Response to Reviewer 1 comments

We thank this reviewer for their supportive and helpful comments. Below we respond to each comment individually.

We note that this reviewer has used line numbers from the version of the manuscript submitted for the technical review, rather than the version of the manuscript openly published in the Discussion phase.

The reviewer’s comments are in italics and blue font, our responses are in normal text.

This is an interesting and generally well-written paper which draws attention to another potentially important source of SOA in the atmosphere. The paper makes nice use of high-quality atmospheric data-sets, and uses not just OA measurements but also other gas and particle data to build confidence in the model. The main conclusions are important: that emissions of IVOC from diesel and their subsequent SOA formation might be much more important than previously assumed.

There are some issues however that I think the authors need to deal with before acceptance for ACP. The main issues I see are:

1. The paper begins (Fig.1, also Sect. 2.5) with an example of a so-called add1.5xPOA approach of illustrating the importance of the 1.5 x PM assumption for IVOC, contrasting with the new addDiesel approach used in this paper (Fig.2). We then see many results from the base and addDiesel cases, but have to wait until almost the very end before seeing some annual-average result from the add1.5xPOA case. The 1.5xPM approach used here though differs from that of previous authors, e.g.Robinson et al. (2007), Shrivastava et al. (2008) or even the Bergstrom et al. (2012) paper. In the add1.5xPOA used here, the authors still seem to assume the same inert POA emission as in the base-case, but add IVOC with a very high C* value. If correct, this is a significant deviation from the other studies, which added SVOC and IVOC across a range of volatilities. If the authors really did just use one IVOC component at C*=1.0e5, then this will lead to more POA close to sources and less downwind compared to e.g. Robinson. This begs the question, would a more ‘traditional’ (Robinson-like) add1.5xPOA scheme give results that might anyway have been better compared to observations than the Base-case used here? This would not mean that diesel-IVOC isn’t important, but it might have qualified the relative importance. Of course, one is adding less IVOC and hence producing less SOA. On the other hand, an 1.5xPOA approach (with both SVOC and IVOC) would have generated a bigger gradient between London and the outlying sites, perhaps in better agreement with the observed gradients.

Concluding, I think they could be better off either (i) re-running with the (dare-I-say) ‘traditional’ 1.5xPOA for SVOC+IVOC, or (ii) just skipping this test altogether.
Response: We have now undertaken the reviewer’s suggestion (i) and run a model experiment identical to Shrivastava et al. 2008. We have amended text accordingly in both the Methods and in the Results (Sect. 3.6, Fig. 18). The relevant text in the Methods now reads:

“Nevertheless, we have also performed a model run using the POA-based IVOC estimate, also including the semivolatile treatment of POA. The emitted semivolatile POA (SVOCs) and 1.5xPOA IVOCs are assigned to 9 VBS bins: 0.03xPOA, 0.06xPOA, 0.09xPOA, 0.14xPOA, 0.18xPOA, 0.30xPOA, 0.40xPOA, 0.50xPOA, 0.80xPOA to the bins 0.01–10⁻⁶, respectively; totalling 2.5xPOA (Shrivastava et al. 2008). Both SVOCs and IVOCs then go through atmospheric ageing with OH (k = 4.0 x 10⁻¹¹ cm³ molecule⁻¹ s⁻¹; Shrivastava et al. (2008)). /…/ SVOCs and IVOCs that have undergone at least one ageing shift and are in the particulate phase are included under SOA (in addition to ASOA and BSOA from VOCs as in the Base case).”

And Sect. 3.6 now reads:

“3.6 Comparison to the previous (IVOCs=1.5xPOA) approach

Figure 18 shows the annual average HOA, SFOA, BSOA and Background OA (BGND OA), and ASOA concentrations at London North Kensington modelled with different assumptions for additional IVOC emissions. As was explained in Sect. 2.4, for the UK, the addDiesel experiment adds 90 Gg of diesel-related IVOCs proportionally to road transport emissions (SNAP7), whereas the IVOCs=1.5xPOA approach only adds 5 Gg to SNAP7 and another 26 Gg to other sectors (mainly to SNAP2: residential and non-industrial combustion). Therefore, our approach creates a considerably larger amount of SOA from IVOCs (and only from diesel-related IVOCs) than the previous method. The 1.5volPOA experiment was undertaken using the semivolatile treatment of POA emissions. This means that the modelled ASOA from this experiment also includes aged semivolatile POA, possibly giving it potential to create more ASOA than the Base or addDiesel experiments (the organic material added to the model in the 1.5volPOA experiment is 1.0xPOA (as SVOCs) + 1.5xPOA (IVOCs) = 2.5xPOA as introduced by Robinson et al. (2007) and Shrivastava et al. (2008)). It can be seen from Figs. 18a, b that treating POA as semivolatile leads to much lower concentrations than the nonvolatile treatment (which already underestimates measured concentrations of HOA and SFOA by -54% and -71%, respectively; Fig.5). This is not surprising since with the semivolatile treatment of POA only 3% + 6% + 9% of the POA is assigned to the three lowest volatility bins with saturation concentrations of 0.01, 0.1 and 1 μg m⁻³ (as given in Sect. 2.4). In a study in Mexico City, Shrivastava et al. (2011) then revised this treatment – assuming much higher total semi- and intermediate volatility POA emissions: 7.5 x the inventory emissions of (particulate) POA. This was justified by the fact that their emission factors of POA were derived from measurements at urban background sites, but, following Robinson et al. (2007), 2/3 of POA would have evaporated by then. (Recently, Shrivastava et al. (2015) also used this factor of 7.5 in global simulations.) Emission factors used in European inventories are, however, taken from tailpipe measurements with concentrations sufficiently high that most of the semivolatiles should still be reported in the particulate phase. Therefore the further underestimation of HOA and SFOA concentrations with the volatile treatment could be due to a number of issues: (i) a systematic underestimation of emissions, but for a different reason than in Shrivastava et al. (2011), (ii) the volatility of POA is overestimated by Robinson et al.
(2007), (iii) the evaporation of semivolatile POA emission is too rapid in the model (instantaneous in our set-up).

Figure 18c shows that the lower HOA and SFOA concentrations lead to a very small negative change for the absorptive partitioning of BSOA. Finally, it can be seen from the annual average concentrations of ASOA in Fig. 18d that including aged SVOCs and IVOCs in the simulation doubles the modelled ASOA concentration compared to the Base case scenario (ASOA from officially reported anthropogenic VOCs), but that the ASOA in our 1.5volPOA experiment is still much lower than simulated with our addDiesel experiment.”

![Figure 18. Simulated annual and seasonal average concentrations of OA components (BGND OA stands for Background OA) for the London North Kensington site of three different model experiments: Base - all emissions as in officially reported emissions inventories, POA is treated as non-volatile; 1.5volPOA - semivolatile treatment of POA + IVOC emissions added as 1.5xPOA; addDiesel - Base + IVOC emissions from diesel traffic added proportionally to VOC emissions from the on-road traffic source sector (SNAP7); both the latter additions as described in the main text.](image)

2. Given that POA are assumed to be inert, this study likely overestimates OA close to sources. The statement that diesel IVOC can explain about 30% of the annual SOA around London would have to change if POA were allowed to evaporate and react in the atmosphere.

Response: We derive the “30%” from the change in normalised mean bias (so in comparison to the measurements), not in comparison to total modelled SOA, and therefore we believe
3. The results presented here make a strong case that most SOA is ASOA. This conclusion contrasts strongly with studies based upon radiocarbon and other tracers. Heal et al. (2011) for example suggest a much stronger component from BSOA in Birmingham, and state that this was consistent with other European studies. Can the authors explain this apparent discrepancy?

Response: It is certainly not our intention to make a case that most SOA in London was ASOA, and we do not believe that our text does imply this claim. From Fig.17a in our manuscript (barplot of modelled annual average ASOA/BSOA/Background OA concentrations), it can be calculated that the ratio of modelled BSOA+Background OA to total SOA is 53%. Therefore even with the additional ASOA generated from the additional IVOC emissions, about half of the simulated SOA is still of biogenic origin, which is therefore not inconsistent with the experimental measurements using the radioisotope of carbon (14C) reported in Heal et al. (2011). It needs also to be noted that whilst 14C is an ideal tracer for distinguishing between fossil and contemporary carbon, it cannot directly distinguish whether the carbon is of primary or secondary origin. Therefore, for a direct comparison with a modelling study like ours, all primary components (such as the HOA and SFOA) would also have to be well represented in the model, but we showed a -71% bias for SFOA. This bias is caused by the fact that the national atmospheric emissions inventory assumes zero domestic wood and coal burning emissions in London, as it is a smoke control area (and therefore residential burning of these solid fuels is not allowed), but recent studies (e.g. Crilley et al. 2015) have concluded that there are indeed local sources of SFOA in London. Furthermore, in the current set-up, we model SFOA as one entity (so wood and coal together), but in the 14C analysis, OA from wood-burning and coal would be apportioned into different categories (contemporary and fossil, respectively).

To emphasise the above point, we have now added the following sentence to the end of Section 3.4: “We note that Fig. 17a shows that in the addDiesel simulation, the modelled BSOA+Background OA still makes up 53% of the SOA, as an annual average.”

4. The mass yields for OH oxidation of the n-pentadecane IVOC products is ca. 0.8 for C* up to 10 ug/m3, after correcting for the assumed density, and the great majority of this is one bin, the 10 ug/m3 bin. The potential for much SOA formation is very clear, but I wonder if the authors are exaggerating the amounts. The mass yields are taken from Presto et al. (2010), but that paper suggested that the yields were the product of multi-generational aging, not of the first reaction step. I wonder if aging should have been ignored for these compounds?

Response: Our view is that ageing should not be ignored for these compounds. Although Presto et al. (2010) report that these yields almost certainly include multigenerational processing during the experiment, they also conclude (in the last paragraph of their paper): “The slow rise in f44 in experimental systems - a few percent over several hours - further indicates that OA constantly evolves over long time scales, on the order of days, and that short chamber experiments likely do not reproduce the complete transformation from emissions to OOA.”
We do acknowledge, however, that the ageing rates and assumptions used in VBS modelling studies can vary quite widely between different studies, as is noted in the Discussion (paragraph beginning “We use an ageing rate of...”).

Responses to smaller points:

1. Abstract: I found the first sentence rather vague (what is high-resolution?), and not so interesting (yet another model study). The abstract would make more impact if it began with a comment on the extent of new emissions which forms the basis for this study.

Response: We do not believe the opening sentence is vague as it provides a number of specific contextual facts to the work; namely, that it is an ACTM modelling study, that the atmospheric process of interest is SOA formation, that the geographical context is the UK, and that the period of study is the full year 2012. However, we accept that the phrasing “high resolution” is not specific and that we could introduce the concept of new estimates of emissions into the opening sentence (although we note the latter is encapsulated in the title of the paper). We have therefore amended the opening sentence to now read: “We present high-resolution (5 km × 5 km) atmospheric chemistry transport model (ACTM) simulations of the impact of new estimates of traffic-related emissions on secondary organic aerosol (SOA) formation over the UK for 2012.”

2. P2, L12. I don’t think the results ‘prove’ that the model has good SOA prediction skill. Even if the comparison with measurements was impressive, there are too many unknowns regarding SOA formation and I don’t think any model can claim good skill. I think that this phrase can be omitted.

Response: We agree that the text referring to good model skill is not appropriate for the abstract and have now omitted this sentence.

3. P3, L64. You need to define the temperature at which these C* values apply.

Response: The temperature of 298 K has now been added.


Response: It was not our intention to imply that national inventories use AMS data. However, on review we note that our original text referring to AMS measurements of the OC content of particles is not relevant to the point being made in this sentence that SVOC and IVOC species are hard to measure, so we have now deleted it. The modified text now reads: “Current emissions inventories, however, only report estimates for VOCs and for the particle fraction of the emissions of species with lower volatilities. The main reason for this is that compounds with intermediate volatility (SVOCs and IVOCs) are difficult to quantify and this is currently not routinely undertaken.”

5. P4, L114. Define PMF. Also, which PMF method was used?
Response: The acronym PMF is defined already in the second paragraph of the Introduction. We have amended a sentence in Sect. 2.6 to include the PMF methods:
“A detailed description of the derivation and optimization of the factors retrieved from the AMS data at Detling can be found in Xu et al. (2016), at London North Kensington in Young et al. (2015a) and Young et al. (2015b) (all three of these analyses were performed with the PMF2 solver), and at Harwell in Di Marco et al. (2015) (using the ME-2 solver).”

6. P5, L132. WRF can be set up in many different ways, with varying impacts on accuracy for air pollutant applications. Please give more details or a suitable reference.

Response: We have added the following sentence: “The WRF configuration was as follows: Lin Purdue for microphysics; Grell-3 for cumulus parametrization; Goddart Shortwave for radiation physics; and Yonsey University (YSU) for planetary boundary layer (PBL) height (see NCAR (2008) for further information).”

7. P5, L137, specify anthropogenic emissions here.

Response: Done.


Response: Done.

9. P5, L142. Why use a paragraph on an NFR system which is not used in this work? Delete.

Response: The text is now deleted.

10. P6, L156. The term SFOA is confusing, and wasn’t used by Bergstrom et al as claimed here. If I understand right SFOA includes biomass burning (which is usually said to give BBOA), but also coal and charcoal.

Response: We have now added the BBOA factor as well as the following sentence in the Introduction explaining the difference between SFOA and BBOA: “The SFOA factor is a more general version of BBOA as it includes (in addition to biomass) other sources such as coal and charcoal.”

11. P6, L173. It could be noted that the Jathar et al. (2014) study also suggested different ratios of IVOC to PM than those of Shrivastava et al.

Response: The emissions of “unspeciated non-methane organic gases” in Jathar et al. (2014) are still based on the same measurements as is Shrivastava et al.’s estimate (although they also included a couple of newer studies and averaged the estimates, so the numbers are slightly different). We believe that our paragraph about Jathar et al. in the Introduction is sufficient and that there is no requirement to add more about this to the Methods.
12. P7, L191. ‘under modeled SOA’ - do you mean when comparing with observations?

Response: Yes, this was what we meant. We have now added “when comparing with observations” to the end of this sentence.

13. P7, L200. Shouldn’t you also mention aromatics and other compounds here.

Response: In the Dunmore et al. (2015) paper, the authors describe a quantification technique which uses the grouping of similar species in a lumped analysis based on carbon number and functionality. Given the separation of VOCs in a two dimensional space (from the use of a comprehensive two dimensional gas chromatography system), the aliphatic and aromatic compounds could be quantified separately. In our analysis, we only include the additional aliphatic IVOC species observed as their dominant emission source is likely from the use of diesel engines. We have now added the word aliphatic to the following sentence in the section “Additional IVOCs from diesel”: “In this study, aliphatic IVOC emissions from diesel vehicles were introduced into the model proportionally to on-road transport VOC emissions,...”


Response: We have now replaced “GC x GC” with “measured by a comprehensive two dimensional gas chromatography (GCxGC) system (Dunmore et al., 2015).”

15. P7, L209. Any reference for the number of studies providing that rate constant?

Response: We have added a citation to the review article by Atkinson and Arey, 2003.


Response: We have added the word “VOCs” so text now reads “biogenic emissions of VOCs”.

17. p9. Sect. 2.6 ‘Comparison with measurements’: This section can be simply renamed ‘Measurements, since that is what it deals with down to L288.

Response: We would like to retain “comparison with measurements” in the section title since (i) the IVOC emissions were also based on measurements, but were not used to compare the model with, (ii) then we can keep the evaluation statistics in the same section, reducing the number of short sub-sections.

18. P9, L259. Why have references to Fig. 3? Give the references after the mention of each site, or add ’site details given in’ or some such phrase.

Response: We have moved this sentence to after the citation of the references.

19. P9, L274. Are you sure that European inventories don’t include cooking OA? I think it may be underestimated, but am not sure it is ignored completely.
Response: We are sure that COA is not included in the UK National emissions inventory (Tim Murrells 2016, personal contact; NAEI 2013) Fountoukis et al. 2016 also claim that COA is not included in the European inventory they use. We have changed the sentence in the manuscript to read: “As our emissions inventory does not include cooking OA (NAEI, 2013), this factor could not be compared with the model.”

20. P9, L276. This sentence was confusing. I can see that two instruments can disagree, but what does it mean if there is just one instrument? Can an AMS and its PMF disagree, or what?

Response: We agree that this sentence was not entirely clear and have amended it to: “When AMS measurements and their PMF apportionments are compared, some disagreement is observed, as shown for the two instruments measuring at the same time at the same location at London North Kensington.”

21. P9, L284 .... what period/site/analysis are these sentences and statistics referring to?

Response: We have added “at the London North Kensington site during the winter IOP” to this sentence.

22. P9, L289 on. This small section on statistical metrics has nothing to do with the discussion of AMS etc which it follows, and could be set in a small section of its own.

Response: This section is called “comparison with measurements” so we believe having the statistics here is appropriate. See also our response to comment no. 17.

23. P9, L291. I don’t think correlation coefficient needs a reference to Carslaw and Ropkins; ’r’ has been used for many many years before that paper was written. Actually, NMGE might need more explanation. All these could usefully be defined in supplementary.

Response: We have removed this citation and added the equation for NMGE. We would like to keep this section in the main paper as it is not taking up much space and we are trying to reduce the number of times the reader is referred to the supplement.


Response: The necessary explanations have now been added.

25. P10, L298. Re-phrase - it sounds as though the measurement mean is better at capturing the variation in measurements than the model.

Response: Now re-phrased to read: “a zero or negative COE implies that the model cannot explain any of the variation in the observations”.

26. P10, L303. This bit about WRF could be moved to Sect. 2.1.

Response: This text has been moved as suggested.
27. P11, around L35. All these numbers for NMB, etc. could be tabulated for easier comparison.

Response: These numbers are also given as labels in Fig. 7 in a consistent, comparable manner. We anticipate this will become clearer and easier for the reader when the paper is properly typeset and the figures located within the text, rather than at the end.

28. P12, L364. This refers to Fig. 12a,b, but there are no a,b labels in Fig.12

Response: There were small labels on Fig. 12 which we have now replaced with larger ones.

29. P12, L389. Re-phrase (or omit). It is obvious from the plots that this background OA is an overestimate for some days at least.

Response: We agree; the sentence referred to has now been omitted.

30. P13, L405 on. The whole discussion here is in terms of SOA and IVOC. But, how did the model perform for NOx and CO for these ‘difficult’ periods - maybe the problem is dispersion rather than IVOC? Or maybe the model’s enthalpy values are wrong, and don’t respond to cold temperatures as they should. I don’t see why problems are blamed on domestic sources either. Wouldn’t for example cold-starts for vehicles also produce more POA/IVOC, or commercial premises use more fuel in cold conditions? Are wood-burning emissions really an issue in London?

Response: The model performance for NOx is already presented in the paper (Figure 6a) and the performance for NOx during this period is much better than that of SFOA. We believe that domestic emissions are the most likely culprit here as local emissions of wood-burning are indeed an issue in London. We have added the following sentences: “Furthermore, London is a smoke control area and therefore no residential emissions of SFOA are assumed by the national emissions inventory for this area. However, recent studies have suggested that there are indeed local sources of SFOA in London (Crilley et al. 2015, Young et al. 2015).”

31. P15, L475. The title says comparison to previous (IVOC=1.5xPOA) approach, but as noted above, the method used here seems to be unique; not that of earlier papers.

Response: We have provided detailed response to this comment where it was first made above (1st major comment). In brief, we have undertaken new model simulation for the 1.5xPOA experiment and have amended the text accordingly.

32. P15-P16. The authors make various policy recommendations, e.g. (P15, L500) ‘refinements should be reported to CEIP’ and a very specific recommendation for PM1, PM1-2.5 and PM2.5-10 on P16, L533. Why not just suggest submission of size distributions? Why no mention of volatility - the VBS approach almost begs for people to submit emissions in different volatility classes. And since this paper is really exploring IVOC and not PM emissions per se, why didn’t the authors focus on those?
Actually, I suggest that the authors don’t try to tell countries what to do, but rather discuss any scientific insights into emission reporting that this study on IVOC reveals.

Response: We have now removed the two sentences that were directly referring to “what other countries should do”. The suggestion for the incorporation of more particle size distribution data in inventories was already in the Discussion (a few paragraphs further down from the point in question). We have also now added the suggested recommendation of categorising PM emissions in terms of volatility classes (to emphasise a comment we made on this already in the Introduction):
“We showed that treating POA as semivolatile and letting it evaporate lead to a great underestimation of HOA and SFOA concentrations compared to measurements at the London North Kensington urban background site. As has been highlighted by a number studies before us (listed in the Introduction), we would also emphasize that a major source of uncertainty in OA modelling is the volatility of primary emissions, an issue that currently not addressed by official emissions inventories.”

33. P16, L509. Can’t small changes in SOA (or any pollutant) also be a reflection of longrange transport? Not all pollution is formed at short time-scales close to source.

Response: Yes, we agree and we now mention long-range transport in the sentence in question: “A relatively small daytime increase of SOA could be explained by the expansion of the boundary layer height (Xu et al. 2015), as well as by contributions from long-range transport.”

34. P16, L520. Where did the value 4.0e(-12) come from for ASOA and BSOA oxidation?

Response: The value came from Lane et al. 2008; we cited this in the Methods section, but have now added the citation to the Discussion as well.

35. P16, L520 on. This section offers a few ‘tuning’ suggestions, but there are always any number of these in the field of SOA formation. For example, recent studies have suggested that SOA formation should be much greater than previously assumed, perhaps by a factor of four or so (Zhang et al., 2014).

Response: We agree; we have simply made some suggestions.

36. P16, L527. I would say that this paper illustrates the potential for a significant contribution, rather than that they can quantify the relative impact. To do the latter, one would need to be sure that all relative impacts are reasonably well known, and that clearly isn’t the case.

Response: Whilst we agree with the general sentiment of this comment, our response to this question is the same as Major Point no.2: We derive the “30%” from the change in NMB (so in comparison to the measurements), not in comparison to total modelled SOA, and therefore we believe this is an appropriate number/value for our statement highlighting the potential contribution this addition has.
37. P17, Sect. 5. The first paragraph simply repeats sentences from Sect. Don’t do that. This section should also mention the results for other pollutants (NOx, O3, etc.), which are the main reason one can have some confidence in the basic modeling system. (Use of such data is one of the strengths of this paper I think.)

Response: The text has been reworded so that it is not a direct copy and the results for other pollutants are mentioned.

38. P17, L563. imported from where?

Response: We have modified the sentence to include “mainland Europe”. It now reads: “...this was caused by an intense pollution plume with a strong gradient of SOA from mainland Europe passing over the rural location...”

39. P17, last paragraph. The interpretation of what contributes to the 90-th percentile is not so easy I think. And I certainly don’t think one can state that 40% is due to missing diesel precursors. SOA is too complex for such simple statements.

Response: We agree that SOA is a complex issue for making delimitative quantitative statements; however, the modelling work points us in a certain direction. We have amended the sentence to state that the influence of missing diesel precursors is even greater on high percentile SOA days than its contribution to annual average SOA (removing the statement of “40%”). We have also removed the statement from the abstract. The sentence in the Conclusions now reads: “Moreover, the 90-th percentile of modelled daily SOA concentrations for the whole year is 3.8 μg m\(^{-3}\), and the influence of missing diesel precursors is even greater on high percentile SOA days that its contribution to annual average SOA.”

40. P20, L635. Expand/explain CEIP. Is this the name of a report, or just a web side?

Response: Expanded, see also the response to question no. 41.

41. P21, L665, EEA, Entec - these references are too short for readers to understand or find. Give proper references, with addresses as necessary.

Response: Thank you for pointing this out. Something happened to these references during the formatting of the manuscript (we did have proper URLs for these references at some point). We will make sure they are properly included in the revised manuscript.

42. P29, Fig. 1. The caption should state early that this is PM25 *and* SVOC/IVOC gases. (The issue of IVOC or SVOC+IVOC is discussed above, but it complicates this figure.)

Response: We think that mentioning IVOCs in the second sentence is appropriate, and early enough.

43. P31. Fig. 5. Explain ‘Base’ as used in legend. P31. Fig. 6(a). NOx is a the sum of both NO and NO2. Were they really summed with own molecular weights, or is this as NO2?
(ppb would have been easier to interpret!). Also, in the captions, add the superscript ion-labels too.

Response: Yes, the sum of NO and NO2 is represented as NO2. We have now added a note about this to the caption: “NOx (as NO2)”, as well as added the superscription labels. We agree that ppb is easier to interpret for gases, but this plot also included particulates and we would like to use the same units for the different panels. Furthermore, European Air Quality Standards (http://ec.europa.eu/environment/air/quality/standards.htm) are also defined in mass units for gases.

44. P33, Fig. 9. This Figure looks very fine on-screen, but when printing out it looks very different - much of the black seems to be over-written with green and/or blue. Please use a different figure format, and check the printout.

Response: We thank the reviewer for this comment. We printed the manuscript out with a few different printers, and Fig. 9 looks fine with all of the printers we tried, but we did notice that some printers had problems printing our scatterplots. We have replaced the shading on Fig. 9 with lines and converted our scatterplots into a more printer friendly format/size.

45. P34, Figs. 11-12. We don’t really need these since Fig. 10 has made the point about gradients well. Move to Supplementary.

Response: We agree that Fig. 10 is enough to make the point about gradient, but we also use Figs. 11 and 12 for showing how spatially variable SOA can be even on daily averaged maps (which include contributions from both import and from very local sources).

46. P35, Fig. 14. Why use log-log plots? Wouldn’t a simple liner plot better display the range of data?

Response: We use log-log plots for 2 reasons: (i) it expands the lower range without losing information in the higher range (as on a linear scatter plot for a large dataset, the plotting symbols will overlap with each other much more in the lower range), (ii) log-log plots are have been used in other recent OA modelling studies (e.g. Fountoukis et al. 2014) making it easier to visually compare some of the results of model evaluation.

47. P36, Fig. 16. Here the addDiesel statistics look worse than the base-results. Are these really consistent with data presented in Table 4?

Response: These values are correct. The difference in NMGE, r and COE values is very small, so not really “worse”. Note the (positive) change in NMB.

48. Figs and color schemes. The colors used for ASOA, addDiesel, etc seem to vary randomly from figure to figure (e.g. green is addDiesel in Fig.9 but observed SOA in Fig. 17. Please harmonize.

Response: Figure colours have been changed so that same colours are no longer used for different variables.
References


Heal, MR et al. Application of $^{14}$C analyses to source apportionment of carbonaceous PM2.5 in the UK, Atmos. Environ., 45, 2341-2348, 2011


Zhang, X., et al., Influence of vapor wall loss in laboratory chambers on yields of secondary organic aerosol, PNAS, 111, no. 16, 5802-5807, 2014
