Ensemble-Based Experimental Design for Targeting Data Acquisition to Inform Climate Models

Oliver R. A. Dunbar1, Michael F. Howland1,2, Tapio Schneider1,2, and Andrew M. Stuart1

1California Institute of Technology, Pasadena, CA, USA; 2Civil and Environmental Engineering, Massachusetts Institute of Technology, Cambridge, MA, USA

Abstract Data required to calibrate uncertain general circulation model (GCM) parameterizations are often only available in limited regions or time periods, for example, observational data from field campaigns, or data generated in local high-resolution simulations. This raises the question of where and when to acquire additional data to be maximally informative about parameterizations in a GCM. Here we construct a new ensemble-based parallel algorithm to automatically target data acquisition to regions and times that maximize the uncertainty reduction, or information gain, about GCM parameters. The algorithm uses a Bayesian framework that exploits a quantified distribution of GCM parameters as a measure of uncertainty. This distribution is informed by time-averaged climate statistics restricted to local regions and times. The algorithm is embedded in the recently developed calibrate-emulate-sample framework, which performs efficient model calibration and uncertainty quantification with only $O(10^5)$ model evaluations, compared with $O(10^5)$ evaluations typically needed for traditional approaches to Bayesian calibration. We demonstrate the algorithm with an idealized GCM, with which we generate surrogates of local data. In this perfect-model setting, we calibrate parameters and quantify uncertainties in a quasi-equilibrium convection scheme in the GCM. We consider targeted data that are (a) localized in space for statistically stationary simulations, and (b) localized in space and time for seasonally varying simulations. In these proof-of-concept applications, the calculated information gain reflects the reduction in parametric uncertainty obtained from Bayesian inference when harnessing a targeted sample of data. The largest information gain typically, but not always, results from regions near the intertropical convergence zone.

Plain Language Summary Climate models depend on dynamics across many spatial and temporal scales. It is infeasible to resolve all of these scales. Instead, the physics at the smallest scales is represented by parameterization schemes that link what is unresolvable to variables resolved on the grid scale. A dominant source of uncertainty in climate predictions comes from uncertainty in calibrating empirical parameters in such parameterization schemes, and these uncertainties are generally not quantified. The uncertainties can be reduced and quantified with data that may have limited availability in space and time, for example, data from field campaigns or from targeted high-resolution simulations in limited areas. But the sensitivity of simulated climate statistics, such as precipitation rates, to parameterizations varies in space and time, raising the question of where and when to acquire additional data so as to optimize the information gain from the data. Here we construct an automated algorithm that finds optimal regions and time periods for such data acquisition, to maximize the information the data provides about uncertain parameters. In proof-of-concept simulations with an idealized global atmosphere model, we show that our algorithm successfully identifies the informative regions and times, even in cases where physics-based intuition may lead to sub-optimal choices.

1. Introduction

Parameterizations of subgrid-scale processes, such as the turbulence and convection controlling clouds, are the principal cause of physical uncertainties in climate predictions (Bony & Dufresne, 2005; Bony et al., 2006; Brient & Schneider, 2016; Cess et al., 1989, 1990; Stephens, 2005; Vial et al., 2013; Webb et al., 2013). Similar parameter uncertainties in principle can be quantified and reduced by calibration with data. High-resolution simulations such as large-eddy simulations (LESs) are able to resolve turbulence and convection in atmosphere and oceans over limited areas (Khairoutdinov et al., 2009; Matheou & Chung, 2014; Pressel et al., 2015, 2017; Schalkwijk et al., 2015; Siebesma et al., 2003; Stevens et al., 2005) and have been used to calibrate climate model parameterizations at selected sites (e.g., Couvreux et al., 2021; de Rooy et al., 2013; GEWEX Cloud System Science Team, 1993; Hohenegger & Bretherton, 2011; Hourdin et al., 2021; Li & Fox-Kemper, 2017; Liu et al., 2001;
of the model in which the parameters appear (see Geyer (Dunbar et al., 1995) to choose data acquisition sites and time periods that are maximally informative about parameters in a model. The experimental-design problem we address has similarities with the problem of how to choose sites for targeted weather observations to optimally improve weather forecasts (Bishop & Toth, 1999; Emanuel et al., 1995; Lorenz & Emanuel, 1998). However, in contrast to the situation in weather forecasting, which focuses on trajectory matching for state estimation, our focus is on minimizing mismatches in time-averaged climate statistics for the estimation of parameters in climate models.

To learn from time-averaged statistics, we adopt a Bayesian inverse problem setting (see, e.g., Kaipio and Somersalo (2006), Tarantola (2005), Stuart (2010), and Drovandi et al. (2013) for reviews). In this setting, parameters (or parametric or nonparametric functions) in parameterizations are treated as having probability distributions. Data (e.g., climate statistics) are used to reduce the uncertainty reflected by these distributions, balancing contributions of the data with that of prior knowledge about parameters (e.g., physical constraints). This results in the joint posterior distribution for parameters, including the correlation structure of uncertainties among parameters. The Bayesian experimental design tools we apply in this paper leverage the posterior distribution to determine regions and times where local data are maximally effective at reducing parameter uncertainties. As is typical in such analyses, we measure the quality of a design (site location or time period) by a scalar utility function. We choose a utility that quantifies the information entropy loss between posterior and prior for each parameter.

A natural question arises: how might we most effectively place such high-resolution simulations? In this paper we address the general task of optimal targeting of data acquisition, and we demonstrate our approach within an idealized GCM setting. We create an automated algorithm based on experimental design criteria (Chaloner & Verdinelli, 1995) to choose data acquisition sites and time periods that are maximally informative about parameters in a model. The experimental-design problem we address has similarities with the problem of how to choose sites for targeted weather observations to optimally improve weather forecasts (Bishop & Toth, 1999; Emanuel et al., 1995; Lorenz & Emanuel, 1998). However, in contrast to the situation in weather forecasting, which focuses on trajectory matching for state estimation, our focus is on minimizing mismatches in time-averaged climate statistics for the estimation of parameters in climate models.

To learn from time-averaged statistics, we adopt a Bayesian inverse problem setting (see, e.g., Kaipio and Somersalo (2006), Tarantola (2005), Stuart (2010), and Drovandi et al. (2013) for reviews). In this setting, parameters (or parametric or nonparametric functions) in parameterizations are treated as having probability distributions. Data (e.g., climate statistics) are used to reduce the uncertainty reflected by these distributions, balancing contributions of the data with that of prior knowledge about parameters (e.g., physical constraints). This results in the joint posterior distribution for parameters, including the correlation structure of uncertainties among parameters. The Bayesian experimental design tools we apply in this paper leverage the posterior distribution to determine regions and times where local data are maximally effective at reducing parameter uncertainties. As is typical in such analyses, we measure the quality of a design (site location or time period) by a scalar utility function. We choose a utility that quantifies the information entropy loss between posterior and prior for each parameter.

Construction of the joint posterior distribution of the parameters is well known to be a computationally intensive task, with commonly used Markov chain Monte Carlo (MCMC) methods typically requiring $O(10^5)$ evaluations of the model in which the parameters appear (see Geyer (2011) for an overview). The recent development of the calibrate-emulate-sample (CES) framework accelerates Bayesian learning by a factor of $O(1)$ (Cleary et al., 2021; Dunbar et al., 2021). The calibration stage uses a variant of ensemble Kalman inversion (Chen & Oliver, 2012; Emerick & Reynolds, 2013; Iglesias et al., 2013; Reich, 2011) to obtain a collection of samples of the model about an optimal set of parameters. The emulation stage features the training of a machine learning emulator, here, a Gaussian process (Kennedy & O’Hagan, 2000, 2001; C. K. Williams & Rasmussen, 2006), to emulate output statistics of the model using the pairs of parameters and model outputs from the calibration stage. The sample stage then samples a posterior distribution with MCMC methods, replacing the computationally expensive model with the cheap emulator. This framework can extend to the learning of data-driven parameterizations or other non-parametric functions, such as structural model errors (e.g., M. E. Levine & Stuart, 2021; Lopez-Gomez et al., 2022; Schneider et al., 2022). Our proposed algorithm builds on CES to incorporate Bayesian experimental design at negligible additional computational expense. In particular, we do not require additional forward model (GCM) evaluations over what is already required in CES to perform uncertainty quantification.

We demonstrate our approach with an idealized moist GCM (Frierson et al., 2006) with modifications introduced by O’Gorman and Schneider (2008b), with which we generate surrogates of local data and in which we calibrate parameters in a quasi-equilibrium convection scheme (Frierson, 2007). We conduct numerical experiments with the idealized GCM in statistically stationary and seasonally varying configurations and show how to determine the utility of data at different sites and in different seasons. These experiments serve as proof-of-concept of the broad-purpose algorithm, which can be applied, for example, to determine optimal sites and times for high-resolution simulations for the calibration and uncertainty quantification of parameterizations.

In Section 2, we define the inverse problems for parameter calibration and the optimal design algorithm; details of efficient uncertainty quantification (CES) are provided in Appendix A. In Section 3, we briefly describe the GCM used for demonstrating the algorithm. Results of the optimal design algorithm are described in Section 4. We end with a summary and discussion of conclusions in Section 5.
2. Methodology

Our goal is to target data acquisition to regions and times at which uncertainty reduction (information gain) is maximized. We do this in two stages. First, we learn the temporally and spatially varying sensitivities of the model statistics with respect to model parameters. Second, we use this knowledge to target data acquisition to regions and times at which the model is maximally sensitive to new data. We work in a framework similar to Dunbar et al. (2021) which focuses on accelerated uncertainty quantification within a GCM.

2.1. Inverse Problem to Learn From Limited-Area Data

We study calibration of parameters in a GCM by formulating parameter learning as a Bayesian inverse problem. Define $G_T(\theta; \nu(0))$ to be the forward map sending the parameters $\theta$ to time-aggregated simulated climate statistics (averaged over a window of length $T > 0$) from an initial state $\nu(0)$. We assume that the aggregation $G_T(\theta; \cdot)$ is statistically stationary, and we refer to samples of such aggregated climate statistics as data throughout, irrespective of whether they are observational or computationally generated. We consider a situation in which data are only locally available, at a particular spatial or spatio-temporal location, indexed by $k$, which we refer to as the design point. We make use of a restriction operation $W_k$ to a point $k$, and define the limited-area forward map, $S_T(\theta; k, \nu(0)) = W_k G_T(\theta; \nu(0))$.

For any given $k$, assume we have data $z_k$ available. For example, $z_k$ could be produced with a simulation with limited spatial or temporal extent, or by running a field campaign. We form an inverse problem for GCM learning from this data as

$$z_k = S_T(\theta; k, \nu(0)) + \delta_k,$$  \hspace{1cm} (1)

where $\delta_k$ is a stochastic term to capture discrepancies between model $S_T(\cdot; k, \cdot)$ and data $z_k$, (e.g., Kennedy & O’Hagan, 2001). The initial condition $\nu(0)$ appears in this formulation but is treated as a nuisance variable. This view is justified in the context of learning about atmospheric parameterizations for climate models, where lower-frequency information (e.g., seasonal variations) is particularly informative (Schneider et al., 2017). Indeed, the time-averaged data filters out the high-frequency information. Following Dunbar et al. (2021), we write $S_T(\theta; k, \nu(0)) \approx S_m(\theta; k) + \sigma_k$, where $\sigma_k \sim N(0, \Sigma(\theta))$ is normal noise, independent from $\delta_k$, with mean zero and with a covariance matrix $\Sigma(\theta)$ reflecting chaotic internal variability. The Gaussian assumption is justified on the basis of a central limit theorem (CLT) applied to the time averages. The inverse problem 1 is thus approximated by the problem

$$z_k = S_m(\theta; k) + \delta_k + \sigma_k, \quad \sigma \sim N (0, W_k \Sigma(\theta) W_k^T).$$  \hspace{1cm} (2)

This is a desirable inverse problem without dependence on the initial condition. It is an approximation due to the CLT, but this approximation should be suitable if $T$ is taken larger than the dynamical system’s Lyapunov timescale (for the atmosphere, this equates to $T \geq 15$ days (F. Zhang et al., 2019)). In our experiments, we take $T = 90$ days (Section 3.2), or $T = 30$ days (Appendix C).

Solving Equation 2 involves finding the posterior distribution of $\theta$ given the data $z_k$, denoted $\theta|z_k$. Although we cannot evaluate $S_m$ directly, the emulate phase of the CES algorithm (Cleary et al., 2021) constructs a surrogate of $S_m$ from carefully chosen evaluations of $S_T$. Details of the algorithm are provided in Appendix A.

2.2. Experimental Design

We consider a situation in which acquiring $z_k$ is associated with large costs. For example, $z_k$ could be data obtained by running a computationally demanding simulation, or running an expensive field campaign. Our starting point is to assume that a limited budget restricts us to evaluate $z_k$ at a single design point $k$ at a time. We want to choose the design point $k$ that leads to the most informative inverse problem 2. We continue using a Bayesian point of view, namely, the optimal $k$ is the one for which the posterior distribution of $\theta|z_k$ learned from the inverse problem 2 has the smallest uncertainty. This perspective is motivated by the downstream goal of minimizing the parametric uncertainty of GCM predictions.
To answer this conclusively, one would need to evaluate \( z_k \) at all design points \( k \), which here is too computationally expensive. Instead, we investigate only the sensitivity of the forward model statistics \( G_r \) to its parameters \( \theta \) to assess the additional information provided at each design point \( k \). This additional information at \( k \) is used as a proxy for the information content that would exist when learning from data \( z_k \). The benefits of this approach are that (a) we do not require any evaluations of \( z_k \) to select the optimal location; (b) the measure of information content is naturally constructed from the uncertainty reflected by the Bayesian posterior distribution; and (c) we can perform this efficiently, and in an embarrassingly parallel fashion, requiring only \( O(100) \) GCM runs, determined by the product of the ensemble size and the number of iterations typically needed in the calibration stage of the CES algorithm (see Appendix A). The approach necessarily will contain a bias from the prior distribution of the parameters.

Each evaluation of the forward map involves a simulation with the GCM and thus depends on an initial condition \( v^{(0)} \) and parameters \( \theta \). Together this gives rise to the definition of time-aggregated model statistics \( y \),

\[
y = G_r(\theta; v^{(0)}).
\]

For sufficiently large \( I \), we use the CLT as in Section 2.1 to approximate this relationship as

\[
y = \mu_w(\theta) + \sigma, \quad \sigma \sim N(0, \Sigma(\theta)),
\]

where \( \Sigma(\theta) \) is the internal variability covariance matrix for parameters \( \theta \). To proceed, we must choose a control value \( \theta^* \) for example, we take the mean of the prior distribution. Fixing \( \theta = \theta^* \), we generate a realization of \( y \).

We now solve a set of inverse problems, with the solution of each providing additional information at a design. Specifically, given \( y \), we temporarily “forget” \( \theta^* \), and for any design point \( k \), we consider

\[
W_k y = W_k \mu_w(\theta) + \sigma_k, \quad \sigma_k \sim N(0, W_k \Sigma(\theta) W_k^T),
\]

where \( W_k \) restricts the data space to \( k \). The posterior distributions of \( \theta | W_k y \) for all \( k \) obtained by solving Equation 5 informs us about the sensitivities of \( \mu_w \) with respect to parameters, when only data at different \( k \) is available. To simplify the solution of the inverse problem, we approximate the internal variability covariance matrix \( \Sigma(\theta) \) by a fixed covariance matrix \( \Sigma(\theta^*) \). This covariance matrix can be obtained by running a collection of control simulations with parameters fixed to (the known) \( \theta^* \) but with different initial conditions.

The utility \( U \) of a design \( W_k \) is a scalar function reflecting the quality of a given design. The design that maximizes the utility function is known as the optimal design. We choose a utility function by measuring information gain (or uncertainty reduction) in \( \theta | W_k y \) relative to the prior, in a form of Bayesian optimal design. We use a common choice of utility function that arises in both the Bayesian and non-Bayesian design literature (e.g., Chaloner & Verdinelli, 1995; Fedorov & Hackl, 1997; Ryan et al., 2016; Schneider & Griffies, 1999), namely, the inverse of the determinant of the information matrix (i.e., the inverse of the posterior covariance matrix),

\[
U(W_k) = (\det (\text{Cov} (\theta | W_k y)))^{-1}.
\]

In practice, the posterior covariance matrix is estimated as the empirical covariance matrix of samples drawn from \( \theta | W_k y \). This utility fulfills the so-called D-optimality criterion; unlike trace-based measures (e.g., A-optimal utility functions), it is invariant under arbitrary linear transformations of the parameters, for example, when parameters have different dimensional scales. It has been used in investigations of linear and nonlinear design (Alexanderian et al., 2016; Alexanderian & Saibaba, 2018; Drovandi et al., 2013; Ryan et al., 2014) and particularly in the context of sensor placement (Uciński, 2000; Uciński & Patan, 2007). For linear forward maps and Gaussian priors, maximization of this utility is equivalent to maximization of the expected Kullback-Leibler divergence (KLD), a relative entropy measure (Cook et al., 2008; Huan & Marzouk, 2013; Kim et al., 2014). While KLD has beneficial mathematical properties, especially for highly non-Gaussian posteriors (Paniński, 2005), it is difficult to evaluate, especially in high-dimensional problems (e.g., Huan & Marzouk, 2013).
2.3. Synthesis: Targeted Uncertainty Quantification Algorithm

The combined algorithm for targeted uncertainty quantification consists of two stages: first, finding an optimal design point \( k \) in a design stage and, second, evaluating parameter uncertainty with data from \( k \) in an uncertainty quantification stage. Let \( D \) be the finite index set for the set of design points, and define \( W_k \) to be the restriction map for any \( k \in D \). The two stages then are as follows:

1. The design stage consists of the following steps:
   
   (a) Generate a sample of GCM simulated data \( y = G_{\theta}(\theta^*; \nu^{(0)}) \), and estimate the internal variability covariance matrix \( \Sigma(\theta^*) \). We approximate \( \Sigma(\theta) \) as \( \Sigma(\theta^*) \).
   
   (b) For each \( k \in D \), solve Equation 5, in parallel, for the posterior of \( \theta|W_k y \), using the CES-type algorithm described in Appendix A.
   
   (c) For each \( k \in D \), calculate the utility \( U(W_k) \) from Equation 6 and choose the optimal design

   \[
   \hat{k} = \arg\max_{k \in D} U(W_k).
   \]

2. The uncertainty quantification stage consists of the following steps:

   (a) At the optimal design point \( \hat{k} \), obtain a sample \( z_{\hat{k}} \).
   
   (b) Solve the inverse problem Equation 2 for the posterior distribution of \( \theta|z_{\hat{k}} \).

This algorithm could be used as one iteration of a workflow loop where, for example, the posterior distribution \( \theta|z_{\hat{k}} \) can be used to inform a new choice of \( \theta^* \).

The complexity of the first stage grows linearly with the candidate design points \( k \) because we only consider one point at a time. However, if one wishes to choose a design composed of \( K \) simultaneous points from a set \( D \), a combinatorial problem arises, with complexity growing like \( |D|^K / (|D|! - |K|!) |K|! \) — a common problem in the related field of sensor placement design (Uciński & Patan, 2007; van de Wal & de Jager, 2001). This will become prohibitively costly to solve by brute force, even in parallel. We focus on the algorithm for single design points \( k \) for now, addressing scaling questions in the discussion section.

3. Idealized GCM and Experimental Setup

3.1. Idealized GCM, Parameters, and Priors

To demonstrate the algorithm in a simplified setting, we use the idealized aquaplanet GCM described by Frierson et al. (2006) and Frierson (2007) with the modifications introduced by O’Gorman and Schneider (2008b). The idealized GCM uses the spectral transform dynamical core of the Flexible Modeling System, developed at the Geophysical Fluid Dynamics Laboratory. We use a coarse spectral resolution of T21 (32 latitude points and 64 longitude points on the Gaussian transform grid). The vertical is discretized with finite differences with 20 equally spaced sigma levels (Simmons & Burridge, 1981). The time discretization uses a second-order leapfrog method with a Robert-Asselin-Williams filter (P. D. Williams, 2011). The GCM’s atmosphere is coupled to a 1-m thick slab ocean, and it uses a two-stream gray radiation scheme. Convection is represented by a simple quasi-equilibrium moist convection scheme, which relaxes temperature and specific humidity toward moist-adiabatic reference profiles with a fixed relative humidity RH (Frierson, 2007). The timescale with which the temperature and specific humidity relax to their respective reference profiles is given by the parameter \( r \). The parameters RH and \( r \) are the focus of this study.

Since the GCM has no topography or other asymmetries at the surface, its statistics are zonally symmetric. With fixed insolation at the top of the atmosphere, the statistics are also statistically stationary. Prescribing seasonally (but not diurnally) varying insolation generates seasonally varying (cyclostationary) statistics, with symmetry between the northern and southern hemisphere (i.e., winter in the northern hemisphere winter is statistically identical to winter in the southern hemisphere) (Bordoni & Schneider, 2008; Howland et al., 2022). Dunbar et al. (2021) and Howland et al. (2022) have shown that the parameters RH and \( r \) of the convection parameterization in the GCM can be calibrated in the stationary and cyclostationary regimes. Here we want to determine optimal designs for learning about these parameters in the two regimes.
The priors for these parameters are taken to be logit-normal and lognormal distributions, \( RH \sim \text{Logitnormal}(0, 1) \) and \( \tau \sim \text{Lognormal}(12 \text{h}, (12 \text{h})^2) \). That is, we define the invertible transformation
\[
\mathcal{T}(RH, \tau) = \left( \text{logit}(RH), \ln\left( \frac{\tau}{1 \text{ s}} \right) \right),
\]
which transforms each parameter to values along the real axis. We label the transformed (or computational) parameters as \( \theta = \mathcal{T}(RH, \tau) \). The untransformed (or physical) parameters (relative humidity and timescale) are uniquely defined by \( \mathcal{T}^{-1}(\theta) \). We apply our calibration methods in the space of the transformed parameters \( \theta \), where priors are unit-free, normally distributed, and unbounded; meanwhile, the idealized GCM uses the physical parameters \( \mathcal{T}^{-1}(\theta) \), with \( RH \in [0, 1] \) and \( \tau \in [0, \infty) \). In this way, the prior distributions enforce physical constraints on the parameters.

3.2. Objective Function for Parameter Learning

We learn from statistics of model output that are known to be sensitive to the convection parameters. We have knowledge about these sensitivities from a body of previous studies that used this idealized GCM (e.g., Bischof & Schneider, 2014; Bordoni & Schneider, 2008; Kaspi & Schneider, 2011, 2013; X. Levine & Schneider, 2015; Merlis & Schneider, 2011; O’Gorman, 2011; O’Gorman & Schneider, 2008a, 2008b, 2009b; Schneider et al., 2010; Wei & Bordoni, 2018; Wills et al., 2017). We know, for example, that the convection scheme primarily affects the atmospheric thermal stratification in the tropics, with weaker effects in the extratropics (Schneider & O’Gorman, 2008). We also know that the relative humidity parameter \( RH \) in the convection scheme controls the humidity of the tropical free troposphere but has a weaker effect on the humidity of the extratropical free troposphere (O’Gorman et al., 2011). Thus, we expect tropical circulation statistics to be especially informative about the parameters in the convection scheme. However, convection plays a central role in intense precipitation events at all latitudes (O’Gorman & Schneider, 2009a, 2009b), so we expect statistics of precipitation intensity to be informative about convective parameters, and in particular to contain information about the relaxation timescale \( \tau \).

As statistics to learn from, we choose averages of the free-tropospheric relative humidity, of the precipitation rate, and of a measure of the frequency of intense precipitation. We use averages over \( T = 90 \) days in both the statistically stationary and seasonal cycle simulations. We exploit the statistical zonal symmetry in the GCM by taking zonal averages in addition to the time averages. The relative humidity data are evaluated at \( \sigma = 0.5 \) (where \( \sigma = p/p_s \) is pressure \( p \) normalized by the local surface pressure \( p_s \)), the precipitation rate is taken daily, and as a measure of the frequency of intense precipitation, we use the frequency with which daily precipitation exceeds the latitude-dependent 90th percentile of precipitation rates in a long (18,000 days) control simulation. We hence have 3 statistics, each a function of the 32 latitude points on the spectral transform grid, resulting in a 96-dimensional output vector \( H_T \). In the statistically stationary case, we take the forward map \( G_T = H_T \).

For the simulations with a seasonal cycle, \( H_T \) is not statistically stationary but is cyclostationary over multiples of a year. The year length in the GCM is 360 days. We stack four 90-day seasons of data together (Howland et al., 2022) and define the forward map
\[
G_T(\theta; v^{(i)}) = \left[ H_T(\theta; v^{(0)}), \ldots, H_T(\theta; v^{(3)}) \right]
\]
over a one-year cycle (360 days), where \( v^{(i)} \) is the model state at the beginning of each 90-day long season labeled \( i = 0, 1, 2, 3 \). With this batching, we have now constructed stationary statistics for the stacked data. The theory of Section 2 applies, and our inverse problems can be formulated in the seasonally varying case.

3.3. Design Choices

In the stationary GCM setting, we aggregate statistics temporally and zonally. Thus, a local design implies a restriction to certain latitudes. Recall our discretization has 32 discrete latitudes. We therefore choose a design space that contains sets of \( \ell \) consecutive discrete latitudes, indexed from south to north poles. In the stationary experiments, we focus on the case \( \ell = 1 \).
In the seasonally varying setting, we still aggregate temporally and zonally, but we also stack the seasons in a vector. We define a local design by indexing both a restriction to a season and a restriction to certain latitudes. We choose a design space that contains sets of \( \ell \) consecutive discrete latitudes, collected season by season, indexed from south to north poles. In the seasonal experiments, we focus on the case \( \ell = 1 \).

For additional design scenarios in the stationary setting, we consider cases with wider design stencils, \( \ell = 3 \), in Appendix B, and we consider cases with shorter averaging periods, \( T = 30 \) days, in Appendix C.

### 3.4. Synthetic Data and Noise

We generate limited-area data \( z_k \) with the idealized GCM itself at a fixed parameter vector \( \theta^* \), adding Gaussian noise \( \delta_k \) with zero mean and covariance matrix \( \Delta \) as in Equation 2. One interpretation of this added noise is that it plays the role of an artificial corruption of \( S_{\gamma}(\theta^*; k) \), with unbiased model error \( \delta_k \) that plays the same role as additional observational noise (Kennedy & O’Hagan, 2001). One can obtain unbiased \( \delta_k \) by inclusion of models for structural model error within \( S_{\gamma} \), for example, learned error models that enforce conservation laws and sparsity (M. E. Levine & Stuart, 2021; Schneider et al., 2022). The inverse problem 2 can be written as

\[
z_k = S_{\gamma}(\theta; k) + \gamma_k, \quad \gamma_k \sim N(0, W_k(\Sigma(\theta) + \Delta)W_k^T).
\]  

We construct the measurement error covariance matrix \( \Delta \) to be diagonal with entries \( d_k^2 = \Delta_{ii} > 0 \), where \( i \) indexes over data type (three observed quantities) and over the discrete latitudes,

\[
\Sigma + \text{diag}(d_k^2) = \Sigma + \Delta.
\]

We choose \( d_i \) so that it is proportional to the mean \( \mu_i \) of the variable in question, with a proportionality factor \( C_{\text{max}} = 0.1 \). To prevent the noise from becoming so large that the variables can cross a physical boundary \( \partial \Omega_i \) (e.g., relative humidity becoming negative), we limit the noise standard deviation to a factor \( C = 0.2 \) times the distance between the approximate 95% noise confidence interval and the physical boundary:

\[
d_i = \min \left( C_{\text{min}} \left( \text{dist} \left( \mu_i + 2 \sqrt{\Sigma_{ii}}, \partial \Omega_i \right), \text{dist} \left( \mu_i - 2 \sqrt{\Sigma_{ii}}, \partial \Omega_i \right) \right), C_{\text{max}} \mu_i \right).
\]

In our proof-of-concept experiments, we generate a sample of ground truth data, \( z_k \), and its variability, by carrying out a set of control simulations, with the parameters fixed to values \( \theta^* \), where \( T^{-1}(\theta^*) = (0.7, 2 \text{ h}) \) are standard values used in previous studies (O’Gorman & Schneider, 2008b). We use this set of control simulations to estimate the restricted covariance matrix \( W_k^T \Sigma(\theta)W_k^T \approx W_k^T \Sigma(\theta^*)W_k^T \) for any \( k \). In the statistically stationary case, we carry out control simulations for 200 windows of length \( T = 90 \) days, after discarding the first 50 months for spin-up, and calculate the sample covariance matrix \( \Sigma(\theta^*) \) from the 200 samples. Here, \( W_k \Sigma(\theta^*)W_k^T \) is a symmetric matrix whose size depends on the design space; it represents noise from internal variability in the 90-day time averages. In the seasonally varying case, we carry out a control simulation for 150 years, discarding the first 4 years for spin-up, and obtain the sample covariance matrix \( \Sigma(\theta^*) \) from the stacked seasonal (\( T = 90 \) days) averages. In the seasonal case, \( W_k \Sigma(\theta^*)W_k^T \) is a symmetric matrix whose size depends on 4 times the design space. We add a small regularization term of \( 10^{-4} \) to the diagonal of \( \Sigma(\theta^*) \) to prevent zero variability, which occurs due to finite-time averages of intense precipitation. In practical implementations of this method, good estimates of the local variability that we represent by \( W_k^T \Sigma(\theta^*)W_k^T \) can be obtained from the observed climatology of the statistics of interest, instead of estimating them from a control simulation of the GCM.

In the data acquisition algorithm, we require a sample of data \( W_k^T \gamma_k \) and its variability, for different \( k \). To obtain this, we use a set of control simulations of the GCM in which we fix the parameters to the prior mean \( \theta^* \), the value used to generate \( y_k \), equivalent to the physical values \( T^{-1}(\theta^*) = (0.5, 7 \text{ h}) \). In the stationary case, the three latitude-dependent fields evaluated at 32 latitude points produce a \( 96 \times 96 \) symmetric matrix \( \Sigma(\theta^*) \), representing noise from internal variability in 90-day averages. Similarly, in the seasonal case, the stacked statistics produce a \( 384 \times 384 \) symmetric matrix \( \Sigma(\theta^*) \), and since \( T = 90 \) days, \( \Sigma(\theta^*) \) represents noise from internal variability in 90-day averages. We again add a small regularization term of \( 10^{-4} \) to the diagonal of \( \Sigma(\theta^*) \). In both cases, we estimate \( \Sigma(\theta) \approx \Sigma(\theta^*) \) in the optimal design stage of the algorithm.
4. Results

4.1. Stationary Statistics

We first apply the optimal design algorithm to the statistically stationary GCM. The logarithm of the utility function is shown in Figure 5. The extent to which hemispheric symmetry of the statistics is broken in Figure 5 is an indication of sampling variability, as the infinite-time GCM statistics are hemispherically symmetric. The design landscape appears surprising, as precipitation and parameterized tendencies from convection are largest in the ITCZ (within ±3° of the equator), and one may expect the optimal region to be in the ITCZ as well. Our algorithm indicates that the equatorial region is indeed a good location, but larger utilities are found at latitude ±19°, near the precipitation minima under the descending branches of the Hadley circulation in this model. Indeed, daily precipitation rates at this subtropical latitude correlate more strongly with the relative humidity parameter in the convection scheme than in the equatorial latitudes (Figure 6). With designs focused on a single latitude (ℓ = 1), this region is indicated to be most informative. With wider design stencils (ℓ = 3), the algorithm's aligns closer with intuition, placing optimal utility near the equator (Figure B1).
We validate our optimal choice by solving Equation 7 at four representative design choices, at latitudes $-19^\circ$, $-3^\circ$, $-8^\circ$, and $-75^\circ$, (in decreasing order of utility) shown as colored discs in Figure 5. The samples of climate statistics used at each latitude are shown in Figure 3 (colored discs). Density plots of the posterior distributions at each latitude are shown in Figure 7. Each panel shows the density contours bounding 50%, 75%, and 99% of the posterior distribution, shaded dark to light; the priors are largely uninformative and have been excluded from the plots. The panels a–d are ordered by decreasing utility from Figure 5, which is a predictor of information content based on uncertainty at the prior mean $\hat{\theta}$. The true utilities of the posterior distributions $\hat{\theta} | z_k$ are 26.4, 13.9, 4.4, and 1.7. Thus, the order of predicted information content reflects the order of actual information content. Visually, we see an increased area covered by the different contours for less informative distributions. However, the prediction of the ordering of utilities does not extend to providing accurate prediction of the actual utility value, due to the additional error inflation present in the true data and sampling error. Physical intuition, positing the equatorial region as the optimal target location, would lead to a reasonable design with a utility of 13.9 (Figure 7b), around half that of the optimal design (26.4). A poor guess, positing high latitudes as the optimal target location, would lead to only moderate improvements relative to the prior (Figure 7d), with a utility of 1.7 that is around a factor 20 smaller than that for the optimal design. With wider design stencils, the optimally informative location is predicted to be closer to the equator (Figure B1) As observed in other investigations (Dunbar et al., 2021), the posterior distributions are subject to variability due to the finite-time sampling and the inflation. However, all distributions capture the true parameter values within 50% of the posterior mass.

Figure 3. Aggregated climate statistics in the statistically stationary control simulation using the ground truth parameters $\theta^\ast$. The mean (gray lines) and 95% confidence intervals (shading) of the data are plotted against latitude. Noise mimicking observational and/or model error is added. Each colored disc represents a 90-day realization of GCM data coming from a different design (latitude) used in the experiment.

Figure 4. Aggregated climate statistics in the seasonally varying control simulation using the ground truth parameters $\theta^\ast$. We added noise mimicking observational and/or model error. The mean (solid lines) and 95% confidence intervals (shading) of the data are plotted against latitude, with the colors indicating different seasons, referenced to the northern hemisphere. The blue vertical line indicates the location and season (northern winter) in which we observe the data for uncertainty quantification; the specific 90-day realization of GCM data for the one-latitude design is given by the blue disc.
When the climate statistics that are used are based on shorter-term averages and hence are noisy, the targeting algorithm, as expected, can become less effective, and parameter posteriors can become more multimodal (Figure C5).

4.2. Seasonally Varying Statistics

In the seasonally varying case, we choose the optimal design with the algorithm in Section 2.3 applied to the data stacked in seasons. Figure 8 shows the logarithm of the utility function. Hemispheric and seasonal asymmetries are evident here. In northern winter, latitudes just south of the equator (−3°) optimize the design, in the vicinity of the ITCZ. Conversely, in northern summer, latitudes just north of the equator (3°) optimize the design, again in the vicinity of the seasonally migrating ITCZ. Additional peaks in the data utility can be seen around 30°, in the summer subtropics and again near the descending branch of the Hadley circulation. The equinox seasons have less utility at the optimal designs (3° and −3°). Because the equinoctial Hadley cells and ascent regions in the ITCZ are less pronounced than the solstitial Hadley cells (Schneider et al., 2010), utility is more spread out across the latitudes.

![Figure 5. Logarithm of the data utility as a function of latitude, with designs corresponding to a single latitude. The colored discs signify the four representative designs indicated in Figure 3, which are used in the uncertainty quantification experiment.](image1)

![Figure 6. Daily precipitation rates at equatorial (left) and subtropical (right) latitudes, plotted against the relative humidity and relaxation timescale in the convection scheme. The scatter plots are generated by sampling independently from the prior distribution for the two parameters and then projecting into each parameter dimension.](image2)
We solve the analogue inverse problem as in the statistically stationary case with a sample of data taken at latitudes of ±3° or ±30°, corresponding to the first and second peaks of utility for the solstice seasons. The posterior distributions are collected in Figure 9, colored by season. In general, the true parameter values lie within 50% of the posterior mass in each case. The utilities at the optimal latitudes in northern summer and winter are 131.9 and 154.7, respectively. In contrast, the utilities corresponding to the secondary peaks in the subtropics are 47.9 and 39.5 for northern summer and winter, respectively. As in the statistically stationary case, the design with highest predicted utility (northern winter at 3°) indeed has highest utility. Visually we see symmetry between these seasons, with qualitatively similar distributions in the opposing hemispheres for northern summer and winter. For the equinox seasons, from data sampled at their respective optimal latitudes of +3° and −3° (Figure 10), we see lower utilities of 89.7 and 54.8 for northern fall and spring, respectively, and we see asymmetry most likely indicating sampling variability, because the infinite-time GCM statistics are hemispherically symmetric. In this seasonally varying setting, we again observe that our targeted data acquisition algorithm is a good predictor of informativeness of additional data for learning about the convection parameters.

5. Conclusions and Discussion

We have presented a novel framework for automated optimal data acquisition to calibrate a global model. The framework can be used with computation-
ally expensive and chaotic (noisy) GCMs, whose derivatives may not be available. The data are assumed to be accessible only at limited locations and at different times of year. Given a global simulation, we use parameter uncertainty information provided by the CES algorithm to guide our choice of design (when and where we target data acquisition). We have demonstrated the efficacy of the algorithm for finding optimally informative locations in perfect-model settings in which we generated data with an idealized GCM and learnt about parameters in a convection parameterization. Using statistically stationary or seasonally varying statistics, we have explored both spatial and spatio-temporal designs.

With the idealized GCM, we have targeted a location and time period at which additional data will produce parameter estimates that minimize uncertainty. In our proof-of-concept with narrow designs consisting of data measured only at a single latitude ($\ell = 1$), the automatically targeted optimal location for new data acquisition was, in the seasonal case, in the vicinity of the seasonally migrating ITCZ, with secondary maxima in the summer subtropics. This is consistent with the fact that the convection scheme in the idealized GCM is most important near the ITCZ (O’Gorman & Schneider, 2008b). In the statistically stationary case, regions near the ITCZ are optimal for data acquisition with wider design stencils ($\ell = 3$, Appendix B). However, in scenarios with narrower designs ($\ell = 1$), the subtropical precipitation minimum turns out to be the optimal location, which we confirmed by calibrating convection parameters at this and other locations. We showed that the optimal targeting is limited in its effectiveness when the available data are very noisy (as shown in Appendix C when both the averaging timescale and stencil sizes are reduced). However, the algorithm provides access to the posterior distributions of the parameters, so that this behavior is both diagnosable a posteriori and actionable with successive iterations of the optimal design process (e.g., using the current posterior as the prior for

![Figure 9](image_url). Posterior distribution obtained from using data at the optimal latitudes ($\pm 3^\circ$, left) and second-optimal latitudes ($\pm 30^\circ$, right). The top row corresponds to data targeted to northern summer in the northern hemisphere, and the bottom row corresponds to data targeted to southern summer in the southern hemisphere. Contours bound 50%, 75%, and 99% of the distribution (in decreasing color saturation). The true utility of the northern summer distributions are (left: 131.9, right: 47.9), and southern summer distributions are (left: 154.7, right: 39.5). The true parameter values in the control simulation are given by the blue circle. The parameters found to be optimal in the calibration scheme (given a single random realization of data) are given by the red star in each case.
We also showed that although the algorithm correctly predicts the ordering of information content of different sites in many scenarios, it does not necessarily provide an accurate estimate of the actual information content at the sites, due to sampling variability and the additional model error inflation.

Our algorithm couples the optimization over the design space to the specific application through the posterior distribution of parameters. Therefore, it captures different applications of targeted data acquisition by modifying only the forward map and data entering the loss function to learn this parameter distribution, without changing the algorithm structure. Our framework is thus immediately applicable to the motivating example of automatically targeting embedded high-resolution simulations such as those in Shen et al. (2020) and Shen et al. (2021) to regions that are maximally informative about parameterizations. One could even consider targeting observational data acquisition, such as informing choices for new field campaigns (e.g., Rauber et al., 2007; Stevens et al., 2003), or new in-situ observatory locations (e.g., Stokes & Schwartz, 1994). However, many additional practical considerations beyond the scope of optimal experimental design also play a role in site selection in such cases.

The current algorithm relies on evaluating utilities naively at all design points. Thus, for moderately sized design spaces, the computational cost is dominated by the cost of running the GCM. In practice, if we want to determine $O(10^3)$ limited-area data acquisition sites optimally within $10^6$ or more possible locations, such naive approaches are inefficient. Instead, one can use more sophisticated optimization algorithms. For determinant based (i.e., $D$-optimal) utilities, this typically requires accelerating the determinant evaluation (and its gradients). Various methods have been developed to do so, for example, using Laplace approximations (Beck et al., 2018; Long et al., 2013; Rue et al., 2009), polynomial chaos surrogates (Huan & Marzouk, 2014), optimization of criteria bounds (Tsiliﬁs et al., 2017), fast random determinant approximation (Alexanderian et al., 2014; Alexanderian & Saibaba, 2018), and Gaussian process surrogates (Buathong et al., 2020; Paglia et al., 2020). The latter, kernel-based approaches are particularly amenable to our setting, as they give sparse representations of the utility function that are independent of the underlying computational grid. They may offer a way forward in the climate modeling setting.

As we have presented it here, the algorithm is directly applicable to comprehensive climate models. It will be interesting to explore to what extent application to comprehensive models yields results such as the ones we have seen in the idealized setting: non-obvious optimal locations for targeting computational or observational data acquisition for reducing uncertainties in convection or other parameterization schemes.

Figure 10. Posterior distribution obtained from uncertainty quantification using data targeted at the optimal latitude ($\pm 3^\circ$) from each equinox season. Contours bound 50%, 75%, and 99% of the distribution (in decreasing color saturation). The northern spring (at latitude $+3^\circ$) distribution has utility 54.8, while northern autumn (at latitude $-3^\circ$) has utility 89.7. The true parameter values in the control simulation are given by the blue circle. The parameters found to be optimal in the calibration scheme (given a single random realization of data) are given by the red star in each case.
Appendix A: Calibrate-Emulate-Sample With Design

Key to the success of this work, is the ability to efficiently calculate the posterior distribution (in particular the covariance), which is needed to calculate the utility function (6) at all designs. We present a methodology: calibrate-extract-emulate-sample, (CEES) which allows for the parallel sampling of the posterior distribution at all designs with a combined total of $O(10^5)$ evaluations of our forward model.

The methodology is based on the calibrate-emulate-sample (CES) algorithm, for full details of the individual stages, see Cleary et al. (2021) and Dunbar et al. (2021), here we present an overview and motivation.

The core purpose of CES is to form a computationally cheap statistical emulator of $G_{\infty}$ from intelligently chosen samples of $G_T$; then one is able to solve the Bayesian inverse problem for the emulated $G_{\infty}$ with a sampling method. We achieve this by using Gaussian process emulators, trained on the samples of the (noisy and expensive) forward map. The Gaussian process mean function is naturally smoother than the data it is trained on (Kennedy & O’Hagan, 2001; Notz et al., 2018), and is capable of representing the noise of the forward model within the covariance function, leading to a smooth likelihood function that is quick to evaluate. The training points for the Gaussian Process are given by applying an optimization scheme, Ensemble Kalman Inversion (EKI) (Chen & Oliver, 2012; Iglesias et al., 2013; Schillings & Stuart, 2017), to the inverse problem in its finite-time averaged form Equation 3. Theoretical work shows that noisy continuous-time versions of EKI exhibit an averaging effect that skips over fluctuations superimposed onto the ergodic averaged forward model (Duncan et al., 2022), and similar effects are observed in practice for EKI, thus it is highly suited to optimization of parameters coming from a noisy, expensive model without derivatives available. Ensemble Kalman methods are scalable to very high dimensional problems (Kalnay, 2003; Oliver et al., 2008) with use of localization and regularization.

Let $D$ index a finite space of designs. Given a time $T > 0$, and prior on $\theta$ with prior mean $\theta^*$. Draw a sample $y = G_T (\theta^*, u^{(0)})$, from initial condition $u^{(0)}$:

1. **Calibrate:** We solve Equation 3 with $y$ using evaluations of $G_T$ in an optimization sense, where we minimize the functional.

$$\Phi_T(\theta, y) = \| y - G_T(\theta) \|_{2\Sigma}^2. \quad (A1)$$

The notation $\| \cdot \|_2 = \| \Sigma^{-\frac{1}{2}} \cdot \|_2$ is the Mahalanobis distance. We drop the notation of the initial conditions, which are drawn at random from the invariant distribution for every evaluation of $G_T$. The weight $2\Sigma$ is the sum of internal variability of $G_T$ and of $y$. The optimization is performed using several iterations the Ensemble Kalman Inversion algorithm. This leads to $\{ \theta_j, G_T(\theta_j) \}_{j=1}^J$ of input-output pairs that are localized around the optimal parameter value.

2. **Extract:** For each design $k \in D$, we apply the restriction mapping $W_k$ to the forward map, $\{ \theta_j, W_k G_T(\theta_j) \}_{j=1}^J$, and apply the following *Emulate(k)* and *Sample(k)* stages.

3. **Emulate(k):** We decorrelate the data space with an SVD on the internal variability covariance $\Sigma$, yielding a change-of-basis matrix $V$. We train Gaussian process emulators, on the pairs $\{ \theta_j, V W_k G_T(\theta_j) \}_{j=1}^J$, yielding $(G_{\text{GP}}(\theta), \Sigma_{\text{GP}}(\theta))$, where $G_{\text{GP}} \approx V W_k G_T(\theta)$ (crucially $G_{\infty}$ and not $G_T$) and $\Sigma_{\text{GP}}(\theta) \approx V W_k \Sigma W_k^TV^T$.

4. **Sample(k):** We now solve the inverse problem Equation 5. This is feasible as the emulator provides us with an approximation of $G_{\infty}$ (not just $G_T$). The posterior distribution associated with Equation 5 is proportional to a product of prior and likelihood contribution from Bayes theorem. Explicitly, for a Gaussian prior $N(m, C)$ on the computational parameters, and the likelihood dependent on the emulator, we write the MCMC objective function (also known as the log-posterior) as

$$\Phi_{\text{MCMC}}(\theta, V W_k y) = \frac{1}{2} \| V W_k y - G_{\text{GP}}(\theta) \|_{\Sigma_{\text{GP}}(\theta)}^2 + \frac{1}{2} \log \det \Sigma_{\text{GP}}(\theta) + \frac{1}{2} \| \theta - m \|_C^2.$$
The posterior is then given by
\[ \mathbb{P}(\theta|V_k y) \propto \exp(-\Phi_{MCMC}(\theta, V_k y)). \]

This can be sampled with a standard random walk metropolis sampling algorithm. In practice we run the algorithm for \(2 \times 10^5\) samples, discarding the first \(10^5\) as spin-up.

The CEES algorithm is illustrated in Figure A1. We then collect the posterior distributions \(\{\theta|W_k y\}_k\), \(\forall k \in D\) and calculate the utility function using Equation 6. In particular the algorithm requires \(J\) model evaluations independent of the number of designs.

The CES algorithm is used to solve Equation 7 at a given design \(\hat{k}\), by calibrating with the corresponding objective function for the limited-area data, followed by emulate and sample stages at \(\hat{k}\).

**Appendix B: Results for Three-Latitude Stencil**

For the statistically stationary case, we increase the stencil size to \(\ell = 3\). Here, we have 30 designs indexed from south to north poles. We plot the logarithm of the utility against the designs in Figure B1. The center of the three-latitude stencil is take as a representative latitude for that design. The colored discs represent the designs centered on latitudes \(-8^\circ, -3^\circ, -19^\circ,\) and \(-75^\circ\), in decreasing order of utility on the plot. The increase in spatial extent smooths the design landscape. We validate the optimal design methodology by taking a data sample at each of these representative designs. We then apply the uncertainty quantification stage of the algorithm for each design to obtain the posterior distributions for the convection parameters given each data. The distributions are displayed in Figure B2; panels a–d are ordered according to decreasing predicted utility given by Figure B1. The true utilities for the distributions a–d are 126.0, 35.3, 97.5, and 2.1. In this case, the algorithm has identified the design with maximal utility centered at \(-8^\circ\) (Figure B1a), where analysis of precipitation and parameterized tendencies would suggest the ITCZ region centered at \(-3^\circ\) that presents the bimodal distribution (Figure B1b).

**Figure A1.** Procedure of the uncertainty quantification framework (blue), to produce output (pink). A restriction operator \(W_1\) extracting a subset of the GCM output (yellow); the subsequent emulate and sample stages may be performed in parallel for all \(W_i\), from a single calibration run.

**Figure B1.** Logarithm of the data utility as a function of latitude, with designs corresponding to a three-latitude stencil, the center of which is plotted. The colored discs signify the four representative designs, which are used in the uncertainty quantification experiment.
Appendix C: Results for 30-Day Time Averages

For the statistically stationary case, we also run a suite of experiments for time-averages of $T = 30$ days. The first control simulation at the prior mean $\theta^*$ produces the 600 samples of 30-day averaged control statistics, with which...
we use to approximate $\Sigma(\theta)$. The statistics are represented in Figure C1. We run an experiment for three-stencil designs ($\ell = 3$). Here we have 30 designs indexed from south to north poles. We plot the logarithm of the utility against the designs in Figure C3. The colored discs represent the designs centered on latitudes $-3^\circ$, $-8^\circ$, $25^\circ$, and $-69^\circ$ (in decreasing order of utility on the plot).

To validate the optimal design methodology, we sample the ground truth data at the designs (Figure C2). We then obtain the posterior distributions for the convection parameters given this data. The distributions are displayed in Figure C4, with panels a–d ordered according to decreasing predicted utility given by Figure C3. The uncertainty of the distributions in panels a–d gives utilities 181.6, 97.5, 66.6, and 1.8. In this case, the automated algorithm has identified the optimal stencil correctly.

**Figure C2.** Aggregated climate statistics in the statistically stationary control simulation using the ground truth parameters. Mean (gray lines) and 95% confidence intervals (shading) of the data are plotted against latitude. Additional inflation noise is added. Each set of colored discs represents a 30-day realization of inflated GCM data coming from a different three-latitude design used in the experiment.

**Figure C3.** Logarithm of the data utility as a function of latitude, with designs represented by a node at the center of each stencil (comprised of three neighboring latitudes). The colored discs signify the four representative designs indicated in Figure 3, which are used in the uncertainty quantification experiment.
With the choices $\ell = 1$ and 2, Figure C5 shows the utility function against the latitude at the center of the stencil and the posterior distribution at the respective optimal designs. The behavior of the utility function is similar to the 90-day time averaged case. For $\ell = 2$, we find optimality around the equator; for $\ell = 1$, we find two additional peaks revealed at $\pm 19^\circ$ (see Figure 5). The posterior distributions are seen to be far broader than in the three-latitude case, as less information is available. The posteriors are multimodal but nevertheless capture the true parameters (blue disc) with high probability. They provide insight into the correlation structure between the parameters at the optimal design location. We observe that for these sparser designs, non-identifiability (multi-
modality) appears only at data from $\mathbf{A} \theta^\dagger$, but not at $\mathbf{A} \theta^\ast$. As a result, the optimal uncertainty is not guaranteed to be found at the location of optimal utility. This is remedied by having a better initial guess through the prior, or by having a less noisy data set from which the parameters are more identifiable.

**Data Availability Statement**

All computer code used in this paper is open source. The code for the idealized GCM, the Julia code for the optimal design algorithm, the plotting tools, and the slurm/bash scripts to run both GCM and design algorithms are available at: https://doi.org/10.5281/zenodo.6679974.

**References**


Zhang, M., Bretherton, C. S., Blossey, P. N., Austin, P. H., Bacmeister, J. T., Bony, S., et al. (2013). CGILS: Results from the first phase of an international project to understand the physical mechanisms of low cloud feedbacks in general circulation models. *Journal of Advances in Modeling Earth Systems*, 3(4), 826–842. https://doi.org/10.1002/2013MS000246