



# Process-based climate model development harnessing machine learning: I. a calibration tool for parameterization improvement

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## Key Points:

- We apply Uncertainty Quantification to Single-Column Model/LES comparison to calibrate free parameters
- We revisit model development strategy with an emphasis on processes for model calibration
- The proposed tuning tool allows to formalize the complementary use of multicasers with various metrics

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This article has been accepted for publication and undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the [Version of Record](#). Please cite this article as [doi: 10.1029/2020MS002217](https://doi.org/10.1029/2020MS002217).

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## Abstract

The development of parameterizations is a major task in the development of weather and climate models. Model improvement has been slow in the past decades, due to the difficulty of encompassing key physical processes into parameterizations, but also of calibrating or tuning the many free parameters involved in their formulation. Machine learning techniques have been recently used for speeding up the development process. While some studies propose to replace parameterizations by data-driven neural networks, we rather advocate that keeping physical parameterizations is key for the reliability of climate projections. In this paper we propose to harness machine learning to improve physical parameterizations. In particular we use Gaussian process-based methods from uncertainty quantification to calibrate the model free parameters at a process level. To achieve this, we focus on the comparison of single-column simulations and reference large-eddy simulations over multiple boundary-layer cases. Our method returns all values of the free parameters consistent with the references and any structural uncertainties, allowing a reduced domain of acceptable values to be considered when tuning the 3D global model. This tool allows to disentangle deficiencies due to poor parameter calibration from intrinsic limits rooted in the parameterization formulations. This paper describes the tool and the philosophy of tuning in single-column mode. Part 2 shows how the results from our process-based tuning can help in the 3D global model tuning.

## 1 Introduction

Atmospheric global or regional circulation models used either for numerical weather prediction (NWP) or climate studies encompass a dynamical core and a physical component. The dynamical core computes the spatio-temporal evolution of atmospheric state variables by solving a discrete version of the fluid dynamic equations. The physical component quantifies the impact on the resolved variables of radiative, thermodynamical and chemical processes, as well as dynamical processes that occur at scales smaller than the computational grid. These processes are handled by a suite of sub-models, most often referred to as parameterizations, which provide source terms in the resolved-scale equations. Parameterizations (e.g., turbulence, convection, radiation, microphysics) are often based on a mixture of physical principles and heuristic description of the involved processes, of their interactions and of their impact on the larger resolved scales. Although it is difficult to trace back the origin of the term “parameterization” in climate model-

ing, it semantically points to the fact that the sub-models summarize the processes as functions of the model state vector  $\mathbf{x}$  (typically the value of zonal and meridional wind, temperature and water phases at each point of the 3D model grid) that depends on some free parameters. These free parameters arise from the simplification of the complex nature of the subgrid processes (e.g., assuming a bulk thermal plume instead of a population of plumes, stationarity). The atmospheric model can be summarized as

$$\frac{\partial \mathbf{x}}{\partial t} = \mathcal{D}(\mathbf{x}) + \sum_p \mathcal{P}_p(\mathbf{x}, \boldsymbol{\lambda}_p) \quad (1)$$

where  $\mathcal{D}$  stands for the discretized form of the fluid dynamic equations,  $\mathcal{P}_p$  for the source term provided by the parameterization of the process  $p$  and  $\boldsymbol{\lambda}_p$  for the associated free parameters. This equation may however be too simplistic, as, in reality, a given parameterization often depends on intermediate variables provided by other parameterizations (e.g., cloud fraction used in radiation, turbulence variance used in the cloud scheme) and computes additional prognostic variables (e.g., turbulence kinetic energy). Nevertheless, with this simplified framework, improving models through parameterization development means both to propose more appropriate functional forms  $\mathcal{P}_p$  and to identify acceptable or better values of the free parameters  $\boldsymbol{\lambda}_p$ .

Among the different parameterizations, those involved in the representation of turbulence, convection and clouds still challenge state-of-the art NWP and climate models (Holtslag et al., 2013; Nam et al., 2012; Nuijens et al., 2015; Klein et al., 2017; Randall et al., 2003; Bony et al., 2015). Innovative and diverse concepts and ideas have been proposed over the past decade to improve this representation (Rio et al., 2019). A detailed understanding of the physical processes leading to the formation of low-level clouds can be obtained by Large-Eddy Simulations (LES) (Guichard & Couvreur, 2017), which reproduce, with high fidelity, the turbulent dynamics within the clouds (e.g., Siebesma & Cuijpers, 1995; Neggers, Duynkerke, & Rodts, 2003; Wang & Feingold, 2009). LES are therefore increasingly used to derive and evaluate the conceptual models at the root of boundary-layer and shallow cloud parameterizations. The choice of the parameterization free parameters is also crucial for the simulation of clouds. Their calibration or “tuning” consists in searching for acceptable or optimal values of these parameters, such that the associated model configuration has a realistic behavior under various conditions and compared to a suite of observations (Mauritsen et al., 2012). Calibration is therefore a fundamental aspect of NWP or climate model development (Bellprat et al., 2012;

Duan et al., 2017; Schmidt et al., 2017). However, it is often conducted without much control on the way it modifies the parameterization behavior at the process level as the calibration focuses more on regional or global constraints, such as the radiative balance of the Earth System for climate models, or performance metrics (e.g. root mean square error, skill scores) for NWP models. Hourdin et al. (2017) compile the tuning strategies of several climate groups and emphasize that most of the parameters used to tune climate models (droplet size, fall velocity, entrainment rate) are related to clouds (see also Golaz et al., 2013), i.e. the most uncertain processes that affect radiation, the primary engine of the atmospheric circulation.

Given the societal needs for reliable climate simulations and weather forecasts, the progress achieved by the global atmosphere modeling community has been found slow (Jakob, 2010). Several systematic errors in state-of-the-art models have been modestly reduced, such as those regarding the surface temperature over the eastern oceans (Richter, 2015), the rainfall distribution in the Tropics (Flato et al., 2013), the variability of the liquid water path (Jiang et al., 2012) and the low clouds (Nam et al., 2012). The deadlock of the cloud parameterization, highlighted by Randall et al. (2003), is still an issue today. This too slow improvement of models can be attributed to remaining deficiencies in the structure of the parameterization itself (the function  $\mathcal{P}_p$ ) but also to the calibration of model parameters that can be considered as a bottleneck in model development. On the one hand, the calibration may not be done efficiently enough, and on the other hand, tuning may induce error compensations that contribute to slow model development. Indeed, a new model development usually starts with a model score degradation by breaking this compensation, as often experienced in the weather prediction centers where strong weight on well-established metrics slows down the implementation of new model development in the operational version (Sandu et al., 2013).

Various avenues have been proposed to get around these difficulties and accelerate climate model improvement. A first avenue seeks to exploit the high resolution, explicitly resolving convection, to reduce the number of involved parameterizations. With the recent increase of computer power, it is nowadays possible to run global kilometer-scale resolution simulations over a few months (Satoh et al., 2008, 2019; Stevens et al., 2019). However, the explicit simulation of the fluid dynamics associated with the life cycle of a cumulus requires grid resolution of the order of several tens of meters. Such resolution will not be accessible in the foreseeable future for climate change projections which



require simulations of the global Earth System covering at least several hundreds of years (model spin-up plus transient simulations in response to anthropogenic forcing). The superparameterization approach (Randall et al., 2003) proposes an intermediate pathway by introducing a convection-permitting model in each column of a conventional general circulation model (GCM) to replace the deep convection parameterization (Khairoutdinov et al., 2005). The use of a large-eddy model instead of a convection-permitting model in such framework further removes the boundary-layer and shallow convection parameterizations (Grabowski, 2016; Parishani et al., 2017). A second avenue recently explored the potential of machine learning approaches, which ultimately envisions to replace some parameterizations by neural networks or similar algorithms, properly trained on convection-permitting model simulations or superparameterized GCM (Krasnopolsky et al., 2013; Brenowitz & Bretherton, 2018; Gentine et al., 2018).

A third proposition consists in retaining parameterizations in models but adjoining new tools relying on machine learning to accelerate model development. This choice is motivated by the fact that parameterizations summarize our current understanding of the dynamics and physics of atmospheric processes and offer the power of interpretation, crucial to build our confidence in the extrapolation beyond observed conditions realized by any climate projections. The ESM2.0, proposed by Schneider et al. (2017), belongs to this category. The authors defend that the major progress in Earth-System model development should come from a more systematic use of global observations and high-resolution simulations thanks to machine learning algorithms. They also underline the importance of climate model calibration. In particular, they stress that their new Earth System modeling framework comes with challenges such as developing innovative learning algorithms, identifying the best metrics, combining information from observations and high-resolution, innovating in the design of parameterizations to more easily benefit from new observations or evolution of the models (e.g., refinement of resolution).

Along the same lines, we propose, in this paper, a new approach which allows the development of the parametrizations and their calibration to be tackled at the same time. We argue that a major slowdown of model improvement resides in the difficulty to clearly identify parameterization deficiencies and to properly disentangle them from the inherent calibration of their adjustable parameters at the process and global scales. It is likely that process-scale parameterization improvements are often hidden by the unavoidable full model re-tuning, required to maintain a reasonable radiative balance or acceptable

scores. In the proposed approach, machine learning is harnessed in a principled way to calibrate parameterizations at process level. We promote a more systematic use of the multi-case comparison between Single-Column Model (SCM) and LES to evaluate and calibrate parameterizations. Such a systematic use is not feasible however without more objective and automatic methods than the traditional trial/error approach used to fix parameter values during the parameterization development. Indeed, this trial/error approach is only applicable to one piece of a particular parameterization and one or two relevant cases at most. Here, we aim at assessing a set of parameterizations  $\mathcal{P}_p$  for a series of test cases, which can be formalized as the question of the existence of a sub-space of the parameters  $\lambda_p$  that allows to match metrics between SCM and LES results for the series of cases, within a given tolerance to error.

Hourdin et al. (2017) reviewed the general practice for climate model calibration and proposed three different levels of calibration in a model development: a first calibration at the level of individual parameterizations, then a calibration of each component of the Earth System model and eventually a calibration of the full Earth System model. Distinguishing those three levels may avoid compensating errors that could arise if the calibration is only done at the last level. In this paper, we propose a methodology to address the first phase, *i.e.* the process-level calibration and defend that it can be part of the elaboration of a well-defined calibration strategy based on solid physical and statistical methodologies. By doing so, we tackle model development and parameter calibration together rather than independently as currently done for most climate model development.

Machine learning has already been proposed to calibrate free parameters (e.g., ensemble Kalman filters as in Schneider et al., 2017). The methodology retained here for model calibration uses history matching with Gaussian processes. History matching is an efficient way to explore and reduce the domain of free parameters  $\lambda_p$  and document how a model physics, namely the suite of functions  $\mathcal{P}_p$ , behaves within this domain. Williamson et al. (2013) applied history matching to tune the Hadley Climate Model and stressed its advantage: it accounts for the various sources of uncertainties in assessing the compatibility of the model with the reference: namely the reference uncertainty itself, the uncertainty introduced by the Gaussian process representation of the parameterization, and the intrinsic ability of the model to represent the reference (often referred to as structural error or model discrepancy). History matching inherently deals with the overcon-

165 fidence issue, which emerges when model calibration is addressed as an optimization prob-  
 166 lem (Salter et al., 2019). It has been widely used to calibrate models in astrophysics (Vernon  
 167 et al., 2010), epidemiology (Andrianakis et al., 2017) and hydrocarbon reservoirs (Craig  
 168 et al., 1996). It has been applied to climate models (Williamson et al., 2015, 2017) and  
 169 is starting to be used to find biases in models (McNeall et al., 2019).

170 Whilst history matching has been applied to calibrate 3D models, it has not been  
 171 harnessed for process-level tuning, as we advocate here through application to SCM/LES  
 172 comparison. The SCM approach provides confidence in the model’s ability to represent  
 173 some of the key processes whereas a direct calibration of the 3D global model targeting  
 174 large-scale constraints may hide compensating errors (as discussed in Williamson et al.,  
 175 2017). SCM calibration is able to reduce the domain of the free parameters for a param-  
 176 eterization, information that can be used for efficiently calibrating the full 3D global model  
 177 (as we demonstrate in part II). The breakthrough proposed here was only possible thanks  
 178 to a strong collaboration between the Uncertainty Quantification community and the at-  
 179 mospheric modelers.

180 The present paper focuses on parameterizations involved in the representation of  
 181 boundary-layer clouds. Indeed, well-established case studies exist for such regimes and  
 182 LES have been shown to realistically represent the main processes. However, this method-  
 183 ology can be easily expanded to other parameterizations and other objectives in the Earth  
 184 System.

185 The paper is organized as follows: the next section describes the SCM/LES frame-  
 186 work highlighting its advantages, recalls the different steps used in the development of  
 187 a parameterization and details the new philosophy advocated here. Section 3 presents  
 188 the statistical tool, with a focus on its philosophy and its main ingredients. Section 4  
 189 presents a guideline for its use based on a simple illustration. The paper ends with con-  
 190 clusions in section 5. A companion paper (part II) illustrates the significant advances  
 191 in model development offered by this tool. It exploits process-based calibration for model  
 192 development and shows how this tool provides guidance for the tuning of a 3D global  
 193 model.

## 2 A systematic use of the SCM/LES comparison

Although observations, especially combinations of observations, nowadays provide detailed information at high temporal and spatial resolution on the characteristics of convection and clouds (Masunaga, 2012; Kumar et al., 2015; Bouniol et al., 2016; Masunaga & Luo, 2016), their use for process-level analysis is still hampered by the difficulty of (i) comparing model output to what the satellite measurements exactly sample (although the model to satellite approach with simulators partly resolves this issue) and (ii) identifying the physical processes responsible for such characteristics. Here, we promote the use of Large-Eddy Simulations for the following reasons. LES have the advantage of providing coherent 3D fields characterizing the dynamical and thermodynamical state of the atmosphere. Of course, LES models include turbulence and microphysics parameterizations and thus contain modeling uncertainties, but they have been shown to reproduce the turbulent dynamics of the clouds with high fidelity (e.g., Neggers, Duynkerke, & Rodts, 2003; Heus et al., 2009). As a result, LES have become a central tool in the development of parameterizations of convection and clouds. Their analysis has helped in building the conceptual models behind several parameterizations (e.g., Neggers et al., 2002; Rio et al., 2010). LES are also used for the evaluation of the parameterizations in particular those involved in the representation of boundary layers and shallow clouds (e.g., Ayotte et al., 1996; Golaz et al., 2002; Hourdin et al., 2002; Neggers et al., 2004; Siebesma et al., 2007; Rio & Hourdin, 2008; Caldwell & Bretherton, 2009; Neggers, 2009; Pergaud et al., 2009; Rio et al., 2010; Suselj et al., 2013; Neggers et al., 2017; Tan et al., 2018; Suselj et al., 2019).

For their evaluation, parameterizations are often tested in a single-column framework, particularly relevant for global circulation model parameterizations, which are fundamentally 1D. SCMs are built by extracting, from a 3D model, a single atmospheric column, which integrates the same set of subgrid parameterizations (boundary-layer, shallow convection, deep convection and microphysics schemes) and is run in a constrained large-scale environment (Zhang et al., 2016). The state vector of the SCM simulation is then a restriction to one column  $\mathbf{x}_c$  of the full 3D state vector  $\mathbf{x}$  and Eq. 1 reduces to Eq. 2. The dynamical term  $\mathcal{D}(\mathbf{x})$  becomes a source term  $\mathcal{F}_c$  specified as a function of time and altitude  $z$ ; we however discard this dependency in the notation for simplicity. It can also depend on the column full state vector,  $\mathcal{F}_c(\mathbf{x}_c)$ , if for instance the large-scale advection is separated between a prescribed horizontal advection and a vertical ad-

vection computed as  $-w\partial\mathbf{x}_c/\partial z$ , where  $w$  is an imposed vertical velocity. During the SCM integration, some parameterizations can be deactivated in which case the corresponding source term is either neglected or included in the forcing  $\mathcal{F}_c$ . It is the case for instance when the radiative heating is imposed rather than being computed interactively by the model radiation scheme or when turbulent surface fluxes are imposed rather than computed by the model bulk parameterizations. What really matters in the SCM/LES approach is that both models use the exact same initial and boundary conditions and forcing terms. In a simplified formalism, the SCM thus corresponds to

$$\frac{\partial\mathbf{x}_c}{\partial t} = \sum_{p \in \mathcal{P}_{\text{activated}}} \mathcal{P}_p(\mathbf{x}_c, \lambda_p) + \mathcal{F}_c(\mathbf{x}_c) \quad (2)$$

and the LES to

$$\frac{\partial\mathbf{y}}{\partial t} = \mathcal{L}(\mathbf{y}) + \mathcal{F}_c^*(\bar{\mathbf{y}}) \quad (3)$$

with

$$\mathbf{x}_c(t=0) = \bar{\mathbf{y}}(t=0) \quad (4)$$

where  $\mathbf{y}$  stands for the full LES state vector,  $\mathcal{L}(\mathbf{y})$  to the LES model equations (which include the LES parameterizations),  $\bar{\mathbf{y}}$  to the horizontal-domain average of the LES state vector and  $\mathcal{F}_c^*$  provides a 3D field but consists of the same forcing as the SCM,  $\mathcal{F}_c$  applied identically on each individual column of the LES. The SCM/LES framework thus provides a rigorous comparison between both simulations, as it removes the uncertainties, which may arise from different initial conditions or large-scale forcing when directly comparing SCM to observations. This constrained framework also avoids the need to disentangle parameterization contributions from their coupling with the large-scale dynamics. Another important aspect of the method is that SCM simulations are computationally very cheap. The joint utilization of LES and SCM was first advocated by Randall et al. (1996); Ayotte et al. (1996) and has been, since then, widely used within the Global Energy and Water Exchanges (GEWEX) Cloud System Study (GCSS; Browning et al. (1993) community, now renamed the Global Atmospheric System Studies, GASS, community). One of the most important legacies of this group for the atmospheric modeling community is an ensemble of test cases that connect observations, LES and SCM, and which sample many typical situations over the globe, thought to be of importance for the climate system (e.g., Siebesma & Cuijpers, 1995; Brown et al., 2002; Duynkerke et al., 2004). As such, this framework has been increasingly used in model development (e.g., Hourdin et al., 2013; Gettelman et al., 2019; Hourdin et al., 2020; Roehrig et al.,

2020), all the more so as SCM simulations have been shown to reproduce uniquely the behavior of their GCM justifying the use of SCM simulations for improving weather and climate models (Hourdin et al., 2013; Neggers, 2015; Gettelman et al., 2019).

Traditionally, parameterizations are often tested over a few specific cases for which high-resolution simulations are available (e.g., Ayotte et al., 1996). Recently, the importance of using a wide benchmark of cases covering the different regimes encountered in reality instead of only a limited number of cases has been stressed (e.g., Neggers et al., 2012). We also highlight here the importance of using an extensive ensemble of cases. The use of multi-case is indeed essential for exploring the various degrees of freedom of the parameterization package. A stable boundary-layer case will constrain the turbulent diffusion; the combination of cloud free and cumulus topped convective boundary layers will ensure that cloud cover is obtained for a good representation of convection; transition cases from stratocumulus to cumulus will ensure the extension to stratocumulus regimes, etc. Combining multi cases and multi metrics is a much more robust assessment of model performance as also highlighted by Neggers et al. (2017). To better use multi-cases, one important technical aspect is a common definition, in a predefined acknowledged format, for the description of the setup of reference cases, to be used both to perform SCM simulations or LES. This definition should include the description of the initial profiles and large-scale forcing but also contain information on the configuration to be used (e.g. the type of surface boundary conditions, the existence of any nudging towards reference vertical profiles, the way large-scale forcing are provided). An international initiative is ongoing to agree on the description of the format for this definition file. Such a standard format to define cases will ease the realization of cases by any model and facilitate the share of new cases. The importance of creating libraries of high-resolution simulations representing different climate is another important aspect already identified as a goal by the GCSS community and stressed in Schneider et al. (2017). A common format and the libraries of LES are an important pre-requisite for the tool presented here. In addition, both will contribute to bringing the process-scale community and the community developing global models more closely together.

When comparing SCM and LES, the modeler has to decide which metrics to consider. Various types of metrics can be used. One can directly compare components of the SCM state vector  $\mathbf{x}_c$  to their equivalent in LES, the horizontal domain-average state vector  $\bar{\mathbf{y}}$  (e.g., vertical profiles of potential temperature, specific humidity and less of-

ten wind components). Assessing the ability of the parameterizations to reproduce the time evolution of  $\mathbf{x}_c$  for a given forcing is indeed the ultimate goal. By doing so, one not only tests the behavior of one particular parameterization but also its coupling with the other parameterizations activated in the SCM. This may make the determination of the behavior of the targeted parameterization more difficult and can hide compensating errors: for example, a given temperature turbulent flux can be obtained by different contributions from organized structures and small-scale turbulence when represented by two different parameterizations such as in the Eddy-Diffusivity Mass-Flux framework (Hourdin et al., 2002; Siebesma et al., 2007; Neggers, 2009; Pergaud et al., 2009). Another type of metrics targets parameterization-oriented variables, such as mass fluxes, heating source associated with one part of the motion only, subgrid-scale distribution of temperature or water, cloud vertical structure, updraft vertical velocity, area fraction or entrainment and detrainment rates. The metric, from the SCM point-of-view, is no-longer derived from the model state variables but corresponds to a variable internal to the parameterizations. However, additional uncertainty arises from the way such variables and associated metrics can be derived from LES. For example, clouds can be characterized in an LES as all the grid cells containing condensed water (e.g., Siebesma & Cuijpers, 1995). Combined with thresholds on the vertical velocity, cloudy updrafts can be separated from cloudy downdrafts. The analysis of the joint distribution of variables or the use of ad-hoc passive tracers can also be used in the LES to identify objects relevant with the conceptual model of the parameterization (e.g., Couvreux et al., 2010; Rio et al., 2010; Chinita et al., 2018; Briant et al., 2019). Such parameterization-oriented diagnostics have helped in the refinement of the conceptual model at the root of the parameterization (e.g., Rio et al., 2010; Jam et al., 2013; Rochetin et al., 2014). However, a question arises if such diagnostics should also be used as metrics in the calibration process. Answering this question on the relative importance to give to one type of metrics or another requires efficient algorithms, as the one proposed here, to explore the various options. Note also that using state vector-based metrics on a large set of cases that are more or less sensitive to one aspect of the parameterization may help avoid the error compensation issue.

In line with Neggers et al. (2012), we advocate that, although not a new approach, the power of SCM/LES comparisons is largely underestimated and under-exploited. Applying history matching to this comparison is a way to fully take advantage of the SCM/LES on a large multi-case ensemble and explore whether there exists a sub-space of the pa-

parameter space for which the SCM is able to reproduce a series of LES simulations within a given uncertainty. Note that the metrics can be different from one case to the other. This tool offers the possibility to revisit the different intercomparison exercises documented in the literature and to benefit from this rich database still underused.

Eventually, a point that becomes crucial when using LES for parameterization evaluation and tuning is the assessment of LES reliability and its uncertainties. Although it has been shown, through the comparison to observations, that LES is able to correctly reproduce boundary-layer processes and shallow clouds (Couvreur et al., 2005; Neggers, Jonker, & Siebesma, 2003; Heus & Jonker, 2008), LES, as in many models, come with uncertainties associated to the advection scheme and the parameterizations still active in such simulations concerning small-scale turbulence, microphysics, radiation and surface fluxes. Sullivan and Patton (2011) have shown that a horizontal resolution of a few tens of meters for convective boundary layers is enough to get convergence for the mean, fluxes and variances but 10m resolution is needed in order to get convergence on skewness. The sensitivity of LES of shallow convection to resolution, size of the domain, sub-grid model and advection scheme has been widely investigated (Brown, 1999; Matheou et al., 2011; Pressel et al., 2017; Zhang et al., 2017; Wurps et al., 2020). In particular, it has been shown that most of the ensemble-averaged turbulence statistics are reasonably insensitive, allowing one to use LES results to develop and evaluate convection parameterizations. However, some characteristics of the cloud fields (e.g. size distribution of individual clouds) are more sensitive to resolution, advection scheme or subgrid-scheme (Brown, 1999; vanZanten et al., 2011; Pressel et al., 2017). For example, LES at 5-10m vertical resolution still have large uncertainties in boundary-layer regimes with sharp inversions where the LES subgrid turbulence parameterization is significantly active. Uncertainty around this reference should be documented so that history matching can explicitly take it into account.

### **3 *High-Tune Explorer* (htexplo), a statistical tool to calibrate model parameters and more**

#### **3.1 Overview**

The present section describes the tool proposed to perform process-based calibration. Its objective is twofold: (i) characterize the domain of the model parameter values that allows the model to appropriately capture process-level metrics and which can



be used for subsequent calibration of the global model, and (ii) identify the model parameters that limit model performance and thus highlight the need for model parameterization revision. The tool relies on history matching approach developed by Vernon et al. (2010) and first used for climate studies by Williamson et al. (2013). This method aims at removing “unphysical” regions of parameter space iteratively, refocusing the search for “acceptably tuned” models at each step. The tool finds the subspace of the model parameter space containing simulations consistent with the reference metrics, acknowledging the various sources of uncertainty. This tool has already been successfully applied to identify the acceptable range of model parameter values in the 3D configuration of the Hadley Centre climate model (Williamson et al., 2013, 2015) or in the NEMO oceanic model (Williamson et al., 2017). It is here used for the first time in the context of the SCM/LES comparison for a given set of cases.

As already stated in the previous section, we focus here on the parameterizations involved in the representation of boundary-layer clouds (turbulence, convection, cloud micro and macrophysics, radiation). However, this methodology can be easily expanded to other parameterizations and other objects of the Earth system as soon as reliable references are available.

Figure 1 sketches the main steps of the *High-Tune Explorer* (htexplo in the following for an explorer to use High-resolution simulation to improve and Tune parameterizations) tool:

- **1. Metric selection and references** First, the cases and associated target metrics are selected. The relevant reference for each metric is then identified and the associated uncertainty is estimated. In the present case, the reference is an LES and the associated uncertainty is based on an LES ensemble. Observations could also be used with an associated error when an LES is not available. This phase is not model-specific and could be shared between different models.
- **2. Selection of model parameters** The model parameters to be calibrated are identified and their possible range of values are determined.
- **3. Experimental design and SCM runs** The experimental design consists of defining the ensemble of experiments (or SCM) to be run. The goal is to optimally sample the parameter space and provide a small set of parameter values for which

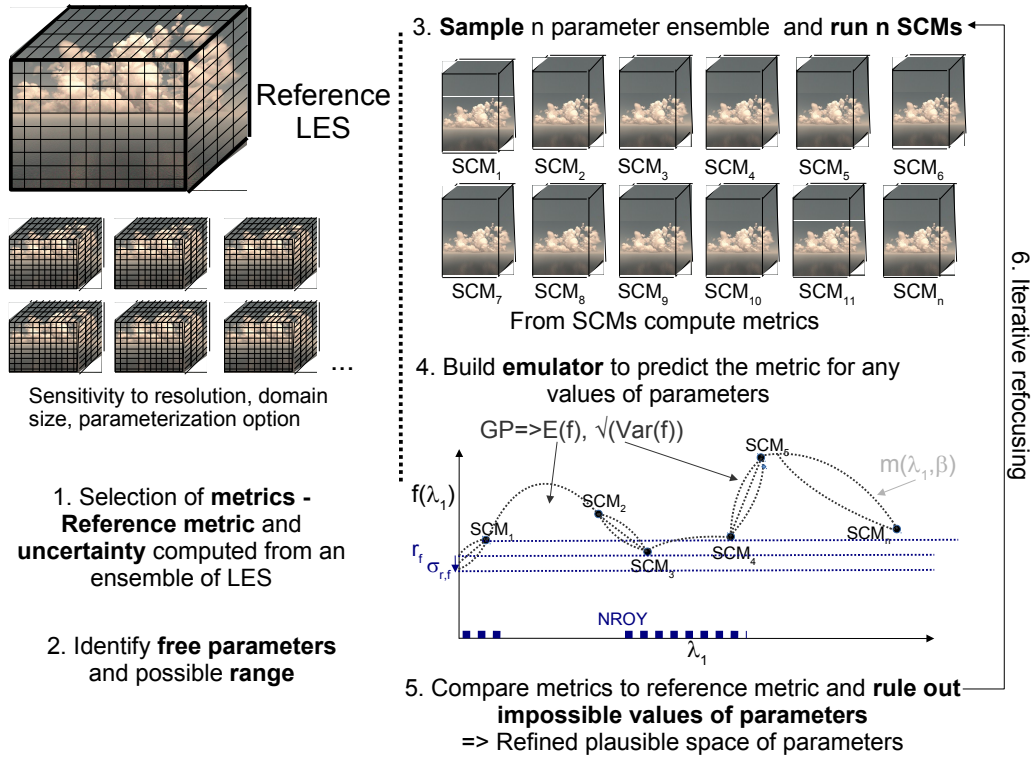
the single-column model will be run. Metrics are computed from each of the SCM simulations and form the training data-set on which emulators are built.

- **4. Building emulators**, i.e. construction of surrogate models, also called “emulators”, one for each metric. Each emulator is based on a Gaussian Process (GP) and predicts the corresponding metric value at any point of the full parameter space, without running the SCM. The GP statistical model also provides a probability distribution of its prediction, thus quantifying the prediction uncertainty for use in calibration.
- **5. History matching** The comparison between the reference metrics and those inferred with the emulators is based on a distance that accounts for reference uncertainty, modeler tolerance to error or model discrepancy (induced by e.g., misrepresentation of specific processes, inaccuracy of numerical solvers, model resolution) and emulator uncertainty. History matching rejects parameter values that lead to unacceptable model behavior (too large distance from the reference) and thus defines a not-ruled out yet (NROY) space, the model parameter space that cannot be further reduced given the sources of uncertainty.
- **6. Iterative refocusing** To reduce the emulator uncertainty, but only where needed, new iterations (or waves) following steps 3 to 5 are performed, sampling the NROY space obtained at the end of the previous wave for the design and only constructing emulators over the NROY domain.

This tool is available freely under: <https://svn.lmd.jussieu.fr/HighTune>. Details on the different steps are given below. For simplicity, we first describe them for the first iteration and only one metric. Subsequent iterations and the addition of other metrics are discussed in section 3.7. This section ends with a discussion about the relationship between the present tool and more common tools used for calibration and sensitivity analysis.

### 3.2 Step 1: Metric selection and references

The metrics used to evaluate the SCM behavior depend on the physical situation considered and the parameterization hypothesis. Scalar metrics based on a dynamical or thermodynamical variable (e.g., potential temperature, water vapor mixing ratio, wind speed, cloud fraction) sampled at a given time can be used, such as the value at a given



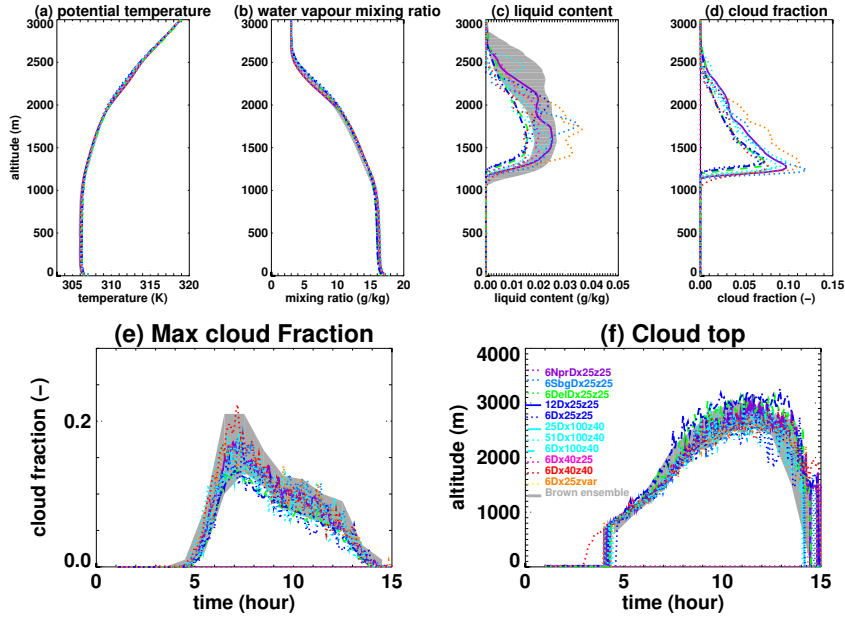
**Figure 1.** Schematic of the different steps of the htxplo tool

vertical level, the average or the maximum over a given layer (e.g., boundary layer, cloud layer), or the maximum over the whole atmospheric column. Radiation-oriented metrics are particularly relevant to enhance the link between the present process-oriented model calibration and the calibration of the corresponding 3D configuration. Ideally, the chosen metric should be as insensitive as possible to the model vertical resolution. In that regard, integrals (or averages) are good candidates for scalar metrics, as will be illustrated in Part II. Root-mean square errors are not encouraged for two reasons, i/ there are usually associated to a smaller signal to noise ratio and ii/ the implausibility (see section 3.6) is already a kind of root-mean square error. The number of metrics to be used is generally of the order of ten, but it can be many more.

More complex metrics such as vertical profiles, time series or spatial fields, can also be considered. In that case, methods are used to reduce the dimensions of the outputs and principal component decomposition is one option (e.g., Salter et al., 2019). However, scalar metrics, taken at a given time, or averaged over a short period of time, seem

often sufficient to robustly constrain most of the SCM simulations. Therefore, in the present paper and in Part II, only scalar metrics will be used.

References and their associated uncertainty are estimated from an LES ensemble. There are a priori two possibilities to build such an ensemble, which can be combined. The first consists in building the ensemble from simulations performed by different large-eddy models, as has been done in several GCSS intercomparison exercises (Brown et al., 2002; Siebesma et al., 2003; Stevens et al., 2005; vanZanten et al., 2011; de Roode et al., 2016). The reference thus corresponds to the LES ensemble mean, while the uncertainty is quantified by the LES ensemble variance. The second option, used in this paper, relies on only one large-eddy model and estimates the uncertainty around the reference model configuration by performing sensitivity experiments to horizontal and vertical resolution, domain size, and parameterization options (e.g., turbulence, microphysics, surface fluxes, radiation). In this study, we have chosen to use the simulation realized with the higher resolution over the largest domain and with the most relevant parameterization options as the reference, but the ensemble mean could also be used. The large-eddy model is the LES-configuration of Meso-NH (Lac et al., 2018). It makes use of a fourth-order centered discretization associated with an explicit fourth-order Runge-Kutta time integration. Figure 2 illustrates the spread obtained from a Meso-NH LES ensemble exploring the sensitivity to horizontal, vertical resolution, domain size and options in the turbulence and cloud schemes for one given case, namely the ARM Cumulus case, which is a golden case for the study of continental cumulus (Brown et al., 2002). Table A2 in the Appendix describes the different simulations used to estimate the uncertainty. Consistently with the literature (Brown et al., 2002; Matheou et al., 2011; vanZanten et al., 2011; Zhang et al., 2017), domain-average conserved thermodynamical quantities are weakly sensitive to changes in resolution, domain size and parameterization choices while the domain-average liquid water content and cloud fraction exhibit more spread. Metrics derived from those latter quantities will therefore be associated to a larger uncertainty. Figure 2 also indicates in grey shading the spread obtained from the LES intercomparison of Brown et al. (2002) highlighting a similar uncertainty estimate between the two methods mentioned above. Similar results are obtained for LES ensembles of other intercomparison exercises (not shown). For a given metric  $f$ ,  $r_f$  is the reference metric value, estimated from the reference LES simulation or the average of the LES ensemble and  $\sigma_{r,f}^2$  is the associated square error estimated from the LES ensemble. Note that, in the absence of available



**Figure 2.** Vertical profile of (a) potential temperature, (b) water vapor mixing ratio, (c) liquid water content and (d) cloud fraction averaged over the horizontal domain at the 10<sup>th</sup> hour of the simulation (1530 LT) and time series of (f) the cloud top and (e) the maximum cloud fraction over the atmospheric column. The grey shading corresponds to the results of the Brown et al. (2002) intercomparison. The different color lines correspond to different sensitivity tests realized with Meso-NH changing either, one by one, the size of the domain, the vertical or horizontal resolution and some option in the cloud scheme, microphysics scheme or turbulence scheme (detailed in Table A2).

LES, observations can also be used as a reference to be compared to the SCM runs as illustrated in Ahmat Younous et al. (2018) but the observation error needs to be quantified.

### 3.3 Step 2: Selection of model parameters

The number of model parameters can be large (generally on the order of 10 for each parameterization). Estimating the prior range of values that needs to be explored for each of them requires the modeler's expertise. The definition of this range is an important step as the results are only valid in this predefined parameter space (Williamson

et al., 2013). So, we advise to choose a range as wide as possible in the absence of physical reasons or numerical concerns for constraining it. Nevertheless, the user might consider some tradeoff as the smaller the ranges, the smaller the space to explore.

As the tool samples any parameter independently from the others (see Step 3), the method remains efficient even though a parameter with no influence on the results was included. A sensitivity analysis (Oakley & O’Hagan, 2004) could be used as a preliminary step in order to reduce the number of selected parameters but may not be a good idea in general (see section 3.8). The user can consider either linear or logarithmic variations of the parameter values.

In the following, we consider a set of parameters  $\boldsymbol{\lambda} = (\lambda_k)$ , where the  $k$  parameters are a subset of the model parameters involved in the different parameterizations (see section 1).

### 3.4 Step 3: Experimental design and SCM runs

Once the model parameters are selected and their range of values defined, an experimental design is built. It corresponds to the selection of a relatively small set of values for the model parameters  $(\lambda_i)_{i=1,\dots,n}$ , usually on the order of ten times the number of parameters, as discussed in Loeppky et al. (2009). It explores the initial (or input) space of the parameter values in the range given for each parameter. An SCM simulation is performed for each of them and provides the state vector  $\mathbf{x}_c(\boldsymbol{\lambda}_i)$ . The objective is to "fill" the parameter space as uniformly as possible maximizing the minimum distance between points. Here, as classically used for the design of computer experiments, a Latin Hypercube (LHC) (Williamson et al., 2015) is used to efficiently sample the input parameter space. Classically, a LHC for a  $n$ -member ensemble uniformly divides each dimension of the input space into  $n$  bins that are sampled once each and only once. All the parameters are thus varied simultaneously in contrast to other sensitivity analysis approaches such as in the Morris sensitivity analysis (Saltelli, 2002), where parameters are varied one by one. The LHC sampling used here maximizes the minimum distance between the selected points of the input space.

More precisely, here we use  $k$ -extended latin hypercubes as proposed by Williamson (2015). It consists in producing several LHCs, added sequentially, which ensure that each additional LHC samples an area of the space that has not been sampled yet by the pre-

vious LHCs. Such a design provides the advantage of being able to robustly check the GP performance on well-designed sub-LHCs.

### 3.5 Step 4: Building emulators

The selected metric (see Step 1) is computed for each SCM simulation, noted  $f(\lambda_i)$  for  $i = 1, \dots, n$ . These numbers serve as a training dataset for the building of an emulator. The emulator is then used to predict the metric values  $f(\lambda)$  for any vector of parameter values  $\lambda$  in the input space. A separate emulator is constructed for each metric.

Specifically, we use a Gaussian process (GP), a well known statistical model which has the advantage of interpolating observed model runs and provides a probabilistic prediction. The emulator gives a probability distribution for  $f$  written as:

$$f(\lambda) \mid \beta, \sigma^2, \delta \sim \text{GP} \left( m(\lambda, \beta), k(\cdot, \cdot, \sigma^2, \delta) \right),$$

where  $m(\lambda, \beta)$  is a prior mean function with parameters  $\beta = (\beta_i)_i$  and  $k$  a specified kernel (a covariance function describing the covariance between any 2 points). The kernel has a parameter that normally controls variance,  $\sigma^2$ , and parameters  $\delta_k$  for each dimension of the input parameter  $\lambda_k$  that control the correlation attributed to each input. To start with, we assume a stationary kernel, i.e., the covariance only depends on the distance between points and not the absolute position. The GP is such that any finite collection  $f(\lambda_1), \dots, f(\lambda_n)$  has a multivariate normal distribution with mean vector  $m(\lambda_1, \beta), \dots, m(\lambda_n, \beta)$ , and variance matrix  $\Sigma$  with  $\Sigma_{ij} = k(\lambda_i, \lambda_j, \sigma^2, \delta)$ . Let the training data be  $F = (f(\lambda_i))_{i=1, \dots, n}$ , then

$$f(\lambda) \mid F, \beta, \sigma^2, \delta \sim \text{GP} \left( m^*(\lambda, \beta), k^*(\cdot, \cdot, \sigma^2, \delta) \right),$$

where there are well-known closed form expressions for  $m^*$  and  $k^*$  (Williamson et al., 2017). Note that  $m^*$  and  $k^*$  are the updated mean and covariance representing what the emulator has ‘learned’ from the data,  $F$ .

Whilst there are many possible prior choices of  $m$  and  $k$ , htxplo uses a 2-phase approach. First, we impose a structured mean surface  $m(\lambda, \beta) = \beta^T g(\lambda)$  as a linear combination of simple functions of the input parameters contained in the vector  $g(\lambda)$  (e.g. monomials, Fourier functions and interaction terms are chosen through the forwards selection and backwards elimination method described in Williamson et al., 2013)). In

the second stage, we use the squared exponential kernel function and Hamiltonian Monte Carlo (HMC, implemented in Stan – Carpenter & Coauthors, 2017) to sample from the posterior distribution of the parameters  $\beta$ ,  $\sigma^2$ , and  $\delta$  given  $\mathbf{F}$  (note that the mean surface  $m(\lambda, \beta)$  is not directly fitted in phase 1, but its structure is chosen, with Bayesian inference ultimately used in fitting for phase 2).

The choice of HMC implemented in Stan was motivated by requiring robust automation of emulator building across many metrics and cases. Stan affords us with the ability to specify flexible and intuitive priors, and we use weakly informative priors as advocated by Gelman (2006). With the exception of the intercept term (which is uniform), our prior for each  $\beta$  is  $N(0, 10)$  and we use the ordinary least squares (OLS) fitted values as starting values for the HMC. We set  $\delta_k \sim \text{Gamma}(4, 4)$  for all  $k$  to allow a wide range of potential correlation structures (this is a weakly informative prior) whilst penalizing very small values that typically have high likelihoods, but lead to emulators with no predictive power (for discussion, see Volodina, 2020). Our prior for  $\sigma^2$  is a truncated Normal (at 0), with mean at the residual from our OLS fits, and variance set using the variability of the ensemble (full details for these choices in Volodina, 2020).

The emulator is then tested using standardized Leave One Out diagnostics (e.g. Rougier et al., 2009) on the training data. These tests remove one point at a time from the training set and use the emulator fitted on the remaining data to predict the removed point. Repeated over the training set, we then check whether the majority of left out points lie within 95% prediction intervals (we would expect 5% to miss). Another check consists in removing a subdesign of the training set and attempting to predict it based on the new reduced training set. If the emulator fails these checks we revisit the computation of the emulator. For example, the procedure described in Volodina and Williamson (2020) (and available in `htexplo`) can be used to derive an appropriate non-stationary kernel  $k$  before refitting the emulator by HMC. Once fitted, the GP expectation  $E[f(\lambda)]$  provides an estimation of the metric for any given  $\lambda$ , and its variance  $\text{Var}[f(\lambda)]$  provides an uncertainty around this estimation.

SCM runs are computationally cheap, but the fitted emulators are even cheaper and thus allow the computation of millions of predictions, with associated uncertainties, in a short time (a few minutes). This enables us to numerically define the space containing acceptable sets of parameters with respect to the chosen metrics and in particular,



to visualize it (Step 5). The choice of Stan has proven effective for this project, though it does not scale well to larger ensembles. Going forward, a new version of the tools defaulting to MAP estimation and using efficient parallel implementation has just been released enabling millions of predictions in just a few seconds (Williamson & Volodina, 2020).

### 3.6 Step 5: History matching

The htexplo tool relies on the history matching technique, which seeks to rule out parameter values from the input space that are “implausible”, given the SCM behavior for these parameter values and the sources of uncertainty. These sources include the reference (observation) error, treated as a random quantity with mean 0 and variance  $\sigma_{r,f}^2$ , and the SCM discrepancy, which has mean 0 (unless the user knows the direction in which the model is biased) and variance  $\sigma_{d,f}^2$  (Sexton et al., 2011). The emulator is used to estimate the model behavior on a much larger sample of the input space than possible with the SCM. To history match the SCM behavior, we introduce the “implausibility” measure for the metric  $f$  (Williamson et al., 2013),  $I_f(\boldsymbol{\lambda})$ , which is a distance between the metric prediction  $f(\boldsymbol{\lambda})$  by the emulator at  $\boldsymbol{\lambda}$ , and the reference metric value,  $r_f$ , with respect to the norm induced by our second-order uncertainty specification, noted  $\|\cdot\|_H$  below. The implausibility reads

$$\begin{aligned} I_f(\boldsymbol{\lambda}) = \|r_f - f(\boldsymbol{\lambda})\|_H &= \frac{|r_f - \mathbb{E}[f(\boldsymbol{\lambda})]|}{\sqrt{\text{Var}[r_f - \mathbb{E}[f(\boldsymbol{\lambda})]]}} \\ &= \frac{|r_f - \mathbb{E}[f(\boldsymbol{\lambda})]|}{\sqrt{\sigma_{r,f}^2 + \sigma_{d,f}^2 + \text{Var}[f(\boldsymbol{\lambda})]}}. \end{aligned} \quad (5)$$

The model discrepancy for the metric  $f$ ,  $\sigma_{d,f}$ , accounts for the model structural error due to the inherent inability of the SCM to reproduce the LES exactly (due to unresolved physics or missing processes, for example). It could be defined as the minimum error possible when exploring the full set of parameters, however, this could permit the SCM to be close to the reference for the wrong reasons and does not account for multiple metrics and cases, so we avoid this definition. Instead it is typically defined to be the uncertainty left in the difference between the SCM metric when the parameters are fixed at their best values (fixed the same for all metrics) and the references. This quantity is perhaps the target of model development in the first place and, as such, is unknown. For example, suppose we want to test the ability of a new parameterization to capture the behavior of the reference. With the standard definition of discrepancy, the uncertainty needed so that the new parameterization captures the behavior of the reference,

it is not clear how to proceed with testing. Our approach instead is to treat model discrepancy as a “tolerance to error” as detailed in Williamson et al. (2017). The tolerance to error is the distance between model results and the reference that the modeler would be satisfied with, enabling modelers to place confidence in certain metrics/parts of their parameterization, and relax restrictions on others as needed. As illustrated in section 4 and Part II, defining this tolerance to error can be a difficult a-priori task; however experimenting with this value provides important insights into the behavior of the model and its inherent limitations. The most attractive feature of this approach to discrepancy is that, for a given tolerance to error, if the induced NROY space is empty it means that the parameterization is not able to reproduce the reference under the given tolerance. Either the tolerance can be relaxed, accepting the limitations of the current set of parameterizations, or the parameterization can be revisited.

The implausibility defines a membership rule for NROY space after the first iteration:

$$\text{NROY}_f^1 = \{\boldsymbol{\lambda} \mid I_f(\boldsymbol{\lambda}) < T\}.$$

where  $T$  is a chosen threshold (or cutoff). For scalar metrics, it is standard to use  $T = 3$  justified using Pukelsheim’s rule that states 95% of the probability density for any unimodal distribution is within 3 standard deviations of the mean (Pukelsheim, 1994). Using this threshold makes it unlikely that good parameter values are ruled out by chance. To measure and visualize NROY space the implausibility  $I_f(\boldsymbol{\lambda})$  is calculated on a random LHC sampling of a large number (on the order of hundreds of thousands or millions) of vectors  $\boldsymbol{\lambda}$ .

Note that  $I_f(\boldsymbol{\lambda})$  can be smaller than the chosen threshold  $T$  either because  $E[f(\boldsymbol{\lambda})]$  is close to the reference or because the sum of the different errors is large. When the uncertainty of the emulator is larger than the tolerance to error and observation error, points that should be ruled out are kept in the NROY. In this case, further iterations are desirable in order to increase the density of the sampling of NROY and hence improve the emulator quality and reduce the associated uncertainty.

### 3.7 Iterative refocusing and multi-metrics

One advantage of this method is to progressively optimize the design of simulations to be run. New simulations are iteratively added only where it is useful to increase the

emulator accuracy. This is performed by iterating the same process previously described several times in "waves", (this is termed "iterative refocusing" and is a fundamental part of the history matching approach). Each new iteration  $n$  starts from the remaining space  $\text{NROY}_f^{n-1}$  estimated at the end of the previous wave. Because of its complex geometry, a LHC sampling, as in the first wave, cannot be applied, and therefore the remaining space is re-sampled uniformly. A new SCM simulation ensemble is performed with this design and is used to proceed with steps 4 and 5. The new emulator is only valid in the new parameter space, namely  $\text{NROY}_f^{n-1}$ . Outside this space, we rely on the emulators from the previous waves. As in Step 5, to measure and visualize  $\text{NROY}_f^n$ , the implausibility is computed over a large number of points in the input space. The threshold  $T$  may be varied between waves, but we advise to keep it to 3 as long as the process has not converged (i.e. the emulator variance within the current NROY space remains large – see also section 4 and Part II). The iterative refocusing stops when the convergence of the sequence  $(\text{NROY}_f^n)_n$  has been qualitatively achieved.

So far, we have considered only one metric, but several metrics  $(f_k)_k$  can be combined at the same time. An implausibility is then computed for each metric and the total  $\text{NROY}^n$  space is the intersection of the  $\text{NROY}_{f_k}^n$  associated with each metric:

$$\text{NROY}^n = \bigcap_k \text{NROY}_{f_k}^n = \{ \lambda \mid \#\{k \mid I_{f_k}^n(\lambda) > T\} \leq \tau \},$$

$\#$  represents the number of metrics fulfilling the condition indicated into brackets (where the implausibility is greater than the threshold) and  $\tau$ , the number of metrics for which the model is allowed to be far from the reference while still kept in the NROY space. If  $\tau = 0$ , all metrics must satisfy our implausibility cutoff. If there are a large number of metrics then  $\tau$  should be increased ( $\tau \geq 1$ ) to avoid multiple testing problems meaning that too many good parameter values are ruled out by chance. If a modeler seeks to prioritize certain metrics, they can either be introduced in early waves, ensuring that the NROY space satisfies priority metrics first before introducing new ones, or the tolerance to error, which is defined for each metric, can be used to impose priorities (a larger tolerance to error induces a less constraining metric).

### 3.8 Sensitivity analysis provided by the tool

The htexplo tool provides its own sensitivity analysis, which, due to the use of multi-wave history matching, is rather different from traditional methods applied to models

throughout the literature (Bastidas et al., 2006; Guo et al., 2014; Johnson et al., 2015). Traditional methods, either derivative-based (Saltelli, 2002), or variation-based (Oakley & O’Hagan, 2004), essentially seek to identify which parameters modify model output. This can help focus further study, model development or even observation collection to help understand these parameters. Note that the htexplo tool provides at the first iteration a sensitivity analysis over the entire space where correlation among parameters is included as the parameters are not varied one at a time.

However, for calibration purposes, once history matching is considered as a valid approach for a given model, the sensitivity analysis should not be done on the full model input space. By using history matching, we acknowledge that there is a large part of the model parameter space that is not useful for understanding reality. The Gaussian processes remove this uninformative space in order to target the space where the model becomes useful. Once we have this useful subspace, the usual and important questions that are posed by sensitivity analysis should be considered. For example, how is the model output changing as we move through parameter space and which parameters are responsible for these changes? As will be illustrated in section 4, the NROY visualization allows us to see, as we move in two dimensions of a parameter space, in addition to the possible values of each parameters, which combinations of parameters it is important to get right. As all models within the NROY space are consistent with our metrics, sensitivity analysis as described here is now really focused on the relevant subspace. Note that sensitivity analysis on the original input space does not answer these questions. Seen through the history matching lens, on the full space, sensitivity analysis is showing us which parameters are responsible for the variability in the space we are about to cut. Whilst informative for helping us cut the space efficiently, sensitivity analysis is not necessary at this stage. Our methods are already efficiently able to do this. As well as all of the benefits we have for tuning, we would argue that history matching is achieving many of the same things that a sensitivity analysis achieves in terms of informing the modeling, but concentrated only on the model input space that is consistent with the observations.

Performing variance-based sensitivity analysis in NROY space is not trivial and we are not aware of any methods that are currently able to do this. Variance-based sensitivity analysis requires independent input spaces (which is what we always start with in Wave 1). But after cutting space, we have complex relationships between the parameters. NROY space may not even be simply connected, and can be highly non-linear. Ef-

efficient methods for calculating sensitivity in these unusual spaces would be interesting to apply for history matching as an avenue for further research.

### 3.9 On the use of history matching and the avoidance of optimization

Whilst history matching is well established and is being used in a growing number of climate studies, other methods of calibration are more popular and we believe should be avoided for process-based model development. Whilst many methods based on optimizing a cost function exist (Hourdin et al., 2017), the most popular in the UQ community is Bayesian calibration (Kennedy & O’Hagan, 2001). Bayesian calibration requires a similar set up to history matching (emulators, observation errors and model discrepancy) and then jointly finds the posterior probability distribution of the “best” value of the input parameters and the model discrepancy (strong prior information on the discrepancy is required to make this sensible, Brynjarsdóttir & O’Hagan, 2014). Optimization methods like these do not afford us with the chance to falsify a parameterization (they always find the best value), nor do they give all parameter values that are consistent with the observations (in our case reference LES) that can then be used when tuning the 3D model (see Part II).

## 4 Illustration of htexplo on a simple case

In this section, the use of htexplo is illustrated for the ARPEGE-Climat 6.3 atmospheric model (Voldoire et al., 2019; Roehrig et al., 2020) based on a single 1D case. More comprehensive exploitation of the tool will be given in Part II.

### 4.1 Model, parameters and case-study

We use the SCM version of ARPEGE-Climat 6.3, the atmospheric component of the CNRM-CM6-1 climate model (Voldoire et al., 2019; Roehrig et al., 2020) and aim at analyzing the importance of the values of free parameters of the turbulence scheme (based on Cuxart et al., 2000) on the simulation of an idealized clear boundary layer. Details on the ARPEGE-Climat atmospheric component, the turbulence scheme and the used configuration are given in the Appendix B. Among the different free parameters of the turbulence scheme, three are selected for this analysis.  $A_\epsilon$  controls the expression of the dissipation length-scale as a function of the mixing length-scale;  $A_U$  and  $A_T$  re-

spectively enter into the expression of the exchange coefficient in Eq. B3 for the wind and the temperature (the same coefficient,  $A_U$ , is used for both the zonal and meridional component of the wind). The range of variation explored for each parameter is indicated in Table 1 and the parameters are varied linearly in those ranges. The turbulence parameterization includes other free parameters but the three most influential parameters for this case have been selected and no free parameters of the mass-flux scheme are considered.

**Table 1.** List of the free parameters of the turbulence scheme that are varied in this example with default values and range of variation

Names	Default	Minimum	Maximum	Parameter Description
$A_U$	0.126	0.01	0.4	Affects the eddy-diffusivity of momentum
$A_\epsilon$	0.85	0.1	3.	Controls the dissipation length-scale
$A_T$	0.14	0.01	1.	Affects the eddy-diffusivity of temperature

To keep the example simple, only one case is used here. This case is a dry idealized case of a convective boundary layer with a constant-in-time large surface sensible heat flux of  $270 \text{ W m}^{-2}$  ( $Q^* = 0.24 \text{ K m s}^{-1}$  in, Ayotte et al., 1996) with a strongly capped boundary layer, called 24SC in the following. The importance of combining different cases will be illustrated in part II.

We first document a sequence of three waves where additional metrics are added at each iteration (Experiment 1). We will then discuss the results obtained when adding all the metrics directly at Wave 1 (Experiment 2), varying the threshold used to determine the NROY (Experiment 3 see also section 3.5), using more SCM runs (Experiment 4), and varying the tolerance to error (Experiments 5 and 6).

#### 4.2 Three consecutive waves adding metrics progressively

For the first iteration (or wave in the following) of Experiment 1, 30 SCM simulations of the 24SC case were realized by varying values for the three parameters exploring at best (using a LHC sampling, see section 3.4) the range of each parameters (Table 1). Figure 3 illustrates that the parameters are randomly sampled as indicated by

the distribution of the black dots along the different x-axes. Three different metrics are used to characterize the turbulent mixing in the boundary layer and are progressively introduced through the successive waves. The first chosen metric is the potential temperature averaged over the layer 400-600 m. It is a good proxy for the boundary-layer potential temperature, which is well mixed between the surface and the boundary-layer top, located around 1300 m. This metric is computed for the 30 SCM runs; these computations serve as training data for the construction of the emulator. The prior mean function (see section 3.5),  $m$ , for this emulator is a sum of linear and quadratic functions of the parameters. The stationary squared-exponential kernel provides a sufficient fit to the data according to the leave-one-out methodology. Figure 3 presents the variation of the metric as a function of the parameters: some first-order relationships appear with the boundary-layer potential temperature increasing with  $A_U$  and  $A_T$  to a lesser extent (due to an increased mixing associated to a larger diffusivity and larger fluxes) and decreasing with  $A_\epsilon$  (due to a reduced mixing because of the increased dissipation). For this metric, we have chosen a tolerance to error of 0.5 K. This may be a bit large for this very idealized case (with no moisture, an already convective initial state) but this is an error we will be satisfied with generally for boundary-layer potential temperature. Given this tolerance to error (indicated by the dashed horizontal grey line), the metric does not provide much constraint on the model behavior and the entire initial parameter space is kept (c.f. Table 2). Note that this tolerance to error is much larger than the uncertainty around the LES ( $\sigma_{r,f} = 0.075$  K) and the emulator (this uncertainty varies across the values of the parameters; it is quantified here as the mean of the standard deviation for all the points of the dataset during the LOO experiment. For wave 1 and the first metric, it is 0.042 K). Section 4.3 details the effect of a reduced tolerance to error.

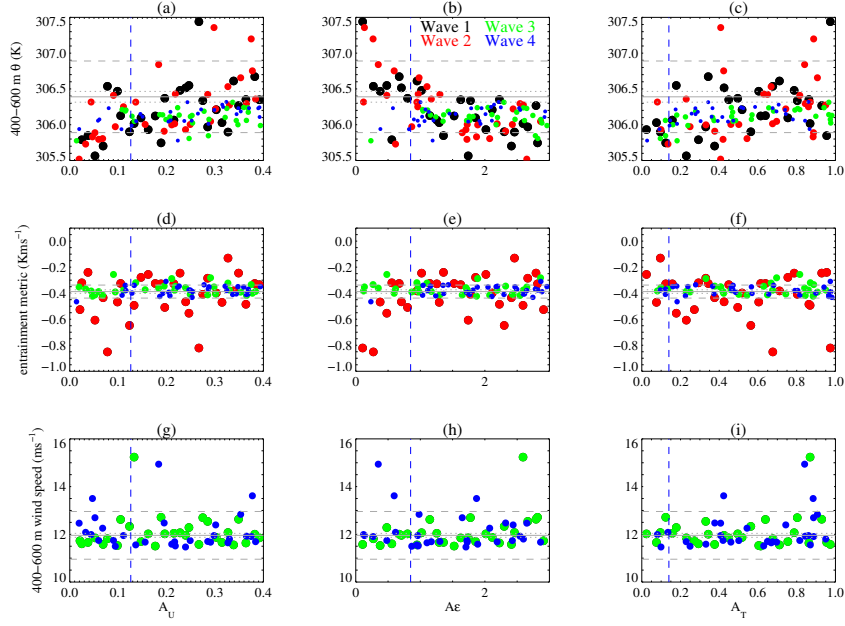
A second wave is realized, with 30 runs sampling the NROY space of the first wave (the previous 30 SCM runs could also have been used for efficiency), which is in fact the entire initial parameter space as the first metric did not constrain the parameter space. Two metrics are computed from those 30 runs: the potential temperature averaged between 400 m and 600 m as in the first wave and the entrainment metric,  $A$ , quantifying the overshoot of the boundary layer relative to the initial profile as defined in Ayotte et al. (1996).  $A$  is computed as:

$$A = \frac{\int_{zi(t_0)}^H (\theta(z, t_f) - \theta(z, t_0)) dz}{t_f - t_0} = \frac{\int_0^H (\max(\theta(z, t_f) - \theta(z, t_0), 0)) dz}{t_f - t_0}$$

**Table 2.** Description of the model discrepancy (Disc.) of the given metric (indicated in the 2<sup>nd</sup>, 3<sup>rd</sup> and 4<sup>th</sup> columns), the Cutoff, threshold used for implausibility (5<sup>th</sup> column) and the Not-Ruled-out-Yet Space (fraction in % of initial space of parameters, 6<sup>th</sup> column) for each metric (7<sup>th</sup> column) for each Experiment and wave.

$N^o$ Expt $N^o$ Wave	$\sigma_{d,\theta_{BL}}$ [K]	$\sigma_{d,Ay\theta}$ [Kms <sup>-1</sup> ]	$\sigma_{d,ws_{BL}}$ [m s <sup>-1</sup> ]	Cutoff	NROY (%)
Exp1-1	0.5	-	-	3	100
Exp1-2	0.5	0.05	-	3	30
Exp1-3	0.5	0.05	1	3	23
Exp1-4	0.5	0.05	1	3	20
Exp1-5	0.5	0.05	1	3	18
Exp2-1	0.5	0.05	1	3	40
Exp2-2	0.5	0.05	1	3	38
Exp2-3	0.5	0.05	1	3	27
Exp2-4	0.5	0.05	1	3	17
Exp3-1	0.5	0.05	1	3	72
Exp3-2	0.5	0.05	1	3	32
Exp3-3	0.5	0.05	1	2.5	22
Exp3-4	0.5	0.05	1	2.	15
Exp4-1	0.5	0.05	1	3	25
Exp4-2	0.5	0.05	1	3	19
Exp5-1	0.25	0.025	0.5	3	32
Exp6-1	0.1	0.01	0.25	3	31





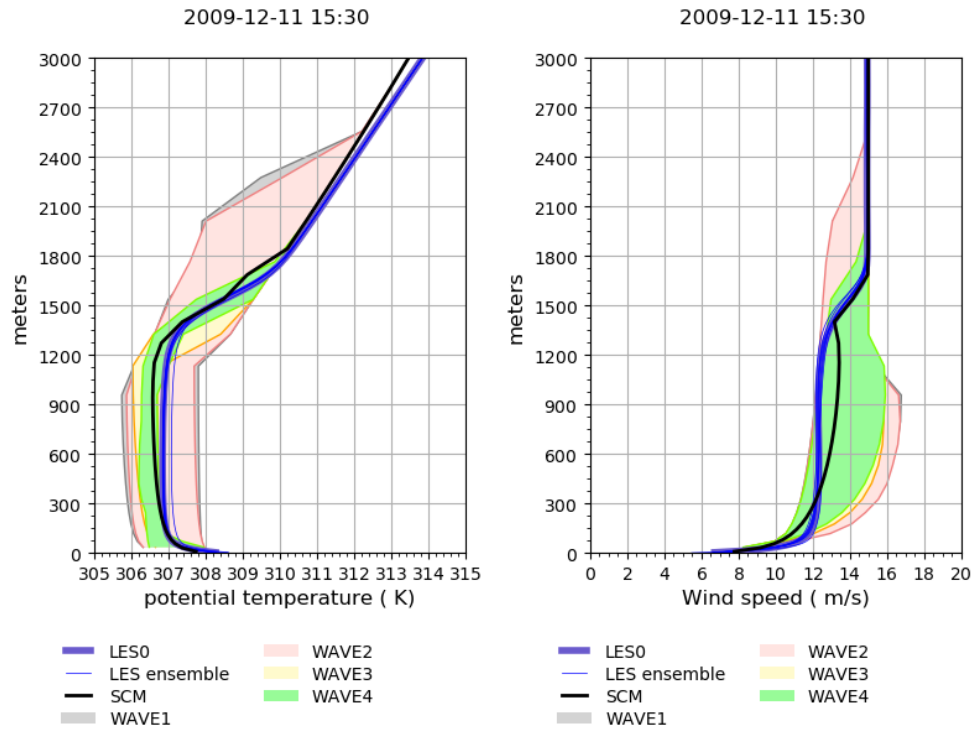
**Figure 3.** The three metrics, boundary-layer potential temperature (a–c), entrainment metric (d–f) and boundary-layer windspeed (g–i) are plotted as a function of the value of each parameter,  $A_U$  (a, d, g),  $A_\epsilon$  (b, e, h) and  $A_T$  (c, f, i). A different color is used for the different waves of Experiment 1 (black for Wave 1, red for Wave 2, green for Wave 3 and blue for Wave 4). The vertical dashed blue line corresponds to the default value of the parameter used in the model, the horizontal thin full grey line correspond to the reference metric and the dotted lines indicates the uncertainty around this reference from the different LES simulations while the dashed lines indicate the tolerance to error around the reference.

$t_0$  being the initial time,  $t_f$  the time at which the metric is computed and  $H$  the top of the model or a level largely above the boundary-layer top. This metric is less commonly used for evaluating models and it was more difficult to specify a tolerance to error, which was taken as  $0.05 \text{ K.m s}^{-1}$ . An emulator is built for each metric. The second metric is more restrictive and the NROY space is now reduced to 30% of the initial parameter space (Table 2). The obtained NROY (not shown) is not very different from the one obtained for the third wave. It excludes values of the parameters that lead to simulations with too large or too small entrainment metric as indicated by the differences between the red dots and the green ones in Fig. 3.

A third wave is realized, with 30 new SCM runs sampling the new NROY. Three metrics are computed from those 30 runs: the two previous ones plus the wind speed averaged between 400 m and 600 m. For this last metric, we fixed the tolerance to error to  $1 \text{ m s}^{-1}$ . After this third iteration, the NROY is 23% of the initial space. As shown in Fig. 4, the spread of the different simulations that sampled the parameter values reduces progressively throughout the different waves and this tool allows to discard values of parameters that induce a too deep boundary layer. The wind-speed profiles did not completely converge and this is associated to the tolerance to error, which has been fixed to  $1 \text{ m s}^{-1}$ .

The uncertainty around the LES obtained from eight different LES runs with slightly different configurations, detailed in the appendix A, is  $0.075 \text{ K}$  for  $\theta_{BL}$ ,  $0.014 \text{ K m s}^{-1}$  for  $A_\theta$  and  $0.083 \text{ m s}^{-1}$  for  $ws_{BL}$ , on the same order of magnitude of the emulator uncertainty. For the first and third metrics, the tolerance to error is much larger than the reference and emulator uncertainties while for the second metric the three uncertainties are of the same order of magnitude.

The final NROY space after the third wave is visualized in Fig. 5. This figure shows, on the upper right side, the two-dimensional density plots of the acceptable parameter space for each pair of parameters. For a given point in each sub-figure the shading indicates the percentage of the domain in the other dimensions ( $n-2$ , here only one as only three parameters are considered) that is acceptable. The metrics tend to reject preferentially low values of  $A_\epsilon$  with high values of  $A_U$  or high values of  $A_\epsilon$  with low values of  $A_U$  underlying some correlation between these two parameters. As a practical tool, those density plots provide their own type of second-order sensitivity analysis. They allow us



**Figure 4.** Vertical profile of (a) potential temperature and (b) wind speed for the last hour of the simulation with the spread of the ensemble of simulations used for the different waves indicated in different color shadings for Exp 1, the default simulation is in black, the reference LES in thick dark blue and the different elements of the LES ensemble in thin blue lines.

to see, as we move in two dimensions of the parameter space, how the shape is changing and, moreover, which combinations of parameters it is important to get right and, not usually included in a sensitivity analysis, how they need to be set in order to get sensible answers. The default values of the parameters are within the NROY space confirming that they correspond to an acceptable calibration of the turbulence scheme, given the chosen tolerance to error and the LES uncertainty. This is also confirmed by the simulations of the last wave having a behavior similar to the default simulation as shown in Fig. 4.

### 4.3 Robustness

In this subsection, we analyze the sensitivity of the results to i) the sequence of introduction of metrics (Experiment 2 uses the three metrics directly at Wave 1), ii) the threshold used to determine the NROY space (Experiment 3), iii) the number of SCM runs used to form the training dataset (Experiment 4), and, iv) the tolerance to error (Experiments 5 and 6).

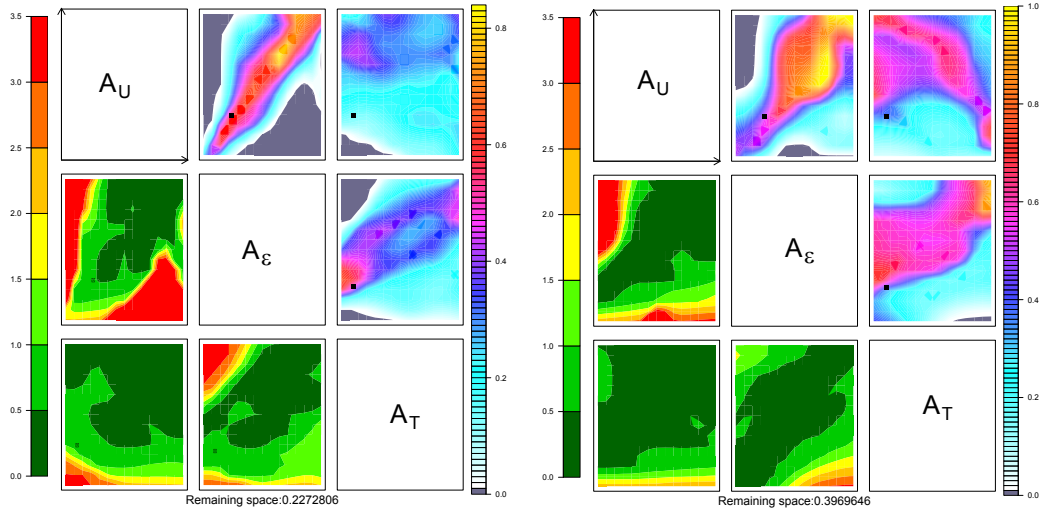
If the three metrics are introduced directly in the first wave (Experiment 2), the NROY space is similar in shape to the one obtained after three waves (see Table 2 and Fig. 5) although the NROY space is larger (40% against 23%). Repeating more waves with the same metrics allows to progressively converge to the same NROY space. Note that a test with only one metric but the most constraining one, namely the entrainment metric, leads to very similar result ( $NROY = 43\%$ ) for the first wave (not shown). Although not illustrated for this case, introducing the metrics one by one, is sometimes important: i/ it can allow us to give some priority among the metrics, first finding a space consistent with the first metric in which the second metric is then used as a constraint and ii/ if one metric has a strong non-linear behavior reducing the initial parameter spaces with other metrics may increase the capacity of the emulator to reproduce the metric behavior. These results also indicate that adding a new metric in the core of the process does not alter the selection, allowing us to add supplementary metrics if one realizes that some behavior of the SCM is not constrained enough, a fundamental aspect of history matching. Defining when to stop the iteration is not easy. We recommend to stop iterations when the NROY stops to significantly decrease. At this stage, one can reduce the cutoff used to define the implausibility and re-iterate with this new cutoff. This is illustrated with more detail in part 2. Here, Table 2 shows that a NROY of 18% is ob-

tained after Wave 5 for Experiment 1, Wave 4 for Experiment 2 or Wave 2 for Experiment 4. We can assume that for this cutoff the convergence is reached at those waves.

In Experiment 3, we first realize two waves as in Experiment 2 and then progressively reduce the threshold used to determine the NROY space from 3 to 2.5 in Wave 3 and from 2.5 to 2 in Wave 4 (see Table 2) to explore the impact of less conservative threshold (a threshold of 3 corresponds to ruling out what exceeds three times the uncertainties and keeps 95% of the probability for any unimodal probability distribution). The differences in the NROY space of the first wave with Exp2-1 indicates that 30 SCM runs are probably not enough to robustly constrain the first iteration and more iterations are needed. Then, reducing the cutoff induces a smaller NROY space but the change is not radical. This was expected from the lower left figures of Fig. 5 that show the minimum value of the implausibility for any variations of the other parameters (here, the third parameter). Indeed, the area with minimum value of  $I_f(\boldsymbol{\lambda}) > 3$  (i.e. the points that are excluded from the NROY space whatever the value of the third parameter) is very similar to the area with minimum value of  $I_f(\boldsymbol{\lambda}) > 2$ .

All of the previous experiments have been realized using a rather small training dataset of 30 SCM runs (ten times the number of parameters). Experiment 4 has tested the impact of using 90 SCM runs instead of 30 for wave 1. This experiment produces directly a smaller NROY space (NROY=25%; Fig. 6) at the first wave than obtained from 30 SCM runs (see Exp3-1 or Exp2-1 in Table 2). A compromise must be found between a larger ensemble of simulations that increases robustness but is more costly.

The sensitivity to the tolerance to error is illustrated in Table 2 and Fig. 6 with Experiments 5 and 6. When reducing the tolerance to error by a factor of two the NROY space is 32% of the initial space in Exp5-1 (using the three metrics at once, so to be compared to 40%). The NROY space (31% of the initial space) is not much reduced further when reducing the tolerance to error twice more (Exp6-1), because the tolerance to error is not anymore the limiting uncertainty. It is interesting to note that even when strongly reducing the tolerance to error, the default values for the three selected parameters are still in the NROY space validating the choice of parameter values used in the control simulation. The lower left panel of the subfigures in Fig. 5 and Fig. 6 indicates the minimum implausibility along the other dimensions of the space and as illustrated in Fig. 6,

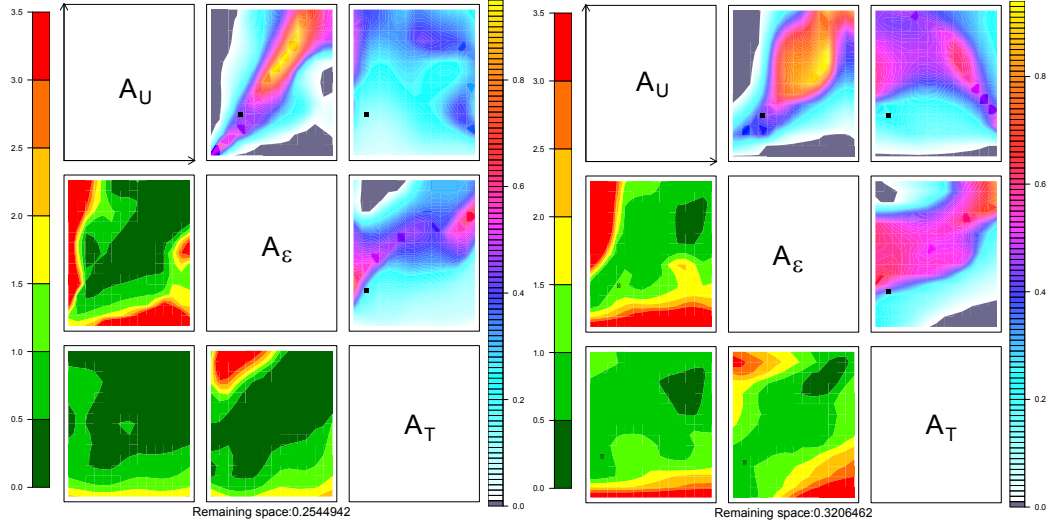


**Figure 5.** The left panel corresponds to the result of Exp1-3 and the right panel to Exp2-1. The upper right triangle contains 3 subfigures showing 2D sub-matrix. Each sub-matrix is a restriction to 2 parameters, the name of which are given in the diagonal of the main figure, and presents in colors the fraction of points with implausibility smaller than the threshold (here a value of 3). This fraction is obtained by fixing the two parameters at values of the x-axis and y-axis of the plotted location and searching the other dimensions (here the third dimension as we have only three parameters) of the parameter space. This allows to visualize in 2-D the full NROY which is 3-D here but can be n-D if n parameters are selected. The lower left triangle (with also 3 subfigures) presents the minimum value of the implausibility when all the parameters (here only one) are varied except those used as x- and y-axis. These plots are orientated the same way as those on the upper triangle, for easier visual comparison. The black dots correspond to the default values used in the model.

reducing the tolerance error (when larger than the other errors) induces a reduction of the denominator in the implausibility and therefore an increase of implausibility.

## 5 Conclusion

In this paper, we make a proposal to accelerate weather and climate model development. Our proposal tackles model development and calibration jointly. For that purpose, we have developed a tool that formalizes a process-based calibration, the *High-Tune Explorer* made available to the other modeling groups. It extensively exploits the SCM/LES comparison on a multicasest, multi-metrics basis and benefits from machine learning tech-



**Figure 6.** Same as Fig. 5 but for the sensitivity to the number of SCM runs (Experiment 4, left panel) and to the tolerance error (Experiment 5, right panel).

812 In contrast with other recent proposals to use machine learning techniques in cli-  
 813 mate modeling, we keep parameterizations as key ingredients of these models because  
 814 they summarize our current understanding of the main physical processes. This choice  
 815 is motivated in particular by the confidence needed when extrapolating the model re-  
 816 sults to a future climate.

817 The tool allows us to define the sub-domain of the parameter values for which SCM  
 818 matches LES on selected metrics for a series of cases within a given uncertainty. The ex-  
 819 ploration of the free-parameter space is facilitated using Gaussian process emulators. These  
 820 emulators, once trained on a limited number of real simulations, predict the SCM with  
 821 uncertainty for any value of the parameters in a much shorter time than required to run  
 822 the SCM. History matching using the emulator is performed iteratively to progressively  
 823 shrink the space of acceptable parameter values. This iterative approach contrasts with  
 824 the more traditional tuning strategy based on optimization, which seeks an individual  
 825 “best” value where the SCM minimizes a cost function computed for a set of given met-  
 826 rics. The latter approach strongly depends on the weights given to each metric and is  
 827 highly sensitive to the choice of metrics. By pursuing a strategy for discarding param-  
 828 eter values, we are left with a free parameter domain that is (i) consistent with the met-  
 829 rics we have chosen, (ii) can be further reduced by introducing new metrics or altering

our tolerance to model error, and (iii) does not claim a single best simulation which may be over-fitted to one or more metrics, needlessly biasing the simulation and potentially leading to less physical behavior than other choices in our not-ruled-out-yet space when the model is projected into different regimes. Our tool formalizes the consideration of the different sources of uncertainties associated to the reference, the statistical tool and the model. For the latter, we take a “tolerance to error” approach, allowing the question of whether a parameterization can match our reference as well as we think it ought to, and enabling us to understand the model’s limitations throughout the process.

In the present study, we present applications of the *High-Tune Explorer* to the SCM/LES framework, focused on the representation of the atmospheric boundary layer. We have illustrated how this tool allows us to objectively verify choices that have been made by model developers for the free-parameter values. Experimenting with the combination of the metrics with this tool allows us to clarify the importance of a given metric, the number or combination of metrics that should be used, and the possible redundancy between metrics all in an efficient way that was not possible before. The tool also enables us to include new metrics at a new iteration so that we can pursue the calibration exercise, even though one realizes an important deficiency of the model is not addressed by the previously selected metrics. Our framework allows a progressive addition of metrics, cases or a gradual reduction of the tolerance to error and is therefore very flexible.

Although this new framework is tested here for the improvement of boundary-layer processes (turbulent transport in Part I and cloud representation in Part II) by running the full atmospheric physics on one model column considering well established test cases for which LES are particularly relevant, it has much broader application. It can be used for instance to calibrate elementary pieces of parameterization (e.g., entrainment formulation) without time integration. This methodology can be easily expanded to other parameterizations as well. The key ingredient for doing this is a reliable reference with documented uncertainty. This reference could come either from a detailed modeling of the process, as done here with LES, or from observations as long as the other sources of discrepancy, as the uncertainty coming from the case definition, are documented. Proposing new relevant metrics and estimation of associated uncertainties will become valuable now that we know how to include them in the model improvement process. An effort is currently done in that direction in parallel to the work presented here, consisting in providing reference radiative transfer computations on the classical cloud test cases currently



used for parameterization development. The development of the parameterization of boundary layer and clouds based on SCM/LES comparisons focused so far on the representation of atmospheric transport and macrophysics of clouds, but the radiative transfer computations run in LES models were often not more reliable than those used in GCM, preventing the use of radiative metrics. By developing fast and accurate radiative tools that account for the full 3D radiative transfer in LES cloud scene, as proposed by Villefranque et al. (2019), we can compute many types of radiative metrics, from monochromatic, local, and directional observable to integrated energetic quantities. The use of such radiative metrics will allow us to tackle calibration of radiative parameterizations but also to better link the calibration realized at the level of the parameterizations itself with the one realized for the final full 3D model calibration, which mainly targets the radiative forcing of the atmospheric general circulation.

To conclude, the application of the *High-Tune Explorer* on SCM/LES comparisons allows us: (i) to quantify the parametric uncertainty at process level, (ii) to identify parameters which limit model performance, whatever their value, and should be replaced by a more physical parameterization (i.e. when combining different cases, it may appear that no value of a parameter is found acceptable for all cases and therefore suggests that this parameter can not be kept constant but instead should depend on environmental conditions), and (iii) to reduce the domain of acceptable values of free parameters used in the final tuning of the global model.

We show indeed in Part II how the tool applied first to SCM/LES comparisons, on a multicas e basis, can be used to reduce the range of acceptable values for the calibration of the complete 3D model configuration and considerably accelerate the resource and time consumption for this step of model development. The final 3D tuning becomes a part of the history matching process, by adding new metrics or constraints using the exact same codes.

We believe that this tool is a breakthrough for model development as it allows us to place the importance of the physical understanding of the processes at the heart of model development, based on an extensive use of the SCM/LES comparison, whilst harnessing important techniques in machine learning and uncertainty quantification. We advocate that the approach presented here leads to a well-defined strategy for calibration of the full model that may result in a significant acceleration in model improvement.

**Table A1.** List of the different LES runs of the Ayotte case used to determine the uncertainty around the reference

Name	Resolution	White noise	Turbulence	Diffusion
Name	Dx, Dz	Standard deviation (K)	length-scale	Timescale
<b>Reference</b>	50 m,nested <25 m	0.01 K	Deardorff length scale	1800 s
WhiteNoise	”	0.1 K	”	”
WhiteNoiseLL	”	0.5 K	”	”
Turb	”	”	size of the grid	”
Difshort	”	”	”	300 s
Diflong	”	”	”	7200 s
Dx	25 m, ”	”	”	”
Dz	”, nested <12.5 m	”	”	”

## Appendix A The different Large-Eddy Simulations

Different simulations have been run with Meso-NH (Lac et al., 2018), varying the resolution, domain size, turbulence formulation, intensity of the white noise introduced at the first level and initial time to trigger turbulence, activation of subgrid condensation and changes in the microphysics scheme for the cloudy cases. The Table A1 lists the different simulations of the Ayotte case used in section 4 to estimate the uncertainty associated to the reference LES and the Table A2 lists the different simulations of the ARMCU case used in section 3 to estimate the uncertainty associated to the reference LES. The reference LES is highlighted in bold.

## Appendix B ARPEGE-Climat 6.3 and its turbulence parameterization

ARPEGE-Climat 6.3 is the atmospheric component of the CNRM-CM6-1 climate model (Voldoire et al., 2019; Roehrig et al., 2020). It has 91 vertical levels, 15 of them below 1500 m. The model time step is 15 minutes. Here, we use its SCM version and focus on its representation of a clear convective boundary layer. To simulate the processes involved in the boundary layer, the model combines a turbulence scheme with a mass-

**Table A2.** List of the different LES runs of the ARMCU case used to determine the uncertainty around the reference; the names indicated in the left column are those used in the legend of Figure 2

Name	Horizontal Resolution	Vertical Resolution	Domain side	Subgrid Condensation	Microphysics	Turbulence mixing length
<b>12Dx25z25</b>	25 m	25 m	12.8 km	No	Warm (ICE3)	Deardorff
6Dx25z25	”	”	6.4 km	”	”	”
6Dx40z25	40 m	25 m	6.4 km	”	”	”
6Dx40z40	40 m	40 m	6.4 km	”	”	”
6Dx25zvar	25 m	stretched grid	6.4 km	”	”	”
6Dx100z40	100 m	40 m	6.4 km	”	”	”
25Dx100z40	100 m	40 m	25.6 km	”	”	”
51Dx100z40	100 m	40 m	51.2 km	”	”	”
6DelDx25z25	25 m	25 m	6.4 km	”	”	$(Dx * Dy * Dz)^{1/3}$
6SbgDx25z25	25 m	25 m	6.4 km	Yes	”	Deardorff
6NprDx25z25	25 m	25 m	6.4 km	No	Only saturation adjustment	”

flux scheme, thus following the Eddy-Diffusivity Mass-Flux framework (e.g. Hourdin et al., 2002; Soares et al., 2004; Siebesma et al., 2007; Pergaud et al., 2009). The mass-flux scheme represents convection in a unified way from the clear convective boundary layer regime to the shallow cumulus and deep convection regimes (Pirou et al., 2007; Gueremy, 2011). In the illustration, we aim at analyzing the importance of the values of free parameters of the turbulence scheme on the simulation of an idealized clear boundary layer. A boundary-layer-top vertical entrainment is activated in the default version of ARPEGE-Climat 6.3 (see Roehrig et al., 2020)). For the sake of simplicity of the present illustration, and also because this parameterization is weakly active in the analyzed case, it is fully deactivated. Similar results are obtained when it is activated.

The turbulence scheme is based on Cuxart et al. (2000) which provides the vertical turbulent fluxes from which the turbulent source term is derived for the prognostic variables (see more details in Roehrig et al., 2020). The scheme relies on a prognostic equation of the grid-scale turbulence kinetic energy,  $\bar{e}$ :

$$\frac{\partial \bar{e}}{\partial t} = \frac{-1}{\rho} \frac{\partial(\rho \overline{w'e'})}{\partial z} - (\overline{w'u'} \frac{\partial \bar{u}}{\partial z} + \overline{w'v'} \frac{\partial \bar{v}}{\partial z}) + \beta \overline{w'\theta'_{vl}} - \frac{\bar{e}^{3/2}}{L_\epsilon} \quad (\text{B1})$$

where the advection terms, the pressure fluctuations and the diffusion transport have been neglected.  $\rho$  is the air density,  $w$  the vertical velocity,  $u$  and  $v$  the zonal and meridional wind components,  $\beta$  is the buoyancy parameter (equal to  $\frac{g}{\theta}$  with  $g$  the gravitational constant,  $\theta$  being the potential temperature),  $\theta_{vl}$  is the liquid virtual potential temperature and  $L_\epsilon$  the dissipation length. Primes indicate fluctuations with respect to the grid-scale values indicated with overbars. The different turbulent vertical fluxes are diagnosed using  $\bar{e}$  following, for any variable  $\varphi$ :

$$\overline{w'\varphi'}(z) = -K_\varphi \frac{\partial \bar{\varphi}(z)}{\partial z} \quad (\text{B2})$$

with

$$K_\varphi = \sqrt{\bar{e}} L_m A_\varphi \Phi_\varphi \quad (\text{B3})$$

with  $\Phi_\varphi$  a stability function also computed at each altitude (for more details see Cuxart et al. (2000)) and  $A_\varphi$  a free parameter. The mixing length,  $L_m$ , is computed following Bougeault and Lacarrere (1989); it consists in computing the vertical displacement an air parcel can travel upwards and downwards with its available turbulence kinetic energy according to the thermal stratification. Also,  $L_\epsilon$  in Eq. B1 is defined by  $L_\epsilon = \frac{1}{A_\epsilon} \times L_m$  with  $A_\epsilon$  another free parameter.

## Acknowledgments

This work received funding from grant HIGH-TUNE ANR-16-CE01-0010. It was supported by the DEPHY2 project, funded by the French national program LEFE/INSU and the GDR-DEPHY. Daniel Williamson was funded by NERC grant: NE/N018486/1. Daniel Williamson and Victoria Volodina were funded by the Alan Turing Institute project Uncertainty Quantification of multi-scale and multiphysics computer models: applications to hazard and climate models as part of the grant EP/N510129/1 made to the Alan Turing Institute by EPSRC. The authors would like to thank Ignacio Lopez-Gomez and an anonymous reviewer for their constructive comments. We also thank S Barbier, T Costablos, S Nicolau and S Richet for their work during a short internship on that subject. Beyond the presentation of a new approach that we think could constitute a break through in climate model improvement, we intend to provide a tool for the climate community. All the programs, scripts and reference LES are publicly available via a Subversion through "svn checkout <http://svn.lmd.jussieu.fr/HighTune>"; a fixed version of this code is provided under <http://doi.org/10.14768/70efa07b-afe3-43a4-8334-050354f9deac>. Note, however, that this tool is a new research tool, and, as such, is still evolving. The code, the SCM runs and the LES used to produce Experiment 1 is available at <http://doi.org/10.14768/29fbfe70-a8e8-41db-914c-b14be9a6f90b>.

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