas and evaluate the importance of emission location and
magnitude, as well as the concept of the absolute regional temperature-change potentials ([ARTP](#)). We perform simulations
with a climate model (NorESM) with a fully coupled ocean and with fixed sea surface temperatures. BC emissions are
increased by a [factor](#) of 10 and 20 in South Asia, North America and Europe, respectively, and by 5 and 10 in East Asia
15 (due to higher emissions there). We find strikingly similar regional surface temperature responses and geographical patterns
per unit BC emission in Europe and North America, but somewhat lower temperature sensitivities for East Asian emissions.
BC emitted in South Asia shows a different geographical pattern [in surface temperatures](#), by changing the Indian monsoon and
cooling the surface. [We find that the ARTP approach rather accurately reproduces the fully-coupled temperature response of
NorESM.](#) Choosing the highest emission rate results in lower surface temperature change per emission unit compared to the
20 lowest rate, but the difference is generally not statistically significant except for the Arctic. An advantage of high-perturbation
simulations is the clearer emergence of regional signals. Our results show that the linearity of normalized temperature effects
of BC is fairly well preserved despite the relatively large perturbations, but that regional temperature coefficients calculated
from high perturbations may be a conservative estimate. Regardless of emission region, BC causes a northward shift of the
ITCZ, and this shift is apparent both with fully coupled ocean and with fixed sea surface temperatures. For these regional BC
25 emissions perturbations, we find that the effective radiative forcing is not a good measure of the climate response.

1 Introduction

There has been a growing interest for reducing black carbon (BC) emissions to slow global warming and improve air quality
([Jacobson, 2002](#); [Shindell et al., 2012](#); [Quinn et al., 2015](#); [Sims et al., 2015](#)). However, estimating the total climate impact
from BC is complicated because BC absorbs solar radiation and therefore rapidly influence heating rates, humidity and clouds
30 in the atmosphere ([Bond et al., 2013](#); [Hansen et al., 1997](#); [Cook and Highwood, 2004](#)). [These effects are often called ‘rapid](#)

adjustments' and are distinct from the direct radiative forcing (DRF), i.e. scattering and absorption of sunlight by BC, and the indirect radiative forcing, i.e. changing the microphysical properties of clouds ((Twomey, 1977; Haywood and Boucher, 2000). The surface warming by BC is highly sensitive to these rapid adjustments ~~in the atmosphere~~ and is dependent on the altitude of the BC layer and the co-location of clouds (Ban-Weiss et al., 2012; Koch and Del Genio, 2010; Johnson et al., 2004; (Jacobson, 2002)). For instance in Johnson et al. (2004) using large-eddy simulations, it was found that BC within the boundary layer heated the air, decreased liquid water path (LWP) and thinned the marine stratocumulus clouds, while BC above the boundary layer increased LWP leading to a more shallow and moist boundary layer. Ramanathan et al. (2001) showed that BC can also reduce the surface solar radiation and increase static stability lowering the surface moisture fluxes. Using prescribed BC in a general circulation model coupled to a mixed-layer ocean Ban-Weiss et al. (2012) showed that BC near the surface caused surface warming and increased precipitation, while BC near the tropopause and in the stratosphere, on the other hand, decreased surface temperatures and decreased precipitation. These rapid adjustments are shown to be negative and offset the direct effect of BC (Stjern et al., 2017; Smith et al., 2018; Samset and Myhre, 2015). BC forcing from outside the region can ~~also~~ contribute to the surface temperature response via transport of heat (Shindell, 2007; Sand et al., 2013; Menon et al., 2002). Coupled Earth System Models (ESMs) now include most of the relevant processes to study the complex feedbacks mentioned above and could in principle be used to analyse and compare the effects of different regional/sectoral BC mitigation options of interest to policy makers. However, the change in emissions for these mitigation options are often too small (e.g. agricultural waste in Europe) to get a statistically significant signal in an ESM without running it for thousands of years, which makes it nearly impossible with state-of-the-art supercomputer clusters. As an alternative, climate metrics provide an easy way to compare emission perturbations, forcing and response. The absolute regional temperature-change potentials (ARTPs) derived by Shindell and Faluvegi (2009) provide a relationship between forcing in one region and a surface temperature response in another region. The relation between forcing and surface temperature response were calculated for four different latitude bands (the Arctic, mid-latitudes, tropics and southern hemisphere (SH)), using a fully coupled climate model (giss modelE). Shindell and Faluvegi (2009) highlighted the importance of remote forcing on certain regions and found that the Arctic and mid-latitudes were especially sensitive to the location of the forcing. For instance, the radiative forcing of BC at mid latitudes strongly influenced the surface temperature in the Arctic. In a follow-up-study Shindell et al. (2010) evaluated the method using transient historical simulations in four climate models and found good agreement. The ARTP coefficients have been used in many studies to estimate the surface temperature response to different forcing perturbations derived from chemistry transport models (Sand et al., 2015a; Collins et al., 2013). For instance, Sand et al. (2015a) estimated the Arctic surface temperature response to emissions of BC, SO₂ and OC from a wide range of sectors and regions by calculating the direct radiative forcing (DRF) in four CTMs and applying the ARTP coefficients.

The ARTP method is quick and efficient, however, there are important simplifications underlying the calculations. As these coefficients cannot easily be validated by observations and have not been calculated in a multi-model experiment, there are considerable uncertainties in these coefficients. Also, the emissions rates are calculated for broad latitude bands and any longitudinal variations in the response is not represented.

65 Lewinschal et al. (2019) evaluated the surface temperature change in response to SO₂ emission perturbations in four major
emission regions using a different climate model (NorESM) and compared it to the temperature response estimated by using
the ARTPs. The ARTP method predicted similar latitudinal temperature response with those estimated by NorESM. Here we
perform simulations of the response to regional BC-emissions with the fully coupled climate model (NorESM) following the
70 ARTP concept. We calculate emission-to-temperature responses by estimating the regional surface temperature response to
BC emissions from four major emission regions; North America, Europe, South Asia, and East Asia. To get a signal of a small
emission perturbation in a coupled (atmosphere-ocean) climate model, it is often necessary to scale up the emissions. We
evaluate this assumed linearity by perturbing the BC emissions with two different emission rates.

2 Methods

75 2.1 NorESM

We have used the Norwegian Earth System model (NorESM1) (Bentsen et al., 2013;Iversen et al., 2013), which is largely
based on the CCSM4.0 framework (Gent et al., 2011) with special features for aerosols and their interaction with radiation and
warm cloud microphysics. (Kirkevåg et al., 2013;Seland et al., 2008). The model uses the finite volume dynamical core for
transport, with a horizontal resolution of 1.9° latitude by 2.5° longitude and 26 levels in the vertical. Unlike CCSM4.0,
80 NorESM1 is run with an elaborated version of the Miami Isopycnic Community Ocean Model (MICOM).

The aerosol life cycle scheme calculates mass concentrations of SO₂, BC, organic matter, sea salt, and mineral dust in up to
four size modes (nucleation, Aitken, accumulation, and coarse mode). BC is emitted in the nucleation, Aitken and accumulation
mode and in the internally mixed Aitken mode with organic matter. BC primary particles are assumed externally-mixed. BC
is internally-mixed with organic matter and SO₄, the latter through either condensation or coagulation. The mass of internally
85 mixed coating species is used in the calculations of the optical properties and cloud droplet number concentration, but it does
not alter wet deposition rates. Water is mixed into the aerosols based on the hygroscopicity and ambient relative humidity. The
calculated gas-phase components are DMS and SO₂. The aerosol concentrations are tagged according to different processes.
The processes include gas phase and aqueous phase chemical production, gas to particle nucleation, condensation on pre-
existing aerosol surfaces, and coagulation of smaller particles onto pre-existing Aitken, accumulation and coarse mode
90 particles. The process-tagged aerosol mass concentrations and relative humidity are given as input to look-up tables that
estimate the optical and physical properties of the aerosols. The aerosols can also act as cloud condensation nuclei (CCN)
based on their size and composition. BC that is mixed with other components and has become hydroscopic can contribute to
the number of CCN. Cloud droplet number concentration (CDNC) and liquid water path are prognostically calculated. CCN
activation is estimated based on supersaturations calculated from Köhler theory (Abdul-Razzak and Ghan, 2000) using a sub-
95 grid scale vertical velocity. The rate of activation depends on the size distribution and the hygroscopicity of the particles. Any
cloud ‘burn-off’-effects due to absorption by BC within individual cloud droplets are not included. Deposited aerosols on

100 snow-cover and bare sea-ice are calculated prognostically and include hydrophobic and hydrophilic black carbon and dust. While the effects of deposition of these light-absorbing aerosols are taken into account, we have not calculated the radiative transfer of BC in snow explicitly. The BC-on-snow effect is included in the model. A comprehensive assessment of the mean model state is available in Bentsen et al. (2013). Here it is shown that the global mean energy fluxes and associated cloud forcing are close to or within the observations range, but that the cloud cover is underestimated. The aerosols fields are evaluated in Kirkevåg et al. (2013). Here it is shown that global BC concentrations is underestimated by -18 % compared to observations. The global anthropogenic aerosol DRF is -0.08 Wm^{-2} and the indirect radiative forcing is -1.2 Wm^{-2} .

2.2 Experimental Set-up

105 Coupled atmosphere/ocean simulations with NorESM have been performed for a set of emission perturbations. BC emissions have been increased in four areas: South Asia, East Asia, North America and Europe. In these regions the emissions have been perturbed by factors of 10 and 20, except East Asia that has been perturbed by 5 and 10 (due to the relative higher emissions in this region). The magnitude of these rates was chosen to get a large enough signal, and at the same time keep global DRF below $\sim 1 \text{ W/m}^2$. Figure S1 shows the increase in total global emissions compared to a baseline simulation for each region. The
110 baseline is run with year 2000 annually repeating constant year 2000 emissions of aerosols and precursors, land use conditions and greenhouse gas concentrations. The aerosol emissions (and their precursors) are the historical emissions of CMIP5 described in Lamarque et al. (2010). The global mean BC emissions for the baseline simulation are 7.7 Tg/yr . Table 1 provides an overview of all the simulations. For each perturbation coupled simulations for 100 years with 3 ensemble members have been run, i.e. a total of 300 years of simulation per emission perturbation. Furthermore, we calculate the DRF as the
115 instantaneous change in energy flux at the top-of-the-atmosphere with a double-call in the radiation routine. The calculations of the DRF does not include the BC-on-snow-forcing. In separate simulations with prescribed sea surface temperatures and sea-ice cover we calculate the effective radiative forcing (ERF) which include rapid tropospheric adjustments, often related to humidity and clouds that may exhibit fast adjustments to the radiative forcing and any changes in CCN. We have calculated the ERF as the annual mean change in the net radiation at TOA after the atmosphere is allowed to respond to the forcing (i.e.
120 rapid adjustments in temperature, relative humidity and clouds), over the last 10 years of total 15 years simulations. The surface temperature response per unit of DRF, ERF and per emission change is then compared for the four emission areas. All numbers reported are annual means from the last 80 years of the simulations (i.e. 80×3 years for each perturbation experiment). -20-
year spin-up was sufficient for equilibrium to occur for BC perturbations (this is also shown in Sand et al., 2015b)

3 Results

125 Figure 1 shows the surface temperature change for the perturbed emission runs compared to baseline. The broad patterns are similar to the well-known warming pattern due to CO_2 forcing, with larger warming over land and in the northern areas. The geographical pattern of warming is comparable for BC emitted in North America, Europe and East Asia with only small

longitudinal differences between those perturbations. For South Asia, however, there is a significant local cooling effect over India. This cooling is linked to a strong decrease in downwelling SW radiation over India caused by the absorption and scattering by BC aerosols, less upward motion and an increase in low level clouds. Northern hemisphere BC emissions cause a northward displacement of the ITCZ that is most pronounced for emissions in South Asia and a corresponding shift in the Indian monsoon. The tropospheric jet is moving northward to adjust for a northward shift in the meridional temperature gradient (not shown). This type of displacement caused by a hemispherically asymmetric heating is a robust feature found in several climate model studies (Chung and Seinfeld, 2005; Jones et al., 2007; Meehl et al., 2008; Allen et al., 2012; Voigt et al., 2017; Kovilakam and Mahajan, 2015).

Figure 2 shows the regional surface temperature change in four latitude bands (The Arctic (60°N-90°N, mid latitude (28°N-60°N), Tropics (28°S-28°N) and SH (90°S-28°S)) normalized by the enhancement in the BC emissions. Table 2 provides the corresponding numbers. For all cases the Arctic region shows the highest sensitivities, followed by the mid latitudes. The temperature sensitivities in the tropics and SH are lower compared to the mid latitudes and the Arctic, but these two regions are also much larger regions compared to the mid latitudes and the Arctic. These zonal mean temperature sensitivities are similar regardless of emission location, especially for BC emitted in North America and Europe. The temperature sensitivities are slightly lower in the Arctic and mid-latitudes for BC emitted in Asia, in particular South Asia. Temperatures in the Arctic increase by 0.04-0.1 K per Tg/yr BC emitted. In the mid latitudes the temperature increases by 0.04-0.08 K per Tg/yr BC emitted.

The two twin-bars in Fig. 2 given for each emission region represent the linearity test. The lighter coloured bars are the perturbations with the lowest emission rates, while the darker colour bars are the runs with the highest rates. In general, the sensitivity decreases with increasing emissions, most pronounced for the response in the Arctic (compare e.g. 20×EU vs. 10×EU, i.e. the right and left columns in Fig. 2 or numbers in Table 2). The decrease in Arctic sensitivities for the highest emissions are significant for BC emitted in Europe and Asia, but not North America ($p < 0.05$). For the mid latitude sensitivities, the changes are small, and only European emissions show significant changes (again, with weaker sensitivities for 20×emissions compared to 10×emissions).

To understand the regional differences and the non-linearities we first investigate the relation between emissions and concentrations. Figure 3 shows the change in regional BC burden normalized to global BC emissions. As expected, the burden changes per global emissions are largest in the latitude band where the emissions are located. The regional burden change normalized to emissions is slightly higher for the highest emission rates. These changes are statistically significant using a 0.05 level of significance (and also using a 0.005 level), except for the changes in the Tropics and SH for North American emissions. The same results are also obtained when we look at total global burden change (not shown). E.g. in Europe, emitting 20× emissions results in a 1217 % increase in global BC burden compared to 10×emissions. The rapid adjustments of BC cause a local warming of the atmosphere and an increased local convection pushing the BC layers higher up in the atmosphere where the BC is less likely to be scavenged. For the simulations with the high emission rate, the relative concentration increase normalized to global emissions (up to 270% increase) is largest at high altitudes above 200 hPa (25-55% increase shown in

zonal mean plots of concentrations normalized to global emissions per altitude ~~are shown~~ in Fig S3-S6). This effect has also been shown in Sand et al. (2015b) for BC perturbed by increasing the BC emission rate compared to increasing the BC concentrations. emission driven vs. concentration driven BC. As in this study, increasing the emission rate ~~resulteds~~ in a relative increase in BC at higher altitudes compared to lower altitudes and a longer BC residence time. This effect may explain why the temperature increase per emissions is largest for the small emission rate. Although BC perturbations at higher altitudes cause a larger DRF per unit burden change (Samset and Myhre, 2011), this is out weighted by a smaller temperature response per unit forcing. As BC is transported higher up in the atmosphere the surface temperature response decrease, as shown in Ban-Weiss et al. (2012). In the Arctic the effect of high-altitude BC is particularly strong, even causing a cooling at the surface for positive regional DRF, because of the strong vertical stability in this area (Flanner, 2013; Sand et al., 2013). A similar weaker temperature efficiency with stronger emission perturbations have also been found in Yang et al. 2019, but for much higher emission perturbations.

Figure 4 a) shows the climate sensitivity in terms of regional surface air temperature change per global DRF, i.e. a similar concept as the ARTPs (but for global forcing). In general, the temperature sensitivities for BC emitted in Europe and North America are similar, and 50% lower for BC emitted in East Asia and South Asia. In the Arctic the temperature increases 0.8-1.7 K per W/m^2 , while in the mid-latitudes the temperature increases by 1 K per W/m^2 . ~~The~~For the Arctic region shows, there are generally a higher climate sensitivity for the lower ~~response~~-emission rate compared to the higher emission rate (although the difference is not significant for North American emissions). The ERF is defined as the net SW minus the net LW for fixed SSTs perturbations. The temperature sensitivities when using ERF instead for DRF (Fig 4b), do not show the same consistent pattern. The rapid adjustments of BC are strongly negative in our model, and partly offset the DRF; resulting in lower ERF ($-0.0004 - 0.2 W/m^2$) compared to DRF ($0.3 - 1.3 W/m^2$). The small value of ERF makes the sensitivities (K/Wm^{-2}) large and uncertain (maps of ERF are shown in S02). For 20×South Asian emissions, the (global mean) ERF is close to 0 (which makes the climate sensitivity defined here undefinedly large). Even though ERF may not be a good predictor of the surface temperature response, only using DRF will hide some of the uncertainties related to atmospheric absorption. The rapid adjustments of BC cause a shift in the large-scale circulation patterns in the atmosphere. This shift is apparent both when using a coupled ocean and fixed SSTs. As an example, the net TOA SW and LW fluxes are plotted for 10×East Asia for both ocean setups in Fig 5. The ERF is positive close to (all) the emission perturbations, but shows negative values linked to the northward shift in the ITCZ.

The rapid adjustments to BC perturbations include changes in cloud cover. Figure 6 shows the global maps of changes in high cloud and low clouds. High cloud cover is slightly reduced in all runs and are surprisingly consistent between the runs. Low cloud cover increases globally for all simulations, especially for marine stratocumulus cloud regions. This increase of low cloud cover outside the west coast of North America is largest for the Asian emission increases and is correlated with surface cooling. In the emission areas local low cloud cover decrease, except for South Asian emissions where local low clouds increase. As in most other models, BC is not included as an ice nuclei in NorESM. This might have an impact on the monsoon

195 response and clouds. The version of NorESM we have used is known to have a relatively large convective transport of aerosols (Allen and Landuyt, 2014), which may cause an overestimation of BC in the upper troposphere.

Simulating the temperature response with coupled ESM requires large computer resources (here we have used 300 model year simulations for each experiment) due to the internal variability of the model. Alternatively, one can estimate the temperature response in broad latitude bands by using the ARTP approach. We compare our annual mean surface temperature estimates
200 from the coupled simulations with estimates using the much simpler ARTP-based approach based on the regional response coefficients (in K/Wm^{-2}) from Shindell and Faluvegi (2009). This is done by calculating the temperature responses in each band by using their response coefficients multiplied with our forcing estimates (i.e. the DRF), see Collins et al., 2013. Please note that the coefficients from Shindell and Faluvegi (2009) are normalized by DRF, and the rapid adjustments in the atmosphere are thus included in the regional response coefficients (as calculated in the coupled runs by Shindell & Faluvegi
205 (2009)). Figure 7 shows the comparison (temperature sensitivities in $\text{K}/(\text{Tg}(\text{BC})\text{yr}^{-1})$ per latitude band) between the ARTP-based method and our coupled model estimates. For the ARTP calculations in the Arctic response region, we have used numbers from Sand et al (2015a), which includes vertically resolved forcing and sea-ice/snow forcing. This is not included in the other latitude bands (as we did not have it available). GISS modelE and NorESM have the same equilibrium climate sensitivity (2.9 K). By reconstructing the temperature response, we obtain a fairly comparable response, even though the ARTP
210 constructed response varies more between the emission regions in each latitude band. Reasons for this might be differences in the vertical distributions of BC and indirect effects.

Lewinschal et al. (2018) calculated the same emission-to-temperature responses using NorESM, but for SO_2 emissions. They found that the temperature response was independent of emission location and as a global average equal to $-0.006 \text{ K/TgSyr}^{-1}$. As in this study, the Arctic was found to have the largest temperature response in all simulations. Here, we find global surface
215 temperature responses per unit BC emitted to vary between the emission regions, with a systematic north-south gradient; 0.029 (or 0.021 with high emission rate) K/Tg yr^{-1} for European emissions; 0.025 (0.023) K/Tg yr^{-1} for North American emissions; 0.019 (0.016) K/Tg yr^{-1} for East Asian emissions; and 0.017 (0.015) K/Tg yr^{-1} for South Asian emissions.

4 Summary and conclusion

We have estimated the temperature responses for BC emitted in four major emission regions in the Northern Hemisphere. The
220 largest temperature response was found in the Arctic, independent of emission region. Generally, the response is similar whether BC is emitted in North America or Europe, and to some extent also East Asia (but with slightly lower response in the Arctic). For [BC emitted in](#) South Asia the response is weaker due to a strong decrease in downwelling solar radiation and local surface cooling due to a displacement of the Indian monsoon. Regardless of emission region, BC cause a northward shift in the ITCZ. This is apparent both when using a coupled ocean and with fixed SSTs.

225 The Arctic temperature change per unit emissions depends on the magnitude of the forcing. The higher the emission rate, the lower the temperature sensitivity. This non-linearity is partly because enhanced absorption in the highest emission cases

increase vertical mixing so that of BC is transported higher up in the atmosphere, which decreases the surface temperature response, a feature also shown in (Ban-Weiss et al., 2012). For considerably lower emissions, the sensitivity could be higher. This result implies that the regional temperature coefficients calculated from high emission perturbations may be a conservative estimate, but in general that the linearity of normalized temperature effects of BC is fairly well preserved in our model. When comparing our global temperature responses per unit BC emitted to the response to SO₂ emissions for the same regions also using the NorESM model (Lewinschal et al., 2019), we find that the BC sensitivities are 3-5 times larger compared to SO₂. The global temperature responses to BC also vary between the emission regions with a systematic north-south gradient. By reconstructing the temperature response using the much simpler ARTP method, we find that the ARTP method works quite well, but that there are regional differences within the latitude bands, especially linked to circulation changes and the Indian monsoon.

Data availability

The model data can be made available upon request by the corresponding author. The source code for NorESM is available at <https://github.com/metno/noresm>.

240 Author contribution

MS performed the simulations, figures/statistics, and lead the writing of the paper. All authors contributed to the design of the study, the analysis of the results and the writing of the paper.

Competing interests

The authors declare that they have no conflict of interest.

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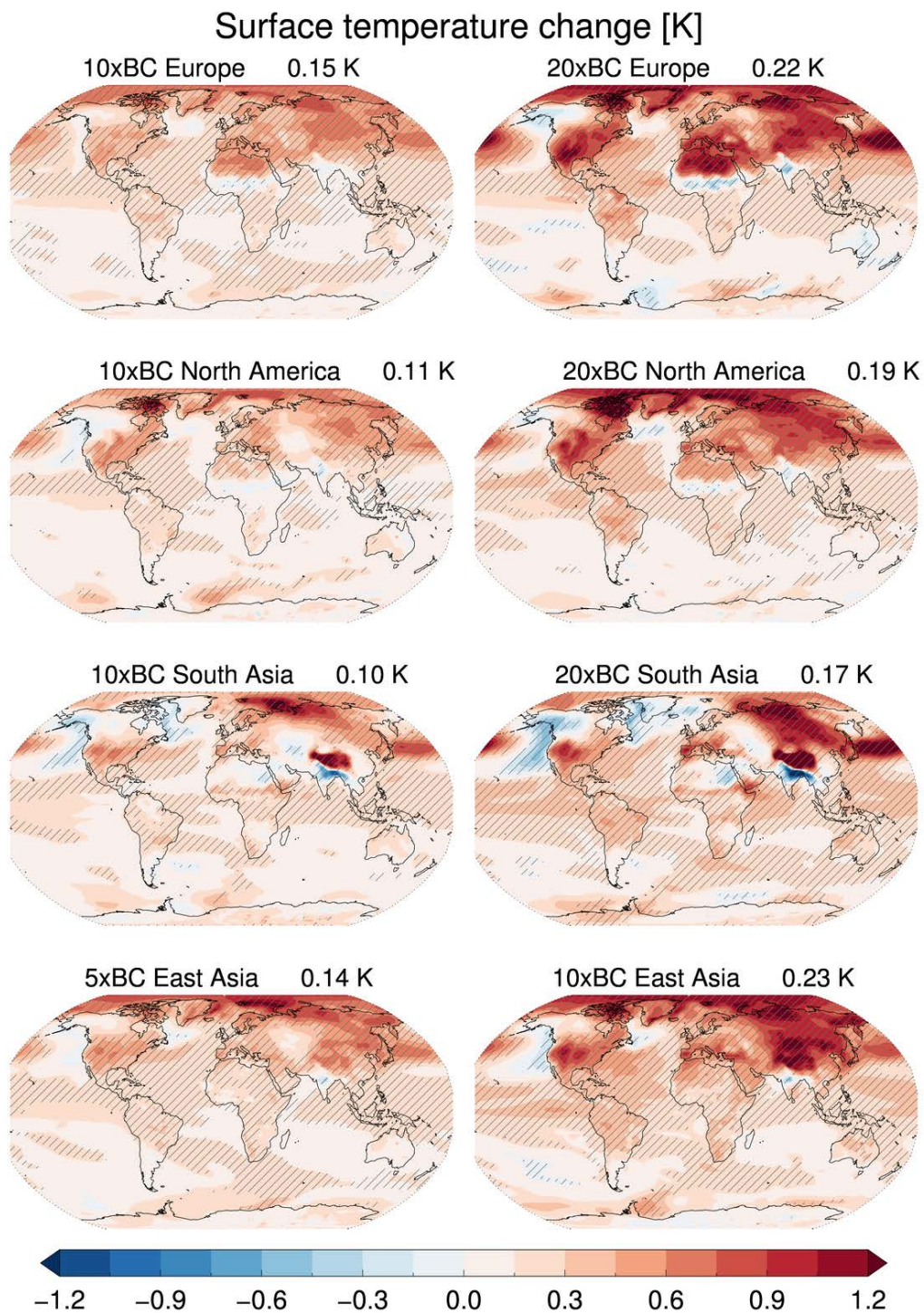
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355 **Figure 1: Surface temperature change (in K) for the perturbed runs minus the baseline. The emission region is given on each panel (from top: Europe, North America, South Asia, East Asia). In the right-side column the emission rate is doubled compared to the left-side column. The stippled areas represent statistically significant changes ($p < 0.05$).**

ΔT per global emissions [$K/Tgyr^{-1}$]

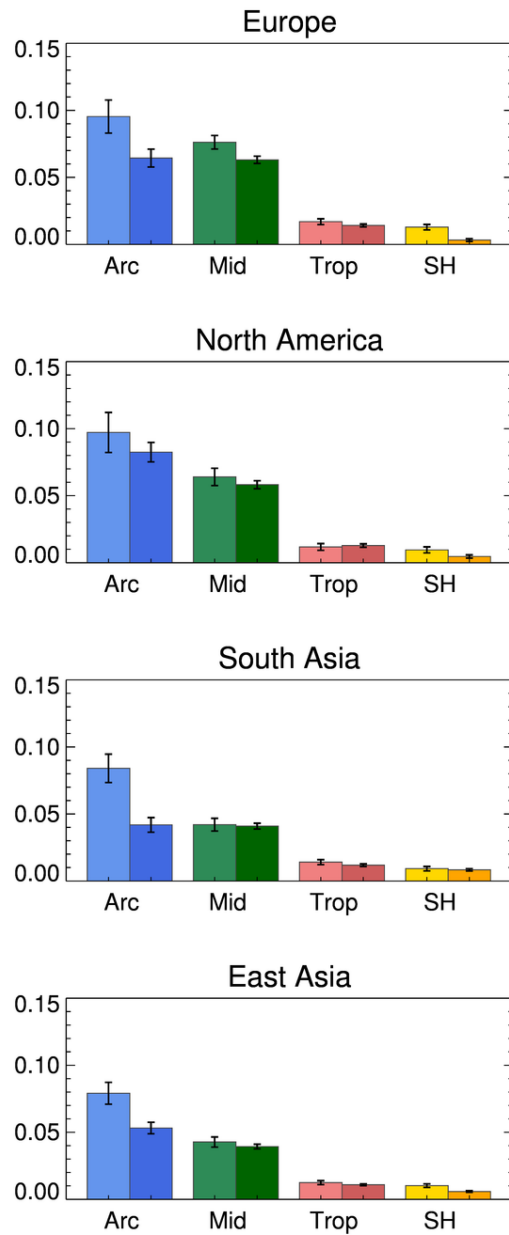
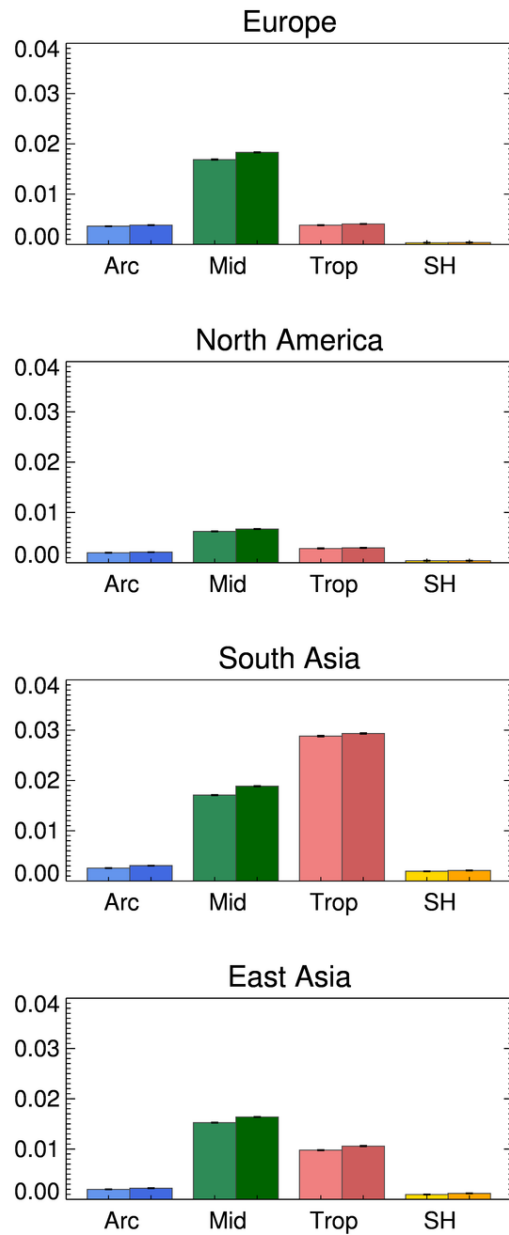


Figure 2: Regional surface temperature response per BC emissions ($K/Tgyr^{-1}$). The emission location (Europe, North America, South Asia, East Asia) is given on top of each plot. The response region (latitude band) is given on the x-axis. The emission rate is doubled on the right-side bars (darker colours) compared to the left-side bars (lighter colours). The error bars represent the standard error of mean.

360

Burden per global emissions [Tg/Tgyr⁻¹]



365 **Figure 3: Regional BC burden change per global BC emissions (Tg/Tgyr⁻¹).** The emission location (Europe, North America, South Asia, East Asia) is given on top of each plot. The response region (latitude band) is given on the x-axis. The emission rate is doubled on the right-side bars (darker colours) compared to the left-side bars (lighter colours). The error bars represent the standard error of mean.

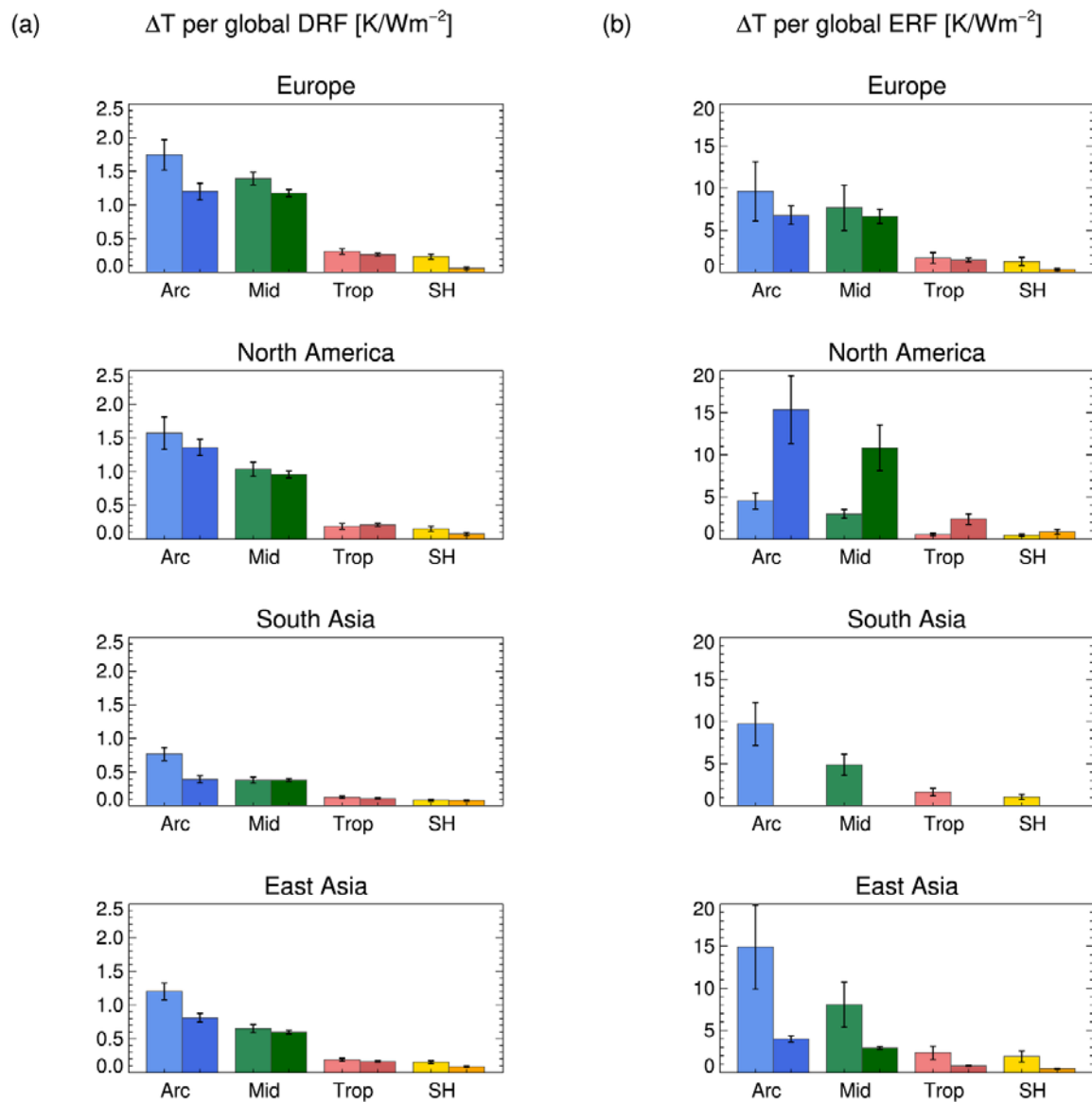
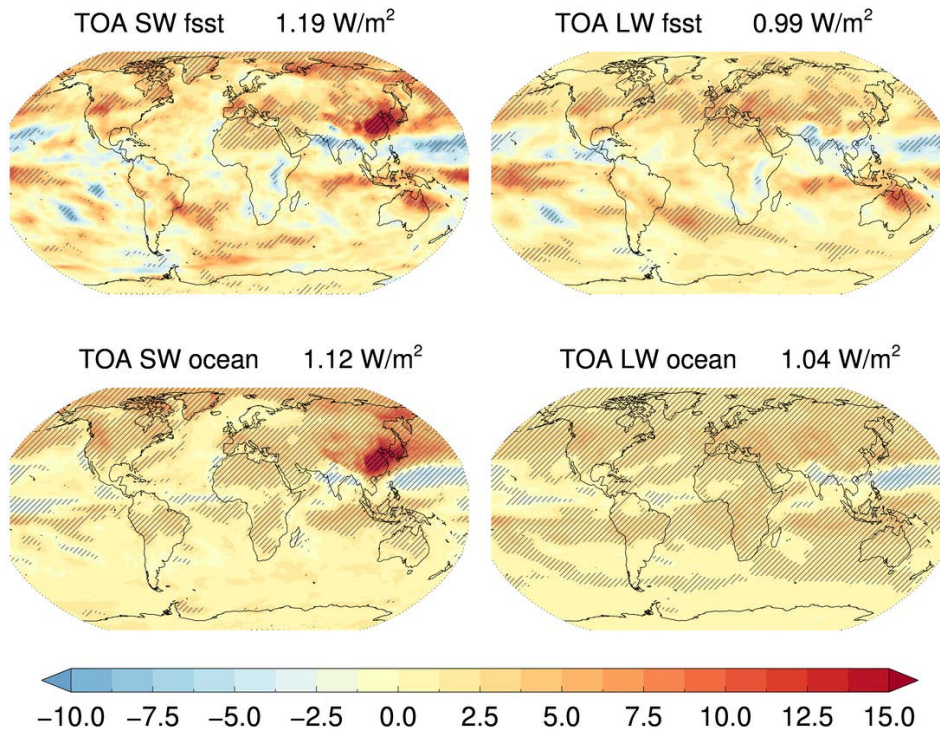


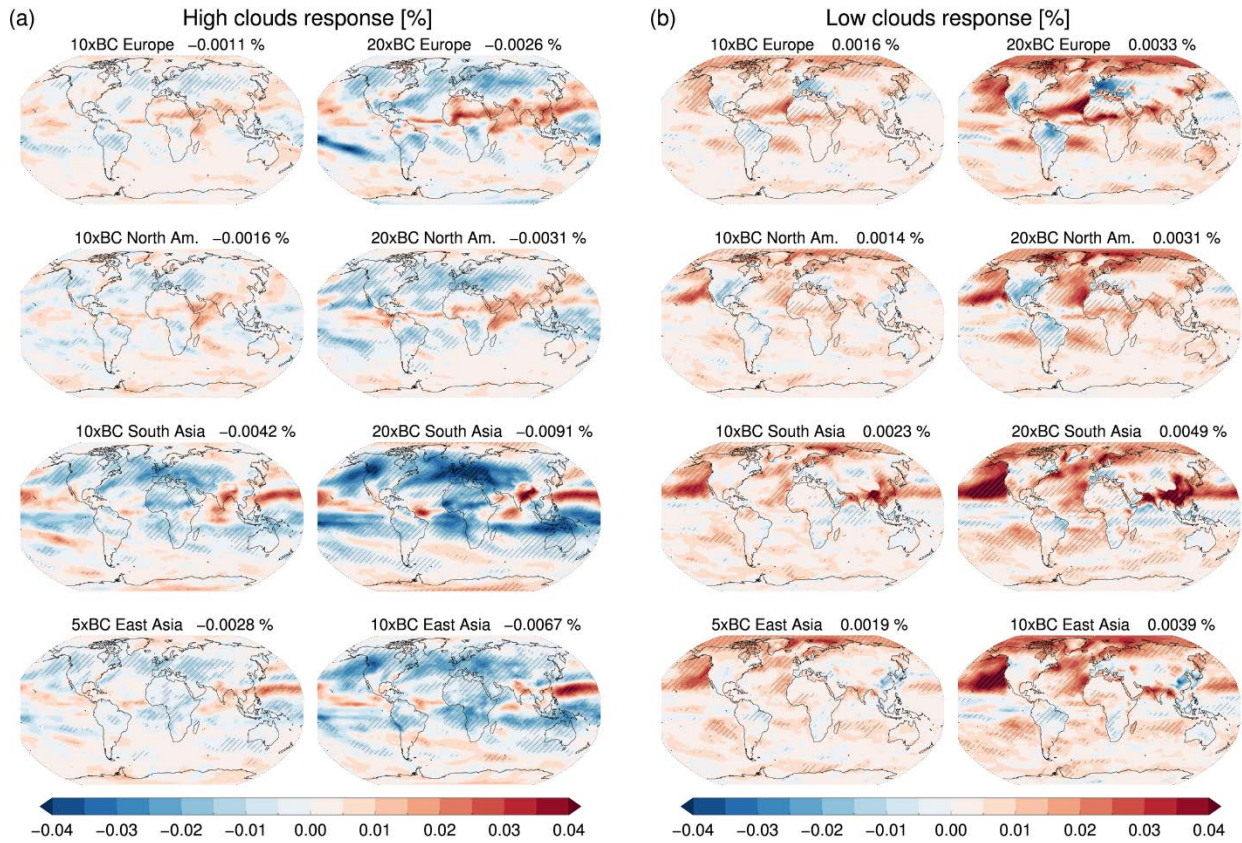
Figure 4: Regional surface temperature response per global BC (a) DRF and (b) ERF (in K/Wm⁻²). The emission location (Europe, North America, South Asia, East Asia) is given on top of each plot. The response region (latitude band) is given on the x-axis. The emission rate is doubled on the right-side bars (darker colours) compared to the left-side bars (lighter colours). The error bars represent the standard error of mean.

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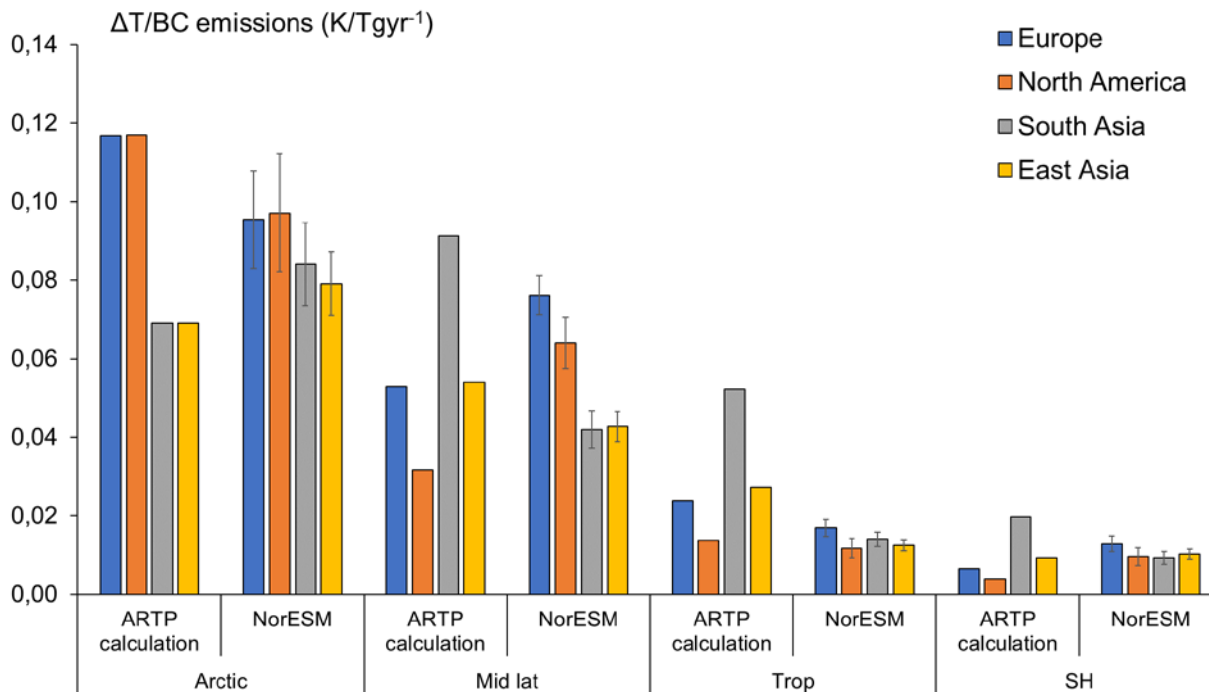
10xBC East Asia



375 | **Figure 5: Change in delta TOA net SW and LW radiative fluxes (net SW = sum of downward and upward) and net LW) for the 10xBC in East Asia perturbation minus the baseline. Top row shows the runs with fixed SSTs, and bottom row shows their runs with fully coupled ocean. The stippled areas represent statistically significant changes (p < 0.05).**



380 **Figure 6: Response in high clouds (located above 400 hPa) (a) and low clouds (surface up to 700 hPa) (b) in (%). The emission location (Europe, North America, South Asia, East Asia) is given on top of each plot. In the right-side column the emission rate is doubled compared to the left-side column.**



385 **Figure 7: A comparison between ARTP calculated and NorESM calculated regional surface temperature response per unit BC emissions (K/Tgyr⁻¹). The lowest emission perturbation is shown here (10×Europe, 10×North America, 10×South Asia, 5×East Asia). The response region (latitude band) is given on the x-axis. The error bars for NorESM represent the standard error of mean.**

Tables

390

Table 1: Overview of the model simulations in this study and the global BC emissions in each simulation (in Tg BC/yr).

Control	Year 2000 control run 3×100 years fully coupled equilibrium simulations	<u>7.7</u>
10 × Europe	Same as control, but with BC emissions in Europe multiplied by 10	<u>12.8</u>
20 × Europe	Same as control, but with BC emissions in Europe multiplied by 20	<u>17.9</u>
10 × North America	Same as control, but with BC emissions in North America multiplied by 10	<u>11.9</u>
20 × North America	Same as control, but with BC emissions in North America multiplied by 20	<u>16.1</u>
10 × South Asia	Same as control, but with BC emissions in South Asia multiplied by 10	<u>13.5</u>
20 × South Asia	Same as control, but with BC emissions in South Asia multiplied by 20	<u>19.3</u>
5 × East Asia	Same as control, but with BC emissions in East Asia multiplied by 5	<u>15.0</u>
10 × East Asia	Same as control, but with BC emissions in East Asia multiplied by 10	<u>22.4</u>

Table 2: Surface temperature change per global BC emissions (in K/Tg yr⁻¹). Bold numbers represent statistically significant differences between the emission rates (using 10 vs. 20, or 5 vs. 10 for East Asia).

	Arctic	Mid lats	Tropics	SH
10 × Europe	0,10	0,08	0,02	0,01
20 × Europe	0,06	0,06	0,01	0,00
10 × North America	0,10	0,06	0,01	0,01
20 × North America	0,08	0,06	0,01	0,00
10 × South Asia	0,08	0,04	0,01	0,01
20 × South Asia	0,04	0,04	0,01	0,01
5 × East Asia	0,08	0,04	0,01	0,01
10 × East Asia	0,05	0,04	0,01	0,01

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Table 3: Surface temperature change per global direct radiative forcing (in K/Wm⁻²). Bold numbers represent statistically significant differences between the emission rates (using 10 vs. 20, or 5 vs. 10 for East Asia).

	Arctic	Mid lats	Tropics	SH
10 × Europe	1,74	1,39	0,31	0,23
20 × Europe	1,20	1,18	0,26	0,06
10 × North America	1,57	1,04	0,19	0,15
20 × North America	1,36	0,96	0,21	0,08
10 × South Asia	0,77	0,38	0,13	0,08
20 × South Asia	0,39	0,39	0,11	0,08
5 × East Asia	1,20	0,65	0,19	0,15
10 × East Asia	0,81	0,60	0,17	0,09

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