Answers to reviewer comments regarding the prepared manuscript 'Coupling aerosol optics to the chemical transport model MATCH (v5.5.0) and aerosol dynamics module SALSA (v1)'

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This document contains answers to the two anonymous referee comments and the short comment given by A. Kerkweg to the manuscript "Coupling aerosol optics to the chemical transport model MATCH (v5.5.0) and aerosol dynamics module SALSA (v1)" prepared for the journal GMD. The first (to anonymous referee #1) answers follows directly after this first page, page 2-15, the second answer (to anonymous referee #2) on page 16-19 and the third answer (to A. Kerkweg) on page 20. The answers will after that be followed by a marked up manuscript, to which all the changes are referred to.

Author response regarding the manuscript 'Coupling aerosol optics to the chemical transport model MATCH (v5.5.0) and aerosol dynamics module SALSA (v1)', to the reviewer 1

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The authors describe the implementation of a new aerosol optics model into the atmospheric chemistry transport model MATCH. They compare simulation results with this new optics model to results with the previously implemented one. In order to assess the significance of the differences in simulated optical properties for estimates of the aerosol radiative forcing and for data assimilation, they discuss simulations with the new optics model in combination with the former bulk aerosol treatment in MATCH and in combination with the more recent aerosol dynamics model SALSA (as part of MATCH). The differences between the simulations with and without aerosol dynamics serve as a reference to evaluate the significance of the impact of the different optics models.

From my point of view, the topic falls well within the scope of GMD, the idea of this study is sound, and it is well motivated. I think the aerosol modeling community will benefit from this study once the following comments to the authors are adequately addressed.

Thank you for taking your time reviewing our paper. Your comments have been most valuable to improve our paper and we specially thank you for being thorough and constructive in your comments. Below you will find your comments in blue together with our answers in black for each comment in major, minor, specific and typo section. Following the comments and answers, a document highlighting the differences between the original manuscript and the revised manuscript is added. To which we also refer in the answers.

1 Major comments

We will answer the first three comments together, since they are interrelated, and since the reviewer presented these comments as three different alternatives to improve the manuscript.

1. In order to assess the significance of the differences between the simulations, I think it is necessary to provide some measure of uncertainty or variability in the simulated aerosol optical properties and radiative effects, depending on the simulated aerosol (component) mass mixing ratios. From my point of view, an analysis of the involved nonlinearities that the authors mention in the Conclusions cannot be completely deferred to a later study, as they also impact the assessment of the significance of differences between simulations with aerosol (optics) models

of different complexity. If, as the authors state, the "test cases [may not be] in any way representative for typical aerosol and black carbon loads", then the study seems to me like a purely academic exercise.

- 2. Additionally (or maybe alternatively, as this would also provide a backdrop against which to gauge the significance of the discussed simulation differences) the study would benefit from comparisons with observations, in order to put the aerosol optics and dynamics impacts into a context of simulation-observation differences. AERONET and various satellite data sets come to mind here, and lidar observations could probably also be used.
- 3. A third option that might help make the conclusions a bit more robust would be to run another simulation with the old optics model in combination with SALSA and check if the MT-SALSA differences (using the old optics model) and the EXT-CGS differences (using SALSA) are similar to those discussed in the manuscript.

We first want to respond to the reviewer's concern about the representativeness of our study (comment 1). Although we picked only four locations and two points in time, we did choose these cases in order to cover as different situations as possible (two points over land, one north and one south of the Alps; two points over water, one in the north, one in the south; one summer and one winter day). The main goal we pursue with our approach is not only to quantify the differences between the two optics models, but also to show where those differences come from. We start our analysis by considering how morphological differences in the model particles cause differences in single-scattering optical properties. Then we investigate how those differences impact the radiative fluxes throughout the atmosphere, and how that, in turn, impacts the TOA radiative forcing. We want to help the reader appreciate that subtle microphysical properties impact radiometric properties of particles, particle populations, and macroscopic media. This chain of physical processes would remain obscure in a statistical analysis; it can only be revealed in a detailed analysis of selected cases. We are aware that this may be mind-boggling and demanding for the reader, and that a simple statistical analysis would be much simpler to follow. But we also think that our approach is quite valuable, because we do not just tell the reader what kind of errors one may introduce by using very simple optics models, but we show why this is

We have deleted the misleading part in Sect. 3.1 regarding the representativeness of the selected cases.

Now we turn to the reviewer's main concern. As we understand it, the reviewer wants more facts to strengthen our main conclusion, namely, that the choice of optics model can have a significant impact on the calculated radiometric properties. The reviewer makes three alternative suggestions, 1. Provide a measure of uncertainty or variability (also mentions analysing the non-linearities of the optics model); 2. Perform comparison with observations; or 3. Perform additional computations with the old optics model implemented into SALSA.

We performed additional computations and analyses that combine some ideas of 1. and 3. More specifically, we implemented, as the reviewer suggested, the old external-mixture/homogeneous-sphere optics model into SALSA and repeated the computations. Further, we computed mean errors for different optical properties over the entire geographic region and averaged over a whole month. The results are presented in a new table (Table 4 in the revised version), and discussed in a dedicated section (Sec. 3.1 in the revised version). The statistical analysis confirms the essential conclusions of this study. We believe that this addresses the reviewer's main con-

cern by supplementing our study with more quantitative information on the differences between the optics models. We also make it clear in the revised version that the analysis of averaged optical properties mainly serves to draw some general conclusions, while the case-studies mainly serve didactic purposes to explain *why* different optics models can give different predictions for radiative fluxes.

We did not analyse the non-linearities of the optics model; even a crude analysis would probably take several months of work. However, we are currently applying for funding for a dedicated project that will deal with just this problem. Also, we do not wish to add a comparison with observations to this study. This would completely change the character of this paper. As chemical transport models have numerous sources of error and biases, a meaningful comparison with measurements comparison would require a comprehensive analysis of all model uncertainties. This would completely shift the focus of this paper away from the optics model toward various other error sources in MATCH and SALSA, which are not the subject of this study.

4. Please check the publications by Jacobson (2000), Matsui et al. (2013), and Klingmüller et al. (2014), and refer to them if and where appropriate.

The paper by Jacobson (2000) is based on the use of the core-shell model, which underestimates the absorption cross section (which was, however, not known until quite recently). We added a reference to this paper in the discussion of our optics model in Sect. 2.3.2 (p. 13, 1. 346). We have also added the references Matsui *et al.* (2013) (p. 15 1.407), Andersson *et al.* (2015) (p.4 1. 104, p. 7 1. 216, p.15 1. 407) and KlingmÃijller & Lelieveld (2014) (p. 4 1. 97) as well as Kokkola *et al.* (2008) (p.4 1. 104).

5. Please give some details on the model setup, e.g., model domain, horizontal and vertical resolution, etc.

This information has now been added to Sect. 2.2 (p. 51. 139-145).

6. As far as I understand, the effects of including more detailed assumptions on internal particle structure are only studied for BC-containing particles, i.e., not for dust-containing particles, for instance. Please clarify this throughout the manuscript.

The paper does have a dedicated methodology section (Section 2). In subsection 2.3.2 we give a very detailed, itemized, and unambiguous description of the optics model, which we believe leaves very little room for misunderstandings. Therefore, we would prefer not to repeat the model description in other parts of the paper, for the sake of conciseness.

- 7. Section 3.1.1 requires improvement.
 - Please expand on the results shown in Table 3:
 - What do you mean by "the same behaviour"?
 - Are the reasons for this "same behaviour" also the same?
 - What about Northern Italy in winter?

We have re-written the whole result section as well as the mentioned paragraph in a, hopefully, clearer way, This paragraph can now be found at p. 22 l. 605-612.

• What is a "dominant feature" (p. 10754, l. 12) if the "differences [...] are almost negligible" (p. 10755, ll. 4 and 5)?

We re-wrote this sentence as follows: "Again, differences in $\Delta F_{\rm net}$ at TOA are mainly caused by corresponding differences in the upwelling diffuse radiative flux $\Delta F_{\rm u}$." See p. 22 1. 618-619.

- The paragraph on Fig. 8 confuses me:
 - How do you arrive at the conclusion that the MT-SALSA difference is "not as prominent" over the Mediterranean as over Northern Italy, given that in the next paragraph you state that this difference is small at all locations except the Mediterranean? This was indeed confusing, because in the first paragraph we referred to absolute differences, but in the following we mean relative differences (but we did not say this clearly). We have now made this distinction throughout the text. In addition to this comment and the next item of this comment, we have changed the paragraph about black carbon forcing in the context of comparing MT and Salsa, completely. Figure 8 is now also referred to as figure 10 according to specific comment 26. See p. 23.
 - The word "difference" appears very often in this paragraph and refers to different contexts. This makes the discussion hard to follow.
 We agree that this paragraph was confusing and little helpful. The main point is that the two models are similar, because the optical properties do not differ appreciable in this case. We emphasised this fact in the revised version, and deleted the rest of the text.
 - Please expand on the "multiple scattering effects".
 Reference to multiple scattering effects has been removed see preceding item.
- Please expand on the differences between summer and winter. How can a larger difference between the MT and SALSA simulations be tied to a larger difference in the BC mixing ratios if the summer ΔF_{net} difference is greater over the Mediterranean, where the BC difference is smaller, than over Northern Italy?
 - When taking a closer look at the concentration differences (not shown), then it becomes clear that the sensitivity of the forcing rate to concentration differences is rather low. Thus, when we have differences by a factor of 2–10, this has a noticeable effect on the forcing rate. But the tiny differences we see in summer between the two models over Northern Italy and the Mediterranean has no clear effect. We have re-written and expanded the discussion of the table accordingly. See p. 23 1. 659-671.
- 8. In many cases the language of the manuscript does not seem precise enough to me. For (some) details, see the Minor comments and Specific comments below

We did our best to improve the preciseness of our language (see our response to the minor comments).

2 Minor comments

1. What are the "known and important effects from using aerosol dynamics"? (Quoted from the Abstract, but also appears in the Conclusions in similar form.)

We rephrased the abstract and conclusion as well as added references to make our statement more elaborated and robust. We also realise that the term "aerosol dynamics" is a broad and non-specific term, therefore we have exchanged this term to the more proper "aerosol microphysics". This will help us address the vague formulations that the referee pointed out.

The statement refers to the fact that the microphysical processes involving aerosol particles (aerosol dynamics) strongly affects their size distributions and mixing state, as well as their direct and indirect radiative forcing. This has been investigated by several studies, among others Matsui *et al.* (2013) that looks at how ageing processes affects radiative processes together with different mixing states and Kokkola *et al.* (2008) as well as Andersson *et al.* (2015) describes and evaluates an aerosol microphysical module in a chemical transport model.

2. In order to avoid confusion concerning different types of atmospheric or climate models as much as possible I would recommend to follow the terminology out- lined in Dameris and Jöckel (2013)

Thank you for noting the terminology of model types. We have now used the term "Chemistry-Climate" model instead of "Earth system" model throughout the text. The changes occur on p. 1 l. 4, p. 2 l. 30, 34 and 50, and p. 53 l. 836.

3. I would suggest to replace most, if not all, occurrences of "aerosols" by an appropriate choice of "aerosol", "aerosol particles", "aerosol populations", "aerosol components" or similar, as the term "aerosol" technically does not only refer to the particulate phase, although it is often used this way in spoken language.

We have searched through the whole document and replaced most occurrences of "aerosols" with either "aerosol particles", "particles" or "aerosol components".

4. Furthermore, there is often a distinction between "aerosol" and "black carbon", although the latter is of course a component of the former.

Black carbon is, of course, part of the aerosol phase. In the text we do talk about both aerosol particles and black carbon particles. In the introduction and methodology sections we introduce and describe the importance of model aerosol particles mass concentrations, size distribution and mixing state on the remote sensing and climate related properties. We continue with distinguishing the black carbon particles, since those are the particles we focus on improving with the new aerosol optics model.

Later on we continue with a distinction between "aerosol particles" and "black carbon particles" when we in the model have "removed" all the aerosol components or only the black carbon components. This to study "aerosol radiative forcing" and "black carbon radiative forcing". We believe that it is this latter distinction we might have caused some confusion to the reader. However, due to the reformulation of "aerosols" (from one of the above comments) we believe the issue will become clearer. We also reformulated appropriately by adding "all aerosol components" instead of just "aerosol components" to p. 19 1. 528, p. 20 1. 551, 554-555 and 555-556, where we start to present the results for the aerosol radiative effects.

5. Please be specific as to what is compared, evaluated, mixed, etc.: is it the model, the simulation output, the size bins, the aerosol components, etc. For instance, "comparing the differences in the optics models to other sources of error" (p. 10739, l. 9) should be replaced by something like "comparing the differences between simulations with the two optics models to errors from other sources".

Thank you for being thorough with our text. We have now changed the text to "comparing the differences in the optics model output to other sources of error", see p. 4 l. 100-101.

6. I would prefer if you referred to table contents rather than to table rows (e.g., p. 10743, ll. 9 and 10; p. 10759, l. 21)

The first example on p. 10743 an Il. 9 in the discussion paper does refer to table contents and not to rows and columns. However, the second example does refer to rows, which we have changed in the revised edition to "in June (summer) for all four geographical locations and in December (winter) for the locations Poland and North Sea", see p. 32 and l. 785-786.

7. Please avoid generalizations. For instance, what is "required" (p. 10737, 1. 19) or "oversimplified" (e.g., p. 10760, 1. 4) certainly depends on the application.

We have looked into the occurrences of the more generalising words and we have changed the following; on p. 2 l. 53. Further, we have removed all instances of "over-simplified".

8. Please define technical terms at first use, e.g. external vs. internal mixture (defined later), backscattering coefficient (defined later), effective radius (not defined at all), etc.

Thank you for pointing this out. We have now restructured and rephrased the text slightly in order to avoid confusion with newly introduced terms. The effective radius is now defined with its corresponding equations together with its introduction at page 21 and line number 590. The external mixture assumption is now explained on p. 31. 68 where it is first mentioned. The internal mixing is now defined on p. 31. 81, and for the optical properties, we have added an appendix (Appendix A, B and C) describing how these parameters are derived in the model set-ups.

 Please keep in mind that an interested reader should in principle be able to reproduce your simulations. For instance, this requires complete information about the size distributions assumed for emitted particles.

Thank you for pointing this out, we have now added appropriate references and extra information. Regarding the different chemical transport versions, i.e. with and without aerosol microphysics, they are fully described in Andersson *et al.* (2013) and Andersson *et al.* (2007), respectively. Regarding the size distribution assumed for the emitted particles, they are written in Table 1 which now also includes the appropriate reference, Andersson *et al.* (2007). For the MATCH set-up with aerosol microphysics, the emission size distribution is directly refereed to with a specific citation, table 4 and figure 6 in Andersson *et al.* (2013), on p. 7 and l. 193-194.

10. I would appreciate if table and figure captions contained more, and more specific information.

We realise that too little information was put in the caption of many of the figures and some of the tables. Therefore, we have now added more specific information regarding table 3, 4 and 5 as well as figure 6, 7, 9, 10 and 11. More information regarding the time of the chosen events as well as the content of the table and plots has been added.

11. SALSA bins are not only distinguished by size, but also by composition. The term "size bin" therefore seems inappropriate, or at least incomplete.

Regarding the terminology we could have used the term "size composition bin", but we have chosen to follow the terminology from Andersson *et al.* (2015). See also the answer to minor comment nr. 13.

12. I find it difficult to understand Table 2 without some further expansion of Sect. 2.2.2. Some of the questions that come to mind are:

- Why are the smallest particles assumed to remain dry?
- Could you give an example for externally mixed PNHx particles?
- Why does PNOx appear only in internally mixed particles in one single size bin?

In order to make Table 2 more understandable, we have extended the section about the aerosol microphysics module SALSA, section 2.2.2, page 6-7. The specific questions are addressed below:

- Table 2, it is stated that the smallest size bin of particles that remains dry, i.e. that are not included in cloud droplet activation is size bin 3 with a radius of 9-25nm. As stated in McFiggans *et al.* (2006), particles with radius r < 40nm are not involved in cloud droplet activation. Our next size bin in MATCH-Salsa is 25-49nm and therefore includes non-dry particles.
- An example for externally mixed PNHx would be pure ammonium sulphate, (NH₄)₂SO₄.
- PNOx only appears in single size bin because the development of PNOx-chemistry in MATCH-SALSA is still in a very early stage. As noted in the specific comments, a more elaborated description of PNOx is desired; we have added a comment to section 2.2.2, p. 71. 190-195.
- 13. Furthermore, it should be mentioned in Sect. 2.2.2 that SALSA explicitly tracks particle number mixing ratios.

What MATCH-Salsa tracks is particle number concentrations, i.e. number of particles per volume of air per size interval. This has been added in the section 2.2.2 on p. 6 l. 186-187.

14. The term "bin" should be used consistently. In Sect. 2.3.1, for instance, three different words are used: "class", "bin", and "mode".

We understand that different terminology might confuse the reader. We have revised the text and consistently use the term "bin" now, which is consistent with the literature, e.g. Andersson *et al.* (2015) and Kokkola *et al.* (2008). However, we still do use the term "mode" when appropriate, i.e., when referring to a modal model, such as one using a log-normal size distribution. The changes occurred at: p. 61. 174, p. 81. 231 and 232, p. 91. 269, and p. 141. 371.

15. Sects. 2.3.1 and 2.3.2: It could be stated more clearly that the external mixture assumption is only used in conjunction with the MT aerosol module and the mixing state-resolved optics model is used both with the MT scheme and with SALSA.

In our new revised edition of the manuscript we have done as the reviewer suggested in major comment nr. 3, i.e., we added a fourth model set-up where we include the MATCH-SALSA CTM version together with the old EXT optics model (Salsa-EXT). This should be clear by the list on page 15 in the revised edition of the manuscript.

16. Please specify what is meant by the "total flux" (e.g., p. 10752, l. 13).

We have now changed the formulation "total flux" to what we actually defined in section 3.1, the "net radiative flux" on page 20 l. 563 and p. 21 l. 1.

17. Vague formulations like "slightly", "somewhat", "quite" should be avoided as much as possible. Quantitative information is preferred.

Thank you for pointing this out, we have now removed vague formulations in the text.

18. Sect. 3.1.2: If possible, please check (the inter-model differences in) ΔF_u as a function of wavelength to support your conclusion. Otherwise, please explain why the IR AOD effect should be "dominant" over the g effect. Furthermore, this seems to contradict what you state in the introduction to Sect. 3.1 (p. 10751, l. 16): "At other wavelengths (not shown) the optical properties behave similarly."

First, we removed the rather imprecise statement in the introduction. Secondly, we have added an appendix (Appendix E) with all the simulated wavelengths for the in depth analysed model set-ups (MT-EXT,MT-CGS and Salsa-CGS), which will support our statement in the section, where we discuss the possible impact on the black carbon radiative fluxes from different wavelengths and the two optics models (see p. 24 l. 706-708).

3 Specific comments

1. Correct citation format (possibly "citep" instead of "citet"): p. 10738, l. 18, p. 10740, l. 25, p. 10744, ll. 2 and 3, p. 10748, l. 7.

Thank you for noticing these errors, they are now corrected for with the right type of citation format. See p. 31. 79, p. 51. 140, p. 9, 1. 245 and 251, as well as p. 141. 377 in the revised edition of the manuscript.

2. p. 10739, l. 7: Please remove the word "rates".

The word "rates" has now been removed, see page 4 line number 99.

3. p. 10739, l. 15: Neither of the optics models is actually evaluated here, at least not in the sense that its output is compared to observations.

We follow the reviewer's advice and use a different formulation since "evaluation" is commonly used in the context of observations. Instead of "evaluation", we now use "comparison".

4. p. 10740, l. 5: Probably "fraction" is supposed to mean "component" here, otherwise please specify the "whole" of which a fraction is discussed here.

What is referred to as "The SIA fraction" is the secondary inorganic aerosol fraction of aerosol particles. This has now been made clearer in the text, see p. 5 l. 153.

5. p. 10741, l. 15: Please summarize briefly how the emissions inventory was generated.

We have added some more detailed information on how we unified the EMEP gridded emissions with the non-gridded BC and OC emissions given by Kupiainen and Klimont in order to obtain gridded OC and BC emissions. See p. 5 l. 163 - p. 6 l. 173.

6. Sect. 2.2.1 vs. Table 1: Are PPM (incl. BC and OC) assigned to three (Table) or four (text) size bins?

Other PPM, BC and OC are only assigned to three of the four available size bins. In the text we have stated that there are four available size bins, and in table 1 we list what is in each of the four size bins.

7. Sect. 2.2.1: The last two sentences should be moved up, where the corresponding modules are mentioned.

Thank you, this has now been edited for. The changes can be found on p. 5 l. 155-157.

8. p. 10742, l. 7: Please summarize briefly what the simplified PNOx description is.

We have added a brief clarification, stating that PNO_x in the current version is simply computed by the mass-transport model, and the mass is assigned to the SALSA-size bin No. 15. See page 7 and 1. 190-192.

9. Section 2.3.1: Please add a reference to Table 1, and add information on the hydrophilic/hydrophobic assumption for "other PPM".

We have added a reference to table 1, Andersson *et al.* (2007), but we are not sure what the reviewer means when asking for more "information on the hydrophilic/hydrophobic assumption for "other PPM". However, we realised that we need to clarify what "other PPM" means. This has been done in the section 2.3.1 together with the description of the gridded EMEP emissions. See page 6 l. 172-173.

10. p. 10741, l. 15: Please state briefly how the Gerber (1985) parametrization is applied.

We added the information that the parametrisation computes the wet radius as a function of dry radius, relative humidity, and temperature. See p. 91. 255-256.

11. p. 10744, l. 14: Please state briefly how the optical properties are interpolated onto intermediate water volume fractions.

They are interpolated linearly, which we have added to the text as well, see page 9, 1. 263.

12. Section 2.3: Please state clearly that optical properties are computed from the (effective) refractive indices.

This is only true for homogeneous internal mixtures. Section 2.3 makes some rather general observations on optics modelling. We have looked through the text and wrote "effective refractive index" wherever appropriate, but we retained "refractive index" whenever no effective-medium assumptions are involved.

13. Sect. 2.3.1, 3rd paragraph: I find this confusing. If I understand it correctly, you explain how to get from mass mixing ratios, which you simulate, to particle number mixing ratios (but not number densities). In case this is correct, please restructure the paragraph accordingly, and add information on how to convert particle mass to particle volume. Furthermore, as single scattering albedo and asymmetry parameter are properties of individual particles (in contrast to aerosol optical depth and backscattering coefficient), please clarify that the particle number concentration/size distribution is required here for the averaging, rather than for the computation of the per-particle properties. Finally, the terms "mean radii", and "variances" are used incorrectly here. Please either give mean radii and variances, or use the appropriate terms: "geometric mean radii" and "geometric standard deviations"

We have added two sections in an appendix (Appendix A and B) that explain in detail how to convert mass mixing ratio to number *density* (not number mixing ratio!), and how to compute ensemble-averaged optical properties. Further, we explain how to compute radiometric properties of the medium, such as optical depth and backscattering coefficient, from the ensemble-averaged optical properties of the particles. The whole averaging procedure is a bit tricky when working with truncated size modes, as we do in the old optics model. For this reason, we did not want to include all this in the text, as it would be a bit of a detour away from the more essential issues. However, the appendix now provides all the equations necessary for the interested

reader who wants to understand the details of the size-averaging process. We hope that this clarifies the reviewer's questions. We would also like to point out that single-scattering albedo and asymmetry parameter are not *exclusively* properties of individual particles, as the reviewer said. They can also be properties of a small *ensemble* of particles. To compute ensemble-averaged single-scattering albedos, asymmetry parameters, etc. the number density/size distribution is required just as much as for computing AOD and other radiometric properties of the medium. This should be clear now when reading the appendix. Finally, we have followed the reviewers advise and now use the terms "geometric mean radius" and "geometric mean standard deviation". See the whole paragraph starting on page 9 l. 267 and finishing on page 10 l. 281.

14. p. 10746, l. 2: Please specify how the averaging over particle orientations is achieved (e.g., analytically over all orientations, or using a sampling technique, etc.)

We have added a sentence specifying that the orientation-averaging has been done analytically, and we added a reference to the paper by Khlebtsov (1992), who first derived the analytic expressions for non-axisymmetric particles. See page 10 and line number 310.

15. p. 10746, l. 15: Please specify "size" (i.e., measured as what).

We have clarified this text by rewriting the sentence to "... were computed in the range of 100-500 nm (volume-equivalent radius)" on page 7 line number 198-201 in the revised edition of the manuscript.

16. p. 10747, l. 24: Please specify the core-shell partitioning parameter, and state whether this was taken from the given reference, or whether it was found in the same way. Furthermore, please comment on whether the parameter is/should be size- and/or composition-dependent.

The parameter was taken from the given reference. It is independent of size, volume-fraction, and optical parameter, but it does depend on wavelength. We have provided this information in the revised text (see page 13 and 14, l. 357-365 blue marked text) and added a table (table 3 on page 14) with the core-shell partitioning parameter as a function of wavelength.

17. Section 2.3.2, item 2: Parts of the third paragraph should be moved to item 3, or to a separate one, as item 2 deals with the treatment of internally mixed BC. Furthermore, it is again unclear here how "other PPM" is treated. Please also state whether BC is treated the same way both in the core and the (gray) shell.

To make this section more clear, we made a fourth item to the enumerated list containing the treatment of all internally mixed particles, except black carbon. See page 14 line number 379-380. We also added information that "other PPM" is interpreted as dust particles, see page 6 l. 172-173 and page 7 l. 211-212.

18. Section 2.3.2: As the optics model is the essential novelty of this study, I think the 28 wavelength bands, 37 discrete BC volume fractions, etc. should be given explicitly in this publication, maybe in an Appendix or in a Supplement. Please also state briefly how the interpolation is performed and add some mathematical formulas to the description of the size-averaging procedure (in the main text), so that it is easier to understand.

The 28 wavelength bands and the refractive indices of each aerosol component are given in an extra table in appendix D, page 37. The 37 discrete BC volume fractions are now given in the text, page 14 line number 383. The interpolation is performed linearly, and the size-averaging is now explained in the new appendix.

19. Section 2.4: As stated above, I think this is not an "evaluation" in a strict sense. Furthermore, the second paragraph and the first sentence of the third paragraph actually belong to the Results section rather than to the description of the method. Please also comment on how clouds might affect your results, either here, or in Sect. 3.1.

We replace "Evaluation" by "Methodology for comparing". We also moved the figure showing AOD over the model domain and the discussion of the figure to the results-section. We have only considered clear-sky situations. As we did not perform any calculations involving clouds, we refrain from speculating on the possible effects these and other complications would have on our results.

20. Sect. 3.1: Please mention that Fs and Fd are counted positive downward, and Fu is counted positive upward.

We have added to the definition on page 20 and line number 534-537 that F_s and F_d are accounted for with a forward direction downwards and F_u upwards.

- 21. Sect. 3.1.1: This section could be more concise. Some sentences could be removed completely (this also applies to Sects. 3.1.2 and 3.2). Other parts should be more precise, or even expanded (cf. Major comment on this section). Please clarify (at least) the following points:
 - The magnitude of ΔF_u would also increase with decreasing altitude in case the AOD per layer was constant.
 - Aerosol extinction does not result in the generation of diffuse flux. Actually, the opposite is the case: the processes that generate the diffuse flux are responsible for the extinction.
 - The difference in "convertible flux" (last sentence of first paragraph) actually increases with altitude; it is its magnitude that decreases.
 - When you discuss "mass" you actually refer to "mass mixing ratios". (This appears in similar form in Sect. 3.3 ("mass densities").)
 - "Number density"" is actually what you call in in Sect. 2.3.1. What you refer to here is more likely "number mixing ratio".
 - Although the assumption is unavoidable in the optics model in connection with the MT module, aerosol number mixing ratio does not necessarily increase with the mass mixing ratio in reality. (This appears again in Sect. 3.3, 2nd paragraph.)
 - Yes, this is, of course true. We meant to say that the *sharp* increase in the magnitude of ΔF_s is caused by the *sharp* increase by the AOD per layer. This is now clarified in the revised text, p. 201. 556-558.
 - We disagree. The process that generates diffuse flux is scattering. Extinction is just the generic term for the combined effect of scattering and absorption. We reformulated the text to preclude misinterpretations; it now reads "extinction in the form of scattering results in the generation of diffuse flux", see page 20 line number 558.
 - Yes, we changed the text accordingly.
 - Yes, we are referring to mass mixing rations, since we use the figure with vertical profiles of aerosol particle distribution with the unit ppb(m). In section 3.3 we now use "mass mixing ratio" instead of "mass densities", page 30 line number 764.

- No, by "number densities", we mean particle number concentration (see answer to specific comment 13).
- Correct. We have rephrased our statement on page 21 l. 598-599.

22. p. 10757, l. 8: Please specify which two locations

Thank you for pointing this out, we have now changed the sentence to "The black carbon forcing in Fig. 9 (Northern Italy) and (Mediterranean) 8 display different behaviours in radiative fluxes, comparing the EXT (blue) and CGS (red) model results.", see page 25 line number 714-716.

- 23. Sect. 3.2: As this still deals with the radiative forcing I would suggest to move it into Sect. 3.1. We have now moved section 3.2 to section 3.1.3.
- 24. Table 1: Please replace "wind blown" by "dust" or similar. We have changed "Wind blown" to "Dust" in Table 1.
- 25. Table 2: I suggest to sort the table by bin size rather than by composition.

Table 2 is sorted according to the number of size bins, which has different size ranges, mixing states and composition. In order to change the table and sort it by the size alone would be rather confusing, since most size ranges appears more than one time. with different mixing states and compositions. We would therefore like to keep Table 2 as it is.

26. Figures:

- I suggest to increase the font size (everything except the titles in Fig. 11).
- As you mainly discuss "TOA" effects, it might be advisable to reduce the number of vertically resolved plots in Figs. 5-8 to those that are actually required to understand the discussion.
- It looks as if the plotted data in Figs. 5-8 was somehow averaged/interpolated in/onto 1 km bins/levels. Please comment on this in the text.
- Why do the lowermost plots in Figs. 7 and 8 show differences between calculations with and without BC instead of just the BC values?
- Please comment on the increase in BC ΔF_u with altitude as simulated with SALSA in the lowest kilometre in Fig. 8.
- Maybe replace "Bulk" by "MT" in Figs. 9 and 10.
- As sulfate and nitrate are not discussed in the text, they can as well be left out of Fig. 10.
- Following the structure of the discussion, I recommend to swap Figs. 7/8 and 9/10.
- The font sizes have now been increased for all the figures (except the titles). The font weight has also been changed to "bold" to make it more readable.
- We believe that all the vertically resolved figures (Fig. 5-8) are helpful for understanding our analysis, so we would prefer to not erase any of them.
- Fig. 5-8 show the radiative fluxes together with the corresponding optical properties aerosol optical depth, single scattering albedo and the asymmetry parameter. The radiative transfer model we have used operates on these fixed levels. The optical properties have therefore been interpolated onto these levels. This is now mentioned in the text, see page 17 line number 466.

- The lowermost plots in Figs. 7 and 8 *do* show just the BC-values, but it is the differences between the solid and dashed lines one should focus on, because this is what we need in order to understand the forcing of BC. This was different for the total aerosol forcing in Figs. 5 and 6: to understand the total forcing of aerosols, we need to look at just the optical properties of aerosols. (In the absence of aerosols, the optical properties of aerosols are, trivially, zero!)
- We have thought about this, but did not find a satisfactory explanation for the increase in ΔF_u with altitude over the lowest kilometre in Fig. 8.
- We have now replaced the occurrences "Bulk" with "MT" for consistency.
- Sulphate is however mentioned in the text, whereas nitrate is not. Therefore we agree with the reviewer to remove nitrate form the figure showing the aerosol mass mixing ratios.
- We now follow the order of the discussion and have placed the figures of the effective radius and the profiles of aerosol mass mixing ratios before the black carbon forcing discussion.

4 Typos

```
• p. 10736, l. 19: effect \rightarrow affect
• p. 10739, l. 14 and l. 15: model \rightarrow models
• p. 10740, l. 7: hydrophillic \rightarrow hydrophilic
• p. 10745, l. 16: was \rightarrow is
• p. 10745, l. 21: calculation \rightarrow calculations
• p. 10751, l. 11: are \rightarrow is
• p. 10751, l. 17: 6 \rightarrow 5
• p. 10752, l. 4: Algier \rightarrow Algiers
• p. 10752, l. 17: at higher altitudes \rightarrow [delete]
• p. 10752, l. 24: then \rightarrow than
• p. 10757, l. 1: downdwelling \rightarrow downwelling
• p. 10757, l. 7: a \rightarrow [delete]
• p. 10760, l. 3: differences \rightarrow difference
• p. 10761, l. 4: an \rightarrow and
• p. 10762, l. 4: extend \rightarrow extent
• p. 10762, l. 9: acknowledge \rightarrow acknowledges
• p. 10764, l. 4: Nousainen \rightarrow Nousiainen
```

We have gone through the typos and corrected them accordingly.

• Table 3: Polen → Poland

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Author response regarding the manuscript 'Coupling aerosol optics to the chemical transport model MATCH (v5.5.0) and aerosol dynamics module SALSA (v1)', to the reviewer 2

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March 24, 2016

Thank you for taking your time to review our paper. Your comments and feedback has helped us improving our paper. Below you will find answers to each of the comments we received, where the your comments are marked with blue and our answers below in black.

1 General comments

- I had the general impression that the authors were crediting substantially the author's work, while a larger relevant literature exists. I recommend the authors to cite more independent work from other research scientists in the field, as it would strength the manuscript itself and the external perception.
 - Since this manuscript is not a review paper, it was not our ambition to provide a complete list of references on black-carbon optics. Rather, this manuscript is a model-development paper, so it is both natural and necessary that we cite those publications that have been relevant to the validation and development of our aerosol-optics model. The present manuscript is, indeed, only the tip-of-the-iceberg of a long-term research project on black carbon, which involved a substantial amount of development work. The citations of our own work in the manuscript reflect this fact. For instance, we have performed exactly the kind of validation studies that the reviewer mentions under specific comments 1 (dependency of MAC and SSA on aggregate size, monomer size and number), and more (dependency of MAC and SSA on refractive index, which is even more important for MAC than the monomer size). However, we agree, of course, that it may be helpful for the reader if we include a more complete reference list, so we added additional citations (see specific comments 1 below).
- 2. Many of the plots are not easy to read, neither in the screen or in a printed version. I would recommend to make labels of the axis with a larger font size. I would also suggest to add a description of the plots in the caption, so a reader does not need to go back a forward in the text to find relevant details. The manuscript does not provide any code to perform calculations, which seems strongly suggested by the journal.

We agree that the quality of the plots was in need of improvements. The plots have been revised along the reviewer's suggestions. We also added a section on code-availability, see page 34 and section 5.

2 Specific comments

1. Page 10745 Please, state T-matrix estimated mass absorption coefficients (MAC) values and how close are to Bond and Bergstrom (2006) recommended values, in addition to SSA values.

The choose of monomers radius of 25nm might be legitimate, nerveless I would recommend a statement addressing the variability in the monomers radius 15-25 nm. Different monomer radii exist due to differences in sources, which mainly depend on burning materials (i.e. Weztner et al., 2003, China et al., 2014., Chakrabarty et al., 2014)

Optical properties of BC aggregates may vary depending on assumptions of BC monomers radius. One key manuscript addressing the issue at one wavelength is Liu et al., 2008, other papers based on observational driven constrains of BC particle aggregates may be worth to be cited, i.e., Scarnato et al., 2013, 2015. The choose of a Df=1.8 can be considered consistent with semi aged BC aggregates, i.e. China et al., 2015, China et al., 2014.

A good fitting of observed values of MAC and SSA with T-matrix simulations reside on the choice of aggregate physical constrains (aggregate size, monomer size and number, other than fractal dimension).

The MAC values of our aggregate model have been validated in detail in Kahnert and Devasthale (2011), which we have cited.

We added, as suggested, that the monomer radius can vary between 10–25 nm with a citation of Bond and Bergstrom (2006) (see page 16 line number 12-13). However, we emphasise that our choice of a=25 nm has been validated in the AST-paper by Kahnert (2010), as explained in the text; it gives the best agreement with measurements for the SSA. This is actually not surprising. It is much more surprising that in the literature authors often seem to tacitly assume that the best choice of the monomer radius a would be some mean value, $\int n(a)ada$, over a normalised monomer-size distribution n(a). We cannot think of any good arguments to support this conjecture. Rather, the monomers are tiny compared to the wavelength, so if anything they should behave similarly to Rayleigh scatterers, for which the absorption cross section scales as a^3 . Thus, in our own humble opinion a meaningful first-guess for an effective radius would be $a_{\rm eff} = \int n(a)a^4da/\int n(a)a^3da$, the ratio of the fourth to the third moment of the size-distribution. I.e., one should weigh $n(a) \cdot a$ with the absorption cross section, which is proportional to a^3 . If doing so, we would expect that the effective monomer radius that gives the most realistic estimates of the optical properties should lie close to the upper end of the size scale 10–25 nm, which is just what we assume.

We also added some of the citations suggested by the reviewer (China et al. (2014) at p. 10 l. 294, Chakrabarty et al. (2014) at p. 10 l. 293, Scarnato et al. (2013) and Scarnato et al. (2015) at p. 1 l. 320). However, we would like to point out that the manuscript by Liu et al. (2008), although highly cited, covered an enormous parameter space that far exceeds the range relevant for atmospheric black carbon (e.g. Df as low as 1.25 and as high as 3; as well as quite unrealistic refractive indices, 2+i and 1.75+0.5i). The AST-paper by Kahnert (2010) considered a considerably more constrained and realistic range of parameters and was much more relevant for the validation of our aerosol-optics model.

We agree that Df=1.8 is typical for rather fresh or "semi-aged" aggregates. Older aggregates may have higher fractal dimensions. However, exactly this question has been discussed in the ACP-paper by Kahnert and Devasthale (2011); this study showed that, despite being rather low, the value of Df=1.8 gives a better agreement of modelled and measured MAC values than Df-values that may be more representative for aged black carbon.

2. The paper mainly focuses on the impact of various treatments of optical modules to chemical transport models. Secondly, the paper discusses how various optical treatments can impact estimates of backscattering coefficients and Angstrom exponent. The authors should consider, at least for a site, to make comparison with observational data (i.e., A comparison between space born lidar backscattering values with those predicted).

In principle we very much agree with the reviewer that comparison with observations are extremely valuable. However, doing comparison with observations is an all-or-nothing business. It would not be very illuminating to simply show a plot of model results and observations. Chemical transport models are notoriously plagued by a large number of errors and biases, such as the emission estimates, deposition velocities, chemical reaction rates, meteorological input data (wind fields, precipitation rates, boundary-layer processes, etc.), land use data, parameterisations of physical processes, such as nucleation, condensation, coagulation, and water-uptake, to name just a few examples. All of these sources of error would effect a comparison of modelled optical properties with measurements. A meaningful comparison would not be possible without performing a comprehensive analysis of *all* involved sources of error. This would completely change the focus of the paper. Our intention with this paper is to educate the chemical-transport modelling community, and raise some awareness for the importance of aerosol optics modelling. This message would have become completely buried if we had set up this study as a CTM validation study.

3. Page 10756: Can you please provide an explanation for the statement: "Over the Mediterranean (Fig.6), the EXT and CGS model have almost identical AOD profiles in the green part of the spectrum. However, at longer wavelengths (not shown) EXT predicts substantially higher AOD values than CGS"

This statement refers to a difference in the AOD at different wavelengths. In the paper, we only show AOD for the green spectrum, i.e. 500(EXT)/532(CGS) nm, but in order to explain certain behaviours, we had to look closer at some of the other wavelengths as well. In the revised edition, we have now added an appendix (Appendix E) where we include figures with optical properties for the other wavelengths. We also added in the text a reference to the appropriate figure that shows ADO at longer wavelengths; this should help to make this statement clearer.

4. Can you please provide an order of magnitude for the statement? "TOA net flux in EXT as compared to the CGS model. Note that the differences in SSA between EXT and CGS are fairly small, while the differences in g are rather large".

The reviewer has a good point; this statement needs to be more precise and quantitative. We have changed the text accordingly and provided concrete numbers for the differences in the asymmetry parameter and SSA between the two models. See page 24 line number 703-706.

5. It might be useful to strength the perception of the paper, to add when possible in the abstract and conclusion a percentage (or order of magnitude) of the impact of different optical modules.

An excellent point. We have taken some of the most important results from the new table 4 (which shows spatio-temporally averaged model differences) and cited them in the revised abstract and conclusions. See the abstract on page 2 line number 24-27 and the Conclusion at page 33 line number 823-827.

Author response regarding the manuscript 'Coupling aerosol optics to the chemical transport model MATCH (v5.5.0) and aerosol dynamics module SALSA (v1)', to the short comment by A. Kerkweg

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March 24, 2016

Thank you for your comment and helping us full-fill the requirements of papers in GMD. We have included your comment in blue here in this document followed your answer in black.

1 Short comment

Dear authors,

In my role as Executive editor of GMD, I would like to bring to your attention our Editorial version 1.1:

http://www.geosci-model-dev.net/8/3487/2015/gmd-8-3487-2015.html This highlights some requirements of papers published in GMD, which is also available on the GMD website in the "Manuscript Types" section:

http://www.geoscientific-model-development.net/submission/manuscript_
types.html

In particular, please note that for your paper, the following requirement has not been met in the Discussions paper:

 Inclusion of Code and/or data availability sections is mandatory for all papers and should be located at the end of the article, after the conclusions, and before any appendices or acknowledgments.

You included the required information at the end of the conclusions sections. Please make the last sentence of your conclusions to an independent section with the title "Code and data availability" upon the revision of your article.

Yours, Astrid Kerkweg

Thank you for pointing this out, we have now added a section about the code availability after the conclusion section. The section is found on page 34 in the attached revised manuscript containing the differences between the previous version and the updated version of the manuscript.

Coupling aerosol optics to the chemical transport model MATCH (v5.5.0) and aerosol dynamics microphysics module SALSA (v1)

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Abstract. Modelling aerosol optical properties is a notoriously difficult task due to the particles' complex morphologies and compositions. Yet aerosols aerosol particles and their optical properties are important for Earth system chemistry-climate modelling and remote sensing applications. Operational optics models often make drastic and non-realistic approximations regarding morphological properties, which can introduce errors. In this study a new aerosol optics aerosol-optics model is implemented, in which more realistic morphologies and mixing states are assumed, especially for black carbon aerosolsparticles. The model includes both external and internal mixing of all chemical species, it treats externally mixed externally-mixed black carbon as fractal aggregates, and it accounts for inhomogeneous internal mixing of black carbon by use of a novel "core-grey shell core-grey-shell" model. Simulated results of radiative fluxes, aerosol optical properties, such as aerosol optical depth, backscattering coefficients and the Angström exponent from, as well as radiative fluxes computed with the new optics model are compared with results from another model simulating an older optics-model version that treats all particles as externally mixed homogeneous spheres. To gauge the impact on the optical properties from the new optics model, the known and important effects from using aerosol dynamics serves For comparison, we perform computations with two different model-versions, one that accounts for aerosol-microphysical processes, and another one that entire neglects these processes. Since it is well understood that aerosol microphysics has a profound impact on aerosol mass- and number-concentrations, their size-distribution, and their size-dependent chemical composition (which, in turn, strongly impact their optical properties), these additional model-runs can serve as a reference against which we can gauge the significance of the morphological assumptions in the optics model. The results show that using a more detailed description of particle morphology and mixing states influences the optical properties to the same

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degree as aerosol dynamics. a degree that is on the same order of magnitude as the corresponding effects of aerosol-microphysical processes. For instance, the aerosol optical depth computed with the two optics models differs over the optical spectrum by -25-18 %, while corresponding differences caused by the inclusion or omission of aerosol microphysics range between -50-37 %. The corresponding differences in the backscattering coefficient are -8-99 % and -47-28 %, respectively. This is an important finding suggesting that over-simplified simple optics models coupled to a chemical transport model can introduce considerable errors; this can strongly effect affect simulations of radiative fluxes in Earth-system chemistry-climate models, and it can compromise the use of remote sensing observations of aerosols aerosol particles in model evaluations and chemical data assimilation.

1 Introduction

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Aerosol optics Aerosol-optics models are employed in large-scale chemical transport models (CTMs) in mainly two contexts, namely, in Earth-system climate modelling chemistry-climate modelling (CCM), and in conjunction with remote sensing observations. In Earth-system modelling a CCM one couples a CTM to an atmosphere-ocean general circulation model (GCM). One purpose is to account for the dynamic effects of aerosols aerosol particles on cloud microphysics. Another is to obtain a better description of the direct effect of aerosols aerosol particles and radiatively active trace gases on the radiative balance. The aerosol optics aerosol-optics model provides a link that converts the aerosol fields delivered by the CTM to the aerosol optical properties that are required as input to the radiative transfer model, with which one computes the radiative energy budget. In remote sensing applications one is faced with the obstacle that the aerosol concentration fields computed with a CTM are not directly comparable to the radiometric quantities that are observed with remote sensing instruments. The aerosol optics aerosol-optics model provides the observation operator that maps the CTM output to radiometric variables that can be compared to satellite observations or satellite retrieval products. This allows us to either employ satellite observations for evaluating CTM model results, or to assimilate satellite data into a CTM-based air-quality forecasting system. It is clear that the aerosol optics aerosol-optics model has a pivotal role in these kinds of applications. It may constitute an additional source of error that could compromise the reliability of Earth-system climate models CCMs, impair the reliability of CTM evaluations, or degrade chemical data assimilation results. It is, therefore, important to better understand this potential source of error, quantify its possible impact on model predictions of aerosol radiometric quantities, and assess the level of morphological detail that is required in aerosol optics might be required in aerosol-optics models coupled to CTMs.

A main difficulty is that aerosols aerosol particles in nature can have a high degree of morphological complexity. For instance, mineral dust particles can have irregular shape, small-scale surface roughness, and inhomogeneous mineralogical composition (e.g. Nousiainen, 2009). Black carbon

aerosols are fractal aggregates particles suspended in air have fractal-aggregate shapes (e.g. Jones, 2006) that can be coated by weakly absorbing liquid-phase components that condense onto the aggregates as they age in the atmosphere (e.g. Adachi and Buseck, 2008). Volcanic ash particles are composed of crustal material in which multiple air vesicles may have been trapped during the generation of the particles. In aerosol optics aerosol-optics models one has to make a choice what level of morphological detail is necessary and affordable. A detailed discussion of this question can be found in Kahnert et al. (2014).

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In environmental modelling practical and computational constraints often force us to invoke drastically simplifying assumptions about aerosol morphology. For instance, one frequently computes aerosol optical properties based on the assumption that all chemical aerosol components are contained in separate particles (externally mixed), and that each such particle can be approximated as a homogeneous sphere. As pointed out in Kahnert (2008); Benedetti et al. (2009), this approach is highly attractive from a practical point of view, because the aerosol optical observation operators, which map mixing ratios to radiometric properties, become linear functions of the mixing ratios of the different chemical species. A linearisation of the observation operator is a prerequisite for most of the commonly used data assimilation data-assimilation methodologies, such as the variational method (e.g. Kahnert, 2008; Benedetti et al., 2009). However, such approximations can also introduce substantial errors. In the remote sensing remote-sensing community awareness for this problem has been growing over the past 1-2 decades. As a result, one has developed retrieval methods for desert dust aerosols aerosol particles that are based on spheroidal model particles (e.g. Dubovik et al., 2006), which can mimic the optical properties of mineral dust particles better than homogeneous spheres Kahnert (2004); Nousiainen et al. (2006) (Kahnert, 2004; Nousiainen et al., 2006). In chemical data assimilation, the problem is still treated rather negligently. A few assimilation studies account for internal mixing (where several aerosol components can be contained within one particle) of different chemical components (e.g. Saide et al., 2013). But the particles are still assumed to be perfectly homogeneous spheres. To the best of our knowledge there are currently no aerosol optical observation operators in chemical transport models that take complex morphological properties of aerosols such as non-sphericity aerosol particles such as non-sphericity or inhomogeneous internal structure into account.

This study describes the coupling of two different aerosol optics aerosol-optics models to a regional CTM. One optics model is based on the simple external-mixture and homogeneous-sphere approximations. The second model takes both external and internal mixing of aerosol components into account. Also, it employs morphologically more realistic models for black carbon aerosolsparticles. Although black carbon contributes, on average, only some 5 % to the mass mixing ratio of particulate matter over Europe, it can have a significant global radiative warming effect. Previous theoretical studies on the optical properties of black carbon aerosols black-carbon particles suggest that the use of homogeneous sphere models can introduce substantial errors in the absorption cross section

and single scattering albedo of such particles (e.g. Kahnert, 2010a; Kahnert et al., 2013). Also, the largest mixing-state sensitivity in both regional and global radiative fluxes comes from black carbon according to Klingmüller et al. (2014).

The main goal of this study is to assess the impact of aerosol morphology and mixing state on radiometric quantities and radiative forcing rates—simulated with a chemical transport model. To this end we compare the two optics models, and we gauge the significance of morphology by comparing the differences in the optics models model output to other sources of error. As a gauge we use the impact of including or omitting aerosol dynamic microphysical processes; this provides us with a reference which is generally agreed to have a significant effect on aerosol transport models (Andersson et al., 2015; Kokkola et al., 2008).

The CTM, its aerosol dynamics microphysic and mass transport modulesset-ups, and the aerosol optics model aerosol-optics models are described in Sect. 2. There we also explain the methodology we employ for evaluation comparison of the optics model models. In Sect. 3 we present and discuss computational results for selected cases and for several radiative and optical parameters. Concluding remarks are given in Sect. 4.

110 2 Model description and methods

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2.1 General considerations and terminology

Aerosol particles typically originate from different emission sources, such as sea-salt particles coming from marine sources, wind-blown dust from dry land surfaces, volcanic ash from magmatic or phreatomagmatic eruptions, or black carbon produced during combustion of fossil fuel, biofuel, or biomass. During atmospheric transport particles from different sources can be mixed, resulting in heterogeneous aerosol populations consisting of particles of different morphologies, sizes, and chemical composition. A mixture in which different chemical species are contained in separate particles is referred to as an external mixture. On the other hand, aerosol dynamic processes, such as nucleation, condensation, and coagulation, give rise to the formation and growth of secondary particles from precursor gases, as well as to the condensation of precursor gases onto existing primary particles. These processes result in particles in which several chemical species are mixed with each other in one and the same particle. Such a population is referred to as an internal mixture. There are two types of internal mixtures. If, e.g., hydrophillic hydrophillic liquid-phase components mix with each other, one can obtain a homogeneous internal mixture of different chemical species. On the other hand, condensation of gas-phase species onto non-soluble primary particles, or cloud processing of aerosols aerosol particles can result in liquid-phase material coating a solid core of, e.g., mineral dust or black carbon. We refer to the latter as an inhomogeneous internal mixture. Aerosol populations in nature are often both externally and internally mixed, i.e., they contain particles that

are composed of a single chemical species as well as other particles that are composed of different chemical species, which can be homogeneously or inhomogeneously internally mixed.

Aerosol optical properties are strongly dependent on not only the size and chemical composition, but also on the mixing state, shape, and internal structure of particles. Therefore, before explaining the aerosol optics aerosol-optics model, we first need to briefly describe the kind of information that can be provided by the aerosol transport model. In particular, we need to understand the level of detail with which the size distribution, size-dependent chemical composition, and the mixing state of the aerosols aerosol particles can be computed in a large-scale model.

2.2 Aerosol transport modelling with MATCH

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As a regional model we employ the Multiple-scale Atmospheric Transport and CHemistry modelling system (MATCH)Andersson et al. (2007). an offline Eulerian model developed by the Swedish Meteorological and Hydrological Institute (Andersson et al., 2007). For this study we have set up the MATCH model over the European domain with a $0.4 \times 0.4^{\circ}$ horizontal resolution and a rotated latitude-longitude grid, covering about 34° longitude and 42° latitude. The model has 40 vertical η layers with varying thickness depending on the topography, and it extends up to about 13 hPa. The meteorological input comes from the numerical weather-prediction model HIRLAM (HIgh-Resolution Limited Area Model) (Unden et al., 2002).

The MATCH model allows us to choose between two aerosol model versions, a simpler mass transport model, and a more sophisticated aerosol dynamic transport model.

2.2.1 Mass transport model

A simple version of the CTM MATCH, which we refer to as the "mass transport mass-transport model", neglects all aerosol dynamic processes. It contains a photochemistry model that computes mass concentrations of secondary inorganic aerosols (SIA), which are formed from precursor gases. The SIA fraction of aerosol particles consists of ammonium sulphate ((NH₄)₂SO₄), ammonium nitrate (NH₄NO₃), other particulate sulphates (PSO_x), and other particulate nitrates (PNO_x). The mass transport model further contains a seasalt sea-salt module that computes NaCl emissions based on the parametrisations described in Mårtensson et al. (2003); Monahan et al. (1986). He More details on the MATCH photochemistry model can be found in Robertson et al. (1999); Andersson et al. (2007); the MATCH sea-salt model is described in Foltescu et al. (2005). The mass transport model also contains a simple wind-blown dust model and a module for transport of primary particulate matter (PPM), i.e., aerosols other than seasalt and wind-blown aerosol particles other than sea-salt and wind-blown dust that are emitted as particles, rather than being formed from gas precursors. The size bins in the PPM model are flexible. In the current model set-up the sea-salt sea-salt and PPM models were run for four size bins as shown in Table ??1. We used gridded EMEP PPM emission data for the year 2007, and, based on those and on Kupiainen and Klimont (2004, 2007), we

Table 1. Size bins (characterised by the radius r) and chemical species in the MATCH mass transport model (Andersson et al., 2007). The labels "p" and "s" refer to primary emitted particles and secondary particles generated from gas precursors.

size				wind-	other				other	other
bin	r (nm)	OC	ВС	blown Dust	PPM	NaCl	$(NH_4)_2SO_4$	NH_4NO_3	PSO_x	PNO_x
1	10-50	p	p		p	p	S	S	s	s
2	50-500	p	p		p	p	S	S	s	s
3	500-1250					p	S	S	S	S
4	1250-5000	p	p	p	p	p	S	S	S	S

generated gridded emission data for black earbon in conjunction with black-carbon (BC), organic earbon-and organic-carbon (OC), and all other PPM. (Here, OC refers to the mass of all organic matter, not just the mass of carbon atoms in organic compounds.) The primary particle emission data by Kupiainen and Klimont (2004, 2007). The latter provide BC and OC emissions per country and emission sector. We distributed these among the grid cells in the model domain according to the EMEP PPM gridded emissions. Thus, the BC and OC emissions vary among grid cells in accordance with the EMEP PPM emissions, while the sum of all BC and OC emissions over all grid cells per country and emission sector agrees with the corresponding BC and OC emissions, respectively, reported in Kupiainen and Klimont (2004, 2007). The remaining emissions (PPM-BC-OC) in each grid cell are interpreted as dust particles. The primary-particle emissions are distributed among the four size classes size bins; during atmospheric transport they remain chemically and dynamically inert in the model. Thus no chemical transformation, mixing processes with other compounds, or other size transformation size-transformation processes are included in the model. The SIA components are given as total mass concentrations without any information about their size distribution. In the optics model a fixed size distribution is assumed to assign the total SIA mass to the four size bins. Water adsorption by particles is computed in the optics model as described in Sect. 2.3.1. More details on the MATCH photochemistry model can be found in Robertson et al. (1999); Andersson et al. (2007). The MATCH seasalt model is described in Foltescu et al. (2005).

2.2.2 Aerosol dynamics modelmicrophysics module - SALSA

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A more realistic description of particles can be achieved by accounting for aerosol dynamic microphysical processes. To this end the Sectional Aerosol module for Large Scale Applications (SALSA) (Kokkola et al., 2008) has recently been coupled to the MATCH photochemistry model (Andersson et al., 2015). This model tracks mass concentrations of different species per size bin, and particle number concentrations. Thus, it provides size-dependent composition and mixing state of aerosol particles. The description of PNO_x , wind-blown dust, and secondary organic aerosols (SOA) is still under development in an early development stage in MATCH-SALSA. A simplified In the current version,

PNO_x description has been included in the model version employed here, while wind-blown is simply computed according to the same photochemistry-scheme as in the mass-transport model, and the PNO_x mass is assigned to size bin 15 (see Tab. 2). Wind-blown dust and SOA are absent . The number and range of size bins is flexible in SALSA. Table ?? in the present model version. The size distributions for the emitted particles can be found in table 4 and figure 6 of Andersson et al. (2013).

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Table 2 shows the current model set-up -

with the number and range of the size bins. As is evident from the this table, MATCH-SALSA accounts for both internally and externally mixed aerosols, aerosol particles. In total, there are 20 different size bins in MATCH-SALSA, each one of them representing a particle size range (volume-equivalent radius, r), mixing state, and composition. Some size bins have the same size range, but different mixing states and/or compositions. For instance, size bins 12, 15, and 18 describe the same size range (350–873350-873 nm), but different internal mixtures of various species. Similarly, bins 4 and 8 have the same size range (25–49 nm), but one describes an internal mixture, the other an external mixture of aerosol species.

Note that water is not directly calculated as a prognostic variable in MATCH-SALSA. Rather, it is a diagnostic variable computed in the MATCH-optics model as explained in Sect. 2.3.2. The table merely indicates which size bins are assumed in the optics model to be internally mixed with adsorbed water. A more detailed description of the MATCH-SALSA model can be found in Andersson et al. (2015).

210 2.3 Aerosol optics modelling

As in the mass-transport model, "other PPM", i.e. primary particles other than BC and OC, are interpreted as dust particles. Note that water is not directly calculated as a prognostic variable in MATCH-SALSA. Rather, it is a diagnostic variable computed in the MATCH-optics model as explained in Sect. 2.3.2. The table merely indicates which size bins are assumed in the optics model to be internally mixed with adsorbed water. A more detailed description of the MATCH-SALSA model can be found in Andersson et al. (2015).

Acrosol optics

2.3 Aerosol-optics modelling

Aerosol-optics models coupled to a CTM have to make consistent use of the information provided by the CTM, while invoking assumptions on optically relevant parameters that are not provided by the CTM. The parameters that influence the particles' optical properties are

- the aerosol size distribution;
- the refractive index of the materials of which the aerosols aerosol particles are composed;

Table 2. Size bins and chemical species in the MATCH-SALSA aerosol dynamic microphysical transport model. An "x" marks that the species is present in a particular size bin.

size		mixing			other				
bin	r (nm)	state	OC	BC	PPM	NaCl	PSO_x	PNO_x	PNH_x
1	1.5-3.8	internal	х				x		X
2	3.8-9.8	internal	x				X		X
3	9.8-25	internal	X				X		X
4	25-49	internal+ H_2O	X	X	X	X	X		X
5	49–96	internal+ H_2O	X	X	X	X	X		X
6	96-187	internal+ H_2O	X	X	X	X	X		X
7	187-350	internal+ H_2O	X	X	X	X	X		X
8	25-49	external	X	X			X		X
9	49–96	external	X	X			X		X
10	96–187	external	X	X			X		X
11	187-350	external	X	X	X		X		X
12	350-873	NaCl+ $\rm H_2O$				X			
13	873-2090	NaCl+ $\rm H_2O$				X			
14	2090-5000	$NaCl+H_2O$				X			
15	350-873	internal+ H_2O	X	X	X		X	X	X
16	873-2090	internal+ H_2O	X		X		X		X
17	2090-5000	internal+ H_2O	X		X		X		X
18	350-873	internal+ $\mathrm{H}_2\mathrm{O}$			X		X		X
19	873-2090	internal+ $\mathrm{H}_2\mathrm{O}$			X		X		X
20	2090-5000	internal+H2O			X		X		X

⁻ the morphology of the particles.

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225 *Morphology* refers to both the overall shape of the particle, and, in case of inhomogeneously mixed particles, the variation of the refractive index inside the particle.

The information provided by the CTM depends on the level of detail in the process descriptions. In the MATCH mass transport model, we have size information for the primary particles, but only the total mass for secondary inorganic aerosols. Thus we have to invoke assumptions about the size distribution of these particles. The MATCH optics models in conjunction with the MATCH mass transport model assume that 10% of the SIA aerosol mass are in the smallest size elass bin (see Table ??1), 60% in the second, 20% in the third, and 10% in the fourth size elass bin. Also, the mass transport model lacks any information about the mixing state of the particles. We therefore have to invoke appropriate assumptions on whether the aerosols aerosol particles are externally or internally mixed. Both the mass transport model and MATCH-SALSA lack information on whether the internally mixed particles are homogeneous or inhomogeneous. Also, neither model provides any information on the shape of the particles. The refractive index of the different chemical components indices of each chemical component in the aerosol phase and their spectral variation is given in are

listed in Appendix D. They can also be found in Fig. 4 of in Kahnert (2010a). That reference also contains detailed information about the different literature sources from which the refractive indices are taken.

2.3.1 Optics model for externally mixed aerosols aerosol particles

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The simplest conceivable optics model assumes that all particles are homogeneous spheres, and that all chemical species are each in separate particles, i.e., externally mixed. As explained in Kahnert (2008) (Kahnert, 2008), the external-mixture assumption results in a linear relation between the mass mixing ratios and the optical properties. Owing to the linearity, this model is particularly attractive for data assimilation applications (e.g. Benedetti et al., 2009), which require linearised observation operators. However, this is also the crudest possible optics model, as it neglects both the effect of internal mixing and of particle morphology on optical properties.

The external-mixture model is implemented in the MATCH mass transport model, where it is primarily being used in the MATCH 3DVAR data assimilation system Kahnert (2008) (Kahnert, 2008). Optical properties are pre-computed for twelve wavelength bands ranging from the UV-C to the mid-IR. Dust and black carbon are assumed to be hydrophobic, while sea salt, OC, and SIA components can each mix internally with water. The water volume fraction depends on temperature and humidity; it is computed by use of the parametrisation given in Gerber (1985), which computes the particle's wet-radius as a function of dry-radius, relative humidity, and temperature. The aerosol/water mixture is assumed to be homogeneous. The dielectric properties of a homogeneous mixture of two or more components are described by a complex effective refractive index $m_{\rm eff}$, which is usually computed by effective medium theory (EMT) (although chemical transport modellers often use simple volume mixing rules, most likely because EMTs are not commonly known in that field). We use Bruggemann's EMT (Bruggemann, 1935). More information of EMT is given in Appendix C. Optical properties are pre-computed for eleven water volume fractions between 0 and 0.98; for intermediate volume fractions the optical properties are linearly interpolated. The optical properties contained in the database are the extinction cross section C_{ext} , the scattering cross section C_{sca} , the value of the phase function in the exact backscattering direction $p(180^{\circ})$, and the asymmetry parameter g.

As explained in Kahnert (2008), size-averaged optical properties are pre-computed by averaging over a log-normal size distribution $n_i(r) = N_i/(\sqrt{2\pi}r\ln\sigma_i)\exp[-\ln^2(r/r_i)/(2\ln^2\sigma_i)]$ $n_i(r) = N_i^0/(\sqrt{2\pi}r\ln\sigma_i)\exp[-\ln^2(r/R_i)/(2\ln^2\sigma_i)]$ for each size class bin i, where N_i represents N_i^0 relates to the number density of particles in size bin i, r denotes the particle radius, $r_1 = 0.022$ $R_1 = 0.022 \, \mu m$, $r_2 = 0.158 \, R_2 = 0.158 \, \mu m$, $r_3 = 0.791 \, R_3 = 0.791 \, \mu m$, $r_4 = 2.5 \, R_4 = 2.5 \, \mu m$ are the geometric mean radii in each size mode, and the variances geometric standard deviation $\sigma_1 = \sigma_3 = \sigma_4 = 1.8$, $\sigma_2 = 1.5$ are based on measurements in Neusüß et al. (2002). The volume per size bin ean be obtained by integrating $(4/3)\pi r^3 n_i(r)$ over the ith size bin interval; this can be used for

converting Appendix A provides detailed explanations of how to convert mass mixing ratios into number densities in each size bin, which, in turn, allows us to compute radiative properties computed in the model into particle number densities, how these are used in computing size-averaged optical properties, and how to obtain radiometric properties of the atmosphere, such as aerosol optical depth (AOD) , single-scattering albedo (SSA), asymmetry parameter g, and or backscattering coefficient
 β_{bak}in each atmospheric grid cell in MATCH – see Kahnert (2008) for details, from the particles' optical properties and from the MATCH aerosol fields.

2.3.2 Optics model for aerosols aerosol particles of different mixing states

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The new MATCH-optics model accounts for both internally and externally mixed aerosolsaerosol particles, and it contains both homogeneously and inhomogeneously mixed aerosolsaerosol particles. Different shapes and morphologies are assumed for different types of particles.

1. Pure, externally mixed black carbon aerosols particles are assumed to have a fractal aggregate morphology as shown in Fig. 1. The fractal morphology can be described by the statistical scaling law $N_{\rm s} = k_0 (R_{\rm g}/a)^{D_{\rm f}}$, where $N_{\rm s}$ denotes the number of spherical monomers in the aggregate, $D_{\rm f}$ and k_0 are the fractal dimension and fractal prefactor, a is the monomer radius, and $R_{\rm g}=\sqrt{\sum_{n=1}^{N_{\rm s}}r_n^2/N_{\rm s}}$ is the radius of gyration, where r_n describes the distance of the nth monomer from the aggregate's centre of mass. We use $D_{\rm f}=1.8,\,k_0=1.3,$ which is based on the review in Bond and Bergstrom (2006). Although in the atmosphere black carbon aggregates may also have higher fractal dimensions (e.g. Adachi et al., 2007; Chakrabarty et al., 2014; China et al., 2014), assuming a lower fractal dimension around 1.8 yields mass absorption eross sections mass-absorption cross sections at 550 nm wavelength that lie closer to experimental data, as was shown in Kahnert and Devasthale (2011). The monomer radius was can vary within a range of 10-25 nm (Bond and Bergstrom, 2006). Here it is assumed to be a=25 nm. This is consistent with field observations (Adachi and Buseck, 2008); also, it was shown (Kahnert, 2010b) that this choice of monomer radius in light scattering computations yields results for the single-scattering albedo of black carbon aggregates consistent with observations (Bond and Bergstrom, 2006).

The calculation calculations in Kahnert and Devasthale (2011) were limited to aggregates up to $N_{\rm s}=600$. In order to cover the size range of externally mixed black carbon in SALSA we had to extend these calculations to aggregate sizes up to $N_{\rm s}=2744$, which corresponds to a volume-equivalent radius of $R_{\rm V}=350\,{\rm nm}$ (compare with Table ??2). We used the multiple-sphere T-matrix code (Mackowski and Mishchenko, 2011), which is based on the numerically exact superposition T-matrix method for solving Maxwell's equations. Figure 2 shows some of the computed black carbon optical properties as a function of particle size volume-equivalent particle radius and wavelength. All optical properties are averaged over particle orientations, where the orientation-averaging is performed analytically (Khlebstov, 1992). The absorption

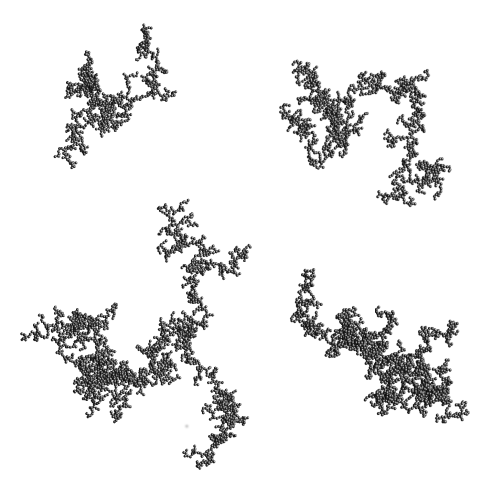


Figure 1. Examples of fractal aggregate model particles for computing optical properties of externally mixed black carbon. The aggregates consist of 1000, 1500, 2000, and 2744 monomers (in clockwise order, starting from upper left).

cross section $C_{\rm abs}$ shows the characteristic decline $\sim 1/\lambda$ at long wavelengths, where the refractive index of black carbon is changing only slowly (Chang and Charalampopoulos, 1990). Also, $C_{\rm abs}$ increases with particle size. For small particle sizes this increase goes as $\sim R_{\rm V}^3$, which is typical for the Rayleigh scattering regime (Mishchenko et al., 2002).

2. Black carbon aerosols particles that are internally mixed with other aerosol components are morphologically very complex. It is technically beyond the reach of our present capabilities to build an aerosol optics aerosol-optics database with the use of morphologically realistic model particles. However, it is possible to employ realistic model particles in reference computations for some selected cases. This has recently been done in Kahnert et al. (2013). In that study different studies Kahnert et al. (2013); Scarnato et al. (2013, 2015). In the study by Kahnert et al. (2013), optical properties of encapsulated aggregate model particles, such as the one shown in Fig. 3 (left), were computed in the size range from range of 100–500 nm

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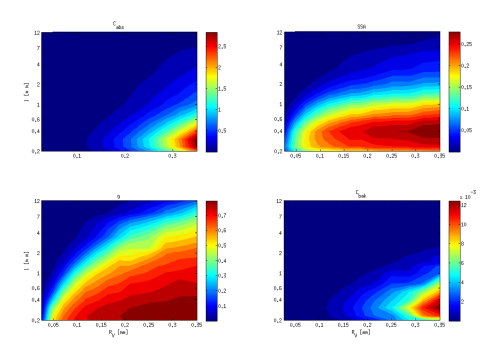


Figure 2. Absorption cross section C_{abs} (upper left), single-scattering albedo SSA (upper right), asymmetry parameter g (bottom left), and backscattering cross section C_{bak} (bottom right) of black carbon aggregates as a function of volume-equivalent radius R_V and wavelength λ .

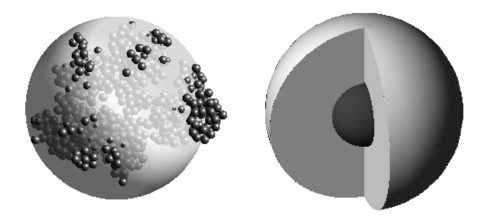


Figure 3. Morphologically realistic encapsulated aggregate model for internally mixed black carbon (left), and core-grey-shell model (right).

(volume-equivalent radius), for different black carbon volume fractions, and for wavelengths from the UV-C to the mid-IR. The morphological parameters characterising these model particles were based on field observations (Adachi and Buseck, 2008); the coating material was sulphate. The computations were performed with the discrete dipole approximation (Yurkin and Hoekstra, 2007).

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In Kahnert et al. (2013) the computational results were compared to those obtained with simple model particles, such as externally mixed externally-mixed homogeneous spheres, internally mixed internally-mixed homogeneous spheres, and concentric core-shell particles with a carbon core and a sulphate shell. The analysis revealed which morphological properties of the encapsulated aggregate particles had the dominant impact on the optical properties. There are two important properties: (1) the amount of carbon mass that interacts with the electromagnetic field has a major impact on the absorption cross section C_{abs} . In a core-shell model as well as in a model based on externally mixed homogeneous spheres, all of the black carbon is concentrated in a single sphere. Owing to absorption of the electromagnetic field does not penetrate deeply into this sphere. Hence much of the carbon mass is shielded from interacting with the field, resulting in an underestimation of $C_{\rm abs}$ compared to the encapsulated aggregates, in which a much larger fraction of the carbon mass can contribute to the absorption of electromagnetic energy. By contrast, in a homogeneous internal mixture internal-mixture model the black carbon is distributed evenly throughout the sulphate host, which allows too much of the carbon mass to interact with the field. This results in an overestimation of C_{abs} . (2) Compared to a bare black bare-black carbon aggregate, a coated aggregate has a larger geometric cross section. Hence more light is intercepted by an internally mixed-internally-mixed particle and focused onto the black carbon inclusion, thus enhancing C_{abs} . This effect is neglected in the external mixture external-mixture model, resulting in an underestimation of C_{abs} .

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Once Note that in earlier studies (e.g. Jacobson (2000)) it was often tacitly assumed that a core-shell model would give the most accurate estimates of the aerosol optical properties, owing to its morphological similarity to encapsulated black-carbon particles. However, the results in Kahnert et al. (2013) have clearly shown the shortcomings of the conventional core-shell model. But once we understand which morphological properties are most essential, and which ones make a minor contribution to the optical properties, we can devise model particles that account for the most important morphological effects, yet are sufficiently simple for computing a look-up table for large-scale modelling. It was proposed in Kahnert et al. (2013) to use a core-shell model (hence accounting for the coating effect) in which only part of the carbon mass is contained in the core, and the remaining part is homogeneously mixed with the shell. The model particle is illustrated in Fig. 3 (right). The core-shell partitioning fraction f_c of the carbon mass located in the core is a free parameter, with which one can interpolate between the two extreme models of the homogeneous mixture $(f_c = 0)$, all carbon mass mixed with the shell) and the regular core-shell model ($f_c = 1$, all carbon mass in the core). This model has been referred to as the concentric core-grey-shell (CGS) model. The tuning of the free parameter f_{c} in the model was done to fit the reference model of encapsulated aggregates as described in Kahnert et al. (2013). It was found that f_c is independent of particle size,

Table 3. Core fraction f_c in the core-grey-shell model as a function of wavelength λ .

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$\frac{\lambda \left[\mu m\right]}{f_{\rm E}}$	0.2000	0.2316	0.3040	0.3400	0.3550	0.3800	0.3932
	0.7	0.7	0.7	0.7	0.7	0.6	0.6
$\frac{\lambda \left[\mu \mathbf{m} \right]}{f_{\mathbf{c}}}$	0.4400	0.5000	0.5320	0.5332	0.6750	0.7016	0.8700
	0.6	0.5	0.5	0.5	0.5	0.5	0.2
$\frac{\lambda \left[\mu \mathbf{m}\right]}{f_{\mathrm{c}}}$	1.0101	1.0200	1.064	1.2705	1.4625	1.7840	2.0460
	0.1	0.1	0.1	0.1	0.1	0.1	0.1
λ[μm]	2.3250	2.7885	3.4615	3.5000	8.0205	10.600	12.195
Le	0.1	0.1	0.1	0.1	0.1	0.1	0.1

black-carbon volume fraction, and of the optical property one wants to fit. However, f_c does depend on the wavelength of light.

The CGS model has been employed in generating the new MATCH-optics look-up table. The shell material can be any mixture of water-soluble components. We use the same values of f_c as those determined in Kahnert et al. (2013). Its dependence on wavelength is given in Table 3.

- 3. In the mass transport model, we assume that all SIA components and all sea salt is internally mixed. We further assume that in size elasses bins 1–4, 0, 70, 70, and 100 %, respectively, of the black carbon, 0, 70, 70, and 70 % of the organic carbon, and 0, 1.3, 1.3, and 1.3 % of the dust are internally mixed; the remaining BC, OC, and dust mass is externally mixed. In SALSA, the mixing state depends on the size bin (see Table ??2), and the mixing proportions are provided by the model results. In both the mass transport model and in MATCH-SALSA, the contribution to the effective refractive index of dust and black carbon is computed by the Maxwell–Garnett EMT Maxwell Garnett (1904) (Maxwell Garnett, 1904), while for all other components we use the Bruggemann EMT (Bruggemann, 1935).
 - 4. All other externally mixed particles not containing black carbon are assumed to be homogeneous spheres in the present version of the look-up table.

The look-up tables contain results for $C_{\rm ext}$, $C_{\rm sca}$, g, and $C_{\rm bak}$ in 28 wavelength bands from the UV-C to the mid-IR. Computations with the CGS model were performed for 37 discrete BC volume fractions, namely, $f_{\rm BC} = 0.00, 0.01, \ldots, 0.20, 0.25, \ldots, 1.00$. For the shell material, as well as for non-carbon containing particles, the table contains (depending on the wavelength band) up to 40 discrete values of the real part and up to 18 discrete values of the imaginary part of the refractive index. The range of the refractive indices varies with wavelength; it is determined by those chemical components that, at each given wavelength, have the most extreme values of the refractive index. The optical properties are pre-averaged over particle sizes for each size bin. Thus we generated one

look-up table each for the mass transport model with its four size bins, and for SALSA with its 20 size bins. It-In Salsa it is assumed that the number density is constant in each size bin.

The MATCH-optics model computes in each grid cell and for each size bin the effective refractive index of the internally mixed material by use of EMT. The corresponding optical properties are obtained by linearly interpolating the closest pre-computed results in the look-up table. Size-averaging is performed by weighing the optical cross sections as well as $g \cdot C_{\text{sca}}$ in each size bin with the number density per bin and adding over all bins. The integrated quantities are then divided by the total particle number density; the integral over $g \cdot C_{\text{sca}}$ is also divided by the size-averaged scattering cross section.

2.4 Evaluation of Methodology for comparing the optics models

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The new internal-mixture optics model with its BC fractal aggregate and core-grey-shell model particles accounts for significant morphological details in aerosolsaerosol particles. The main question we want to address is whether or not this high level of detail is really necessary, i.e., if it has any *significant* impact on optical properties modelled with a CTM. By *significant* we mean an impact that is comparable to other effects whose importance is well understood. Thus to make such an assessment we need to pick a well-understood effect that can serve as a gauge, i.e., to which we can compare the impact of particle morphology on optical properties. We take the effect of aerosol dynamics microphysics as a gauge. As aerosol microphysics is well-known to have a substantial impact on aerosol concentrations and size distributions (Andersson et al., 2015; Matsui et al., 2013), this effect will provide us with a reference to which we can compare the impact of the morphological assumptions made in the aerosol-optics model. Thus we compute aerosol optical properties

- 1. with the MATCH mass-transport model (i.e., with aerosol dynamics microphysics switched off), in conjunction with the old optics model (abbreviated by MT-EXT, "mass-transport external mixture");
 - 2. with the MATCH mass-transport model in conjunction with the new optics model (MT-CGS, "mass-transport core-grey-shell");
- 3. with the MATCH-SALSA model (i.e., with aerosol dynamics microphysics switched on), in conjunction with the old optics model (abbreviated SALSA-EXT, "MATCH-SALSA external mixture");
 - 4. with the MATCH-SALSA model, in conjunction with the new optics model (SALSA-CGS, "MATCH-SALSA core-grey-shell").
- 420 Comparison of 1. and 2. will allow We first perform a comparison of monthly and geographically averaged differences in aerosol optical properties. More specifically, comparison of model set-ups MT-EXT with MT-CGS, or SALSA-EXT with SALSA-CGS allows us to assess the impact of the

morphological assumptions in the aerosol optics model. Comparison of 2. and 3. MT-EXT with SALSA-EXT, or MT-CGS with SALSA-CGS will give us an estimate of how much the inclusion or omission of aerosol dynamic microphysical processes impacts modelling results of aerosol radiometric properties. As aerosol dynamics is well-known to have a substantial impact on aerosol concentrations and size distributions, this effect will provide us with a reference to which we can compare the impact of the morphological assumptions made in the aerosol optics model.

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As an example, Fig. 4 shows the extinction aerosol optical depth (AOD) over the European model domain computed on 22 December 2007 at 12:00 with MT-EXT (left), MT-CGS (centre), and SALSA-CGS (right). The general spatial patterns are similar, as they should, since all three runs used the same EMEP emissions and HIRLAM meteorological data. However, the magnitude of the AOD results can differ significantly among the three model runs (note the semi-logarithmic scale!). Interestingly, differences While statistical analyses can uncover general trends, it is difficult to understand the underlying physical reasons for model differences from an analysis of temporally and geographically averaged model results. Therefore, we also consider a few case studies in some more detail. We take the optical properties modelled with different MATCH-versions as input to a radiative transfer model and analyse the total aerosol radiative forcing and the black-carbon radiative forcing. The main goal is to understand how differences in single-scattering optical properties between the two optics models (left and centre) are roughly on the same order as those between the mass-transport and aerosol dynamic models (centre and right).

Aerosol optical depth over Europe on 22 December 2007, 12:00 (noon). Results are shown for the mass transport model in conjunction with the old external-mixture optics model (left), and with the new internal-mixture/core-grey-shell/fractal BC aggregate model (center), as well as for the MATCH-SALSA model in conjunction with the new optics model (right). The circles indicate the four locations used for radiative transfer studies. Note the semi-logarithmic colour scale!

Thus a first inspection of computed fields of aerosol optical properties suggests that the level of detail in the morphological assumptions of the aerosol optics model may be significant for the modelling results. In a next step we quantify differences in modelled aerosol radiative forcing among the three model versions. To this end we pick impact the outcome of the radiative transfer simulations. To keep the case studies within manageable bounds, we restrict ourselves to four geographic locations that are indicated by circles in Fig. 4; (two over land, two over the ocean), two instances (one representing low-BC summer concentrations, one representing high-BC winter conditions), and we limit ourselves to comparing the model set-ups MT-EXT, MT-CGS, and SALSA-CGS.

More specifically, we consider one site over Northern Italy (45.0° N, 8.5° E), one over the Mediterranean Sea (37.5° N, 5.5° E), one over Poland (52.6° N, 21.0° E), and one over the North Sea (52.0° N, 2.7° E). We further pick two instances representing low-BC summer and high-BC winter conditions, namely, For the two instances, we pick 22 June 2007 12:00 UTC, and 22 December 2007 12:00 UTC. Radiative transfer calculations are performed for each of these four sites and for

both instances. Vertical profiles of the aerosol optical depth per layer, the single-scattering albedo, and the asymmetry parameter are used as input to the libRadtran radiative transfer package (Kylling et al., 1998), assuming a plane-parallel atmosphere. For the surface albedo of the ocean we assume a spectrally constant value of 0.065, while for the spectrally varying surface albedo of the two land locations we used MODIS observations for each of the two instances. The results were spectrally integrated to obtain the broadband radiative fluxes. The radiative transfer simulations were repeated for corresponding profiles of optical properties (with a 1 km resolution) in the absence of black carbon, as well as for clear-sky conditions. This allows us to compute differences in broadband radiative fluxes, i.e., the radiative effect of black carbon, and the radiative effect of all aerosolsaerosol components. The results of this radiative transfer study are discussed in Sect. 3.2.

To further investigate the significance of the optics model on radiometric properties, we look at remote sensing-related optical properties also look at optical properties relevant for remote sensing, namely, backscattering coefficient and Ångström exponent. These results are discussed in Sects. 3.3 and 3.4, respectively.

3 Results

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475 3.1 Aerosol optical properties in MATCH

Figure 4 gives us a first impression of the differences among the four model configurations. The extinction AOD is shown for MT-EXT (1st from the left), MT-GCS (2nd), Salsa-CGS (3rd) and Salsa-EXT (4th). The overall spatial patterns are similar. This is expected, since all model configurations used the same EMEP emissions and HIRLAM meteorological data. However, the magnitude of the AOD results can differ significantly among the four model runs (note the semi-logarithmic scale!). It is also remarkable that the differences between the two optics models depend on whether we make this comparison within the mass-transport model (compare MT-EXT to MT-CGS), or within Salsa (compare SALSA-CGS and Salsa-EXT). It can also vary geographically. This merely confirms the complexity of aerosol-optics modelling. Replacing one optics model by another will not simply offset the resulting optical properties by some common factor; it will introduce a significant change in modelled optical properties, of which the magnitude and even the sign can be dependent on local conditions, such as the size-distribution and the chemical composition of the aerosol particles.

This is also evident from a comparative analysis of geographically and temporally averaged aerosol optical properties. Table 4 shows aerosol optical depth (AOD), backscattering coefficient (BSCA), single scattering albedo (SSA) and asymmetry parameter (g), each at three different wavelengths (355, 532, and 1064 nm), and each averaged over the whole model domain and over one month (December 2007). The columns show relative differences; for instance, MT(EXT,CGS)=(MT-EXT – MT-CGS)/(MT-CGS) is the relative difference of MT-model results obtained with the EXT and CGS optics models.

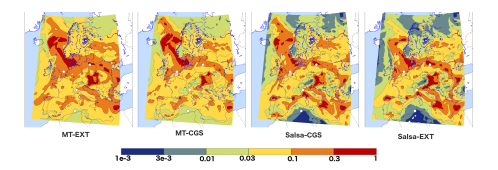


Figure 4. Aerosol optical depth over Europe on 22 December 2007, 12:00 UTC (noon). Results are shown for the mass transport model in conjunction with the old external-mixture optics model (1st to the left), and with the new internal-mixture/core-grey-shell/fractal BC aggregate model (2nd to the left), as well as for the MATCH-SALSA model in conjunction with the new optics model (3rd to the left) and old optics model (4th to the left). The circles indicate the four locations used for radiative transfer studies. Note the semi-logarithmic colour scale!

Table 4. Averaged relative difference in aerosol optical depth (AOD), backscattering coefficient (BSCA), single scattering albedo (SSA) and asymmetry parameter (g), among the different model set-ups for December 2007. The average has been performed over a whole month and over all grid-cells (horizontally for AOD, horizontally and vertically for all other optical properties). Each number corresponds to a relative difference between two model set-ups. For instance, MT(EXT,CGS) = (MT-EXT – MT-CGS)/(MT-CGS) compares results obtained with the mass transport model (MT) by using the two different optics models (EXT and CGS).

X	MT(EXT,CGS)	Salsa(EXT,CGS)	CGS(MT,Salsa)	EXT(MT,Salsa)
AOD ₃₅₅	0.16	~-0.28~	-0.50	-0.19
$\underbrace{\text{AOD}_{532}}_{}$	0.08	-0.14	0.00	0.25
$\underbrace{\text{AOD}_{1064}}$	0.18	-0.03	0.14	0.37
BSCA ₃₅₅	0.44	~0.01	-0.47	-0.23
$\underbrace{BSCA_{532}}_{BSCA_{532}}$	0.26	-0.08	-0.19	0.11
BSCA ₁₀₆₄	0.99	~0.01	-0.36	0.28
SSA ₃₅₅	-0.02	0.04	0.03	-0.03
SSA_{532}	-0.02	0.05	0.04	-0.02
SSA ₁₀₆₄	-0.07	0.08	0.08	-0.07
£ 355	0.06	~-0.01	-0.10	-0.03
£ 532	0.10	-0.00	-0.06	0.04
g 1064~	<u>0.17</u>	~-0.02	-0.11	0.06

Comparison of the columns "MT(EXT,CGS)" and "Salsa(EXT,CGS)" illustrates the differences between the optics models in the absence and presence of aerosol-microphysical processes. As we already suspected from inspection of Fig. 4, differences in the optics models defy simplistic explanations; both the magnitude and sign of these difference can be strongly dependent on the size-dependent chemical composition and mixing state of the aerosols, hence on the model version with which the optics models are being compared. In our case, we see that compared to the CGS model, the EXT-optics model predicts higher values of AOD, BSCA, and g in the MT model, and lower values of SSA. In Salsa the roles of the CGS and EXT model are reversed.

The column "CGS(MT,Salsa)" shows differences between optical properties computed in the absence and presence of aerosol-microphysical processes, where the optics-computations have been performed with the CGS model. The column "EXT(MT,Salsa)" shows an analogous comparison, where the optics-computations have been performed with the EXT model. If we compare the magnitudes of the entries in the columns "MT(EXT,CGS)" and "Salsa(EXT,CGS)" with the corresponding magnitudes of the entries in the columns "CGS(MT,Salsa)" and "EXT(MT,Salsa)", then we see that the differences between the two optics models (EXT,CGS) are roughly on the same order as the differences between the two aerosol models (MT,Salsa). Thus, the main observation is that the choice of aerosol-optics model can have an effect on modelled optical properties that is of comparable magnitude as the level of detail in the description of aerosol-microphysical processes.

While spatio-temporally averaged model results allows us to draw some general conclusions, it is difficult to understand the reasons for the observed differences from such an analysis. We will, therefore, complement this investigation in the following sections with a more detailed analysis of some selected case studies.

3.2 Optical properties and radiative forcing

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In Sec. 2.3.2 we have discussed how morphological properties of aerosol particles can impact their optical properties. We now take this one step further and discuss how the optical properties of particles can impact the radiative properties of an aerosol-laden atmosphere. We will show results for a single wavelength near the maximum of the solar spectrum. The comparison will focus on three model set-ups, MT-EXT, MT-CGS, and Salsa-CGS. We will use MT-CGS as a reference and compare it first to Salsa-CGS in order to investigate the impact of aerosol-microphysics (MT versus Salsa). Next, we compare MT-CGS to MT-EXT in order to investigate the impact of the optics model (CGS versus EXT).

The result for the optical properties obtained with the three model versions (AOD per layer, SSA and g) at the wavelength 532(CGS)/500(EXT) nm, together with the radiative forcing for aerosols all aerosol component and black carbon, respectively, can be seen in Figs. 5–10 for Northern Italy and the Mediterranean on 22 June 2007. Each figure shows the differences in direct solar flux $\Delta F_{\rm s}$ (top left), diffuse downwelling flux $\Delta F_{\rm d}$ (top right), diffuse upwelling flux $\Delta F_{\rm u}$ (centre

left), and net radiative flux $\Delta F_{\rm net} = \Delta F_{\rm s} + \Delta F_{\rm d} - \Delta F_{\rm u}$ (centre right), where either the difference between aerosol-laden and clear sky conditions are considered (Figs. 5 and 6), or the difference between fluxes in the presence and absence of black carbon (Figs. 9 and 10). Top-of-atmosphere Here, *downwelling* fluxes are obtained by integrating the radiance over all azimuthal angles, and over polar angles from 90° to 180°, where the positive z-axis is directed from the ground to the top-of-atmosphere (TOA). Similarly, the *upwelling* flux is obtained by integrating the radiance over all azimuthal angles, and over polar angles from 0 to 90°. TOA results for the other geographical locations are summarized in Table 5 in terms of aerosol forcing in Table 5 and black carbon forcing ($\Delta F_{\rm net} = F_{\rm net}$ (with aerosol particles) – $F_{\rm net}$ (no aerosol particles)), and in Table 6 in terms of black-carbon forcing ($\Delta F_{\rm net} = F_{\rm net}$ (with black carbon) – $F_{\rm net}$ (no black carbon)). The wavelengths 532(CGS)/500(EXT) nm are near the maximum of the solar spectrum. At other wavelengths (not shown) the optical properties behave similarly. Each figure has a vertical span of 65 km, which comprises that part of the troposphere where almost all aerosols most aerosol particles are concentrated in the cases we picked.

Before starting our analysis, we note that the magnitude of the radiative fluxes generally depends on the concentration of aerosols. As we cannot claim that the test cases we happened to pick are in any way representative for *typical* aerosol and black carbon loads, we are not focusing on the *magnitude* of the radiative fluxes here. Rather, we want to compare the *differences* in radiative fluxes among the three model versions.

3.2.1 Comparison of aersol dynamics aerosol microphysics and mass-transport model

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We start by comparing radiative fluxes in the presence and absence of aerosolsall aerosol components, which we refer to as the "aerosol radiative effects". Figures 5 and 6 show the aerosol radiative effects modelled over Northern Italy and over the Mediterranean north of Algier Algiers, respectively. The general patterns in both plots can be understood as follows. In the presence of aerosolsall aerosol components, optical extinction is stronger than in clean air. Thus the presence of aerosols all aerosol components gives a reduction $\Delta F_{\rm s}$ of the direct solar flux (upper left). As the aerosol optical depth per atmospheric layer strongly increases near the ground, the magnitude of $\Delta F_{\rm s}$ increases sharply with decreasing altitude. Further, aerosol extinction extinction in the form of scattering results in the generation of diffuse flux; the downwelling diffuse flux accumulates downward, resulting in an increasing excess of downwelling flux $\Delta F_{\rm d}$ in the presence of aerosols aerosol components as one approaches the surface. The upwelling flux $F_{\rm u}$ is generated by scattering in the atmosphere and reflection at the surface. Since aerosol extinction reduces the total net radiative flux as one approaches the surface, less upwelling diffuse flux is generated at low altitudes; hence the difference in upwelling flux $\Delta F_{\rm u}$ between an aerosol-laden and a clear sky atmosphere is negative near the surface. However, it increases with altitude, because at higher altitudes the magnitude of the difference (aerosol –

clear sky) in the total net radiative flux that can be converted into upwelling diffuse flux decreases at higher altitudes.

If we focus now on differences in the radiative net flux $\Delta F_{\rm net}$ at high altitudes, i.e., the radiative forcing effect of aerosolsaerosol particles, then we see significant differences between the mass transport model (MT, red) and SALSA (green) at both geographic locations. It is evident that the main cause for these are corresponding differences in the diffuse upwelling flux $\Delta F_{\rm u}$.

At both locations the diffuse upwelling flux is smaller for SALSA then for MT, but for different reasons. Over the Mediterranean (Fig. 6), the AOD is significantly smaller for SALSA than for MT, resulting in less extinction of the direct flux, hence less generation of diffuse flux, and a smaller radiative cooling effect for SALSA.

Over Northern Italy (Fig. 5), there is almost no difference in AOD between the two models. It can be seen from the AOD profile that the majority of aerosols aerosol particles reside in the lowest 1 km near the surface. However, above 1 km the results of $\Delta F_{\rm u}$ obtained with SALSA and MT diverge with increasing altitude. This is a result of the reflection by the near-surface aerosol layer, which is slightly different in the two models. In MT the SSA is higher than in SALSA, resulting in more scattering, thus in more diffuse radiation. The asymmetry parameter is slightly larger in MT than in SALSA; correspondingly, the partitioning in MT between downwelling and upwelling radiation is somewhat shifted in favour of the former. However, this only partially counteracts the generation of a higher amount of diffuse upwelling radiation in MT due to the higher SSA. The net effect is a higher value of $\Delta F_{\rm u}$ in MT, hence a larger radiative cooling effect at higher altitudes.

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To further analyse the difference in optical properties between MT and SALSA, we look at the aerosol masses and the relative sizes of the particles. Figure 7 shows vertical profiles of the effective radius of the size distributions $r_{\rm eff}$ defined according to Eq. 1, in SALSA (green) and the MT model (black) over Northern Italy (left) and over the Mediterranean (right).

$$r_{\text{eff}} = \frac{\int_0^\infty n(r)\pi r^2 \cdot r dr}{\int_0^\infty n(r) \cdot \pi r^2 dr}$$
 (1)

Figure 8 shows profiles for the total aerosol mass (1st row), BC (black carbon) (2nd row), sulphate (3rd row), and nitrate (4th row) for both Northern Italy (1st column) and Mediterranean (2nd column). We focus on the total aerosol mass, which is expected to impact the aerosol optical depth. The aerosol optical depth is dependent on the number density (which, in turn, increases with the mass mixing ratio), as well as on the extinction cross section (which generally increases with the effective radius of the particles). Over Northern Italy, the SALSA model predicts a larger mass mixing ratio than the MT model (Fig. 8, upper left) , but also and a smaller particle size (Fig. 7, left). This results in a higher number density but a smaller extinction cross section. These two effects cancel almost exactly, resulting in nearly identical aerosol optical depths predicted with the two models (Fig. 5, bottom left). By contrast, over the Mediterranean the two models predict similar mass mixing ratios (Fig. 8, upper right), while SALSA predicts a much lower effective radius than the MT model

(Fig. 7, right). As a consequence, the optical depth is significantly lower in SALSA than in MT (Fig. 6, bottom left). The SSA is lower in SALSA than in MT. This is mainly caused by the fact that the effective radius is smaller in SALSA than in MT, since SSA is usually increasing with size.

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The radiative impacts For the other two geographical locations and the second time event, the TOA results are summarised in Table 5show the same behaviour as Figs. 5 and 6 at three of the four geographical locations. The Polish site deviates, since SALSA produces a much larger radiative cooling than the MT model. Over the North Sea, Northern Italy, and the Mediterranean the TOA forcing is strongest in the MT-EXT model (mass transport with old optics model), it is smaller in the MT-CGS model (mass transport with new optics model), and weakest in the SALSA-CGS model (aerosol-microphysics with new optics model). However, over Poland the negative forcing in SALSA-CGS is strongest among the three models in the summer, and second strongest in the winter. This can be explained by SALSA having a larger amount of aerosols aerosol particles throughout the column at that site, especially more sulphate, which, when externally mixed, contributes to a larger amount of scattering and therefore a higher SSA and a larger diffuse upwelling radiative flux.

We now compare radiative fluxes in the presence and absence of black carbon, which we refer to as the "black carbon radiative effect". Figures 9 and 10 show the radiative effect of black carbon together with the optical properties with and without black carbon. Again, the dominant feature of differences in $\Delta F_{\rm net}$ at TOA comes from are mainly caused by corresponding differences in the upwelling diffuse radiative flux $\Delta F_{\rm u}$. In these figures, we have to focus on the difference in the optical properties when analysing the radiative fluxes. The general pattern can be seen in Fig. 9, which shows the differences in radiative fluxes and in the optical properties over Northern Italy. The direct solar flux (upper left) decreases with decreasing altitude owing to extinction. The magnitude of the decrease mainly reflects the differences in optical depth in the presence and absence of black carbon (bottom left), which is slightly-larger in SALSA than in MT. The decrease in solar flux does not automatically result in an increase in the downwelling diffuse flux with decreasing altitude (upper right), as it was in the comparison of fluxes in the presence and absence of all aerosolsaerosol components. The situation is more complex now. Near the surface, where the optical depth is largest, the difference in SSA in the presence and absence of black carbon is quite large in the MT model (bottom centre, red lines), and slightly smaller in SALSA (green lines). As a result, absorption contributes more to the total extinction in the MT model than in SALSA (at least near the surface). Hence, the portion of the downwelling flux that is absorbed on its way downward is larger in the MT model than in SALSA, resulting in a decrease of the diffuse downwelling flux with decreasing altitude (upper right, red line). The differences between the two models in the diffuse upwelling flux are very small (centre left, red and green lines). This is the result of cancelling effects; for instance, there is less direct solar flux, but more diffuse downwelling flux in SALSA that can be converted into diffuse upwelling flux through scattering. As a result, the differences between both models in the net flux (centre right, red and green lines) are almost negligible.

Figure 10 shows the radiative effect of black carbon over the Mediterranean. Again, the dominant feature of differences in $\Delta F_{\rm net}$ at TOA comes from are mainly caused by corresponding differences in the upwelling diffuse radiative flux $\Delta F_{\rm u}$. The difference between MT and SALSA are not as prominent here, mainly due to the fact that the differences in the optical properties similar, especially in the SSA. There is still a small difference for the Mediterranean, where SALSA has a larger radiative cooling than MT. This difference is very small, but it may come from the slight difference in the AOD combined with the difference in SSA from above 1, where SALSA has a marginally larger difference than MT, creating the difference in the net radiative flux above 1. Another possibility is that these small differences are caused by multiple scattering effects, which are notoriously difficult to understand by an intuitive approach. If studying the differences in AOD and SSA for the twelve wavelengths used in this study (not shown), the differences for SALSA AOD and SSA are smaller for the long-wave (LW) part of only differ by a few mW/m². The main reason is that the optical properties, especially the spectrum (533.2–3461.5). This results in less LW extinction and scattering and slightly more radiative cooling for SALSASSA, differ in the presence and absence of BC by almost the same amount in both models.

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Table 6 shows the black carbon forcing for all the four geographical locations and both months. In June the difference at the TOA is very small between MT and SALSA for all the locations, but larger for the Mediterranean as noted in Fig. 10. The month of December shows a pattern where SALSA has a higher radiative heating over land, and smaller over the ocean compared to MT. This is strongly coupled to the larger difference in BC amounts over land than over the oceans, where the two models have similar values all through the column, see FigAs a general trend, large differences in BC concentrations are visible as corresponding differences in BC forcing. For instance, near-surface BC winter- concentrations in Northern Italy are about a factor of 10 higher than in summer; summer-concentrations over the Mediterranean are more than a factor of 2 higher than in winter; in Northern Italy in winter, Salsa predicts more than 2 times higher BC-concentrations than the mass-transport model, while over the Mediterranean in winter the role of the two models is reversed (not shown). All of this corresponds with the respective BC-forcing rates in Table 6. However, when the differences in BC-concentrations are small, then there is no clear relation between the concentration-differences and the corresponding differences in BC-forcing rates. For instance, as we see in Fig. 8, there is almost no difference between the BC-concentrations computed with the two models over the Mediterranean in summer. But the table shows that the mass-transport model predicts a slightly higher forcing rate than SALSA. A possible cause is the difference in the size-distributions in SALSA and the mass-transport model. 8 for Northern Italy and Mediterranean. Poland and North Sea are not shown here, but show similar behaviour.

3.2.2 Comparison of the two optics models

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The comparison between SALSA and the MT model in the previous section served two purposes. First, it helped us to develop a basic understanding for the effects of aerosols aerosol particles and black carbon on radiative fluxes. Second, it provided us with a gauge for assessing the importance of the aerosol optics aerosol-optics model, which will be the subject of this section.

Aerosol forcing and optical properties at 532(CGS)/500(EXT) nm for Northern Italy in June. Aerosol forcing and optical properties at 532(CGS)/500(EXT) nm for the Mediterranean in June. The aerosol forcing at the top of the atmosphere (TOA), Δ*F*_{net,TOA} W, for the four different geographical locations, one summer and one winter event, and three model versions. MT-EXT MT-CGS SALSA-CGS SummerPoland – 0.21 – 0.21 – 0.77 North Sea – 0.34 – 0.29 – 0.24 Northern Italy – 0.06 – 0.05 0.01 Mediterranean – 1.20 – 0.99 – 0.30 WinterPolen – 4.41 – 2.00 – 2.18 North Sea – 2.85 – 1.21 – 0.86 Northern Italy – 1.15 – 0.53 – 0.57 Mediterranean – 0.09 – 0.04 – 0.03

We compare the old EXT (blue line) and the new CGS (red line) optics models in Figs. 5 and 6, each used in conjunction with the MT-version of MATCH. The net radiative flux ΔF_{net} in the CGS model shows a weaker TOA cooling effect than the EXT model, both over Northern Italy and over the Mediterranean. Again, the upwelling flux has the dominant impact on the behaviour of the TOA net radiative flux. Over Northern Italy (Fig. 5) the diffuse upwelling flux is larger for the EXT model above 1 km, whereas it is smaller below 1 km and at the bottom of the atmosphere (BOA). The AOD profile reveals that in the EXT model extinction is stronger than in the CGS model throughout the tropospheric column. As a result, there is more diffuse downwelling flux being generated in EXT than in CGS. At the bottom of the atmosphere (BOA) extinction of the direct flux is stronger than generation of diffuse downwelling flux; hence less downwelling flux is reflected by the surface, resulting in less BOA upwelling diffuse flux in EXT than in CGS. Higher up in the troposphere, the upwelling diffuse flux is mostly generated by atmospheric scattering rather than reflection from the surface. As the SSA in EXT is higher than in CGS, more diffuse flux is generated, resulting in a stronger radiative cooling effect in EXT than in CGS.

Over the Mediterranean (Fig. 6), the EXT and CGS model have almost identical AOD profiles in the green part of the spectrum. However, at longer wavelengths (not shown) EXT predicts substantially higher AOD values than CGS (see (see Appendix E, Fig. 16). For instance, at $\lambda = 1020$ nm the near-surface AOD per layer computed with the EXT-model is about twice as high as that computed with the CGS-model. This explains the larger amount of diffuse broadband radiation generated in the EXT model, which results in a stronger negative TOA net flux in EXT as compared to the CGS model. Note that the differences in SSA between EXT and CGS are fairly smallless than 0.03, while the differences in g are rather large as much as 0.07. The higher values of g in EXT may contribute to the large amount of diffuse downdwelling downwelling radiation in that model; however, the dominant effect is likely to be the high optical depth at red and IR wavelengths (see Appendix E).

Table 5. The aerosol forcing at the top of the atmosphere (TOA), $\Delta F_{\text{net,TOA}}$ [W m⁻²], for the four different geographical locations, one summer (2007-06-22, 12:00) and one winter (2007-12-22, 12:00) event, and three model versions, MT-EXT, MT-CGS and Salsa-CGS.

		MT-EXT	MT-CGS	SALSA-CGS
Summer	Poland	<u>-0.21</u>	<u>-0.21</u>	<u>-0.77</u>
	North Sea	<u>-0.34</u>	-0.29	~ 0.24
	Northern Italy	-0.06	-0.05	0.01
	Mediterranean	-1.20	-0.99	-0.30
Winter	Poland	-4.41	-2.00	-2.18
	North Sea	-2.85	-1.21	~ 0.86
	Northern Italy	<u>-1.15</u>	-0.53	~ 0.57
	Mediterranean	-0.09	-0.04	-0.03

Table 5 summarises the TOA net radiative flux at all four geographical locations for both June and

December. The largest differences among the models are seen in December at the two northernmost locations, i.e., Poland and the North Sea. At these two places, the total aerosol amount (not shown) is significantly higher than at the other two locations farther south, giving rise to a-larger absolute changes in the aerosol forcing.

The black carbon forcing looks rather different at the two geographical locations in Fig. 9 (Northern Italy) and 10 (Mediterranean) display different behaviours in radiative fluxes, comparing the EXT (blue) and CGS (red) model results. Over Northern Italy, the net black carbon forcing is more significant than over the Mediterranean in Fig. 10 due to higher levels of BC, see Fig. 8. As can be seen in Fig. 9, the differences in optical properties computed with and without black carbon are larger in the CGS model than in the EXT model, particularly for the SSA. This means that in the CGS model the presence of black carbon causes more absorption than in the EXT model, thus generating less diffuse down- and upwelling flux by scattering. As a result, the CGS model predicts more radiative warming, i.e., a higher TOA radiative net flux than the EXT model. The reason for this is that (i) the CGS model treats externally mixed soot as aggregates, which have a lower SSA than the massive black carbon spheres in the EXT model; and (ii) the CGS model treats internally mixed soot as a coated core-grey-shell model, which accounts for focusing of electromagnetic radiation onto the carbon core, thus enhancing absorption, i.e., lowering the SSA, while the EXT model treats *all* black carbon as externally mixed.

Effective radius for the two chemical transport model versions MT and SALSA over Northern Italy and Mediterranean in June.

Vertical distribution of aerosols at Northern Italy and Mediterranean in June.

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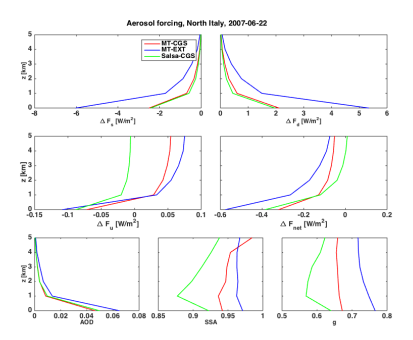


Figure 5. Aerosol forcing and optical properties at 532(CGS)/500(EXT) nm over Northern Italy in June. Results are shown for the three model versions MT-EXT (blue), MT-CGS (red) and Salsa-CGS (green). The aerosol forcing is derived by taking the difference in radiative fluxes between an aerosol laden and a clear sky. The difference in direct ($\Delta F_{\rm s}$) and diffuse ($\Delta F_{\rm d}$) downwelling, as well as the diffuse upwelling ($\Delta F_{\rm u}$) and the net radiative flux (aerosol forcing) ($\Delta F_{\rm net} = \Delta F_{\rm s} + \Delta F_{\rm d} - \Delta F_{\rm u}$) are shown in the first four figures (1st and 2nd row of plots). The optical properties aerosol optical depth (AOD), single scattering albedo (SSA) and asymmetry parameter (g) are shown in the 3rd row of plots.

Table 6. The Same as table 5, but for black carbon forcing at the top of the atmosphere (TOA), $\Delta F_{\text{net,TOA}}$ W, for the four different geographical locations, one summer and one winter event, and three model versions.

		MT-EXT	MT-CGS	SALSA-CGS
Summer	Poland	1.02×10^{-2}	1.16×10^{-2}	1.20×10^{-2}
	North Sea	1.71×10^{-2}	1.54×10^{-2}	1.49×10^{-2}
	Northern Italy	4.61×10^{-2}	7.77×10^{-2}	7.86×10^{-2}
	Mediterranean	8.54×10^{-3}	6.45×10^{-3}	2.41×10^{-3}
Winter	Poland	4.03×10^{-2}	3.56×10^{-2}	6.83×10^{-2}
	North Sea	1.95×10^{-2}	2.28×10^{-2}	4.97×10^{-3}
	Northern Italy	6.73×10^{-2}	1.08×10^{-1}	1.46×10^{-1}
	Mediterranean	6.03×10^{-4}	1.34×10^{-3}	3.13×10^{-4}

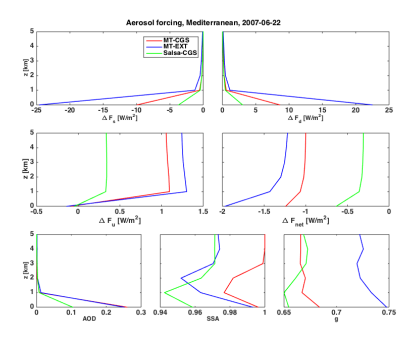


Figure 6. Same as figure 5 but over the Mediterranean.

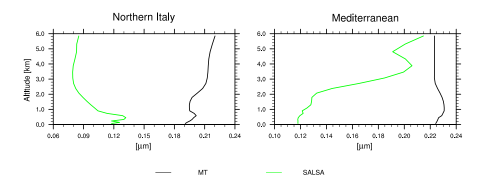


Figure 7. Black carbon forcing and optical properties at 532(CGS)/500(EXT) nm Effective radius, $r_{\rm eff}$, for the two chemical transport model versions MT and SALSA over Northern Italy in June and Mediterranean the 2007-06-22 at 12:00.

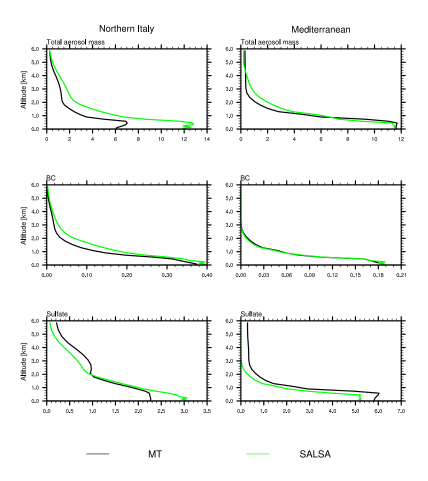


Figure 8. Vertical distribution of aerosol particles in Northern Italy and over the Mediterranean on 2007-06-22 at 12:00.

3.3 Gauging the significance of aerosol optics modelling

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Now that we have understood the impact of the acrosol optical properties on radiative fluxes, we finally turn Let us now return to the main question of this study. We ask if , namely, whether or not the level of detail in acrosol optics modelling has a significant impact on observable radiometric properties. As a gauge we consider the changes in radiometric properties. We already saw in Table 4 that, on average, the effect of including acrosol microphysics on optical properties is of comparable magnitude as the effect caused by the inclusion or omission of acrosol dynamic processes. Thus we morphological assumptions in the acrosol optics model. However, we also saw that the magnitude and sign of these impacts can be quite variable and depending on several factors. We find this confirmed in the analysis of our radiative-transfer study. More specifically, we can compare in Figs. 5–10 the differences in radiative forcing between the MT-CGS and Salsa-CGS (red and green lines) to the corresponding differences in the between MT-CGS and MT-EXT (red and blue

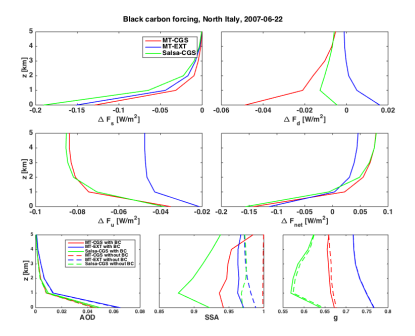


Figure 9. Black carbon forcing and optical properties at 532(CGS)/500(EXT) nm over Northern Italy in June. Results are shown for the three model versions MT-EXT (blue), MT-CGS (red) and Salsa-CGS (green). The black carbon forcing is derived by taking the difference in radiative fluxes between an aerosol laden including black carbon and an aerosol laden sky omitting black carbon. The difference in direct (ΔF_s) and diffuse (ΔF_d) downwelling, as well as the diffuse upwelling (ΔF_u) and the net radiative flux (aerosol forcing) ($\Delta F_{net} = \Delta F_s + \Delta F_d - \Delta F_u$) are shown in the first four figures (1st and 2nd row of plots). The optical properties aerosol optical depth (AOD), single scattering albedo (SSA) and asymmetry parameter (g) are shown in the 3rd row of plots.

lines). We see that in some cases the choice of optics model has a stronger effect than the inclusion of aerosol dynamics microphysics (e.g. Fig. 9), while in other cases it is the other way round (e.g. Fig. 6). We can also inspect Tables 5 and 6 and arrive at the same result. On average, the effect of including aerosol dynamics on the TOA radiative forcing is of comparable magnitude as the effect caused by employing a more realistic aerosol optics model.

In the following two subsections, we will focus on the selected case-studies and discuss the significance of the optics model for radiometric quantities that are relevant for remote sensing applications.

750 3.3 Backscattering coefficient

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From ground-based and space-borne $\frac{\text{lidars-lidar}}{\text{lidars-lidar}}$ measurements one can obtain the aerosol backscattering coefficient β , which is proportional to the backscattering cross section C_{bak} of the particles and the aerosol number density. Figure 11 shows vertical profiles of β computed at two locations

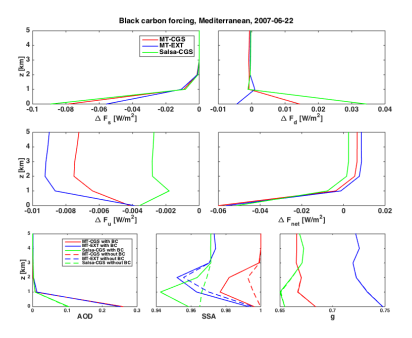


Figure 10. Black carbon forcing and optical properties at 532(CGS)/500(EXT) nm for Same as Figure 9, but over the Mediterraneanin June.

and at two instances, as indicated in the figure headings. Each panel shows computational results obtained with the three different model versions. The figure shows results for the second Nd:YAG harmonic of 532 nm. Corresponding results computed for wavelengths of 355 and 1064 nm lead to similar conclusions.

We saw in Fig. 8 for June over Northern Italy (upper left) that SALSA predicts an aerosol mass mixing ratio, hence a particle number density, that is higher than that in the MT model. But we also saw in Fig. 7 (left) that SALSA predicts lower values of $r_{\rm eff}$. This results in lower values of $C_{\rm bak}$. We see in Fig. 11 (upper left) that the effect on β of the higher number density dominates over the effect of the lower $r_{\rm eff}$, resulting in values of β that are about 30 % higher in SALSA (green line) than in the MT model (red line). Over the Mediterranean, both SALSA and the MT model predict similar mass densities mixing ratios (Fig. 8, upper right); but SALSA still predicts substantially lower values of $r_{\rm eff}$ (Fig. 7, right). The result is that β computed with the MT model (red line) is almost twice as high as the corresponding results obtained with SALSA (green line) (Fig. 11, upper right).

A similar comparison of the two optics models (red and blue lines in Fig. 11) shows that the new CGS optics model consistently predicts substantially lower values of β than the old EXT optics model. This agrees with the comparison shown in Kahnert et al. (2013) between encapsulated black carbon aggregates and externally mixed homogeneous spheres. (In a retrieval algorithm, an optics model that overestimates the backscattering cross section would result in underestimated retrieval

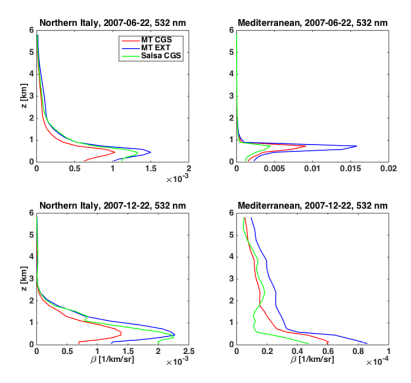


Figure 11. Backscattering coefficient at a wavelength of 532 nm at two different locations. Northern Italy and Mediterranean and at two different instancestime events (22/6 and 22/12-2007 at 12.00), computed with the three model versions, MT-EXT (blue), MT-CGS (red) and Salsa-CGS (green).

results for the particle number density.) The differences between the two optics models are on the same order of magnitude (and often even slightly larger) than the corresponding differences between the SALSA and the MT versions of the aerosol transport model.

775 3.4 Ångström exponent

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The Ångström exponent α in a wavelength interval $[\lambda_1, \lambda_2]$ is defined as

$$\alpha = -\frac{\log(\tau(\lambda_1)/\tau(\lambda_2))}{\log(\lambda_1/\lambda_2)},\tag{2}$$

where τ denotes the extinction optical depth. This quantity is often used for obtaining particle size information (usually, the smaller the particle size, the larger α). Table 7 shows values of α for our different test cases computed with the three model versions in the wavelength interval 532–1064 nm. If we compare the columns labelled MT-CGS and SALSA-CGS, then we see that the mass-transport model consistently gives lower values of α . This is related to the high values of $r_{\rm eff}$ in that model, which we noted earlier. On the other hand, if we compare the columns labelled MT-EXT and MT-CGS, then we see that the new optics model (CGS) predicts higher values of α than the

Table 7. Ångström exponent in the wavelength region 532–1064 nm for the four different geographical locations, one summer (2007-06-22, 12:00) and one winter (2007-12-22, 12:00) event, and three model versions, MT-EXT, MT-CGS and Salsa-CGS.

		MT-EXT	MT-CGS	SALSA-CGS
Summer	Poland	0.32×10^{0}	0.12×10^{1}	0.28×10^{1}
	North Sea	0.80×10^{0}	0.12×10^{1}	0.21×10^{1}
	Northern Italy	0.11×10^{1}	0.11×10^{1}	0.15×10^{1}
	Mediterranean	0.36×10^{0}	0.12×10^{1}	0.21×10^{1}
Winter	Poland	0.80×10^0	0.12×10^1	0.22×10^1
	North Sea	0.79×10^0	0.11×10^1	0.14×10^1
	Northern Italy	0.13×10^1	0.10×10^1	0.12×10^1
	Mediterranean	0.13×10^1	0.98×10^{0}	0.14×10^1

old model (EXT) in the first six rows, and lower values in the last two rows June (summer) for all four geographical locations and in December (winter) for the locations Poland and North Sea. This indicates that the errors introduced by the simple external-mixture model in computing α are quite unpredictable, even the sign of the error. When used in a size retrieval algorithm the retrieval errors caused by the EXT model would be equally hard to predict. The difference between the MT and SALSA model is somewhat larger, but not much larger, than the differences between the old and new optics models. Note that the performance of the MT model could be improved in comparison to SALSA by modifying the assumed size distribution in the MT model. By contrast, the differences between the two optics models is are rather fundamental; it is caused by the over-simplified simple treatment of aerosol morphology in the EXT model.

795 4 Conclusions

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We have implemented a new acrosol optics acrosol-optics model in a regional chemical transport model. The new model differs from an earlier optics model described in Kahnert (2008) in three essential points. (i) While the old model treats all chemical components as externally mixed, the new model accommodates both external and internal mixtures of acrosol species. (ii) The old model treats black carbon acrosols particles as homogeneous spheres; the new model assumes a fractal aggregate morphology with fractal parameters based on observations. Mass absorption cross sections and single scattering albedos computed with this model have previously been evaluated by comparison with measurements (Kahnert, 2010b). (iii) The new model describes internally mixed black carbon acrosols particles by a recently developed "core-grey-shell" model (Kahnert et al., 2013). This model accounts for the inhomogeneous internal mixing state of black carbon aggregates en-

capsulated in a shell of liquid-phase material. The model has been evaluated by comparison with reference computations based on observation-derived realistic models for encapsulated fractal aggregates (Kahnert et al., 2013). Item (i) has been incorporated in other CTMs earlier (e.g. Saide et al., 2013); however, to the best of our knowledge, items (ii) and (iii) go significantly beyond the current state-of-the-art of aerosol optics aerosol-optics models employed in CTMs. The main question of the present study is whether or not such a substantial level of detail in the description of aerosol morphology and optical properties is needed in a CTM.

To this end we compare radiative fluxes, backscattering coefficients, and Ångström exponents modelled with the old and new optics models We first performed a comparison of optical properties averaged over the entire model domain and over one month. To gauge the differences we observe between the new and the old optics model, we further compare two model versions of the CTM with different levels of detail in the aerosol process descriptions, namely, one version that includes aerosol dynamic aerosol-dynamic processes, and one a simpler mass-transport model, in which aerosol dynamics-microphysics is switched off. The importance of aerosol dynamics-microphysics is well understood an can therefore serve as a reference. The comparison showed that both for radiative fluxes, and for backscattering coefficients, and for Ångström exponents the differences. We found that the differences in optical properties between the two optics models are on the same order as those between the versions that include and exclude microphysical processes. For example, the aerosol optical depth computed with the two optics models differs by -25-18 %; differences obtained by inclusion or omission of aerosol microphysics are between -50-37 %. Corresponding differences in the backscattering coefficient are -8-99 % and -47-28 %, respectively. Analogous observations can be made for other radiometric properties.

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We further wanted to understand how the differences in optical properties impact radiative transfer processes in an aerosol-laden atmosphere. To this end we compare radiative fluxes modelled with the old and new optics models. The comparison showed that the differences in radiative net-fluxes between the two different optics models are of similar magnitude as corresponding differences between an aerosol dynamics and a the aerosol-microphysics and the mass-transport model. This strongly suggests that over-simplified aerosol optics models are likely to models.

These results strongly suggest that simplifications in the assumptions on aerosol morphology in the optics model can introduce substantial errors in modelled radiative fluxes and remote sensing-relevant observables. In Earth-system observables relevant to remote sensing. In chemistry-climate models such errors would enter into the simulation of the direct aerosol radiative forcing effect and add to all other sources of error in the model. In model evaluations that make use of remote sensing observations these errors would complicate the comparison between model results and observations.

The modifications to the optics model studied here morphology-assumptions in the optics model were limited to black carbon aerosols black-carbon particles. There are many other aerosols particles with complex morphological properties, such as mineral dust, which our optics model

still treats by an over-simplified homogeneous sphere a simple homogeneous-sphere model. The findings of our study should be an incentive for improving the description of dust and volcanic ash-volcanic-ash optical modelling in CTMs. A recent review of our current state of knowledge state-of-knowledge on aerosol morphology and aerosol optics for a variety of different aerosols has recently been reviewed aerosol particles can be found in Kahnert et al. (2014).

The findings of this study are likely to have implications for chemical data assimilation. In data assimilation one employs an *observation operator* that maps the model results to observable quantities. In case of satellite-based observations of aerosol optical properties, the observation operator is just our aerosol optics aerosol-optics model, possibly coupled to a radiative transfer model. Many data assimilation data-assimilation methodologies, such as the variational method, require a linear (or, at least, linearised) observation operator. In the old optics model, which assumes externally mixed aerosolsexternally-mixed aerosol particles, the observation operator is, indeed, linear (Kahnert, 2008). This largely explains why external-mixture optics models are widely used in chemical data assimilation systems (e.g. Kahnert, 2008; Benedetti et al., 2009; Liu et al., 2011). However, the new optics model we introduced here does not provide us with a linear map from the aerosol concentrations to the optical parameters. To what extend one could linearise this model and make use of its Jacobian in a data assimilation system mainly depends on the degree of nonlinearity, which would need to be investigated thoroughly.

All datasets used in this study, the MATCH model data and the aerosol optics data

5 Code availability

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The aerosol microphysics code SALSA is distributed under the Apache 2.0 license, while the chemistry transport model MATCH and the aerosol-optics data base are available upon request contacting the second author. from SMHI.

Appendix A: Size-averaged optical properties in the external-mixture optics model

The external-mixture optics model is based on using four size bins that cover the dry-radius intervals $[r^{\min}, r^{\max}]$ =[0.01, 0.05] μ m, [0.05, 0.5] μ m, [0.5, 1.25] μ m, and [1.25, 5] μ m. The geometric mean radius $R = \sqrt{r^{\min}r^{\max}} = 0.5(\log r^{\min} + \log r^{\max})$ is given in each of these intervals by R_1 =0.022 μ m, R_2 =0.158 μ m, R_3 =0.791 μ m, and R_4 =2.5 μ m. In each size bin it is assumed that the particle number density is given by a log-normal distribution

$$n_i(r) = N_i^0 / (\sqrt{2\pi}r \ln \sigma_i) \exp[-\ln^2(r/R_i) / (2\ln^2 \sigma_i)],$$
 (A1)

where $\sigma_1 = \sigma_3 = \sigma_4 = 1.8$, $\sigma_2 = 1.5$ are based on measurements in Neusüß et al. (2002). Here, N_i^0 would be the total number density per mode if each size-mode extended from r = 0 to $r = \infty$. However, since each mode is truncated at the bin-boundaries r^{\min} and r^{\max} , the number density N_i

of particles per size bin is obtained from integration over this finite interval, i.e.

$$N_{i} = \int_{r_{i}^{\min}}^{r_{i}^{\max}} n_{i}(r) dr = N_{i}^{0} \frac{1}{2} \left[\operatorname{erf}(x_{i}^{\max}) - \operatorname{erf}(x_{i}^{\min}) \right], \tag{A2}$$

where $x_i^{\text{max}} = \ln(r_i^{\text{max}}/R_i)/(\sqrt{2}\ln\sigma_i)$, and similarly for x_i^{min} . Analogously, one obtains the particle-mass density M_i in each size bin by integrating over the truncated log-normal mode, which yields

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$$M_i = \frac{4}{3}\pi\rho_i \int_{r_i^{\min}}^{r_i^{\min}} n_i(r)r^3 dr$$

$$= \frac{4}{3}\pi R_i^3 \rho_i N_i^0 \exp\left(\frac{9}{2}\ln^2 \sigma_i\right) \frac{1}{2} \left[\operatorname{erf}(y_i^{\max}) - \operatorname{erf}(y_i^{\min}) \right], \tag{A3}$$

where $y_i^{\text{max}} = x_i^{\text{max}} - 3\ln\sigma_i/\sqrt{2}$, and similarly for y_i^{min} , and where ρ_i is the density of the aerosol particles in the *i*th size bin. From this we obtain the desired relation for converting the mass-density M_i into the number-density N_i :

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$$N_i = \frac{M_i}{\frac{4}{3}\pi R_i^3 \rho_i} \cdot \frac{\operatorname{erf}(x_i^{\max}) - \operatorname{erf}(x_i^{\min})}{\exp\left(\frac{9}{2}\ln^2 \sigma_i\right) \left[\operatorname{erf}(y_i^{\max}) - \operatorname{erf}(y_i^{\min})\right]}.$$
 (A4)

Mass-mixing ratios X_i are simply converted into mass densities M_i according to $M_i = X_i \rho_{air}$, where ρ_{air} denotes the density of air.

In the external-mixture optics-database, optical properties are pre-computed by integrating optical properties at discrete sizes over the truncated log-normal size distribution. This integration is done numerically with a high size-resolution. The computation is performed for different refractive indices m, optical wavelengths λ , and for each size bin i. Thus, one obtains, e.g., extinction cross sections $C_{\text{ext}}(\lambda, m, i)$, which can be saved in a look-up table.

Secondary inorganic aerosols as well as organic aerosols and sea salt are assumed to be hydrophilic. We use the parameterisation by Gerber (1985) to compute the wet-radius r_w from the aerosol dry radius R, from which we obtain the volume-fraction of water $f_w = (r_w^3 - R^3)/R^3$. The effective refractive index m of the aerosol-water mixture is computed from that of the dry aerosol, m_a , and of water, m_w by use of effective-medium theory. In each grid cell, we obtain from the MATCH model, for each size bin i and for each aerosol component k, the number density $N_i(k)$. From that we compute the ensemble-averaged extinction cross section

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$$\bar{C}_{\text{ext}}(\lambda) = \frac{1}{N} \sum_{k} \sum_{i=1}^{4} N_i(k) C_{\text{ext}}(\lambda, m(k), i),$$
 (A5)

where the total number density is given by

$$N = \sum_{k} \sum_{i=1}^{4} N_i(k).$$
 (A6)

Note that the ensemble-average involves an average over both size and chemical composition. The ensemble-averaged scattering cross section $\bar{C}_{\rm sca}(\lambda)$ is computed analogously. From this we obtain the averaged single-scattering albedo 905

$$\bar{\omega}(\lambda) = \frac{\bar{C}_{\text{sca}}(\lambda)}{\bar{C}_{\text{ext}}(\lambda)}.$$
(A7)

The phase function $p(\Theta)$, hence its first Legendre-moment, known as the asymmetry parameter g are normalised quantities. Here Θ denotes the scattering angle. To average these quantities, one first needs to "de-normalise" by multiplying them with the scattering cross section. Thus

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$$\bar{p}(\Theta; \lambda) = \frac{1}{N\bar{C}_{\text{sca}}(\lambda)} \sum_{k} \sum_{i=1}^{4} N_i(k) C_{\text{sca}}(\lambda, m(k), i) p(\Theta, m(k), i; \lambda)$$
 (A8)

$$\bar{g}(\lambda) = \frac{1}{N\bar{C}_{\text{sca}}(\lambda)} \sum_{k} \sum_{i=1}^{4} N_i(k) C_{\text{sca}}(\lambda, m(k), i) g(m(k), i; \lambda). \tag{A9}$$

Once the ensemble-averaged optical properties in each grid cell of the model domain have been computed, one can compute radiometric observables, such as the extinction aerosol optical depth

$$\tau_{\text{ext}}(\lambda) = \sum_{z} N(z)\bar{C}_{\text{ext}}(\lambda, z)\Delta z,\tag{A10}$$

or the backscattering coefficient 915

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$$\beta_{\text{bak}}(\lambda, z) = \frac{1}{4\pi} N(z) \bar{C}_{\text{sca}}(\lambda, z) \bar{p}(180^{\circ}; \lambda, z), \tag{A11}$$

where z labels grid cells in the vertical column, and Δz denotes the layer-thickness.

Appendix B: Size-averaged optical properties in the internal-mixture model

In SALSA the number-density as a function of particle radius, n(r), is given by a step-function with $n_i(r) = \text{const}_i$ in each size bin i. This makes the pre-integration of optical properties over each size bin rather simple. On the other hand, we no longer assume that all aerosol components are externally mixed. Thus the ensemble-average over different chemical components k is no longer given by a simple summation $\sum_{k \dots n}$ as it was in the external-mixture model. Rather, for each size bin in which several aerosol components are internally mixed one computes an effective refractive index, $m_{\rm eff}$, by 925 use of effective-medium theory. One then reads the optical properties for that refractive index from the look-up table. Finally, one computes ensemble-averaged optical properties by summing over all size bins, $\sum_{i} \cdots$

Appendix C: Effective-medium theory

In effective-medium theory (EMT) one considers a composite material consisting of two materials with refractive indices m_1 and m_2 and volume fractions f_1 and $f_2 = 1 - f_1$. One then invokes assumptions about the topology of the mixture and derives a formula for the effective refractive index, $m_{\rm eff}$, of the composite material. For instance, it is often the case that $f_1 >> f_2$. In this case one can regard the first material as a host matrix that contains inclusions of the second material. This is the basis of the Maxwell-Garnett EMT (Maxwell Garnett, 1904). The resulting expression for $m_{\rm eff}$ is

$$m_{\text{eff}} = \sqrt{m_1^2 \frac{m_1^2 (2 - 2f_2) + m_2^2 (1 + 2f_2)}{m_1^2 (2 + f_2) + m_2^2 (1 - f_2)}}.$$
 (C1)

In the Bruggemann EMT (Bruggemann, 1935) one treats both materials more symmetrically; both components are assumed to be embedded in a host matrix with an effective refractive index given by

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$$m_{\text{eff}} = \sqrt{\frac{1}{4}m_1^2(2-3f_2) + m_2^2(3f_2-1) + \sqrt{\frac{1}{16}\left[m_1^2(2-3f_2) + m_2^2(3f_2-1)\right]^2 + \frac{1}{2}m_1^2m_2^2}}$$
 (C2)

Although not immediately manifest, this equation is symmetric under exchange of the two materials.

The volume-fraction is obtained from the mass concentrations M_1 and M_2 computed in the transport model, i.e. $f_2 = M_2/(M_1 + M_2)$. In SALSA, we apply the Maxwell-Garnett rule for an internal mixtures of mineral-dust inclusions in a host matrix of soluble compounds. Also, in the core-grey-shell model the effective refractive index of the grey shell, i.e., the homogeneous mixture of black carbon with soluble compounds, is computed with Maxwell-Garnett EMT. For mixtures of soluble compounds (sulphate, nitrate, ammonium, sea salt, organic compounds, and water) we use the Bruggemann EMT. If more than two components are mixed with each other, then the mixing rule is applied iteratively.

Appendix D: Refractive indices

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The refractive indices that are used in the new optics model (and in the effective-medium calculations) are listed in Table 8.

Appendix E: Optical properties at different wavelengths in the considered case-studies

955 Figures 12–35 show vertical profiles of AOD, single-scattering albedo, and asymmetry parameter at the four geographic location, for the summer and winter incident, and for 12 different optical wavelengths.

Table 8. Refractive index *m* for each wavelength in the aerosol-optics look-up table and for various aerosol components.

A Land 0.2000					
MIRC	$\stackrel{\lambda}{\sim} \stackrel{[\mu \mathrm{m}]}{\sim}$	0.2000	0.2316	0.3040	0.3400
	$\overset{m(SO_4)}{\sim}$	1.4840+0.1000E-07 i	1.4840+0.1000E-07 i	1.4676+0.1000E-07 i	1.4554+0.1000E-07 i
	∞ (BC)	0.9400+0.3500E+00 i	1.0717+0.5817E+00 i	1.3314+0.7523E+00 i	1.4471+0.7214E+00 i
Miles	∞	1.5300+0.5000E-02 i	1.5300+0.5000E-02 i	1.5300+0.5000E-02 i	1.5300+0.5000E-02 i
Miles	∞ (NaCl)	1.5100+0.5000E-05 i	1.5100+0.5000E-05 i	1.5100+0.1866E-05 i	1.5100+0.6592E-06 i
Marie	∞ (Dust)	1.5190+0.2070E-01 i	1.5190+0.2070E-01 i	1.5240+0.1947E-01 i	1.5272+0.1683E-01 i
	$\overset{m(\mathrm{H_2O})}{\sim}$	1.4517+0.1101E-06 i	1.4094+0.1092E-07 i	1.3701+0.3879E-08 i	1.3604+0.2758E-08 i
	N F 1	0.2550	0.2000	0.2022	0.4400
MIRC		$\sim\sim$	\sim	~~~	~~~
		~~~~~~	~~~~~~		
1,5000+0,2946-06    1,5000+0,1476-06    1,5014+0,6998-07    1,5000+0,2544-07    1,5000+0,2544-07    1,5000+0,2544-07    1,5000+0,2544-07    1,5000+0,2544-07    1,5000+0,2544-07    1,5000+0,2544-07    1,5000+0,2544-07    1,5000+0,2544-07    1,5000+0,2544-07    1,5000+0,2544-07    1,5000+0,2544-07    1,5000+0,2544-07    1,5000+0,2544-07    1,5000+0,2544-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,5000+0,2500-07    1,50	,0000	,00000000	~~~~~~	~~~~~	~~~~~
1,5339+0,1250E011	,000	~~~~~~	~~~~~~	~~~~~	~~~~~
Milbo		~~~~~~	~~~~~	~~~~~	~~~~~~
	,000	,00000000	,00000000	,00000000	,00000000
M(SO ₂ )	$\overset{m(\mathrm{H}_2\mathrm{O})}{\sim}$	1.3572+0.2416E-08 i	1.3528+0.1944E-08 i	1.3508+0.1702E-08 i	1.3449+0.9324E-09 i
Table   Tabl	$\lambda [\mu m]$	0.5000	0.5320	0.5332	0.6750
m(BC		1.4310+0.1000E-07 i	1.4304+0.1000E-07 i	1.4303+0.1000E-07 i	1.4285+0.1860E-07 i
1,5300+0,5500E-021   1,5300+0,5500E-021   1,5300+0,5500E-021   1,5300+0,05500E-021   1	,000	~~~~~	~~~~~	~~~~~	~~~~~
M(SC)   15000+0.1550E-071   15000+0.1198E-071   15000+0.1198E-071   14900+0.1212E-061   15160+0.1200E-021   15160+0.1200E-021   15160+0.1200E-021   15170+0.9818E-031   15170+0.9918E-031   15170+0.9918E-0	$\sim\sim$	~~~~~	~~~~~	~~~~~	~~~~~
Miles	,000	~~~~~	~~~~~	~~~~~	~~~~~
MITTO   1.339440.9243E-091   1.337140.1818E-081   1.337040.1850E-081   1.329740.2187E-071	$\sim\sim$	~~~~~	~~~~~	~~~~~	~~~~~
\( \lambda \)   \( \lambda			.00000000	,00000000	,00000000
m(SO ₄ )         1.4280+0.2214E-07;         1.4253+0.2044E-06;         1.4216+0.1749E-05;         1.4211+0.1963E-05;           m(BC)         1.8175+0.5730E+00;         1.8752+0.564SE+00;         1.9210+0.562E+00;         1.9219+0.564SE+00;           m(DC)         1.5300+0.7333E-02;         1.5300+0.327E-01;         1.5300+0.1327E-01;         1.5300+0.1327E-01;           m(NaCl)         1.4900+0.2282E-06;         1.4800+0.3027E-04;         1.4700+0.1498E-03;         1.4700+0.1584E-03;           m(Dust)         1.5170+0.9333E-03;         1.5184+0.8000E-03;         1.5190+0.7347E-03;         1.5190+0.7261E-03;           m(H ₂ O)         1.3287+0.3624E-07;         1.3243+0.3714E-06;         1.3215+0.2657E-05;         1.3213+0.2380E-05;           λ [µm]         1.0640         1.2705         1.4625         1.7840           m(SO ₄ )         1.4197+0.2915E-05;         1.4122+0.1621E-04;         1.4045+0.1030E-03;         1.3213+0.2380E-03;           m(BC)         1.9261+0.5738E+00;         1.9457+0.6188E+00;         1.9639+0.6597E+00;         1.9943+0.7290E+00;           m(NaCl)         1.4700+0.1966E-03;         1.4692+0.3754E-03;         1.4615+0.5332E-03;         1.4800+0.1337E-01;           m(Dust)         1.5190+0.683E-03;         1.5188+0.6418E-03;         1.5180+0.8000E-03;         1.5180+0.9990E-03;           m(H ₂ O)<	~~~		~~~~~	~~~~~	
m(BC)         1,8175+0.5730E+001         1,8752+0.5645E+001         1,9210+0.562E+001         1,9210+0.562E+001           m(OC)         1,5300+0.7333E+02 i         1,5300+0.9409E+02 i         1,5300+0.137E+01 i         1,5300+0.1370E+01 i           m(NsCl)         1,4900+0.2282E+06 i         1,4800+0.3027E+04 i         1,4700+0.1498E+03 i         1,4700+0.1584E+03 i           m(Dust)         1,5170+0.9335E+03 i         1,5184+0.8000E+03 i         1,5190+0.7347E+03 i         1,5190+0.7241E+03 i           M(H₂O)         1,3287+0.3624E+07 i         1,3243+0.3714E+06 i         1,3215+0.2657E+05 i         1,3213+0.2380E+05 i           M(BC)         1,926+0.5738E+00 i         1,9457+0.6188E+00 i         1,9639+0.6597E+00 i         1,9943+0.7290E+00 i           m(BC)         1,926+0.5738E+00 i         1,9457+0.6188E+00 i         1,9639+0.6597E+00 i         1,9943+0.7290E+00 i           m(Dust)         1,5190+0.0853E+03 i         1,5190+0.1721E+01 i         1,5088+0.1864E+01 i         1,4801+0.1337E+01 i           m(Dust)         1,5190+0.0853E+03 i         1,5180+0.2373E+03 i         1,4615+0.5332E+03 i         1,5180+0.9990E+03 i           m(BC)         2,0292+0.7884E+00 i         2,3250         2,7885         3,4615           m(BC)         2,0192+0.7884E+00 i         2,010+0.48453E+00 i         1,3144+0.2569E+01 i         1,3600+0.1779E+00 i	$\frac{\lambda}{\sim}$ [ $\mu$ m]	0.7016	0.8700	1.0101	1.0200
MOC	$\underset{\sim}{\text{m(SO_4)}}$	,00000000	1.4253+0.2044E-06 i	1.4216+0.1749E-05 i	1.4213+0.1963E-05 i
m(NaCl)         1.4900+0.2282E-061         1.4800+0.3027E-041         1.4700+0.198E-031         1.4700+0.188E-031           m(Dost)         1.5170+0.9335E-031         1.5184+0.8000E-031         1.5190+0.734TE-031         1.5190+0.724TE-031           m(H ₂ O)         1.3287+0.3624E-071         1.3243+0.3714E-061         1.3215+0.2657E-051         1.3213+0.2380E-051           λ [µm]         1.0640         1.2705         1.4625         1.7840           m(SO ₄ )         1.4197+0.2915E-051         1.412240.1621E-041         1.4045+0.1030E-031         1.3926+0.5308E-031           m(BC)         1.9261+0.5738E+001         1.9457+0.6183E+001         1.9639+0.6597E+001         1.9943+0.7290E+001           m(BC)         1.5285+0.1515E-011         1.5179+0.1721E-011         1.5066+0.1864E-011         1.4801+0.1337E-011           m(NaCl)         1.4700+0.1966E-031         1.4692+0.3754E-031         1.4615+0.5382E-031         1.4500+0.7944E-031           m(Dust)         1.5190+0.8833E-031         1.5188-0.6418E-031         1.5180+0.8000E-031         1.5180+0.9990E-031           m(H ₂ O)         1.3205+0.1279E-051         1.31674-0.1990E-041         1.3128+0.3528E-031         1.3040+0.1270E-031           λ [µm]         2.0460         2.3250         2.7885         3.4615           m(SO ₄ )         1.3803+0.1490E-021	$\underbrace{m(\mathrm{BC})}_{}$	1.8175+0.5730E+00 i	1.8752+0.5645E+00 i	1.9210+0.5622E+00 i	1.9219+0.5643E+00 i
m(Dbs)         1,5170+0,9335E-031         1,5184+0,8000E-031         1,5190+0,7347E-031         1,5190+0,7247E-031           m(H₂O)         1,3287+0,3624E-071         1,3243+0,3714E-061         1,3215+0,2657E-051         1,3213+0,2380E-051           λ [µm]         1,0640         1,2705         1,4625         1,7840           m(SO4)         1,41974-0,2915E-051         1,4927+0,6183E+001         1,9457+0,6183E+001         1,4455+0,5382E+001         1,4450+0,734E+031         1,450+0,734E+031         1,450+0,734E+031         1,450+0,734E+031         1,450+0,734E+031         1,450+0,734E+031         1,450+0,734E+031         1,5180+0,990E-031         1,15180+0,990E-031         1,13180+0,8452E+031         1,3140+0,560E+011 <td>$\infty$ (OC)</td> <td>1.5300+0.7333E-02 i</td> <td>1.5300+0.9409E-02 i</td> <td>1.5300+0.1327E-01 i</td> <td>1.5300+0.1370E-01 i</td>	$\infty$ (OC)	1.5300+0.7333E-02 i	1.5300+0.9409E-02 i	1.5300+0.1327E-01 i	1.5300+0.1370E-01 i
M(H2O)   13287403624E-071   1324340.3714E-061   1321540.2657E-051   1321340.2380E-051	$\underbrace{m(\text{NaCl})}$	1.4900+0.2282E-06 i	1.4800+0.3027E-04 i	1.4700+0.1498E-03 i	1.4700+0.1584E-03 i
\$ [µm]         1,0640         1,2705         1,4625         1,7840           m(SO4)         1,447+0,2915E-05i         1,4422+0.1631E-04i         1,4045+0.1030E-03i         1,3926+0.5308E-03i           m(BC)         1,9261+0.5738E+00j         1,9457+0.6183E+00j         1,9639+0.6597E+00j         1,993+0.7290E+00j           m(OC)         1,5285+0.1515E-01i         1,5179+0.1721E-01i         1,5068+0.1864E-01i         1,4801+0.1337E-01i           m(NaCl)         1,4700+0.1966E-03i         1,5189+0.47354E-03i         1,4500+0.7944E-03i         1,5180+0.8000E-03i         1,5180+0.9990E-03i           m(H₂O)         1,3205+0.1279E-05i         1,3167+0.199E-04i         1,3128+0.3528E-03i         1,3460+0.796E-03i           λ [µm]         2,0460         2,3250         2,7885         3,4615           m(SO4)         1,3803+0.1490E-02i         1,3380+0.2885E-02i         1,3146+0.5669E-01i         1,3669+0.1579E+00i           m(BC)         2,019240.7854E+00i         2,0510+0.8453E+00i         2,1994-0.944E+00i         2,1955+0.1088E+01i           m(BC)         1,4613+0.1000E-01i         1,4554+0.9641E-02i         1,4800+0.7724E-02i         1,4800+0.7794E-02i           m(So4)         1,3180+0.1492E-02i         1,4770+0.2950E-02i         1,5180+0.9077E-02i         1,5180+0.2805E-01i           m(BC)         1,2947+0.	$\underbrace{m({ m Dust})}$	1.5170+0.9335E-03 i	1.5184+0.8000E-03 i	1.5190+0.7347E-03 i	1.5190+0.7261E-03 i
m(SO ₄ )         1.4497+0.2915E-05 i         1.4422+0.1621E-04 i         1.4484+0.1030E-03 i         1.3926+0.5308E-03 i           m(BC)         1.9261+0.5738E+00 i         1.9457+0.6183E+00 i         1.9639+0.6597E+00 i         1.9943+0.7290E+00 i           m(DC)         1.5285+0.1515E-01 i         1.5179+0.1721E-01 i         1.5008+0.1864E-01 i         1.4801+0.1337E-01 i           m(NaCl)         1.4700+0.1966E-03 i         1.4692+0.3754E-03 i         1.4615+0.5382E-03 i         1.4500+0.7944E-03 i           m(Dust)         1.5199+0.0853E-03 i         1.5188+0.6418E-03 i         1.5180+0.8000E-03 i         1.5180+0.9990E-03 i           m(H₂O)         1.3205+0.1279E-05 i         1.3167+0.1090E-04 i         1.3128+0.3528E-03 i         1.3040+0.1270E-03 i           λ [μm]         2.0460         2.3250         2.7885         3.4615           m(SO ₄ )         1.3803+0.1490E-02 i         1.3380+0.2885E-02 i         1.3146+0.5669E-01 i         1.3669+0.1579E+00 i           m(BC)         2.0192+0.7854E+00 i         2.0510+0.8453E+00 i         2.1099+0.9444E+00 i         2.1955+0.1888E+01 i           m(OC)         4613+0.1090E-02 i         1.4554+0.9641E-02 i         1.4800+0.7724E-02 i         1.4800+0.1757E-02 i           m(Dust)         1.5180+0.1492E-02 i         1.5180+0.2805E-00 i         1.5180+0.2805E-00 i         1.1278+0.1055E+00 i	$\stackrel{m({\rm H_2O})}{\sim\!\!\!\sim\!\!\!\sim}$	1.3287+0.3624E-07 i	1.3243+0.3714E-06 i	1.3215+0.2657E-05 i	1.3213+0.2380E-05 i
m(SO ₄ )         1.4497+0.2915E-05 i         1.4422+0.1621E-04 i         1.4045+0.1030E-03 i         1.3926+0.5308E-03 i           m(BC)         1.9261+0.5738E+00 i         1.9457+0.6183E+00 i         1.9639+0.6597E+00 i         1.993+0.7290E+00 i           m(OC)         1.5285+0.1515E-01 i         1.5179+0.1721E-01 i         1.5068+0.1864E-01 i         1.4801+0.1337E-01 i           m(NaCl)         1.4700+0.1966E-03 i         1.4692+0.3754E-03 i         1.4615+0.5382E-03 i         1.4500+0.7944E-03 i           m(Dust)         1.5199+0.6853E-03 i         1.5188+0.6418E-03 i         1.5180+0.8000E-03 i         1.5180+0.9990E-03 i           m(H₂O)         1.3205+0.1279E-05 i         1.3167+0.1090E-04 i         1.3128+0.3528E-03 i         1.3040+0.1270E-03 i           λ [μm]         2.0460         2.3250         2.7885         3.4615           m(SO ₄ )         1.3803+0.1490E-02 i         1.3880+0.2885E-02 i         1.3146+0.5669E-01 i         1.3669+0.1579E+00 i           m(BC)         2.0192+0.7854E+00 i         2.0510+0.8453E+00 i         2.1099+0.9444E+00 i         2.1955+0.1088E+01 i           m(OC)         1.4613+0.1000E-01 i         1.4554+0.9641E-02 i         1.4800+0.7724E-02 i         1.4800+0.1757E-02 i           m(Dust)         1.5180+0.1492E-02 i         1.5180+0.2904E-02 i         1.5180+0.2904E-02 i         1.5180+0.2904E-02 i	λ [μm]	1.0640	1.2705	1.4625	1.7840
m(BC)         1.9261+0.5738E+001         1.9457+0.6183E+001         1.9639+0.6597E+001         1.9943+0.7290E+001           m(OC)         1.5285+0.1515E-01i         1.5179+0.1721E-01i         1.5068+0.1864E-01i         1.4801+0.133TE-01i           m(NaCl)         1.4700+0.1966E-03i         1.4692+0.3754E-03i         1.4615+0.5332E-03i         1.4500+0.7944E-03i           m(Dust)         1.5190+0.6853E-03i         1.5188+0.6418E-03i         1.5180+0.8000E-03i         1.5180+0.9990E-03i           m(H₂O)         1.3205+0.1279E-05i         1.3167+0.1090E-04i         1.3128+0.3528E-03i         1.3040+0.1270E-03i           λ [μm]         2.0460         2.3250         2.7885         3.4615           m(SO ₄ )         1.3803+0.1490E-02i         1.3380+0.2885E-02i         1.3144+0.5669E-01i         1.3669+0.1579E+00i           m(BC)         2.0192+0.7854E+00i         2.010+0.8453E+00i         2.1099+0.9444E+00i         2.1955+0.1088E+01i           m(SO ₄ )         1.3480+0.1490E-02i         1.4554+0.9641E-02i         1.4800+0.7734E-02i         1.4800+0.1757E-02i           m(NaCl)         1.4482+0.1276E-02i         1.5180+0.290E-02i         1.5180+0.807TE-02i         1.5180+0.2805E-01i           m(H₂O)         1.2947+0.7103E-03i         1.2756+0.5344E-03i         1.1278+0.1055E+00i         1.3913+0.1237E-01i           λ [μm] <td></td> <td>1.4197+0.2915E-05 i</td> <td>1.4122+0.1621E-04 i</td> <td>1.4045+0.1030E-03 i</td> <td>1.3926+0.5308E-03 i</td>		1.4197+0.2915E-05 i	1.4122+0.1621E-04 i	1.4045+0.1030E-03 i	1.3926+0.5308E-03 i
m(OC)         1.5285+0.1515E-01;         1.5179+0.1721E-01;         1.5068+0.1864E-01;         1.4801+0.1337E-01;           m(NaCl)         1.4700+0.1966E-03;         1.4692+0.3754E-03;         1.4615+0.5382E-03;         1.4500+0.7944E-03;           m(Dust)         1.5190+0.6853E-03;         1.5188+0.6418E-03;         1.5180+0.8000E-03;         1.5180+0.9990E-03;           m(H₂O)         1.3205+0.1279E-05;         1.31674-0.1990E-04;         1.3128+0.3528E-03;         1.3040+0.1270E-03;           λ [µm]         2.0460         2.3250         2.7885         3.4615           m(SO4)         1.3803+0.1490E-02;         1.3380+0.2885E-02;         1.3146+0.5669E-01;         1.3669+0.1579E+00;           m(CC)         1.4613+0.1000E-01;         1.4554+0.9641E-02;         1.4800+0.7724E-02;         1.4800+0.7000E-02;           m(NaCl)         1.4482+0.1276E-02;         1.4370+0.2950E-02;         1.5339+0.7462E-02;         1.4800+0.1757E-02;           m(Dust)         1.5180+0.1492E-02;         1.5180+0.2610E-02;         1.5180+0.8077E-02;         1.5180+0.2805E-01;           λ [µm]         3.5000         8.0205         10.6000         12.1950           m(SO4)         1.3760+0.1580E+00;         1.1641+0.5511E+00;         1.7200+0.3400E+00;         1.7858+0.2517E+00;           m(SO4)         1.3760+0.1580E+00;		,00000000	,00000000	,00000000	,00000000
m(NaCl)         1.4700+0.1966E-031         1.4692+0.3734E-031         1.4615+0.5382E-031         1.4500+0.7944E-031           m(Dast)         1.5190+0.6853E-031         1.5188+0.6418E-031         1.5180+0.9000E-031         1.5180+0.9900E-031           m(H₂O)         1.3205+0.1279E-051         1.31674-0.1900E-041         1.3128+0.3528E-031         1.3040+0.1270E-031           λ [µm]         2.0460         2.3250         2.7885         3.4615           m(SO ₄ )         1.3803+0.1490E-021         1.3380+0.2885E-021         1.3146+0.5669E-011         1.3669+0.1579E+001           m(BC)         2.0192+0.7854E+001         2.0510+0.8453E+001         2.1099+0.9444E+001         2.1955+0.1088E+011           m(OC)         1.4413+0.1000E-011         1.4554+0.9641E-021         1.4800+0.7724E-021         1.4800+0.7734E-021           m(NaCl)         1.4482+0.1276E-021         1.4370+0.2950E-021         1.5339+0.7462E-021         1.4800+0.775E-021           m(Dast)         1.5180+0.1492E-021         1.5180+0.2805E-011         1.2756+0.534E-031         1.1278+0.1055E+001         1.3913+0.1237E-011           λ [µm]         3.5000         8.0205         10.6000         12.1950           m(SO ₄ )         1.3760+0.1580E+001         1.1641+0.5511E+001         1.7200+0.3400E+001         1.7858+0.2517E+001           m(SO ₄ )		,00000000	.00000000	,00000000	,00000000
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		,00000000	.00000000	,00000000	,00000000
$\begin{array}{cccccccccccccccccccccccccccccccccccc$			.00000000	,00000000	,00000000
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	$\sim\sim$	~~~~~	~~~~~	1.3128+0.3528E-03 i	1.3040+0.1270E-03 i
m(SO ₄ )         1,3803+0.1490E-02 i         1,3800+0.2885E-02 i         1,3146+0.5669E-01 i         1,3669+0.1579E+00 i           m(BC)         2,0192+0.7884E+00 i         2,0510+0.8453E+00 i         2,1099+0.944E+00 i         2,1955+0.1088E+01 i           m(CC)         1,4613+0.1000E-01 i         1,4554+0.9641E-02 i         1,4800+0.7724E-02 i         1,4800+0.7000E-02 i           m(NaCl)         1,4482+0.1276E-02 i         1,4370+0.2950E-02 i         1,5339+0.7462E-02 i         1,4800+0.1757E-02 i           m(Dust)         1,5180+0.1492E-02 i         1,5180+0.2610E-02 i         1,5180+0.8077E-02 i         1,5180+0.2805E-01 i           m(H ₂ O)         1,2947+0.7103E-03 i         1,2756+0.534E-03 i         1,1278+0.1055E+00 i         1,3913+0.1237E-01 i           λ [μm]         3,5000         8,0205         10,6000         12,1950           m(SO ₄ )         1,3760+0.1580E+00 i         1,1641+0.5511E+00 i         1,7200+0.3400E+00 i         1,7858+0.2517E+00 i           m(BC)         2,2004+0.1097E+01 i         2,6572+0.1742E+01 i         2,9285+0.2063E+01 i         3,0719+0.2210E+01 i           m(NaCl)         1,4800+0.1600E-02 i         1,4080+0.1581E-01 i         1,5000+0.1400E-01 i         1,6332+0.5117E-01 i           m(Dust)         1,5180+0.2973E-01 i         1,1798+0.1015E+00 i         1,9100+0.2500E+00 i         1,7614+0.4543E+00 i </td <td>~~~</td> <td></td> <td></td> <td></td> <td></td>	~~~				
m(BC)         2.0192407854E+001         2.051040.8455E+001         2.109940.944E+001         2.109540.1088E+011           m(CC)         1.461340.1000E-011         1.455440.9641E-021         1.480040.7724E-021         1.480040.7000E-021           m(NsCl)         1.448240.1276E-021         1.437040.2950E-021         1.533940.7462E-021         1.480040.1757E-021           m(Dust)         1.518040.1492E-021         1.518040.2600E-021         1.518040.8077E-021         1.518040.2805E-011           m(H ₂ O)         1.294740.7103E-031         1.275640.5344E-031         1.127840.1055E+001         1.391340.1237E-011           λ [µm]         3.5000         8.0205         10.6000         12.1950           m(SO ₄ )         1.3760+0.1580E+001         1.1641+0.5511E+001         1.7200+0.3400E+001         1.7858+0.2517E+001           m(BC)         2.2004+0.1097E+011         2.6572+0.1742E+011         2.9285+0.2063E+011         3.0719+0.2210E+011           m(NaCl)         1.4800+0.1600E-021         1.4080+0.1581E-011         1.5000+0.1400E-011         1.6352+0.5117E-011           m(Dust)         1.5180+0.2973E-011         1.179840.1015E+001         1.9100+0.2500E+001         1.7614+0.4543E+001	$\sim$ $\sim$	$\sim\sim$	~~~	$\sim\sim$	$\sim\sim$
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\sim\sim$	~~~~~	~~~~~	~~~~~	~~~~~
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\overset{m{m}(\mathrm{BC})}{\sim\!\!\!\sim\!\!\!\sim}$	~~~~~	~~~~~	~~~~~	~~~~~
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\stackrel{m({ m OC})}{\sim}$	1.4613+0.1000E-01 i	~~~~~	1.4800+0.7724E-02 i	1.4800+0.7000E-02 i
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\underbrace{m( ext{NaCl})}$	1.4482+0.1276E-02 i	1.4370+0.2950E-02 i	1.5339+0.7462E-02 i	1.4800+0.1757E-02 i
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	$ \underbrace{m(\text{Dust})} $	,00000000	,00000000	,00000000	,00000000
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\overset{m(\mathrm{H}_2\mathrm{O})}{\sim}\!\!\!\!\!\sim$	1.2947+0.7103E-03 i	1.2756+0.5344E-03 i	1.1278+0.1055E+00 i	1.3913+0.1237E-01 i
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	λ [μm]	3.5000	8.0205	10.6000	12.1950
m(BC)         2,2004+0.1097E+01 i         2,6572+0.1742E+01 i         2,9285+0.2063E+01 i         3,0719+0.2210E+01 i           m(OC)         1,4800+0.7000E+02 i         1,1237+0.7906E+01 i         1,7600+0.7000E+01 i         1,6352+0.5117E+01 i           m(NaCl)         1,4800+0.1600E+02 i         1,4080+0.1581E+01 i         1,5000+0.1400E+01 i         1,4383+0.1539E+01 i           m(Dust)         1,5180+0.2973E+01 i         1,1798+0.1015E+00 i         1,9100+0.2500E+00 i         1,7614+0.4543E+00 i		,000	~~~	~~~	~~~
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m(NaCl)     1.4800+0.1600E-02 i     1.4800+0.1581E-01 i     1.5000+0.1400E-01 i     1.4833+0.1539E-01 i       m(Dust)     1.5180+0.2973E-01 i     1.1798+0.1015E+00 i     1.9100+0.2500E+00 i     1.7614+0.4543E+00 i		~~~~~~	~~~~~~	~~~~~	~~~~~~
m(Dust) 1.5180+0.2973E-01 1.1798+0.1015E+00i 1.9100+0.2500E+00i 1.7614+0.4543E+00i	,0000	~~~~~~	~~~~~~	~~~~~	~~~~~~
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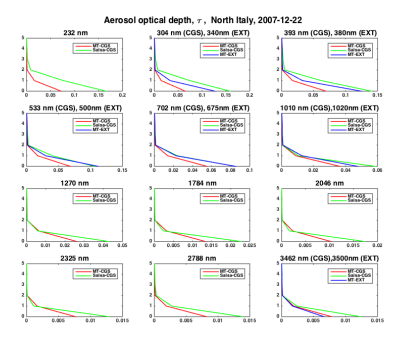


Figure 12. Aerosol optical depth over North Italy at 22-12-2007 for the 12 wavelengths in the CGS optics model and 5 of the 7 wavelengths in the EXT model. The wavelengths do not exactly overlap, but the EXT wavelengths that lies within 40 nm of the CGS wavelength are plotted in the same graph.

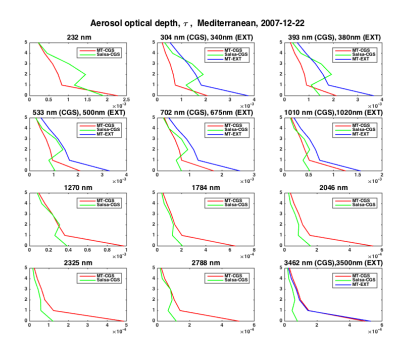


Figure 13. Same as Fig. 12, but over the Mediterranean.

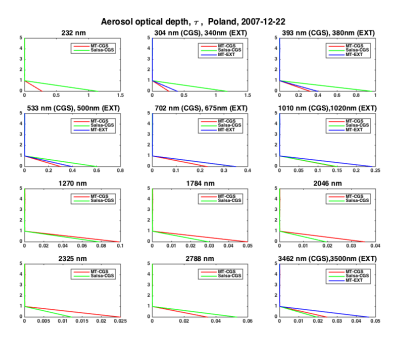


Figure 14. Same as Fig. 12, but over Poland.

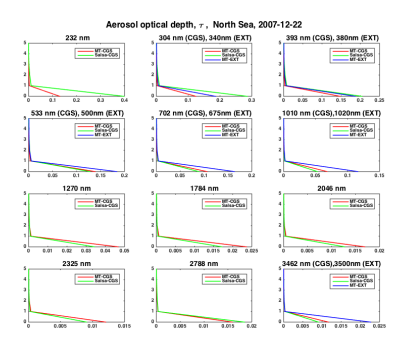


Figure 15. Same as Fig. 12, but over the North Sea

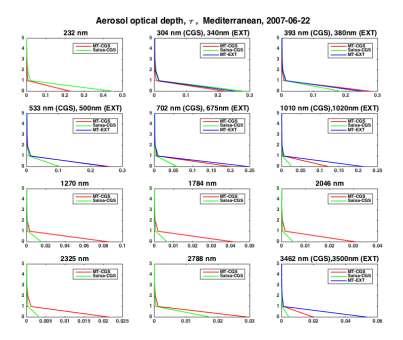


Figure 16. Same as Fig. 12, but 2007-06-22.

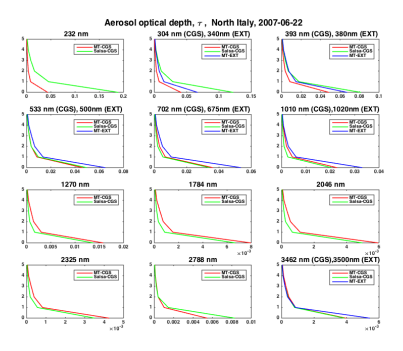


Figure 17. Same as Fig. 12, but 2007-06-22 and over the Mediterranean.

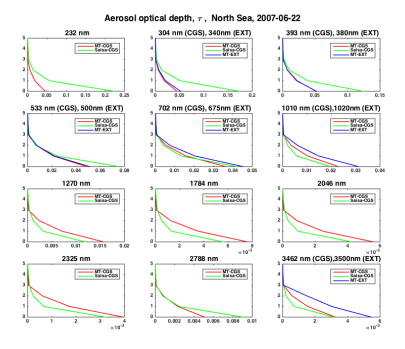


Figure 18. Same as Fig. 12, but 2007-06-22 and over Poland.

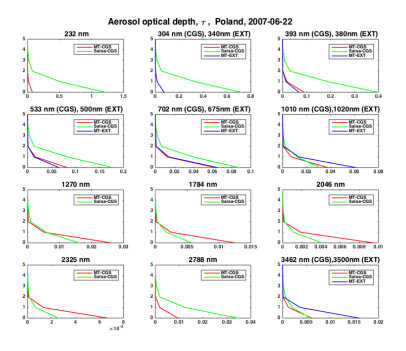


Figure 19. Same as Fig. 12, but 2007-06-22 and over the North sea.

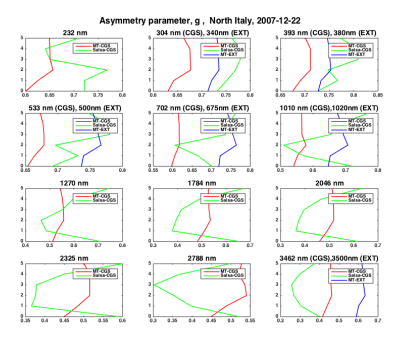


Figure 20. Asymmetry parameter over North Italy at 22-12-2007 for the 12 wavelengths in the CGS optics model and 5 of the 7 wavelengths in the EXT model. The wavelengths do not exactly overlap, but the EXT wavelengths that lies within 40 nm of the CGS wavelength are plotted in the same graph.

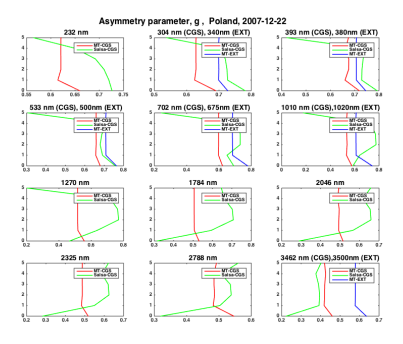


Figure 21. Same as Fig. 20, but over the Mediterranean.

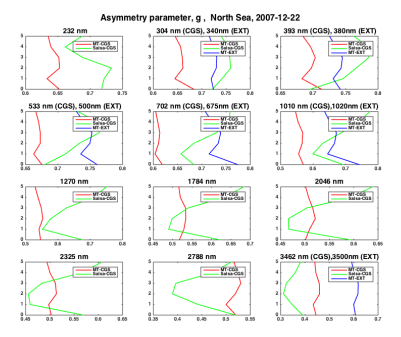


Figure 22. Same as Fig. 20, but over Poland.

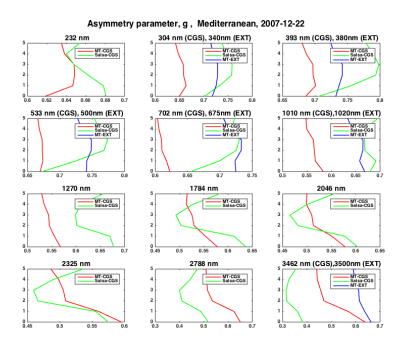


Figure 23. Same as Fig. 20, but over the North sea.

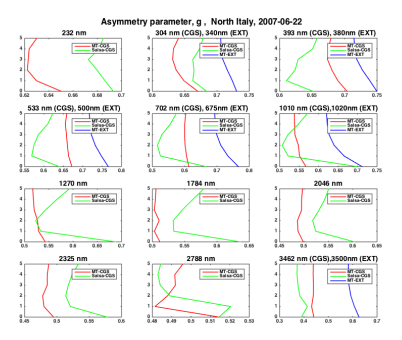


Figure 24. Same as Fig. 20, but 2007-06-22.

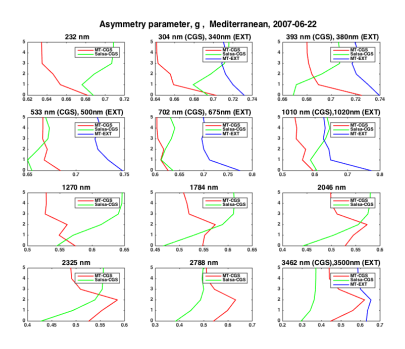


Figure 25. Same as Fig. 20, but 2007-06-22 and over the Mediterranean.

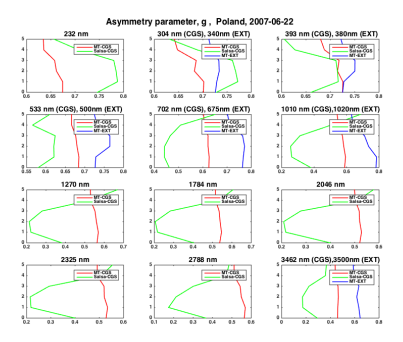


Figure 26. Same as Fig. 20, but 2007-06-22 and over Poland.

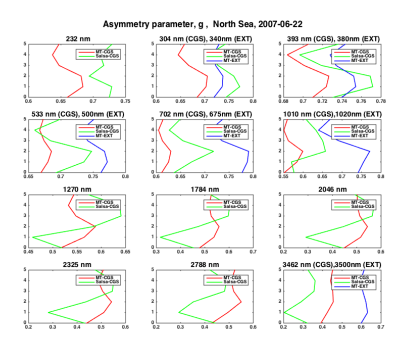


Figure 27. Same as Fig. 20, but 2007-06-22 and over the North sea.

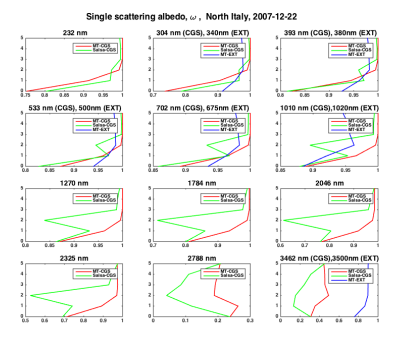


Figure 28. Single scattering albedo over North Italy at 22-12-2007 for the 12 wavelengths in the CGS optics model and 5 of the 7 wavelengths in the EXT model. The wavelengths do not exactly overlap, but the EXT wavelengths that lies within 40 nm of the CGS wavelength are plotted in the same graph.

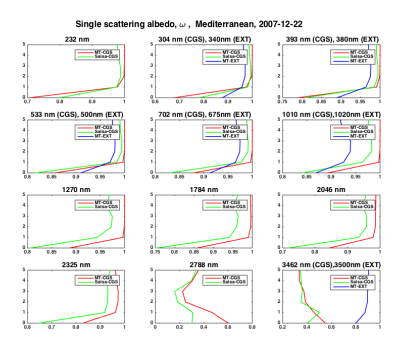


Figure 29. Same as Fig. 28, but over the Mediterranean.

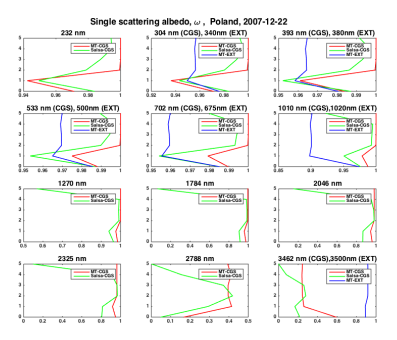


Figure 30. Same as Fig. 28, but over Poland.

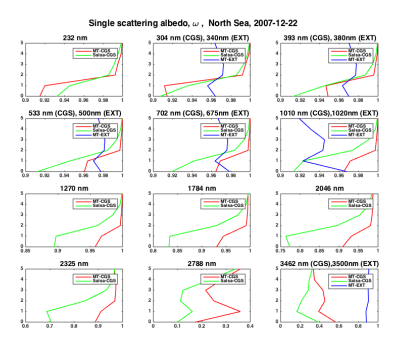


Figure 31. Same as Fig. 28, but over the North sea.

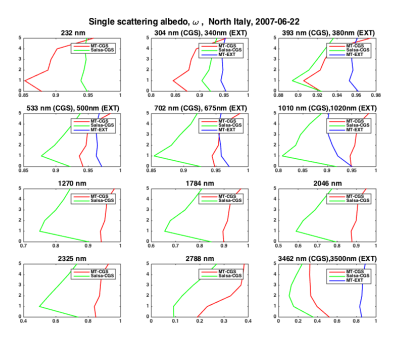


Figure 32. Same as Fig. 28, but 2007-06-22.

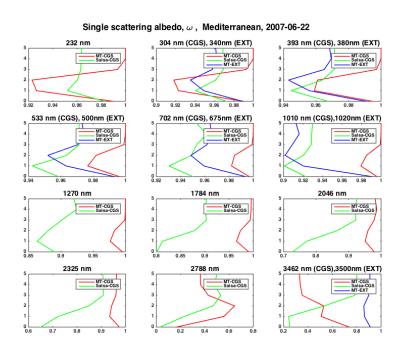


Figure 33. Same as Fig. 28, but 2007-06-22 and over the Mediterranean.

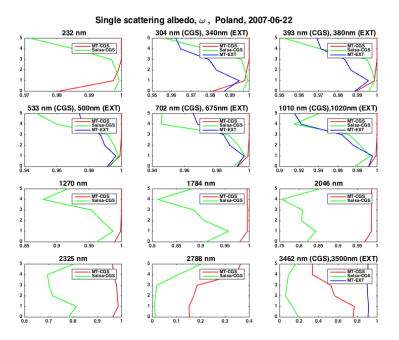


Figure 34. Same as Fig. 28, but 2007-06-22 and over Poland.

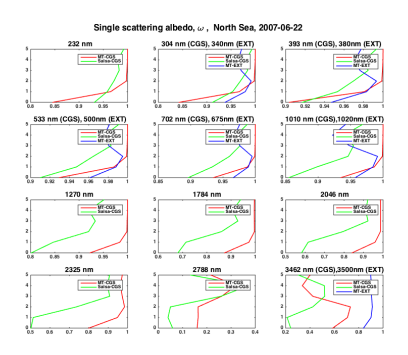


Figure 35. Same as Fig. 28, but 2007-06-22 and over the North sea.

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