1	Calibrating a global atmospheric chemistry transport model using Gaussian process
2	emulation and ground-level concentrations of ozone and carbon monoxide
3	Edmund Ryan <sup>1,2</sup> , Oliver Wild <sup>1</sup> *
4	
5	<sup>1</sup> Lancaster Environment Centre, Lancaster University, UK.
6	<sup>2</sup> Now at: Corndel, London, UK.
7	
8	
9	*Corresponding author:
10	Lancaster Environment Centre,
11	Lancaster University,
12	Lancaster,
13	LA1 4YW
14	o.wild@lancaster.ac.uk
15	
16	Keywords: atmospheric chemistry transport model, model calibration, surface ozone,
17	measurement representativeness, Gaussian Process emulation, Markov Chain Monte Carlo, Just
18	Another Gibbs Sampler.
19	
20	For submission to: Geoscientific Model Development
21	
22	

# 1 Abstract

2 Atmospheric chemistry transport models are important tools to investigate the local, regional and global controls on atmospheric composition and air quality. To ensure that these models 3 represent the atmosphere adequately it is important to compare their outputs with measurements. 4 5 However, ground based measurements of atmospheric composition are typically sparsely distributed and representative of much smaller spatial scales than those resolved in models, and 6 7 thus direct comparison incurs uncertainty. In this study, we investigate the feasibility of using observations of one or more atmospheric constituents to estimate parameters in chemistry 8 transport models and to explore how these estimates and their uncertainties depend upon 9 10 representation errors and the level of spatial coverage of the measurements. We apply Gaussian process emulation to explore the model parameter space and use monthly averaged ground-level 11 concentrations of ozone  $(O_3)$  and carbon monoxide (CO) from across Europe and the US. Using 12 synthetic observations we find that the estimates of parameters with greatest influence on  $O_3$  and 13 CO are unbiased, and the associated parameter uncertainties are low even at low spatial coverage 14 or with high representation error. Using reanalysis data, we find that estimates of the most 15 influential parameter - corresponding to the dry deposition process - are closer to its expected 16 value using both O<sub>3</sub> and CO data than using O<sub>3</sub> alone. This is remarkable because it shows that 17 18 while CO is largely unaffected by dry deposition, the additional constraints it provides are 19 valuable for achieving unbiased estimates of the dry deposition parameter. In summary, these 20 findings identify the level of spatial representation error and coverage needed to achieve good 21 parameter estimates and highlight the benefits of using multiple constraints to calibrate atmospheric chemistry models. 22

23

- 2 -

# 1 Introduction

2 Changes in atmospheric composition due to human activities make an important contribution to 3 Earth's changing climate (Stocker et al., 2013) and to outdoor air pollution, which is currently responsible for about 4.2 million deaths worldwide each year (Cohen et al., 2017), with 365,000 4 5 deaths due to surface ozone (DeLang et al., 2021). Chemistry transport models (CTMs) simulate the production, transport, and removal of key atmospheric constituents, and are important tools 6 for understanding variations in atmospheric composition across space and time. They permit 7 investigation of future climate and emission scenarios that fully account for the interactions and 8 feedbacks that characterise physical, chemical and dynamical processes in the atmosphere. For 9 practical application, CTMs need to reproduce the magnitude and variation in pollutant 10 11 concentrations observed at a wide range of measurement locations. Where biases occur, these can often be reduced by improving process representation through adjusting model parameters so 12 that the CTM matches the measurements to a sufficient level of accuracy (e.g. Menut et al., 13 2014). While estimation of model parameters is common in many fields of science, and has 14 successfully been applied to climate models (e.g. Chang and Guillas, 2019; Couvreux et al., 15 2021), it is rarely attempted with atmospheric chemistry models because they are 16 computationally expensive to run and it is thus burdensome to perform the large number of 17 model runs required to explore model parameter space. Instead, data assimilation has become a 18 standard method for ensuring that model states are consistent with measurements, usually 19 treating model parameters as fixed (Khattatov et al., 2000, Bocquet et al., 2015, van Loon et al., 20 2000, Emili et al., 2014). 21

In this study, we explore computationally efficient ways of estimating parameters in
chemistry transport models, focusing on two important tropospheric constituents, ozone (O<sub>3</sub>) and

- 3 -

carbon monoxide (CO). Ozone is a major pollutant that is produced in the troposphere by 1 oxidation of precursors such as CO and hydrocarbons, which are emitted during combustion 2 3 processes from vehicular, industrial and residential sources. Ozone is harmful to human health and has been shown to damage vegetation and reduce crop yields (Goldsmith and Landaw, 1968, 4 Kampa and Castanas, 2008, Van Dingenen et al., 2009, van Zelm et al., 2008). A recent 5 6 assessment of surface O<sub>3</sub> was carried out for the Tropospheric Ozone Assessment Report 7 (TOAR) based on measurements from an extensive network of 10,000 sites around the world (Schultz et al., 2017). A simple statistical model of changes in surface  $O_3$  between 2000 and 8 9 2014 showed that significant decreases of 28% and 6% have occurred in Eastern North America and Europe, respectively, but increases of 20% and 45% in south-east and east Asia (Chang et 10 al., 2017). In recent decades, a similar pattern of decreases in CO in Europe and North America 11 and increases over parts of Asia has also been observed (Granier et al., 2011). To fully explain 12 and attribute these changes, a thorough understanding of the processes controlling these 13 14 pollutants is needed.

To assess the performance of CTMs, it is essential to compare simulations of 15 tropospheric chemical composition with measurements. A comprehensive evaluation of 15 16 17 global models found that they broadly matched measured  $O_3$ , but that modelled  $O_3$  was biased high in the northern hemisphere and biased low in the southern hemisphere (Young et al., 2018). 18 19 The models were unable to capture the long-term trends in tropospheric  $O_3$  observed at different 20 altitudes. Similar biases were found in an independent study of long-term trends involving three chemistry climate models (Parrish et al., 2014). While identification of these model biases is 21 22 informative, correcting the deficiencies is challenging because it is often unclear why different 23 models perform well at certain times and for certain places, but poorly elsewhere (Young et al.,

- 4 -

2018). A practical solution is to perform global sensitivity analysis to identify the parameters or
 processes that influence the model results most and then to calibrate the model to estimate these
 parameters and their uncertainties by comparing model predictions with measurements in a
 statistically rigorous way. This provides insight into the physical processes causing model biases
 that is typically unavailable from simpler approaches.

6 The principal challenge with performing global sensitivity analysis and model calibration is that they may require thousands of model runs, and this is infeasible for a typical global CTM 7 that may require 12-24 hours to simulate a year on high performance computing facilities. This 8 9 can be overcome by replacing the model with a surrogate function such as a Gaussian process emulator that is computationally much faster to run (Johnson et al., 2018, Ryan et al., 2018, Lee 10 et al., 2013). Sensitivity analysis and model calibration can then be performed based on 11 thousands of runs with the emulator rather than the CTM. Since the first application of 12 emulation methods for model calibration (Kennedy and O'Hagan, 2001), these approaches have 13 14 been extended to models with highly multivariate output (Higdon et al., 2008). Examples include an earth system model (Wilkinson, 2010), an aerosol model (Johnson et al., 2015), an ice 15 sheet model (Chang et al., 2016) and a climate model (Salter et al., 2018). In this study, we 16 17 apply these approaches to models of tropospheric ozone for the first time to demonstrate the feasibility of parameter estimation. 18

We identify three issues that need to be addressed for successful atmospheric model calibration. 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. These errors associated with spatial representativeness may be important

- 5 -

even for satellite measurements which provide information at a 10 km scale (Boersma et al.,
2016, Schultz et al., 2017). This representation error is distinct from instrument error, which is
often relatively narrow and better understood. The effect of representation errors was explored
in simple terrestrial Carbon model by Hill et al. (2012), who found that as these errors decreased,
the accuracy of parameter estimates improved.

Secondly, the spatial coverage of atmospheric composition measurements is typically
relatively poor, and this limits our ability to estimate parameters accurately. Thus, it is important
to explore how the spatial coverage of measurements affects estimates of model parameters and
their associated uncertainties.

Thirdly, evaluation of atmospheric chemistry models is typically performed for different 10 variables independently (e.g., Stevenson et al., 2006, Fiore et al., 2009). However, atmospheric 11 constituents such as O<sub>3</sub>, CO, NOx, and VOC are often closely coupled through interrelated 12 chemical, physical and dynamical processes. Evaluation of a model with measurements of a 13 14 single species neglects the additional process information available from accounting for species relationships. Lee et al. (2016) highlight the limitation of using a single observational constraint 15 on modelled aerosol concentrations, finding that this resulted in reduced uncertainty in 16 17 concentrations but not in the associated radiative forcing. The benefits of using multiple constraints have been highlighted previously. For example Miyazaki et al. (2012) used the 18 19 Ensemble Kalman Filter and satellite measurements of NO<sub>2</sub>, O<sub>3</sub>, CO and HNO<sub>3</sub> to constrain a 20 CTM, resulting in a significant reduction in model bias in NO<sub>2</sub> column, O<sub>3</sub> and CO concentrations simultaneously. Nicely et al. (2016) used aircraft measurements of O<sub>3</sub>, H<sub>2</sub>O and 21 22 NO to constrain a photochemical box model, and found estimates of column OH that were 12-23 40% higher than those from unconstrained CTMs. They also found that although the CTMs

- 6 -

simulated O<sub>3</sub> well, they underestimated NOx by a factor of two, explaining the discrepancy in
 column OH.

To address these gaps in knowledge, we estimate the probability distributions of eight 3 parameters from a CTM, given surface O<sub>3</sub> and CO concentrations from the USA and Europe. 4 5 We focus on model calibration with a limited number of parameters as a proof of concept, but 6 show how this could be expanded to a much wider range of parameters in future. To overcome the excessive computational burden of running the model a large number of times, we replace the 7 8 model with a fast surrogate using Gaussian process emulation. After evaluation of the emulator 9 to ensure that it is an accurate representation of the input-output relationship of the CTM, we investigate how well model parameters can be estimated from chemical measurement data. We 10 quantify the impacts of measurement representation error and spatial coverage on the bias and 11 uncertainty in the estimated model parameters and highlight the extent to which parameter 12 estimates can be improved using measurements of different variables simultaneously. 13

14 **2. Materials and methods** 

#### 15 2.1 Atmospheric Chemical Transport Model

Chemistry transport models simulate the changes in concentration of a range of atmospheric 16 17 constituents (e.g. O<sub>3</sub>, CO, NO<sub>x</sub>, CH<sub>4</sub>) with time over a specified three-dimensional domain. They 18 represent many of the physical and chemical processes involved, usually in a simplified form, 19 but a detailed understanding is often incomplete. Key processes include the emission of trace 20 gases into the atmosphere, photochemical reactions that result in chemical transformations, 21 transport by the winds, convection and turbulence, and removal of trace gases from the atmosphere through deposition processes. In this study, we apply the Frontier Research System 22 23 for Global Change version of the University of California, Irvine chemical transport model, the

- 7 -

FRSGC/UCI CTM (Wild and Prather, 2000; Wild et al, 2004). We focus on eight important 1 processes affecting tropospheric oxidants that were chosen based on one-at-a-time sensitivity 2 3 studies with the model (Wild, 2007) and that have been used in previous global sensitivity analyses of tropospheric ozone burden and methane lifetime (Ryan et al., 2018; Wild et al., 4 2020). These processes include the surface emissions of nitrogen oxides (NOx), lightning 5 6 emissions of NO, biogenic emissions of isoprene, wet and dry deposition of atmospheric constituents, atmospheric humidity, cloud optical depth and the efficiency of turbulent mixing in 7 8 the boundary layer, see Table 1. These do not encompass all sources of uncertainty in the model, 9 but are broadly representative of major uncertainties across a range of different processes. To provide a simple and easily interpretable approach to calibration, we define a global scaling 10 factor for each process that spans the range of uncertainty in the process and that is applied 11 uniformly in space and time. These scaling factors form the parameters that we aim to calibrate. 12 The choice of parameters and uncertainty ranges are described in more detail in Wild et al. (2020). 13 14 For this study, we focus on monthly-mean surface  $O_3$  and CO distributions at the model native grid resolution of 2.8°×2.8° and compare with observations over North America and Europe for 15 model calibration (Fig. 1). The model uses meteorological driving data for 2001, a relatively 16 17 typical meteorological year without strong climate phenomena such as El Nino (Fiore et al. 2009). 18

# 19 2.2 Surface $O_3$ and CO data

Ground-based observations of  $O_3$  are relatively abundant in Europe and North America, where there are ~1800 individual sites that have continuous long-term measurements of  $O_3$  (Chang et al., 2017, Schultz et al., 2017). Measurements of CO are made at fewer locations, but reliable long-term data are available from 57 sites that are part of the Global Atmospheric Watch

- 8 -

1	network (Schultz et al., 2015). To allow more thorough testing of the effects of spatial coverage
2	over these regions, we use CAMS interim reanalysis data of surface O <sub>3</sub> and CO from the
3	European Centre for Medium-Range Weather Forecasts (ECMWF) which has been tuned to
4	match measurements using 4D-Var data assimilation (Flemming et al. 2017). This reanalysis
5	reproduces observed O <sub>3</sub> and CO distributions relatively well, and biases at surface measurement
6	stations are generally small (Huijnen et al., 2020). The dataset also has the benefit of complete
7	global coverage, allowing us to test the importance of measurement coverage directly.
8	Reanalysis data for $O_3$ and CO are available for 2003–2015, and we average the data by
9	month across this period to provide a climatological comparison. The control run of the
10	FRSGC/UCI model matches CO from the reanalysis data reasonably well (Fig. 2), but
11	overestimates surface O <sub>3</sub> . Overestimation of O <sub>3</sub> in continental regions has been noted in previous
12	studies and is partly a consequence of rapid photochemical formation from fresh emissions that
13	is magnified at coarse model resolution (Wild and Prather, 2006). For this exploratory study we
14	bias-correct the modelled surface O <sub>3</sub> by reducing it by 25%, following the approach taken by
15	Shindell et al. (2018), so that it matches the reanalysis data (Fig. 2a). This adjustment accounts
16	for the effect of chemical processes and model resolution which are not explored in this study,
17	and provides a firmer foundation for investigating the effects of other processes.

## 18 2.3 Representation error

The "representation error" describes how well measurements made at a single location represent a wider region at the spatial scale of the model (2.8°×2.8° for this study). The error may be reduced by averaging measurements made at different stations within a model grid box, although atmospheric measurements may be too sparse to permit this (Schultz, 2016). The representation error is sometimes taken as the mean of the spatial standard deviation of different measurements

- 9 -

within a grid-box (Sofen et al. 2016). However, this measure quantifies the spatial variability of
 measured O<sub>3</sub> within a grid-box and may not match the representation error

To test the effect on parameter estimates of varying this representation error, we use synthetic data from the control run of the model using parameters set to their nominal default values. Synthetic O<sub>3</sub> and CO data were generated by adding different levels of representation error for each level of spatial coverage. In mathematical terms:

$$data_i = m_i(x_{control}) + \varepsilon_i, \quad \varepsilon_i \sim N(0, \sigma_i^2)$$
(1)

7 where for the *i*th point in space or time, *data<sub>i</sub>* refers to the synthetic data for O<sub>3</sub> or CO,
8 m<sub>i</sub>(x<sub>control</sub>) is the O<sub>3</sub> or CO from the model control run, and ε<sub>i</sub> is generated from a Normal

9 distribution with mean of zero and standard deviation  $\sigma_i$  that is directly proportional to the

10 magnitude of  $m_i(x_{control})$ . In this case,  $\sigma_i = p \times m_i(x_{control})$  where p is a scaling factor that

11 provides a measure of the representation error. We used the reanalysis data to estimate *p* 

12 alongside the model parameters, and found posterior values of p that were in the range 0.16-0.19.

13 We therefore selected four values of p (0.01, 0.1, 0.2 and 0.3) to explore the importance of

14 representation error when using the synthetic data.

#### 15 *2.4 Global sensitivity analysis*

Sensitivity analysis was carried out to determine the sensitivity of the simulated surface O<sub>3</sub> and CO to changes in each of the eight parameters. This allows us to identify which of the parameters are most important in governing surface O<sub>3</sub> and CO. We use global sensitivity analysis (GSA), varying each input while averaging over the other inputs. This provides a more integrated assessment of uncertainty than the traditional one-at-a-time approach varying each input in turn while fixing the other inputs at nominal values. We use the extended FAST method (Saltelli et al., 1999), a common and robust approach to GSA in which the sensitivity indices are

- 10 -

quantified by partitioning the total variance in the model output (i.e. modelled surface O<sub>3</sub> or CO)
into different sources of contribution from each input. Like most sensitivity analysis methods,
this approach requires several thousand executions of the model, which would be
computationally expensive for the CTM used here. This is overcome by replacing the CTM with
a Gaussian process (GP) emulator. Further details of the implementation of GSA are described
in Ryan et al. et al. (2018).

#### 7 2.5 Gaussian Process Emulation - theory

8 We replace the CTM with a surrogate model that maps the inputs of the CTM (the eight 9 parameters listed in Table 1) with its outputs (surface  $O_3$  and CO). We employ a surrogate model based on Gaussian process (GP) emulation for three reasons. Firstly, due to the attractive 10 mathematical properties of a GP, the emulator needs very few runs of the computationally 11 12 expensive model to train it, typically less than 100. This is in contrast to methods based on 13 neural networks, which often have a large number of parameters that necessitate thousands of 14 training runs. Secondly, a GP emulator is an interpolator and so predicts the output of the model with no uncertainty at the input points it is trained at. Thirdly, it gives a complete probability 15 distribution, as a measure of uncertainty, for estimates of the model output at points it is not 16 17 trained at.

18 A GP is an extension of the multivariate Gaussian distribution, where instead of a mean 19 vector  $\mu$  and covariance matrix  $\Sigma$ , mean and covariance functions given by E(f(x)) and 20  $\operatorname{cov}(f(x), f(x'))$  are used (Rasmussen, 2006). Here,  $f(\cdot): \chi \in \mathbb{R}^q \to \mathbb{R}^{q'}$  represents the 21 computationally expensive model and  $\chi$  denotes the input space given by  $x = (x_1, \dots, x_q) \in$ 22  $\chi_1 \times \dots \times \chi_q = \chi \subset \mathbb{R}^q$ , and q is the number of input variables. GP emulators within a Bayesian 23 framework were first developed in the 1990s and early 2000s (O'Hagan, 2006, Oakley and

- 11 -

O'Hagan, 2004, Kennedy and O'Hagan, 2000, Currin et al., 1991). The simplest and most 1 common GP emulator is one where the outputs to be emulated are scalar. Thus, if the 2 3 computationally expensive model is given by  $f(\cdot)$ , then the one-dimensional output y is calculated by y = f(x). This means that if the model output is multidimensional – e.g. a global 4 map or a time-series – then we need to build a separate emulator for each point in the output 5 space. To build the emulator requires training runs from the expensive model. In general, we 6 choose *n* training inputs, denoted by  $\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_n$ , based on a space filling design such as a 7 8 Maximin Latin Hypercube design (Morris and Mitchell, 1995). The number of training points is 9 based on the rule of thumb  $n = 10 \times q$  (Loeppky et al., 2012). Denoting the scalar outputs by  $y_1 = f(\mathbf{x}_1), y_2 = f(\mathbf{x}_2), ..., y_n = f(\mathbf{x}_n)$ , we then build 10 an emulator  $\hat{f}(\cdot)$  given by  $\hat{y} = \hat{f}(x)$ , where  $\hat{y}$  is the estimated output from the emulator. If x 11 represents one of the training inputs (i.e.  $x = \mathbf{x}_i, 1 \le i \le n$ ), then  $\hat{y}$  is equal to the output from 12  $f(\cdot)$  with no uncertainty (i.e.  $\hat{y} = y$ ). If x represents an input the emulator is not trained at, then 13

\$\heta\$ has a probability distribution represented by a mean function \$m(x)\$ and a covariance function
\$V(x, x')\$, where \$x'\$ is a different input. The mean function is given by:

$$m(x) = h(x)^T \hat{\beta} + t(x)^T \mathbf{A}^{-1} \big( \mathbf{y} - H \hat{\beta} \big),$$
<sup>(2)</sup>

16 where  $h(x)^{T}$  is a 1×(q+1) vector given by  $(1, x^{T})$ ,  $\hat{\beta}$  is a vector of coefficients determined by 17  $\hat{\beta} = (H^{T}\mathbf{A}^{-1}H)^{-1}H^{T}\mathbf{A}^{-1}\mathbf{y}, t(x)^{T} = (C(x, x_{1}; \psi), ..., C(x, x_{n}; \psi))$ , and  $\mathbf{A}$  is a matrix whose ele-18 ments are determined by  $\mathbf{A}_{i,j} = C(\mathbf{x}_{i}, \mathbf{x}_{j}; \psi), \mathbf{y} = [f(\mathbf{x}_{1}), ..., f(\mathbf{x}_{n})]^{T}, H = [h(\mathbf{x}_{1}), ..., h(\mathbf{x}_{n})]^{T}$ . 19 Here,  $C(x, x'; \psi)$  is a correlation function that represents our prior belief about how the inputs x20 and x' are correlated. A common choice is a Gaussian correlation function which takes the form: 21  $C(x, x'; \psi) = exp(-(x - x')^{T}\mathbf{B}(x - x'))$ , where  $\mathbf{B}$  is a  $p \times p$  matrix with zeros in the off-22 diagonals and diagonal elements given by the roughness parameters  $\psi = (\psi_{1}, ..., \psi_{q})$ . The roughness parameters give an indication of whether the input-output relationship for each input
 variable, given the training data, should be linear. Low values reflect a linear (or smooth)
 relationship, whereas high values (e.g. > 20) suggest a non-linear (or non-smooth) response
 surface. For implementation purposes we express the correlation function as C(x, x'; ψ) =

5 
$$\sum_{j=1}^{q+1} \exp\left(-\psi_j(x_j - x_j')^2\right)$$
, where  $x = (x_1, \dots, x_q)$  and  $x' = (x_1', \dots, x_q')$ . The formula for the  
6 covariance function  $V(x, x')$  is given in appendix A.

A final issue to resolve is how to estimate the roughness parameter since the posterior
distribution of f(·) is conditional on these emulator parameters. A Bayesian approach would be
to integrate out these emulator parameters in the formulation of the GP emulator. This would
require highly informative priors, but in most cases such informative priors do not exist.
Kennedy and O'Hagan (2001) propose using maximum likelihood to provide a point estimate of
the emulator parameters and to use these in the formulae for the mean and covariance functions
of the GP emulator. We adopt this approach in this study.

## 14 2.6 Gaussian Process Emulation - implementation

22

Using the Loeppky rule we choose n = 80 different training inputs for our eight-parameter 15 calibration study. In total, we emulate two variables (surface  $O_3$  and CO) over 12 months at 272 16 spatial locations, and so require 6528 different GP emulators. To estimate the model parameters 17 we evaluate each of the GP emulators tens of thousands of times. Although emulation is 18 computationally fast, this presents a substantial computational burden, even for more 19 computationally efficient versions of the emulator (Marrel et al., 2011, Roustant et al., 2012). 20 21 We overcome this by computing parts of equation (2) prior to these evaluations. Specifically, we compute the vectors  $\hat{\beta}$ ,  $m_{LP}$  and  $\psi$  for all points in the output space, where  $m_{LP}$  denotes 22  $\mathbf{A}^{-1}(\mathbf{y} - H\hat{\boldsymbol{\beta}})$ , the last part of m(x) from equation (2). We store these three objects as three 23

1 matrices  $\hat{\beta}_{ALL}$ ,  $m_{LP.ALL}$  and  $\psi_{ALL}$ . Evaluated at a new input  $x_{new}$ , the mean function of the 2 emulator (equation 1) can now be expressed as:

$$m_{i}(x_{new}) = h(x_{new})^{T} \hat{\beta}_{ALL}[i,:] + t_{i}(x_{new})^{T} m_{LP.ALL}[i,:],$$

$$t_{i}(x_{new})^{T} = (C(x_{new}, x_{1}; \psi_{ALL}[i,:]), \dots, C(x_{new}, x_{n}; \psi_{ALL}[i,:])),$$
(3)

where i (1 ≤ i ≤ 6528) denotes the ith point in the output space. The equivalent formula for
V(x, x') is given in appendix A.

To the test the accuracy of GP emulation, we ran each of the 6528 emulators at 20 sets of parameters which were not used for training the emulators. The estimated  $O_3$  and CO values from the emulators for all spatial locations and months closely match the simulated  $O_3$  and CO output from the FRSGC/UCI model for these validation runs, with  $R^2 > 0.995$  for each variable, see Fig. 3.

10 Finally, we recognise that principal component analysis (PCA) could be used to reduce 11 the dimensionality of the output space and hence the number of emulators required (Higdon et 12 al., 2008). In a previous study we found that a PCA-emulator hybrid approach resulted in similar 13 performance compared to using separate emulators for each point in the output space, and 14 reduced the number of emulators required from 2000 to 40 or fewer (Ryan et al., 2018). However, for this study, we choose an emulator-only approach because it is much simpler to 15 16 demonstrate. Nonetheless, future emulation-calibration studies could benefit from the 17 computational savings of applying a PCA-emulator hybrid approach. Other approaches for dealing with high dimensional output are also available, such as low rank approximations 18 19 (Bayerri et al.,2007).

#### 20 2.7 Parameter Estimation

We estimate the eight model parameters using Bayesian statistics via the software package Just Another Gibbs Sampler (Plummer, 2003). This uses Gibbs sampling, which is an approach based on Markov Chain Monte Carlo (MCMC) that we use to determine multi-dimensional posterior probability distribution of the model parameters (Gelman et al., 2013). Gibbs sampling is an extension of the more traditional Metropolis-Hasting variant of MCMC, and uses conditional probability to sample from the marginal distribution when moving around the multidimensional parameter space.

To find the posterior distribution, the MCMC algorithm searches the parameter space 8 9 using multiple sets of independent chains. Here, a chain refers to a sequence of steps in the parameter space that the algorithm takes. A new proposed parameter set in this search is 10 accepted on two conditions: (1) the set is consistent with the prior probability distribution, which 11 for our study was a set of Uniform distributions with the lower and upper bounds given by the 12 defined ranges in Table 1; and (2) the resulting modelled values using the proposed set of 13 14 parameters are consistent with measurements, which is assessed using the following Gaussian likelihood function: 15

$$L(\theta) = \prod_{i=1}^{N} \frac{1}{\sqrt{2\pi\sigma_i}} exp\left(\frac{f_i(\theta) - m_i}{\sigma_i^2}\right)^2,\tag{4}$$

where *N* is the number of measurements used,  $f_i(\theta)$  is the *i*th model output  $(1 \le i \le N)$  using the proposed parameter set  $\theta$ ,  $m_i$  is the measurement corresponding to the *i*th model output and  $\sigma_i$  is the representation error for measurement  $m_i$ . We note that although separate emulators are used for each of the spatial and temporal locations in the model output, there is still only a single likelihood function. Hence, evaluating all of the emulators for a specific set of values of the scaling parameters is equivalent to evaluating the CTM once at those values of the parameters.

We ran three parallel chains for 10,000 iterations each. After discarding the first half of these iterations as 'burn in', we thinned the chains by a factor of five to reduce within-chain autocorrelation. Convergence was assessed using the Brooks-Gelman-Rubin diagnostic tool (Gelman et al., 2013). This produced 3000 independent samples from the posterior distribution for each parameter, which we summarize using their posterior means and 95% credible intervals (CIs) defined by the 2.5<sup>th</sup> and 97.5<sup>th</sup> percentiles (Gelman et al., 2013). We used the R language to code up our configuration of the MCMC algorithm.

#### 8 *2.8 Model discrepancy*

9 It has been suggested that a model discrepancy term should be included when carrying out model
10 calibration involving Gaussian process emulators (e.g. Kennedy and O'Hagan, 2001;

11 Brynjarsdóttir and O'Hagan, 2014). The discrepancy term represents the processes missing in 12 the model. However, in this demonstration study we have chosen not to include a discrepancy 13 term for two reasons. Firstly, for scenarios where we use synthetic data, no discrepancy term is 14 required because the synthetic data is generated by adding noise and spatial gaps to the emulator output for the control run. Secondly, for scenarios involving reanalysis data, there is no simple 15 16 and defensible method to estimate the term. When performing model calibration by applying 17 MCMC directly, a discrepancy term would not be included. Since the purpose of the emulator here is to estimate the output of the model for a given set of parameter values, we argue that it is 18 not necessary to include a discrepancy term into the calibration formulation. However, we agree 19 20 that including such a term may be helpful in situations where there is good prior information.

To investigate the importance of a discrepancy term, we repeat the experiment to estimate the eight scaling parameters using surface ozone reanalysis data and assuming a discrepancy term that is 10% of the magnitude of the observation. We find that there is almost no difference

- 16 -

in the marginal posterior distribution when we include the discrepancy term compared with when
 we omit it (see Figure S16 in the supplemental material). We therefore choose to omit the term
 for our study.

# 4 2.9 Experimental approach

We first perform a global sensitivity analysis to identify the parameters which have the greatest 5 influence on the two variables we consider. We then perform parameter estimation using surface 6 7 concentration data over the regions of North America and Europe shown in Fig 1 and focus our 8 analysis on the parameters which have the greatest influence. To provide a demonstration of the approach we first use "synthetic" measurement data drawn from the control run of the CTM 9 which was not used to train the emulators, adding increasing levels of noise to represent 10 11 measurement representation errors of 1, 10, 20 and 30% (p = 0.01, 0.1, 0.2 and 0.3), and varying 12 the spatial coverage of these measurements over the regions considered over a wide range: 2.5, 5, 10, 20, 40 and 100%. We focus on surface O<sub>3</sub> only, surface CO only and then both variables 13 14 together. We then use the reanalysis data to represent the measurements, focussing on the effects of spatial coverage alone, and estimating the representation error p from this independent dataset. 15 16 The 90 different scenarios we consider are summarised in Table 2. We discuss the implication 17 of these results and the limitations of considering a simple eight-parameter system rather than all sources of model uncertainty in Section 4. 18

# 19 **3. Results**

#### 20 *3.1 Global sensitivity analysis*

Results from global sensitivity analysis reveal that over the continental regions of Europe and
North America considered here, the simulated monthly mean concentrations of surface O<sub>3</sub> are

- 17 -

most sensitive to dry deposition and, to a lesser extent, to isoprene emissions (Fig. 4). This is not 1 unexpected, given the importance of direct deposition of ozone to the Earth's surface, and the 2 role of isoprene as a natural source of ozone in continental regions. The simulated surface CO is 3 most sensitive to isoprene emissions, which represent a source of CO, and to boundary layer 4 mixing, which influences the transport of CO from polluted emission regions. We thus identify 5 6 the scaling parameters corresponding to dry deposition, isoprene emissions and boundary layer 7 mixing as the most important of the eight considered here to estimate accurately to reduce the 8 bias in modelled surface  $O_3$  and CO. For completeness, we show the geographical distribution 9 of sensitivity indices in Figs 5 and 6, which reveal the importance of humidity in governing O<sub>3</sub> over oceanic regions and highlight the very different responses of surface O<sub>3</sub> and CO to the 10 major driving processes. 11

### 12 *3.2 Estimation of scaling parameters using synthetic data*

13 We next use synthetic observation data to calibrate the model and estimate scaling parameters. 14 For synthetic data we use the model control run with a specified level of representation error (Table 2), and the default model parameters define the true scaling that we aim to retrieve. 15 16 Prescribing surface O<sub>3</sub> with very little error (p = 0.01) gives an estimate of the dry deposition 17 scaling parameter, which has the largest influence on modelled surface O<sub>3</sub>, close to its true value and the uncertainty is small even when the spatial coverage of measurements is only 2.5% (Fig. 18 7, column 1). As the representation error is increased to p = 0.1, the parameter uncertainty is 19 20 larger at low spatial coverage but the mean estimate remains unbiased (Fig. 7, column 2). The uncertainty at all levels of spatial coverage becomes larger as p increases to 0.2 and 0.3, but the 21 means remain very close to the true values (Fig. 7, columns 3 and 4). Surface CO is largely 22 unaffected by dry deposition, and thus provides very little constraint on the scaling parameter. 23

- 18 -

The effect of prescribing surface CO and O<sub>3</sub> together is very similar to that of using surface O<sub>3</sub>
alone.

3	Using surface CO alone with very little representation error ( $p = 0.01$ ), the mean estimate
4	of the isoprene emission scaling parameter is equal to the true value with very little uncertainty,
5	regardless of the spatial coverage (Fig. 8, column 1). When the representation error is increased
6	to $p = 0.1$ , the estimate remains very close to the true value, but the uncertainty is substantially
7	higher at low spatial coverage (2.5% and 5%) than at higher coverage (40% and 100%) (Fig. 8,
8	column 2). The estimates deviate further from the truth at higher levels of representation error ( $p$
9	= 0.2 and 0.3) and the uncertainty is greater (Fig. 8, columns 3 and 4). Estimates of the isoprene
10	scaling parameter are less accurate than those of the dry deposition scaling parameter as the
11	posterior means are further from the true value of the parameter and the uncertainty intervals are
12	wider (Fig. 8 vs Fig. 7). As with our findings for dry deposition, the posterior means and the
13	lengths of the uncertainty intervals for the isoprene scaling parameter remain relatively
14	unchanged when surface O <sub>3</sub> data is prescribed at the same time.
15	Our findings for the boundary layer mixing scaling parameter follow a similar pattern to
16	the other two parameters (Fig. 9). In all combinations of representation error and spatial
17	coverage, we find that the mean estimates are unbiased. Furthermore, we find that the parameter
18	uncertainty is significantly smaller when the spatial coverage is 10% or higher when $p = 0.1$ ,
19	20% or higher when $p = 0.2$ , and 40% or higher when $p = 0.3$ (Fig. 9, Table 2). It is clear from
20	these results that the scalings for these three model parameters can be successfully estimated
21	from synthetic data with low uncertainty when the representation error is low, and that the
22	estimates remain good, albeit with higher uncertainty, at higher representation error if the spatial
23	coverage is relatively good.

- 19 -

## 1 3.3 Estimation of scaling parameters using reanalysis data

We consider next the CAMS interim reanalysis data for surface O3 and CO which are based on 2 assimilated concentrations from the ECMWF model and are thus independent of the 3 FRSGC/UCI model. The reanalysis is representative of similar spatial scales to the FRSGC/UCI 4 5 model, and thus we ignore the representation error and vary the spatial coverage only. However, 6 we are able to estimate the representation error factor p by treating it as a parameter to estimate. With 100% spatial coverage, this error term is estimated with the MCMC algorithm to be p =7  $0.168 \pm 0.004$  and  $p = 0.191 \pm 0.005$  for surface O<sub>3</sub> and CO, respectively. Although we do not 8 9 know the true values of the parameters in this case, the good agreement between the control run of the FRSGC/UCI model and the reanalysis data suggests that they lie close to their true values. 10 Using the reanalysis data for surface O<sub>3</sub> alone, we find that the posterior means and 11 uncertainty for the dry deposition parameter are in the upper half of the range defined, indicating 12 that the real dry deposition flux is greater than that calculated with the FRSGC/UCI model. This 13 14 is largely as expected, as the FRSGC/UCI model overestimates surface O<sub>3</sub> at these continental sites and greater deposition would bring the model into better agreement with the reanalysis. As 15 the spatial coverage is increased, the estimate of the scaling factor increases to around 1.4 and 16 17 the uncertainty is reduced (Fig. 10a). In contrast, using surface O<sub>3</sub> and CO together results in an estimate closer to 1 and an additional reduction in uncertainty (Fig. 10g). Inclusion of surface 18 19 CO measurements, as an additional constraint to surface  $O_3$ , results in an estimate of the dry 20 deposition parameter closer to that modelled. Using surface CO alone, estimates of the isoprene scaling parameter lie in the central part 21

of the defined range, whilst estimates of the boundary layer mixing scaling parameter lie in the
upper half of the defined range (Fig 10e,f). For both parameters, increasing the spatial coverage

- 20 -

leads to a reduction in uncertainty. Unlike for dry deposition, inclusion of surface O<sub>3</sub> when
 estimating either of these parameters results in very little difference in the magnitude of the
 estimate or in the associated uncertainty (Fig. 10e vs 10h; Fig. 10f vs 10i).

# 4 *3.4 Evaluation of surface O<sub>3</sub> following calibration*

We demonstrate the benefit of the calibration by evaluating the emulators using the values of the 5 scaling parameters sampled from the prior and posterior distributions. As an example, we show 6 7 surface  $O_3$  before and after calibration using the calibration runs involving synthetic data at 20% spatial coverage and a representation error of p=0.2 (Figure 11). Despite the calibration 8 9 involving only 20% spatial coverage, we apply the resulting parameter values to all grid squares. We can clearly see that the prior surface O<sub>3</sub> concentrations are unbiased but have large 10 11 uncertainty, especially at high values. In contrast the calibrated O<sub>3</sub> concentrations have a small 12 uncertainty, demonstrating that even with 20% spatial coverage in the calibration data we are 13 able to achieve improved predictions for all model grid boxes.

# 14 **4. Discussion**

#### 15 4.1 Representation error

Our results show the impact of the size of the representation error on the accuracy of estimated model parameters. The parametric uncertainty (i.e. the size of the credible intervals in Figs 7-9) increases at an approximately linear rate as the representation error increases from p = 0.01 to p= 0.3. This is consistent with Hill et al. (2012) who estimated the parameters and uncertainties of a simple terrestrial carbon model under varying levels of measurement error.

For the reanalysis data, we treat the representation error as a parameter for the MCMC
algorithm to estimate along with the eight model parameters. This is possible because we

- 21 -

assume that the measured value of  $O_3$  is proportional to the simulated value from a forward run 1 of the FRSGC/UCI model, although such an assumption may not be possible in other situations. 2 3 An alternative approach to estimate the representation error would be to carry out an intensive measurement campaign to determine whether the average O<sub>3</sub> from different measuring stations 4 5 within a grid-square is representative of the true average. Satellite products of the terrestrial 6 biosphere are checked for accuracy using this type of approach (De Kauwe et al., 2011). Although measurement campaigns at these large spatial and temporal scales would be 7 8 challenging and costly, they may not be need to continue for long periods of time since we might 9 expect representation error to decrease as the temporal scale increases (Schutgens et al., 2016).

#### 10 *4.2 Spatial coverage*

11 We find that as the volume of measurements increase, the estimates of the model parameters are 12 closer to the truth and the width of the credible intervals decrease. This is particularly clear for 13 the dry deposition and isoprene emission scaling parameters when using both  $O_3$  and CO 14 concentrations (Figs 8 and 9). While this highlights the value of good spatial coverage, we note 15 that the benefits are greatly reduced if the representation error is relatively high. For the 16 boundary layer mixing parameter, we find little decrease in the credible intervals using synthetic 17 CO data with the highest representation error (p = 0.3), where the spatial coverage is less than 20% (Fig. 9, row 2). In contrast, at the p = 0.1 level, a large decrease in uncertainty is seen 18 between the 2.5% and 20% coverage. Similar effects are seen, to a lesser extent, for the dry 19 20 deposition and isoprene scaling parameters as the spatial coverage increases.

Our results using synthetic data show that while the size of the uncertainty intervals vary substantially depending on the spatial coverage or representation error, the posterior means are for the most part very close to the true values. Deviation from these typically occurs when the

- 22 -

measurements contain less information either due to low spatial coverage or high representation
error. However, the uncertainty intervals include the true values of the parameters for all the
experimental scenarios considered here, unlike in Hill et al. (2012). This gives strong confidence
in the reliability of the MCMC method used to estimate the parameters.

## 5 *4.3 Applying multiple constraints*

The importance of multiple constraints was most apparent for scenarios involving the 6 7 reanalysis data. For the dry deposition scaling parameter, which explains much of the variance 8 in surface  $O_3$  (Fig. 4), we found that using  $O_3$  data alone results in mean estimates that are in the 9 upper half of the range of possible values (Fig 10a). However, including CO data brought the mean estimates into the central part of the range where we would expect the true value to lie 10 11 (Fig. 10g). This is remarkable given that dry deposition is not an important process for 12 controlling CO, and highlights the coupling between processes that permits constraints on one 13 process from one variable to influence those on another. However, it is consistent with previous 14 studies exploring the uncertainty in estimates of key parameters in an aerosol-chemistry-climate model (Johnson et al., 2018). For the isoprene emission and boundary layer mixing scaling 15 parameters, there was little difference in the mean estimates or the size of the uncertainty 16 17 intervals when using O<sub>3</sub> and CO together rather than a single constraint. This reveals that the importance of using multiple constraints is dependent on the process and on the variable 18 constrained. A judicious choice of these could allow a particular process to be targeted. 19 20 Overall, our estimates of the dry deposition and isoprene emission scaling parameters are close to a priori values from the FRSGC/UCI CTM, with respect to the independent reanalysis data. In 21 contrast, our estimates of the boundary layer mixing scaling parameter are substantially larger 22

than those from the model, suggesting that this process is not represented well in the model, or
that other processes not considered here may be influencing the result.

#### 3 *4.4 Towards constraint with real surface measurements*

Our results have demonstrated the feasibility of using measurement data to constrain model 4 parameters under the right conditions. We have chosen to use synthetic data as they have allowed 5 us to vary the spatial coverage and to investigate the effects of representation error which is 6 7 poorly characterised when using real measurements data. Quantifying this type of error for real 8 measurements is difficult because measurement sites are relatively sparse and are often 9 representative of a limited area rather than the larger area typical of a model grid-square. 10 However, this study has allowed us to estimate the representation error associated with the 11 reanalysis data, and in the absence of more information these values could be used as a guide 12 when applying surface measurements as a constraint.

13 The reanalysis data provide a more critical test, as they are independent of the 14 FRSGC/UCI CTM used here. Although we do not know the true values of the scaling parameters, we expect them to lie close to those used in the control run given the relatively good 15 16 agreement for O<sub>3</sub> and CO concentrations. For the dry deposition parameter, we expect scaling 17 values to be close to 1, but using surface O<sub>3</sub> reanalysis data alone we found posterior mean scaling parameters approaching 1.4, with credible intervals that did not include 1 (Fig. 10a). 18 This likely reflects overestimation of surface O<sub>3</sub> in continental regions in the CTM and may 19 20 reflect uncertainties and biases in other processes not considered here, most notably in the chemical formation and destruction of O3 and in model transport processes. In the absence of 21 consideration of the uncertainty in these processes in this feasibility study, the dry deposition 22 parameter is used as a proxy process to reduce O<sub>3</sub> concentrations. This is an example of 23

- 24 -

equifinality, where different sets of parameters can result in model predictions that give equally
good agreement with observations (Beven et al., 2001). Applying simultaneous constraints to
CO goes some way to addressing this, but does not remove the problem. Before applying real
surface measurements to constrain the CTM, we propose a more comprehensive assessment of
model uncertainties with a wider range of parameters so that the constraints can more directly
inform process understanding and model development.

## 7 Conclusion

8 We have demonstrated the use of surface O<sub>3</sub> and CO concentrations to constrain a global 9 atmospheric chemical transport model and generate accurate and robust estimates of model parameters. This would normally be prohibitive for such a model given that thousands of model 10 runs are required. Our approach is to replace the CTM with a surrogate model using Gaussian 11 process emulation and then estimate the parameters using the emulator in place of the CTM. In 12 13 this feasibility study we have shown that surface  $O_3$  has a large sensitivity to dry deposition, and that surface CO is most sensitive to isoprene emissions and boundary layer mixing processes, as 14 expected. We find that estimates of the scaling parameters for these processes are dependent on 15 16 the spatial coverage and representation error of the surface  $O_3$  and CO data. Our parameter estimates become less uncertain as coverage increases and as the representation error decreases, 17 whilst remaining unbiased. Furthermore, we show that using two separate data constraints, in 18 19 this case surface  $O_3$  and CO, instead of a single one can result in mean parameter estimates that 20 are much closer to their likely true values. However, this is dependent on the processes considered and constraints applied, and while it is effective for dry deposition here, we find 21 22 relatively little improvement in the estimates or uncertainties for isoprene emission or boundary 23 layer mixing processes that are also considered here.

- 25 -

The approach we adopt here provides a means of constraining atmospheric models with 1 observations and identifying sources of model error at a process level. Our results based on the 2 independent reanalysis data suggest that dry deposition and isoprene emissions are represented 3 relatively well in the FRSGC/UCI CTM but that boundary layer mixing processes may be 4 somewhat underestimated. However, we have explored the effect of only eight parameters in 5 6 this study and consideration of a more complete set of processes, including those governing photochemistry and dynamics, is needed to generate more realistic constraints for key pollutants 7 such as O<sub>3</sub>. We aim to expand this study to investigate a more extensive range of parameters and 8 9 processes and to constrain with a wider range of observation data. The emulator-based approach for estimating parameters that we have successfully demonstrated here can be applied to any 10 model where evaluating the model the required number of times is too computationally 11 demanding. 12

## 13 Code and data availability

The R code used for building and validating the emulators and estimating the posterior
distribution of the model parameters using the Markov Chain Monte Carlo algorithm is available
from the Zenodo data repository via the link: <u>https://zenodo.org/record/4537614</u>. The
FRSGC/UCI model output used for training the emulators is available from the CEDA data
repository via the link: <u>https://catalogue.ceda.ac.uk/uuid/d5afa10e50b44229b079c7c5a036e660</u>.

# 19 Appendix A

20 The formula for the covariance function V(x, x') from §2.2 is given by:

21 
$$V(x, x') = \sigma^{2} [C(x, x'; \psi) - t(x)^{T} \mathbf{A}^{-1} t(x)]$$

22 + 
$$(h(x)^T + t(x)^T \mathbf{A}^{-1} H)(H^T \mathbf{A}^{-1} H)^{-1}(h(x')^T + t(x')^T \mathbf{A}^{-1} H)^T]$$

- 26 -

1 where,

2 
$$\sigma^{2} = \frac{\mathbf{y}^{\mathrm{T}} (\mathbf{A}^{-1} - \mathbf{A}^{-1} H (H^{\mathrm{T}} \mathbf{A}^{-1} H)^{-1} H^{\mathrm{T}} \mathbf{A}^{-1}) \mathbf{y}}{n - q - 1}$$

To compute the variance or uncertainty of a prediction x we use the formula for V(x, x') with x' = x, which results in  $C(x, x; \psi) = 1$ . Since we need to evaluate a large number of emulators for each MCMC iteration step (because we have a separate emulator for every dimension of the model output), it is more computationally efficient to compute the parts of the above formula prior to using the emulator. Hence, the above formula can be replaced with:

8 
$$V_i(x_{new}, x_{new}) = \sigma_{ALL}^2[i, 1] [(1 - t_i(x_{new})^T V_{i,1} t_i(x_{new}))]$$

9 + 
$$(h(x_{new})^T + t(x_{new})^T V_{i,2}) V_{i,3} (h(x_{new})^T + t(x_{new})^T V_{i,2})^T$$

10 where:

- 11  $i (1 \le i \le r)$  denoted the *i*th point in the *r*-dimensional simulator output.
- 12  $\sigma_{ALL}^2$  is a  $r \times 1$  vector that stores the values of  $\sigma^2$  for all r outputs.
- 13  $V_{i,1}$  is the  $n \times n$  matrix  $\mathbf{A}^{-1}$  corresponding to the *i*th point in the simulator's output. It is 14 stored as the *i*th block of the  $nr \times n$  matrix  $V_1$  defined by:

15 
$$V_1 = \begin{pmatrix} V_{1,1} \\ V_{2,1} \\ \vdots \\ V_{r,1} \end{pmatrix}$$

•  $V_{i,2}$  is the  $n \times q$  matrix  $\mathbf{A}^{-1}H$  corresponding to the *i*th point in the simulator's output. It is stored as the *i*th block of the  $nr \times q$  matrix  $V_2$  defined by:

18 
$$V_2 = \begin{pmatrix} V_{1,2} \\ V_{2,2} \\ \vdots \\ V_{r,2} \end{pmatrix}$$

*V<sub>i,3</sub>* is the *q* × *q* matrix (*H*<sup>T</sup>**A**<sup>-1</sup>*H*)<sup>-1</sup> corresponding to the *i*th point in the simulator's output. It is stored as the *i*th block of the *qr* × *q* matrix *V*<sub>3</sub> defined by:

$$V_3 = \begin{pmatrix} V_{1,3} \\ V_{2,3} \\ \vdots \\ V_{r,3} \end{pmatrix}$$

# 4 Author contributions

- 5 ER and OW designed the study. ER carried out the statistical analyses, and OW ran the
- 6 FRSGC/UCI model and provided the outputs that were used to train and validate the emulators.
- 7 ER wrote the paper with input from OW.

# 8 Acknowledgements

- 9 This work was supported by the Natural Environment Research Council [grant number
- 10 NE/N003411/1]. We thank Karl Hennermann at the ECMWF for making CAMS interim
- reanalysis data for O<sub>3</sub> and CO available. We also thank Lindsay Lee from Sheffield Hallam
- 12 University for her feedback and comments on an early version of this paper.

# 13 References

14 BARET, F., WEISS, M., ALLARD, D., GARRIGUES, S., LEROY, M., JEANJEAN, H., FERNANDES, R., MYNENI, R., PRIVETTE, J. & MORISETTE, J. 2005. VALERI: a network of sites and a 15 methodology for the validation of medium spatial resolution land satellite products. 16 17 Remote Sensing of Environment, 76, 36-39. 18 Bayarri, M., Berger, J., Cafeo, J., Garcia-Donato, G., Liu, F., Palomo, J., Parthasarathy, R., Paulo, 19 R., Sacks, J., and Walsh, D. (2007). Computer model validation with functional output. Ann. Statist., 35(5):1874–1906 20 21 BERG, B. A. 2005. Introduction to Markov chain Monte Carlo simulations and their statistical analysis. Markov Chain Monte Carlo Lect Notes Ser Inst Math Sci Natl Univ Singap, 7, 1-22 23 52. 24 BEVEN, K. and FREER, J., 2001. Equifinality, data assimilation, and uncertainty estimation in mechanistic modelling of complex environmental systems using the GLUE 25 26 methodology. Journal of hydrology, 249(1-4), pp.11-29.

BOCQUET, M., ELBERN, H., ESKES, H., HIRTL, M., ŽABKAR, R., CARMICHAEL, G., FLEMMING, J., 1 INNESS, A., PAGOWSKI, M. & PÉREZ CAMAÑO, J. 2015. Data assimilation in atmospheric 2 3 chemistry models: current status and future prospects for coupled chemistry meteorology models. Atmospheric Chemistry and Physics, 15, 5325-5358. Brynjarsdóttir, J., & 4 O'Hagan, A. (2014). Learning about physical parameters: The importance of model 5 discrepancy. Inverse problems, 30(11), 114007. 6 7 BOERSMA, K., VINKEN, G. & ESKES, H. 2016. Representativeness errors in comparing chemistry transport and chemistry climate models with satellite UV-Vis tropospheric column 8 9 retrievals. Geoscientific model development, 9, 875. CHANG, K.-L., PETROPAVLOVSKIKH, I., COOPER, O. R., SCHULTZ, M. G. & WANG, T. 2017. 10 Regional trend analysis of surface ozone observations from monitoring networks in 11 12 eastern North America, Europe and East Asia. Elem Sci Anth, 5. 13 CHANG, W., HARAN, M., APPLEGATE, P. & POLLARD, D. 2016. Calibrating an ice sheet model using high-dimensional binary spatial data. Journal of the American Statistical 14 Association, 111, 57-72. CHANG K. L., & GUILLAS, S. (2019). Computer model calibration 15 16 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. 17 18 COHEN, A. J., BRAUER, M., BURNETT, R., ANDERSON, H. R., FROSTAD, J., ESTEP, K., ... & FEIGIN, V. 2017. Estimates and 25-year trends of the global burden of disease attributable to 19 ambient air pollution: an analysis of data from the Global Burden of Diseases Study 20 2015. The Lancet, 389(10082), 1907-1918.COUVREUX, F., HOURDIN, F., WILLIAMSON, 21 22 D., ROEHRIG, R., VOLODINA, V., VILLEFRANQUE, N. et al. (2021). Process-based climate model development harnessing machine learning: I. a calibration tool for 23 24 parameterization improvement. Journal of Advances in Modeling Earth Systems, 13(3), 25 e2020MS002217. CURRIN, C., MITCHELL, T., MORRIS, M. & YLVISAKER, D. 1991. Bayesian prediction of 26 27 deterministic functions, with applications to the design and analysis of computer experiments. Journal of the American Statistical Association, 86, 953-963. 28 DE KAUWE, M. G., DISNEY, M., QUAIFE, T., LEWIS, P. & WILLIAMS, M. 2011. An assessment of 29 30 the MODIS collection 5 leaf area index product for a region of mixed coniferous forest. 31 Remote Sensing of Environment, 115, 767-780. Delang, M. N., et al., 2021, Mapping Yearly Fine Resolution Global Surface Ozone through the 32 33 Bayesian Maximum Entropy Data Fusion of Observations and Model Output for 1990-34 2017, Environ. Sci. Technol. 2021, 55, 8, 4389–4398, doi:10.1021/acs.est.0c07742. 35 EMILI, E., BARRET, B., MASSART, S., LE FLOCHMOEN, E., PIACENTINI, A., EL AMRAOUI, L., PANNEKOUCKE, O. & CARIOLLE, D. 2014. Combined assimilation of IASI and MLS 36 37 observations to constrain tropospheric and stratospheric ozone in a global chemical transport model. Atmospheric Chemistry and Physics, 14, 177-198. 38 FIORE, A. M., DENTENER, F., WILD, O., CUVELIER, C., SCHULTZ, M., HESS, P., TEXTOR, C., 39 SCHULZ, M., DOHERTY, R. & HOROWITZ, L. 2009. Multimodel estimates of 40 intercontinental source-receptor relationships for ozone pollution. Journal of 41 Geophysical Research: Atmospheres, 114. 42 43 FLEMMING, J., BENEDETTI, A., INNESS, A., ENGELEN, R. J., JONES, L., HUIJNEN, V., REMY, S., PARRINGTON, M., SUTTIE, M. & BOZZO, A. 2017. The CAMS interim Reanalysis of Carbon 44

Monoxide, Ozone and Aerosol for 2003-2015. Atmospheric Chemistry and Physics, 17, 1 2 1945. 3 GELMAN, A., CARLIN, J. B., STERN, H. S., DUNSON, D. B., VEHTARI, A. & RUBIN, D. B. 2013. 4 Bayesian data analysis, CRC press. 5 GOLDSMITH, J. R. & LANDAW, S. A. 1968. Carbon monoxide and human health. Science, 162, 6 1352-1359. 7 GRANIER, C., BESSAGNET, B., BOND, T., D'ANGIOLA, A., VAN DER GON, H. D., FROST, G. J., HEIL, 8 A., KAISER, J. W., KINNE, S. & KLIMONT, Z. 2011. Evolution of anthropogenic and 9 biomass burning emissions of air pollutants at global and regional scales during the 1980–2010 period. *Climatic Change*, 109, 163. 10 Higdon, D., Gattiker, J., Williams, B., & Rightley, M. (2008). Computer model calibration using 11 12 high-dimensional output. Journal of the American Statistical Association, 103(482), 570-13 583. 14 HILL, T. C., RYAN, E. & WILLIAMS, M. 2012. The use of CO2 flux time series for parameter and 15 carbon stock estimation in carbon cycle research. Global Change Biology, 18, 179-193. Huijnen, V., Miyazaki, K., Flemming, J., Inness, A., Sekiya, T., and Schultz, M. G.: An 16 intercomparison of tropospheric ozone reanalysis products from CAMS, CAMS interim, 17 TCR-1, and TCR-2, Geosci. Model Dev., 13, 1513–1544, https://doi.org/10.5194/gmd-13-18 19 1513-2020, 2020. 20 JOHNSON, J. S., REGAYRE, L. A., YOSHIOKA, M., PRINGLE, K. J., LEE, L. A., SEXTON, D. M., 21 ROSTRON, J. W., BOOTH, B. B. & CARSLAW, K. S. 2018. The importance of comprehensive parameter sampling and multiple observations for robust constraint of 22 23 aerosol radiative forcing. Atmospheric Chemistry and Physics, 18, 13031-13053. JOHNSON, J.S., CUI, Z., LEE, L.A., GOSLING, J.P., BLYTH, A.M. and CARSLAW, K.S., 2015. 24 25 Evaluating uncertainty in convective cloud microphysics using statistical emulation. Journal of Advances in Modeling Earth Systems, 7(1), pp.162-187. 26 KAMPA, M. & CASTANAS, E. 2008. Human health effects of air pollution. Environmental 27 28 pollution, 151, 362-367. KENNEDY, M. C. & O'HAGAN, A. 2000. Predicting the output from a complex computer code 29 when fast approximations are available. *Biometrika*, 87, 1-13. 30 KENNEDY, M. C. & O'HAGAN, A. 2001. Bayesian calibration of computer models. Journal of the 31 Royal Statistical Society: Series B (Statistical Methodology), 63, 425-464. 32 KHATTATOV, B. V., LAMARQUE, J. F., LYJAK, L. V., MENARD, R., LEVELT, P., TIE, X., BRASSEUR, G. 33 34 P. & GILLE, J. C. 2000. Assimilation of satellite observations of long-lived chemical 35 species in global chemistry transport models. Journal of Geophysical Research: 36 Atmospheres, 105, 29135-29144. LANDRIGAN, P. J., FULLER, R., ACOSTA, N. J., ADEYI, O., ARNOLD, R., BALDÉ, A. B., BERTOLLINI, 37 R., BOSE-O'REILLY, S., BOUFFORD, J. I. & BREYSSE, P. N. 2017. The Lancet Commission on 38 pollution and health. The Lancet. 39 LEE, L., PRINGLE, K., REDDINGTON, C., MANN, G., STIER, P., SPRACKLEN, D., PIERCE, J. & 40 CARSLAW, K. 2013. The magnitude and causes of uncertainty in global model 41 simulations of cloud condensation nuclei. Atmos. Chem. Phys, 13, 8879-8914. 42

1 LEE, L. A., REDDINGTON, C. L. & CARSLAW, K. S. 2016. On the relationship between aerosol 2 model uncertainty and radiative forcing uncertainty. Proceedings of the National 3 Academy of Sciences, 113, 5820-5827. 4 LOEPPKY, J. L., SACKS, J. & WELCH, W. J. 2012. Choosing the sample size of a computer 5 experiment: A practical guide. *Technometrics*. 6 MALLEY, C. S., HENZE, D. K., KUYLENSTIERNA, J. C., VALLACK, H. W., DAVILA, Y., ANENBERG, S. 7 C., TURNER, M. C. & ASHMORE, M. R. 2017. Updated global estimates of respiratory 8 mortality in adults>/= 30Years of age attributable to long-term ozone exposure. 9 Environmental Health Perspectives, 2017, vol. 125, num. 8, p. 087021. MARREL, A., IOOSS, B., JULLIEN, M., LAURENT, B. & VOLKOVA, E. 2011. Global sensitivity 10 analysis for models with spatially dependent outputs. *Environmetrics*, 22, 383-397. 11 MENUT, L., BESSAGNET, B., KHVOROSTYANOV, D., BEEKMANN, M., BLOND, N., COLETTE, A., 12 13 COLL, I., CURCI, G., FORET, G. & HODZIC, A. 2014. CHIMERE 2013: a model for regional 14 atmospheric composition modelling. *Geoscientific model development*, 6, 981-1028. MIYAZAKI, K., ESKES, H., SUDO, K., TAKIGAWA, M., VAN WEELE, M. & BOERSMA, K. 2012. 15 Simultaneous assimilation of satellite NO2, O3, CO, and HNO3 data for the analysis of 16 tropospheric chemical composition and emissions. Atmos. Chem. Phys, 12, 9545-9579. 17 MORRIS, M. D. & MITCHELL, T. J. 1995. Exploratory designs for computational experiments. 18 19 Journal of statistical planning and inference, 43, 381-402. 20 NICELY, J. M., ANDERSON, D. C., CANTY, T. P., SALAWITCH, R. J., WOLFE, G. M., APEL, E. C., ARNOLD, S. R., ATLAS, E. L., BLAKE, N. J. & BRESCH, J. F. 2016. An observationally 21 constrained evaluation of the oxidative capacity in the tropical western Pacific 22 23 troposphere. Journal of Geophysical Research: Atmospheres, 121, 7461-7488. 24 O'HAGAN, A. 2006. Bayesian analysis of computer code outputs: a tutorial. Reliability 25 Engineering & System Safety, 91, 1290-1300. 26 OAKLEY, J. E. & O'HAGAN, A. 2004. Probabilistic sensitivity analysis of complex models: a Bayesian approach. Journal of the Royal Statistical Society: Series B (Statistical 27 28 *Methodology),* 66, 751-769. PARRISH, D., LAMARQUE, J. F., NAIK, V., HOROWITZ, L., SHINDELL, D., STAEHELIN, J., DERWENT, 29 30 R., COOPER, O., TANIMOTO, H. & VOLZ-THOMAS, A. 2014. Long-term changes in lower tropospheric baseline ozone concentrations: Comparing chemistry-climate models and 31 observations at northern midlatitudes. Journal of Geophysical Research: Atmospheres, 32 33 119, 5719-5736. 34 PLUMMER, M. JAGS: A program for analysis of Bayesian graphical models using Gibbs sampling. Proceedings of the 3rd international workshop on distributed statistical computing, 35 36 2003. Technische Universit at Wien, 125. 37 RASMUSSEN, C. E. 2006. Gaussian processes for machine learning. RICHARDSON, A. D., WILLIAMS, M., HOLLINGER, D. Y., MOORE, D. J., DAIL, D. B., DAVIDSON, E. 38 A., SCOTT, N. A., EVANS, R. S., HUGHES, H. & LEE, J. T. 2010. Estimating parameters of a 39 40 forest ecosystem C model with measurements of stocks and fluxes as joint constraints. Oecologia, 164, 25-40. 41 ROUSTANT, O., GINSBOURGER, D. & DEVILLE, Y. 2012. DiceKriging, DiceOptim: Two R packages 42 43 for the analysis of computer experiments by kriging-based metamodeling and 44 optimization.

1	DVAN E MUD O VOUCADAKIS A 8 LEE L 2018 Fast consitivity analysis matheds for
1 2	RYAN, E., WILD, O., VOUGARAKIS, A. & LEE, L. 2018. Fast sensitivity analysis methods for computationally expensive models with multi-dimensional output. <i>Geoscientific model</i>
2	development, 11, 3131-3146.
4	SALTELLI, A., TARANTOLA, S. & CHAN, KS. 1999. A quantitative model-independent method for
4 5	global sensitivity analysis of model output. <i>Technometrics</i> , 41, 39-56.
6	SALTER, J. M., WILLIAMSON, D. B., SCINOCCA, J. & KHARIN, V. 2018. Uncertainty quantification
7	for spatio-temporal computer models with calibration-optimal bases. <i>arXiv preprint</i>
8	arXiv:1801.08184.
9	SCHULTZ, M. G. 2016. Cluster analysis of European surface ozone observations for evaluation of
10	MACC reanalysis data. Atmospheric Chemistry and Physics, 16, 6863.
11	SCHULTZ, M. G., AKIMOTO, H., BOTTENHEIM, J., BUCHMANN, B., GALBALLY, I. E., GILGE, S.,
12	HELMIG, D., KOIDE, H., LEWIS, A. C. & NOVELLI, P. C. 2015. The Global Atmosphere
13	Watch reactive gases measurement network. <i>Elem Sci Anth,</i> 3.
14	SCHULTZ, M. G., SCHRÖDER, S., LYAPINA, O., COOPER, O., GALBALLY, I., PETROPAVLOVSKIKH, I.,
15	VON SCHNEIDEMESSER, E., TANIMOTO, H., ELSHORBANY, Y. & NAJA, M. 2017.
16	Tropospheric Ozone Assessment Report: Database and metrics data of global surface
17	ozone observations. <i>Elem Sci Anth,</i> 5.
18	SCHUTGENS, N. A., GRYSPEERDT, E., WEIGUM, N., TSYRO, S., GOTO, D., SCHULZ, M. & STIER, P.
19	2016. Will a perfect model agree with perfect observations? The impact of spatial
20	sampling. Atmospheric Chemistry and Physics, 16, 6335-6353.
21	SHINDELL, D., FALUVEGI, G., SELTZER, K. and SHINDELL, C., 2018. Quantified, localized health
22	benefits of accelerated carbon dioxide emissions reductions. Nature climate
23	<i>change, 8</i> (4), pp.291-295.
24	SOFEN, E., BOWDALO, D., EVANS, M., APADULA, F., BONASONI, P., CUPEIRO, M., ELLUL, R.,
25	GALBALLY, I., GIRGZDIENE, R. & LUPPO, S. 2016. Gridded global surface ozone metrics
26	for atmospheric chemistry model evaluation. <i>Earth System Science Data</i> , 8, 41.
27	STEVENSON, D., DENTENER, F., SCHULTZ, M., ELLINGSEN, K., VAN NOIJE, T., WILD, O., ZENG, G.,
28	AMANN, M., ATHERTON, C. & BELL, N. 2006. Multimodel ensemble simulations of
29	present-day and near-future tropospheric ozone. <i>Journal of Geophysical Research:</i>
30	Atmospheres, 111.
31 22	STOCKER, T. F., QIN, D., PLATTER, G. K., TIGNOR, M., ALLEN, S. K., BOSCHUNG, J., & MIDGLEY,
32	P. M. 2013. Climate change 2013: The physical science basis. <i>Contribution of working</i>
33 24	group I to the fifth assessment report of the intergovernmental panel on climate
34 35	<i>change, 1535.</i> VAN DINGENEN, R., DENTENER, F. J., RAES, F., KROL, M. C., EMBERSON, L. & COFALA, J. 2009.
36	The global impact of ozone on agricultural crop yields under current and future air
30 37	quality legislation. Atmospheric Environment, 43, 604-618.
38	VAN LOON, M., BUILTJES, P. J. & SEGERS, A. 2000. Data assimilation of ozone in the atmospheric
39	transport chemistry model LOTOS. Environmental Modelling & Software, 15, 603-609.
40	VAN ZELM, R., HUIJBREGTS, M. A., DEN HOLLANDER, H. A., VAN JAARSVELD, H. A., SAUTER, F. J.,
41	STRUIJS, J., VAN WIJNEN, H. J. & VAN DE MEENT, D. 2008. European characterization
42	factors for human health damage of PM10 and ozone in life cycle impact assessment.
43	Atmospheric Environment, 42, 441-453.

1 2	WILD, O. 2007. Modelling the global tropospheric ozone budget: exploring the variability in current models. <i>Atmospheric Chemistry and Physics Discussions</i> , 7 (1), pp.1995- 2035.
3	ffhal-00302576f
4	WILD, O. and PRATHER, M.J., 2006. Global tropospheric ozone modeling: Quantifying errors due
5	to grid resolution. Journal of Geophysical Research: Atmospheres, 111(D11).
6	WILD, O., Voulgarakis, A., O'Connor, F., Lamarque, J. F., Ryan, E., & Lee, L. 2020. Global
7	sensitivity analysis of chemistry-climate model budgets of tropospheric ozone and OH:
8	Exploring model diversity. Atmospheric Chemistry and Physics, 20, 4047-4058.
9	WILKINSON, R. D. 2010. Bayesian calibration of expensive multivariate computer experiments.
10	John Wiley & Sons.
11	WILLIAMS, M., RICHARDSON, A., REICHSTEIN, M., STOY, P., PEYLIN, P., VERBEECK, H.,
12	CARVALHAIS, N., JUNG, M., HOLLINGER, D. & KATTGE, J. 2009. Improving land surface
13	models with FLUXNET data. <i>Biogeosciences</i> , 6, 1341-1359.
14	YOUNG, P. J., NAIK, V., FIORE, A. M., GAUDEL, A., GUO, J., LIN, M., NEU, J., PARRISH, D., RIEDER,
15	H. & SCHNELL, J. 2018. Tropospheric Ozone Assessment Report: Assessment of global-
16	scale model performance for global and regional ozone distributions, variability, and
17	trends. <i>Elem Sci Anth,</i> 6.
18	
19	
20	
21	
22	
23	
24	
25	
26	
27	
28	
29	
30	
31	
32	
33	

# **Figures and Tables for**

# 'Calibrating a global atmospheric chemistry transport model using Gaussian process emulation and ground-level concentrations of ozone and carbon monoxide'



**Figure 1.** Annual mean surface ozone mixing ratio (in ppb) from the FRSGC/UCI CTM showing the regions considered here and the 272 grid cells used for model calibration.



Figure 2. Monthly mean surface O<sub>3</sub> (panel a) and surface CO (panel b) over Europe and North America
 simulated with the FRSGC/UCI CTM compared with ECMWF reanalysis data.



Figure 3. Simulated surface O<sub>3</sub> (panel a) and surface CO (panel b) from the FRSGC/UCI CTM versus
those predicted from the Gaussian process emulators. The simulated and emulated concentrations were
generated using 20 sets of model parameters that were not used for training the emulators.



- 7 Figure 4. Sensitivity indices representing the percentage of the variance in surface O<sub>3</sub> and CO over the
- 8 USA and Europe in the FRSGC/UCI model output due to changes in each parameter.



Figure 5. Sensitivity indices representing the percentage of the variance in surface O<sub>3</sub> in the FRSGC/UCI
model output due to changes in each input parameter. The four parameters displayed here have the
highest sensitivity indices and the largest effect on simulated surface O<sub>3</sub>. Maps of sensitivity indices
corresponding to the other four parameters are shown in Figure S2 of the supplementary material.

6 7



# 8

9 Figure 6. Sensitivity indices representing the percentage of the variance in surface CO in the

10 FRSGC/UCI model output due to changes in each input parameter. Maps of sensitivity indices for the

11 other four parameters are shown in Figure S3 of the supplementary material.



2 Figure 7. Means and 95% credible intervals of 3000 samples of the Dry Deposition scaling parameter

3 from posterior distributions using the MCMC algorithm based on synthetic datasets from scenarios 1-72

4 (table 1). *Control* refers to the FRSGC/UCI model control run surface concentration for each output point.



2 Figure 8. Means and 95% credible intervals of 3000 samples of the Isoprene emission scaling parameter

3 from posterior distributions using the MCMC algorithm based on synthetic datasets from scenarios 1-72

4 (table 1). *Control* refers to the FRSGC/UCI model control run surface concentration for each output point.



2 Figure 9. Means and 95% credible intervals of 3000 samples of the Boundary Layer Mixing scaling

3 parameter from posterior distributions using the MCMC algorithm based on synthetic datasets from

4 scenarios 1-72 (table 1). Control refers to the FRSGC/UCI model control run surface concentration at

5 each output point. The scaling parameter values are given here on a  $log_{10}$  scale.





Figure 10. Means and 95% credible intervals of 3000 samples of the Dry Deposition, Isoprene and
 Boundary Layer Mixing scaling parameters from posterior distributions using the MCMC algorithm

4 based on reanalysis datasets from scenarios 73-90 (table 1). The first and second rows show these

5 parameters estimated using one stream of data ( $O_3$  for the first row and CO for the second row), while the

- 6 third row shows estimates using two data streams ( $O_3$  and CO).
- 7





9

Figure 11. Emulator predictions of surface O<sub>3</sub>, evaluated at values of the scaling parameters sampled
 from the prior distribution (panel a) and posterior distribution (panel b) showing the effects of calibration.
 In panel b, the outputs correspond to the scenario where the calibration involved synthetic O<sub>3</sub> data, a

13 representation error of p = 0.2 and a spatial coverage of 20% (table 2). The predictions shown here are

14 carried out for all model grid boxes, i.e. 100% spatial coverage.

Number	Model process	Control run value	Scaling parameter values
1	Global surface NOx emissions (TgN/year)	40	0.75 - 1.25
2	Global lightning NO emissions (TgN/year)		0.40 - 1.60
3	Global isoprene emissions (TgC/year)	500	0.40 - 1.60
4	Dry deposition rates	model value	0.40 - 1.60
5	Wet deposition rates	model value	0.40 - 1.60
6	Humidity	model value	0.80 - 1.20
7	Cloud optical depth	model value	0.33 - 3.00
8	Boundary Layer mixing	model value	0.10 - 10.0

**1 Table 1.** Model processes and associated scaling parameter ranges used in this study.

2

3 Table 2. Summary of the 90 different MCMC scenarios carried out for this study. The scenarios involved

4 varying: (i) the type of data (synthetic or reanalysis); (ii) the representation error used for the synthetic

5 data (p) where  $m_i(x_{control})$  is the control run output of the CTM and  $\sigma_i$  is the amount of statistical noise

6 added; (iii) the percentage coverage of grid-squares in the USA and Europe. For the synthetic data the 24

- 7 scenarios correspond to a full factorial combination of four levels of representation error and six levels of
- 8 spatial coverage, while for the reanalysis data the six scenarios correspond to the six levels of spatial
- 9 coverage.

Scenarios	Dataset	Representation error, p	Spatial coverage
		$\left(\sigma_i = p \times m_i(x_{control})\right)$	
1-24	Synthetic O <sub>3</sub>	0.01, 0.1, 0.2, 0.3	2.5%, 5%, 10%, 20%, 40%, 100%
25-48	Synthetic CO	0.01, 0.1, 0.2, 0.3	2.5%, 5%, 10%, 20%, 40%, 100%
49-72	Synthetic O <sub>3</sub> & CO	0.01, 0.1, 0.2, 0.3	2.5%, 5%, 10%, 20%, 40%, 100%
73-78	Reanalysis data (O <sub>3</sub> )	Parameter to be estimated	2.5%, 5%, 10%, 20%, 40%, 100%
79-84	Reanalysis data (CO)	Parameter to be estimated	2.5%, 5%, 10%, 20%, 40%, 100%
85-90	Reanalysis data (O <sub>3</sub> & CO)	Parameter to be estimated	2.5%, 5%, 10%, 20%, 40%, 100%