Reply to referee 2

We thank the referee for the thorough and constructive review. We have revised the manuscript accordingly. Please find below the point by point reply to the individual comments.

General comment
This manuscript investigates how microphysical and chemical interactions between dust and pollution alters the properties of aerosols and their direct radiative effects (DRE). The experimental design is simple and effective. A set of four simulations have been used to model the properties of aerosols when dust and pollution are either emitted separately, or together so that they interact. By contrasting results from these simulations the study reveals the “interaction” term showing how the properties of the dust and the pollution-related aerosol change due to two-way microphysical and chemical interactions. It is interesting to explore such interactions and the changes in aerosol radiative effects are not trivial, so worth noting.

The main result is that the dust-pollution interactions lead to a ~0.05 Wm$^{-2}$ change in net flux at TOA, dominated by increased SW reflection. This occurs mainly due to increases in accumulation-mode aerosol mass and AOD. The AAOD also drops slightly, though it is not completely clear why, though it is perhaps related to a drop in coarse-mode dust. As climate models and Earth-system models are increasingly adopting more complex microphysical aerosol schemes it is worthwhile understanding what happens as such interactions are enabled. To my knowledge this manuscript is novel and I would judge it to be relevant and worth publishing in ACP. The text is generally well written, well structured and concise. However, significant improvements in the analysis and interpretation of the results are required for the study to be published.

Major comments
The main difficulty with this manuscript is that it is not very clear what has caused the negative change in aerosol DRE. The text interprets this as a change in dust forcing, or an “anthropogenic radiative forcing associated with dust”. However, the dust-pollution interaction is a two-way process and changes both the dust and the fine-mode anthropogenic aerosols. Figures S11, S12, S13 indicate significant changes to the “pollution-related” aerosol once the dust and pollution are emitted together so one can not attribute the change in DRE entirely to the dust.

Indeed the interaction is a two-way process and technically we treat both directions equally as manifested by Eq. (2). The interpretation as a change in dust forcing is motivated by the fact that historically the dust was there before the pollution and therefore the pollution modified the already existing dust forcing. In contrast, the opposite case where dust is added to pollution never occurred on global scale. Therefore and to make the discussion of the results more comprehensible, we use this interpretation as guiding principle but nevertheless discuss the impact of dust on pollution. The term “anthropogenic radiative forcing associated with dust” is supposed to reflect that the effect is linked to (but not solely attributed to) dust and at the same time anthropogenic because of the anthropogenic origin of the pollution.
The main cause of the negative change in aerosol DRE seems to be the increases in accumulation-mode aerosol, but it is not very clear from the study which aerosol components have contributed to this increase. More information is required to show how the aerosol properties have changed, including changes in aerosol mass, chemical composition, hygroscopity, and possibly particle size across the relevant size modes.

The increase of the accumulation-mode burden is caused by mineral dust due to the reduced coagulation with coarse particles in the presence of pollution. The accumulation mode-burden of other components is generally reduced due to the increase of coagulation in the presence of dust, exceptions are the ions (Fig. S11, now S12). The increase of accumulation mode dust dominates in dusty regions, resulting in a net-increase of the accumulation-mode burden. We have added figures showing the interaction effect on dust, BC, SS and water burdens separately to the supplement and discuss them in the main text.

Related to this, it is not very clear what has caused the changes in AAOD and SSA. Presumably AAOD reduces in dusty regions due to a drop in coarse-mode mass? The AAOD increases across the Sahel are apparently due to increases in BC and OC mass (lifetime) but it would be good to see the evidence to support this. However, it isn’t clear why the SSA drops in non-dusty regions that are quite remote from dust sources. Presumably there is a relative reduction in non-absorbing aerosol components such as sulphate and/or nitrate, but why does this occur in regions very remote from dust. There are some clues in Figure S11 but there are many competing effects and the information is not comprehensive enough to understand what it going on.

The AAOD reduction in dusty regions is mainly caused by the change of the accumulation mode composition in the presence of dust which transfers absorbing carbonaceous components from the accumulation to the coarse mode. This becomes more clear from the new aerosol component burden plots in the supplement. On the other hand the increased accumulation mode dust increases the AAOD, this effect dominates south of the Sahel to produce a net AAOD increase. In contrast, over Asia where not only the burden of absorbing components but also the AOD decreases, both effects reduce the AAOD so that a negative net effect is obtained even relatively remote from dust sources. Additionally, as you mention, in dusty regions the coarse mode mineral dust and hence absorbing component is reduced, which further contributes to the AAOD decrease.

The other major concern I have is the short duration of the simulations (only one year). Given the episodic nature of dust emissions there is likely to be considerable interannual variability in dust loadings and in how these interact with pollution outbreaks. This could affect both the magnitude and spatial patterns of the results. I would recommend extending the simulations to at least a 10 years, unless the authors can provide evidence that a single year is sufficient to gauge the magnitude and characteristics of the dust-pollution interactions.

Despite the episodic nature of dust emissions the global annual averages presented here are relatively robust regarding interannual variations which is confirmed by a lower resolution sim-
ulation over 10 years that yields coefficients of variation (CV) for our main results below 10%. Therefore, even though the results are explicitly presented for 2011, they are considered representative for recent years. We have added a paragraph to the methodology section and CV estimates to the caption of Fig. 6.

### Minor comments

**Abstract:** The abstract is short and direct but needs to be altered to reflect the concerns above. In particular, the change in aerosol DRE is described as a “radiative forcing” here and throughout the manuscript. This could be confusing or misleading as the term “forcing” is usually used to indicate a change in radiation balance due to a perturbation in aerosol emissions. The ∆F is really more “the change in aerosol DRE due to dust-pollution interactions”. This could be given a label such as ∆DREint to avoid using this long definition each time in the text.

While a “forcing” is often attributed to emissions, it is not uncommon to assign a forcing to other effects such as changes in the solar irradiance, surface albedo changes or cloud adjustments due to aerosols. Especially the latter, the forcing of aerosol-cloud interactions, bears similarities to the forcing by dust-pollution interactions so that we consider using the term “forcing” in our context to be not exceptionally confusing.

**P1 L5:** Please spell out EMAC.

We have expanded the acronym.

**P1 L7:** Whilst the magnitude of the change in TOA radiation balance is worth noting, I would not describe it as large. In fact it is quite small compared to the total DRE of aerosol in present-day climate (∼ -2 to -3 Wm-2).

We have deleted “large”.

**P1 L10:** Please quantify this “considerable fraction”.

We have quantified the fraction (40%).

**Methods:** **P3 L10:** It is not quite clear what the term “prognostic radiative-transfer calculations” means. Presumably this is the radiative-transfer calculations that are used to calculate fluxes and heating rates in the simulations. Using “prognostic” is a bit confusing since the prognostic aerosols are clearly not used in the radiation scheme, and a radiation scheme is not itself prognostic.

We have reformulated to “the aerosol radiative effect on the dynamics is computed using the extinction, single scattering albedo and asymmetry factor from the Tanre aerosol climatology”.
P₃ L13: Is the dynamical evolution of the atmosphere (wind, temperature, moisture, cloud) identical in all four simulations? I would have thought so if the prognostic aerosols neither interact with the clouds or the radiation scheme.

Yes, they are identical which is now mentioned in the discussion of Eqs. (1) and (2) in section 2.

P₄ L17-18: I think that you need to swap F₂ – F₄ and F₃ – F₄ in this sentence. From my reading of the text F₃ – F₄ corresponds to the dust radiative effect and F₂ – F₄ corresponds to the pollution.

This has been fixed in the revision.

P₄ L23: So it sounds like a radiation double-call procedure has been used, as outlined in Ghan et al. (2012). https://journals.ametsoc.org/doi/10.1175/JCLI-D-11-00650.1 If so it might be useful to reference Ghan et al. (2012) as this paper outlines the double-call concept fully.

We have added the reference.

Results: P₅ L20 / Fig 1: Does the mass shown in Fig 1 include the aerosol water content?

Yes (we have added a note in the caption of Fig. 1).

Figures 1 – 3: Exactly how are the differences in mass and optical properties calculated? Are these calculated using the same logic as in equation 1?

Figures 1 and 2 are calculated using Eq. (1). Fig. 3 shows the difference of the SSA results from the simulation with (simulation 1) and without pollution (simulation 3) (both simulations include dust). Equation (1) cannot be used to study the effect on the SSA since the SSAs of dust and pollution are not additive when neglecting the interaction.

P₃ L24: Would it be possible to provide a figure for the dust mass loading, or at least refer to Figure S1 here so the author can see where the “dust affected regions” are simulations.

We have added the reference to Fig. S1.
Section 3: I found this section difficult to follow (particularly the top half of P6) and it did not provide a full explanation of how / why the aerosol properties changed.

P6 L7-9: This argument needs explaining more fully. It is clear that the the coarsemode dust is removed more rapidly due to secondary aerosol forming on the particles leading to more rapid wet deposition. However, what happens to the accumulation mode dust? Wouldn’t the same process also speed the removal of the accumulation mode dust compared to the simulation where pollution was not emitted with the dust? From figure S11 it looks like the overall mass of sulphate and nitrate aerosol in the accumulation-mode has decreased. So this would tend to decrease accumulation-mode mass and the hygroscopicity, yet total accumulation-mode mass and AOD have increased. Presumably the mass of BC and OC must have increased dramatically to compensate the decreases in sulphate and nitrate. Is there evidence of this? It would be good to see all the relevant mass components in figure S11 and have the full story explained.

Compared to the coarse mode, an additional sink for the accumulation mode is the coagulation with coarse particles, which is less efficient in the polluted simulation 1 than in simulation 3 without pollution (due to the more efficient removal of coarse particles). Therefore simulation 1 produces more accumulation mode dust particles than simulation 3 despite a more efficient deposition. This difference dominates Eq. (1) resulting in the burden increase displayed in Fig. (i). We have added the corresponding plots for mineral dust, black carbon, sea salt and water individually in Fig. S12 in the supplement, showing that the accumulation mode burden increase is due to dust.

P6 L25-26: The SSA has reduced in most non-dusty regions. Is this due to an increase in the relative proportion of BC and/or OC versus sulphate and nitrate? Why would this have occurred even in regions very remote from dust? It would be good to provide a table listing the global-mean values and global mean changes in relevant quantities, such as AOD, AAOD, radiative flux changes and the various aerosol mass components.

Please note that Fig. 3 does not show the result of Eq. (1) but the SSA difference between simulation 1 with pollution and simulation 3 without pollution. The negative values very remote from dust result from adding pollution including absorbing BC and OC to the mostly non-absorbing natural background aerosol (e.g., sea salt, water). Since the natural aerosol burden can be very low compared to the burden from pollution, the SSA difference in remote regions is not very relevant (hence the colour scale cut-off at -0.012). Because global means integrate compensating effects and large areas unaffected by dust, regional values are more informative (the relevance of the global TOA forcing for the climate’s energy budget make it an exception), accordingly we have added a table with regional mean values and the corresponding contributions of the dust-pollution interactions for relevant quantities.

P6 L20: How has the aerosol become more reflective? I suspect the drop in AAOD is dominated by the decline in coarse-mode dust mass so I would omit “due to the higher reflectance” from this sentence.

The water which is taken up by the aged hygroscopic dust particles reduces the average imaginary
refractive index of the particles, increasing the reflectance.

P6 L21. The AAOD is presumably increased in the Sahel due to increased BC mass (and brown carbon if this is included in the model). The explanation that this is due to AOD increase doesn’t make sense, only one can say that both may be increasing for similar reasons.

The AAOD is proportional to the AOD (for constant SSA), thus an increase of the AOD increases the AAOD.

Conclusions: The main concerns given above need to be addressed throughout the conclusions. The change in aerosol DRE can not be interpreted as a change in dust forcing as it is caused by both changes in dust and pollution aerosol properties. The conclusions section is very concise, which is good, but a bit more discussion is required to explain what has caused the changes in aerosol DRE, and how these are linked to aerosol processes and changes in aerosol properties (mass, hygroscopicity, optical properties etc).

While we agree that the impact of dust on pollution is crucial which is now emphasized more throughout the revised manuscript, the change in the forcing is by definition the change of the dust forcing which occurs when pollution is added to the natural scenario (see Eq. (i)). Mathematically this is identical to the change of the pollution forcing when adding dust to a dust free scenario ((F1 - F3) - (F2 - F4)). However, as mentioned above, this never occurred in the real world on a global scale, therefore we consider the interpretation as change in dust forcing to be more instructive. We have added a paragraph summarising the causes of the changes in aerosol forcing.

P8 L20: The maximum impact to the south of the Sahel is given as -2.5 Wm-2 here but -2 Wm-2 in the abstract. From reading closer I see this is because the -2.5 Wm-2 quoted here corresponds to the surface forcing and the abstract gives the TOA forcing. Please use the same headline result in abstract and conclusions.

We have included the TOA result in the revised conclusions.

Figures: There are a lot of additional figures in the supplementary material and some offer an unnecessary level of detail on the spatial and seasonal variability of aerosol radiative effects (S5, S9, S15, S19 - 23). Given that the spatial distributions and seasonality of results are probably very specific to the model and the meteorological evolution in this specific set of simulations, this level of detail is not particularly useful and could be misleading. The 3D visualizations of aerosol heating in particular are not at all useful.

It is true that due to the shorter underlying time period the seasonal results have a higher statistical uncertainty and are less representative for the same season of other years. But since the meteorology is nudged towards reanalysis data, the meteorological evolution in the simulations agrees well with observations and so do the aerosol concentrations which is supported by many previous studies (e.g., Pozzer et al. 2012, Abdelkader et al. 2015, 2017, Metzger et al. 2016, Kling-mueller et al. 2018). Therefore we consider the seasonal results to be not specific to the sim-
ulations but well representative for the year 2011. Given the seasonality of dust emissions we assume the seasonal results are valuable to some readers and prefer to retain most of the figures while removing the 3D plots.

Figure S11: As expressed above, to really understand how the dust-pollution interaction affects aerosol properties this figure needs to include changes in OC, BC, dust, water and sea salt. The figure may need expanding to two or three figures to give a complete summary of the changes in aerosol mass and composition.

Figure S11 focuses on the main aerosol ions which interact with the mineral cations in the dust and are in exchange with precursor gases shown in the same figure and therefore play a distinct role. The revised supplement additionally includes the Figures for dust, BC, SS and water burdens separately (Fig. S11) and a discussion in the main text. Qualitatively the same effect as for BC is obtained for OC but no OC coarse mode model output is available.

Figure S12 & S13: It is quite interesting to have this kind of information, but it was rather difficult to interpret how the mass of dust, pollution, water and natural aerosols change between the three scenarios. It would be clearer to have just one bar for each emission scenario (all emission, no pollution, no dust) and have each bar stacked showing the relevant mass components (dust, pollution, water and other natural aerosol). This way it would be totally clear how each mass contribution has changed depending on what has been emitted. It is still very interesting to provide this separately for both the accumulation and coarse-modes. Would it be possible to produce this kind of figure also for the global mean changes? As this analysis is really quite important to the story of the paper it would also be worth considering moving this (especially a plot with global-mean changes) to the main article.

Figures S12 and S13 are closely related to Eq. (2), the “No dust” and “No pollutions” bars are stacked to allow a direct comparison of the term corresponding to (F2 - F4) + (F3 - F4) with the term corresponding to F1 - F4. We have straightened the figure by stacking the mass contribution bars, but still stacking the “No dust” and “No pollution” values and moved it to the main article. This is complemented by a new figure in the supplement showing regional burdens (which are more useful than global averages) with one bar per simulation as proposed.

Figure 6: This graph really emphasizes the interpretation that the dust-pollution interaction has strengthened dust forcing, which is misleading since the interaction has also altered the strength of the anthropogenic (pollution) aerosol forcing. For a more balanced summary it would be good to include a bar for the forcing from pollution and bars for “Dust + pollution with interaction”, and “Dust + pollution without interaction” instead of the bar with “Dust interacting with pollution”.

As argued above, while we agree that the interpretation as a change in dust forcing is not the only interpretation possible, we consider it to be a valid and well motivated perspective. Aside from this the comparison with the dust forcing works better due the similar magnitude. The underlying reason is that for both, the interaction and the dust forcing, regional contributions
compensate each other. The pollution forcing is discussed in section 2.