

Geosci. Model Dev. Discuss., author comment AC2  
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## Reply on RC2

Edmund Ryan and Oliver Wild

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Author comment on "Calibrating a global atmospheric chemistry transport model using Gaussian process emulation and ground-level concentrations of ozone and carbon monoxide" by Edmund Ryan and Oliver Wild, Geosci. Model Dev. Discuss., <https://doi.org/10.5194/gmd-2021-39-AC2>, 2021

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### Major Comments

**(1) Reviewer's comment:** When I read the abstract and introduction, I have an impression that the author will apply the calibration framework developed by Kennedy and O'Hagan (2001), but it turns out that it is not the case. The author should have explained in the beginning why they do not include model discrepancy in the methodology or GP model. It is well known that model discrepancy is very important for model calibration (Brynjarsdóttir and O'Hagan, 2014), and it is obvious there is no CTM that can perfectly predict the true process, even with the optimised inputs. Brynjarsdóttir, J. and O'Hagan, A.: Learning about physical parameters: The importance of model discrepancy, *Inverse Problem*, 30, 114007, 2014

**Author's response:** Many thanks for this comment. We have specifically chosen not to include a discrepancy term for two reasons:

- For the scenarios where we use synthetic data, no discrepancy term is required because the synthetic data is generated by adding noise and spatial gaps to the emulator output for the control run.
- For the scenarios involving reanalysis data, there is no simple and defensible method to estimate the discrepancy term.

Neither of the papers cited in the reviewer's comment explain the need for the term for anything other than the most simple model system. The discrepancy represents the missing processes in the model. However we often don't know what these missing processes are or how to estimate them. When performing a regular model calibration without an emulator (e.g., applying MCMC on the original model) we would not include a discrepancy term. It is therefore not clear why we need to include a discrepancy term into the calibration formulation when using an emulator. We highlight that in the abstract of the Brynjarsdóttir and O'Hagan paper it states: *"The challenge with incorporating model discrepancy in statistical inverse problems is being confounded with calibration parameters, which will only be resolved with meaningful priors."* If we wish to estimate the discrepancy term as part of the calibration process, then this confounds the parameters we are trying to estimate. Hence, the paper makes the point that you need

highly informative priors. The paper does not address what to do if you do not have these, as is the case here.

However, to investigate the importance of this term we adopt the simple rule of thumb that the discrepancy term is 10% of the magnitude of the observation (Jeremy Oakley, personal communication). We repeated the experiment to estimate the eight scaling parameters and the SD term using surface ozone reanalysis data at 2.5% spatial coverage. We find that there is almost no difference in the marginal posterior distribution when we include the discrepancy terms in the formulation compared with when we omit them (see Figure 1 in the supplemental pdf).

**(2) Reviewer's comment:** I used to think the calibration won't work on tropospheric ozone, since accurate tropospheric ozone simulation relies on accurate regional emission inventory and meteorology. These inputs need to be continuously updated, rather than calibrated. As far as I am aware, the existing calibration approaches assume the parameters are "fixed initial condition", rather than time-varying or spatial-varying settings. Would it be making more sense to consider such input, e.g. emission inventory, as a time series? Morris, M. D. (2012). Gaussian surrogates for computer models with time-varying inputs and outputs. *Technometrics*, 54(1), 42-50.

**Author's response:** The parameters we are estimating are global scaling parameters applied uniformly to major input variables, e.g. global surface NO<sub>x</sub> emissions. These variables already vary in space and time, and our parameters merely scale the global annual magnitude. It would certainly be interesting in a future study to explore the effects of seasonality through temporally varying factors, as the reviewer suggests, although this comes at the cost of a greatly increased number of parameters to calibrate. However, for the current study our aim is to demonstrate the feasibility of calibration and thus we have attempted to keep the problem as simple as possible, and focus only on a single, globally uniform scaling parameter for a given input. We anticipate that using the approaches successfully demonstrated here, atmospheric modellers will be able to explore the more complex spatial and temporal sensitivities for different variables in future.

**Changes made to the manuscript:** We have rephrased our introduction to the scaling factors in section 2.1 to highlight that they are global factors applied uniformly to the particular process considered.

**(3) Reviewer's comment:** The authors split the total output into 6528 separated GP emulators (p13). Since each emulator is treated separately, how can they be certain all the emulators yield the same calibrated inputs?

- what if the calibrated inputs from different emulators are diverse?
- even if the calibrated inputs from different emulators are similar, how the optimised inputs are determined?

**Author's response:** The estimates and associated uncertainties of the eight scaling parameters are determined using a common type of Markov Chain Monte Carlo (MCMC) algorithm called Gibbs sampling. In its basic form, Gibbs sampling is a special case of the Metropolis-Hastings algorithm. A key component of the algorithm is a likelihood function which quantifies the mismatch between the measurements and the model predictions for a given set of values for the parameters. The 6528 separate emulators correspond to 12 monthly measurements at each of the 272 spatially varying grid cells, for each of the 2 variables (ozone and carbon monoxide). In short, the number of emulators corresponds to the dimension of the output space. For the scenario involving all 6528 emulators, the likelihood function incorporates all 6528 modelled values (where the emulator is used in place of the chemistry model) and the corresponding 6528 measurement values. Thus

while the emulators are treated separately, only a single likelihood function is considered, not 6528 separate likelihood functions as the reviewer's comment suggests. We thus end up with a single set of calibrated inputs that has already been optimised over all times and locations. We have made changes to our description of the approach used to make this clearer.

**Changes made to the manuscript:** We recognise that in the current description of the MCMC algorithm is brief and in particular we do not mention Gibbs sampling. In the revised manuscript we give more detail about what Gibbs sampling is and cite well known sources such as the 'Bayesian Data Analysis' book by Gelman et al.

**(4) Reviewer's comment:** Given the fact that the authors build an emulator separately for each location and month, I am actually concerned whether this study is actually useful. If the inputs are only calibrated at a specific month/location, and might not be valid for other months/locations, then this message is not really useful for modelers. If they cannot run the model at a single setting of the input parameters, the calibration does not consider to be working. It is also contradictory to its title "Calibrating a global atmospheric chemistry transport model", because the authors have not taken any spatial and temporal correlations into account in their GP model.

**Author's response:** As noted in our previous response, the likelihood function is applied across all independent emulators so that we derive a single set of calibrated inputs across all times and places. We agree that independent calibration of each emulator would not be useful, as the reviewer suggests, but we hope that we have now addressed this misunderstanding through the changes we have made to the text in response to the comment above. Our approach generates a single set of scaling parameters and thus allows us to effectively calibrate the model.

**Changes made to the manuscript:** We have improved the clarity of the text in the methods section of the manuscript to make this point clearer, as noted in the previous response.

**(5) Reviewer's comment:** A collection of 272 locations is not strictly prohibited for GP emulation/calibration, why building 24 emulators is not an option?

**Author's response:** As noted above, we build separate emulators for each variable (ozone and CO) for each month at each of the 272 grid cells. It would be possible to reduce the number of emulators needed through application of principal component analysis methods, which we have demonstrated in a previous study (Ryan et al., 2018), but we have chosen to generate separate emulators here both for reasons of simplicity and because emulator generation is not the most computationally demanding aspect of this study.

**Changes made to the manuscript:** As noted above we have improved the clarity of the text in the methods section of the manuscript to address this.

**(6) Reviewer's comment:** Calibration is not just about estimating the optimised inputs, but also about estimating (under reasonable assumptions) the potential simulator misspecification, or discrepancy. However, this latter estimation is not explored in the studies. This should be explored more carefully, even if the model discrepancy is not accounted for. For example, does the final calibrated model outperform than the ensemble mean output from their Latin hypercube design with respect to synthetic or real observations?

**Author's response:** The reviewer makes a good point here about estimating the model misspecification. To address this point, we have created a new figure that compares the

emulator predictions of the surface concentrations using the prior and posterior values for the inputs (see figure 2 in supplemental pdf). The left hand plot shows the mean and 95% prediction interval of the surface O<sub>3</sub> as predicted by the emulators, using 1000 samples of the inputs from the prior distribution. For the right hand plot, we took 1800 samples of the inputs from the posterior distribution, based on the calibration run involving only surface O<sub>3</sub> synthetic data, 20% spatial coverage, and a representation error factor of  $p=0.2$  (the third level of representation error). We then ran all 1800 posterior samples through each emulator. For each panel there are 272\*12 points, which are made up of 272 spatial pixels and 12 months for each pixel. Note that although the calibration run involved 20% of the data, these predictions involve 100% of the data. We can clearly see that the prior distribution values of the predicted surface ozone are biased (more so at high values) and have large uncertainty (again, more so at high values). In contrast the posterior distribution values of the predicted ozone have only a tiny bias which is indicated by the very small value of the median absolute difference (MAD).

**Changes made in manuscript:** A new figure has been added to the manuscript as described above. This is a very useful addition to the paper and we thank the reviewer for suggesting it.

**Reviewer's comment:** How do the authors choose the settings of their prior distributions for the hyperparameters in GP mean and covariance functions? This is not discussed anywhere.

**Author's response:** The hyperparameters for each emulator are estimated by maximum likelihood using the DiceKriging R package (Roustant et al., 2012). As Kennedy & O'Hagan (2001) point out, in order to integrate out the hyperparameters in the formulation of the GP emulator, we would require highly informative priors. In most cases, such informative priors do not exist. Hence, Kennedy & O'Hagan (2001) propose to provide a point estimate of the hyperparameters and to use these in the formulae for the mean and covariance functions of the GP emulator. We have adjusted the text in section 2.6 to make our approach here clearer.

**Changes made in manuscript:** We have improved the clarity of the text in the methods section of the manuscript with regard to this point.

**Reviewer's comment:** The authors often attribute the variation between irregularly distributed measurements and gridded output to representative error throughout the paper. This is unfair and inadequate, because model simulated output at gridded points does not imply model output is more representative at a broader scale, but simply because the model is too limited or too coarse to represent all the fine structures.

**Author's response:** The model output represents concentrations averaged over a coarse model grid square (not at a "gridded point"), while measured concentrations are indeed at specific points. There is a fundamental incommensurability in this comparison, given that the model cannot resolve fine structures, while the measurements may or may not sample them. To calibrate the model we need a measurement based assessment of concentrations at the model grid scale, ideally, but there are insufficient measurement sites per grid square to generate this. Our comparison is thus dependent on how well the available measurement sites represent the wider area, and we have defined this as the representative error in the paper. We note that this is a function of the grid scale considered as much as a property of the measurement due to its siting.

**Changes made in manuscript:** We have rephrased our definition of representative error on page 5 in the introduction to make this clearer. See also our response to the specific comments about this point, below.

## Minor Comments

**Reviewer's comment:** P2, l5 and P6, l4, I understand that this is a standard statement to say poor spatial coverage of atmospheric composition measurements, but in your case the surface ozone measurements are very dense in Europe and most of the US, compared to the 2.8 degree resolution of their model output.

**Author's response:** While there are certainly more surface measurements in Europe and the US than elsewhere around the globe, the adequacy of the coverage must be judged on the spatial variability of the variable of interest, not the model resolution. While the lifetime of ozone is long in the free troposphere, the short timescales for chemical and dynamical processes controlling surface ozone drive much greater spatial variability. Given that the current network is still far from sufficient to capture this, we feel fully justified in describing the measurements as sparse.

**Changes made in manuscript:** No changes were made.

**Reviewer's comment:** p3, l4, this reference should be updated to the most recent GBD 2019 study. 4.2m premature deaths are referring to the pm2.5 estimates. This study is about ozone, so the latest estimate of 365 thousand premature deaths is more appropriate.

**Author's response:** We agree that this would be more relevant here and thank the reviewer for pointing this out.

**Changes made in manuscript:** We have updated the reference and amended the text to include this figure, as suggested.

**Reviewer's comment:** p3, l14-16, this statement seems unfair, since several attempts were already made for calibrating/emulating this type of model output. See following references Chang, K. L., & Guillas, S. (2019). Computer model calibration with large non-stationary spatial outputs: application to the calibration of a climate model. *Journal of the Royal Statistical Society: Series C (Applied Statistics)*, 68(1), 51-78. Couvreur, F., Hourdin, F., Williamson, D., Roebrig, R., Volodina, V., Villefranque, N. et al. (2021). Process-based climate model development harnessing machine learning: I. a calibration tool for parameterization improvement. *Journal of Advances in Modeling Earth Systems*, 13(3), e2020MS002217. Guan, Y., Sampson, C., Tucker, J. D., Chang, W., Mondal, A., Haran, M., & Sulsky, D. (2019). Computer model calibration based on image warping metrics: an application for sea ice deformation. *Journal of Agricultural, Biological and Environmental Statistics*, 24(3), 444-463. Karagiannis, G., Konomi, B. A., & Lin, G. (2019). On the Bayesian calibration of expensive computer models with input dependent parameters. *Spatial Statistics*, 34, 100258. Salter, J. M., Williamson, D. B., Scinocca, J., & Kharin, V. (2019). Uncertainty quantification for computer models with spatial output using calibration-optimal bases. *Journal of the American Statistical Association*.

**Author's response:** The point we are making here refers specifically to atmospheric chemistry models, as stated in the text. However, we acknowledge that calibration has been successfully applied to climate models, as the reviewer highlights, and we have therefore modified the text to note this.

**Changes made in manuscript:** We have updated text and cited some of the papers mentioned by the reviewer.

**Reviewer's comment:** P5, l5 and p22, l21 "require thousands of model runs" appear to

be exaggerating, since the authors are aware that will be dependent on how many parameters to be turned (and they only use 80 runs).

**Author's response:** The point that we are making here is that global sensitivity analysis (e.g. extended FAST) and model calibration (e.g. MCMC) require thousands of model evaluations. This is true whether we use a computationally expensive model or a surrogate model. The process is currently only feasible with a surrogate model, and this is our motivation for using Gaussian Process emulation. Generating the emulator requires only a small number of executions of the expensive model (80 in our case), but we still need to carry out 1000s of runs with the emulator to conduct sensitivity analysis and model calibration. However, we acknowledge the reviewer's concern and have rephrased the first occurrence to avoid the appearance of exaggeration.

**Changes made in manuscript:** We have updated the text to make the point clearer, noting specifically that sensitivity analysis and model calibration "may require many thousands of model runs".

**Reviewer's comment:** P5, l10, "Since the first application of emulation methods for model calibration (Kennedy and O'Hagan, 2001), [...] In this study, we apply these approaches to models of tropospheric ozone for the first time to demonstrate the feasibility of parameter estimation." This is inappropriate because the authors do not consider model discrepancy and measurement uncertainty in their emulator component (see my major comment 1).

**Author's response:** As noted in our responses above, we have chosen to omit the discrepancy term for this demonstration of feasibility, as our exploratory tests showed that the effect is small, but measurement uncertainty is included as a component of representation error. We have applied the same emulation methods as Kennedy and O'Hagan, despite considering a very different system, and therefore we feel that it is fully appropriate to credit them with introducing it.

**Changes made in manuscript:** No changes were made.

**Reviewer's comment:** P5, l12, Higdon et al. (2008) should be cited, since this is the first paper successfully extending the calibration framework into the "highly multivariate output". Higdon, D., Gattiker, J., Williams, B., & Rightley, M. (2008). Computer model calibration using high-dimensional output. *Journal of the American Statistical Association*, 103(482), 570-583.

**Author's response:** Thank you for this reference; we now cite this here as suggested.

**Changes to be made in manuscript:** This paper is now cited in the manuscript.

**Reviewer's comment:** P5, l18, "Firstly, ground-level composition measurements are usually made at a single location which may not be representative of a wider region at the grid-scale of the model. Global chemistry transport models typically have a spatial scale of the order of 100 km." This statement is somehow misleading, because dense ground based measurements (especially in Europe) reflect the local fine variations that can not be solved by coarse model resolution.

**Author's response:** The reviewer is correct to point out that this is phrased somewhat awkwardly in the paper from the perspective of the model (the observations aren't representative of the model grid square) rather than of the measurements (the model is unable to represent observed variations below the grid scale). However, even over Europe where there are more observations than in other parts of the world, measurement sites are insufficiently dense to fully characterise the spatial variability that would be needed to

integrate them reliably to the model grid scale. To avoid any potential confusion, we have now rephrased this sentence in the manuscript.

**Changes made in manuscript:** We have rephrased these sentences to read: "Firstly, global chemistry transport models typically have grid scales of the order of 100 km which is insufficient to resolve spatial variability in many atmospheric constituents. Surface measurements made at a single location may not be representative of the spatial scales resolved in the model."

**Reviewer's comment:** P8, l6, As I mentioned earlier, emissions are also dynamic in time and space. So the authors should comment if these parameters only represent the initial conditions.

**Author's response:** As noted in our response to the earlier comment, we are applying globally uniform scaling factors that do not vary in space and time. These are applied to the processes continuously, and are independent of the initial conditions. As noted above, the introductory text in Section 2.1 has been adjusted to make this clearer.

**Changes made in manuscript:** As mentioned in the earlier comments

**Reviewer's comment:** P8, l19, I believe the reference is Chang et al. (2017).

**Author's response:** Thank you, and apologies for getting the publication date wrong here!

**Changes made in manuscript:** The reference has been corrected in the manuscript.

**Reviewer's comment:** P11, l23, There are a few alternative approaches, such as principal components (Higdon et al. 2008; Holden et al., 2015) or low rank approximations (Bayarri et al., 2007; Bowman and Woods 2016; Chang and Guillas, 2019), that are proposed to tackle high dimensional output. Bayarri, M., Berger, J., Cafeo, J., Garcia-Donato, G., Liu, F., Palomo, J., Parthasarathy, R., Paulo, R., Sacks, J., and Walsh, D. (2007). Computer model validation with functional output. *Ann. Statist.*, 35(5):1874–1906 Bowman, V. E. and Woods, D. C. (2016). Emulation of multivariate simulators using thin-plate splines with application to atmospheric dispersion. *J. Uncertainty Quant.*, 4(1):1323–1344. Holden, P. B., Edwards, N. R., Garthwaite, P. H., and Wilkinson, R. D. (2015). Emulation and interpretation of high-dimensional climate model outputs. *J. Appl. Statist.*, 42(9):2038–2055.

**Author's response:** In a previous paper using the same model, we implemented the principal component approach to carry out global sensitivity analysis (Ryan et al., 2018, cited in the manuscript). This gives a similar answer to building separate scalar-output emulators for each dimension of the output space, and is thus a useful way to reduce the dimensionality. We have considered multivariate outputs, but feel that unless the outputs of the emulator are being used as inputs for another emulator it is fine to use the scalar output emulator approach. However, we agree that it would be useful to reference other approaches to dealing with multivariate output when implementing emulators, and have now done this.

**Changes made in manuscript:** We have added references to other approaches of dealing with multivariate output when implementing emulators, including those suggested by the referee.

**Reviewer's comment:** P12, l16, If "where B is a  $p \times p$  matrix with zeros in the off diagonals" how the different input parameters can be correlated?

**Author's response:** As explained in section 2.5, B is a diagonal matrix where the elements are roughness parameters that describe the linearity of the input-output relationship. The matrix does not describe the relationship between the input terms as the reviewer suggests here. This is part of a standard explanation for the description of the Gaussian process emulator (e.g. see papers by Jeremy Oakley and Tony O'Hagan).

**Changes made in manuscript:** We will investigate to see if any changes can be made to make this part of the text read clearer.

**Reviewer's comment:** P14, l1, The emulators used in this study are not taken into account measurement uncertainty or model discrepancy, so it merely represents the "output interpolator", I do not see why the authors should report R2.

**Author's response:** The uncertainty in the measurements is typically substantially less than that in their representativeness of the model grid scale, and it is thus effectively included as a small component of the representation error that we do fully consider. We do not include a model discrepancy term as explained above. We implement a Gaussian process emulator as described in O'Hagan (2006), which also quantifies the uncertainty at points in the output spaces where there are no training data. We therefore feel that it is appropriate to quote an  $R^2$  for the comparison. O'Hagan (2006) Bayesian analysis of computer code outputs: a tutorial. Reliability Engineering & System Safety, 91, 1290-1300

**Changes made in manuscript:** The  $R^2$  term has been retained in the manuscript.

Please also note the supplement to this comment:

<https://gmd.copernicus.org/preprints/gmd-2021-39/gmd-2021-39-AC2-supplement.pdf>