We would like to thank the editor and the two anonymous reviewers, whose constructive comments helped improve the manuscript. In the following pages, we provide detailed answers to each comment. In summary, the following three main changes are applied to the manuscript:

1. We define the term ”acceptance ratio” better.
2. We describe the initial sampling.
3. We better explain the non-intuitive result concerning the higher $P$-value cases.

To aid the reading, the original comments by the reviewers are displayed in black, while our replies are both indented and blue. Line numbers in the reviewers comments refers to the original submission, while line numbers in our replies refers to the revised manuscript.
The authors present a sampling method to select parameters leading to plausible model results. The
manuscript has been revised by 2 independent reviewers. All reviewers consider the study of (poten-
tial) interest for HESS after moderate revisions. All reviewers provide detailed comments and sug-
gestions on diverse aspects of the manuscript. One aspect that the authors should consider during the
revision of their work is to further elucidate the metric used for the global sensitivity analysis per-
formed, GSA. Although Sensitivity is an intuitive concept, a variety of approaches/metrics have been
proposed in the literature. Since each metric focuses on a different property of the model response(s),
diverse metrics may lead to different results (Razavi and Gupta 2016, Water Resources Research, 51, 5,
doi: 10.1002/2014WR016527; Dell’Oca et al. 2017, Hydrology and Earth System Science, 21,12, doi:

We would like to thank the editor for taking time to handle our manuscript. As for our choice
of GSA-metric, the editor is right that we did not discuss this point in the manuscript. The
suggested articles are highly relevant for this purpose, and we have added them together with
a brief discussion on the definition of the used GSA-metric. However, the main purpose of this
manuscript is the development of the selection method within the sampling scheme, and in the
context of the paper the GSA-analysis itself is merely one way of comparing the results of the
sampling. The following text has been added (lines 50-57):

"It should be noted that there are different global sensitivity methods with different metrics that
may give different results (e.g. Razavi and Gupta, 2015; Dell’Oca et al., 2017). In principle,
nothing speaks against computing another global-sensitivity metric for the sample selected by
our active-subspace based sampling scheme, as long as computing the metric is based on a
random sample. For practical reasons and for a direct comparison with our previous work, we
use the activity score in the present study. For the interested reader, a longer discussion about
the current metric in relation to the specific application is given by Erdal and Cirpka (2019),
and more general discussions have been presented by Saltelli et al. (2008); Song et al. (2015);
Pianosi et al. (2016), among others.”
Anonymous Referee #1

We thank the reviewer for taking the time to review our manuscript and providing constructive comments. In the following we address all individual comments one-by-one.

Content Comments

1. Page 5, Figure 2 - Consider labeling the red line “Behavioral Limit Line” for clarity. Can one assume the point has to be above the limit line to be considered acceptable behavior? Could Figure 2 be moved so that it is after Line 115?
   We use the improved label suggested by the reviewer. The figure is also moved as suggested, but the final typesetting is not in our control. The reviewer is correct: a point has to be above the limit line to be considered acceptable. This is also further highlighted in the figure caption. Figure with new caption is found between lines 136-137 on page 6 in the revised manuscript.

2. Page 5, Line 105 – Where does the active subspace come from that the initial candidate parameter sets (say, the first 1-99) are projected onto? Line 113 states that the active subspace is recalculated after adding 100 state-1 accepted parameter sets—but, how do you start?
   We thank the reviewer for highlighting this point, as the information is clearly missing. The initial active subspace is created from a random set of 50 parameter sets drawn from the unconditional prior using Latin Hypercube sampling. The following text is added to the manuscript (lines 115-117):
   "As in the original sampling scheme, we start with a set of 50 candidate parameter sets, sampled using a Latin Hypercube setup, which are per definition directly stage-1 accepted.”

3. Page 5, Line 106 – Can you provide any insight about how the values/criteria (e.g., 5 closest neighbors plus 1% radius) were selected for this work that would be beneficial for another researcher trying to implement this method?
   As the reviewer correctly points out, this is an important aspect of the sampling scheme. In our case, the values were chosen based on prior tests. However, the results were not highly dependent on the choice. The 1% was chosen just to ensure that the 5 neighbors are not so close to the candidate point that relevant uncertainties are neglected. We would consider both values to be applicable also for other model setups. In the text, the following is added (lines 123-125): ”The number of neighbors selected and the radius of the ellipse are tuning parameters, here chosen based on a few prior tests. However, we believe they are applicable also for other applications, at the very least as good starting points.”

4. Page 6, Line 121 – Is the “acceptance ratio” the ratio of candidates that are stage-1 accepted to the total number of candidate parameter sets (stage-1 accepted + rejected)? Or, is the “acceptance ratio” the ratio of candidates that are stage-1 accepted to those that are stage-2 accepted (i.e., the amount of pre-accepted candidates that become accepted). This clarification would also help interpret Figure 3.
   A very relevant comment that was not clearly explained. Alternative number 2 is what we meant (the ratio of candidates that are stage-1 accepted to those that are stage-2 accepted). In the text, the following is added (new text within ”) (line 138): ”... shows the acceptance ratios ’(number of stage-2 accepted samples divided by the number of stage-1 accepted samples)’ or the original sampling scheme ...”

5. Page 6, Line 121 and 136 – Intuitively, I am struggling to understand why P=0.75 is the fastest when it should, in my mind, be the most difficult to achieve. And, along those lines, why P=0.75 sampling results in a significantly different distribution from the unbiased pure Monte-Carlo scheme. Do you have any insight into why this is occurring?
We understand the reviewers difficulty, indeed one would think that the sampling scheme which requires the highest number of correct neighbours would be most difficult and therefore most correct. This is also true when we look at stage-1 acceptances only. Here, \( P=0.75 \) results in many more discharged candidate points, and of the stage-1 accepted ones, many are also stage-2 accepted (high ratio in Figure 3). The drawback, however, is that this sampling behavior effectively avoids sampling the boundaries between the behavioral and non-behavioral parameter space (it samples only ”safe” parameter sets). This leads to a poor match when comparing to the pure Monte Carlo sampling (which samples everything and one uses stage-2 acceptance).

In the text, the following is added (lines 164-168): "While it may seem counter-intuitive that the highest \( P \)-values gets the highest acceptance ratio and the poorest match of the marginal distributions, it is worth noting that a higher \( P \)-value means that the requirement for stage-1 acceptance is higher. Hence, at high \( P \)-values we only sample the interior of the behavioral parameter space and avoid the boundaries where the behavioral status of a candidate parameter set is more uncertain. This results in the bias clearly seen in Figure 4."

5a. Furthermore, do you think the \( P \) value selected is dependent on the model/application? Based on your experience, is the exercise of comparing different \( P \) values and selecting one necessary for another researcher trying to implement this method, or do you think the \( P=0.55 \) scheme is broadly applicable?

In this work, \( =0.55 \) has been shown to be the best comprise between efficiency and accuracy, while also the \( P=0.15 \) case could be considered a good choice. We believe that either of these two, or a value in between is generally applicable for any sampling scheme, at least as a starting point.

The following text is added to the text (new text within ”) (lines 176-177): "... captured by the faster sampling schemes may be an acceptable trade-off between speed and accuracy, depending on the individual application. "Based on the experience gained within this project, a recommended starting \( P \)-value for our presented sampling scheme is \( P=0.55.\)"

**Grammar Comments**

1. Page 4, Line 67 – Line states that the model considers 6 observations, but there are only 5 listed below this sentence. Should 6 be changed to 5?
2. Page 4, Line 67 – Consider revising the sentence to state “...observations that define acceptable behavioral performance...”
3. Page 4, Lines 69-73 – Make the list style consistent in regard to the period placement at the end of each list item (or remove them all).
4. Page 4-5, Lines 87-93 – Add period after list item number 4.
5. Page 6, Lines 109 and 111 – Remove hyphen between “parameter-set”.
6. Page 6, Line 125 – Present the acceptance ratio at 0.005 (not a percent) since the acceptance ratios are shown as decimal values on the y-axis of Figure 3.

We thank the reviewer for carefully reading our manuscript and suggesting grammatical correction. All suggestions are followed in the revised manuscript.
The current work presents a sampling strategy for those cases when some values of the investigated model output(s) are classified as unfeasible/unacceptable and the corresponding parameters sets are labelled as non-behavioral. A key step is the transformation from the original N-parameters space into the space spanned by the n-most relevant eigenvectors (here two are considered, i.e., n = 2). Then, in this reduced dimensional space an active region is identified (i.e., the active subspace) is identified (see Fig. 2 where all the space above the red line is the active subspace). Then a set of parameters is chosen to be behavioral or not (i.e., the associated output(s) belongs to the active subspace or not) in a two stages approach: (1) a surrogate model of dimension n in the space spanned by the n-most influential eigenvector is built and then used to check if a parameter set is behavioral or not, as a ‘first approximation’; (2) if a parameters set passes stage-1 the full model is run for that parameters set a second check on being or not behavioral is done. Then only stage-2 parameter sets are retained for successive analysis. The main gains here are due to the reduction of the dimension (from N to n) and the use of a surrogate model in the n-dimensional space to skim those parameters sets that are not behavioral. The improvement/modifications proposed in the current work are during stage-1, where an additional constrain is added: a parameters set passes stage-1, if in its neighborhood there is a certain fraction P of parameter sets that have already passed stage-2. The paper is of interest and well written. There are some unclear (at least to me) points which I would like to be addressed before publication, hoping for a more clear and more accessible work after revision.

We would like to thank the reviewer for her/his positive view on our work. All comments are addressed below.

Comment 1. In both approaches, after 100 parameters sets passed stage-1 the eigen-vector decomposition is re-done, and so the surrogate in the n-space dimensions is built again. My understanding is that the output(s) values associated with these 100 samples are obtained through the n-dimensional surrogate model (before adding the 100 samples), right? If this is the case, isn’t there the risk of ‘guiding’/’move’/’bias’ the active subspace toward the results of the surrogate model? For example, in Fig. 2 the new extra 100 stage-1 accepted points will all falls along the purple curve (along its branch above the red line). This could be an issue if the surrogate is doing a poor job. Am I wrong? Why not use stage-2 accepted sample (even though they require full model runs) to update the eigenvectors/eigenvalues? This will avoid the issues associated with a possibly poor surrogate modelling.

There seems to be a slight misunderstanding which may require a better explanation from our side. The reviewer is right that all surrogate-model samples will lie on the purple line in Fig.2 (or, in reality a surface in 2-D). However, this information is only used to compare against the user defined limit (red line in Fig.2) to decide if the parameter set is to be run in the full model or not. If the surrogate model is not doing a good job, the new points will be far away from the purple line. 100 of these full flow model runs are required before the update is performed. Hence, all samples used to train the active subspace and the surrogate model are full model runs. We think this might be part of the confusion and we stress this point much clearer in the revised manuscript (see changes applied below).

Using just the stage-2 accepted samples would not be very beneficial for our purpose, which is to explore the full behavioral parameter space, since the surrogate model would only be good within the behavioral space, but rather poor at the boundary. This point, however, has well been discussed in our preceding HESS publication (Erdal & Cirpka, 2019), on which the present technical note is based. In order to keep the technical note brief, we avoid to discuss it here again.

To increase the clarity of the manuscript we add the following:

Description of surrogate model (lines 92-94): "Also, as the surrogate model is only used as a preselection filter, all results and the training of the surrogate model are based exclusively on full-flow model simulations.”

Sampling scheme point 4 (line 106): "Hence, the surrogate-model is based on all currently available full-flow model simulations.”
Comment 2. How is the algorithm initialized? Which is the size of the sample to build the first n-dimensional subspace? How is relevant? For example, in Fig. 2 there are previously analyzed parameters samples/output, they should come from a set of full model runs (then they are updated after 100-samples pass stage-1t, see the previous comment).

As also pointed out in comment 2 from reviewer 1, this information was clearly missing. The first 50 parameter sets, which are also the first 50 full model runs, are sampled randomly from the unconditional prior using Latin Hypercube sampling. After the first 50 parameter sets are run, the first active subspace/surrogate model is built based on these runs, and then subsequently updated in 100 full model run intervals. In principle, we do not think that the method of initialization is that relevant, just as long as a reasonable coverage of the parameter space is achieved.

We have improved the manuscript by the following addition (lines 116-119): "As in the original sampling scheme, we start with a set of 50 candidate parameter sets, sampled using a Latin Hypercube setup, which are per definition directly stage-1 accepted. Hence we run the full flow model 50 times to initialize the sampling scheme. The actual number is not critical, and should be chosen with consideration to the number of unknown parameters."

Comment 3. Acceptance ratio: this the ratio between the stage-2 accepted sample and the drawn samples, right? Why is it a function of the stage-2 accepted samples (see Fig. 3)? I don’t see this aspect being used in the algorithm (both previous and current versions) at any step. I would have expected a dependence on the stage-1 accepted samples. Moreover, as P (i.e., the fractions of neighborhood accepted, at least at stage-1, samples) increases I would expect lower acceptance ratios, i.e., it becomes harder for a sample set to be accepted as a larger fraction of its neighbors have to be in the active subspace (i.e., P increases). (see also lines 116-117 that go along this line of reasoning). Please clarify.

We see the reviewers confusion, as this was probably not explained in the manuscript (see also comment 4 from reviewer 1). The acceptance ratio is the ratio between the number of full model runs that are stage-2 accepted, and the total number of full model runs (which is the same as the number of stage-1 accepted model runs). In the manuscript we did not report about the total number of drawn parameters-sets, but this number is much (much!) higher than the number of stage-1 accepted samples. Further, the number of stage-1 rejected parameter-sets is by far the largest in the high P case. This results in a collection of stage-1 accepted samples that poorly explores the behavioral parameter space, but where a majority of the them are stage-2 accepted. Hence, the high-P case has a high acceptance ratio.

The difference between rejected and stage-1 accepted samples and their influence on the result has been clarified in the manuscript:
1) See answers to comments 4 and 5 from reviewer 1
2) Added in the introduction of the sampling scheme (lines 91-92): "Hence, one of the beauties of the surrogate-assisted sampling is its ability to quickly discharge large quantities of non-behavioral parameter-set without running the full flow model for each one (i.e. stage-1 rejected samples)."

and

3) Added to the results section (lines 142-147): "It should be noted here that the acceptance-ratio as a statistic only shows the ratio between the runs that are behavioral after running the full-flow model (stage-2 accepted) versus the number of full-flow model runs (stage-1 accepted). This, however, does not reflect the number of stage-1 rejected parameter sets, which is not reported in this work, but is by far the largest for the higher P-values. Hence, the acceptance-ratio is a measure of computational efficiency rather than a measure of search efficiency (which here is simple Monte Carlo and, hence, comparably inefficient)."

Comment 4. Isn’t that, since P is the exact fraction (not an exceedance fraction) of good neighbors, as P increases the active subspace is updated (on top of 100 samples that pass stage-1) by favoring those regions of the active subspace that are the most distant from the threshold condition (e.g., upper left part of Fig. 3a) where it is more easy to have P high than low? Then, the n-dimensional surrogate will be
update by favoring these far-from threshold condition regions leading to a poor behavior (due to its global character) in those regions close to the threshold conditions (e.g., lower right region in Fig. 3). This is then reflected in the decreased quality of the behavioral parameters pdfs as shown in Fig. 4. Or maybe, I am just speculating too much here. It could be of interest to see how the n-dimensional surrogates evolve as a function of $P$, for example after some updates are conducted to see if there is this tendency or not.

We are not quite sure we fully understand what the reviewer means here. The $P$-value states the minimum fraction of neighbors that has to be behavioral, and is hence in our view an exceedance number. We do, however, agree with the reviewer that the higher-$P$ cases (i.e. requiring more neighbors to become stage-1 accepted) leads to a sampling that poorly samples the boundary regions (e.g. around the reg line in fig. 2). This is, as the reviewer also points out, clear from the results in figure 4, where the $P = 0.75$ case does not sample the margins of the histogram particularly well. However, we do not find any intuitive and easily understandable way of showing how the surrogate evolves, other than the clear results in Figure 4. Hence, based on this comment, no changes will be applied to the manuscript. However, if the reviewer has a clear suggestion we happy to learn about it!

Comment 5. Since a surrogate model is used to mimic also the full model response (see Sec. 2.2) I would suggest to refer to this as ‘full-model-surrogate’ in order to mark the distinction with the surrogate model build in the n-dimensional space.

We see the reviewer’s point, however, we rather like to avoid confusion by not naming the GPE-surrogate-model used as our virtual truth a surrogate. We will add this information to Section 2.2 and hope it makes the nomenclature clearer (lines 73-75):

"In order to avoid confusion we would like to point out that, in this paper, the term full-flow model means the GPE-model, while the term surrogate model is, outside of this paragraph, exclusively used for the surrogate model used to improve the sampling schemes."

7
Technical Note: Improved Sampling of Behavioral Subsurface Flow Model Parameters Using Active Subspaces

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**Abstract.** In global sensitivity analysis and ensemble-based model calibration it is essential to create a large enough sample of model simulations with different parameters, which all yield plausible model results. This can be difficult if a-priori plausible parameter combinations frequently yield non-behavioral model results. In a previous study (Erdal and Cirpka, 2019), we developed and tested a parameter-sampling scheme based on active subspace decomposition. While in principle this scheme worked well, it still implied testing a substantial fraction of parameter combinations that ultimately had to be discarded because of implausible model results. This technical note presents an improved sampling scheme and illustrates its simplicity and efficiency by a small test case. The new sampling scheme can be tuned to either outperform the original implementation by improving the sampling efficiency while maintaining the accuracy of the result, or by improving the accuracy of the result while maintaining the sampling efficiency.

1 Introduction

Global sensitivity analysis (e.g., Saltelli et al., 2004, 2008) is an established technique for quantifying the importance of uncertain parameters of a model. It has also gained popularity within hydrological sciences, with many different methods to choose from (e.g., Mishra et al., 2009; Song et al., 2015; Pianosi et al., 2016). An increasingly popular global-sensitivity approach is the method of active subspaces (e.g. Constantine et al., 2014; Constantine and Diaz, 2017). While been designed for engineering applications (e.g. Constantine et al., 2015a, b; Hu et al., 2016; Glaws et al., 2017; Constantine and Doostan, 2017; Hu et al., 2017; Grey and Constantine, 2018; Li et al., 2019), it has recently been used with good performance in hydrology (e.g., Gilbert et al., 2016; Jefferson et al., 2015, 2017; Teixeira Parente et al., 2019), including a recent study of ourselves (Erdal and Cirpka, 2019).

A key issue when conducting a global sensitivity analysis, is the requirement of a large enough sample of model simulations with parameters ranging over the full parameter space. Simulations showing unrealistic behavior (e.g., wells or rivers running dry in the model, while they in reality always have water) should be removed from the sample. Already in moderately complex models this may result in many model trials that must be discarded on the level of a plausibility check. This leads to the contradictory requirements of sampling the entire space of parameters defined by preset wide margins to capture the entire distribution while exploring only the part of the parameter space yielding plausible results. One way of easing the computational
burden, is to make use of a simpler model (i.e. surrogate/proxy/emulator model), discussed, e.g., in the comprehensive reviews of Ratto et al. (2012), Razavi et al. (2012), Asher et al. (2015), and Rajabi (2019). A common sampling approach is to use a two-stage acceptance sampling scheme, in which a candidate parameter set is first tested with the surrogate model, and only if the surrogate model predicts the parameter set to be behavioral, it is applied in the full model. This idea has been applied to groundwater modelling by Cui et al. (2011), Laloy et al. (2013), and the authors of the current study (Erdal and Cirpka, 2019). In the latter study, we used a response surface fitted to the first two active subspaces as the surrogate model in a sampling scheme for a subsurface catchment-scale flow model. The scope of the current technical note is to present an improvement of this scheme and compare it to the original one.

2 Methods

In the following subsections we briefly describe the active-subspace method and the base flow model. More details are given by Erdal and Cirpka (2019).

2.1 Active Subspaces

In this section we briefly repeat the basic derivation of active subspaces for a generic function \( f(\tilde{x}) \), in which \( \tilde{x} \) is the vector of scaled parameters \( x \) with a scaling to the range between 0 and 1. An active subspace is defined by the eigenvectors of the following matrix \( C \), computed from the partial derivatives of \( f \) with respect to \( \tilde{x} \), evaluated over the entire parameter space (Constantine et al., 2014), here shown with its eigen-decomposition and Monte Carlo approximation (Constantine et al., 2016; Constantine and Diaz, 2017):

\[
W \Lambda W^{-1} = C = \int \nabla f(\tilde{x}) \otimes \nabla f(\tilde{x}) \rho(\tilde{x}) d\tilde{x} \approx \frac{1}{M} \sum_{i=1}^{M} \nabla f(\tilde{x}_i) \otimes \nabla f(\tilde{x}_i)
\]  

(1)

in which \( \otimes \) denotes the matrix product, \( \rho \) is a probability density function, the integration is performed over the entire parameter space, \( W \) is the matrix of eigenvectors, \( \Lambda \) is the diagonal matrix of the corresponding eigenvalues, and \( M \) is the number of samples used. The \( n \)-dimensional active subspace is spanned by the eigenvectors with the \( n \) highest eigenvalues. In our application, we use \( n = 2 \) as we could detect very little improvement with higher numbers.

In a global sensitivity analysis using active subspaces, the activity score \( a_i \) of parameter \( i \) is defined by:

\[
a_i = \sum_{j=1}^{n} \lambda_j w_{i,j}^2.
\]  

(2)

in which \( \lambda_j \) is the \( j \)-th eigenvalue and \( w_{i,j} \) the element relating to parameter \( i \) in the \( j \)-th eigenvector. In the following, we consider the square root of the activity score to obtain a quantity that has the same unit as the target variable \( f \). It should be noted that there are different global sensitivity methods with different metrics that may give different results.
(e.g., Razavi and Gupta, 2015; Dell’Oca et al., 2017). In principle, nothing speaks against computing another global-sensitivity metric for the sample selected by our active-subspace based sampling scheme, as long as computing the metric is based on a random sample. For practical reasons and for a direct comparison with our previous work, we use the activity score in the present study. For the interested reader, a longer discussion about the current metric in relation to the specific application is given by Erdal and Cirpka (2019), and more general discussions have been presented by Saltelli et al. (2008); Song et al. (2015); Pianosi et al., among others.

### 2.2 Model Application

In our application we consider a model of the small Käsbach catchment in south-west Germany. The model has 32 unknown parameters, including material properties, boundary-condition values, and geometrical parameters of subsurface zones. Originally, Erdal and Cirpka (2019) simulated subsurface flow in the domain using the model-software HydroGeoSphere (Aquanty Inc., 2015), which solves the 3-D Richards-equation, here using the Mualem-van-Genuchten (Van Genuchten, 1980) parameterization for unsaturated flow. Figure 1 illustrates the model domain. Details, including the governing equations, are given by in Erdal and Cirpka (2019).

![Figure 1](image.png)

**Figure 1.** Illustration of the model domain. Left: shape of the domain and topography; right: example of a geological realization.
In a related study, we constructed a surrogate model using Gaussian Process Emulation (GPE) from roughly 4,000 parameter sets. In the GPE model, the model response \( f(\tilde{x}_i) \) at the scaled parameter location \( x_i \) is constructed by interpolation from the existing set of parameter realizations using kriging in parameter space with optimized statistical parameters. The GPE-model is constructed with the Small Toolbox for Kriging (Bect et al., 2017). In the present work, we use the GPE-model instead of the full HydroGeoSphere flow model as our virtually true model response. The prime reason for this is that we can perform pure Monte Carlo sampling of behavioral parameter sets with the GPE model, requiring about 600,000 model evaluations to create a set of 3,000 behavioral parameter-sets, which would be unfeasible with the original HydroGeoSphere model. That is, we use a surrogate model (the GPE model) to judge the performance of other surrogate models (based on active-subspace decomposition) in creating ensembles of plausible parameter sets. In order to avoid confusion we would like to point out that, in this paper, the term full-flow model means the GPE-model, while the term surrogate model is, outside of this paragraph, exclusively used for the surrogate model used to improve the sampling schemes.

Like in our prior work (Erdal and Cirpka, 2019), the model considers 6 observations defining a 5 observations that define acceptable behavioral performance (for locations see Figure 1):

- Limited Flooding: maximum of \( 2 \times 10^{-3} \) m\(^3\)/s of water leaving the domain on the top but outside of the streams.
- Division of water: between 25-60% of incoming recharge leaves the domain via the streams.
- Gage C: minimum flow of \( 5 \times 10^{-3} \) m\(^3\)/s.
- Stream A: maximum flow \( 3 \times 10^{-3} \) m\(^3\)/s.
- Stream B: minimum flow \( 5 \times 10^{-6} \) m\(^3\)/s.

With the aim of keeping this technical note rather concise, we will not discuss individual parameters or their meaning in the model. To this end, we address all parameters by a parameter index (1-32) instead of a name, and the resulting histograms refer to the the scaled parameters, ranging from 0 to 1.

### 2.3 Sampling Schemes using Active-Subspace Decomposition

The basic idea of using a surrogate-assisted sampling scheme is to use the (very fast) surrogate model to first evaluate a candidate parameter-set. If the surrogate model predicts the parameter set to be behavioral, it is stage-1 accepted and will be ran with the full model. If accepted also after running the full model, a parameter-set is stage-2 accepted. Only the stage-2 accepted parameter sets are used in the global sensitivity analysis, whereas the stage-1 accepted ones are used to improve the surrogate model. Hence, one of the beauties of the surrogate-assisted sampling is its ability to quickly discharge large quantities of non-behavioral parameter-set without running the full flow model for each one (i.e. stage-1 rejected samples). Also, as the surrogate model is only used as a preselection filter, all results and the training of the surrogate model are based exclusively on full-flow model simulations.

For each observation considered, we need to perform an active-subspace decomposition. In our our previous work (Erdal and Cirpka, 2019), a decision on whether to accept or reject a parameter set is made in the following way:
1. A third-order polynomial surface is fitted in the active subspace spanned by the two major active variables.

2. These polynomial surfaces are used to predict the observations of a candidate parameter-set.

3a If all predicted observations are acceptable, the candidate is stage-1 accepted.

3b If any predicted observation is between the acceptance point and a user-defined outer point, we assign a probability of being stage-1 accepted by linear interpolation between 0 (at the outer point) and 1 (at the acceptance point), draw a random number from a uniform distribution, and stage-1 accept the parameter set if the assigned probability is larger than the random number.

3c If any predicted observation is outside of the outer point, we reject the sample, draw a new candidate, and return to (2).

4. After adding 100 stage-1 accepted parameter sets, we recalculate the active subspace using all stage-1 accepted parameter sets collected to this point. Hence, the surrogate-model is based on all currently available full-flow model simulations.

Two critical points can be seen with this scheme. First, the polynomial surface is fitted through all stage-1 accepted points across the entire parameter space. However, locally, where we wish to make a prediction, it could still be strongly biased. Second, the user needs to prescribe the outer-points, which should not only cover our uncertainty about the acceptance point, but also implicitly addresses the error by using the active-subspace decomposition. As we project 32 dimensions to two, the potential for an imperfect decomposition is rather high (that is, two close points in active subspace may have different behavioral status). As we have no rigorous and yet simple method to address this uncertainty, the choice of the outer point becomes fairly subjective.

Illustration of the two active-subspace sampling schemes, shown for a 1-D test. The right plot shows a zoom in into the left plot. Blue dots: previously analyzed points; magenta line: fitted polynomial surrogate model; red dot: candidate parameter in active subspace (x-value) with the assigned polynomial prediction (y-value) of the original sampling scheme; green dots: neighbors considered in the new scheme, which are chosen exclusively by the active variable value; red line: acceptance criterion.

To overcome these issues, we here suggest a modified sampling scheme, with fewer tuning parameters and less sensitivity to local biases. As with the original scheme, we require one active subspace decomposition per observation and use the first two active variables to create the two-dimensional active subspace. The process is as in the original sampling scheme, we start with a set of 50 candidate parameter sets, sampled using a Latin Hypercube setup, which are per definition directly stage-1 accepted. Hence we run the full flow model 50 times to initialize the sampling scheme. The actual number is not critical, and should be chosen with consideration to the number of unknown parameters. The new sampling scheme then proceeds as follows:
1. The candidate parameter set is projected into the active subspace.

2. The closest neighbors in the active subspace are sought. In this work we use the 5 closest neighbors plus all neighbors that fall within an ellipse around the candidate point that has a radius of 1% of the total range of each active subspace, in each of the two dimensions. The number of neighbors selected and the radius of the ellipse are tuning parameters, here chosen based on a few prior tests. However, we believe they are applicable also for other applications, at the very least as good starting points.

3. For each observation, a candidate parameter set is pre-accepted if a certain ratio ($P$) of its neighbours are behavioral (i.e., stage-2 accepted).

4. The candidate parameter set is stage-1 accepted if it was pre-accepted for all observations, otherwise it is rejected.

5. If rejected, draw a new candidate parameter set and return to (1).

Like before, we recalculate the active subspace after adding 100 stage-1 accepted parameter sets. The two approaches are illustrated in Figure 2, although just for a 1-D illustrative example. As can be seen in the figure, the original sampling scheme suggests that the candidate is behavioral (red dot is above the red line). With the new sampling scheme, on the other hand, it becomes a matter of the $P$-value chosen. At $P = 0.15$ and $P = 0.55$, the candidate would have been stage-1 accepted (60% of the green dots are behavioral), while at $P = 0.75$ the candidate would have been rejected. In this work, we consider the ratios $P = 0.15$, $P = 0.55$ and $P = 0.75$, and compare the performance of the sampling scheme with that used in the previous study (Erdal and Cirpka, 2019).

3 Results and Discussion

Figure 3 shows the acceptance ratios (number of stage-2 accepted samples divided by the number of stage-1 accepted samples) for the original sampling scheme and the new sampling scheme with three different $P$-values, together with a pure Monte-Carlo sampler without preselection, applied to the Käsbach GPE-model with 32 parameters. As can be seen, the new scheme with $P = 0.75$ is the fastest, while the original scheme and the new scheme with $P = 0.15$ show rather comparable behavior with lower acceptance rates. For comparison, the pure Monte Carlo sampling has an acceptance ratio of $\approx 0.5\% - 0.005$. It should be noted here that the acceptance-ratio as a statistic only shows the ratio between the runs that are behavioral after running the full-flow model (stage-2 accepted) versus the number of full-flow model runs (stage-1 accepted). This, however, does not reflect the number of stage-1 rejected parameter sets, which is not reported in this work, but is by far the largest for the higher $P$-values. Hence, the acceptance-ratio is a measure of computational efficiency rather than a measure of search efficiency (which here is simple Monte Carlo and, hence, comparably inefficient).

While high acceptance rates are favorable in light of computational efficiency, we also want to avoid introducing a bias by the preselection scheme. We evaluate such bias, by considering the marginal parameter distributions of the stage-2 accepted...
Figure 2. Illustration of the two active-subspace sampling schemes, shown for a 1-D test. The right plot shows a zoom-in into the left plot. Blue dots: previously analyzed points; magenta line: fitted polynomial surrogate model; red dot: candidate parameter in active subspace (x-value) with the assigned polynomial prediction (y-value) of the original sampling scheme; green dots: neighbors considered in the new scheme, which are chosen exclusively by the active-variable value; red line: Behavioral Limit Line. Here, points above the red line are considered to have acceptable behavior.

Figure 3. Acceptance ratios of the different sampling schemes, plotted as a function of the number of stage-2 accepted samples.
samples, which should agree with the distribution obtained by the (inefficient) pure Monte-Carlo sampler. Figure 4 shows the resulting histograms for the three parameters with the most complex marginal distributions. We quantified the agreement of the marginal distributions of the sampling schemes with preselection and the pure Monte-Carlo sampling by the Cramér–von Mises metric \( \omega^2 \):

\[
\omega^2 = \int_0^1 \left( \hat{P}_{ss}(\tilde{x}_i) - \hat{P}_{MC}(\tilde{x}_i) \right)^2 d\tilde{x}_i
\]

in which \( \hat{P}_{ss}(\tilde{x}_i) \) is the marginal cumulative probability of the scaled parameter \( \tilde{x}_i \) for a tested sampling scheme and \( \hat{P}_{MC}(\tilde{x}_i) \) is the same quantity for pure Monte-Carlo sampling. The corresponding values of \( \omega^2 \) are reported in the subplots of Figure 4.

**Figure 4.** Histograms of the three parameters with the most complicated posterior marginal distributions. Each row shows a parameter and each column a sampling scheme. Blue bars: histograms from pure Monte Carlo sampling (i.e. true distribution); brown bars: sampling schemes with preselection; numbers: Cramér–von Mises metric \( \omega^2 \) for the distance between the two distributions, here shown multiplied with 1000 for increased readability.

From the histograms in Figure 4 and the values of the Cramér–von Mises metric \( \omega^2 \) it becomes obvious that the fast new sampling with \( P = 0.75 \) results in marginal distributions that significantly differ from those of the unbiased pure Monte-Carlo scheme. The new scheme with \( P = 0.55 \) results in marginal distributions that are comparable to those of the original scheme,
but that have been achieved by a sampling scheme with twice the acceptance rate and thus half the computational effort. By contrast, the new scheme with \( P = 0.15 \), which caused a computational effort similar to the original scheme, resulted in a marginal posterior distribution that is very similar to that obtained by pure Monte-Carlo sampling. Hence, we can conclude that the proposed sampling scheme is superior to the old one: either it has much better sampling accuracy for the same efficiency \((P = 0.15)\), or it has a much better efficiency with a very comparable accuracy \((P = 0.55)\). While it may seem counter-intuitive that the highest \( P \)-values gets the highest acceptance ratio and the poorest match of the marginal distributions, it is worth noting that a higher \( P \)-value means that the requirement for stage-1 acceptance is higher. Hence, at high \( P \)-values we only sample the interior of the behavioral parameter space and avoid the boundaries where the behavioral status of a candidate parameter set is more uncertain. This results in the bias clearly seen in Figure 4.

Figure 5. Square-root of activity scores of the 10 most influential parameters for the target variable stream flow at gage C resulting from applying the active-subspace based global sensitivity analysis to the posterior distributions using the different sampling schemes.

Figure 5 shows the square-root of the activity score for a selected target variable, computed by the active-subspace based global sensitivity analysis and using the different sampling schemes, which confirms the impression of the histograms shown in Figure 4. The pure-MC scheme and the new scheme with \( P = 0.15 \) show almost identical activity scores, while the score-patterns increasingly differ with increasing \( P \)-values. Similarly, the original sampling scheme differed in the activity scores compared to the pure-MC scheme. Nonetheless, all sampling schemes correctly identified the two most important parameters and the correct set of the ten most important parameters. That the order of the parameters within the set of the most important parameters is not captured by the faster sampling schemes may be an acceptable trade-off between speed and accuracy, de-
pending on the individual application. **Based on the experience gained within this project, a recommended starting P-value for our presented sampling scheme is P=0.55.**

In the current study, we have used Gaussian process emulation (GPE) as a proxy of the full HydroGeoSphere model, putting the question forward whether a GPE model could not also be used as surrogate model for preselection in an advanced sampling scheme. This is indeed possible, and we are currently developing such schemes, achieving acceptance ratios between 70-90%. Hence, GPE-based sampling schemes can be notably more efficient than the new scheme presented in this work. Nonetheless, we see a clear value in using the less efficient active-subspace based sampling schemes. The key word is simplicity. The full active subspace-sampling scheme is implemented in-house, and the most complicated step is likely the eigenvalue decomposition, which is a standard tool in any programming environment. Hence, we have full control over the entire selection procedure. Further, the active-subspace based sampling scheme presented here has a single tuning coefficient $P$ with an easily comprehensible meaning, and the resulting active subspace can easily be visualized for an intuitive understanding of the method. This is quite different with GPE-based methods which require choosing a covariance function in parameter space with coefficients that needs to be estimated from the current set of training data. In our application, we have 32 original parameters, requiring one variance and 32 integral scales as covariance coefficients to be estimated every time the GPE-model is re-trained. Estimating 33 covariance parameters from $O(1000)$ parameter sets is time consuming, and the integral scales in non-sensitive parameter directions are not well constrained by the data at all. Finally, to train a GPE model we need to rely on third-party codes which remain black boxes to a large extent, and usually involve a rather decent amount of work until they do what they are supposed to do. Hence, we clearly see a benefit of using the simpler active-subspace based sampling schemes even if they are computationally less efficient.

4 Conclusions

In this work we have presented an improved sampling scheme to obtain ensembles of parameter sets that lead to plausible model results. Like in the preceding study of Erdal and Cirpka (2019), the sampling scheme makes use of an active-subspace based preselection scheme that reduces the number of full model runs that need to be discarded. In contrast to the preceding method, we don’t perform a polynomial fit over the entire parameter space anymore, neither do we have to set fuzzy boundaries of the target variables to define the behavioral status. Instead, the preselection of a parameter set is simply based on the behavior of surrounding trial solutions. The new scheme outperforms the preceding one by either achieving a higher accuracy in the resulting posterior parameter distributions for the same sampling efficiency, or by having a much higher sampling efficiency for a comparable accuracy. We hence conclude that the new scheme presented here should be used instead of the original one.

**Code availability.** All own-developed codes necessary to run the Stochastic Engine used in this work are available via http://hdl.handle.net/10900.1/6a66361b-b713-4312-819b-18f82f27aa18

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