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Title: An evaluation framework for input variable selection algorithms for environmental data-driven models

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Abstract: Input Variable Selection (IVS) is an essential step in data-driven modelling and is particularly relevant in environmental applications, where potential input variables are often collinear and redundant. While methods for IVS continue to emerge, each has its own advantages and limitations and no single method is best suited to all datasets and modelling purposes. Rigorous evaluation of IVS methods would allow their effectiveness to be properly identified in various circumstances. However, such evaluations are largely neglected due to the lack of guidelines to facilitate consistent and standardised assessment. This work proposes a new evaluation framework, which consists of benchmark datasets with the typical properties of environmental data, a recommended set of evaluation criteria and a website for sharing data and code. The framework is demonstrated on four IVS algorithms commonly used in environmental modelling studies. The results indicate interesting differences in the algorithms' performance that have not been identified previously.

Response to Reviewers: Editor

I have now received reviews of the above paper and these lead me to recommend that revision according to all the reviewers' comments is necessary. I may not send it back to reviewers, trusting that you will cut it down, otherwise few people will not bother reading it.

Response to Editor comment No. 1. We significantly reduced the manuscript length by mostly focusing on Section 2 and 3, as also suggested by reviewer #2. Where possible, we also tried to reduce Section 5. Overall, we obtained a reduction of about 6 pages (from the introduction to the conclusion) with respect to the previous version of the manuscript. Furthermore, we removed Appendix A, since this material can be directly accessed from the framework website. This gives an overall reduction of 21 pages.

Another issue is that I'd like it to fit better with EMS being a generic journal and so link to our key outputs. Most citations to EMS papers are to the authors themselves! Just one way to do this is to link with/refer to other key modelling concepts and issues in the journal. For example see the next paragraph.

On model evaluation: that it is credible and addressed well. In this connection, I would like you to justify, and if pertinent expand or comment upon, your choice of evaluation metrics and methods among the ones, for example, in the recent EMS Position paper of Bennett et al (2013) on performance evaluation (they propose a 5-step procedure for evaluating the performance of models). You could also add/comment on visual methods and quantitative measures used to examine model quantities and residuals, including visual inspection. There are several other evaluation issues you could address/compare as well and the paper by Robson and cited below presents an excellent example in Section 13 of that paper. One of our aims for EMS is to strengthen the credibility and relevance of the modelling reported and do this whatever the environmental problem sector. That way your paper is more suited to our journal.

Bennett ND, Croke BFW, Guariso G, Guillaume JHA, Hamilton SH, Jakeman AJ, Marsili-Libelli S, Newham LTH, Norton JP, Perrin C, Pierce SA, Robson B, Seppelt R, Voinov AA, Fath BD and Andreassian V (2013) Characterising performance of environmental models. *Environmental Modelling & Software* 40: 1-20.

Robson, Barbara J (in press) State of the art in modelling of phosphorus in aquatic systems: Review, criticisms and commentary. *Environmental Modelling & Software*, In Press, Corrected Proof, Available online 6 February 2014.

Response to Editor comment No. 2. We linked our work with other key contributions in EMS by discussing about evaluation metrics and other issues related to the use of IVS algorithms in environmental modelling problems (as also suggested by reviewer #1). This discussion is contained in the newly added Section 6.3:

“Unlike the synthetic data here considered, a key aspect of real-world environmental modelling problems is that the true underlying function is unknown, and IVS is thus used to reduce the uncertainty in the model development process by selecting a subset of relevant and non-redundant input variables. This opens some relevant theoretical and practical issues that are highlighted below:

- Most of the IVS algorithms currently available select a unique subset of input variables, although the structural uncertainty in the inputs to be used often results in the possibility of choosing different, but equally informative, subsets. An attempt to account for this issue was recently made by Sharma and Chowdhury (2011), who proposed a PMI-based heuristic approach to select five different subsets of predictors in the context of medium-term hydro-climatic forecasting. The approach ensures that the cross-dependence between these subsets is limited, while the predictions of the resulting models are eventually combined with ensemble averaging.
- In many practical situations, input variables can be characterised by errors, due, for example, to the interpolation of data in space and time or to the conversion of point measurement into areal values. Whilst methods exist for assessing the impact of input errors on parameter estimation procedures (Chowdhury and Sharma, 2007; Woldemeskel et al., 2012), IVS algorithms cannot take into account the change in the uncertainty associated with the different inputs.
- A benefit of IVS is the improvement in the performance of the model being identified. Although the manner in which such performance is characterised depends on the specific domain of interest and the model objectives (Jakeman et al., 2006), two important aspects should always be considered when dealing with quantitative testing. First, the use of observational data for comparison must rely on appropriate data-division methods, such as cross-validation or bootstrapping, that allow for testing the ability of the model to generalise. Data division can account for both temporal and spatial dimensions, so it is suitable for spatial modelling as well (see Chowdhury and Sharma (2009) for an application to hydrological modelling problems). Second, an exhaustive quantitative evaluation should rely on a set of metrics focussing on different aspects in order to test the ability of the model in reproducing all the important features of the system. The reader is referred to Bennett et al. (2013) for a comprehensive

review of techniques available for both data-division and quantitative evaluation, and to Robson (2014) for a more general assessment of environmental models.”

In preparing the review we used the following rules: references to line numbers, equations and figures are all to the original manuscript; authors’ reply are in blue.

Reviewer #1

It was a joy reading this paper - very nicely put together. I just had four additions to include to what has been written here.

172 - I think one other issue needs to be added here. Most of these algorithms assume a unique input variable set exists. In my experience, a natural system can be equally well described using alternate predictor sets. This represents the structural uncertainty in specifying any one predictive model. I have attempted to highlight this issue in an invited seasonal forecasting paper (Sharma, A., and S. Chowdhury (2011), Coping with model structural uncertainty in medium-term hydro-climatic forecasting, *Hydrology Research*, 42(2-3), 113, doi:10.2166/nh.2011.104.) where we select 5 plausible predictor sets, but ensure the cross-dependence between them is not too high (so they can be argued to represent independent predictive models). These when combined using some model averaging rationale, lead to significant improvements in the stability of the predictive model. My argument is - for a practical problem when you wouldn’t like the model to issue unstable predictions, I would pursue this option any day over selecting a single unique model. The input variable selection problem never allows for reference datasets where multiple predictive models are plausible. One of these needs to be included in any evaluation framework that is proposed (maybe just as a mixture model having two different (psuedo-independent) predictor sets).

Response to Reviewer comment No. 1. As explained in our reply to the Editor’s comments, we introduced a new section to discuss the most important issues related to the use of IVS algorithms in real-world environmental modelling problems (Section 6.3). In this case, the true underlying function is unknown and different, but equally informative, subsets could indeed exist. We highlighted this aspect in Section 6.3, where we also referred to the paper mentioned above.

“Most of the IVS algorithms currently available select a unique subset of input variables, although the structural uncertainty in the inputs to be used often results in the possibility of choosing different, but equally informative, subsets. An attempt to account for this issue was recently made by Sharma and Chowdhury (2011), who proposed a PMI-based heuristic approach to select five different subsets of predictors in the context of medium-term hydro-climatic forecasting. The approach ensures that the cross-dependence between these subsets is limited, while the predictions of the resulting models are eventually combined with ensemble averaging.”

1458 - I would add another dataset to this list that we needed to create to highlight the importance of the predictive algorithm when coupled to input variable selection in Sharma and Mehrotra 2014 - I suggest this as the typical datasets listed would not be able to differentiate between situations where the partial weights associated with each predictor variable are dramatically different - something that was pointed out to us in the review process of the above mentioned paper. Please see equation 22 of the paper.

Response to Reviewer comment No. 2. Since we preferred not to highlight the importance of the predictive algorithm (and the corresponding predictive performance), we decided not to include the dataset within the framework. Furthermore, we notice that datasets 6-8 and 11-18 are characterized by similar properties.

1500 - I think the selection metrics being considered could be expanded. For instance, if I am developing a predictive model to make predictions in space, the assessment can be done by leaving data points out one at a time (the usual leave-one out cross-validation) or entire blocks (I think this is called block cross-validation but not sure). If model is making prediction over time, the same thing applies along with an independent sample, the blocks here representing longer periods of time to account for persistence that may create bias with L1CV measures. An example of this is in one of my seasonal forecasting papers - Chowdhury, S., and A. Sharma (2009), Multisite seasonal forecast of arid river flows using a dynamic model combination approach, *Water Resources Research*, 45(10), doi:10.1029/2008wr007510. What I like most about this paper is the very extensive cross-validation that was performed towards the end, which showed the differences when using one cross-validation measure versus another.

Response to Reviewer comment No. 3. As explained in Section 3.2.1, we believe that the predictive performance should not be used when dealing with synthetic data (such as those proposed in this framework), since the accuracy depends on different factors, e.g. choice of the model or calibration method. This said, we understand that the predictive accuracy becomes important in case of real-world applications, so we included a discussion about this aspect in Section 6.3.

“A benefit of IVS is the improvement in the performance of the model being identified. Although the manner in which such performance is characterised depends on the specific domain of interest and the model objectives (Jakeman et al., 2006), two important aspects should always be considered when dealing with quantitative testing. First, the use of observational data for comparison must rely on appropriate data-division methods, such as cross-validation or bootstrapping, that allow for testing the ability of the model to generalise. Data division can account for both temporal and spatial dimensions, so it is suitable for spatial modelling as well (see Chowdhury and Sharma (2009) for an application to hydrological modelling problems). Second, an exhaustive quantitative evaluation should rely on a set of metrics focussing on different aspects in order to test the ability of the model in reproducing all the important features of the system. The reader is referred to Bennett et al. (2013) for a comprehensive review of techniques available for both data-division and quantitative evaluation, and to Robson (2014) for a more general assessment of environmental models.”

Last point - I have not published this yet - but my PMI code also takes into account the change in the uncertainty associated with the predictor variable over time. Again - this was included as typical seasonal forecasting problems have markedly different standard errors depending on when the data was collected. A good example of implications of this changing error on predictions is in Chowdhury, S., and A. Sharma (2007), Mitigating Parameter Bias in Hydrological Modelling due to Uncertainty in Covariates, *Journal of Hydrology*, 340(doi:10.1016/j.jhydrol.2007.04.010), 197-204. But a better example of how these standard errors can be ascertained (varying over space and time, in this case for GCM simulations) is in Woldemeskel, F. M., A. Sharma, B. Sivakumar, and R. Mehrotra (2012), An error estimation method for precipitation and temperature projections for future climates, *Journal of Geophysical Research-Atmospheres*, 117, doi:Artn D22104Doi 10.1029/2012jd018062. Strongly feel predictor identification needs to offer a sensible basis of including such variations in data quality over time. This should be stated somewhere in this paper.

Response to Reviewer comment No. 4. This aspect is discussed in Section 6.3.

“In many practical situations, input variables can be characterised by errors, due, for example, to the interpolation of data in space and time or to the conversion of point measurement into areal values. Whilst methods exist for assessing the impact of input errors on parameter estimation procedures (Chowdhury and Sharma, 2007; Woldemeskel et al., 2012), IVS algorithms cannot take into account the change in the uncertainty associated with the different inputs.”

On the whole, this is a great paper, that can be quite useful to people who identify predictor variables for use in different prediction problems. Well done folks!

Response to Reviewer comment No. 5. We thank the reviewer for the comment.

Reviewer #2

The authors propose a framework in three points to evaluate input variable selection (IVS) algorithms:

- 1) 26 benchmark synthetic datasets
- 2) a set of evaluation criteria
- 3) website for sharing data and results

Four IVS algorithms are compared and evaluated according to the proposed framework and discussed thoroughly. The idea is really interesting and I think frameworks of this type are more and more developed and are necessary to help research scientist be more systematic in their evaluation and comparison of new and existing methodologies. The paper is generally well written but is very long (91 pages with appendices and 56 pages before the reference section). I think it could be shortened without diminishing its coherence. I will suggest some possible ways to shorten it below.

Response to Reviewer comment No. 1. We understand that the paper is a bit lengthy, so we shortened it by removing some marginal elements (while improving some specific aspects). The revised version is 21 pages shorter (including the appendices).

I have one major disappointment: I could not find the website at the address mentioned on p. 31 (www.ivs4em.deib.polimi.it) only www.deib.polimi.it works but from there, I cannot find the framework webpage. I also tried some Google searches unsuccessfully. I think this is a limitation of the paper, if one does not have access to the benchmark datasets and cannot have a look at the web page, the whole paper remains at the stage of a good idea. Also, I think the paper could include snapshot images of the website to illustrate its functionalities for sharing results for instance. I would also expect the authors to include a functionality in the website which would allow the computation of the recommended criteria automatically. If not directly in the website, some R code could be shared to compute the criteria easily and people could contribute to new criteria.

Response to Reviewer comment No. 2. We fixed this problem. The website is now accessible, and the updated url (<http://ivs4em.deib.polimi.it>) is included in the revised version of the manuscript. From the website it is possible to download the 26 datasets (with their corresponding description), the source code of each IVS algorithm and an R script to compute the evaluation criteria. We have also included a functionality to upload algorithms, datasets and evaluation criteria.

We agree with the reviewer that the paper could include snapshot images of the website; however, we decided not to include them in order to limit the manuscript length.

Other comments

Section 2 describes the background on IVS methods. I found this section both a little long and not so easy to understand. I had to read the Guyon and Elisseeff (2003) paper to understand more clearly the three IVS categories. In particular, filters method are basically ranking methods (as described in Guyon and Elisseeff (2003)) and I think it's more intuitive to present them by mentioning ranks.

Response to Reviewer comment No. 3. Following the reviewer's suggestion, we shortened Section 2 and we clarified all the unclear aspects. Furthermore, we included at the beginning of Section 2.2.1 a brief explanation of filters that explicitly refers to ranking methods.

I would suggest to start the description of each class of IVS by a typical algorithm from this class. This would help to understand the definition of the class. For the filters, the method of ranking in terms of correlation between one input and the output, for instance. For wrapper, the GA-ANN method used in the application of the framework could be described rapidly here. And for embedded algorithms, I would think of the LASSO algorithm which is quite popular.

Response to Reviewer comment No. 4. We agree with the reviewer that a simple description of each class of IVS would simplify the understanding of Section 2.2. Hence, we included a brief explanation of filters, wrappers and embedded algorithms at the beginning of Section 2.2.1, 2.2.2 and 2.2.3, respectively. We think that this approach is more effective than describing a typical algorithm for each class.

P. 10, lines 202-204: I have some trouble to understand why the ACF and PACF are useful for input selection. As far as I know, these techniques are used for time series analyses to choose the proper coefficients in an ARMA model. I would suggest to mention here the partial correlation and partial mutual information which are used in the application of the framework later.

Response to Reviewer comment No. 5. ACF and PACF can be used to measure the (linear) correlation between inputs and output, and then rank the former according to the pairwise correlation. As such, they can be seen as filters. Following the reviewer's suggestion, we also mentioned the Partial Correlation Input Selection algorithm.

P. 12 lines 245-250: the description of the Gamma near-neighbour test was not clear to me. I am wondering if it is useful since this method is not used in the comparison of IVS algorithms and the purpose of the paper is not to review the state-of-the-art on IVS algorithms.

Response to Reviewer comment No. 6. The description of the Gamma near-neighbour test was shortened as suggested.

Section 3 describes the evaluation framework. Basically, as far as I am concerned, two things are missing: some real datasets and a performance criterion based on predictive accuracy. I understand the point made by the authors for the synthetic datasets: it is the only way to know the "true" inputs and their performance criteria SA are based on this knowledge. However, from a practical point of view, I am, most of the time, mainly interested to evaluate if the model selection I performed yield the best model in terms of predictive power. I think that some real datasets along with a predictive accuracy criterion would be complementary to the framework. This could be similar in spirit with the Delve datasets mentioned by the authors: some datasets are used for development and other for assessment. The real datasets could serve the later goal.

Response to Reviewer comment No. 7. We understand the reviewer's suggestion, but we believe that including some real datasets and one, or more, performance of predictive accuracy may be counterproductive. This opinion is supported by the following reasons: 1) There exists a variety of filters that do not rely on any underlying model (induction or learning algorithm), so it is not possible to evaluate the accuracy of such algorithms in terms of predictive accuracy. This would be against the rationale of the IVS framework, which is aimed at supporting the quantitative (and qualitative) evaluation of any input selection algorithm; 2) The predictive accuracy is 'biased' by several factors, such as the choice of the underlying model and calibration (and validation) algorithm. Minimizing such bias would require introducing an exhaustive comparison of different models (e.g. neural networks,

regression trees, linear models, support vector machines etc.) and calibration methods, but this would dramatically affect the length of the manuscript; 3) The same reasoning applies to the inclusion of some real datasets. Indeed, the comparison of different input selection algorithms on some real datasets could only be run by comparing the predictive accuracy of some underlying models; furthermore 4) The inclusion of a few real datasets prevents an exhaustive assessment of the IVS algorithms against the statistical properties described in Section 3.1.

P. 18 lines 399-402. The sentence "Finally, the use of synthetic data enable previously analysed datasets ... would provide very little information about algorithm performance" is not clear to me.

Response to Reviewer comment No. 8. The sentence has been removed.

P. 19 line 417 : "a universal approximator", like an artificial neural network ? or the authors have something else in mind ?

Response to Reviewer comment No. 9. Yes, a feed-forward neural network (with a single hidden layer containing a finite number of neurons) could indeed serve as a universal approximator (Cybenko, 1989). We clarified this aspect in the revised version of the manuscript.

"The amount of noise in the output is defined as the fraction of the variance that would remain unexplained if a universal approximator, such as an artificial neural network (Cybenko, 1989), were used on an infinite training set."

Section 3.2.1. Selection accuracy: do we really need SA in addition to SAe and SAc? I find the later two sufficient since SA is computed from them. Moreover, SA requires to set a parameter which controls the tradeoff between SAe and SAc and it seems not necessary to make such a choice.

Response to Reviewer comment No. 10. We believe that the three scores (i.e. SA, SAc and SAe) are important, since they serve two different purposes: 1) The Selection Accuracy (SA) makes the comparison between different algorithms quite fast and straightforward, since it quantifies the degree to which a model has been correctly or incorrectly specified. Furthermore, the presence of the parameter γ allows the user to weight the importance of missing a relevant input against choosing an extraneous one; 2) The SAc and SAe allow for a more in-depth analysis, since they quantify the proportion of correct and extraneous inputs that have been selected. The single SA score also allows a simple and direct trade-off between selection accuracy and runtime.

Computational efficiency: I have a tendency to think that the total runtime is enough as a measure of computational efficiency. I understand it is not directly comparable across platforms and programming languages but I am not sure if that really matters that much. What basically matters is the order of magnitude: does it take a couple of seconds or a couple of days?

Response to Reviewer comment No. 11. The total runtime provides simple, 'practical' information that is certainly useful to most users and practitioners. For this reason, the results in terms of runtime are reported within the text, while the analysis of computational complexity is reported in Appendix C (now Appendix B). Although only few readers may be interested in it, we believe that such analysis can have both theoretical (e.g. determining the growth rate of the runtime) and practical (e.g. planning the execution of several IVS experiments) implications, particularly as enables platform independent comparisons of the computational efficiency of different IVS algorithms. This will become increasingly important as researchers will add the performance of different algorithms to the website, as these measures will enable computational efficiency to be compared in an objective manner.

P. 29 lines 657-658: how does the framework provides a theoretical measure of computational complexity? as far as I know, this has to be computed for each IVS algorithm by considering the computation steps involved. This would be a kind of $O(NP)$ classification for instance, am I right ?

Response to Reviewer comment No. 12. Yes, the theoretical measure of computational complexity is determined for each algorithm by evaluating the computational steps involved at each iteration (see Appendix C). This concept has been further clarified in the revised version of the manuscript.

“In particular, the analysis of computational complexity is determined for each algorithm by evaluating the computational steps involved at each iteration, and it is aimed at producing a theoretical classification that estimates the increase in run-time as a function of the input dimensionality N and P .”

Experimental setup

I found it difficult to follow the explanations on the IVS algorithms and on their performance without further explanations on their mechanisms which are given in the appendix. This is why I am suggesting to use the space in the section 2 to already introduce the IVS algorithms which will be compared.

Response to Reviewer comment No. 13. We understand that Section 4 may appear unclear without reading the appendix, but, at the same time, we think that Section 2 should contain a general description of IVS approaches and not a detailed description of the IVS algorithms adopted in this study. In order to solve this problem, we included a brief description of each algorithm in Section 4, and we tightened the connection between Section 4 and the appendix.

I am wondering if it is useful to include 4 IVS algorithms since this means that all of them should be described in details for the reader to understand what is going on. For instance, p. 32 line 743, I am wondering how the Gaussian reference bandwidth is set and line 749, how do you compute the "correlation between inputs and output and a multiple linear regression".

Response to Reviewer comment No. 14. The presence of four algorithms is critical to demonstrate why the framework can be useful to identify the pros and cons of different types and classes of IVS algorithms. For example, the comparison between PCIS and PMIS shows the effect due to the presence of nonlinearities, while the one between PMIS and IIS is used to discuss the effects of non-Gaussian data. Furthermore, the comparison between filters (PCIS, PMIS and IIS) and wrappers (GA-ANN) allows discussing the computational demands of different methodologies. Limiting the comparison to two algorithms would not allow for this exhaustive analysis. This said, we understand the reviewer's concern, so we clarified all these technical aspects in Section 4 (please refer to the previous reply as well).

It would probably be possible to retain 2 distinct IVS algorithms and to compare them in order to illustrate the framework. The paper would be easier to read then since the goal is not so much to inform on IVS algorithms than to present to framework.

Response to Reviewer comment No. 15. Please refer to the previous reply.

Other questions on IVS algorithms: p. 33 line 755 what are SISO models? I found the explanation later in the appendix. In general, the explanation of the IIS algorithm was fairly obscure to me.

Response to Reviewer comment No. 16. The description of the IIS algorithm has been improved as suggested.

p. 33 lines 767: a 1 hidden unit neural network do not have much non-linear capability. I understand it takes time to tune the number of hidden units of a neural network but otherwise, they do not have much predictive power.

Response to Reviewer comment No. 17. Yes, we totally agree with this remark (which is indeed commented on in Section 6.1). The adoption of such architecture, however, can easily serve our purpose: we aim at practically demonstrating the pros and cons of wrappers (and filters), rather than providing a definitive answer as to which of the algorithms performs best.

p. 33 line 771: since the number of hidden units is fixed, what is the use of k-fold cross-validation?

Response to Reviewer comment No. 18. The k-fold cross-validation is used to quantify the accuracy of the ANN. We clarified this aspect in the revised version of the manuscript.

“The accuracy of the ANN is measured in terms of out-of-sample AIC, computed using a k-fold cross-validation (with $k = 5$).”

p.36 lines 841-842 : " ... all four combinations of SAc and SAe were obtained for the combination of IVS algorithm and datasets..." this sentence needs to be rephrased.

Response to Reviewer comment No. 19. The sentence has been rephrased as suggested.

“Furthermore, Figure 4 shows that different values of SAc and SAe were obtained for the combination of IVS algorithms and datasets considered.”

Regarding Figs 4-5 and Figs 6-7, I think they could be re-organized; as it is, they are redundant. The authors could either choose to show the SA scores for datasets which yield contrasted results for the four IVS algorithms or to group datasets according to their properties (as it is done in the text in section 5.1.2).

Response to Reviewer comment No. 20. Following the reviewer’s comment, we removed Figure 4 and 6, since the most of the information in Figure 4 (or 6) is available from Figure 5 (or 7). The reason for maintaining Figure 5 and 7 is that they allow organizing the results by dataset (Figure 5) and by algorithm (Figure 7). The former highlights the performance of the four IVS algorithms on the same modelling conditions, while the latter provides insight into the way different dataset properties impact on the behavior of a specific algorithm.

The discussion on the results could be more condensed: p.44 lines 1021-1033: I found pretty evident that larger N helps model selection, I would suggest to shorten lines 1024-1033.

Response to Reviewer comment No. 21. Section 5.1.3 (‘Effect of N and P on algorithm performance’) has been revised and shortened. In general, the entire Section 5 has been thoroughly revised and condensed.

Computational efficiency

I found it difficult to follow the discussion on where the computations take more time for each IVS algorithm since I was not very familiar with them. I kept wondering: what is exactly Extra-trees, GRNN, PCIS, IIS... By retaining just 2 IVS algorithms and providing more detailed explanations would probably help to benefit from the kind of discussion in this section.

Response to Reviewer comment No. 22. Section 5.2 gives two different types of information about ‘Computational efficiency’. The first is based on the total runtime (Table 2) and it is built on the concept

that filter algorithms (such as PMIS, PCIS and IIS) are computationally efficient, while wrappers (such as the GA-ANN algorithm) require more computing resources. This information is directly accessible by any reader, and does not require being familiar with the algorithms considered. The second information, which is based on the analysis of complexity (Table 3), requires an in-depth knowledge of the algorithms, so we believe that the improvements to Section 4 (plus the presence of a dedicated appendix) will allow the interested readers in understanding the technical aspects of such analysis.

I found the qualitative criteria section quite long. I understand the interest in these type of criteria but it could probably be shortened.

Response to Reviewer comment No. 23. The section was shortened as recommended.

Reviewer #3

The objective of this paper is to create a framework for evaluating and comparing input subset selection (IVS) algorithms for environmental modeling applications. IVS for environmental systems modeling is an extremely challenging task because of the vast number of possible explanatory variables given the space/time correlation of the processes being modeling. However, for the same reason there is also the possibility for significant colinearity of input variables. For this reason IVS is an important first step for any environmental modeling project. Unfortunately, as noted by the authors, there has been little research into what makes a "good" IVS algorithm, as most IVS algorithm research has been focused at a particular dataset or modeling task. The proposed framework would create a repository of data sets and algorithms that would permit comparison of the existing or newly proposed IVS algorithms to identify which perform well in general, thus providing guidance on which algorithm to select for new modeling projects.

This is a well written paper describing project of great interest to the environmental modeling community. The discussion of existing IVS methods is thorough given the scope and length of the paper, and the explanation of the evaluation criteria and the benchmark synthetic data sets is thorough. I recommend this paper be published as-is by Environmental Modelling and Software.

Response to Reviewer comment No. 1. We thank the reviewer for the comment.

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August 14, 2014

A.J. Jakeman (Editor in Chief)

Dear Tony,

We would like to thank you and the reviewers for the thorough and very helpful review. We took all the suggestions into consideration and revised the manuscript accordingly.

In particular, we improved the manuscript by focusing on three main aspects:

- We reduced the manuscript length by about 6 pages (from the introduction to the conclusion) by shortening Section 2, 3 and 5. Furthermore, we removed Appendix A. This gives an overall reduction of 21 pages with respect to the previous version of the manuscript;
- We linked our work with other key contributions in EMS, and included a discussion about the challenges related to the use of IVS algorithms in environmental modelling problems (see Section 6.3);
- We updated the link to the website url (<http://ivs4em.deib.polimi.it>). The website is now complete, and it also includes the scripts for calculating the evaluation criteria, as suggested by reviewer #2.

Further details and relevant information can be found in our response to the reviewers' comments.

Should you need further information, please do not hesitate to contact me.

Sincerely,

Stefano Galelli, PhD
(Corresponding author)



Reply to reviewers about paper ENVSOFT-S-14-00494

An evaluation framework for input variable selection algorithms for environmental data-driven models

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Editor

I have now received reviews of the above paper and these lead me to recommend that revision according to all the reviewers' comments is necessary. I may not send it back to reviewers, trusting that you will cut it down, otherwise few people will not bother reading it.

We significantly reduced the manuscript length by mostly focusing on Section 2 and 3, as also suggested by reviewer #2. Where possible, we also tried to reduce Section 5. Overall, we obtained a reduction of about 6 pages (from the introduction to the conclusion) with respect to the previous version of the manuscript. Furthermore, we removed Appendix A, since this material can be directly accessed from the framework website. This gives an overall reduction of 21 pages.

Another issue is that I'd like it to fit better with EMS being a generic journal and so link to our key outputs. Most citations to EMS papers are to the authors themselves! Just one way to do this is to link with/refer to other key modelling concepts and issues in the journal. For example see the next paragraph.

On model evaluation: that it is credible and addressed well. In this connection, I would like you to justify, and if pertinent expand or comment upon, your choice of evaluation metrics and methods among the ones, for example, in the recent EMS Position paper of Bennett et al (2013) on performance evaluation (they propose a 5-step procedure for evaluating the performance of models). You could also add/comment on visual methods and quantitative measures used to examine model quantities and residuals, including visual inspection. There are several other evaluation issues you could address/compare as well and the paper by Robson and cited below presents an excellent example in Section 13 of that paper. One of our aims for EMS is to strengthen the credibility and relevance of the modelling reported and do this whatever the environmental problem sector. That way your paper is more suited to our journal.

Bennett ND, Croke BFW, Guariso G, Guillaume JHA, Hamilton SH, Jakeman AJ, Marsili-Libelli S, Newham LTH, Norton JP, Perrin C, Pierce SA, Robson B, Seppelt R, Voinov AA, Fath BD and Andreassian V (2013) Characterising performance of environmental models. *Environmental Modelling & Software* 40: 1-20.

Robson, Barbara J (in press) State of the art in modelling of phosphorus in aquatic systems: Review, criticisms and commentary. *Environmental Modelling & Software*, In Press, Corrected Proof, Available online 6 February 2014.

We linked our work with other key contributions in EMS by discussing about evaluation metrics and other issues related to the use of IVS algorithms in environmental modelling problems (as also suggested by reviewer #1). This discussion is contained in the newly added Section 6.3:

“Unlike the synthetic data here considered, a key aspect of real-world environmental modelling problems is that the true underlying function is unknown, and IVS is thus used to reduce the uncertainty in the model development process by selecting a subset

of relevant and non-redundant input variables. This opens some relevant theoretical and practical issues that are highlighted below:

- Most of the IVS algorithms currently available select a unique subset of input variables, although the structural uncertainty in the inputs to be used often results in the possibility of choosing different, but equally informative, subsets. An attempt to account for this issue was recently made by Sharma and Chowdhury (2011), who proposed a PMI-based heuristic approach to select five different subsets of predictors in the context of medium-term hydro-climatic forecasting. The approach ensures that the cross-dependence between these subsets is limited, while the predictions of the resulting models are eventually combined with ensemble averaging.*
- In many practical situations, input variables can be characterised by errors, due, for example, to the interpolation of data in space and time or to the conversion of point measurement into areal values. Whilst methods exist for assessing the impact of input errors on parameter estimation procedures (Chowdhury and Sharma, 2007; Woldemeskel et al., 2012), IVS algorithms cannot take into account the change in the uncertainty associated with the different inputs.*
- A benefit of IVS is the improvement in the performance of the model being identified. Although the manner in which such performance is characterised depends on the specific domain of interest and the model objectives (Jakeman et al., 2006), two important aspects should always be considered when dealing with quantitative testing. First, the use of observational data for comparison must rely on appropriate data-division methods, such as cross-validation or bootstrapping, that allow for testing the ability of the model to generalise. Data division can account for both temporal and spatial dimensions, so it is suitable for spatial modelling as well (see Chowdhury and Sharma (2009) for an application to hydrological modelling problems). Second, an exhaustive quantitative evaluation should rely on a set of metrics focussing on different aspects in order to test the ability of the model in reproducing all the important features of the system. The reader is referred to Bennett et al. (2013) for a comprehensive review of techniques available for both data-division and quantitative evaluation, and to Robson (2014) for a more general assessment of environmental models.”*

In preparing the review we used the following rules: references to line numbers, equations and figures are all to the original manuscript; authors' reply are in blue.

Reviewer #1

It was a joy reading this paper - very nicely put together. I just had four additions to include to what has been written here.

172 - I think one other issue needs to be added here. Most of these algorithms assume a unique input variable set exists. In my experience, a natural system can be equally well described using alternate predictor sets. This represents the structural uncertainty in specifying any one predictive model. I have attempted to highlight this issue in an invited seasonal forecasting paper (Sharma, A., and S. Chowdhury (2011), Coping

with model structural uncertainty in medium-term hydro-climatic forecasting, *Hydrology Research*, 42(2-3), 113, doi:10.2166/nh.2011.104.) where we select 5 plausible predictor sets, but ensure the cross-dependence between them is not too high (so they can be argued to represent independent predictive models). These when combined using some model averaging rationale, lead to significant improvements in the stability of the predictive model. My argument is - for a practical problem when you wouldn't like the model to issue unstable predictions, I would pursue this option any day over selecting a single unique model. The input variable selection problem never allows for reference datasets where multiple predictive models are plausible. One of these needs to be included in any evaluation framework that is proposed (maybe just as a mixture model having two different (psuedo-independent) predictor sets).

As explained in our reply to the Editor's comments, we introduced a new section to discuss the most important issues related to the use of IVS algorithms in real-world environmental modelling problems (Section 6.3). In this case, the true underlying function is unknown and different, but equally informative, subsets could indeed exist. We highlighted this aspect in Section 6.3, where we also referred to the paper mentioned above.

“Most of the IVS algorithms currently available select a unique subset of input variables, although the structural uncertainty in the inputs to be used often results in the possibility of choosing different, but equally informative, subsets. An attempt to account for this issue was recently made by Sharma and Chowdhury (2011), who proposed a PMI-based heuristic approach to select five different subsets of predictors in the context of medium-term hydro-climatic forecasting. The approach ensures that the cross-dependence between these subsets is limited, while the predictions of the resulting models are eventually combined with ensemble averaging.”

1458 - I would add another dataset to this list that we needed to create to highlight the importance of the predictive algorithm when coupled to input variable selection in Sharma and Mehrotra 2014 - I suggest this as the typical datasets listed would not be able to differentiate between situations where the partial weights associated with each predictor variable are dramatically different - something that was pointed out to us in the review process of the above mentioned paper. Please see equation 22 of the paper.

Since we preferred not to highlight the importance of the predictive algorithm (and the corresponding predictive performance), we decided not to include the dataset within the framework. Furthermore, we notice that datasets 6-8 and 11-18 are characterized by similar properties.

1500 - I think the selection metrics being considered could be expanded. For instance, if I am developing a predictive model to make predictions in space, the assessment can be done by leaving data points out one at a time (the usual leave-one out cross-validation) or entire blocks (I think this is called block cross-validation but not sure). If model is making prediction over time, the same thing applies along with an independent sample, the blocks here representing longer periods of time to account for persistence that may create bias with L1CV measures. An example of this is in one of my seasonal forecasting papers - Chowdhury, S., and A. Sharma (2009), Multisite seasonal forecast of arid river flows using a dynamic model combination

approach, *Water Resources Research*, 45(10), doi:10.1029/2008wr007510. What I like most about this paper is the very extensive cross-validation that was performed towards the end, which showed the differences when using one cross-validation measure versus another.

As explained in Section 3.2.1, we believe that the predictive performance should not be used when dealing with synthetic data (such as those proposed in this framework), since the accuracy depends on different factors, e.g. choice of the model or calibration method. This said, we understand that the predictive accuracy becomes important in case of real-world applications, so we included a discussion about this aspect in Section 6.3.

“A benefit of IVS is the improvement in the performance of the model being identified. Although the manner in which such performance is characterised depends on the specific domain of interest and the model objectives (Jakeman et al., 2006), two important aspects should always be considered when dealing with quantitative testing. First, the use of observational data for comparison must rely on appropriate data-division methods, such as cross-validation or bootstrapping, that allow for testing the ability of the model to generalise. Data division can account for both temporal and spatial dimensions, so it is suitable for spatial modelling as well (see Chowdhury and Sharma (2009) for an application to hydrological modelling problems). Second, an exhaustive quantitative evaluation should rely on a set of metrics focussing on different aspects in order to test the ability of the model in reproducing all the important features of the system. The reader is referred to Bennett et al. (2013) for a comprehensive review of techniques available for both data-division and quantitative evaluation, and to Robson (2014) for a more general assessment of environmental models.”

Last point - I have not published this yet - but my PMI code also takes into account the change in the uncertainty associated with the predictor variable over time. Again - this was included as typical seasonal forecasting problems have markedly different standard errors depending on when the data was collected. A good example of implications of this changing error on predictions is in Chowdhury, S., and A. Sharma (2007), Mitigating Parameter Bias in Hydrological Modelling due to Uncertainty in Covariates, *Journal of Hydrology*, 340(doi:10.1016/j.jhydrol.2007.04.010), 197-204. But a better example of how these standard errors can be ascertained (varying over space and time, in this case for GCM simulations) is in Woldemeskel, F. M., A. Sharma, B. Sivakumar, and R. Mehrotra (2012), An error estimation method for precipitation and temperature projections for future climates, *Journal of Geophysical Research-Atmospheres*, 117, doi:Artn D22104Doi 10.1029/2012jd018062. Strongly feel predictor identification needs to offer a sensible basis of including such variations in data quality over time. This should be stated somewhere in this paper.

This aspect is discussed in Section 6.3.

“In many practical situations, input variables can be characterised by errors, due, for example, to the interpolation of data in space and time or to the conversion of point measurement into areal values. Whilst methods exist for assessing the impact of input errors on parameter estimation procedures (Chowdhury and Sharma, 2007;

Woldemeskel et al., 2012), IVS algorithms cannot take into account the change in the uncertainty associated with the different inputs.”

On the whole, this is a great paper, that can be quite useful to people who identify predictor variables for use in different prediction problems. Well done folks!

We thank the reviewer for the comment.

Reviewer #2

The authors propose a framework in three points to evaluate input variable selection (IVS) algorithms:

- 1) 26 benchmark synthetic datasets
- 2) a set of evaluation criteria
- 3) website for sharing data and results

Four IVS algorithms are compared and evaluated according to the proposed framework and discussed thoroughly. The idea is really interesting and I think frameworks of this type are more and more developed and are necessary to help research scientist be more systematic in their evaluation and comparison of new and existing methodologies. The paper is generally well written but is very long (91 pages with appendices and 56 pages before the reference section). I think it could be shorten without diminishing its coherence. I will suggest some possible ways to shorten it below.

We understand that the paper is a bit lengthy, so we shortened it by removing some marginal elements (while improving some specific aspects). The revised version is 21 pages shorter (including the appendices).

I have one major disappointment: I could not find the website at the address mentioned on p. 31 (www.ivs4em.deib.polimi.it) only www.deib.polimi.it works but from there, I cannot find the framework webpage. I also tried some Google searches unsuccessfully. I think this is a limitation of the paper, if one does not have access to the benchmark datasets and cannot have a look at the web page, the whole paper remains at the stage of a good idea. Also, I think the paper could include snapshot images of the website to illustrate its functionalities for sharing results for instance. I would also expect the authors to include a functionality in the website which would allow the computation of the recommended criteria automatically. If not directly in the website, some R code could be shared to compute the criteria easily and people could contribute to new criteria.

We fixed this problem. The website is now accessible, and the updated url (<http://ivs4em.deib.polimi.it>) is included in the revised version of the manuscript. From the website it is possible to download the 26 datasets (with their corresponding description), the source code of each IVS algorithm and an R script to compute the evaluation criteria. We have also included a functionality to upload algorithms, datasets and evaluation criteria.

We agree with the reviewer that the paper could include snapshot images of the website; however, we decided not to include them in order to limit the manuscript length.

Other comments

Section 2 describes the background on IVS methods. I found this section both a little long and not so easy to understand. I had to read the Guyon and Elisseeff (2003) paper to understand more clearly the three IVS categories. In particular, filters method are basically ranking methods (as described in Guyon and Elisseeff (2003)) and I think it's more intuitive to present them by mentioning ranks.

Following the reviewer's suggestion, we shortened Section 2 and we clarified all the unclear aspects. Furthermore, we included at the beginning of Section 2.2.1 a brief explanation of filters that explicitly refers to ranking methods.

I would suggest to start the description of each class of IVS by a typical algorithm from this class. This would help to understand the definition of the class. For the filters, the method of ranking in terms of correlation between one input and the output, for instance. For wrapper, the GA-ANN method used in the application of the framework could be described rapidly here. And for embedded algorithms, I would think of the LASSO algorithm which is quite popular.

We agree with the reviewer that a simple description of each class of IVS would simplify the understanding of Section 2.2. Hence, we included a brief explanation of filters, wrappers and embedded algorithms at the beginning of Section 2.2.1, 2.2.2 and 2.2.3, respectively. We think that this approach is more effective than describing a typical algorithm for each class.

P. 10, lines 202-204: I have some trouble to understand why the ACF and PACF are useful for input selection. As far as I know, these techniques are used for time series analyses to choose the proper coefficients in an ARMA model. I would suggest to mention here the partial correlation and partial mutual information which are used in the application of the framework later.

ACF and PACF can be used to measure the (linear) correlation between inputs and output, and then rank the former according to the pairwise correlation. As such, they can be seen as filters. Following the reviewer's suggestion, we also mentioned the Partial Correlation Input Selection algorithm.

P. 12 lines 245-250: the description of the Gamma near-neighbour test was not clear to me. I am wondering if it is useful since this method is not used in the comparison of IVS algorithms and the purpose of the paper is not to review the state-of-the-art on IVS algorithms.

The description of the Gamma near-neighbour test was shortened as suggested.

Section 3 describes the evaluation framework. Basically, as far as I am concerned, two things are missing: some real datasets and a performance criterion based on predictive accuracy. I understand the point made by the authors for the synthetic datasets: it is the only way to know the "true" inputs and their performance criteria SA

are based on this knowledge. However, from a practical point of view, I am, most of the time, mainly interested to evaluate if the model selection I performed yield the best model in terms of predictive power. I think that some real datasets along with a predictive accuracy criterion would be complementary to the framework. This could be similar in spirits with the Delve datasets mentioned by the authors: some datasets are used for development and other for assessment. The real datasets could serve the later goal.

We understand the reviewer's suggestion, but we believe that including some real datasets and one, or more, performance of predictive accuracy may be counterproductive. This opinion is supported by the following reasons: 1) There exists a variety of filters that do not rely on any underlying model (induction or learning algorithm), so it is not possible to evaluate the accuracy of such algorithms in terms of predictive accuracy. This would be against the rationale of the IVS framework, which is aimed at supporting the quantitative (and qualitative) evaluation of any input selection algorithm; 2) The predictive accuracy is 'biased' by several factors, such as the choice of the underlying model and calibration (and validation) algorithm. Minimizing such bias would require introducing an exhaustive comparison of different models (e.g. neural networks, regression trees, linear models, support vector machines etc.) and calibration methods, but this would dramatically affect the length of the manuscript; 3) The same reasoning applies to the inclusion of some real datasets. Indeed, the comparison of different input selection algorithms on some real datasets could only be run by comparing the predictive accuracy of some underlying models; furthermore 4) The inclusion of a few real datasets prevents an exhaustive assessment of the IVS algorithms against the statistical properties described in Section 3.1.

P. 18 lines 399-402. The sentence "Finally, the use of synthetic data enable previously unanalysed datasets ... would provide very little information about algorithm performance" is not clear to me.

The sentence has been removed.

P. 19 line 417 : "a universal approximator", like an artificial neural network ? or the authors have something else in mind ?

Yes, a feed-forward neural network (with a single hidden layer containing a finite number of neurons) could indeed serve as a universal approximator (Cybenko, 1989). We clarified this aspect in the revised version of the manuscript.

"The amount of noise in the output is defined as the fraction of the variance that would remain unexplained if a universal approximator, such as an artificial neural network (Cybenko, 1989), were used on an infinite training set."

Section 3.2.1. Selection accuracy: do we really need SA in addition to SAe and SAc? I find the later two sufficient since SA is computed from them. Moreover, SA requires to set a parameter which controls the tradeoff between SAe and SAc and it seems not necessary to make such a choice.

We believe that the three scores (i.e. SA, SAc and SAe) are important, since they serve two different purposes: 1) The Selection Accuracy (SA) makes the comparison between different algorithms quite fast and straightforward, since it quantifies the degree to which a model has been correctly or incorrectly specified. Furthermore, the presence of the parameter γ allows the user to weight the importance of missing a relevant input against choosing an extraneous one; 2) The SAc and SAe allow for a more in-depth analysis, since they quantify the proportion of correct and extraneous inputs that have been selected.

The single SA score also allows a simple and direct trade-off between selection accuracy and runtime.

Computational efficiency: I have a tendency to think that the total runtime is enough as a measure of computational efficiency. I understand it is not directly comparable across platforms and programming languages but I am not sure if that really matters that much. What basically matters is the order of magnitude: does it take a couple of seconds or a couple of days?

The total runtime provides simple, ‘practical’ information that is certainly useful to most users and practitioners. For this reason, the results in terms of runtime are reported within the text, while the analysis of computational complexity is reported in Appendix C (now Appendix B). Although only few readers may be interested in it, we believe that such analysis can have both theoretical (e.g. determining the growth rate of the runtime) and practical (e.g. planning the execution of several IVS experiments) implications, particularly as enables platform independent comparisons of the computational efficiency of different IVS algorithms. This will become increasingly important as researchers will add the performance of different algorithms to the website, as these measures will enable computational efficiency to be compared in an objective manner.

P. 29 lines 657-658: how does the framework provides a theoretical measure of computational complexity? as far as I know, this has to be computed for each IVS algorithm by considering the computation steps involved. This would be a kind of $O(NP)$ classification for instance, am I right ?

Yes, the theoretical measure of computational complexity is determined for each algorithm by evaluating the computational steps involved at each iteration (see Appendix C). This concept has been further clarified in the revised version of the manuscript.

“In particular, the analysis of computational complexity is determined for each algorithm by evaluating the computational steps involved at each iteration, and it is aimed at producing a theoretical classification that estimates the increase in run-time as a function of the input dimensionality N and P .”

Experimental setup

I found it difficult to follow the explanations on the IVS algorithms and on their performance without further explanations on their mechanisms which are given in the appendix. This is why I am suggesting to use the space in the section 2 to already introduce the IVS algorithms which will be compared.

We understand that Section 4 may appear unclear without reading the appendix, but, at the same time, we think that Section 2 should contain a general description of IVS approaches and not a detailed description of the IVS algorithms adopted in this study. In order to solve this problem, we included a brief description of each algorithm in Section 4, and we tightened the connection between Section 4 and the appendix.

I am wondering if it is useful to include 4 IVS algorithms since this means that all of them should be described in details for the reader to understand what is going on. For instance, p. 32 line 743, I am wondering how the Gaussian reference bandwidth is set and line 749, how do you compute the "correlation between inputs and output and a multiple linear regression".

The presence of four algorithms is critical to demonstrate why the framework can be useful to identify the pros and cons of different types and classes of IVS algorithms. For example, the comparison between PCIS and PMIS shows the effect due to the presence of nonlinearities, while the one between PMIS and IIS is used to discuss the effects of non-Gaussian data. Furthermore, the comparison between filters (PCIS, PMIS and IIS) and wrappers (GA-ANN) allows discussing the computational demands of different methodologies. Limiting the comparison to two algorithms would not allow for this exhaustive analysis. This said, we understand the reviewer's concern, so we clarified all these technical aspects in Section 4 (please refer to the previous reply as well).

It would probably be possible to retain 2 distinct IVS algorithms and to compare them in order to illustrate the framework. The paper would be easier to read then since the goal is not so much to inform on IVS algorithms than to present to framework.

Please refer to the previous reply.

Other questions on IVS algorithms: p. 33 line 755 what are SISO models? I found the explanation later in the appendix. In general, the explanation of the IIS algorithm was fairly obscure to me.

The description of the IIS algorithm has been improved as suggested.

p. 33 lines 767: a 1 hidden unit neural network do not have much non-linear capability. I understand it takes time to tune the number of hidden units of a neural network but otherwise, they do not have much predictive power.

Yes, we totally agree with this remark (which is indeed commented on in Section 6.1). The adoption of such architecture, however, can easily serve our purpose: we aim at practically demonstrating the pros and cons of wrappers (and filters), rather than providing a definitive answer as to which of the algorithms performs best.

p. 33 line 771: since the number of hidden units is fixed, what is the use of k-fold cross-validation?

The k -fold cross-validation is used to quantify the accuracy of the ANN. We clarified this aspect in the revised version of the manuscript.

“The accuracy of the ANN is measured in terms of out-of-sample AIC, computed using a k-fold cross-validation (with k = 5).”

p.36 lines 841-842 : " ... all four combinations of SA_c and SA_e were obtained for the combination of IVS algorithm and datasets..." this sentence needs to be rephrased.

The sentence has been rephrased as suggested.

“Furthermore, Figure 4 shows that different values of SA_c and SA_e were obtained for the combination of IVS algorithms and datasets considered.”

Regarding Figs 4-5 and Figs 6-7, I think they could be re-organized; as it is, they are redundant. The authors could either choose to show the SA scores for datasets which yield contrasted results for the four IVS algorithms or to group datasets according to their properties (as it is done in the text in section 5.1.2).

Following the reviewer’s comment, we removed Figure 4 and 6, since the most of the information in Figure 4 (or 6) is available from Figure 5 (or 7). The reason for maintaining Figure 5 and 7 is that they allow organizing the results by dataset (Figure 5) and by algorithm (Figure 7). The former highlights the performance of the four IVS algorithms on the same modelling conditions, while the latter provides insight into the way different dataset properties impact on the behavior of a specific algorithm.

The discussion on the results could be more condensed: p.44 lines 1021-1033: I found pretty evident that larger N helps model selection, I would suggest to shorten lines 1024-1033.

Section 5.1.3 (‘Effect of *N* and *P* on algorithm performance’) has been revised and shortened. In general, the entire Section 5 has been thoroughly revised and condensed.

Computational efficiency

I found it difficult to follow the discussion on where the computations take more time for each IVS algorithm since I was not very familiar with them. I kept wondering: what is exactly Extra-trees, GRNN, PCIS, IIS... By retaining just 2 IVS algorithms and providing more detailed explanations would probably help to benefit from the kind of discussion in this section.

Section 5.2 gives two different types of information about ‘Computational efficiency’. The first is based on the total runtime (Table 2) and it is built on the concept that filter algorithms (such as PMIS, PCIS and IIS) are computationally efficient, while wrappers (such as the GA-ANN algorithm) require more computing resources. This information is directly accessible by any reader, and does not require being familiar with the algorithms considered. The second information, which is based on the analysis of complexity (Table 3), requires an in-depth knowledge of the algorithms, so we believe that the improvements to Section 4 (plus the presence of a dedicated appendix) will allow the interested readers in understanding the technical aspects of such analysis.

I found the qualitative criteria section quite long. I understand the interest in these type of criteria but it could probably be shortened.

The section was shortened as recommended.

Reviewer #3

The objective of this paper is to create a framework for evaluating and comparing input subset selection (IVS) algorithms for environmental modeling applications. IVS for environmental systems modeling is an extremely challenging task because of the vast number of possible explanatory variables given the space/time correlation of the processes being modeled. However, for the same reason there is also the possibility for significant colinearity of input variables. For this reason IVS is an important first step for any environmental modeling project. Unfortunately, as noted by the authors, there has been little research into what makes a "good" IVS algorithm, as most IVS algorithm research has been focused at a particular dataset or modeling task. The proposed framework would create a repository of data sets and algorithms that would permit comparison of the existing or newly proposed IVS algorithms to identify which perform well in general, thus providing guidance on which algorithm to select for new modeling projects.

This is a well written paper describing project of great interest to the environmental modeling community. The discussion of existing IVS methods is thorough given the scope and length of the paper, and the explanation of the evaluation criteria and the benchmark synthetic data sets is thorough. I recommend this paper be published as-is by Environmental Modelling and Software.

We thank the reviewer for the comment.

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A framework for the evaluation of Input Variable Selection algorithms is proposed.

The framework consists of assessment criteria and twenty-six datasets.

The framework is supported by a dedicated website (<http://ivs4em.deib.polimi.it>).

Four popular IVS algorithms are considered for evaluation purposes.

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An evaluation framework for input variable selection algorithms for environmental data-driven models

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Abstract

Input Variable Selection (IVS) is an essential step in the development of data-driven models and is particularly relevant in environmental modelling. While new methods for identifying important model inputs continue to emerge, each has its own advantages and limitations and no single method is best suited to all datasets and modelling purposes. Rigorous evaluation of new and existing input variable selection methods would allow the effectiveness of these algorithms to be properly identified in various circumstances. However, such evaluations are largely neglected due to the lack of guidelines or precedent to facilitate consistent and standardised assessment. In this paper, a new framework is proposed for the evaluation and inter-comparison of IVS methods which takes into account: (1) a wide range of dataset properties that are relevant to

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9 real world environmental data, (2) assessment criteria selected to highlight
10 algorithm suitability in different situations of interest, and (3) a website for
11 sharing data, algorithms and results (<http://ivs4em.deib.polimi.it/>). The
12 framework is demonstrated on four IVS algorithms commonly used in en-
13 vironmental modelling studies and twenty-six datasets exhibiting different
14 typical properties of environmental data. The main aim at this stage is to
15 demonstrate the application of the proposed evaluation framework, rather
16 than provide a definitive answer as to which of these algorithms has the best
17 overall performance. Nevertheless, the results indicate interesting differences
18 in the algorithms' performance that have not been identified previously.

19 *Keywords:* Input variable selection, Data-driven modelling, Evaluation
20 framework, Large environmental datasets, Artificial neural networks
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33 **1 Software and data availability**

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36 **2 Software**

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- 38 Name of software: PMIS_PCIS, IIS, GA_ANN.
40 Developers (PMIS_PCIS, GA_ANN): Greer B. Humphrey, Holger R. Maier,
42 Graeme C. Dandy, Matthew S. Gibbs.
44 Developers (IIS): Stefano Galelli, Andrea Castelletti.
46 Year first available: 2014.
48 Hardware required: PC or MAC.
50 Software required: R (PMIS_PCIS and GA_ANN), MatLab (IIS).
52 Program language: R (PMIS_PCIS and GA_ANN), MatLab (IIS).
54 Program size: 41 KB (PMIS_PCIS), 135 KB (IIS), 172 KB (GA_ANN).

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12 *Data*

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11 Name of dataset: IVS Framework datasets.
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14 Developers: Greer B. Humphrey.
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16 Form of repository: zipped files.
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18 Size of archive: 239.3 MB.
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20 Access form: public Dropbox folder.
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23 Contact address: Pillar of Engineering Systems and Design, Singapore Uni-
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27 Telephone: + 65 6499 4786.
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29 E-mail: stefano_galelli@sutd.edu.sg.
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31 Url: <http://ivs4em.deib.polimi.it>.
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33 Availability: software and data are available on the IVS framework website.
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35 Cost: free of charge.
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38 **1. Introduction**
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40 In data-driven modelling, such as the application of Artificial Neural Net-
41 works (ANNs), determining which inputs are most useful for predicting a
42 variable of interest can be one of the most critical decisions in the model
43 development process. The input variables (or predictors) contain the infor-
44 mation necessary for defining, albeit, in a simplified manner, the underlying
45 process that generated the data. However, the set of *candidate* inputs usu-
46 ally includes variables which might be either *irrelevant* to the problem or
47 *redundant*. Irrelevant input variables are uninformative about the underly-
48 ing process and only serve to add noise and complexity into the model, while
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36 the inclusion of redundant, but relevant, inputs increases the dimensionality
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37 of the model identification problem without providing any additional predic-
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12 tive benefit. The omission of relevant input variables, on the other hand,
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38 leads to an inaccurate model, where part of the output behaviour remains
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39 unexplained by the selected input variables. Thus, the appropriate selection
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40 of both relevant and non-redundant inputs can mean the difference between
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41 a reliable and parsimonious model, which generalises well to the underlying
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42 process, and a model that produces nonsensical outputs (garbage in, garbage
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43 out), is slower to run, and more difficult to interpret. The challenge of Input
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44 Variable Selection (IVS) is, therefore, to select the fewest input variables that
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45 best characterise the underlying input-output relationship while minimising
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46 variable redundancy (Guyon and Elisseeff, 2003).
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34 While the task of IVS is not unique to environmental modelling, it can
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49 be a particularly difficult one when it comes to environmental systems, since
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50 many of the underlying processes are often partially understood. Further-
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51 more, as environmental systems vary in space and time, potentially important
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52 inputs may include observations of causal variables at different locations and
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53 time lags, as well as lagged observations of the dependent variable of interest
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54 (Maier and Dandy, 2000). As a result, the number of potentially important
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55 inputs can be very large; a problem which has been exacerbated in recent
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56 years by the emergence of new types of data, including remotely sensed, GIS
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57 and reanalysis data. To further complicate matters, the correlated nature of
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58 such input variables induces redundancy and collinearity in the input pool
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59 (Galelli and Castelletti, 2013b), while the non-linearity and inherent com-
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61 plexity associated with environmental systems make it ineffective to apply
10 well established analytical variable selection methods, such as correlation
11 analysis (May et al., 2011). As such, the development and adaptation of
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61 plexity associated with environmental systems make it ineffective to apply
62 well established analytical variable selection methods, such as correlation
63 analysis (May et al., 2011). As such, the development and adaptation of
64 IVS methods for environmental modelling applications is an important and
65 active field of research, which has further been stimulated by reviews of envi-
66 ronmental modelling procedures discussing the need for improved and more
67 rigorous IVS (see, for example, Araújo and Guisan (2006); Elith and Leath-
68 wick (2009); Maier et al. (2010); Abrahart et al. (2012); Wu et al. (2014)).

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70 However, despite recent efforts to improve IVS in environmental mod-
71 elling, studies in this field tend to draw overly general conclusions about the
72 performance of the IVS approaches used. They are usually conducted with a
73 single focus (e.g. to select the inputs for a particular case study of interest)
74 and the evaluation of IVS methods is summarised accordingly (e.g. based on
75 the predictive performance of the resulting model). Such evaluations make
76 it difficult to determine how the performance of one IVS method, either new
77 or existing, compares with that of another, and, ultimately, are of limited
78 value to users wishing to select an IVS method that is most appropriate for
79 a particular problem. As noted by Elshorbagy et al. (2010) in relation to the
80 development of data-driven modelling techniques in hydrology, “one of the
81 fundamental means to assess a modelling technique is to evaluate it against
82 other modelling techniques”, yet “comparative studies are usually impaired
83 due to the less-than-comprehensive approach adopted”. The same can be
84 said about the assessment and comparison of IVS methods, where current
85 studies tend to:

- 86 • select a limited number of data sets which do not adequately encompass
87 the range of properties typical of environmental data (e.g. nonlinear,
88 non-Gaussian, high redundancy);
- 89 • select case studies for which the “true” inputs are unknown and thus
90 do not enable selection accuracy to be properly assessed;
- 91 • consider limited assessment criteria, often based on the predictive per-
92 formance of the constructed model, which is complicated by the chosen
93 functional form of the model and calibration performance;
- 94 • lack rigorous implementation (e.g. no repeated experiments), thus pre-
95 venting the statistical significance of any observed results to be evalu-
96 ated; and
- 97 • only consider a single algorithm without comparison with other algo-
98 rithms.

99 In order to address these shortcomings, a generic framework for the stan-
100 dardised and rigorous comparative analysis of IVS algorithms is introduced
101 in this paper. The framework is comprised of three main components: (1)
102 a set of benchmark data; (2) a recommended set of evaluation criteria; and
103 (3) a website for sharing data and results. The datasets are synthetically
104 generated to have, to different degrees, the typical properties of real envi-
105 ronmental data, while the evaluation criteria are designed to quantitatively
106 and comprehensively assess selection accuracy and computational complex-
107 ity. To demonstrate the application of the framework, four IVS algorithms
108 commonly adopted in environmental modelling exercises and representative

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109 of different IVS approaches are comparatively analysed. It is hoped that this
110 framework will facilitate collaboration by researchers developing new IVS al-
111 gorithms and modellers wishing to select an appropriate IVS method.
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113 The remainder of this paper is structured as follows: Section 2 provides
114 a background on IVS methods, with particular focus on those used to date
115 in environmental modelling studies. In Section 3, the proposed IVS evalua-
116 tion framework is presented, while Section 4 describes the application of the
117 framework to four IVS algorithms. Results of these evaluations and compar-
118 isons are presented in Section 5, while discussion and conclusions are given
119 in Sections 6 and 7.

120 **2. Background on IVS Methods in Environmental Modelling**

121 In recent years, the use of automatic and systematic IVS algorithms has
122 been shown to improve prediction accuracy and produce more parsimonious
123 models in numerous applications when compared with empirical IVS methods
124 or the inclusion of all available input data (e.g. Bowden et al. (2005b);
125 D’heygere et al. (2006); Yang and Ong (2011); Wan Jaafar et al. (2011);
126 Tirelli and Pessani (2011)). Comprehensive discussions on the taxonomy of
127 such IVS methods can be found in Blum and Langley (1997), Liu and Motoda
128 (1998), Guyon and Elisseeff (2003) and May et al. (2011). A brief overview
129 is provided here for the purpose of highlighting the relative differences and
130 merits of the various IVS approaches. Figure 1 is adapted from Dash and
131 Liu (1997), who outline the basic steps of any automatic IVS algorithm. As
132 can be seen, such methods involve three main steps: (1) generating a subset

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10 133 of inputs from the candidate input pool; (2) evaluating the subset of inputs
11 134 in terms of their ability to predict the output; and (3) assessing whether the
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13 135 selected set of inputs is optimal using a pre-specified stopping criterion.
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15 16 136 *2.1. Input subset generation*

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18 137 The generation of input subsets is determined by the method used to
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20 138 search the space of all possible input subsets. An exhaustive search of the
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22 139 space is generally infeasible, as there exist $2^P - 1$ possible subsets of input
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24 140 variables, where P is the dimension of the candidate input pool. Instead,
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26 141 search strategies applied to IVS algorithms seek to balance the trade-off
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28 142 between finding the optimal subset of input variables and computational
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30 143 efficiency. These strategies may be classified as *global*, where many combi-
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32 144 nations of input subsets are considered; or *local*, where the search begins
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34 145 at a defined starting point and moves through the search space sequentially
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36 146 (Maier et al., 2010). For example, local search strategies that begin with an
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38 147 empty input set and successively add individual variables are called *forward*
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40 148 *selection*, while those that start with all possible input variables and succes-
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42 149 sively remove them are known as *backward elimination* (Blum and Langley,
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44 150 1997). Both of these search strategies are *greedy*, in that they make locally
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46 151 optimal decisions with the hope that a globally optimum solution will be
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48 152 found; and once such a decision has been made, it cannot be undone (i.e. an
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50 153 input added (eliminated) in the early stages of the search can not later be
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52 154 eliminated (added)). *Stepwise selection* involves the successive addition or
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54 155 elimination of input variables, but allows an earlier decision to be retracted,
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56 156 potentially allowing more optimal subsets to be identified. However, deci-
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10 158 on the already selected inputs. *Random* or *probabilistic* search strategies are
11 159 more adept at finding (near) globally optimum input subsets through their
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13 160 combined use of random subset generation with some mechanism to increase
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15 161 the focus of the search in regions of the search space that lead to good solu-
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17 162 tions. However, due to their random nature, these strategies search through
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19 163 many more solutions than their sequential counterparts and are, thus, less
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21 164 efficient than sequential search algorithms (Kohavi and John, 1997), yet still
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23 165 provide no guarantee that a globally optimal solution will be found.
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25 166 *2.2. Input subset evaluation*

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28 167 The evaluation step in Figure 1 involves determining which inputs should
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30 168 be added to the ‘selected’ input set and which should be discarded, based
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32 169 on their relevance. Automatic IVS algorithms can be broadly categorised as
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34 170 *filter*, *wrapper* or *embedded* approaches according to the way in which this
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36 171 input relevance is measured (Guyon and Elisseeff, 2003). Filter IVS methods
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38 172 are described as being *model-free*, as the entire IVS process is independent of
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40 173 the chosen induction or learning algorithm. Both embedded and wrapper IVS
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42 174 approaches, on the other hand, are *model-based*, relying on the performance
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44 175 of a predetermined underlying model to select the most appropriate inputs.
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46 176 *2.2.1. Filter algorithms*

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48 177 Filter techniques rely on the intrinsic properties of the data (e.g. dis-
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50 178 tance, information, dependency, or consistency) to measure the relevance of
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52 179 the input variables, which are then ranked according to some a-priori defined
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54 180 criteria (Liu and Motoda, 1998). As such, filters tend to be computationally
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56 181 simple and scale easily to high-dimensional datasets. However, as filters are
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10 182 independent of the learning algorithm, they have the disadvantage of disre-
11 183 garding how the selected variable subset will affect the performance of the
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13 184 resulting model (Miller, 2002). In addition, they are typically univariate,
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15 185 which has the disadvantage that the relevance between each potential input
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17 186 and the output variable is considered separately (Saeys et al., 2007). Not
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19 187 only does this necessitate that each input-output relationship be evaluated,
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21 188 but it also means that input variable interactions are ignored. Consequently,
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23 189 an input may be found individually to be irrelevant when, in fact, it is very
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25 190 relevant when combined with other inputs.
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29 192 Linear correlation-based filter algorithms are among the most commonly
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31 193 used IVS methods in environmental studies (see Maier et al. (2010) and
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33 194 Wu et al. (2014) for a review). An example of such an approach is used in
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35 195 the popular Box-Jenkins time-series analysis methodology (Box and Jenk-
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37 196 ins, 1976), where identification of the most important auto-regressive and
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39 197 moving-average parameters is based on the autocorrelation and partial au-
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41 198 tocorrelation function. Another popular linear correlation-based approach
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43 199 is the Partial Correlation Input Selection (PCIS) introduced by May et al.
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45 200 (2008a). Yet, despite their popularity and simplicity, such methods are likely
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47 201 to be inappropriate for nonlinear systems.

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49 202 In recent years, information theoretic-based dependency measures, such as
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51 203 mutual information, have become more popular in IVS, since such measures
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53 204 make no assumptions regarding the structure of the dependence between
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55 205 two variables (i.e. they can estimate both linear and nonlinear dependence)
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57 206 (May et al., 2008a). For example, the Partial Mutual Information (PMI)

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207 based IVS method developed by Sharma (2000) and modified by Bowden
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208 et al. (2005a); May et al. (2008a); Fernando et al. (2009) has been applied in
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209 several studies for identifying the most relevant inputs for predicting rainfall
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210 (Sharma et al., 2000), streamflow (Wu et al., 2013), salinity (Bowden et al.,
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211 2005b; Fernando et al., 2009), water quality (Kingston et al., 2006; May et al.,
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212 2008b) and stormwater runoff (He et al., 2011). Other PMI-based metrics
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213 have been recently proposed by Chen et al. (2013) and Sharma and Mehrotra
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214 (2014). Another well established filter IVS technique based on mutual infor-
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215 mation is the minimum redundancy maximum relevance (mRMR) algorithm
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216 developed by Peng et al. (2005). A more computationally efficient version
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217 of this algorithm was proposed by Hejazi and Cai (2009), who applied it to
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218 selecting the most significant inputs to predict daily reservoir releases. Re-
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219 cently, Galelli and Castelletti (2013b) proposed the Iterative Input variable
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220 Selection (IIS) algorithm, where a tree-based ranking method is used in place
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221 of an information-theoretic measure to estimate the information gained from
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222 the data. This algorithm has been employed to select the most relevant in-
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223 put variables for daily streamflow prediction (Galelli and Castelletti, 2013b),
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224 prediction of phytoplankton biovolume (Fornarelli et al., 2013), prediction of
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225 spatially distributed hydro-ecological data (SurrIDGE et al., 2014) and model
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226 reduction problems (Castelletti et al., 2012).
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47 Another popular filter method is the Gamma (or near-neighbour) test (Končar,
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49 1997; Stefánsson et al., 1997), which uses distance, rather than variable de-
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51 pendence or information gain, in the evaluation of input relevance. This
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53 method was first employed by Chuzhanova et al. (1998) and has recently
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55 become more popular for IVS in the field of environmental modelling. The
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10 232 Gamma test has been used, for example, to select the best inputs for solar
11 233 radiation prediction (Remesan et al., 2008; Ahmadi et al., 2009), runoff mod-
12 234 elling (Remesan et al., 2009), flood regionalisation (Wan Jaafar et al., 2011;
13 235 Wan Jaafar and Han, 2012) and downscaling climate variables for precipita-
14 236 tion forecasting (Ahmadi and Han, 2013).

237 2.2.2. *Wrapper algorithms*

238 Wrapper methods use the learning algorithm itself as part of the IVS
239 procedure, treating the model as a black box, while searching for the subset
240 of inputs that yields the best model performance (Kohavi and John, 1997).
241 Unlike filter methods, wrapper approaches take into account interactions
242 and dependencies between input variables. However, since the learning al-
243 gorithm must be called (and calibrated) for each input subset considered,
244 wrapper methods can be very computationally intensive (Blum and Lan-
245 gley, 1997). They are also more susceptible to overfitting than filters, as
246 most of these approaches focus on finding inputs that maximise predictive
247 performance, rather than those that are both relevant and nonredundant.
248 Consequently, it is particularly important when employing wrapper IVS al-
249 gorithms to adopt an objective function or optimality criterion that penalises
250 model complexity and, hence, overfitting. Wrapper IVS methods tend to be
251 defined by the search strategy employed to generate input subsets. By far
252 the most commonly used wrapper IVS methods in environmental modelling
253 studies are those that involve the sequential (forward, backward or stepwise)
254 selection of inputs (see Olden and Jackson (2000), Mac Nally (2000) and
255 Ssegane et al. (2012) and references therein). In recognising the limitations
256 of sequential search techniques, a number of relatively recent studies have

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10 257 employed random search strategies, such as evolutionary algorithms. For ex-
11 258 ample, Abrahart et al. (1999); Schleiter et al. (2001); Bowden et al. (2005b);
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13 259 D’heygere et al. (2006) and Tirelli et al. (2009) used a genetic algorithm to
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15 260 select the best inputs for rainfall-runoff modelling, water quality modelling
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17 261 and the prediction of species presence/absence using ANNs and decision trees
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19 262 as the induction algorithms.
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21 263 *2.2.3. Embedded algorithms*

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23 264 In embedded IVS techniques, the search for an optimal subset of inputs
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25 265 and calibration of the underlying model occurs concurrently. Thus, the en-
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27 266 tire IVS process is part of the model training procedure. The basic principle
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29 267 behind embedded IVS algorithms is to specify an objective function for con-
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31 268 structing a model consisting of both a goodness-of-fit term and a term that
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33 269 penalises model complexity (Guyon and Elisseff, 2003). Similar to wrap-
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35 270 per methods, embedded techniques account for interactions between inputs
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37 271 and are specific to the chosen learning algorithm, meaning that they can
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39 272 yield high prediction accuracy (the inputs selected will be those that op-
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41 273 timise model performance), but at the cost of decreased generalisation on
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43 274 other learning algorithms (Guyon et al., 2006). Unlike wrapper methods,
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45 275 however, only a single model is trained, since the evaluation of input subsets
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47 276 occurs within the training algorithm. Thus, embedded methods are usually
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49 277 far less computationally intensive than wrapper methods (Guyon and Elis-
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51 278 seeff, 2003). Furthermore, embedded algorithms consider the impact of each
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53 279 individual input on the performance of the model, and adjust the associated
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55 280 model parameters accordingly. A disadvantage of embedded IVS approaches
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57 281 is the lack of algorithms available for directly minimising the number of input
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10 282 variables for nonlinear predictors (Guyon and Elisseeff, 2003). Embedded al-
11 283 gorithms often rely on regularisation methods, which balance model fit and
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13 284 model complexity during the calibration of a model. Using such methods, the
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15 285 penalty term in the objective function is replaced by a regularisation term,
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17 286 which shrinks parameters associated with irrelevant inputs toward zero or sets
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19 287 them equal to zero (Tikka, 2009). There are a number of available regularisa-
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21 288 tion methods, which differ mainly in the way model complexity is measured
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23 289 and, hence, penalised. Ridge regression (Hoerl and Kennard, 1970) and the
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25 290 Lasso algorithm (least absolute shrinkage and selection operator, Tibshirani
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27 291 (1996)) are among the most popular. For an application of such methods to
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29 292 environmental modelling problems, see, for example, Reineking and Schröder
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31 293 (2006); Phatak et al. (2011); Dormann et al. (2013).

32 33 294 *2.3. Stopping criterion*

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35 295 The definition of a suitable stopping criterion is another key consideration
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37 296 in IVS as it can significantly influence selection accuracy and computational
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39 297 efficiency. Stopping criteria may be related to either the search strategy or
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41 298 the evaluation method used in the IVS process. For example, stopping crite-
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43 299 ria related to the search strategy may include whether a predefined number of
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45 300 relevant variables has been selected or whether a predefined number of itera-
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47 301 tions has been reached (Dash and Liu, 1997). Stopping criteria based on the
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49 302 evaluation of input subsets may include whether the addition or elimination
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51 303 of any inputs produces a better (or worse) subset (using, for example, cross-
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53 304 validation error or parsimonious model selection criteria, such as Akaike's
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55 305 information criterion (AIC) or the Bayesian information criterion (BIC)),
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57 306 or whether selected inputs are relevant, as defined by some threshold value

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307 or significance level (using classical statistical tests, such as the t-test, F-
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308 test and chi-squared test or resampling methods such as bootstrapping, for
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13 309 example) (Dash and Liu, 1997).
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310 **3. IVS Evaluation Framework**

311 The IVS evaluation framework proposed in this paper is designed such
312 that it will be generally applicable to all IVS algorithms, producing easy
313 to interpret and unbiased results and minimising any duplication of effort.
314 As well as aiding comparative analyses of IVS approaches, it should also be
315 useful for investigating parameter effects within individual IVS algorithms or
316 selecting appropriate stopping criteria. As mentioned previously, the basic
317 framework is comprised of three main components: (1) a set of benchmark
318 data; (2) a recommended set of evaluation criteria; and (3) a website for
319 sharing data and results. These are represented in Figure 2 and discussed in
320 detail in the following sections.
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321 *3.1. Benchmark datasets*

322 A total of 26 synthetic datasets, summarised in Table 1, were generated
323 for benchmarking the performance of IVS algorithms. These datasets ex-
324 hibit, to different degrees, the following properties, which are considered to
325 reflect the features of real environmental data: nonlinearity in the underlying
326 function, collinearity amongst input variables, non-Gaussian input/output
327 variables, noise in the output, incomplete input information, and interde-
328 pendence of input variables. The benchmark datasets were generated to
329 have different sample sizes and dimensionalities to allow scalability and com-
330 putational efficiency to be assessed on datasets of different sizes. This also
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9 331 enables an investigation of the sensitivities of IVS methods to the relative
10 332 proportion of irrelevant candidate inputs and of the abilities of IVS methods
11 333 to identify important input-output relationships within datasets of varying
12 334 lengths. In Table 1, sample size is denoted by N , K is the number of relevant
13 335 inputs (those that contain important and non-redundant information about
14 336 the output) and P is the total number of candidate inputs (the total pool of
15 337 potentially relevant inputs from which to select from). The $P - K$ candidate
16 338 inputs that are included in the datasets but contain no (or only redundant)
17 339 information about the outputs are primarily lagged values of the true inputs
18 340 or inputs drawn from distributions resembling those of the true inputs. The
19 341 ratio N/P is also given in Table 1, as this value is indicative of the risk of
20 342 retaining irrelevant or redundant inputs (a small value of N/P suggests a
21 343 greater likelihood of overfitting). This risk increases with increasing correla-
22 344 tion between the candidate inputs.
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346 While synthetic data may be considered somewhat unrealistic and lack-
347 ing in substance, their use for IVS algorithm benchmarking is necessary since
348 such data provide the only means for adequately assessing the performance
349 of IVS algorithms using quantitative approaches. Firstly, and most impor-
350 tantly, the use of synthetic data enables selected inputs to be compared to
351 the known set of “true” input variables. This allows ‘selection accuracy’ to be
352 evaluated without relying on prediction accuracy, which can be complicated
353 by a number of factors, including the choice of model, calibration method,
354 error model and calibration criteria, among others. Secondly, with synthetic
355 data it is relatively easy to systematically vary features such as those listed

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10 356 above in order to achieve a balanced design for the comprehensive evaluation
11 357 of IVS techniques. With real data this would be far more difficult and would
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13 358 rely on methods for quantifying the above properties without knowledge of
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15 359 the true underlying function. While synthetic data may be somewhat sim-
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17 360 plistic, it would be reasonable to assume that an IVS method which fails to
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19 361 select the correct inputs from data generated from a rather simplistic model
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21 362 would be unlikely to have good selection accuracy when applied to real data.
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23 363 However, in order to ensure that the true characteristics of real environmen-
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25 364 tal data were captured in the benchmark datasets at least to some extent,
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27 365 several of the benchmark sets are only partially synthetic, where the input
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29 366 data are real, while only the outputs are modelled. Whether a benchmark
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31 367 dataset is fully or partially synthetic is also indicated in Table 1.

32 33 368 *3.1.1. Properties of the datasets*

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35 369 To define the degrees of noise and nonlinearity associated with the bench-
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37 370 mark data, the following DELVE (Rasmussen et al., 1996) definitions were
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39 371 used:

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41 372 • Noise: The amount of noise in the output is defined as the fraction of
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43 373 the variance that would remain unexplained if a universal approxima-
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45 374 tor, such as an artificial neural network (Cybenko, 1989), were used on
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47 375 an infinite training set. If this residual variance is less than 0.25% the
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49 376 noise is “low”. If it lies between 1% and 5% the noise is “moderate”.
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51 377 If it exceeds 25% the noise is “high”.
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53 378 • Nonlinearity: A dataset is classified as “fairly linear” if a linear method
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55 379 would leave less than 5% residual variance on an infinite training set.

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380 It is “highly non-linear” if the linear method would leave more than
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11 381 40% residual variance.
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14 382 The degree of collinearity was simply defined according to the number of
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16 383 pairs of candidate inputs with correlation greater than 0.7. This is similar
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18 384 to the definition Amasyali and Ersoy (2009) used for defining the degrees
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20 385 of collinearity associated with their Friedman datasets, which they subse-
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22 386 quently donated to the WEKA project. For the purposes of the proposed
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24 387 IVS framework, a dataset is considered to have high collinearity if the num-
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26 388 ber of pairs of candidate inputs with correlation > 0.7 divided by the total
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28 389 number P of candidates is greater than or equal to one, i.e. there are at least
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30 390 as many pairs of highly correlated inputs as there are candidate inputs.
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33 392 A number of common synthetic data generators have been used for eval-
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35 393 uating IVS algorithms in previous environmental modelling studies (see, for
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37 394 example, Sharma (2000); Bowden et al. (2005a); May et al. (2008a); Hejazi
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39 395 and Cai (2009); Fernando et al. (2009) and Galelli and Castelletti (2013b)).
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41 396 These include two linear auto-regressive (AR) models, two nonlinear thresh-
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43 397 old autoregressive (TAR) models and a nonlinear (NL) model. For consis-
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45 398 tency, these test problems have been included within this framework. As
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47 399 can be seen in Table 1, the datasets corresponding to these test problems
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49 400 only cover a small subset of the possible combinations of the first four binary
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51 401 dataset properties (i.e. non-Gaussian outputs, nonlinearity, collinearity and
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53 402 noise). Furthermore, N , K and P are fairly uniform amongst these datasets.
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55 403 The extended set of benchmark datasets covers all possible combinations of
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57 404 these dataset properties and includes much greater variation in the size and
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9 405 dimensionality of the datasets. It also covers two special cases where there is
10 406 incomplete information about the target data and where there is interdepen-
11 407 dence of inputs (i.e. there exist inputs that are only relevant when combined
12 408 with other inputs). To account for any variability in algorithm performance
13 409 that may result from variability in the data, 30 replicates of each benchmark
14 410 dataset are provided. This enables the statistical significance of comparison
15 411 results to be considered. These datasets are described in further detail on
16 412 the framework website.

25 413 *3.2. Evaluation criteria*

27 414 There are a number of different factors to consider when evaluating and
28 415 comparing IVS methods including, for example, accuracy; efficiency; scala-
29 416 bility and ease of use. However, objective metrics that enable the general and
30 417 standardised intercomparison of IVS methods are not necessarily straightfor-
31 418 ward to define. Firstly, not all of the criteria that are useful for evaluating
32 419 IVS algorithms can be expressed as quantitative, objective metrics; some
33 420 can be also qualitative. Secondly, it is important that the number of met-
34 421 rics used for intercomparisons be minimised, while the information gained
35 422 from them is maximised. However, the metrics used for evaluation and com-
36 423 parison must also provide sufficient information such that any differences in
37 424 algorithm performance are discernible. Thirdly, the metrics should be sim-
38 425 ple, easy to compute and interpret and have general applicability across a
39 426 wide range of different IVS methods. Individual IVS methods may produce
40 427 information specific to those techniques that can be useful for diagnosing
41 428 algorithm performance. However, such information will have limited value
42 429 when compared with algorithms that do not output the same information.

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430 Finally, it is preferable that the metrics can be expressed probabilistically,
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11 such that the stability and robustness of algorithms can be assessed and the
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13 432 statistical significance of results computed.

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15 433 As mentioned, the majority of IVS algorithms seek to balance the trade-off
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17 434 between finding the optimal subset of input variables and computational effi-
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19 435 ciency. As such, it is important to be able to evaluate algorithms against these
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21 436 criteria in a quantitative and objective way. Details of the proposed quanti-
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23 437 tative performance measures are given in Section 3.2.1. In contrast, criteria
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25 438 related to an algorithm’s ease of use, flexibility or explanation capability, for
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27 439 example, are more difficult to define in such a manner and therefore it is rec-
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29 440 ommended that these be treated as qualitative evaluation criteria. Details of
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31 441 some of these qualitative criteria are given in Section 3.2.2. When assessing
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33 442 and comparing the performance of IVS algorithms, it is recommended that all
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35 443 of the proposed quantitative metrics be used, while the proposed qualitative
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37 444 criteria should be considered and remarked upon where appropriate.

445 3.2.1. Quantitative metrics

446 **Selection accuracy**

447 A selection accuracy (SA) score which expresses the degree to which a
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45 448 selected input subset matches the true input subset is recommended for use
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47 449 in this framework. The proposed SA score is based on the similarity score
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49 450 proposed by Molina et al. (2002), but unlike the original version, it makes
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51 451 no distinction between irrelevant and redundant inputs and simply treats all
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53 452 unnecessary inputs as extraneous. The proposed SA score is given as follows:

$$SA = \gamma \frac{k}{K} + (1 - \gamma) \left(1 - \frac{p}{P - K} \right) \quad (1)$$

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10 453 where K is the total number of relevant inputs; k is the number of relevant
11 454 inputs selected; p is the number of extraneous (irrelevant or redundant) in-
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13 455 puts selected; P is the total number of inputs in the candidate input pool
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15 456 (hence $P - K$ is the total number of extraneous inputs) and γ is a weight
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17 457 ranging from 0 to 1, which influences the penalty applied to the selection
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19 458 of extraneous inputs in relation to the gain achieved from each correctly se-
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21 459 lected input. This score can range from 0 to 1, where $SA = 1$ corresponds to
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23 460 a correctly specified model, while $SA = 0$ corresponds to a completely mis-
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25 461 specified model, with no relevant inputs and all extraneous inputs selected.
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27 462 An advantage of this score is that information about the degree to which a
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29 463 model has been correctly or incorrectly specified is combined into a single
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31 464 metric, which makes for the straightforward comparison of IVS algorithm
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33 465 selection accuracy.

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36 467 The SA score requires the choice of an appropriate value for γ . This
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38 468 choice is subjective and depends on how much one favours accuracy over
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40 469 parsimony, or vice versa. As suggested by Molina et al. (2002), a suitable
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42 470 value for γ should reflect the fact that choosing an extraneous input is usu-
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44 471 ally better than missing a relevant one, which can be achieved by selecting
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46 472 γ such that $\gamma/K > (1 - \gamma)/(P - K)$. However, γ should not be so large
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48 473 that there is no appreciable penalty applied to unnecessary model complex-
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50 474 ity (for example, for $\gamma = 1$, the selection of extraneous inputs would not
51
52 475 be penalised at all). Figure 3 illustrates the effect of γ on the SA score
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54 476 for a theoretical example with 10 inputs in the candidate input pool: 5
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56 477 relevant and 5 irrelevant (or redundant). As can be seen in Figure 3 (a),

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 9 when $\gamma = 0.5$ (i.e. $\gamma/K = (1 - \gamma)/(P - K)$), the penalty incurred for the
 10 selection of extraneous inputs is weighted equally to any improvement in
 11 accuracy gained from the selection of relevant inputs (as evidenced by the
 12 lack of variation in the SA score in the diagonal direction). This would
 13 usually be undesirable given that, in terms of prediction accuracy, the conse-
 14 quences of under-specification (greater bias) are generally more severe than
 15 those of over-specification (greater variance). Conversely, when $\gamma = 0.9$ (i.e.
 16 $\gamma/K \gg (1 - \gamma)/(P - K)$), there is very little reduction in the SA score
 17 for increasing values of p , as can be seen in Figure 3 (c), indicating that un-
 18 necessary complexity is under penalised. As shown in Figure 3 (b), a value
 19 of $\gamma = 0.7$ results in the selection of extraneous inputs being appreciably
 20 penalised; however, missing a relevant input is assigned greater importance
 21 than the selection of an irrelevant or redundant input (as evidenced by the
 22 variability in the SA score in both the vertical and diagonal directions). For
 23 all of the benchmark datasets included in the proposed IVS evaluation frame-
 24 work, a value of $\gamma = 0.7$ satisfies $\gamma/K > (1 - \gamma)/(P - K)$, while still being
 25 sufficiently less than 1 to appropriately penalise unnecessary complexity.
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 496 While values of $SA < 1$ denote over- or under-specification, a limitation
 497 of the SA score is that it does not indicate where the selected input subset
 498 is deficient; for example, whether too many or too few inputs have been
 499 selected. To overcome this limitation, the SA score given by eq. 1 can be
 500 broken into two sub-scores:

$$SA_c = \frac{k}{K} \tag{2}$$

$$SA_e = 1 - \frac{p}{P - K} \tag{3}$$

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10 502 where SA_c indicates the proportion of correct inputs that have been selected
11 503 and SA_e is based on the proportion of extraneous inputs that has been se-
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13 504 lected. Unlike the overall SA score given by eq. 1, these sub-scores do not
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15 505 trade off one measure of accuracy against another; therefore, they do not
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17 506 require the γ parameter. Both of these terms can range from 0 to 1, where a
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19 507 value closer to 1 denotes a better model. In particular, the following combina-
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21 508 tions of SA_c and SA_e are relevant to the analysis of algorithm performance:

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24 509 • $SA_c = 1$ and $SA_e = 1$, i.e. perfect specification ($SA = 1$);
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27 510 • $SA_c = 1$ and $SA_e < 1$, i.e. over-specification of some extraneous inputs
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29 511 ($SA < 1$);
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32 512 • $SA_c < 1$ and $SA_e = 1$, i.e. under-specification of relevant inputs
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34 513 ($SA < 1$) (according to the definitions used by May et al. (2008a) and
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36 514 Galelli and Castelletti (2013b));
- 37
38 515 • $SA_c < 1$ and $SA_e < 1$, i.e. under-specification of relevant inputs and
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40 516 over-specification of some extraneous inputs ($SA < 1$).

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43 517 The advantage of these scores is that they express the degree to which
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45 518 a model is over- or under-specified, which is important for differentiating
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47 519 between IVS algorithm results. Furthermore, the sub-scores can be useful
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49 520 for investigating parameter effects within individual IVS algorithms. For ex-
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51 521 ample, if a method consistently results in over- or under-specification over a
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53 522 range of different datasets, this may signify that the stopping criterion used
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55 523 to terminate the IVS method is inappropriately penalising model complexity
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57 524 or that the threshold or significance level used to determine the relevance of

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9 525 an input has been inappropriately set or computed.
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12 13 527 **Computational efficiency**

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15 528 Under the proposed framework, two quantitative measures of computa-
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17 529 tional efficiency are recommended. The first is total run-time, namely the
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19 530 time required by an IVS algorithm to perform an input selection task. This
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21 531 metric provides a rough estimate of the time it may take to execute a particu-
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23 532 lar algorithm on a given dataset, but depends on the software implementation
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25 533 and the adopted hardware. For this reason, the framework also includes a
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27 534 thorough analysis of computational complexity, which provides a theoretical,
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29 535 platform-independent estimate of the resources needed by an IVS algorithm.
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31 536 In particular, the analysis of computational complexity is determined for each
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33 537 algorithm by evaluating the computational steps involved at each iteration,
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35 538 and is aimed at producing a theoretical classification that estimates the in-
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37 539 crease in run-time as a function of the input dimensionality N and P . This
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39 540 classification allows calculation of the time that would be required by an IVS
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41 541 algorithm to perform a certain task, and is thus useful when planning the
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43 542 execution of several IVS experiments. Moreover, it allows calculation of the
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45 543 growth rate of the run-time for the worst case scenario (for example, when a
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47 544 forward selection algorithm is run over P iterations to evaluate all candidate
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49 545 inputs).

50 51 546 *3.2.2. Qualitative criteria*

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53 547 Where appropriate, it is suggested that the following qualitative assess-
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55 548 ment criteria be commented on when evaluating IVS algorithms; however, it
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57 549 is not recommended that they be used in the intercomparison of algorithm

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550 performance.

551 1. Ease of use and robustness

552 The ease of use of an IVS algorithm relates to how many parameters
553 need to be tuned and how robust the algorithm’s performance is, given
554 a default set of parameter values. An IVS algorithm that can be applied
555 without significant user expertise can be highly desirable, particularly
556 for a potential user trying to select the most appropriate IVS algorithm
557 for a problem at hand. Therefore, where possible, it is recommended
558 that some information be provided about which parameters affect the
559 performance of the algorithm and how readily robust values can be
560 selected for these parameters.

561 2. Explanation capability

562 Forward selection IVS methods and algorithms that utilise an input
563 ranking approach provide information about the order of input rele-
564 vance (i.e. inputs are sorted from most to least relevant) and possibly
565 the relative magnitude of the influence these inputs have on the out-
566 put. Such information can provide useful insight into the underlying
567 mechanisms by which the data were generated and can be said to have
568 some explanation capability. On the other hand, methods that evalu-
569 ate the relevance of an input subset as a whole generally do not provide
570 information about the relevance of individual inputs. Thus, while such
571 algorithms may return the optimum input subset for a particular prob-
572 lem, it may be difficult to determine how the individual inputs relate
573 to the output.

574 3. Flexibility

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10 575 An IVS algorithm represents a single combination of the three main
11 576 components shown in Figure 1. However, if it is found that the perfor-
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13 577 mance of an algorithm is limited by only one of these components, it
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15 578 would be advantageous if this component could be easily substituted
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17 579 with an alternative. The flexibility of an IVS algorithm relates to the
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19 580 ease with which components of the algorithm can be interchanged with
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21 581 other methods to suit user preferences or to overcome identified short-
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23 582 comings.

24 25 583 *3.3. Framework website*

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27 584 The website (<http://ivs4em.deib.polimi.it>) is an ‘open platform’ for shar-
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29 585 ing datasets, code and results. At the current stage, it contains all 26 bench-
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31 586 mark datasets (30 replicates of each), the source code for the four IVS algo-
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33 587 rithms used in this study and the code for performance evaluation. Moreover,
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35 588 the website includes a functionality for uploading new datasets, algorithms
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37 589 and results to build up a comprehensive database for IVS in environmental
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39 590 modelling.

40 41 42 591 **4. Experimental setup**

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45 592 The proposed framework was used for the evaluation and comparison of
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47 593 four IVS algorithms. The aim was not to provide a definitive answer as
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49 594 to which of the algorithms performed best, but rather to demonstrate the
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51 595 application of the proposed framework and how the results obtained may be
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53 596 used for evaluating and gaining greater insight into algorithm performance.
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55 597 The family of filter methods is represented by the PMI-based input Selection
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57 598 (PMIS) algorithm (in the form modified by May et al. (2008a)), the IIS

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599 algorithm (Galelli and Castelletti, 2013b) and the PCIS algorithm introduced
10 by May et al. (2008a). The family of wrapper methods is represented by a
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13 601 GA-ANN algorithm, which adopts a Genetic Algorithm (GA) to select the
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15 602 subset of input variables that maximises the performance of an ANN. Each of
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17 603 the four IVS algorithms considered was implemented on 30 replicates of the
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19 604 26 benchmark datasets, resulting in 780 runs for each algorithm. For all four
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21 605 algorithms, the same parameter sets were used for all case studies. Details of
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23 606 the parameters used and how their values were determined are given below
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25 607 (the reader is referred to Appendix A for a more detailed description of the
26
27 608 algorithms):

- 29 • *PMIS and PCIS algorithms.* The PMIS algorithm adopts a forward
30 selection strategy (i.e. one variable is selected at each iteration) based
31
32 610 on the estimation of the PMI, which measures the partial dependence
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34 611 between each input variable and the output, conditional on the in-
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36 612 puts that have already been selected. To estimate the PMI, the algo-
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38 613 rithm adopts a kernel density estimator, whose accuracy depends on the
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40 614 value of a smoothing parameter (or bandwidth) λ . Similarly to Sharma
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42 615 (2000) and Bowden et al. (2005a), the Gaussian reference bandwidth
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44 616 (Scott, 1992) is adopted, because of its simplicity and computational
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46 617 efficiency. The calculation of PMI also requires estimation of residual
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48 618 information in the input variables once the effect of the already selected
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50 619 inputs has been taken in consideration: this is done through the iden-
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52 620 tification of a General Regression Neural Network (GRNN), which is
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54 621 a nonlinear and nonparametric regression method (Li et al., 2014). In
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56 622 addition to λ , the other parameter to be set is the stopping criterion,
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624 which is based on the coefficient of determination R^2 : the PMIS algo-
625 rithm is stopped when the selection of a further input variable leads to
626 a decrease of R^2 in the underlying GRNN being identified.

627 The PCIS algorithm adopts the same structure as the PMIS algorithm,
628 but it uses the Pearson correlation coefficient to estimate the strength
629 of the relationship between inputs and output and a multiple linear re-
630 gression based on least squares in place of PMI and GRNN, respectively.
631 As such, the algorithm does not require any tuning. PCIS is terminated
632 when the selection of additional inputs no longer results in an improve-
633 ment (increase) in the Bayesian information criterion (BIC), calculated
634 based on the output variable residual, which provides a trade-off be-
635 tween goodness-of-fit and model complexity.

- 636 • *IIS algorithm.* Similarly to PMIS and PCIS, the IIS algorithm pro-
637 ceeds by selecting one input variable at each iteration, but the partial
638 dependence between each input variable and the output relies on a
639 tree-based ranking method, instead of an information-theoretic mea-
640 sure. Furthermore, the relative importance of the p ranked variables
641 is refined through the identification of p Single-Input Single-Output
642 (SISO) models, with the best performing input being added to the set
643 of selected variables. The selection process continues until the accu-
644 racy of an underlying Multi-Input Single-Output (MISO) model, eval-
645 uated with a k -fold cross-validation, does not significantly improve.
646 The number p and k of SISO models evaluated at each iteration and
647 of the folds used in the k -fold cross-validation process is set to 5, while
648 the algorithm tolerance ε is equal to 0.01, which was empirically found

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10 649 to provide an appropriate balance between accuracy and over fitting
11 650 (Galelli and Castelletti, 2013b). This means that the IIS algorithm is
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13 651 stopped when the selection of a further variable leads to an increase
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15 652 of R^2 lower than 0.01. Extra-Trees, a regression based method intro-
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17 653 duced by Geurts et al. (2006), are used for both ranking and modelling.
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19 654 As for the Extra-Trees setting, default values for the parameters M ,
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21 655 K and n_{min} are set according to Galelli and Castelletti (2013a). The
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23 656 number M of trees in an ensemble is 500, the number K of alternative
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25 657 cut-directions is equal to the number of candidate inputs and n_{min} , the
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27 658 minimum cardinality for splitting a node, is 5.

- 29 659 • *GA-ANN algorithm.* The ANN is a 1-hidden node multilayer percep-
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31 660 tron, with the transfer functions used at the hidden and output nodes
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33 661 being the hyperbolic tangent and linear functions, respectively, which
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35 662 are commonly adopted in environmental modelling problems (Maier
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37 663 and Dandy, 2000). The accuracy of the ANN is measured in terms
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39 664 of out-of-sample AIC, computed using a k -fold cross-validation (with
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41 665 $k = 5$). As for the GA algorithm, the population size and maximum
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43 666 number of generations are equal to 50 and 100,000, respectively. The
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45 667 algorithm is terminated based either on the maximum number of eval-
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47 668 uations or convergence of the fitness function (i.e. when the difference
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49 669 in the fitness between one generation and the next remains below a
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51 670 tolerance of 10^{-8} times the previous best value for 20 consecutive gen-
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53 671 erations).

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55 672 All experiments for PMIS, GA-ANN and PCIS are carried out in the R
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57 673 environment running on 12-core 2.6 GHz CPUs AMD with 2.7 GB RAM

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674 per core, while the experiments for the IIS algorithm are carried out using a
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11 675 compiled C++ package running on 8-core 2.2 GHz CPUs Intel Xeon with 8
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13 676 GB RAM per core.

16 677 **5. Results**

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19 678 The results obtained by evaluating the four IVS algorithms with the pro-
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21 679 posed framework are presented and discussed in this section, and organised in
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23 680 terms of selection accuracy, computational efficiency and qualitative criteria
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25 681 in accordance with the framework presented in Section 3.

27 682 *5.1. Selection accuracy*

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30 683 Each of the proposed selection accuracy scores (i.e. SA , SA_c and SA_e)
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32 684 was computed as the average over the 30 replicates for each of the four IVS
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34 685 algorithm and 26 benchmark datasets.

36 686 *5.1.1. Overall accuracy*

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39 687 The results for the SA metric reported in Figure 4 show that the PMIS,
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41 688 IIS and GA-ANN algorithms share a similar range of variation for the SA
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43 689 score, which varies from 1 (corresponding to correctly specified models) to
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45 690 about 0.4. On the other hand, the SA values for the PCIS algorithm vary
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47 691 from 1 to 0, where a value of 0 corresponds to a completely mis-specified
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49 692 model. The cases at the extreme ends of these ranges correspond to the
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51 693 AR1 and Miller datasets: all algorithms are capable of selecting the only
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53 694 relevant variable in the AR1 dataset without choosing any other extraneous
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55 695 input, while all algorithms have difficulties in selecting the correct inputs
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57 696 for the Miller dataset, with the PCIS algorithm unable to select any of the

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697 correct inputs. Unsurprisingly, the performance of the four algorithms varies
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698 depending on dataset properties. For instance, even though the AR1 and
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699 AR9_500 datasets are characterised by high noise and high collinearity, the
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700 fact that there are only a few relevant inputs (one and three, respectively,
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701 see Table 1) and that the N/P ratio is high, makes the input selection task
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702 relatively simple for all algorithms, as indicated by the high SA scores (see
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703 Figure 4 (AR1)-(AR9_500)). However, a variation in just one of the proper-
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704 ties of the data, such as the N/P ratio in the AR9_70 dataset, which differs
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705 from the AR9_500 dataset in the number of observations (70 instead of 500),
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706 affects the performance of all algorithms.
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708 Furthermore, Figure 4 shows that different values of SA_c and SA_e were
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709 obtained for the combination of IVS algorithms and datasets considered.
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710 The case of perfect specification (i.e. $SA = 1$) was obtained for a number
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711 of different datasets, such as AR1, AR9_500, TAR1 and TAR2, for which
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712 all algorithms were capable of only selecting the relevant variables. As com-
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713 mented above, these datasets have high noise and high collinearity, which are
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714 somehow ‘compensated’ for by the N/P ratio, equal to 33.3 (see Table 1).
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715 This high ratio between the number of observations N and candidate inputs
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716 P allows all algorithms to limit the bias in the estimation of the strength
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717 of dependence between inputs and output due to the presence of noise in
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718 the observational dataset (*regression dilution*, Frost and Thompson (2000)).
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719 A decrease in the number of observations, as in the AR9_70 dataset, had
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720 a negative impact on algorithm performance, causing under-specification of
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721 relevant inputs (i.e. $SA_c < 1$ and $SA_e = 1$) or both under-specification
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10 722 of relevant and over-specification of extraneous inputs (i.e. $SA_c < 1$ and
11 723 $SA_e < 1$). For example, use of the PMIS and GA-ANN algorithms did not
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13 724 result in the selection of extraneous inputs ($SA_e = 1$), but over the 30 repli-
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15 725 cates of the dataset, they show an average SA_c score of about 0.65, indicating
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17 726 that the proportion of correct inputs that has been selected is 65%. This re-
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19 727 sults in a SA score of about 0.75, as shown in Figure 4 (AR9.70). For the
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21 728 same dataset, the IIS algorithm results in a SA_c score of about 0.65, but the
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23 729 overall performance is affected negatively by the over-specification of some
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25 730 extraneous inputs, with SA_e equal to 0.80, which means that the proportion
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27 731 of extraneous inputs that has been selected is 20%. On the other hand, the
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29 732 PCIS algorithm shows perfect specification, with both SA_c and SA_e equal to
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31 733 1. The over-specification of extraneous inputs (i.e. $SA_c = 1$ and $SA_e < 1$)
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33 734 was observed for the Miller dataset, where all relevant variables were selected
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35 735 when using the IIS and GA-ANN algorithms ($SA_c = 1$), but the only extra-
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37 736 neous input was also included, resulting in a value of SA_e equal to 0. Worse
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39 737 results were obtained for the PMIS and PCIS algorithms: the former resulted
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41 738 in a SA_c score of 0.5 and a SA_e score of 0, while the latter performed poorly
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43 739 with respect to both indicators (SA_c and SA_e equal to 0).

44 740 *5.1.2. Effect of dataset properties on algorithm performance*

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47 741 The impact of dataset properties on IVS algorithm performance in terms
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49 742 of SA , SA_c and SA_e is shown in Figure 5 and described below for each of
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51 743 the datasets in turn. A discussion of the findings is provided in Section 6.

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53 744 • *AR and TAR datasets.* As discussed in the previous section, these
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55 745 datasets have high noise and collinearity, but a high number N of

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10 746 observations and a reduced number P of total candidate inputs. The
11 747 ratio N/P is therefore high, allowing all of the algorithms to identify the
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13 748 K relevant inputs ($SA_c = 1$), without including any extraneous inputs
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15 749 ($SA_e = 1$). The only exception is the AR9_70 dataset: in this case the
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17 750 number of observations decreases from 500 to 70, and the N/P ratio
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19 751 from 33.3 to 4.7, with an associated greater likelihood of overfitting.
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21 752 This is empirically demonstrated by the IIS algorithm, which indeed
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23 753 shows a value of SA_e lower than 1.

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25 754 • *NL datasets.* The NL_500 dataset is not characterised by high noise
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27 755 or high collinearity, but is highly nonlinear and has a non-Gaussian
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29 756 output. This combination seems to have a negative impact on the per-
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31 757 formance of the PMIS, ANN-GA and PCIS algorithms, as indicated
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33 758 by a significant decrease in SA_c (i.e. greater under-specification of
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35 759 the relevant inputs). The reason for this may be due to the specific
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37 760 characteristics of each algorithm. PMIS, for example, can accurately
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39 761 model nonlinear relationships, but the Gaussian reference bandwidth
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41 762 used in the estimation of the PMI is known to result in reduced perfor-
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43 763 mance in cases where the data follow a non-Gaussian distribution (May
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45 764 et al., 2008a). As expected, the performance of the PCIS algorithm is
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47 765 worse than that of the PMIS algorithm, since it is based on partial
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49 766 linear correlation, and is therefore unable to account for the nonlinear-
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51 767 ity in the data. Finally, the low performance shown by the GA-ANN
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53 768 algorithm may be due to the simple ANN architecture adopted (i.e.
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55 769 1-hidden node multi-layer perceptron), which might not be fully capa-
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57 770 ble of characterizing the highly nonlinear behaviour of NL_500 dataset.

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10 771 The IIS algorithm, based on Extra-Trees, is capable of selecting all
11 772 relevant inputs ($SA_c = 1$) without including any extraneous inputs
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13 773 ($SA_e = 1$), probably because Extra-Trees are capable of accounting for
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15 774 nonlinear relationships and do not require any assumption about the
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17 775 statistical properties of the dataset. However, as seen for the AR9_70
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19 776 dataset, IIS is sensitive to a decrease in the N/P ratio: while PMIS,
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21 777 ANN-GA and PCIS show similar performance for both the NL_500
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23 778 and NL_70 datasets, the performance of IIS decreases for the second
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25 779 dataset. Finally the high noise and collinearity (in addition to non-
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27 780 Gaussian output and nonlinearity) characterising the NL2 dataset do
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29 781 not seem to affect the performance of the PMIS, ANN-GA and PCIS
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31 782 algorithms, when compared with the performance of these algorithms
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33 783 on the NL_500 dataset. This seems to be in line with the results for the
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35 784 AR and TAR datasets, and empirically demonstrates that the overall
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37 785 performance of these algorithms is more sensitive to non-Gaussian out-
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39 786 puts and/or nonlinearity. This is not the case for the IIS algorithm, the
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41 787 performance of which decreases in terms of both SA_c and SA_e when
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43 788 including high noise and collinearity.

- 44 789 • *Bank datasets.* Unlike any other dataset, these four datasets are charac-
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46 790 terised by incomplete information about the output data, which seems
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48 791 to have a negative effect on all algorithms, with all values of SA lower
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50 792 than 1. It is important to note that the sub-optimal values in SA are
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52 793 due to sub-optimal values in SA_c only (SA_e is always equal to 1), indi-
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54 794 cating that the algorithms do not have sufficient information to single
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56 795 out the relevant inputs. The degree of nonlinearity of the underlying

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10 796 function does not seem to significantly affect SA_c (and therefore SA),
11 797 while the presence of noise (datasets Bank_fh and Bank_nh) appears to
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13 798 have a greater negative impact on the ANN-GA and PCIS algorithms.
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16 799 • *Friedman datasets.* These datasets are characterised by a combination
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18 800 of nonlinearity, noise, collinearity and different numbers P of candi-
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20 801 date inputs. For Friedman_c0_10_m and Friedman_c0_50_m, the PMIS
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22 802 and IIS algorithms result in a value of SA equal to 1, as they are
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24 803 both capable of dealing with nonlinear datasets. On the other hand,
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26 804 the GA-ANN and PCIS algorithms have slightly lower performances,
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28 805 most likely due to their lower efficiency in characterising highly non-
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30 806 linear functions, as discussed previously. Other potential sources of
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32 807 failure include the parameterization of the GA (e.g. insufficient ex-
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34 808 ploration of the search space or insufficient numbers of generations).
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36 809 The addition of high noise (Friedman_c0_10_h and Friedman_c0_50_h)
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38 810 reduces the selection accuracy of PMIS and IIS, while it does not af-
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40 811 fect that of GA-ANN and PCIS algorithms. It thus seems that the
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42 812 presence of noise flattens the modelling conditions, with all algorithms
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44 813 showing similar performance. The addition of high collinearity in the
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46 814 Friedman_c25_10_m and Friedman_c25_50_h datasets causes a decrease
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48 815 in SA for all algorithms, with the combination of high noise and high
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50 816 collinearity being particularly critical. It is interesting to note that all
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52 817 algorithms show a value of SA_e equal to 1, with SA_c and SA lower
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54 818 than 1. This means that in the presence of nonlinearity, high noise and
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56 819 collinearity, the algorithms under-specify the relevant inputs, but do
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58 820 not select extraneous inputs.
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- 821 • *Salinity datasets.* The salinity datasets have a relatively large number
822 of candidate inputs (80 or 160), including time lagged values (5 or
823 10) of 16 variables, resulting in high collinearity in the input data.
824 The presence of high collinearity and 80 candidate inputs (Salinity_5_l
825 dataset) slightly affects the performance of the PCIS and GA-ANN
826 algorithms ($SA_e < 1$), which is further reduced by the addition of 80
827 extra inputs (Salinity_10_l dataset). This may be due to the difficulty
828 the GA has in finding the correct combination of input variables among
829 a set of 160 highly correlated inputs. On the contrary, PMIS and IIS
830 are capable of determining all relevant inputs for both datasets. When
831 moderate noise is added to these data (Salinity_5_m and Salinity_10_m
832 datasets), IIS and PCIS maintain the same performance (i.e. perfect
833 specification and a slight over-specification of some extraneous inputs,
834 respectively), while GA-ANN shows a further decrease. The PMIS
835 algorithm shows a pronounced under-specification of relevant inputs
836 ($SA_e < 1$), which may be due to some difficulties in estimating the
837 correct values of PMI in the presence of noise. Finally, the addition
838 of high noise (Salinity_5_h and Salinity_10_h datasets) to both high
839 collinearity and a large number of input variables has a negative effect
840 on all algorithms, which show a decrease in SA_e and hence in SA .
 - 841 • *Kentucky dataset.* Similar to the Salinity datasets, this dataset is also
842 characterised by a large number P of candidate inputs defined as time
843 lagged values of flow and rainfall observations, causing high collinearity
844 in the data. The presence of random noise is limited, but the output
845 is non-Gaussian. As was found for the NL and Bank datasets, the

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10 846 presence of a non-Gaussian output particularly affects the performance
11 847 of PMIS and GA-ANN: the former has a SA_c score equal to about 0.50
12 848 (meaning that the proportion of correct inputs that has been selected is
13 849 only 50%), while the latter is capable of selecting the relevant variables,
14 850 but tends to include extraneous inputs (SA_e equal to 0.80). On the
15 851 other hand, IIS and PCIS seem to be less affected by the non-Gaussian
16 852 output.

- 17 853 • *Miller dataset.* This dataset has a non-Gaussian output and three can-
18 854 didate inputs only: x_1 and x_2 have a strong inter-dependency (i.e. they
19 855 jointly influence the output, while having little influence on the output
20 856 individually), while the extraneous input x_3 has the highest (spuri-
21 857 ous) correlation with the output. This last characteristic makes the
22 858 input selection exercise particularly challenging for forward selection
23 859 methods (i.e. PMIS, IIS and PCIS) that select only one input at each
24 860 iteration, as evidenced by SA scores of less than 1. PMIS and PCIS
25 861 are particularly affected by the inter-dependency. The SA_e score is
26 862 equal to 0 for both algorithms, meaning that they always select the
27 863 extraneous input x_3 , and the SA_c score is respectively equal to 0.5 and
28 864 0, resulting in very low values for the SA score. Surprisingly, the IIS
29 865 algorithm achieves the same performance as the GA-ANN algorithm
30 866 (the only wrapper method adopted in this study), with SA_e equal to 0,
31 867 but SA_c equal to 1. That is both IIS and GA-ANN select all candidate
32 868 inputs. The unexpectedly good performance of the IIS algorithm may
33 869 be due to the algorithm tolerance (i.e. $\varepsilon = 0.01$), which could cause a
34 870 slight tendency to over-specify models and allow the algorithm to select

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9 additional variables beyond the first ‘most relevant’ input.

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12 5.1.3. *Effect of N and P on algorithm performance*

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14 873 As discussed in the previous section, an increase in the number N of
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16 874 observations increases the information available for the IVS algorithm, thus
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18 875 positively impacting selection accuracy (and vice versa for a decrease in N).
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20 876 As far as the number P of candidates is concerned, an increase in P should
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22 877 increase the overall complexity of the IVS problem, but, in practice, the
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24 878 results show that the correspondence between P and SA is neither univocal
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26 879 nor monotonic. Indeed, different values of SA are found for the same value
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28 880 of P (see, for instance, the AR datasets), and SA does not decrease with
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30 881 P . For example, the average performance of the four algorithms on the
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32 882 Miller dataset ($P = 3$) is lower than the average performance on the Salinity
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34 883 datasets, where P is either 80 or 160. The ratio N/P is probably a better
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36 884 indicator of IVS problem complexity, since its value is indicative of the risk of
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38 885 retaining extraneous inputs (i.e. likelihood of overfitting). High values of this
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40 886 ratio are generally associated with high values of SA , and vice versa. The
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42 887 ratio N/P should be evaluated together with the other properties of a dataset,
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44 888 and collinearity in particular, as the likelihood of overfitting increases with
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46 889 the correlation between the candidate inputs. Indeed, it can be observed that
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48 890 when the ratio N/P falls below a critical threshold (about 5), most of the
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50 891 considered IVS algorithms have a SA score of less than 1. In other words,
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52 892 the ratio N/P appears to be a ‘limiting factor’, where low values limit the
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54 893 capability of any IVS algorithm to select the relevant input variables.
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9 894 *5.2. Computational efficiency*

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11 895 From Table 2, it is evident that when the IVS algorithms are applied
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13 896 to the benchmark datasets, two different behaviours, in terms of average
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15 897 run-time, are observed. First, the PMIS and IIS algorithms show similar
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17 898 run-times, especially for the first eighteen datasets. These datasets are char-
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19 899 acterised by a limited number of observations and candidate inputs N and P
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21 900 (ranging from 70 to 500 and 10 to 50, respectively), so both algorithms per-
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23 901 form the input selection task in a time ranging from a few seconds to about
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25 902 one minute. Such computational efficiency is due to their forward selection
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27 903 nature (i.e. one variable is selected at each iteration), which only requires
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29 904 a small number of iterations and therefore few calibrations of GRNNs and
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31 905 Extra-Trees. On the other hand, the application of these two algorithms to
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33 906 the Salinity and Kentucky datasets, which are characterised by much larger
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35 907 numbers of samples and candidate inputs, increases the run-time up to about
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37 908 2.5 hours, but with IIS faster than PMIS. Apart from the adopted server and
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39 909 the specific implementation (a C++ executable may be faster than the R en-
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41 910 vironment), the reason behind this difference stands in the good scalability
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43 911 of Extra-Trees to large datasets. Indeed, as further discussed in Appendix
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45 912 B, the Extra-Trees run-time increases linearly with P and superlinearly with
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47 913 N , while the time required to calibrate a GRNN does not scale well to large
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49 914 datasets. Moreover, Extra-Trees are used in the IIS algorithm a smaller
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51 915 number of times than GRNNs in the PCIS algorithm, resulting in a run-time
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53 916 order that is quadratic in P and superlinear in N , whilst the run-time order
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55 917 of PMIS is $O(P^4 \cdot N^2 + P^5)$ (Table 3). The PCIS algorithm has the smallest
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57 918 run-time, which varies from a few tenths of a second to about 3 minutes

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10 919 for the most complex datasets. Similarly to PMIS and IIS, this algorithm
11 920 also has a forward selection nature, requiring only few estimates of the Pear-
12 921 son correlation coefficient and calibrations of a linear regression model. The
13 922 latter have a high computational efficiency, thus reducing PCIS run-time in
14 923 comparison with that for PMIS and IIS. Apart from this specific difference,
15 924 these three filters are characterised by similar growth rates of the run-time:
16 925 PMIS and PCIS find a solution in an expected number of $O(P^4 \cdot N^2 + P^5)$
17 926 and $O(P^4 \cdot N + P^5)$ steps, respectively, while IIS requires $O(T \cdot P^2)$ steps
18 927 (where T is the time required for cross-validating and ensemble of Extra-
19 928 Trees). That is, the growth rate for the three filters is polynomial in P and
20 929 N (see Appendix B for further details).

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30 930 Second, the GA-ANN algorithm has a run-time that is almost two orders
31 931 of magnitude higher than that of PMIS and IIS. This is due to its wrapper
32 932 nature, which requires several ANN calibration runs at each iteration of the
33 933 GA. As a consequence, the application of this algorithm to the first eigh-
34 934 teen datasets takes a time ranging from a few minutes to about 30 minutes,
35 935 while the time required to analyse the largest datasets (e.g. Salinity_10_l)
36 936 takes almost 80 hours. Notice that such run-times are obtained by adopt-
37 937 ing a relatively-simple ANN, i.e. a 1-hidden node multilayer perceptron, so
38 938 higher run-times would be required if a more complex network architecture
39 939 were adopted. Unlike PMIS, IIS and PCIS, it is much harder to determine
40 940 the growth rate of the run-time of the GA-ANN algorithm, since the GA is
41 941 a stochastic optimisation algorithm, the computational complexity of which
42 942 depends on different factors, such as the (randomly generated) initial popu-
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9 944 run over the whole number of generations, the run-time is proportional to
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11 945 $G \cdot I$, where G and I represent the number of generations and the population
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13 946 size, respectively.
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15 947 5.3. Qualitative criteria

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18 948 In addition to the quantitative criteria, the four IVS algorithms are also
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20 949 assessed in terms of ease of use, explanation capability and flexibility.
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23 950 • *Ease of use.* The three filter algorithms adopted in this study (i.e.
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25 951 PMIS, IIS and PCIS) are easy to use, in terms of both the number of
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27 952 parameters to be tuned and the robustness of the results with respect
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29 953 to the default set of parameter values. The PMIS and PCIS algorithms
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31 954 require the tuning of two and one parameter, respectively (see Section
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33 955 4). The IIS algorithm has a larger number of parameters, but these
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35 956 can be easily tuned via a trial-and-error procedure (Galelli and Castel-
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37 957 letti, 2013b). Furthermore, the accuracy of the PMIS, IIS and PCIS
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39 958 algorithms is very robust with respect to the adopted (default) parame-
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41 959 terization. On the other hand, the accuracy of the GA-ANN algorithm
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43 960 is sensitive to the parameterization of both GA and ANN. In particular,
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45 961 the results described in Section 5 show that the selection of the ANN
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47 962 architecture appears to be critical. This is a common feature of wrap-
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49 963 per algorithms, which generally require an accurate, dataset-dependent
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51 964 tuning of the underlying model (Guyon and Elisseeff, 2003).
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53 965 • *Explanation capability.* Another advantage of the PMIS, IIS and PCIS
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55 966 algorithms, and of filter methods in general, is that they provide in-
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57 967 formation about the relative importance of each selected input. The
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9 968 same information may be obtained from a wrapper algorithm, but this
10 requires an ex-post interpretation of the data-driven model structure.
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14 970 • *Flexibility.* The PCIS and PMIS algorithms are very flexible, as their
15 structure is identical, but with (1) the PMI criterion used in place of
16 971 the partial linear correlation coefficient, and (2) a GRNN in place of
17 972 the partial linear correlation coefficient, and (2) a GRNN in place of
18 973 a linear model. On the other hand, the IIS algorithm relies on Extra-
19 974 Trees for both ranking the candidate input variables and assessing the
20 975 significance of the selected ones. This reduces the overall flexibility of
21 976 the algorithm, although it may increase the accuracy of the underlying
22 977 model. Finally, it could be argued that the GA-ANN algorithm exhibits
23 978 a high level of flexibility, as both the optimization algorithm (GA) and
24 979 the underlying data-driven model (ANN) can be interchanged with
25 980 other methods. However, such high flexibility comes at a price, since
26 981 the adoption of complex, highly parameterized components may neg-
27 982 atively impact the algorithm’s easy of use, explanation capability and
28 983 computational efficiency.

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33 984 **6. Discussion, recommendations and issues in environmental mod-**
34 985 **elling**

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46 986 *6.1. Discussion*

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49 987 The effect of the properties of the benchmark datasets on IVS algorithm
50 988 performance can be summarised as follows:

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54 989 • *Non-Gaussian Output.* Unsurprisingly, it is found that a non-Gaussian
55 990 output inhibits a high level of performance for the PMIS algorithm,

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991 which assumes Gaussian data when estimating the PMI. This tendency
992 is accentuated when this property is combined with other limiting prop-
993 erties (e.g. incomplete information or inter-dependency as in the Bank
994 or Miller datasets). The IIS and PCIS algorithms, which do not assume
995 Gaussian data, appear to be unaffected by a non-Gaussian output, and
996 can indeed achieve a perfect specification, with $SA = 1$, when this
997 property is not combined with other limiting properties (e.g. NL_500
998 for IIS and Kentucky dataset for PCIS).

999 • *High Nonlinearity.* The presence of a highly nonlinear relationship be-
1000 tween inputs and output can be effectively handled by IIS and PMIS,
1001 which rely on regression methods capable of characterising such re-
1002 lationships (Extra-Trees and GRNN, respectively). This is demon-
1003 strated on the Friedman_c0_10m and Friedman_c0_50m datasets, which
1004 are characterised by this property only. As mentioned above, IIS can
1005 simultaneously deal with non-Gaussian outputs and highly nonlinear
1006 input-output relationships, if enough observations are available (see
1007 NL_500 and NL_70 datasets). On the other hand, both GA-ANN and
1008 PCIS are affected by highly nonlinear datasets. Indeed, the former
1009 relies on a simple 1-hidden node ANN, which is effective with weakly
1010 nonlinear functions only, while the latter is based on linear partial cor-
1011 relation.

1012 • *High Noise.* The presence of high noise affects the performance of all
1013 IVS algorithms, but only when combined with certain other properties
1014 of the data. For example, the combination of high noise with high

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1015 nonlinearity decreases the signal-to-noise ratio, which deteriorates the
1016 performance of both the PMIS and IIS algorithms on the Friedman
1017 datasets. Similarly, the presence of high noise is critical when evaluated
1018 in relation to the number N of observations. Indeed, a decrease in N
1019 affects the signal-to-noise ratio, as illustrated by the deterioration in
1020 performance when reducing the number of observations from 500 to 70
1021 in the AR_70 dataset.

1022 • *High Collinearity.* This property is normally due to the presence of time
1023 lagged values of some input variables, as in the AR, TAR, Salinity and
1024 Kentucky datasets. Similar to the presence of high noise, collinearity
1025 can also be effectively handled by all algorithms, even in the presence
1026 of many inputs, such as in the Salinity_5.1 and Salinity_10.1 datasets,
1027 where P is equal to 80 and 160, respectively. However, when high noise
1028 is introduced into the dataset, good performance can only be achieved
1029 if the number P of candidate inputs is limited, as is the case for the
1030 AR and TAR datasets.

1031 • *Inter-dependency.* As explained in Section 5, inter-dependency between
1032 input variables (Miller dataset) is generally problematic for filter, for-
1033 ward selection methods that evaluate one input variable at each it-
1034 eration. Indeed, PMIS and PCIS exhibit low accuracy on the Miller
1035 dataset (SA equal to about 0.40 and 0, respectively), while IIS is capa-
1036 ble of achieving greater accuracy, probably because of the pre-selected
1037 exit condition. Unsurprisingly, the GA-ANN algorithm, the only wrap-
1038 per method considered in this study, achieves the best performance.

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1039 • *Incomplete information.* This property has a significant impact on all
11 IVS algorithms, as evidenced by the inability of any of the algorithms
12 to select the relevant input variables on the Bank dataset.
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1042 • N , P and N/P . The ratio N/P is a further limiting factor that re-
17 duces the accuracy of all algorithms when it drops to values below 5.
18 In particular, IIS seems to be more sensitive to drops in the ratio N/P ,
19 as found for the AR_70 and the NL_70 datasets. In addition, both N
20 and P have a strong impact on the computational performance of IVS
21 algorithms. While filter methods, such as PMIS and IIS, exhibit good
22 scalability with respect to input dataset dimensionality, the run-time
23 of wrapper methods is particularly sensitive to an increase in N and
24 P , up to the point where their adoption becomes impractical for large
25 datasets. For example, the GA-ANN algorithm requires more than 3
26 days of computation for solving an input selection problem with 4115
27 observations and 160 candidate input variables, while PMIS and IIS
28 require 2.5 and 1.5 hours, respectively.
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1056 Finally, it is interesting to highlight that the presence of two properties of
44 the data, namely inter-dependency and incomplete information, have a strong
45 impact on the accuracy of all algorithms, irrespective of the presence/absence
46 of other properties. Non-Gaussian output and a highly nonlinear input-
47 output relationship can only be fully handled by some algorithms (i.e. IIS
48 and PCIS, and PMIS and IIS respectively), and they require the adoption of
49 specific metrics and regression methods that can deal with such properties.
50 Finally, high noise and high collinearity are not a problem per se, but their
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1064 presence adds a further level of complication when they are combined with
1065 other properties, such as non-Gaussian outputs or nonlinear datasets.

1066 *6.2. Recommendations*

1067 Although the results reported here are primarily discussed for the pur-
1068 poses of the demonstration of the framework, they can be used to develop
1069 some preliminary guidelines in relation to the relative importance of differ-
1070 ent properties of the data and the corresponding performance of the four IVS
1071 algorithms.

- 1072 • The presence of a *non-Gaussian* output is a potential limiting factor,
1073 which requires the adoption of IVS algorithms that do not assume
1074 Gaussian data when estimating the relative importance of each input or
1075 when building a regression model. A similar recommendation is valid in
1076 case of a *highly nonlinear* relationship between input and output. The
1077 only algorithm capable of selecting the correct inputs in the presence
1078 of both properties is IIS.

- 1079 • As discussed in Section 6.1, the presence of *high noise* is problematic
1080 only when combined with certain other properties of the data (e.g.
1081 non-Gaussian output or inter-dependency), so the choice of the most
1082 appropriate IVS algorithm should be based on its capability of dealing
1083 with such properties. This guideline still holds in the presence of *high*
1084 *collinearity*.

- 1085 • In the presence of *inter-dependency* between input variables, it is ad-
1086 visable to adopt a wrapper method, which can handle this property

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1087 through the selection of multiple inputs at each iteration. This cannot
11 be done by filters, unless the candidate input set is enlarged to include
12 features that are combinations of the original input variables.
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16 1090 • *Incomplete information* within the dataset affects the performance of
17 all IVS algorithms, so the most suitable algorithm should be chosen
18 according to its capability of dealing with the other properties charac-
19 terising the dataset in hand.
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24 1094 • The *ratio* N/P is a limiting factor when it drops to values approxi-
25 mately below 5: in this case it is recommendable to use IVS algorithms
26 that rely on simple metrics and regression techniques, such as PCIS,
27 which is based on the Pearson correlation coefficient and linear regres-
28 sion. Indeed, more advanced algorithms (e.g. PMIS and IIS) require
29 the identification of complex data-driven models, whose performance
30 increases in the presence of more observations.
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- 38 1101 • While the properties and recommendations above should be considered
39 before any IVS experiment, the size of the dataset and the computa-
40 tional performance of an IVS algorithm matter only in the presence of
41 limited computational resources (or limited time available to conduct
42 the experiments). In general, if the maximum computing time that is
43 available for each experiment is in the order of a few hours, it is advis-
44 able to adopt a filter method. On the other hand, if more or unlimited
45 time is available, a wrapper can be a viable solution. However, it must
46 be remembered that the tuning of a wrapper is a time consuming task,
47 which requires an accurate parameterisation of both the optimisation
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9 1111 algorithm and the architecture of the data-driven model.

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12 1112 *6.3. Issues in environmental modelling*

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14 1113 Unlike for the synthetic data considered here, a key aspect of real-world
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16 1114 environmental modelling problems is that the true underlying function is
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18 1115 unknown, and IVS is thus used to reduce the uncertainty in the model de-
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20 1116 velopment process by selecting a subset of relevant and non-redundant input
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22 1117 variables. This opens some relevant theoretical and practical issues that are
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24 1118 highlighted below:

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27 1119 • Most of the IVS algorithms currently available select a unique subset
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29 1120 of input variables, although the structural uncertainty in the inputs
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31 1121 to be used often results in the possibility of choosing different, but
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33 1122 equally informative, subsets. An attempt to account for this issue was
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35 1123 recently made by Sharma and Chowdhury (2011), who proposed a PMI-
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37 1124 based heuristic approach to select five different subsets of predictors in
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39 1125 the context of medium-term hydro-climatic forecasting. The approach
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41 1126 ensures that the cross-dependence between these subsets is limited,
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43 1127 while the predictions of the resulting models are eventually combined
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45 1128 with ensemble averaging.
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47 1129 • In many practical situations, input variables can be characterised by
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49 1130 errors, due to, for example, the interpolation of data in space and
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51 1131 time or the conversion of point measurement into areal values. Whilst
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53 1132 methods exist for assessing the impact of input errors on parameter
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55 1133 estimation procedures (Chowdhury and Sharma, 2007; Woldemeskel

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9 1134 et al., 2012), IVS algorithms cannot take into account the change in
10 the uncertainty associated with the different inputs.
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14 1136 • A benefit of IVS is the improvement in the performance of the model
15 being identified. Although the manner in which such performance is
16 1137 characterised depends on the specific domain of interest and the model
17 1138 objectives (Jakeman et al., 2006), two important aspects should always
18 1139 be considered when dealing with quantitative testing. First, the use
19 1140 of observational data for comparison must rely on appropriate data-
20 1141 division methods, such as cross-validation or bootstrapping, that allow
21 1142 for testing the ability of the model to generalise. Data division can
22 1143 account for both temporal and spatial dimensions, so it is suitable for
23 1144 spatial modelling as well (see Chowdhury and Sharma (2009) for an
24 1145 application to hydrological modelling problems). Second, an exhaus-
25 1146 tive quantitative evaluation should rely on a set of metrics focussing on
26 1147 different aspects in order to test the ability of the model in reproducing
27 1148 all the important features of the system. The reader is referred to Ben-
28 1149 nett et al. (2013) for a comprehensive review of techniques available for
29 1150 both data-division and quantitative evaluation, and to Robson (2014)
30 1151 for a more general assessment of environmental models.
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48 1153 7. Closure

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50 1154 In this work we present a framework for the comparative analysis of IVS
51 algorithms in environmental modelling problems. The framework consists
52 1155 of a set of benchmark datasets with the typical properties of environmental
53 data, a recommended set of evaluation criteria and a website for sharing
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1158 data, code and results. Since the data and criteria proposed here cannot
1159 exhaustively represent all modelling contexts encountered by developers and
1160 users, it is hoped that the presence of a dedicated website will increase the
1161 flexibility of this framework and facilitate collaboration between researchers.
1162 For example, the benchmark datasets are currently limited to those that have
1163 both continuous input and output variables; however, it is intended that this
1164 set will be extended to include datasets comprised of nominal and categorical
1165 variables. In addition, as this framework is applied to an increasing number
1166 of IVS algorithms and datasets, it is hoped that guidelines for the adoption of
1167 the most appropriate IVS algorithms for datasets with particular properties
1168 can be developed.

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1174 algorithms.

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1412 **Appendix A**

1413 *A.1 PMIS algorithm*

1414 The PMIS algorithm is a filter IVS method developed by Sharma (2000)
1415 and later modified by Bowden et al. (2005a) and May et al. (2008a), where the
1416 relevance of potential inputs is evaluated based on the mutual information
1417 (MI) between each input variable and the output. While MI is a useful
1418 measure of dependence between a potential input variable \mathbf{x} and a dependent
1419 variable \mathbf{y} , it cannot account for redundancy in the candidate input pool,
1420 $X = \{\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_P\}$. To account for such redundancy, the PMI criterion,
1421 which measures the *partial* dependence between a potential input variable
1422 and the output, conditional on any inputs that have already been selected,
1423 is instead used in this algorithm. This criterion is analogous to the partial

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9 correlation coefficient and can be formulated as:

$$\text{PMI} = \frac{1}{N} \sum_{i=1}^N \log \left[\frac{f(x'_i, y'_i)}{f(x'_i) f(y'_i)} \right] \quad (4)$$

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15 where

$$\mathbf{x}' = \mathbf{x} - E[\mathbf{x}|Z]; \text{ and } \mathbf{y}' = \mathbf{y} - E[\mathbf{y}|Z] \quad (5)$$

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20 represent the residual information in variables \mathbf{x} and \mathbf{y} once the effect of the
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22 already selected inputs, Z , has been taken into consideration. In eq. (4), x'_i
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24 and y'_i are the i -th residuals in a sample dataset of size N and $f(x'_i)$, $f(y'_i)$
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26 and $f(x'_i, y'_i)$ are the respective marginal (univariate) and joint (bivariate)
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28 probability density functions (pdfs).
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1432 Calculation of the PMI criterion in eq. (4) requires estimation of the
1433 marginal and joint pdfs of \mathbf{x} and \mathbf{y} . For the PMIS algorithm, Sharma (2000)
1434 proposed the use of a non-parametric kernel density estimation based on the
1435 Gaussian kernel function (Silverman, 1986). The accuracy of this kernel es-
1436 timator is largely dependent on the choice of the smoothing parameter (or
1437 bandwidth) λ , with its optimal value depending on the distribution of the
1438 available data sample (May et al., 2008a). A value of λ that is too large
1439 will result in an over-smoothed probability density, while a value that is too
1440 small can lead to density estimates overly influenced by individual data points
1441 (under-smooth). Sharma (2000) adopted the Gaussian reference bandwidth
1442 (Scott, 1992) due to its simplicity and computational efficiency. Calculation
1443 of the PMI criterion also requires the appropriate estimation of the condi-
1444 tional expectation $E[\cdot]$ of \mathbf{x} and \mathbf{y} on Z . Bowden et al. (2005a) proposed
1445 the use of a General Regression Neural Network (GRNN) to compute these

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10 1446 conditional expectations. GRNNs are very similar in their underlying philos-
11 1447 ophy to kernel regression, where a non-parametric estimate of the pdf of the
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13 1448 observed data, similar to that given by eq. (4), is utilised in the estimation
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15 1449 of $E[\cdot]$, rather than assuming any particular form for the regression function.
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17 1450 At each iteration, the PMIS algorithm seeks to find the variable \mathbf{x}_s which
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19 1451 maximises the PMI with respect to \mathbf{y} , conditional on the inputs that have
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21 1452 been selected in previous iterations, Z . If \mathbf{x}_s is found to be relevant (based
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23 1453 on some stopping criterion), it is added to the selected subset Z and the
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25 1454 selection continues; otherwise, the algorithm is terminated since there are no
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27 1455 more relevant candidate inputs remaining. For the purposes of this study,
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29 1456 the stopping criterion utilised was the coefficient of determination, R^2 , of the
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31 1457 output variable residual, \mathbf{y}' (see May et al. (2008a) for an analysis of different
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33 1458 stopping criteria).

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36 1460 The advantages of using a GRNN in the PMIS algorithm include: accu-
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38 1461 racy in modelling the nonlinear relationships between the inputs and output,
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40 1462 computationally efficient model calibration, and fixed model structure that
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42 1463 does not have to be tuned on each specific dataset (Bowden et al., 2005a).
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44 1464 On the other hand, a limitation of the PMIS algorithm is that, although it
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46 1465 is a filter method, it can still be relatively computationally expensive due
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48 1466 to the use of kernel based approaches for estimating the PMI criterion and
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50 1467 the conditional expectations $E[\cdot]$. While such approaches give efficient and
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52 1468 reliable density estimates for smaller data sets, their computational efficiency
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54 1469 decreases dramatically with increasing sample size (Fernando et al., 2009).
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56 1470 Furthermore, the Gaussian reference bandwidth, which is utilised in the cal-

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9 1471 culation of the marginal and joint pdfs of x and y , as well as in the GRNN
10 1472 estimates of $E[x|Z]$ and $E[y|Z]$, can tend to over-smooth and its optimality
11 1473 might be questionable if the data are not Gaussian (May et al., 2008a). For
12 1474 further details of this algorithm, see Sharma (2000); Bowden et al. (2005a)
13 1475 and May et al. (2008a).

1476 *A.2 IIS algorithm*

1477 The IIS algorithm is a hybrid filter-wrapper IVS method introduced by
1478 Galelli and Castelletti (2013b). Similar to the PMIS algorithm, IIS adopts
1479 a forward selection approach to iteratively select the most significant inputs,
1480 but uses a tree-based ranking method instead of an information-theoretic
1481 measure to estimate the relative contribution of each candidate input. At
1482 each iteration, all the input variables are ranked according to their relative
1483 contribution to the building of an underlying model of the output. The rel-
1484 ative significance of the first p ranked variables is then assessed against the
1485 output by identifying p Single Input-Single Output (SISO) models. Even-
1486 tually, the best performing input among the p considered (according to a
1487 preselected measure of accuracy) is added to the set of the selected variables.
1488 At the first iteration of the IIS algorithm, both ranking and SISO models are
1489 run on a data set composed of time series of the candidate input variables
1490 and the associated output values. At the subsequent iterations, the original
1491 output values are replaced by the residuals of the underlying model built
1492 at the previous iteration. The re-evaluation of ranking and SISO models
1493 every time an input is selected (i.e., at each iteration) ensures that all the
1494 candidate inputs that are highly correlated with the selected input are dis-
1495 carded, thus minimizing the redundancy of the final set of selected inputs.

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1496 The IIS algorithm terminates when the accuracy of the model built upon the
1497 selected variables, as evaluated with a k -fold cross validation (Allen, 1974),
1498 starts decreasing (or when it does not significantly improve). As discussed
1499 in Wan Jaafar et al. (2011), this process is aimed at minimizing the risk of
1500 overfitting the data, since it estimates the ability of the model to capture the
1501 behavior of unseen or future observations.

1502
1503 In the present study the underlying model performance is computed
1504 with the coefficient of determination R^2 , while both the ranking and model
1505 building algorithm are based on Extremely Randomized Trees (Extra-Trees,
1506 Geurts et al. (2006); Galelli and Castelletti (2013a)). Similar to the PMIS
1507 algorithm, the idea of exploiting the underlying model residuals provides ro-
1508 bustness against redundant inputs, while the adoption of Extra-Trees allows
1509 accounting for non-linear interactions and computational efficiency (with re-
1510 spect to sample size N and the number P of candidate inputs). Furthermore,
1511 the tree-based ranking method does not require any specific assumption re-
1512 garding the structure of the dependence between input and output variables.
1513 However, as any other forward selection method, the IIS algorithm does not
1514 account for the inter-dependency between candidate input variables. For fur-
1515 ther technical details the reader is referred to Galelli (2010) and Galelli and
1516 Castelletti (2013b).

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1518 *A.3 GA-ANN algorithm*

1519 The algorithm described herein is one particular implementation of a
1520 combination of a Genetic Algorithm (GA) search procedure with an Artifi-

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9 1521 cial Neural Network (ANN) model. In particular, a simple 1-hidden node
10 1522 multilayer perceptron was utilised in this algorithm¹. The model training
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12 1523 process is performed by means of a simulated annealing algorithm (Belisle,
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14 1524 1992), which is used each time a new combination of inputs is evaluated.

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17 1525 The GA here adopted is a relatively simple variant which is outlined in
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19 1526 Goldberg (1989). In this implementation, solutions representing different
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21 1527 subsets of inputs, are encoded as binary strings, called ‘chromosomes’. Each
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23 1528 bit, or ‘gene’, in these chromosomes represents a candidate input variable,
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25 1529 where a ‘1’ denotes that the input will be included in the model, while a
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27 1530 ‘0’ denotes its omission from the model. The objective function used to de-
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29 1531 termine whether one subset of inputs is better (fitter) than another was the
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31 1532 out-of-sample AIC, computed using a k -fold cross-validation. This objective
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33 1533 function was also used as a stopping criterion to terminate the GA-ANN
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35 1534 algorithm.

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38 1536 The main drawback of this implementation of the GA-ANN algorithm is
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40 1537 that the complexity of the learning algorithm, and hence its ability to accu-
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42 1538 rately model complex functions, is limited by the choice of an ANN with a
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44 1539 single hidden node. However, this model should still provide an improvement
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46 1540 over a simple linear mapping when applied to nonlinear datasets.

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49 ¹Ideally, the structure and complexity of the ANN model would be optimised to suit
50 the problem at hand; however, when evaluating a general algorithm across a number of
51 different datasets this can become impractical.
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9 1541 *A.4 PCIS algorithm*

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11 1542 The partial correlation input selection (PCIS) algorithm (May et al.,
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13 1543 2008a) is based on partial correlation analysis, which aims to find the linear
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15 1544 correlation between two variables after removing the effects of other variables.
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17 1545 The PCIS algorithm is structured the same as the PMIS algorithm, but with
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19 1546 the partial linear correlation coefficient used in place of the PMI criterion
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21 1547 for measuring the relevance of inputs. This coefficient is calculated as Pear-
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23 1548 son’s correlation between the residuals \mathbf{x}' and \mathbf{y}' , given by eq. (5), once the
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25 1549 effect of the already selected inputs, Z , has been taken into consideration.
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27 1550 In this case, the conditional expectation $E[\cdot]$ is a linear regression of \mathbf{x} and \mathbf{y}
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29 1551 with Z . The regression is based on a least-squares approach, which implies
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31 1552 a Gaussian distribution of the residuals. The PCIS algorithm is terminated
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33 1553 when the selection of additional inputs no longer results in an improvement
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35 1554 (increase) in the BIC, calculated based on the output variable residual \mathbf{y}' ,
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37 1555 which provides a trade-off between goodness-of-fit and model complexity.

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40 1556 **Appendix B**

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42 1557 *B.1 PMIS algorithm*

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44 1558 The computing time $t_{PMIS,i}$ associated with the i -th iteration of the PMIS
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46 1559 algorithm is the combination of the time $t_{PMIS,T1}$ required to calibrate a
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48 1560 GRNN to estimate the output based on the selected inputs, the time $t_{PMIS,T2}$
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50 1561 required to calibrate a GRNN to estimate each (non-selected) input based on
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52 1562 the selected inputs, and the time $t_{PMIS,T3}$ for computing the PMI between
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54 1563 the residual of each model. Knowing that the run-time order to calibrate
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56 1564 a GRNN is $O(K^2 \cdot N^2 + K^3)$ (where K and N are the number of inputs

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9 and observations, respectively), and that the run-time order to estimate the
10 PMI is $O(N^2)$, the time $t_{PMIS,T1}$, $t_{PMIS,T2}$ and $t_{PMIS,T3}$ can be estimated as
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13 follows:

$$t_{PMIS,T1} = c \cdot ((i-1)^2 \cdot N^2 + (i-1)^3) \quad (6a)$$

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17 where c is a constant, machine-dependent parameter and i the iteration num-
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19 ber.

$$t_{PMIS,T2} = c \cdot (P - (i-1)) \cdot ((i-1)^2 \cdot N^2 + (i-1)^3) \quad (6b)$$

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23 where P is the number of candidate input variables.

$$t_{PMIS,T3} = c \cdot (P - (i-1)) \cdot N^2 \quad (6c)$$

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29 Therefore, the time $t_{PMIS,i}$ associated with the i -th iteration is equal to

$$t_{PMIS,i} = t_{PMIS,T1} + t_{PMIS,T2} + t_{PMIS,T3} \quad (7a)$$

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33 while the time $t_{PMIS,n}$ required to perform n iterations is

$$\begin{aligned} t_{PMIS,n} &= c \cdot \sum_{i=1}^n ((i-1)^2 \cdot N^2 + (i-1)^3) + \\ &+ c \cdot \left[\sum_{i=1}^n (P - (i-1)) \cdot ((i-1)^2 \cdot N^2 + (i-1)^3) + \sum_{i=1}^n (P - (i-1)) \cdot N^2 \right] = \\ &= c \cdot \sum_{i=1}^n ((i-1)^2 \cdot N^2 + (i-1)^3) + \\ &+ c \cdot \sum_{i=1}^n (P - (i-1)) \cdot ((i-1)^2 \cdot N^2 + (i-1)^3 + N^2) \end{aligned} \quad (7b)$$

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54 In the worst case scenario, the PMIS algorithm is run over P iterations
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10 1574 to evaluate all candidate inputs. In this case, the total computing time is

$$\begin{aligned}
 t_{PMIS}(P) &= c \cdot \sum_{i=1}^P ((i-1)^2 \cdot N^2 + (i-1)^3) + \\
 &+ c \cdot \sum_{i=1}^P (P - (i-1)) \cdot ((i-1)^2 \cdot N^2 + (i-1)^3 + N^2)
 \end{aligned} \tag{8}$$

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19 1575 so the run-time order is $O(P^4 \cdot N^2 + P^5)$.

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21 1576 *B.2 IIS algorithm*

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23 1577 The computing time $t_{IIS,i}$ associated with the i -th iteration of the IIS
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25 1578 algorithm is the combination of the time $t_{IIS,T1}$ required to run the ranking
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27 1579 method, the time $t_{IIS,T2}$ for evaluating the accuracy of p SISO models and
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29 1580 the time $t_{IIS,T3}$ for evaluating the underlying MISO model. Knowing that
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31 1581 the computing time of Extra-Trees grows superlinearly in the number N of
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33 1582 observations, and linearly in the number K and M of inputs and trees, the
34
35 1583 time $t_{IIS,T1}$, $t_{IIS,T2}$ and $t_{IIS,T3}$ can be estimated as follows:

$$t_{IIS,T1} = c \cdot (N \cdot \log(N)) \cdot M \cdot P \tag{9a}$$

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40 1584 where c is a constant, machine-dependent parameter, and P the number of
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42 1585 candidate input variables.

$$\begin{aligned}
 t_{IIS,T2} &= c \cdot p \cdot k \cdot \left(\left(\frac{N}{k} \cdot (k-1) \right) \cdot \log \left(\frac{N}{k} \cdot (k-1) \right) \right) \cdot M \cdot 1 = \\
 &= c \cdot p \cdot T
 \end{aligned} \tag{9b}$$

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49 1586 where k is the number of folds in the k -fold cross-validation process and T
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51 1587 is equal to $k \cdot \left(\left(\frac{N}{k} \cdot (k-1) \right) \cdot \log \left(\frac{N}{k} \cdot (k-1) \right) \right) \cdot M$.

$$\begin{aligned}
 t_{IIS,T3} &= c \cdot k \cdot \left(\left(\frac{N}{k} \cdot (k-1) \right) \cdot \log \left(\frac{N}{k} \cdot (k-1) \right) \right) \cdot M \cdot i = \\
 &= c \cdot T \cdot i
 \end{aligned} \tag{9c}$$

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9 where i is the iteration number, which can range from 1 to P .

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1588 Therefore, the time $t_{IIS,i}$ associated with the i -th iteration is equal to
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$$t_{IIS,i} = t_{IIS,T1} + t_{IIS,T2} + t_{IIS,T3} \quad (10a)$$

1591 while the time $t_{IIS,n}$ required to perform n iterations is

$$t_{IIS,n} = n \cdot t_{IIS,T1} + n \cdot t_{IIS,T2} + \sum_{i=1}^n c \cdot T \cdot i \quad (10b)$$

1592 In the worst case scenario, the IIS algorithm is run over P iterations to
1593 evaluate all candidate inputs. In this case, the total computing time is

$$\begin{aligned} t_{IIS}(P) &= P \cdot t_{IIS,T1} + P \cdot t_{IIS,T2} + c \cdot T \cdot [1 + 2 + \dots + (P - 1) + P] = \\ &= P \cdot t_{IIS,T1} + P \cdot t_{IIS,T2} + c \cdot T \cdot \left[\frac{1}{2} \cdot (P^2 + P) \right] \end{aligned} \quad (11)$$

1594 so the run-time order is $O(T \cdot P^2)$, that is $O(k \cdot ((\frac{N}{k} \cdot (k - 1)) \cdot \log(\frac{N}{k} \cdot (k - 1)))) \cdot$
1595 $M \cdot P^2)$.

1596 B.3 PCIS algorithm

1597 The computing time $t_{PCIS,i}$ associated with the i -th iteration of the PCIS
1598 algorithm is the combination of the time $t_{PCIS,T1}$ required to build a linear
1599 model to estimate the output based on the selected inputs, the time $t_{PCIS,T2}$
1600 required to build a linear model to estimate each (non-selected) input based
1601 on the selected inputs, and the time $t_{PCIS,T3}$ for computing the Pearson
1602 correlation between the residual of each model. Knowing that the run-time
1603 order to build a linear model is $O(K^2 \cdot N + K^3)$, and that the run-time order

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1604 to estimate the Pearson correlation is $O(N)$, the time t_{PCIS,T_1} , t_{PCIS,T_2} and
1605 t_{PCIS,T_3} can be estimated as follows:

$$t_{PCIS,T_1} = c \cdot ((i-1)^2 \cdot N + (i-1)^3) \quad (12a)$$

1606 where c is a constant, machine-dependent parameter and i the iteration num-
1607 ber.

$$t_{PCIS,T_2} = c \cdot (P - (i-1)) \cdot ((i-1)^2 \cdot N + (i-1)^3) \quad (12b)$$

1608 where P is the number of candidate input variables.

$$t_{PCIS,T_3} = c \cdot (P - (i-1)) \cdot N \quad (12c)$$

1609 Therefore, the time $t_{PCIS,i}$ associated with the i -th iteration is equal to

$$t_{PCIS,i} = t_{PCIS,T_1} + t_{PCIS,T_2} + t_{PCIS,T_3} \quad (13a)$$

1610 while the time $t_{PCIS,n}$ required to perform n iterations is

$$\begin{aligned} t_{PCIS,n} &= c \cdot \sum_{i=1}^n ((i-1)^2 \cdot N + (i-1)^3) + \\ &+ c \cdot \left[\sum_{i=1}^n (P - (i-1)) \cdot ((i-1)^2 \cdot N + (i-1)^3) + \sum_{i=1}^n (P - (i-1)) \cdot N \right] = \\ &= c \cdot \sum_{i=1}^n ((i-1)^2 \cdot N + (i-1)^3) + \\ &+ c \cdot \sum_{i=1}^n (P - (i-1)) \cdot ((i-1)^2 \cdot N + (i-1)^3 + N) \end{aligned} \quad (13b)$$

1611 In the worst case scenario, the PCIS algorithm is run over P iterations

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10 to evaluate all candidate inputs. In this case, the total computing time is
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$$\begin{aligned}
 t_{PCIS}(P) = c \cdot \sum_{i=1}^P ((i-1)^2 \cdot N + (i-1)^3) + \\
 + c \cdot \sum_{i=1}^P (P - (i-1)) \cdot ((i-1)^2 \cdot N + (i-1)^3 + N)
 \end{aligned}
 \tag{14}$$

19 so the run-time order is $O(P^4 \cdot N + P^5)$.
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Table 1: Benchmark dataset properties

Dataset	N	K	P	N/P	Fully/Partially Synthetic	Non-Gaussian Output	Highly Nonlinear	High Noise	High Collinearity	Inter-dependency	Incomplete Information
1 AR1	500	1	15	33.3	Fully			X	X		
2 AR9_500	500	3	15	33.3	Fully			X	X		
3 AR9_70	70	3	15	4.7	Fully			X	X		
4 TAR1	500	1	15	33.3	Fully			X	X		
5 TAR2	500	2	15	33.3	Fully			X	X		
6 NL_500	500	3	15	33.3	Fully	X	X				
7 NL_70	70	3	15	4.7	Fully	X	X				
8 NL2	500	3	15	33.3	Fully	X	X	X	X		
9 Bank_fm	400	8	32	12.5	Fully	X					X
10 Bank_fh	400	8	32	12.5	Fully	X		X			X
11 Bank_nm	400	8	32	12.5	Fully	X	X				X
12 Bank_nh	400	8	32	12.5	Fully	X	X	X			X
13 Friedman_c0_10_m	250	5	10	25	Fully		X				
14 Friedman_c0_10_h	250	5	10	25	Fully		X	X			
15 Friedman_c0_50_m	250	5	50	5	Fully		X				
16 Friedman_c0_50_h	250	5	50	5	Fully		X	X			
17 Friedman_c25_10_m	250	5	10	25	Fully		X		X		
18 Friedman_c25_10_h	250	5	10	25	Fully		X	X	X		
19 Salinity_5_l	4120	3	80	51.5	Partially				X		
20 Salinity_5_m	4120	3	80	51.5	Partially				X		
21 Salinity_5_h	4120	3	80	51.5	Partially			X	X		
22 Salinity_10_l	4115	3	160	25.7	Partially				X		
23 Salinity_10_m	4115	3	160	25.7	Partially				X		
24 Salinity_10_h	4115	3	160	25.7	Partially			X	X		
25 Kentucky	4739	4	21	225.7	Partially	X			X		
26 Miller	200	2	3	66.7	Fully	X				X	

Table 2: Average run-time [sec] for the PMIS, IIS, GA-ANN and PCIS algorithms over the 26 benchmark datasets.

Dataset	N	K	P	PMIS	IIS	GA-ANN	PCIS
1 AR1	500	1	15	16.80 ± 2.87	9.49 ± 2.70	1491.22 ± 560.28	0.16 ± 0.05
2 AR9_500	500	3	15	38.84 ± 3.29	26.73 ± 6.64	1973.36 ± 864.35	0.38 ± 0.13
3 AR9_70	70	3	15	2.22 ± 0.43	3.39 ± 0.91	378.49 ± 199.71	0.38 ± 0.13
4 TAR1	500	1	15	14.02 ± 1.22	11.66 ± 3.92	841.69 ± 330.93	0.23 ± 0.09
5 TAR2	500	2	15	26.13 ± 3.46	8.08 ± 0.16	1630.37 ± 860.63	0.40 ± 0.27
6 NL_500	500	3	15	23.41 ± 3.72	24.51 ± 3.39	878.56 ± 272.30	0.20 ± 0.08
7 NL_70	70	3	15	1.82 ± 0.66	4.93 ± 1.79	185.83 ± 117.37	0.25 ± 0.25
8 NL2	500	3	15	24.26 ± 4.94	15.66 ± 5.02	850.15 ± 468.98	0.33 ± 0.15
9 Bank_fm	400	8	32	31.03 ± 6.18	44.41 ± 6.22	1544.37 ± 341.93	1.23 ± 0.36
10 Bank_fh	400	8	32	35.07 ± 7.45	25.52 ± 4.08	1754.10 ± 775.55	0.99 ± 0.27
11 Bank_nm	400	8	32	48.33 ± 17.61	41.94 ± 2.12	1732.64 ± 288.63	1.55 ± 0.50
12 Bank_nh	400	8	32	34.50 ± 12.82	30.59 ± 3.65	1667.77 ± 634.60	1.29 ± 0.37
13 Friedman_c0_10_m	250	5	10	13.36 ± 1.10	9.64 ± 0.45	609.90 ± 215.67	0.31 ± 0.07
14 Friedman_c0_10_h	250	5	10	10.79 ± 1.41	6.68 ± 0.44	710.52 ± 331.96	0.56 ± 0.58
15 Friedman_c0_50_m	250	5	50	46.61 ± 3.42	60.59 ± 4.85	2074.70 ± 564.22	1.26 ± 0.35
16 Friedman_c0_50_h	250	5	50	38.72 ± 6.58	57.39 ± 6.45	1832.76 ± 593.04	1.33 ± 0.55
17 Friedman_c25_10_m	250	5	10	10.31 ± 2.62	10.40 ± 2.47	325.50 ± 87.30	0.26 ± 0.18
18 Friedman_c25_10_h	250	5	10	7.78 ± 1.87	3.17 ± 0.56	303.35 ± 146.60	0.22 ± 0.05
19 Salinity_5_l	4120	3	80	5,017.90 ± 477.32	1,872.05 ± 24.43	100,393.26 ± 24,601.97	73.99 ± 26.89
20 Salinity_5_m	4120	3	80	3,672.10 ± 521.62	1,595.38 ± 66.56	83,150.26 ± 19,792.51	53.32 ± 21.84
21 Salinity_5_h	4120	3	80	3,574.37 ± 394.79	1,451.59 ± 130.43	52,066.80 ± 10,753.36	11.16 ± 4.72
22 Salinity_10_l	4115	3	160	9,024.13 ± 515.44	5,427.50 ± 81.49	287,687.47 ± 58,411.79	143.65 ± 45.33
23 Salinity_10_m	4115	3	160	7,995.80 ± 1,541.41	5,457.86 ± 149.86	226,791.40 ± 47,583.28	143.83 ± 52.24
24 Salinity_10_h	4115	3	160	7,877.20 ± 2,832.16	6,005.77 ± 840.08	147,507.80 ± 25,653.14	18.09 ± 9.16
25 Kentucky	4739	4	21	1,860.37 ± 107.22	800.58 ± 92.86	106,725.55 ± 27,214.61	7.65 ± 1.88
26 Miller	200	2	3	2.65 ± 0.27	0.98 ± 0.55	5664.92 ± 1561.83	0.14 ± 0.08

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Table 3: Run-time order of PMIS, IIS, GA-ANN and PCIS algorithms. P and N represent the number of candidate inputs and observations, respectively, while T is equal to $k \cdot ((\frac{N}{k} \cdot (k - 1)) \cdot \log(\frac{N}{k} \cdot (k - 1))) \cdot M$ (where k is the number of folds in the k -fold cross-validation process and M is the number of trees in an ensemble). See Appendix B for further details.

IVS algorithm	PMIS	IIS	GA-ANN	PCIS
Run-time order	$O(P^4 \cdot N^2 + P^5)$	$O(T \cdot P^2)$	-	$O(P^4 \cdot N + P^5)$

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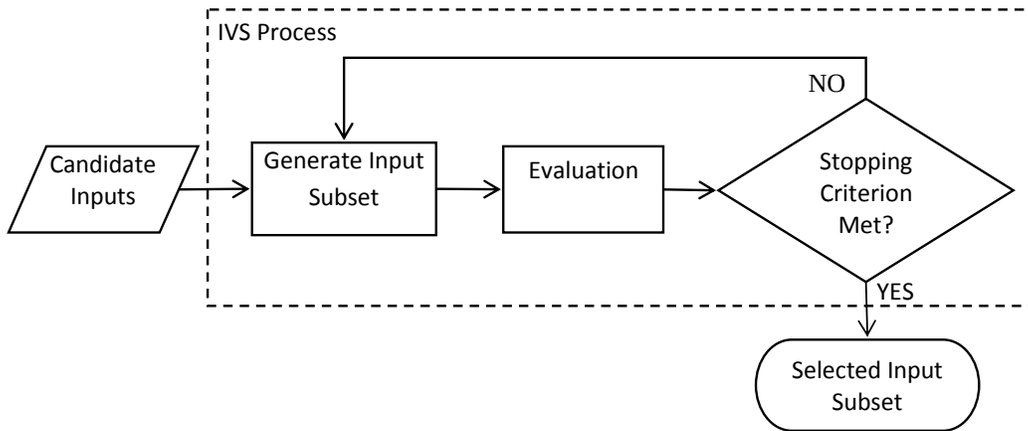


Figure 1: The generic IVS process (adapted from Dash and Liu (1997)).

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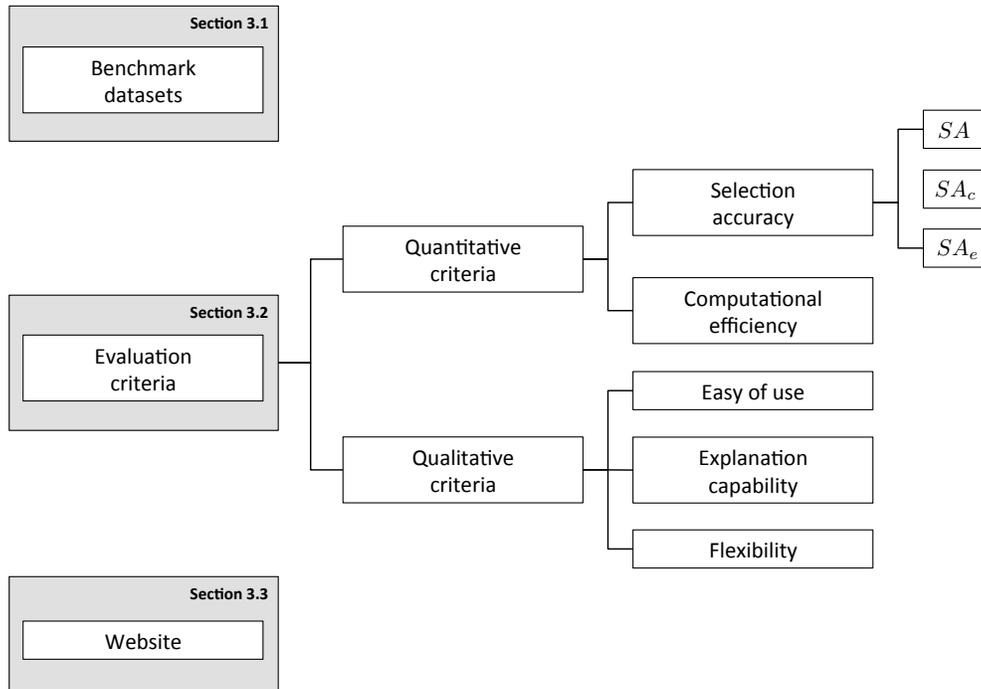


Figure 2: Schematic representation of the IVS framework components.

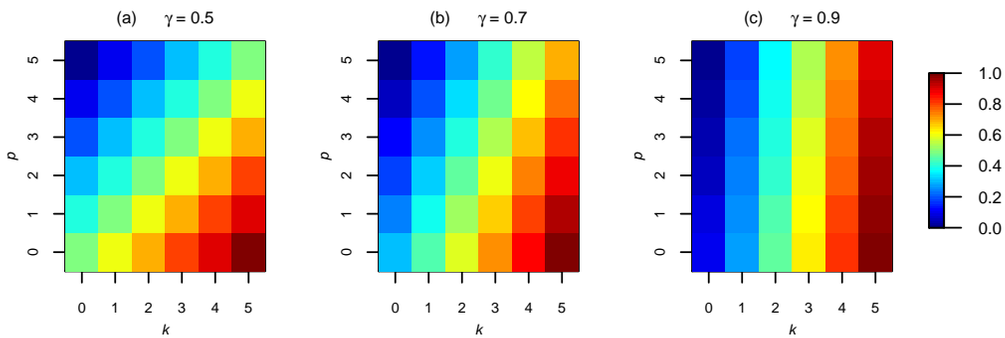


Figure 3: The effect of γ on SA score.

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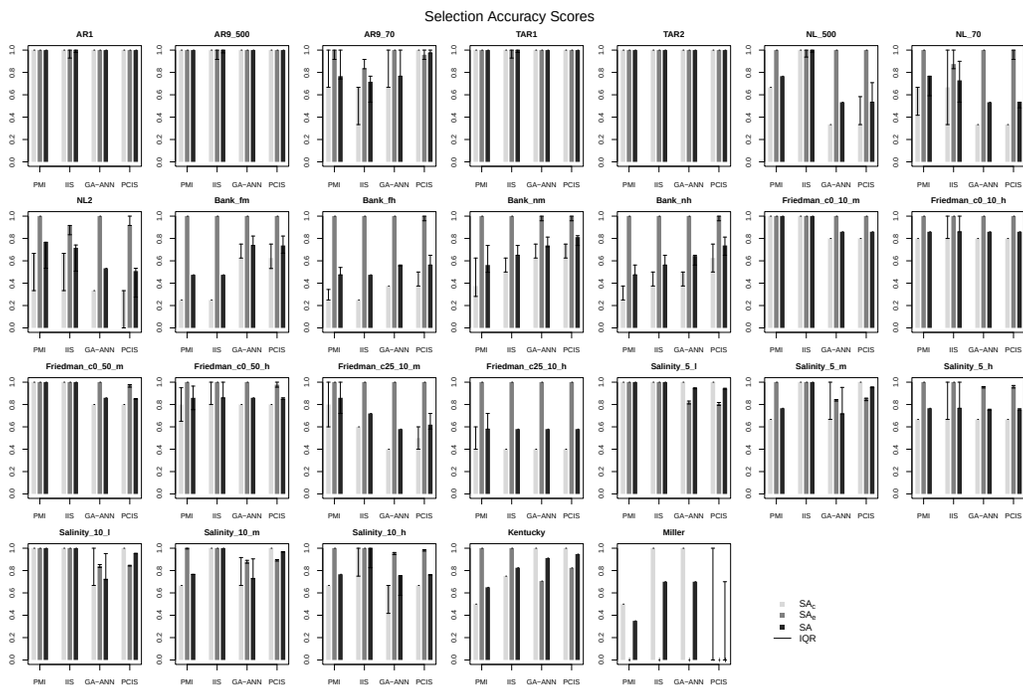


Figure 4: Bar charts representing the values of the scores SA , SA_c and SA_e obtained by running the PMIS, IIS, GA-ANN and PCIS algorithms on the 26 benchmark datasets.

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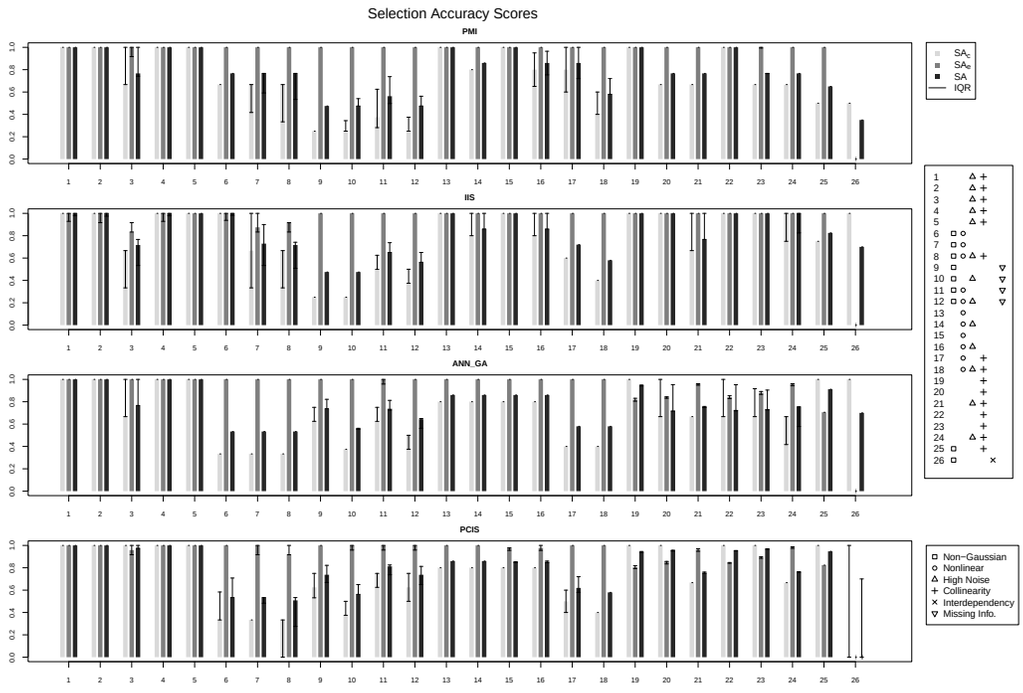


Figure 5: Bar charts representing the values of the scores SA , SA_c and SA_e obtained by running the PMIS, IIS, GA-ANN and PCIS algorithms on the 26 benchmark datasets. The datasets properties are described on the right-hand side.

LaTeX Source Files

[Click here to download LaTeX Source Files: IVS_framework_revised.tex](#)

Figure 1

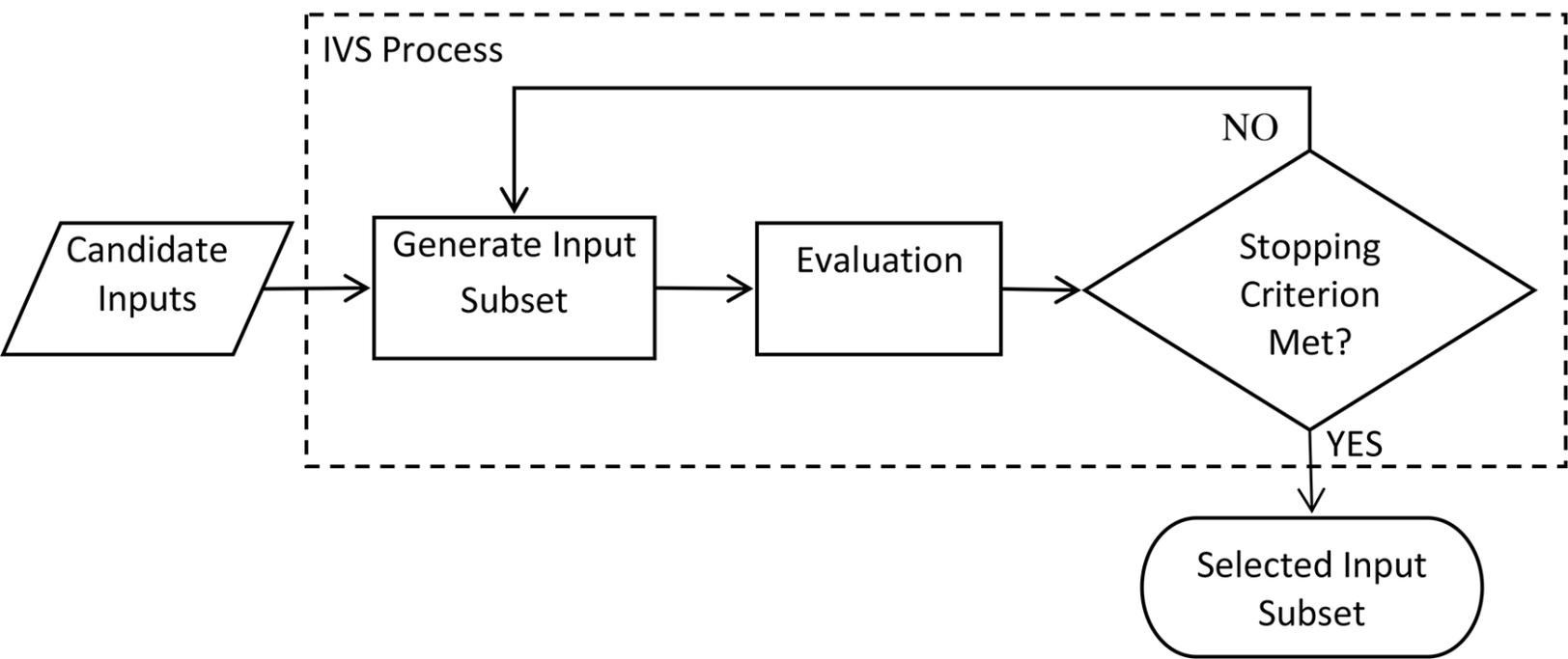


Figure 2

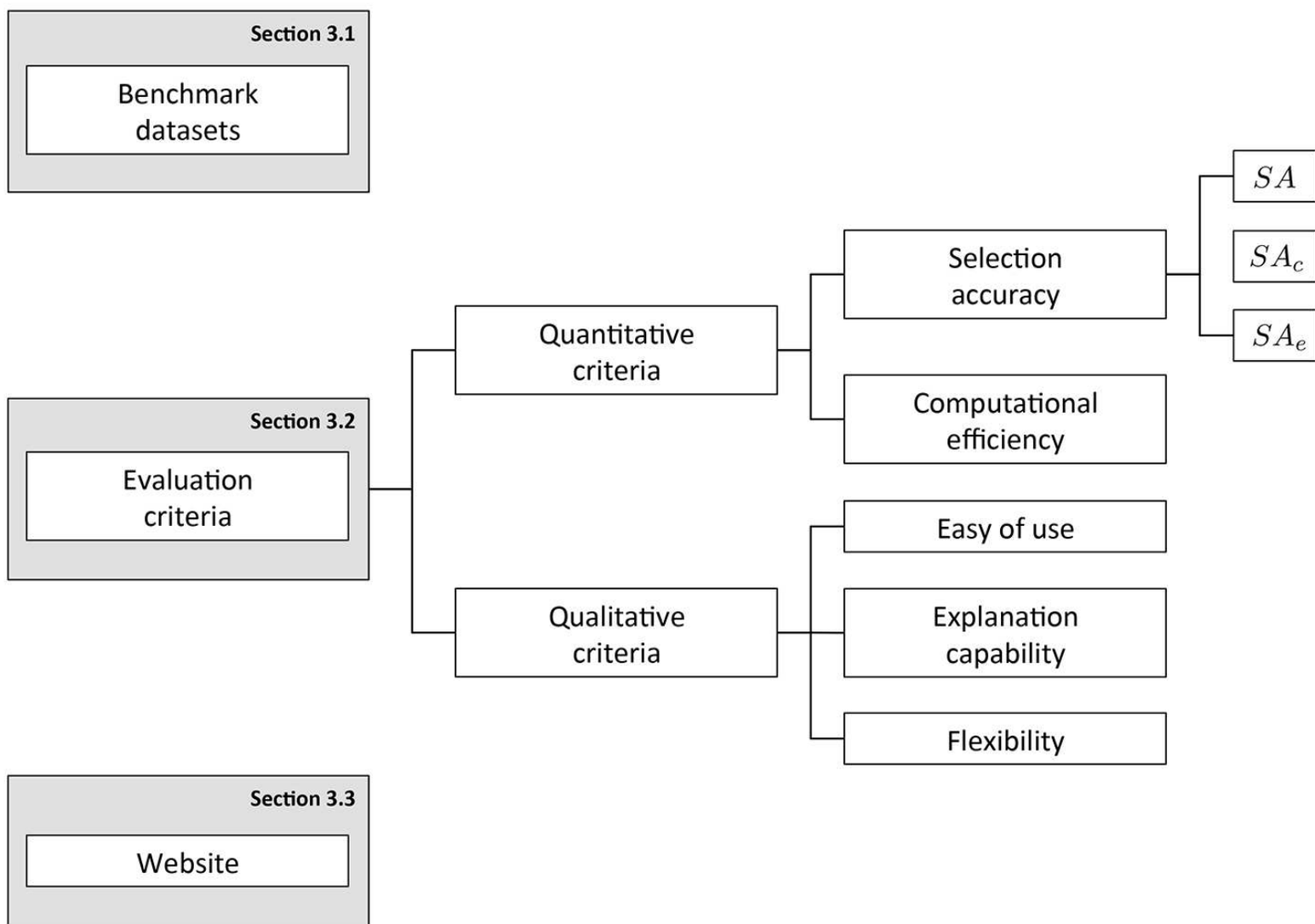


Figure 3

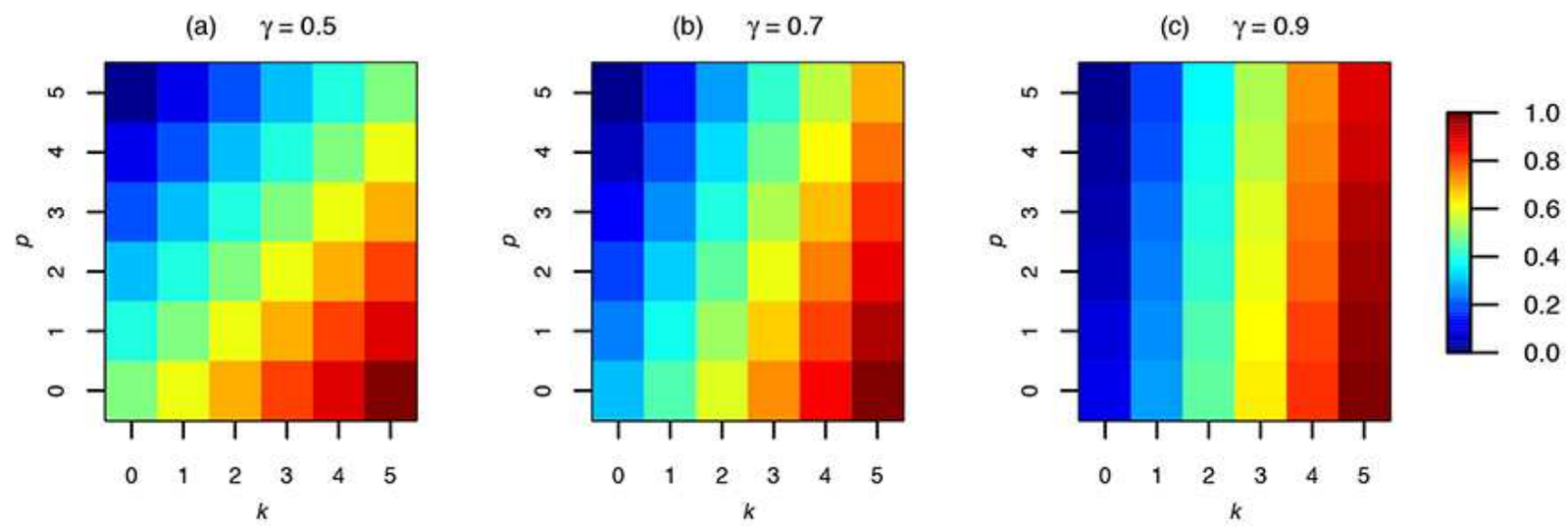


Figure 4

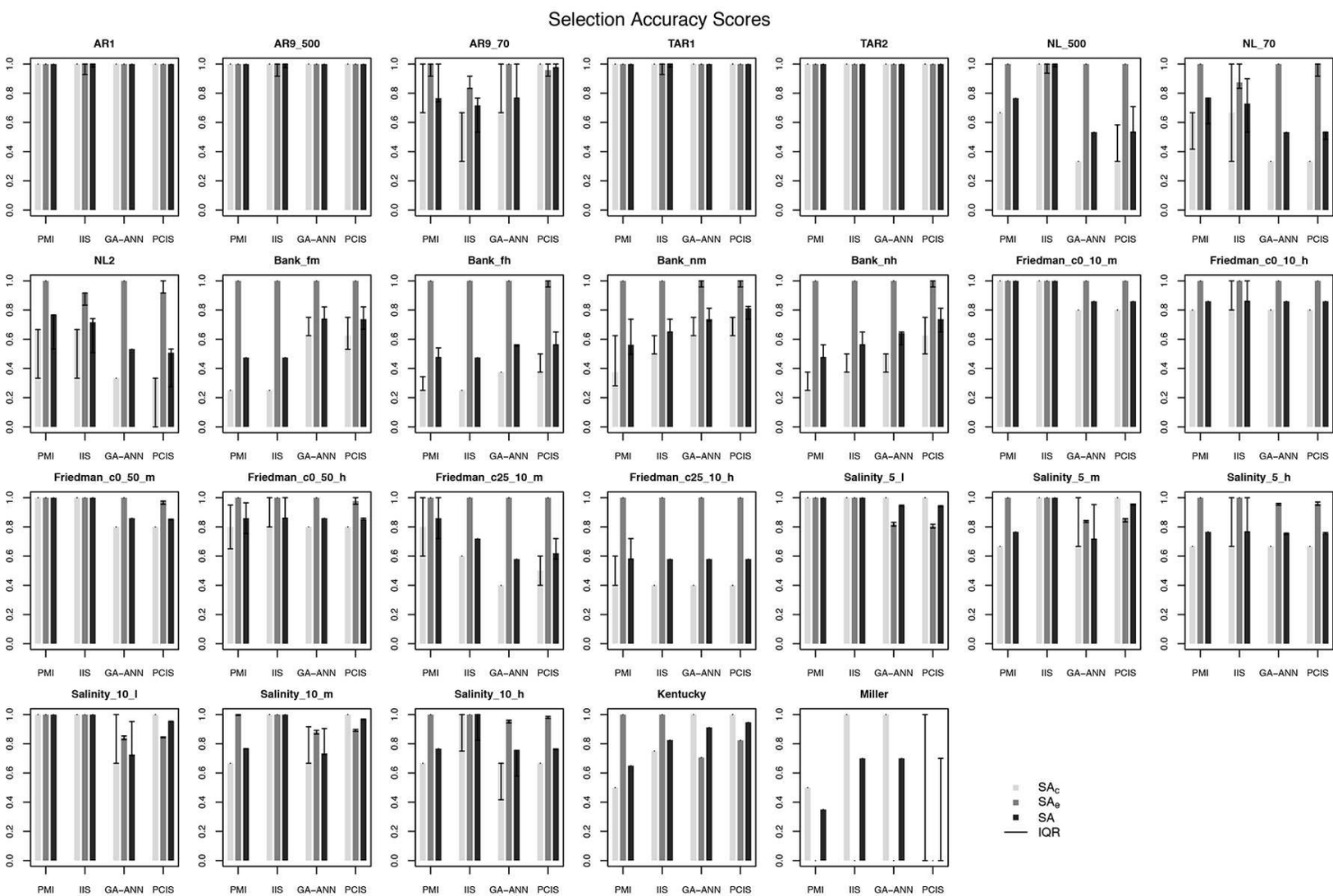


Figure 5

