Application of all relevant feature selection for the failure analysis of parameter-induced simulation crashes in climate models -
Response to reviewers.
Wiesław Paja, Mariusz Wrzesień, Rafał Niemieć and Witold Rudnicki

We would like to thank the reviewers for the valuable comments. The response to this comments helped us to improve the manuscript in many ways. In our response we first quote entire review of Referees and then answer to issues raised by them one by one, quoting the appropriate fragments from the manuscript. The manuscript quotes are enclosed between >> signs. The modified fragments that address issues raised by the referees are marked in bold red face.

**Referee #1**

This manuscript describes a study that used machine learning methods to analyze climate simulation failure (model crash) caused by perturbations in uncertain model parameters. The dataset and the goal of analysis were the same as those of Lucas et al. (2013). However, different methods were applied, which led to the conclusion that some of the parameters deemed important by the analysis of Lucas et al. (2013) were redundant or irrelevant.

Simulation failure analysis is a relevant topic for climate model development, and a more accurate identification of important parameters is beneficial. Hence, the results of this study are potentially useful. On the other hand, I would recommend a serious revision so that the manuscript can be made more informative - and the messages more convincing - for climate model developers and users.

My main difficulty with the manuscript is that it might not have been written with geo- scientific model developers as the target audience. As such, I wonder whether the manuscript is more suited for a statistics or computer science journal.

From the perspective of a climate modeler, I think the manuscript does not provide sufficiently detailed descriptions of the methods and analysis procedure to allow many readers of GMD to reproduce the results or apply the same methods to analyze other datasets.

Comparing this study’s results to those of Lucas et al. (2013), it is worth noting that the 3 strongly relevant parameters identified in this study were listed among the top-4 important parameters by Lucas et al. The benefit of this study’s methods thus seems marginal. It would also be interesting to know what price one has to pay, in terms of algorithm complexity and computing time, in order to do the cross-validation and estimate the statistical uncertainty of the results.

**Response**

Firstly, we would like to thank the reviewer for the general opinion that the subject and results of the study may be useful for the climate modelling community.
As computer scientists developing and applying machine learning methods we are very much interested in making fellow scientist in different disciplines of science aware of the applicability of our tools and methods for their research problems. We have strived to make our article both reasonably succinct and accessible for non-specialists. Apparently, we have not achieved the second goal.

Answers to the more specific issues raised by the referee #1 are given below.

1
My main difficulty with the manuscript is that it might not have been written with geo-scientific model developers as the target audience. As such, I wonder whether the manuscript is more suited for a statistics or computer science journal.

Answer
Our intended audience is the climate model developers. We believe that the results of the study will be interesting for this community. We wanted to achieve several goals.
Firstly, we wanted to strengthen the message from the Lucas et al. that machine learning methods can be a useful tool for diagnosing problems in model development.
Additionally, we introduced methodology that is more robust and in the same time easier to apply.
In comparison with SVM used by Lucas et al. the Random Forest algorithm is much easier to apply – it gives reasonably good results with default parameters. Similarly, the Boruta algorithm simply returns a list of relevant variables with relative ranking of their importance. On the other hand, to achieve good results with SVM one has to select an appropriate kernel function and perform optimisation of the parameters.
Finally, we have improved results of the original work. We elaborate on this in the answer to the third issue raised by the referee #1.

2
From the perspective of a climate modeler, I think the manuscript does not provide sufficiently detailed descriptions of the methods and analysis procedure to allow many readers of GMD to reproduce the results or apply the same methods to analyze other datasets.

Answer
We have improved the manuscript in this respect. We have introduced the more detailed description of the tools used, extended
the description of both tests performed in the study, and added the graphical summaries of both tests. Firstly, we have included the following description of tools used in the study:

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We have used a different classification algorithm, namely Random Forest (Breiman, 2001) and instead of the sensitivity analysis we have applied the all-relevant feature selection algorithm Boruta (Kursa et al., 2010). All computations were performed in R environment for statistical modelling (R Development Core Team, 2008), using the randomForest package for classification (Liaw and Wiener, 2002) and the Boruta package for feature selection. (Kursa and Rudnicki, 2010).

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Then we have rewritten and extended description of the first test. We have also added a graphical representation of the protocol used in new Figure 1. We have explicitly named the packages and functions used in the protocol. The modified paragraph is shown below.

>>

The climate simulations dataset is highly biased towards successful completion of simulation. Only 46 cases out of 540 are failures. Such unbalanced datasets are often difficult for classification because the automatic selection of the majority class results in good, but useless, classification accuracy. In such a case no information is gained and hence one cannot perform feature selection. In the first test of the current study this problem was avoided by application of the following protocol, see Fig 1. Firstly eleven balanced subsamples of training set were constructed, each subsample consisted of all objects from minority class (failed simulations) and 1/11th of majority class (successful simulations). In order to check specificity of the feature selection each dataset was extended by contrast variables. To this end each original variable was duplicated and its values were randomly permuted between all objects. In this way a set of shadow variables that were non-informative by design was added to the original variables. Then the feature selection procedure was performed on each subsample with the help of the all-relevant feature selection algorithm, implemented in Boruta function of the Boruta package. The procedure was repeated 60 times. Altogether all relevant feature selection was performed 660 times. The number of times when the artificially constructed shadow variables were selected as important gives an estimate of the expected level of false discovery. The variables that were selected as important significantly more often than random were examined further, using different test.

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For the second test we included the information about the R package and function used to build models. We have also added the graphical summary of the protocol for the second test in new Figure 2.

The test was performed similarly to the one reported in the original study, see Figure 2. The data set was randomly split into a training set containing 360 objects and a validation set containing 180 objects. The split was performed separately for the minority and majority class, so the number of minority class objects in each training set was 32 and in the validation set it was 14. The randomForest function from the identically named R package was used to perform classification and error estimate. The procedure was repeated 30 times and results of 30 repetitions were analysed.

Finally, we discussed the selection of the value for the key parameter of the Random Forest, namely the number of trees in the ensemble:

The number of trees in the forest was set to 5000 both for feature selection and classification tasks. In both cases the number of variables examined for each split was equal to the square root of the total number of variables. In our experience these settings are fairly robust, we have examined them internally over multiple datasets (Rudnicki et al., 2015). Moreover, we have checked whether they influence results in the initial trials. The number of trees used was 10 times higher than default, to assure that importance estimate in Random Forest converge to their asymptotic values, the number of trees for classification was the same for consistency.

Comparing this study’s results to those of Lucas et al. (2013), it is worth noting that the 3 strongly relevant parameters identified in this study were listed among the top-4 important parameters by Lucas et al. The benefit of this study’s methods thus seems marginal.

Answer

Regarding the observation of the referee, that the main result is not much different from the results published by Lucas et al. and hence the value of the contribution is marginal. We believe that there are at least three reasons why the work is valuable for the climate modellers.

Firstly, the confirmation of the main results of the previous work by a different methodology is valuable by itself. We have shown, that application of more rigorous and computationally demanding
methods confirmed the importance of 6 out of 8 parameters, and we concluded that 3 out of them were non-redundant. We don’t say that the 4-th parameter is not important, but that nearly all variance in data could be explained by models built on 3 main parameters.

This result can be directly useful for modellers since it reduces the effort required to improve the simulation codes. While we don’t have experience with climate models, the general experience with software shows that the difficulty of improving the code grows combinatorially with the number of free parameters. Hence the reduction from four to three may in some cases turn problem from very hard to a reasonably hard and help to get solution quicker and with less effort.

Secondly, we have shown that some conclusions of the original work were far-fetched and not supported by data. In particular Lucas et al. analysed minute effects of different thresholds on the AUC obtained on the single split of data between training and test set. We have shown, that this effect is dwarfed by variance due to the composition of the training and test sets, and hence is irrelevant. This part of the paper shows that any conclusion drawn from application of machine learning methods must be supported by a solid cross-validation study.

Finally, we have shown a very simple methodology for establishing the importance of variables for complex and obscure phenomena. Here it was applied to analysing the influence of the selected parameters for the simulation crashes, however, it can be equally well used for analysis of the standard simulations and exploring unexpected relations between variables.

#4
It would also be interesting to know what price one has to pay, in terms of algorithm complexity and computing time, in order to do the cross-validation and estimate the statistical uncertainty of the results.

The complexity of the algorithm is not increased significantly by cross-validation, it simply requires repeating entire modelling procedure several times, using different splits of data between training set and test set. It is straightforward to implement this in a script that performs all modelling. The additional benefit is that cross-validation procedure imposes rigour on the modelling
procedure – it needs to be defined in a repetitive script. Therefore, the research is easily reproducible.

Regarding the time required for modelling. We have added the following paragraph at the end of the results section that deals with this issue:

\[
\text{A single run of the Boruta algorithm in the first test took 2 minutes on a server equipped with Intel Xeon E5620@2.4GHz CPU. The entire protocol took less than 24 hours of single CPU core. The second test is far less computationally demanding. A single run of the randomForest function takes less than 20 seconds and the same CPU, therefore, computations for the entire protocol take less than 10 minutes. This effort is negligible in comparison with the time required to run 540 simulations of the climate model itself.}
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**Referee #2.**

Summary Statement:
The main purpose of the technical note by Paja et al. is to re-evaluate the climate model failure data reported in Lucas et al. (2013). In particular, Paja et al. use a feature selection technique based on random forests, instead of sensitivity analysis, to identify parameters that influence simulation failures. Their results largely agree with those in the original paper. Lucas et al. determined that 4 parameters account for most of the variance in the failures (about 90%), which are the same four parameters identified by Paja as having the largest feature scores. Paja et al. also show that the feature scores of the less influential parameters (i.e., those ranked lower than the top 4) depend on the train/test split. Their results are reasonable and not surprising because the raw data displayed in figure 2 of Lucas et al. shows that the relationship between failures and parameter values degrades significantly going from higher to lower ranked parameters. It is less clear how much value the geoscientific community can take from the Paja et al. study because it re-evaluates an existing paper and reaches similar conclusions. I am not inclined to recommend the paper for publication in GMD as an original manuscript, but as a technical note it could suffice after addressing the items and comments listed below. I leave it to the discretion of the editor to decide if it passes this bar.

Item 1. The presentation of the material is still rough around the edges in terms of readability and language. I recommend that the authors work with someone to improve the readability.

Item 2. There is a mistake on page 5420 line 23. It should say 540 simulations, not 480 simulations. On the same line, "randomly" is probably a better word than "systematically".

Item 3. On page 5421 line 24, the authors state that the setup used by Lucas et al. "precludes estimation of statistical uncertainty". This is not strictly true, as bootstrapping estimates the distribution of failure probability due to different train/test splits and as a function of input parameter values. Even though they did not report the uncertainty in their sensitivity indices, Lucas et al. used bootstrapping for the sensitivity analysis.
Item 4. Page 5423 describes the general random forest algorithm, but doesn't provide the values used for the control parameters. One potential problem with random forests is the tradeoff between bias and variance during fitting. Can the authors comment on how they determined the values of the control parameters, whether they controlled for bias or variance, and what the impact of their choice is on the feature ranking? Item 5. As shown in figure 1, the importance scores using the Boruta algorithm have values that range from about -10 to +120. How do these translate into sensitivity indices? The latter are fractions between 0 and 1, and thus define the amount of variance explained by the parameters. Can a similar interpretation be made for the Boruta scores? Item 6. Random forests can also have difficulties with correlated features, whereas polynomial chaos expansions, by design, explicitly decompose the sensitivities into various combinations of features. The authors should make some assessment of the potential effects of correlated features on their results. Furthermore, it is important to point out that a climate model parameter may be considered to be important even if it has a low feature score by itself but is correlated with parameters having high scores. This situation is analogous to parameters in figure 10 that have relatively small nodes but large edges. Item 7. Paja et al. should also be aware that some of the co-authors of the Lucas et al paper were co-authors on a related paper that computed sensitivity information using random forest feature scores (doi:10.1002/2014JD022507).

Response
We would like to thank the referee #2 for the detailed review of our contribution. We agree that this is more a technical note than the regular article. This is why we tried to make it as brief as possible when explaining methodology. Regarding the specific issues raised by referee #2

Item 1. The presentation of the material is still rough around the edges in terms of readability and language. I recommend that the authors work with someone to improve the readability.

Answer
We have sought advice of the professional proofreader and improved language and readability. Stylistic and grammatical changes were introduced in numerous places in the manuscript.

Item 2. There is a mistake on page 5420 line 23. It should say 540 simulations, not 480 simulations. On the same line, "randomly" is probably a better word than "systematically".

Answer
Corrected

Item 3. On page 5421 line 24, the authors state that the setup used by Lucas et al. "precludes estimation of statistical uncertainty". This is not strictly true, as bootstrapping estimates the distribution of failure probability due to different train/test splits and as a function of input parameter values. Even though they did not report the
uncertainty in their sensitivity indices, Lucas et al. used bootstrapping for the sensitivity analysis.

**Answer**

After careful rereading our statement we think that indeed this statement is too general. It pertains to the analysis of the influence of the choice of particular decision function for the AUC of the predictive model, however it is not true for the sensitivity analysis. Therefore we modified manuscript in two ways. Firstly we modified the paragraph in the following way:

In the original paper the authors performed true prediction and achieved a high degree of accuracy, therefore showing the true predictive power of this approach. On the other hand, this setup **precludes estimation of statistical uncertainty for some of their findings. In particular, the discussion of the prediction accuracy in sections 4.4 and 4.5 is based on a single split of data between training and test sets and ignores possibility that effects may depend on the particular split.**

We have also modified the following paragraph, to stress that Lucas et al. have used cross-validation in the original study for sensitivity analysis.

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The results of individual trials will differ in most cases, allowing one to draw conclusions not only about mean values but also about variance and even shape of probability distribution. **Lucas et al. have used this approach for the sensitivity analysis, utilising ensembles of SVM (Vapnik, 1995) learners for classification.** Each member of the ensemble was obtained using different subsample of the training set. The classifier was then used for prediction of the simulation result for the validation set.

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We believe that in this way we describe correctly limitations of the original work by Lucas et al. The entire discussion of the prediction accuracy in sections 4.4 and 4.5 is based on the single split of 540 experiments between the training set and the test set. In particular, authors discuss how a selection of the decision criteria can influence the result and result in a prediction quality measured by AUC score to vary between 0.963 and 0.966. In this case the range of changes is 0.003. We have shown that different splits of the same data between training and test set can result in AUC that varies between 0.848 and 0.990 – the range of results is 0.142 which is nearly two orders of magnitude higher. It is possible, that using different criteria for a decision would indeed improve results for most splits or even all
splits, but the discussion in sections 4.4 and 4.5 simply ignores the effect that is nearly two orders of magnitude higher and can significantly influence the results.

Item 4. Page 5423 describes the general random forest algorithm, but doesn’t provide the values used for the control parameters. One potential problem with random forests is the tradeoff between bias and variance during fitting. Can the authors comment on how they determined the values of the control parameters, whether they controlled for bias or variance, and what the impact of their choice is on the feature ranking?

**Answer**

The balance between bias and variance in random forest can be to some extent regulated by the number of variables tried for performing the split – the \textit{mtry} parameter and by the depth of the trees. In our experience the default value, which for classification is a root square of the total number of variables, gives good compromise between bias and variance. What is more, small changes of the default value usually don’t influence the classification error. When random forest is used in Boruta, the number of variables may vary drastically between initial and final stages of the algorithm. Therefore, unless there are very good reasons to modify the default variables, the default value of mtry parameter is the best choice. Moreover, initially we have checked whether modification of the mtry parameter would give better classification results but without any apparent changes in the outcome, so we decided to use the default value also in this work.

On the other hand, we have not modified the depth of the trees. This is technically possible, but it is generally against the spirit of the Random Forest algorithm and one should avoid modifications of this parameter unless everything else fails. In our case Random Forest gave results very similar to those obtained with the help of SVM ensembles by Lucas et al. and we were quite happy with this performance.

We have introduced following modifications in the paper to account for the issues discussed above:

Firstly, we have mentioned which software tools were used:

\begin{verbatim}
All computations were performed in R environment for statistical modelling, using the randomForest package for classification (Liaw and Wiener, 2002) and the Boruta package for feature selection (Kursa and Rudnicki, 2010).
\end{verbatim}
Secondly, we have modified paragraph introducing Random Forest in the following way:

>>> Random Forest is an ensemble algorithm based on decision trees. To ensure the low correlation between elementary learners, each tree is grown using a different random subsample of the original data set. Moreover, each split in the tree is built using only a random subset of the predictor variables. The number of variables in this subset influences the balance between bias and variance for the training set. The default value for classification tasks is a square root of the total number of variables and it is usually a very robust selection. Random Forest is a robust “of the shelf” algorithm that is easily applicable to various classification and regression tasks. It has only few control parameters and usually it does not need fine tuning for the particular problem under scrutiny.

Finally, at the end of methods section we have added information of the number of trees that was used both in standalone Random Forest algorithm as well as in the Random Forest used by Boruta algorithm for feature selection:

>>> The number of trees in the forest was set to 5000 both for feature selection and classification tasks, In both cases the number of variables examined for each split was equal to the square root of the total number of variables. In our experience these settings are fairly robust, we have examined them internally over multiple datasets (Rudnicki et al., 2015). Moreover, we have checked whether they influence results in the initial trials. The number of trees used was 10 times higher than default, to assure that importance estimate in Random Forest converge to their asymptotic values, the number of trees for classification was the same for consistency.

Item 5. As shown in figure 1, the importance scores using the Boruta algorithm have values that range from about -10 to +120. How do these translate into sensitivity indices? The latter are fractions between 0 and 1, and thus define the amount of variance explained by the parameters. Can a similar interpretation be made for the Boruta scores?

**Answer**
The importance measure in Boruta is an average permutation importance obtained from multiple iterations of random forest. Importance for a single variable is related to the decrease of classification accuracy of elementary classifiers in ensemble, when information about given variable is removed. It is not directly
proportional to the amount of variance explained by this variable in the model, although there is strong correlation between these two. We have added the following note after discussion of the Figure 3 (Figure 1 in the previous version of manuscript):

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One should note, that the importance returned by Boruta is the averaged importance obtained from the underlying Random Forest algorithm. It is not directly interpretable in terms of the fraction of variance explained by given variable.
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Item 6. Random forests can also have difficulties with correlated features, whereas polynomial chaos expansions, by design, explicitly decompose the sensitivities into various combinations of features. The authors should make some assessment of the potential effects of correlated features on their results.

**Answer**
We don’t agree with this statement. Random Forest deals very well with correlated features – it can build useful models for systems with multiple correlated features. When there are multiple highly correlated features in the system Random Forest will assign similar importance to all these features. This is because importance is measured only for these base classifiers that use given feature. The sensitivity of the feature selection may go down if many correlated features are present since fewer trees uses any of these features. However, this can be circumvented by using higher number of trees – what we did in the study. This property of the Random Forest is exploited by Boruta for finding all relevant feature selection.

Item 6 – continued.
Furthermore, it is important to point out that a climate model parameter may be considered to be important even if it has a low feature score by itself but is correlated with parameters having high scores. This situation is analogous to parameters in figure 10 that have relatively small nodes but large edges.

**Answer**
We agree with this statement wholeheartedly, this is the underlying idea of the all-relevant feature selection algorithm. We identify all the variables that are more informative than random contrast variables and return all available information to the user of the algorithm. It is up to her/his domain-specific knowledge to take best advantage of the whole information.
We think that the results of two analyses diverge for less important variables, due to more non-linear character of the underlying base classifiers. Lucas et al. have used SVM that, despite using a kernel trick, may be less suited for highly non-linear and non-continuous systems than decision trees. As a simple example – the decision tree can handle simple XOR problem easily in two dimensions, whereas SVM requires additional dimension – either by direct extension of the feature space or by some kernel trick. Hence, the decision trees may find solutions that require less variables than SVM. In this sense the extra variables are useful for SVM, whereas they are not truly informative in the sense of perfect Bayesian classifier. Moreover, in our analysis the apparent importance of the variables was compared with that of the contrast variables, and only those variables that were declared important significantly more often than contrast variables were declared important. On the other hand such analysis was not performed in the original work.

We have added the following paragraph at the end of the Results and Discussion to accommodate these considerations:

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The results of the study are mostly in good agreement with the results of Lucas et al., however, importance of the variables is not identical. The most important difference is the importance of the variable V13 in both studies. This variable is more important than V14 in the SVM-based model by Lucas et al., whereas our analysis deems it relevant but redundant. However, one should note that in the first test V13 was deemed relevant in nearly 90% of cases, only slightly less than in the case of V14. Only the second test revealed that V13 contains mostly redundant information and on average it does not improve quality of Random Forest predictions. The difference is most likely due to the underlying classifier used in each approach. The SVM is essentially a linear classifier, which can be applied to nonlinear problems using some nonlinear, continuous kernel transformation. On the other hand the Random Forest is based on nonlinear and discrete decision trees. Figure 2 in the Lucas et al. suggests that the decision space of the system under scrutiny is non-continuous. The Random Forest can treat such systems more efficiently using less variables, whereas SVM needs higher dimensional spaces to build hyper-plane separating two classes. We have observed such effects in other systems, for example in our earlier study of the recognition of musical instruments, (Kursa et al. 2009). The other differences are less important, since they involve variables with marginal relevance.

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Item 7. Paja et al. should also be aware that some of the co-authors of the Lucas et al paper were co-authors on a related paper that computed sensitivity information using random forest feature scores (doi:10.1002/2014JD022507).

Answer
We would like to thank the referee for this information. We were not aware of this article. Indeed Boyle et al. use Random Forest for estimation of the importance of variables, and although the approach used in their paper is far simpler, but it shows, that Random Forest is a useful tool for analysis of climate simulations and in particular for finding the importance of parameters. We believe that the procedure applied in our paper could be used for analysis of this data as well and return a little bit more information – namely the criterion separating truly informative variables from the non-informative ones. We have included reference to the article in the description of methods, at the end of the following paragraph:

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We have used a different classification algorithm, namely the Random Forest (Breiman, 2001) and instead of the sensitivity analysis we have applied the all-relevant feature selection algorithm Boruta (Kursa et al., 2010). All computations were performed in R environment for statistical modelling (R Development Core Team, 2008), using the randomForest package for classification (Liaw and Wiener, 2002) and the Boruta package for feature selection. (Kursa and Rudnicki, 2010). Interestingly, some of the authors of Lucas et al. have recently used Random Forest in their analysis of the results of the CAM5 model applied for study of Madden Julian Oscillation. It was applied to analyse the influence of the model parameters on selected diagnostic variables.

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Application of all relevant feature selection for the failure analysis of parameter-induced simulation crashes in climate models

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Abstract

The climate models are extremely complex pieces of software. They reflect the best knowledge on the physical components of the climate, nevertheless, they contain several parameters, which are too weakly constrained by observations, and can potentially lead to a crash of simulation. Recently a study by Lucas et al. has shown that machine learning methods can be used for predicting which combinations of parameters can lead to the crash of simulation, and hence which processes described by these parameters need refined analyses. In the current study we re-analyse the dataset used in this research using different methodology. We confirm the main conclusion of the original study concerning the suitability of machine learning for the prediction of crashes. We show that only three of the eight parameters indicated in the original study as relevant for prediction of the crash are indeed strongly relevant, three other are relevant but redundant, and two are not relevant at all. We also show that the variance due to the split of data between training and validation sets has
a large influence both on the accuracy of predictions and relative importance of variables, hence only cross-validated approach can deliver a robust prediction of performance and relevance of variables.

1 Introduction

The development of realistic models of climate is one of the most important areas of research due to the dangers posed by global warming. It is by no means a trivial task since it involves the parameterisation of many processes that are not directly solved within the model. It has been shown by (Lucas et al., 2013) that certain combinations of these parameters, lead to failure of a model, despite each individual parameter having a reasonable value. Authors of this study performed 540 simulations with randomly varied combinations of 18 parameters of the Parallel Ocean Program (POP2) (Smith et al., 2010) module in the Community Climate System Model Version 4 (CCSM4) (UCAR, 2010). About 10 percent of these simulations crashed due to numerical instabilities. Then they have applied machine learning methods to attribute failures to the parameters of the model. To this end they had used the support vector machine (SVM) (Vapnik, 1995) classification to quantify and predict the probability of failure as a function of the values of 18 from POP2 parameters. The causes of the simulation failures were determined through a global sensitivity analysis. Combinations of 8 parameters related to ocean mixing and viscosity from three different POP2 parameterizations were then determined as the major sources of the failures. These 8 parameters were indicated as targets for more detailed research.

These results are somewhat disappointing, since the number of parameters is still rather high. Hence we decided to check whether more elaborate method for analysis could decrease this number further. We have observed potential weak points of the analysis performed by Lucas and co-workers, namely, they have not fully taken into account that the apparent importance of a variable for classification may be in fact the result of a spurious fluctuation. The problem is most acute when a sample used for machine learning algorithm is small. In such a case random fluctuation may introduce spurious correlations within data, which can be utilized by the classification algorithm for model building. The appropriate procedure should be applied to minimize the influence of such random correlations on the final results.

Lucas and co-workers have also analyzed the impact of the decision variable that is used for the classification on the quality of results. While the models were built as an ensemble of learners built on the bootstrap samples of the training set, the evaluation of the classification
performance was based on a single split of data between training set and test set. This setup was due to the construction of the study – simulations for the validation set were performed after the predictions have been made. While this is a very honest method for verification of the predictions, however, it precludes the estimation of the statistical uncertainty of the result. In particular, it is impossible to say whether the observed differences between classification accuracy observed for different decision functions are significant or do they arise due to statistical fluctuations.

The current study is devoted to the reanalysis of the data. It aims at minimizing the influence of random fluctuations on the final results. Our aim was to establish all variables that truly contribute to the final result of the simulations, i.e. whether the simulation was finished successfully or it crashed. To this end we use contrast variables that carry no information on the decision variable, apply Boruta algorithm for all-relevant feature selection and extensive Monte Carlo sampling. We also compare the quality of classification for several subsets of variables used for prediction of simulation result, to perform a parallel check of relevance of variables.

2 Methods

Similarly to the original work, we rely on machine learning algorithms to identify parameters that critically influence the fate of the simulation. The fundamental idea is that when the classification algorithm can predict result of the simulation, i.e. the successful completion of simulation or the crash, using only the information on the values of certain combinations of selected parameters, then these parameters are indeed responsible for the result. In the original paper the authors performed true prediction and achieved a high degree of accuracy, therefore showing the true predictive power of this approach. On the other hand, this setup precludes estimation of statistical uncertainty for some of their findings. In particular, the discussion of the prediction accuracy in sections 4.4 and 4.5 is based on a single split of data between training and test sets and ignores possibility that effects may depend on the particular split.

In the current study we know all results beforehand, thus we are limited to virtual prediction only. In this approach we split the entire dataset into training and validation sets. We then build a model using the training set and check its quality by performing virtual prediction on the validation set and comparing the predicted results with the true ones. One can take advantage of virtualisation to obtain information about the probability distribution of results.
To this end one can perform multiple virtual experiments, with different splits between training and validation sets, and perform classification experiment on each of these splits. The results of individual trials will differ in most cases, allowing one to draw conclusions not only about mean values but also about variance and even shape of probability distribution. Lucas et al. have used this approach for the sensitivity analysis, utilizing ensembles of SVM (Vapnik, 1995) learners for classification. Each member of the ensemble was obtained using different subsample of the training set. The classifier was then used for prediction of the simulation result for the validation set. We have used a different classification algorithm, namely the Random Forest (Breiman, 2001) and instead of the sensitivity analysis we have applied the all-relevant feature selection algorithm Boruta, (Kursa et al., 2010). All computations were performed in R environment for statistical modelling (R Development Core Team, 2008), using the randomForest package for classification (Liaw and Wiener, 2002) and the Boruta package for feature selection, (Kursa and Rudnicki, 2010). Interestingly, some of the authors of Lucas et al. have recently used Random Forest in their analysis of the results of the CAMS model applied for study of Madden Julian Oscillation. It was applied to analyse the influence of the model parameters on selected diagnostic variables.

Random Forest is an ensemble algorithm based on decision trees. To ensure the low correlation between elementary learners, each tree is grown using a different random subsample of the original data set. Moreover, each split in the tree is built using only a random subset of the predictor variables. The number of variables in this subset influences the balance between bias and variance for the training set. The default value for classification tasks is a square root of the total number of variables and it is usually a very robust selection. Random Forest is a robust “of the shelf” algorithm that is easily applicable to various classification and regression tasks. It has only few control parameters and usually it does not need fine tuning for the particular problem under scrutiny. In many cases it has a performance comparable or even better than state of the art classifiers and it rarely fails. A big advantage of the algorithm is that it estimates both the estimate of the classification error and of the importance of variables by internal cross-validation. To estimate the latter it measures how much the accuracy of base learners is decreased when information about variable in question is removed from the system.

Kursa and Rudnicki (2010; Kursa et al., 2010)
R Development Core Team, 2008
Liaw and Wiener, 2002
Boruta (Kursa et al., 2010)
Vapnik, 1995
Kursa and Rudnicki, 2010
Kursa and Rudnicki, 2010; Kursa et al., 2010
Random Forest
randomForest package
Liaw and Wiener, 2002
Boruta package
Kursa and Rudnicki, 2010
Madden Julian Oscillation
random subset of the predictor variables.
Random Forest is a robust “of the shelf” algorithm that is easily applicable to various classification and regression tasks. It has only few control parameters and usually it does not need fine tuning for the particular problem under scrutiny. In many cases it has a performance comparable or even better than state of the art classifiers and it rarely fails. A big advantage of the algorithm is that it estimates both the estimate of the classification error and of the importance of variables by internal cross-validation. To estimate the latter it measures how much the accuracy of base learners is decreased when information about variable in question is removed from the system.
The Boruta algorithm for all-relevant feature selection uses the Random Forest importance measure to infer their relevance. To this end it extends the information system by variables that are non-informative by design – the so-called contrast variables. It then compares the apparent importance of the original variables with that of the non-informative ones. It performs this multiple times using different realizations of the non-informative variables and performs a statistical test. The algorithm finds both strongly and weakly relevant variables. The notions of strong and weak relevance were introduced by (Kohavi and John, 1997) in the context of the ideal classification algorithm. The features are strongly relevant when removing them from the description always results in decreased classification accuracy. Features are weakly relevant, when their removal in some cases may decrease classification accuracy. For a more detailed discussion of relevance and the Boruta algorithm see (Kohavi and John, 1997; Rudnicki et al., 2015). Algorithm has been used in different fields, including bioinformatics, remote sensing, bacteriology and medicine (Aagaard et al., 2012; Ackerman et al., 2013; Buday et al., 2013; Duro et al., 2012; Herrera and Bazaga, 2013; Leutner et al., 2012; Ma et al., 2014; Menikarachchi et al., 2012; Saulnier et al., 2011; Strempel et al., 2013).

The climate simulations dataset is highly biased towards successful completion of simulation. Only 46 cases out of 540 are failures. Such unbalanced datasets are often difficult for classification, because the automatic selection of the majority class results in good, but useless, classification accuracy. In such a case no information is gained and hence one cannot perform feature selection. In the first test of the current study this problem was avoided by application of the following protocol, see Fig 1. Firstly eleven balanced subsamples of training set were constructed, each subsample consisted of all objects from minority class (failed simulations) and 1/11th of majority class (successful simulations). In order to check specificity of the feature selection each dataset was extended by contrast variables. To this end each original variable was duplicated and its values were randomly permuted between all objects. In this way a set of shadow variables that were non-informative by design was added to the original variables. Then the feature selection procedure was performed on each subsample with the help of the all-relevant feature selection algorithm, implemented in Boruta function of the Boruta package. The procedure was repeated 60 times. Altogether all relevant feature selection was performed 660 times. The number of times when the artificially constructed shadow variables were selected as important gives an estimate of the expected level of false discovery. The variables that were selected as important significantly more often than random were examined further, using different test.
The second test probing the importance of variables was performed by analysing the influence of variables used for model building on the prediction quality. The first experiment revealed four variables that were classified as important by Boruta in all, or nearly all, of 660 trials. These variables were considered to form a core variable set, and the model built using these variables was used as a reference. We examined whether removing one of the core variables and whether adding another variable respectively decreases or increases the classification quality measured by AUC. The extension of the core test was examined for three variables that were classified in the first test as important significantly more often than the randomised variables.

The test was performed similarly to the one reported in the original study, see Figure 2. The data set was randomly split into a training set containing 360 objects and a validation set containing 180 objects. The split was performed separately for the minority and majority class, so the number of minority class objects in each training set was 32 and in the validation set it was 14. The randomForest function from the identically named R package was used to perform classification and error estimate. The procedure was repeated 30 times and results of 30 repetitions were analysed.

The number of trees in the forest (parameter ntree both in randomForest and in Boruta functions) was set to 5000 both for feature selection with Boruta and classification with randomForest. In both cases the number of variables examined for each split was equal to the square root of the total number of variables. In our experience these settings are fairly robust, we have examined them internally over multiple datasets (Rudnicki et al., 2015). Moreover, we have checked whether they influence results in the initial trials. The number of trees used was 10 times higher than default, to assure that importance estimate in Random Forest converge to their asymptotic values, the number of trees for classification was the same for consistency.

3 Results and Discussion

The summary of the results of the study is presented in the Table 1. The V1 and V2 variables were deemed important in all 660 cases. Variables V13 and V14 were deemed important in nearly all cases — 593 and 623 cases, respectively. All these variables were also indicated as most important by Lucas et al. However, the results do not agree so well for other variables. Lucas et al. indicated variables V4, V5, V16 and V17 as important but their influence on the
final result was much weaker than that of the first group. In the current study the variables V4 and V16 were deemed important by Boruta for 44 and 66 subsamples, respectively. In both cases the number is significantly higher than the average for the random variables, which was obtained as 25±9. On the other hand variables V5 and V17 were deemed important for 19 and 17 subsamples respectively, and these numbers are lower than the average for random variables. Moreover, variable V9, which was not indicated as important by Lucas et al., was deemed important for 62 subsamples.

Hence the first experiment confirmed the importance of variables V1 and V2, has shown that importance of V13 and V14 is nearly universal, it also confirmed the weak importance of variables V4 and V16. On the other hand the importance of variables V5 and V17 was not confirmed with our method, instead variable V9 was found to be weakly important. The example result of the Boruta run for an interesting sample is presented in Figure 3. In this sample the importance was confirmed for variables V9 and V16, whereas variable V13 was deemed irrelevant. The importance of V4 was higher than that of highest random variable, but only barely so, and hence the final decision of Boruta was “tentative”. One should note, that the importance returned by Boruta is the averaged importance obtained from the underlying Random Forest algorithm. It is not directly interpretable in terms of the fraction of variance explained by given variable.

One should note, that Boruta is an all-relevant feature selection algorithm that aims at finding both strongly and weakly relevant variables, as defined by Kohavi and John. The second test aimed at discerning between strongly and weakly relevant variables. In the case of V1, V2 the removal of the variable from the core dataset resulted in a dramatic drop of AUC, confirming that these variables are truly informative, see Table 1 and Figure 2. In the case of V14 the difference in AUC – referenced further as Δ(AUC) – was smaller, but still statistically significant, whereas for the V13 the Δ(AUC) was much smaller than the standard deviation. Similarly, adding either of the three remaining variables, namely V4, V9 and V16, to the core set, lead to an increase of the AUC by insignificant amount, see Table 1 and Figure 4. Another auxiliary metric that can be used to evaluate the relevance of variables is the number of samples in which the AUC for the model containing the variable is higher than that for the model built without that variable. The results of this metric are consistent with results for the Δ(AUC) – it is 30 for both V1 and V2 and 26 for V14 and these are the only results that are significantly different from random ones. Therefore one can conclude, that only three
variables, namely V1, V2 and V14 are *strongly relevant*, whereas the remaining variables are *weakly relevant*.

One should note that the results of the second test were highly variable and largely dependent on the split of data between test and validation sets. It is illustrated in Figure 5 and examples of the results from several samples are given in Table 2. The highest AUC obtained in the experiment was 0.990 for model built using core variables and V16 in sample #12. In the same sample the AUC for model built from core-V2 was 0.888. On the other hand for sample #1 the highest AUC was obtained for the model built on core+V9 and it was 0.879. Also the relative importance of variables depends strongly on the test sample. For example adding variable V4 to the core set can improve AUC by as much as 0.032 (sample #22) or decrease it by 0.006 (sample #6). Similarly for V16 AUC can decrease by 0.016 (sample #6) or increase by 0.016 (sample #22). Most interestingly removing variable V13, which was deemed relevant by Boruta in nearly 90% of samples, can either decrease the AUC by 0.011 (sample #6) or increase it by 0.030 (sample #22). This results show that one cannot rely on a single split between the training set and test set for the estimate of influence of parameters, and that only the average over sufficiently large number of alternative splits can give robust estimates.

The average of the cross-validated AUC obtained for three strongly important variables, namely V1, V2 and V13, was 0.924. The highest average AUC was obtained for model built using five variables, namely {V1, V2, V9, V13, V14}, nevertheless the value AUC=0.931 was not significantly higher than the value obtained for simpler model built using only three variables. The small differences in AUC arise due to small improvements for assigning the probability of failure of the simulation. Such improvement results in small shift in the ranking from least probable to most probable to fail, without actually improving the error rate at the cost of including two more variables in the model.

A single run of the Boruta algorithm in the first test took 2 minutes on a server equipped with Intel Xeon E5620@2.4GHz CPU. The entire protocol took less than 24 hours of single CPU core. The second test is far less computationally demanding. A single run of the randomForest function takes less than 20 seconds on the same CPU, therefore, computations for the entire protocol take less than 10 minutes. This effort is negligible in comparison with the time required to run 540 simulations of the climate model itself.

The results of the study are mostly in good agreement with the results of Lucas et al., however, importance of the variables is not identical. The most important difference is the
importance of the variable V13 in both studies. This variable is more important than V14 in
the SVM-based model by Lucas et al., whereas our analysis deems it relevant but redundant.
However, one should note that in the first test V13 was deemed relevant in nearly 90% of
cases, only slightly less than in the case of V14. Only the second test revealed that V13
contains mostly redundant information and on average it does not improve quality of Random
Forest predictions. The difference is most likely due to the underlying classifier used in each
approach. The SVM is essentially a linear classifier, which can be applied to nonlinear
problems using some nonlinear, continuous kernel transformation. On the other hand the
Random Forest is based on nonlinear and discrete decision trees. Figure 2 in the Lucas et al.
suggests that the decision space of the system under scrutiny is non-continuous. The Random
Forest can treat such systems more efficiently using less variables, whereas SVM needs
higher dimensional spaces to build hyper-plane separating two classes. We have observed
such effects in other systems, for example in our earlier study of the recognition of musical
instruments, (Kursa et al. 2009). The other differences are less important, since they involve
variables with marginal relevance.

Conclusions
Our reanalysis of the results of 540 simulations is in general qualitative agreement with the
results of Lucas et al. The results of the simulation can be predicted with fairly good accuracy
using the machine learning approach, and the two different methods give very close results.
The cross-validated AUC reported by Lucas et al. by ensemble of SVM classifiers was 0.93.
In the current study the average of the cross-validated AUC obtained for three strongly
important variables, was 0.924.

We have shown by cross-validation that the AUC reported for the prediction experiment
performed by Lucas et al. falls within the range of values that can be expected in such a
prediction, however, one should not assign any weight to the particular value obtained. If the
split between the training set and test set was set differently the resulting AUC for prediction
could be any number between 0.88 and 0.99.

The three most important conclusions for the climate modelling community are following.
Firstly, the efforts on improving the numerical stability of simulations should be concentrated
on 3 parameters of the CCSM4 parallel ocean model, namely vconst_corr, vconst_2, and
bckgrnd_vdc1, that were earlier reported as most important by Lucas et al. The remaining

parameters indicated as important in that study are either redundant or not relevant. Secondly – the machine learning methods in general, and all-relevant feature selection in particular are useful tools for analysis of influence of simulation parameters on the final outcome. Finally, application of machine learning should involve cross-validation, and all important modelling steps should be included in the cross-validation loop.

Author contributions. W. Paja performed most computations and drafted the first version of the manuscript. M. Wrzesien and R. Niemiec performed computations and contributed to the writing. W. R. Rudnicki designed the experiments, and wrote the manuscript.
References


Table 1. Summary of results.

The variables indicated as important by Lucas et al. are marked with *, the variables that were indicated as important in the first test are highlighted in bold face. $\Delta$(AUC) is given in 0.0001 units.

Three values are reported, the number of times the variable was deemed relevant, mean difference in AUC due to adding variable to set of variables and number of times AUC was improved by adding variable to set of variables. The first value is reported for all variables, two other are reported only for these variables that were deemed relevant significantly more often than randomised variables. The unit for $\Delta$(AUC) is 0.0001.

<table>
<thead>
<tr>
<th>Variable</th>
<th>V1*</th>
<th>V2*</th>
<th>V3</th>
<th>V4*</th>
<th>V5*</th>
<th>V6</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td># relevant</td>
<td>660</td>
<td>660</td>
<td>0</td>
<td>44</td>
<td>19</td>
<td>33</td>
<td>25±9</td>
</tr>
<tr>
<td>Mean $\Delta$(AUC)</td>
<td>905±80</td>
<td>749±90</td>
<td>—</td>
<td>20±70</td>
<td>—</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td># improved</td>
<td>30</td>
<td>30</td>
<td>—</td>
<td>16</td>
<td>—</td>
<td>—</td>
<td></td>
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<table>
<thead>
<tr>
<th>Variable</th>
<th>V7</th>
<th>V8</th>
<th>V9</th>
<th>V10</th>
<th>V11</th>
<th>V12</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td># relevant</td>
<td>2</td>
<td>17</td>
<td>62</td>
<td>11</td>
<td>3</td>
<td>5</td>
<td>25±9</td>
</tr>
<tr>
<td>Mean $\Delta$(AUC)</td>
<td>—</td>
<td>—</td>
<td>60±70</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td># improved</td>
<td>—</td>
<td>—</td>
<td>22</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Variable</th>
<th>V13*</th>
<th>V14*</th>
<th>V15</th>
<th>V16*</th>
<th>V17*</th>
<th>V18</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td># relevant</td>
<td>593</td>
<td>623</td>
<td>26</td>
<td>67</td>
<td>19</td>
<td>2</td>
<td>25±9</td>
</tr>
<tr>
<td>Mean $\Delta$(AUC)</td>
<td>11±60</td>
<td>180±80</td>
<td>—</td>
<td>6±60</td>
<td>—</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td># improved</td>
<td>16</td>
<td>26</td>
<td>—</td>
<td>14</td>
<td>—</td>
<td>—</td>
<td></td>
</tr>
</tbody>
</table>
Table 2. Results of experiment 2.

Average AUC obtained for all tested models, as well as examples for five interesting cases. #1 – the sample with lowest AUC from core model, #12 the sample with highest AUC obtained in the study, samples #6, #22 and #30 – samples with core model close to the mean that show variance of AUC for other models.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Variable set</th>
<th>#1</th>
<th>#6</th>
<th>#22</th>
<th>#30</th>
<th>#12</th>
<th>Average</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>core</td>
<td>0.865</td>
<td>0.921</td>
<td>0.922</td>
<td>0.928</td>
<td>0.983</td>
<td>0.925 ± 0.006</td>
</tr>
<tr>
<td></td>
<td>core+V4</td>
<td>0.879</td>
<td>0.915</td>
<td>0.954</td>
<td>0.930</td>
<td>0.982</td>
<td>0.927 ± 0.007</td>
</tr>
<tr>
<td></td>
<td>core+V9</td>
<td>0.866</td>
<td>0.923</td>
<td>0.945</td>
<td>0.919</td>
<td>0.989</td>
<td>0.931 ± 0.006</td>
</tr>
<tr>
<td></td>
<td>core+V16</td>
<td>0.848</td>
<td>0.906</td>
<td>0.938</td>
<td>0.927</td>
<td>0.990</td>
<td>0.926 ± 0.007</td>
</tr>
<tr>
<td></td>
<td>core-V14</td>
<td>0.823</td>
<td>0.907</td>
<td>0.926</td>
<td>0.919</td>
<td>0.967</td>
<td>0.907 ± 0.007</td>
</tr>
<tr>
<td></td>
<td>core-V13</td>
<td>0.877</td>
<td>0.910</td>
<td>0.952</td>
<td>0.921</td>
<td>0.968</td>
<td>0.924 ± 0.006</td>
</tr>
<tr>
<td></td>
<td>core-V1</td>
<td>0.745</td>
<td>0.821</td>
<td>0.806</td>
<td>0.823</td>
<td>0.910</td>
<td>0.835 ± 0.007</td>
</tr>
<tr>
<td></td>
<td>core-V2</td>
<td>0.808</td>
<td>0.808</td>
<td>0.825</td>
<td>0.840</td>
<td>0.888</td>
<td>0.850 ± 0.009</td>
</tr>
</tbody>
</table>
Figure 1. Protocol of the first test.
Figure 2. Protocol of the second test.
Figure 3. Summary of results of the Boruta run. Importance of the variables is shown. The variables are sorted by increasing importance. The variables coloured in green are these, which were classified as relevant. Variables coloured in red are these, which are irrelevant. The blue boxes correspond to respectively minimal (sMin), median (sMed) and maximal (sMax) importance achieved in each run by contrast variables. One can observe wide range of maximal importance values that can be achieved by random variables. In particular in many iterations it can be higher than importance of truly relevant variables.
Figure 4. AUC obtained in simulations study grouped by subset of variables used for model building. The labels are coded in the following way:

- C – core set of variables \{V_1, V_2, V_{13}, V_{14}\};
- C+X – the core set was extended by adding variable VX, where X is one of \{4,9,16\};
- C-X – the variable VX was removed from the core set, with X = \{1,2,13,14\}. 
Figure 5. AUC obtained in simulations study grouped by split between training and validation set.