Pitfalls and Potentials in Simulation Studies

Questionable research practices in comparative simulation studies allow for spurious claims of superiority of any method

Samuel Pawel*†, Lucas Kook*, Kelly Reeve

Epidemiology, Biostatistics and Prevention Institute

Center for Reproducible Science

University of Zurich, Hirschengraben 84, CH-8001 Zurich
{samuel.pawel, lucasheinrich.kook, kelly.reeve}@uzh.ch

Abstract

Comparative simulation studies are workhorse tools for benchmarking statistical methods, but if not performed and reported transparently they may lead to overoptimistic or misleading conclusions. The current publication requirements adopted by statistics journals do not prevent questionable research practices such as selective reporting. There have been numerous suggestions and initiatives to improve on these issues but little progress can be seen to date. In this paper we discuss common questionable research practices which undermine the validity of findings from comparative simulation studies. To illustrate our point, we invent a novel prediction method with no expected performance gain and benchmark it in a pre-registered comparative simulation study. We show how easy it is to make the method appear superior over well-established competitor methods if no protocol is in place and various questionable research practices are employed. Finally, we provide researchers, reviewers, and other academic stakeholders concrete suggestions for improving the methodological quality of comparative simulation studies, most importantly the need for pre-registered simulation protocols.

Keywords: benchmarking studies, Monte Carlo experiments, overoptimism, reproducibility, replicability, transparency

1 Introduction

"The first principle is that you must not fool yourself and you are the easiest person to fool. So you have to be very careful about that. After you've not fooled yourself, it's easy not to fool other scientists."

Feynman (1974, p. 12)

Simulation studies are to a statistician what experiments are to a scientist (Hoaglin and Andrews, 1975). They have become a ubiquitous tool for the evaluation of statistical methods, mainly because simulation can be used for studying the statistical properties of methods under conditions that would be difficult or impossible to study theoretically. In this paper we focus on simulation studies where

Preprint. Version September 1, 2022. Licensed under CC-BY.

^{*}Contributed equally.

[†]Corresponding author: samuel.pawel@uzh.ch

the objective is to compare the performance of two or more statistical methods (*comparative simulation studies*). Such studies are needed to ensure that previously proposed methods work as expected under various conditions, as well as to identify conditions under which they fail. Moreover, evidence from comparative simulation studies is often the only guidance available to data analysts for choosing from the plethora of available methods (Boulesteix et al., 2013, 2017). Proper design and execution of comparative simulation studies is therefore important, and results of methodologically flawed studies may lead to misinformed decisions in scientific and medical practice.

Just like non-simulation based studies, comparative simulation studies require many decisions to be made, for instance: How will the data be generated? How often will a simulation condition be repeated? Which statistical methods will be compared and how are their parameters specified? How will the performance of the methods be evaluated? The degree of flexibility, however, is much higher for simulation studies than for non-simulation based studies as they can often be rapidly repeated under different conditions at practically no additional cost. This is why numerous guidelines and best practices for design, execution, and reporting of simulation studies have been proposed (Hoaglin and Andrews, 1975; Holford et al., 2000; Burton et al., 2006; Smith and Marshall, 2010; O'Kelly et al., 2016; Monks et al., 2018; Elofsson et al., 2019; Morris et al., 2019; Boulesteix et al., 2020). We recommend Morris et al. (2019) for an introduction to state-of-the-art simulation study methodology.

Despite wide availability of such guidelines, statistics articles often provide too little detail about the reported simulation studies to enable quality assessment and replication (see *e.g.*, the literature reviews in Burton et al., 2006; Morris et al., 2019). Journal policies sometimes require the computer code to reproduce the results, but they rarely require or promote rigorous simulation methodology (*e.g.*, the preparation of a simulation protocol). This leaves researchers with considerable flexibility in how they conduct and present simulations studies. As a consequence, readers of statistics papers can rarely be sure of the quality of evidence that a simulation study provides.

Unfortunately, there are many questionable research practices (QRPs) which may undermine the validity of comparative simulations studies and which can easily go undetected under current standards. There is often a fine line between QRPs and legitimate research practices. For instance, there are good reasons to modify the data-generating process of a simulation study based on the observed results, *e.g.*, if the initially considered data-generating process results in many missing or non-convergent simulations. However, it is then important that such *post hoc* modifications are transparently reported. These practices only become questionable when they serve to confirm the hopes and beliefs of researchers regarding a particular method. Consequently, the results and conclusions of the study will be biased in favor of this method (Nießl et al., 2021).

The aim of this paper is to raise awareness about the issue of QRPs in comparative simulation studies, and to highlight the need for the adoption of higher standards. While researchers may make decisions that can make the conclusions of simulation studies misleading, we are not accusing them of doing so intentionally or maliciously but highlighting how this can happen and how to prevent it. External pressures, *e.g.*, to publish novel and superior methods (Boulesteix et al., 2015) or to concisely report large amounts of simulation results, may also lead honest researchers to (unknowingly) employ QRPs. As we will argue, it is not only up to the researchers but also other academic stakeholders to improve on these issues.

This article is structured as follows: We first give an illustrative list of QRPs related to comparative simulation studies (Section 2). With an exemplary simulation study, we then show how easy it is to present a novel, made-up method as an improvement over others if QRPs are employed and *a priori*

simulation plans remain undisclosed (Section 3). The main inspiration for this work is drawn from similar illustrative studies which have been conducted by Jelizarow et al. (2010), Nießl et al. (2021), and Ullmann et al. (2022) in the context of benchmarking studies with real data sets and by Simmons et al. (2011) in the context of p-hacking in psychological research. In Section 4, we then provide concrete suggestions for researchers, reviewers, editors, and funding bodies to alleviate the issues of QRPs and improve the methodological quality of comparative simulation studies. Section 5 closes with a discussion of the results and concluding remarks.

2 Questionable research practices in comparative simulation studies

There are various QRPs which threaten the validity of comparative simulation studies (see Table 1 for an overview). QRPs can be categorized with respect to the stage of research at which they can occur and which other QRPs they are related with (Wicherts et al., 2016). Typically, QRPs becomes more problematic if they are combined with related QRPs. For example, adapting the data-generating process to achieve a desired outcome (E2) is more problematic when the results based on the adapted process are selectively reported (R2) compared to reporting the results based on both the original and the adapted process. In the following, we describe QRPs from all phases of a simulation study, namely, design, execution, and reporting.

2.1 QRPs in the design of comparative simulation studies

The *a priori* specification of research hypotheses, study design, and analytic choices is what separates *confirmatory* from *exploratory* research. Evidence from confirmatory research is typically considered more robust because study hypotheses, design, and analysis are independent of the observed data (Tukey, 1980). The line between the two types of research is, however, blurry in simulation studies since they are often iteratively conducted, with each iteration including newly simulated data and building on the results of the previous study. The first simulation study in a sequence of studies may thus be exploratory whereas the subsequent studies may be confirmatory. Yet, one may argue that in many cases a single confirmatory simulation study which is carefully designed and whose design is justified based on external knowledge provides more relevant evidence than a sequence of simulation studies which are iteratively tweaked based on previous results.

To allow readers to distinguish between confirmatory and exploratory research, many non-methodological journals require pre-registration of study design and analysis protocols. For instance, pre-registration is common practice in randomized controlled clinical trials (Angelis et al., 2004), and increasingly adopted in experimental psychology (Nosek et al., 2018) and epidemiology (Lawlor, 2007; Loder et al., 2010). It is also generally recommended to write and pre-register simulation protocols in simulation studies (Morris et al., 2019). Well-defined study aims and methodology are arguably even more important in simulation studies compared to non-simulation based studies because the space of possible design and analysis choices is typically much larger (Hoffmann et al., 2021). In contrast, if researchers are vague or fail to define the study goals (D1), the data-generating process (D2), the methods under investigation (D3), the estimands of interest (D4), the evaluation metrics (D5), or how missing values should be handled (D6) a priori a high number of researcher degrees of freedom (Simmons et al., 2011) are left open. Researchers can then generate a multiplicity of possible results which may foster overoptimistic impressions if they report only the subset of results aligning with their hopes and belief (R2), and for which they can find plausible justifications post hoc (R1).

Table 1: Types of questionable research practices (QRPs) in comparative simulation studies at different stages of the research process. A QRP becomes more problematic if combined with a related QRP, especially a reporting QRP.

Tag	Related	Type of QRP
Design		
D1	E1, R1	Not/vaguely defining objectives of simulation study
D2	E2, R1	Not/vaguely defining data-generating process
D3	E3, E4, R1	Not/vaguely defining which methods will be compared and how their parameters are specified
D4	E1, E5, R1	Not/vaguely defining estimands of interest
D5	E1, E5, R1	Not/vaguely defining evaluation criteria
D6	E6, R1	Not/vaguely defining how to handle missing values (e.g., due to non-convergence of methods)
D7	E7, E8, R3	Not computing required number of simulations to achieve adequate precision
Execution		
E1	D1, R2	Changing objective of the study to achieve desired outcomes
E2	D2, R2	Adapting data-generating process to achieve desired outcomes
E3	D3, R2	Adding/removing comparison methods to achieve desired outcomes
E4	D3, R2	Selective tuning of method hyperparameters to achieve desired outcomes
E5	D4, D5, R2	Choosing evaluation criteria to achieve desired outcomes
E6	D6, R2	Adapting inclusion/exclusion/imputation rules to achieve desired outcomes
E7	D7, R3	Choosing number of simulations to achieve desired outcomes
E8	D7, R3	Choosing random seed to achieve desired outcomes
Reporting		
R1	D1-D6	Justifying design decisions which lead to desired outcomes post hoc
R2	E1-E6	Selective reporting of results from simulations that lead to desired outcomes
R3	D7, E7, E8	Failing to report Monte Carlo uncertainty
R4		Failing to assure computational reproducibility (<i>e.g.</i> , not sharing code and sufficient details about computing environment)
R5		Failing to assure replicability (e.g., not sufficiently reporting design and execution methodology)

Another crucial part of rigorous design is simulation size calculation (see Section 5.3 in Morris et al. (2019) for an overview). A thorough planning of the number of simulations in terms of expected precision of the primary estimand is important. While an arbitrarily chosen, often too small, number of simulations can be executed faster, they yield noisier results. If claims of superiority are based on *e.g.*, a confidence interval for the difference in method performance excluding zero, "true" differences in method performance are more likely to remain undetected for undersized studies. Furthermore, detected differences in method performance from undersized simulation studies are more likely to be the in the wrong direction and their magnitude is more likely to be overestimated. These drawbacks parallel the undesirable properties of underpowered non-simulation based studies (increased type II, type S, and type M error, see *e.g.*, Gelman and Tuerlinckx, 2000; Gelman and Carlin, 2014; van Zwet and Cator, 2021). By failing to conduct a simulation size calculation (D7), researchers are thus at a higher risk of drawing the wrong conclusions (if their sample size is too small), or wasting computer resources (if their sample size is too large).

2.2 QRPs in the execution of comparative simulation studies

During the execution of a simulation study researchers may (often unknowingly) engage in various QRPs that can lead to overoptimism. For instance, the objective of the simulation study may be changed depending on the outcome (E1), e.g., an initial comparison of predictive performance may be changed to comparing estimation performance if the results suggest that the favored method performs better at estimation tasks rather than prediction. The data-generating process may also be adapted until conditions are found in which the favored method appears superior (E2). For example, the noise levels, the number of covariates, or the effect sizes could be changed. Competitor methods that are superior to the proposed method may also be excluded from the comparison altogether, or methods which perform worse under the (adapted) data-generating process may be added (E3). The methods under comparison may come with hyperparameters (e.g., regularization parameters in penalized regression models). In this case, the hyperparameters of a favored method may be tuned until the method appears superior, or the hyperparameters of competitor methods may be tuned selectively, e.g., left at their default values (E4). Finally, the evaluation criteria for comparing the performance of the investigated methods may also be changed to make a particular method look better than the others (E5). For example, even though the original aim of the study may have been to compare predictive performance among methods using the Brier score, the evaluation criterion of the simulation study may be switched to area under the curve if the results suggest that the favored method performs better with respect to the latter metric. This QRP parallels the well-known "outcome-switching" problem in clinical trials (Altman et al., 2017). It is usually not difficult to find reasonable justification for such modifications and then present them as if they were specified during the planning of the study (R1). As emphasized earlier, iteratively changing simulation goals, conditions, methods under comparison, and evaluation criteria can be part of finding out how a method works. These practices become mostly problematic if only the simulations in line with the researchers hopes and beliefs are reported (R2).

There are, however, practices which are considerably more problematic on their own. For instance, in some simulations a method may fail to converge and thus produce missing values in the estimates. If it is not pre-specified how these situations will be handled, different inclusion/exclusion or imputation strategies may be tried out until a favored method appears superior (E6). Choosing an inadequate strategy can result in systematic bias and misleading conclusions. If no *a priori* simulation size calculation was conducted, the simulation size may also be changed until favorable results are obtained (E7). If in that case the number of simulations is too small, true performance differences are more likely to be missed, their estimated direction is more likely to be incorrect, and their magnitude is more likely overestimated, as explained previously. Finally, if only few simulations are conducted (*e.g.*, because the methods under investigation are computationally very expensive), the initializing seed for generating random numbers may have a substantial impact on the result. A particularly questionable practice in this situation is to tune the seed until a value is found for which a preferred method seems superior (E8).

2.3 QRPs in the reporting of comparative simulation studies

In the reporting stage, researchers are faced with the challenge of reporting the design, results, and analyses of their simulation study in a digestible manner. Various QRPs can occur at this stage. For instance, reporting may focus on results in which the method of interest performs best (R2). Failing to mention conditions in which the method was inferior (or at least not superior) to competitors creates overoptimistic impressions, and may lead readers to think that the method uniformly outperforms com-

petitors. Similarly, presenting simulation conditions which were added based on the observed results as pre-planned and justified (R1) fosters overconfidence in the results.

Another crucial aspect of reporting is to adequately show the uncertainty related to the simulation results (Hoaglin and Andrews, 1975; Van der Bles et al., 2019). Failing to report Monte Carlo uncertainty (R3), *e.g.*, error bars or confidence intervals reflecting uncertainty in the simulation, hampers the readers' ability to assess the accuracy of the results from the simulation study and it allows one to present random differences in performance as if they were systematic.

Finally, by failing to assure computational reproducibility of the simulation study (R4), for example, by not sharing code and software versions to run the simulation, it is more likely that coding errors remain undetected. By not reporting the design and execution of the study in enough detail (R5), other researchers are unable to replicate and expand on the simulation study. Unclear reporting makes it also harder for readers to identify potentially overoptimistic statements. For instance, if it is reported that all but one method are left at their default parameters, readers can better contextualize this method's apparent superior performance.

3 Empirical study: The Adaptive Importance Elastic Net (AINET)

To illustrate the application of QRPs from Table 1 we conducted a simulation study. The objective of the study was to evaluate the predictive performance of a made-up regression method termed the "adaptive importance elastic net" (AINET). The main idea of AINET is to use variable importance measures from a random forest for a weighted penalization of the variables in an elastic net regression model. The hope is that this *ad hoc* modification of the elastic net model improves predictive performance in clinical prediction modeling settings where penalized regression models are frequently used. Superficially, AINET may seem sensible, however, for the data-generating process considered in our simulation study no advantage over the classical elastic net is expected. For more details on the method, we refer the reader to the simulation protocol (Appendix A). We report the per-protocol simulation study results in Appendix B. As expected, the performance of AINET was virtually identical to standard elastic net regression. AINET also did not yield any improvements over logistic regression for the data-generating process that we considered sensible *a priori* (*i.e.*, it was specified based on typical conditions in clinical prediction modeling and simulation studies from other researchers).

We now show how application of QRPs changes the above per-protocol conclusions. Figure 1 illustrates different types of QRPs sequentially applied to simulation-based evaluation of AINET. The top row depicts the per-protocol differences in Brier score (x-axis) between AINET and competitor methods (y-axis) for a representative subset of the simulation conditions. A negative difference indicates superior performance of AINET. In the second row, the arrows depict the change in the per-protocol results after changing the data-generating process (E2). The third row shows the result after removal of the elastic net competitor (E3). Finally, the bottom row shows the end result where selective reporting of simulation conditions and competitor methods (R2) is applied to give a more favorable impression of AINET. We will now discuss these QRPs in more detail.

Altering the data-generating process (E2) We could not detect a systematic performance benefit of AINET over standard logistic regression, elastic net regression, or random forest for the scenarios specified in the protocol. For this reason, we tweaked the data-generating process by adding different

¹We use the term "per-protocol" throughout to refer to simulation analyses conducted as pre-specified in the protocol.

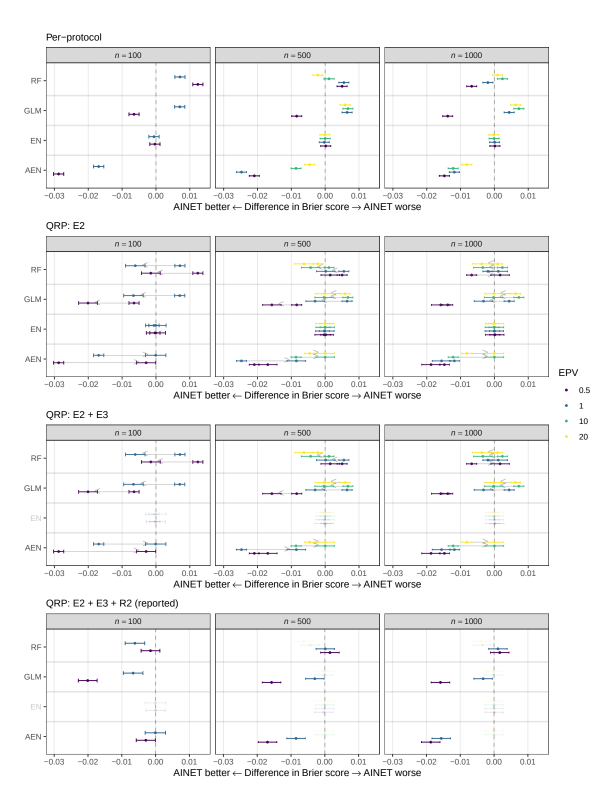


Figure 1: Differences in Brier score with 95% adjusted confidence intervals between AINET and random forest (RF), logistic regression (GLM), elastic net (EN), and adaptive elastic net (AEN) are shown for representative simulation conditions (correlated covariates $\rho=0.95$, prevalence prev = 0.05, a range of sample sizes n and events per variable (EPV), in each simulation the Brier score is computed for 10'000 test observations; for details see Appendix A). The top row depicts the per-protocol results in which AINET does not outperform any competitor uniformly, except AEN. In the second row, we apply QRP E2: altering the data-generating process by adding a non-linear effect and sparsity. The gray arrows point from the per-protocol result to the results under the tweaked simulation. In the third row, QRP E3 is applied: EN is removed as a competitor. In the bottom row, selective reporting R2 is applied: only low EPV settings are reported to give a more favorable impression for AINET. Arrows are depicted only for non-overlapping confidence intervals.

sparsity conditions and a non-linear effect. We then found that AINET outperforms logistic regression under the following conditions: only few variables being associated with the outcome (sparsity), a non-linear effect, and a low number of events per variable (EPV). Figure 1 (second row) shows the changes in Brier score difference between the pre-registered and the tweaked simulation. As can be seen, the tweaked data-generating process leads to AINET being superior to competitors in some conditions, and at least not inferior in others.

Removing competitor methods (E3) Despite the adapted data-generating process, we still observed only minor (if any) improvements of AINET over the elastic net. In order to present AINET in a better light we could omit the comparisons with the elastic net (E3), as shown in Figure 1 (third row). This could be justified, for example, by arguing that for neutral comparison it is sufficient to compare a less flexible method (logistic regression, which has no tuning parameters and captures linear effects), a more flexible method (random forest, which has tuning parameters and captures nonlinear relationships), and a comparably flexible method (adaptive elastic net, which has the same tuning parameters as AINET, but differs in the way the penalization weights are chosen).

Selective reporting of simulation results (R2) After the removal of the competitor elastic net, there are still some simulation conditions under which AINET is not superior to the remaining competitors. To make AINET appear more favorable, we thus report only simulation conditions with low EPV, as shown in Figure 1 (fourth row). This could be justified by the fact that journals require authors to be concise in their reporting. Moreover, further conditions with low EPV values could be simulated to make the results seem more exhaustive. Focusing primarily on low EPV settings could be justified in hindsight by framing AINET as a method designed for high-dimensional data (low sample size relative to the number of variables).

4 Recommendations

The previous sections painted a rather negative picture of how undisclosed changes in simulation design, analysis, and reporting may lead to overoptimistic conclusions. In the following, we summarize what we consider to be practical recommendations for improving the methodological quality of simulation studies; see Table 2 for an overview. Our recommendations are grouped with regards to which stakeholder they concern.

4.1 Recommendations for researchers

Adopting pre-registered simulation protocols is arguably the most important measure that researchers can take to prevent themselves from subconsciously engaging in QRPs. Pre-registration enables readers to distinguish between confirmatory and exploratory findings, and it lowers the risk of potentially flawed methods being promoted as an improvement over competitors. While pre-registered simulation protocols may at first seem disadvantageous due to the additional work and possibly lower chance of publication, they provide researchers with the means to differentiate their high-quality simulation studies from the numerous unregistered and possibly untrustworthy simulation studies in the literature. Platforms such as GitHub (https://github.com/), OSF (https://osf.io/), or Zenodo (https://zenodo.org/) can be used for archiving and time-stamping documents. Moreover, pre-registration can also

Table 2: Recommendations for improving quality of comparative simulation studies and preventing QRPs.

Researchers

- Adopt (pre-registered) simulation protocols
- Adopt good computational practices (e.g., code review, packaging, unit-tests)
- Share code and data (possibly in intermediate/summary form to enable secondary analysis)
- Report the process of the simulation study fully and transparently (e.g., time-stamped protocol
 amendments to disclose pilot studies and post hoc modifications)
- Perform simulation analysis in a blinded manner
- Collaborate with other research groups (with possibly "competing" methods)
- Disclose multiplicity and uncertainty of results (e.g., with sensitivity analyses)
- Teach simulation study methodology in statistics (post)graduate courses

Editors and reviewers

- Require/encourage exploration of conditions where methods should be inferior or break down
- Require/encourage (pre-registered) simulation protocols
- Provide enough space for description of simulation methodology

Journals and funding bodies

- Provide incentives for rigorous simulation studies (e.g., badges on papers)
- Require code and data
- Enforce adherence to reporting guidelines
- Adopt reproducibility checks
- Promote/fund research and software to improve simulation study methodology
- Deincentivize outperforming state-of-the-art methods

save researchers from some work later on, e.g., they can usually copy most parts of the simulation methodology description from the protocol to the final manuscript.

When pre-registering and conducting simulation studies, we recommend using a robust computational workflow. Such a workflow encompasses packaging the software, writing unit tests, and reviewing code (see *e.g.*, Schwab and Held, 2021). Other researchers and the authors themselves then benefit from improved computational reproducibility and less error-prone code. Of course, there are also certain practical limits to computational reproducibility. For instance, if a simulation study requires high performance computing and/or several weeks of running time, the authors should not expect reviewers and journals to replicate their simulation study from scratch. The authors should nevertheless provide the code to run the simulation and, if possible, they should also provide intermediate simulation results (*e.g.*, fitted model objects) so that the simulation study can at least be partially reproduced. Similarly, authors can share the simulated data, either in raw and/or some summarized form (*e.g.*, sharing simulated data sets and parameter estimates of fitted models). This allows interested readers and reviewers to do additional analyses. Unlike experiments with human subjects, there are no privacy concerns for sharing simulation data. Furthermore, online tools, such as INTEREST (INteractive Tool for Exploring REsults from Simulation sTudies, Gasparini et al., 2021), can be used for interactive exploration of the data set.

While planning a simulation study, it is impossible to think of all potential weaknesses or problems that may arise when conducting the planned simulations. In turn, researchers may be reluctant to tie their hands in a pre-registered protocol. However, a transparently conducted and reported preliminary simulation can obviate most of these problems. We recommend researchers to disclose preliminary results and any resulting changes to the protocol, *e.g.*, in a revised and time-stamped version of the protocol. This approach is similar to conducting a small pilot study, as is often done in non-simulation based research. Even if researchers realize that further changes are required after the main simulation study has begun, transparent reporting of when and why *post hoc* modifications were made allows the reader to better assess the quality of evidence provided by the study. Researchers designing simulation studies may draw inspiration from clinical trials by tracking their protocol modifications and time-stamping versions of their protocol.

A different approach for making *post hoc* changes to the protocol is to use blinding in the analysis of the simulation results (Dutilh et al., 2019). Blinded analysis is a standard procedure in particle physics to prevent data analysts from biasing their result towards their own beliefs (Klein and Roodman, 2005), and it lends legitimacy to *post hoc* modifications of the simulation study. For instance, researchers might shuffle the method labels and only unblind themselves after the necessary analysis pipelines are set in place. An alternative blinding approach is to carry out data generation and analysis by different researchers. For instance, the study from Kreutz et al. (2020) involved two independent research groups, one who simulated and one who analyzed the data. A related way for improving simulation studies is to collaborate with other researchers, possibly ones familiar with "competing" methods. This helps to design simulation studies which are more objective and whose results are more useful for making a decision about which method to choose.

We also recommend researchers to disclose the multiplicity and uncertainty inherent to the design and analysis of their simulation studies (Hoffmann et al., 2021). For instance, researchers can report sensitivity analyses that show how the study results change for different analysis decisions (*e.g.*, Table 4 in van Smeden et al. (2016) shows how the evaluation metrics for different estimators change depending on how convergence of a method is defined). Methods from multivariate statistics can be used for visualizing the influence of different design choices, *e.g.*, a multidimensional unfolding approach as shown by Nießl et al. (2021).

One reason for the low standards of simulation studies in the statistics literature may be that rigorous simulation methodology is usually not taught in graduate or postgraduate courses (with a few exceptions, *e.g.*, the course "Using simulation studies to evaluate statistical methods" from the MRC Clinical Trials Unit). To improve training of current and future generations of statisticians, researchers who are involved in teaching should therefore also include simulation study methodology in their curricula. The standards of simulation studies in many statistics related fields (*e.g.*, machine learning, psychometrics, econometrics, or ecology) are arguably not much different. One possible avenue for future research is thus to also provide education and adaption of simulation study methodology for the special needs in these fields.

4.2 Recommendations for editors and reviewers

Peer review is an important tool for identifying QRPs in research results submitted to methodological journals. For instance, reviewers may demand researchers to include competitor methods which are not part of their comparison yet (or which might have been excluded from the comparison). However, reviewers can only identify a subset of all QRPs since some types are impossible to spot if no preregistered simulation protocol is in place (*e.g.*, a reviewer cannot know whether the evaluation criterion

was switched). Even QRPs which can be detected by peer review may be difficult to spot in practice. It is thus important that reviewers and editors demand that authors make simulation protocols and computer code available alongside the manuscript. Moreover, by providing enough space and encouraging authors to provide detailed descriptions of their simulation studies, replicability of the simulation studies can be improved. Finally, reviewers should not be satisfied with manuscripts showing that a method is uniformly superior; they should also encourage authors to explore conditions in which their method is expected to be inferior to other methods or to break down entirely.

4.3 Recommendations for journals and funding bodies

Journals and funding bodies can improve on the status quo by either actively requiring or passively incentivizing more rigorous and neutral simulation study methodology. Actively, journals can make (pre-registered) simulation protocols mandatory for all articles featuring a simulation study. A more passive and less extreme measure would be to indicate with a badge whether an article contains a preregistered simulation study, or to introduce article types dedicated to neutral comparison studies. Such an approach rewards researchers who take the extra effort. Similar initiatives have led to a large increase in the adoption of pre-registered study protocols in the field of psychology (Kidwell et al., 2016). Another measure could be to require standardized reporting of simulation studies, e.g., the "ADEMP" reporting structure proposed by Morris et al. (2019). Journals may also employ reproducibility checks to ensure computational reproducibility of the published simulation studies. This is already done, for example, by the Journal of Open Source Software or the Journal of Statistical Software. Moreover, journals and funding bodies can promote or fund research and software to improve simulation study methodology. For instance, a journal might have special calls for papers on simulation methodology. Similarly, a funding body could have special grants dedicated to software development that facilitates sound design, execution, and reporting of simulation studies (as, for example, White, 2010; Gasparini, 2018; Chalmers and Adkins, 2020). Finally, journals and funding bodies often exert a strong incentive on researchers to publish novel and superior methods. This may lead to articles with non-systematic simulation studies that mainly highlight settings beneficial to the proposed methods. We believe that the above recommendations can shift the incentive structure towards more neutral and transparent conduct and reporting of simulation studies.

5 Conclusions

Simulation studies should be viewed and treated analogously to (empirical) experiments from other fields of science. Transparent reporting of methodology and results is essential to contextualize the outcome of such a study. As in other empirical sciences, QRPs in simulation studies can obfuscate the usefulness of a novel method and lead to misleading and non-replicable results.

By deliberately using several QRPs we were able to present a method with no expected benefits and little theoretical justification – invented solely for this article – as an improvement over theoretically and empirically well-established competitors. While such intentional engagement in these practices is far from the norm, unintentional QRPs may have the same detrimental effect. We hope that our illustration will increase awareness about the fragility of findings from simulation studies and the need for higher standards.

While this article focused on comparative simulation studies, many of the issues and recommendations also apply to neutral comparison studies with real data sets as discussed in Nießl et al. (2021). Some

of the noted problems even exist in theoretical research; due to the incentive to publish positive results, researchers often selectively study optimality conditions of methods rather than conditions under which they fail.

Again, it is imperative to note that researchers rarely engage in QRPs with malicious intent but because humans tend to interpret ambiguous information self-servingly, and because they are good at finding reasonable justifications that match their expectations and desires (Simmons et al., 2011). As in other domains of science, it is easier to publish positive results in methodological research, i.e., novel and superior methods (Boulesteix et al., 2015). Thus, methodological researchers will typically desire to show the superiority of a method rather than to disclose its strengths and weaknesses. Aligning incentives for individual researchers with rigorous simulation research will require a range of actions involving various stakeholders in the research community. We have provided some recommendations that, we believe, could help achieve this goal. Most importantly, we think that reviewers, journals, and funders need to raise the standards for simulation studies by requiring pre-registered simulation protocols and rewarding researchers who invest the extra effort. Although there is evidence for the effectiveness of protocols in preventing QRPs from other fields, it is unclear whether this effect generalizes to simulation studies. Regardless of their effectiveness, we think that pre-registered protocols are a critical step toward improving simulation studies since they ensure a minimum degree of transparency and credibility when studies are conducted honestly. Of course, they cannot prevent fraudulent behaviors, such as researchers engaging in QRPs until they find their desired results and only then writing and registering a protocol. Pre-registered protocols alone are thus not sufficient to solve the issue of QRPs in simulation studies, but we believe they are a necessary step to improve them.

Software and data

The simulation study was conducted in the R language for statistical computing (R Core Team, 2020) using the version 4.1.1. The method AINET is implemented in the **ainet** package and available on GitHub (https://github.com/LucasKook/ainet). We provide scripts for reproducing the different simulation studies on the GitHub repository (https://github.com/SamCH93/SimPaper). Due to the computational overhead, we also provide the resulting data so that the analyses can be conducted without rerunning the simulations. We used **pROC** version 1.18.0 to compute the AUC (Robin et al., 2011). Random forests were fitted using **ranger** version 0.13.1 (Wright and Ziegler, 2017). For penalized likelihood methods, we used **glmnet** version 4.1.2 (Friedman et al., 2010; Simon et al., 2011). The **SimDesign** package version 2.7.1 was used to set up simulation scenarios (Chalmers and Adkins, 2020).

Acknowledgements

We thank Eva Furrer, Malgorzata Roos, and Torsten Hothorn for helpful discussion and comments on the simulation protocol and drafts of the manuscript. We also thank the anonymous referees and the associate editor for constructive and valuable comments that improved the manuscript substantially. Our acknowledgement of these individuals does not imply their endorsement of this article. The authors declare that they do not have any conflicts of interest. SP acknowledges financial support from the Swiss National Science Foundation (Project #189295). The funder had no role in study design, data collection, data analysis, data interpretation, decision to publish, or preparation of the manuscript.

Appendix A Simulation protocol

Below, we include an excerpt of the final version of the protocol for the simulation-based evaluation of AINET. All time-stamped versions of the protocol are available at https://doi.org/10.5281/zenodo.6364575.

A.1 Aims

The aim of this simulation study is to systematically study the predictive performance of AINET for a binary prediction task. The simulation conditions should resemble typical conditions found in the development of prediction models in biomedical research. In particular we want to evaluate the performance of AINET conditional on

- low- and high-dimensional covariates
- (un-)correlated covariates
- small and large sample sizes
- varying baseline prevalences

AINET will be compared to other (penalized) binary regression models from the literature, namely

- Binary logistic regression: the simplest and most popular method for binary prediction
- Elastic net: a generalization of LASSO and ridge regression, the most widely used penalized regression methods
- Adaptive elastic net: a generalization of the most popular weighted penalized regression method (adaptive LASSO)
- Random forest: a popular, more flexible method. This method is related to AINET, see Section A.4.

These cover a wide range of established methods with varying flexibility and serve as a reasonable benchmark for AINET. There are many more extensions of the adaptive elastic net in the literature (see *e.g.*, the review by Vidaurre et al., 2013). However, most of these extensions focus on variable selection and estimation instead of prediction, which is why we restrict our focus only on the four methods above.

A.2 Data-generating process

In each simulation $b=1,\ldots,B$, we generate a data set consisting of n realizations, i.e., $\{(y_i,x_i)\}_{i=1}^n$. A datum (Y,X) consists of a binary outcome $Y \in \{0,1\}$ and p-dimensional covariate vector $X \in \mathbb{R}^p$. The binary outcomes are generated by

$$Y \mid x \sim \text{Bernoulli}\left(\text{expit}\left\{ eta_0 + x^ op oldsymbol{eta}
ight\}
ight)$$

with $expit(z) = (1 + exp(-z))^{-1}$ and the covariate vectors are generated by

$$X \sim N_{v}(0,\Sigma)$$

with covariance matrix Σ that may vary across simulation conditions (see below). The baseline prevalence is prev = expit(β_0). The coefficient vector $\boldsymbol{\beta}$ is generated from

$$\beta \sim N_p(0, Id)$$

once per simulation. Finally, the simulation parameters are varied fully factorially (except for the removal of some unreasonable conditions) as described below, leading to a total of 128 scenarios, see below.

Sample size

The sample size used in the development of predictions models varies widely (Damen et al., 2016). We will use $n \in \{100, 500, 1000, 5000\}$, which span typical values occurring in practice. Note that previous simulation studies usually chose sample size based on the implied number of events together with the number of covariates in the model for easier interpretation (van Smeden et al., 2018; Riley et al., 2018). We will use this approach in reverse to determine the dimensionality of the parameters below.

Dimensionality

Previous simulation studies showed that events per variable (EPV) rather than the absolute sample size n and dimensionality p influences the predictive performance of a method. We will therefore define the dimensionality p via EPV by

$$p = \frac{n \cdot \text{prev}}{\text{EPV}}$$

and $2 \le p \le 100$. If the above formula gives non-integer values, the next larger integer will be used for p. When the formula gives values above 100 or below 2, this simulation condition will be removed from the design. This is done because prediction models are in practice only multivariable models ($p \ge 2$), but at the same time the number of predictors is rarely larger than $p \ge 100$ (Kreuzberger et al., 2020; Seker et al., 2020; Wynants et al., 2020). The exception are studies considering complex data, such as images, omics, or text data which are not the focus here. The values EPV $\in \{20, 10, 1, 0.5\}$ are chosen to cover scenarios with small to large number of covariates (cf. van Smeden et al., 2018).

Collinearity in X

We distinguish between no, low, medium and high collinearity. The diagonal elements of Σ are given by $\Sigma_{ii}=1$ and the off-diagonal elements are set to $\Sigma_{ij}=\rho$, $\rho\in\{0,0.3,0.6,0.95\}$. These values cover the typical (positive) range of correlations.

Baseline prevalence

Different baseline prevalences $expit(\beta_0) \in \{0.01, 0.05, 0.1\}$ are considered, reflecting a reasonable range of prevalences for rare to common diseases/adverse events.

Test data

In order to test the out-of-sample predictive performance, we generate a test data set of $n_{\text{test}} = 10000$ data points in each simulation b.

A.3 Estimands

We will estimate different quantities to evaluate overall predictive performance, calibration, and discrimination, respectively. All methods will be evaluated on independently generated test data.

A.3.1 Primary estimand

• Brier score. We compute the Brier score as

$$\overline{\mathrm{BS}} = n_{\mathrm{test}}^{-1} \sum_{i=1}^{n_{\mathrm{test}}} (y_i - \hat{y}_i)^2,$$

where $\hat{y} = \widehat{\mathbb{P}}(Y = 1 \mid x)$. Lower values indicate better predictive performance in terms of calibration and sharpness. A prediction is well-calibrated if the observed proportion of events is close to the predicted probabilities. Sharpness refers to how concentrated a predictive distribution is (e.g., how wide/narrow a prediction interval is), and the predictive goal is to maximize sharpness subject to calibration (Gneiting, 2008). The Brier score is a proper scoring rule, meaning that it is minimized if a predicted distribution is equal to the data-generating distribution (Gneiting and Raftery, 2007). Proper scoring rules thus encourage honest predictions. The Brier score is therefore a principled choice for our primary estimand.

A.3.2 Secondary estimands

• Scaled Brier score. The scaled Brier score (also known as Brier skill score) is computed as

$$\overline{BS}^* = 1 - \overline{BS} / \overline{BS_0}$$

with $\overline{BS}_0 = \bar{y}(1-\bar{y})$ and \bar{y} the observed prevalence in the data set. The scaled Brier score takes into account that the prevalence varies across simulation conditions. Hence, the scaled Brier score can be compared between conditions (Schmid and Griffith, 2005; Steyerberg et al., 2019).

• Log-score. We compute the log-score on independently generated test data,

$$\overline{\text{LS}} = -n_{\text{test}}^{-1} \sum_{i=1}^{n_{\text{test}}} \{ y_i \log(\hat{y}_i) + (1 - y_i) \log(1 - \hat{y}_i) \},$$

will be used as a secondary measure of overall predictive performance. Lower values indicate better predictive performance in terms of calibration and sharpness. The log-score is a strictly proper scoring rule, however, it is more sensitive to extreme predicted probabilities compared to the Brier score (Gneiting and Raftery, 2007).

• AUC. The AUC is given by

$$AUC = \max\{PI, 1 - PI\}$$

with

$$PI = \widehat{\mathbb{P}}(Y_i \geq Y_j \mid x_i, x_j), i, j = 1, \dots, n_{test},$$

where Y_i and Y_j denote case and non-case, respectively. The AUC is related to the area under the receiver-operating-characteristic (ROC) curve (Steyerberg et al., 2019). It will be used as a

measure of discrimination and values closer to one indicate better discriminative ability. Discrimination describes the ability of a prediction model to discriminate between cases and non-cases. Other discrimination measures, such as accuracy, sensitivity, specificity, etc., are not considered because we want to evaluate predictive performance in terms of probabilistic predictions instead of point predictions/classification.

• Calibration slope \hat{b} . The calibration slope \hat{b} is obtained by regressing the test data outcomes y_{test} on the models' predicted logits $\text{logit}(\hat{y})$, *i.e.*,

$$logit \mathbb{E}[Y \mid \hat{y}] = a + b \, logit(\hat{y}).$$

This measure will be used to assess calibration and deviations of \hat{b} from one indicate miscalibration (Steyerberg et al., 2019).

• Calibration in the large \hat{a} . We inspect calibration in the large \hat{a} on independently generated test data, from the model

$$logit \mathbb{E}[Y \mid \hat{y}] = a + logit(\hat{y}).$$

This measure will also be used to assess calibration and deviations of \hat{a} from zero indicate miscalibration (Steyerberg et al., 2019).

To facilitate comparison between simulation conditions, all estimands will also be corrected by the oracle version of the estimand, e.g., the Brier score will be computed from the ground truth parameters and the simulated data x, subsequently the oracle Brier score will be subtracted from the estimated Brier score.

A.4 Methods

A.4.1 AINET

We now present the mock-method and give a superficial motivation why it could lead to improved predictive performance: Choosing the vector of penalization weights in the adaptive LASSO becomes difficult in high-dimensional settings. For instance, using absolute LASSO estimates as penalization weights omits the importance of several predictors by not selecting them, especially in the case of highly correlated predictors (Algamal and Lee, 2015). The adaptive importance elastic net (AINET) circumvents this problem by employing a random forest to estimate the penalization weights via an *a priori* chosen variable importance measure. In this way, the importance of all variables enter the penalization weights simultaneously.

The penalized log-likelihood for AINET for a single observation (y, x) is defined as

$$\ell_{\text{AINET}}(\beta_0, \boldsymbol{\beta}; y, \boldsymbol{x}, \alpha, \lambda, \boldsymbol{w}) = \ell(\beta_0, \boldsymbol{\beta}; y, \boldsymbol{x}) + \lambda \left(\alpha \sum_{j=1}^p w_j |\beta_j| + \frac{1}{2} (1 - \alpha) \sum_{j=1}^p w_j \beta_j^2 \right)$$

where

$$\ell(\beta_0, \boldsymbol{\beta}; y, \boldsymbol{x}) = y \log \left(\exp it \left\{ \beta_0 + \boldsymbol{x}^{\top} \boldsymbol{\beta} \right\} \right) + (1 - y) \log \left(1 - \exp it \left\{ \beta_0 + \boldsymbol{x}^{\top} \boldsymbol{\beta} \right\} \right)$$

denotes the log-likelihood of a binomial GLM and w is derived from a random forest variable importance

measure IMP as

$$w_j = 1 - \left(\frac{\mathrm{IMP}_j}{\sum_{k=1}^p \mathrm{IMP}_k}\right)^{\gamma},$$

where we transform IMP to be non-negative via

$$IMP_j = \max\{0, \widetilde{IMP_j}\}$$

and γ is a hyperparameter for the influence of the weights similar to γ hyperparameter of the adaptive elastic net. AINET is fitted by maximizing its penalized log-likelihood assuming i.i.d. observations $\{(y_i, x_i)\}_{i=1}^n$, i.e.,

$$\arg \max_{\beta_0, \beta} \sum_{i=1}^n \ell_{AINET}(\beta_0, \beta; y_i, x_i, \alpha, \lambda, w).$$

Per default, we choose mean decrease in the Gini coefficient for $\widehat{\text{IMP}}$. Hyperparameters of the random forest are not tuned, but kept at their default values (e.g., mtry, ntree). The hyperparameter $\gamma = 1$ will stay constant for all simulations.

AINET is supposed to seem like a reasonable method at first glance. However, AINET cannot be expected to share desirable theoretical properties with the usual adaptive LASSO, such as oracle estimation (Zou, 2006). This is because the penalization weights w do not meet the required consistency assumption. Also in terms of prediction performance, AINET is not expected to outperform methods of comparable complexity.

A.4.2 Benchmark methods

- **Binary logistic regression** (McCullagh and Nelder, 2019) with and without ridge penalty for highand low-dimensional settings, respectively. In case a ridge penalty is needed, it is tuned via 5-fold cross-validation by following the "one standard error" rule as implemented in **glmnet** (Friedman et al., 2010).
- Elastic net (Zou and Hastie, 2005), for which the penalized log-likelihood is given by

$$\ell_{\mathrm{EN}}(\beta_0, \boldsymbol{\beta}; y, \boldsymbol{x}, \boldsymbol{\alpha}, \lambda) = \ell(\beta_0, \boldsymbol{\beta}; y, \boldsymbol{x}) + \lambda \left(\boldsymbol{\alpha} \| \boldsymbol{\beta} \|_1 + \frac{1}{2} (1 - \boldsymbol{\alpha}) \| \boldsymbol{\beta} \|_2^2 \right).$$

Here, α and λ are tuned via 5-fold cross-validation by following the "one standard error" rule.

• Adaptive elastic net (Zou, 2006), with penalized loss function

$$\ell_{\text{adaptive}}(\beta_0, \boldsymbol{\beta}; y, \boldsymbol{x}, \boldsymbol{\alpha}, \lambda, \boldsymbol{w}) = \ell(\beta_0, \boldsymbol{\beta}; y, \boldsymbol{x}) + \lambda \left(\alpha \sum_{j=1}^p w_j |\beta_j| + \frac{1}{2} (1 - \alpha) \sum_{j=1}^p w_j \beta_j^2 \right).$$

Here, the penalty weights w are inverse coefficient estimates from a binary logistic regression

$$\hat{w}_i = |\hat{\vartheta}_i|^{-\gamma},$$

where λ and α are tuned via 5-fold cross-validation by following the "one standard error" rule. The hyperparameter $\gamma=1$ will stay constant for all simulations. In case p>n, we estimate the penalty weights using a ridge penalty, tuned via an additional nested 5-fold cross-validation by following the "one standard error" rule.

• **Random forests** (Breiman, 2001) for binary outcomes without hyperparameter tuning. The default parameters of **ranger** will be used (Wright and Ziegler, 2017).

A.5 Performance measures

The distribution of all estimands from Section A.3 will be assessed visually with box- and violin-plots that are stratified by method and simulation conditions. We will also compute mean, median, standard deviation, interquartile range, and 95% confidence intervals for each of the estimands. Moreover, instead of "eye-balling" differences in predictive performance across methods and conditions, we will formally assess them by regressing the estimands on the method and simulation conditions (cf. Skrondal, 2000). To do so, we will use a fully interacted model with the interaction between the methods and the 128 simulations conditions, *i.e.*, in R notation: estimand ~ 0 + method:scenario. We will rank pairwise comparison between two methods within a single condition by their p-values, to more easily identify conditions where methods show differences in predictive performance. The choice of a significance level at which a method is deemed superior will be determined based on preliminary simulations. We set this level to 5%, where p-values will be adjusted using the single-step method (Hothorn et al., 2008) within a single simulation condition for comparisons between AINET and any other method.

A.6 Determining the number of simulations

We determine the number of simulation B such that the Monte Carlo standard error of the primary estimand, the mean Brier score \overline{BS}/B , is sufficiently small. The variance of \overline{BS}/B is given by

$$\operatorname{Var}\left(\overline{\mathrm{BS}}/B\right) = \frac{\operatorname{Var}\left\{(y - \hat{y})^{2}\right\}}{B \cdot n_{\mathrm{test}}}$$

and $\operatorname{Var}\left\{(y_{ib}-\hat{y}_{ib})^2\right\}$ could be decomposed further (Bradley et al., 2008). However, the resulting expression is difficult to evaluate for our data-generating process as it depends on several of the simulation parameters. We therefore follow a similar approach as in Morris et al. (2019) and estimate $\widehat{\operatorname{Var}}\left\{(y_{ib}-\hat{y}_{ib})^2\right\} < V$ from an initial small simulation run with 100 simulations per condition to get an upper bound V for worst-case variance across all simulation conditions. Therefore, the number of simulations is then given by

$$B = \frac{V}{n_{\text{test Var}}(\overline{\text{BS}})}.$$

Since $\overline{BS} \in [0,1]$ we decide that we require the Monte Carlo standard error of \overline{BS} to be lower than four significant digits, 0.0001.

The initial simulation run led to an estimated worst case variance of $\widehat{V}=0.2$. Therefore, we compute that

$$B = 0.2/(10000 \times 0.0001^2) = 2000$$

replications are required to obtain Brier score estimates with the desired precision.

A.7 Handling exceptions

It is inevitable that convergence issues and other problems will arise in the simulation study. We will handle them as follows:

- If a method fails to converge, the simulation will be excluded from the analysis. The failing simulations will not be replaced with new simulations that successfully converge as convergence may be impossible for some scenarios.
- We will report the proportion of simulations with convergence issues for each method and discuss the potential reasons for their emergence.
- In case of severe convergence issues or other problems (more than 10% of the simulations failing within a setting), we may adjust the simulation parameters post hoc. This will be indicated in the discussion of the results.
- Convergence may be possible for certain tuning parameters of a method (e.g., cross-validation of LASSO may fail for some values λ while it could work for others). In this case we will choose a parameter value where the method still converges, as one would usually do with a real data set.

Appendix B Per-protocol results

Here, we describe the outcomes of the preregistered simulations. Overall, the performance of AINET was virtually identical to elastic net regression. The adaptive penalization weights of AINET do not seem to make a difference for the data generating mechanism considered in our simulations. Moreover, since the data were generated under a process equivalent to a logistic regression model, it is no surprise that for reasonably large sample sizes, logistic regression also performed the best. The only exception are conditions with small sample size and low number of events per variable. Here, AINET and elastic net led to more stable and better calibrated predictions than logistic regression. The random forest was outperformed by AINET in most simulation conditions, with exception of very small sample size and prevalence, as well as when a high correlation between covariates was present. Finally, the performance of the adaptive elastic net was generally worse compared to AINET and elastic net. In the following, we summarize the results for each estimand.

B.1 Brier score (primary estimand)

Figure 2 shows the differences in mean Brier score between AINET and the other methods stratified by simulation conditions. We see that there is hardly any difference between AINET and the elastic net (EN) across all simulation conditions meaning that predictive performance of both methods seems to be very similar in the investigated scenarios.

The random forest (RF) shows better predictive performance than AINET in conditions with very low sample size (n=100) and prevalence (prev = 0.01). For increasing sample size and prevalence, the performance of AINET seems to become more similar or improve over RF when the correlation of the covariates is not too large ($\rho \leq 0.6$) especially for low events per variable (EPV ≤ 1). For highly correlated covariates ($\rho = 0.95$), the performance of AINET is similar or worse across most simulation conditions.

Logistic regression (GLM) showed better predictive performance compared to AINET in most simulation conditions. An exception are the conditions with small sample size (n=100), medium to large prevalence (prev ≥ 0.05) and low events per variable (EPV ≤ 1), where AINET performed better than GLM.

The adaptive elastic net (AEN) method performed worse than AINET in almost all simulation conditions. Only in conditions with very large sample size (n = 5000), very small prevalence (prev = 0.01), and high events per variable (EPV = 20), AEN showed predictive performance on par with AINET.

B.2 Scaled Brier score (secondary estimand)

Figure 3 shows the differences in scaled Brier score between AINET and the other methods stratified by simulation conditions. The scaled Brier score is useful to compare the actual values of Brier scores across conditions with different prevalence, but not so much to compare Brier scores of different methods within a simulation condition with fixed prevalence.

We see that for most conditions the plots look like a flipped version of the original Brier scores from Figure 2. Therefore, conclusions are mostly the same. For very small sample sizes coupled with low prevalence and low events per variable (the topleft plots), the scaled Brier score indicates superiority of AINET over RF and GLM, which is opposite the conclusion based on the raw Brier score. We advise to interpret these conditions cautiously since the prevalence prediction which is used for scaling is based on the much larger test data set.

B.3 Log-score (secondary estimand)

Figure 4 shows the differences in log-score between AINET and the other methods stratified by simulation conditions. We see that in certain conditions, the error bars of certain methods are much larger. This is due to the log-score's sensitivity to extreme predictions, which often happen under the RF (and sometimes under the GLM). Despite the larger variability of the log-score, conclusion regarding the comparison between AINET and the other methods are largely the same as under the Brier score.

B.4 Area under the curve (secondary estimand)

Figure 4 shows the differences in area under the curve (AUC) between AINET and the other methods stratified by simulation conditions. As with the other estimands, AINET shows virtually identical performance as EN regression across all simulation conditions. AINET seems to outperform RF across most simulation conditions, with the exception of a conditions with low sample size (n=100), medium prevalence (prev = 0.05), and low events per variable (EPV \leq 1). GLM, typically outperforms AINET conditions with small to medium sample size ($n\leq500$), and also in conditions with larger sample size when the events per variable is normal to high (EPV ≥10) and the prevalence is small (prev = 0.01). Finally, the AEN is worse with respect to AUC than AINET across all simulation conditions.

B.5 Calibration slope (secondary estimand)

Figure 5 shows boxplots of calibration slopes stratified by simulation condition and method. For each condition the percentage of simulations where no estimate could be obtained is indicated. This usually happened because of extreme (close to zero or one) predictions, or non-convergence of the method itself. We caution against interpretation of the random forest (RF) calibration slopes because this method often resulted in predicted probabilities of zero or one, so that a calibration slope could not be fitted.

We see that logistic regression (GLM) shows on average optimal calibration slopes in most simulation condition. In cases where it is off one, its calibration slopes are usually too small indicating overoptimistic predictions. In general, worse calibration slopes are obtained for lower event per variable (EPV).

The penalized methods (AINET, EN, AEN) show a more stable behavior, and on average larger calibration slopes than GLM. This is likely confounded by the simulation conditions in which no GLM calibration slope can be estimated, but estimation of the penalized methods' calibration slope is still possible. Among the penalized method's AINET and EN shows relatively similar calibration slopes whereas the AEN shows worse calibration slopes that are more off the value of one.

B.6 Calibration in the large (secondary estimand)

Figure 6 shows boxplots of calibration in the large estimates stratified by simulation condition and method. For each condition also the percentage of simulations where no estimate could be obtained is indicated. This usually happened because of extreme (close to zero or one) predictions.

We see that the number of simulations with non-estimable calibration is substantially larger when the sample size is small, whereas it decreases for larger sample sizes. An exception is the RF where the number of non-estimable calibrations stays high across most conditions.

While all methods seem to be marginally well calibrated, the penalized methods (AINET, EN, and AEN) show lower numbers of simulations with non-estimable calibration compared to GLM, especially for low to medium sample sizes and low events per variables.

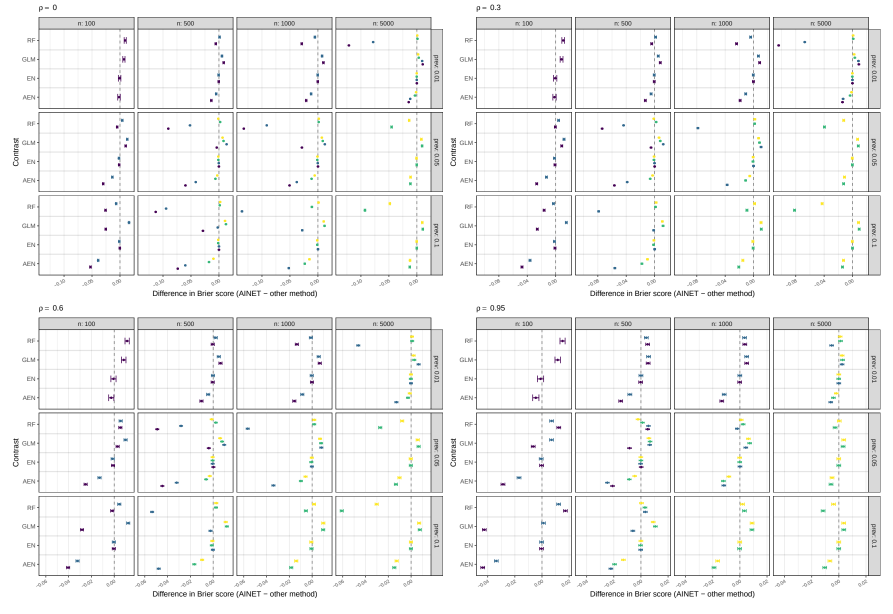


Figure 2: Tie-fighter plot for the difference in Brier score between any method on the *y*-axis and AINET. The 95% confidence intervals are adjusted per simulation condition using the single-step method. Lower values indicate better performance of AINET.

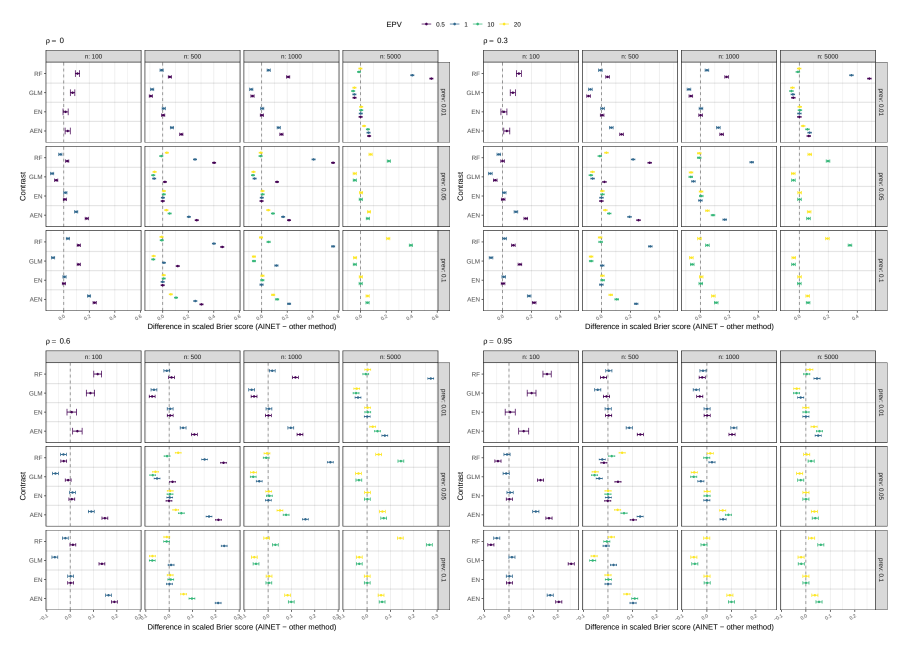


Figure 3: Tie-fighter plot for the difference in scaled Brier score between any method on the *y*-axis and AINET. The 95% confidence intervals are adjusted per simulation condition using the single-step method. Larger values indicate better performance of AINET.

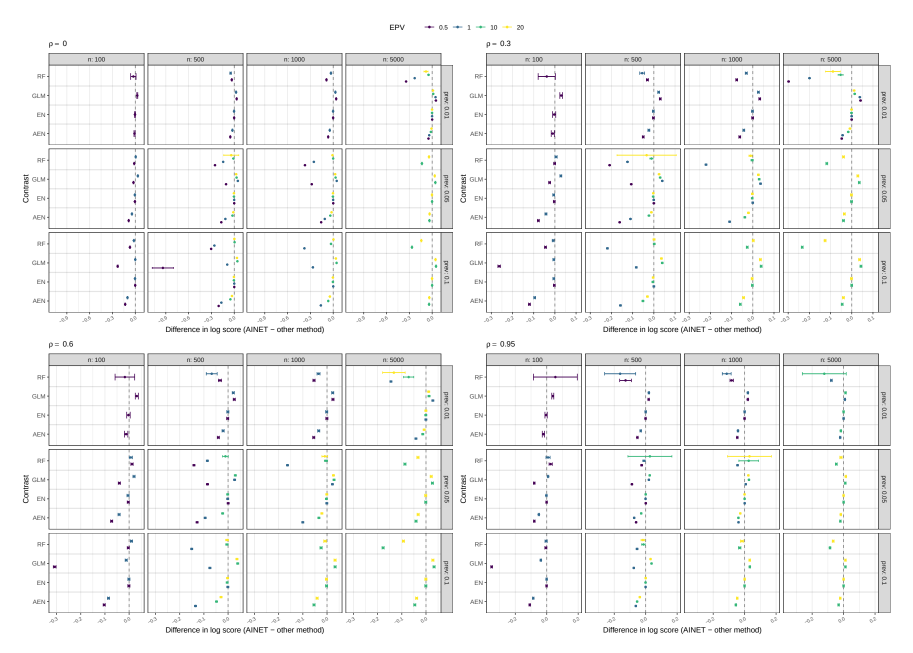


Figure 4: Tie-fighter plot for the difference in log-score between any method on the *y*-axis and AINET. The 95% confidence intervals are adjusted per simulation condition using the single-step method. Lower values indicate better performance of AINET.

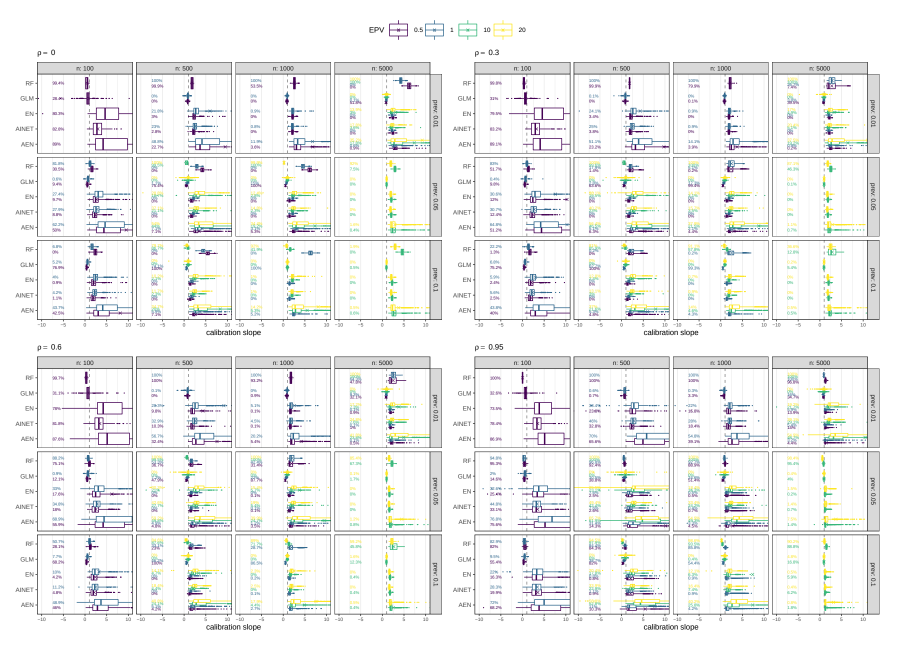


Figure 5: Boxplots of calibration slopes stratified by method and simulation conditions. Mean calibration slope is indicated by a cross. A value of one indicates optimal calibration. Percentage of simulations where calibration slope could not be estimated (due to extreme predictions or complete separation) are also indicated.

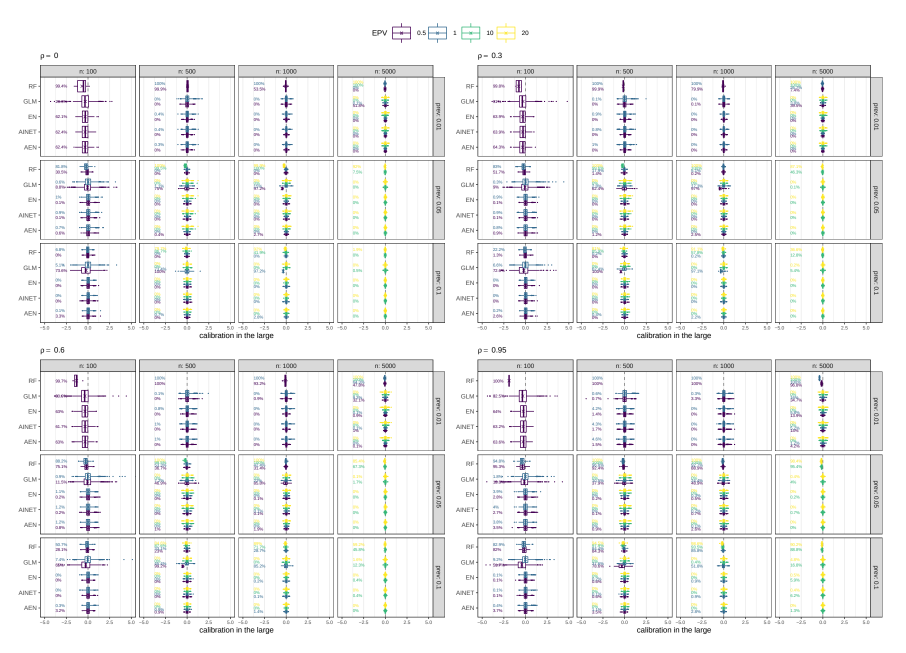


Figure 6: Boxplots of calibration in the large stratified by method and simulation conditions. Mean calibration in the large is indicated by a cross. A value of zero indicates optimal calibration in the large. Percentage of simulations where calibration in the large could not be estimated (due to extreme predictions or complete separation) are also indicated.

References

- Algamal, Z. Y. and Lee, M. H. (2015). Penalized logistic regression with the adaptive LASSO for gene selection in high-dimensional cancer classification. *Expert Systems with Applications*, 42(23):9326–9332. doi:10.1016/J.ESWA.2015.08.016.
- Altman, D. G., Moher, D., and Schulz, K. F. (2017). Harms of outcome switching in reports of randomised trials: CONSORT perspective. *BMJ*, page j396. doi:10.1136/bmj.j396.
- Angelis, C. D., Drazen, J. M., Frizelle, F. A., Haug, C., Hoey, J., Horton, R., Kotzin, S., Laine, C., Marusic, A., Overbeke, A. J. P., Schroeder, T. V., Sox, H. C., and Weyden, M. B. V. D. (2004).
 Clinical trial registration: A statement from the international committee of medical journal editors.
 New England Journal of Medicine, 351(12):1250–1251. doi:10.1056/nejme048225.
- Boulesteix, A.-L., Binder, H., Abrahamowicz, M., and Sauerbrei, W. (2017). On the necessity and design of studies comparing statistical methods. *Biometrical Journal*, 60(1):216–218. doi:10.1002/bimj.201700129.
- Boulesteix, A.-L., Groenwold, R. H., Abrahamowicz, M., Binder, H., Briel, M., Hornung, R., Morris, T. P., Rahnenführer, J., and Sauerbrei, W. (2020). Introduction to statistical simulations in health research. *BMJ Open*, 10(12):e039921. doi:10.1136/bmjopen-2020-039921.
- Boulesteix, A.-L., Lauer, S., and Eugster, M. J. A. (2013). A plea for neutral comparison studies in computational sciences. *PLoS ONE*, 8(4):e61562. doi:10.1371/journal.pone.0061562.
- Boulesteix, A.-L., Stierle, V., and Hapfelmeier, A. (2015). Publication bias in methodological computational research. *Cancer Informatics*, 14s5:CIN.S30747. doi:10.4137/cin.s30747.
- Bradley, A. A., Schwartz, S. S., and Hashino, T. (2008). Sampling uncertainty and confidence intervals for the Brier score and Brier skill score. *Weather and Forecasting*, 23(5):992–1006. doi:10.1175/2007waf2007049.1.
- Breiman, L. (2001). Random Forests. *Machine Learning*, 45(1):5–32. doi:10.1023/a:1010933404324.
- Burton, A., Altman, D. G., Royston, P., and Holder, R. L. (2006). The design of simulation studies in medical statistics. *Statistics in Medicine*, 25(24):4279–4292. doi:10.1002/sim.2673.
- Chalmers, R. P. and Adkins, M. C. (2020). Writing effective and reliable Monte Carlo simulations with the SimDesign package. *The Quantitative Methods for Psychology*, 16(4):248–280. doi:10.20982/tqmp.16.4.p248.
- Damen, J. A. A. G., Hooft, L., Schuit, E., Debray, T. P. A., Collins, G. S., Tzoulaki, I., Lassale, C. M., Siontis, G. C. M., Chiocchia, V., Roberts, C., Schlüssel, M. M., Gerry, S., Black, J. A., Heus, P., van der Schouw, Y. T., Peelen, L. M., and Moons, K. G. M. (2016). Prediction models for cardiovascular disease risk in the general population: systematic review. *BMJ*, page i2416. doi:10.1136/bmj.i2416.
- Dutilh, G., Sarafoglou, A., and Wagenmakers, E.-J. (2019). Flexible yet fair: blinding analyses in experimental psychology. *Synthese*. doi:10.1007/s11229-019-02456-7.
- Elofsson, A., Hess, B., Lindahl, E., Onufriev, A., van der Spoel, D., and Wallqvist, A. (2019). Ten simple rules on how to create open access and reproducible molecular simulations of biological systems. *PLOS Computational Biology*, 15(1):e1006649. doi:10.1371/journal.pcbi.1006649.

- Feynman, R. P. (1974). Cargo cult science. Engineering & Science, 37:10-13.
- Friedman, J., Hastie, T., and Tibshirani, R. (2010). Regularization Paths for Generalized Linear Models via Coordinate Descent. *Journal of Statistical Software*, 33(1):1–22. doi:10.18637/jss.v033.i01.
- Gasparini, A. (2018). rsimsum: Summarise results from monte carlo simulation studies. *Journal of Open Source Software*, 3(26):739. doi:10.21105/joss.00739.
- Gasparini, A., Morris, T. P., and Crowther, M. J. (2021). INTEREST: INteractive tool for exploring REsults from simulation sTudies. *Journal of Data Science, Statistics, and Visualisation*, 1(4). doi:10.52933/jdssv.v1i4.9.
- Gelman, A. and Carlin, J. (2014). Beyond power calculations: Assessing type S (sign) and type M (magnitude) errors. *Perspectives on Psychological Science*, 9(6):641–651. doi:10.1177%2F1745691614551642.
- Gelman, A. and Tuerlinckx, F. (2000). Type S error rates for classical and Bayesian single and multiple comparison procedures. *Computational Statistics*, 15(3):373–390. doi:10.1007/s001800000040.
- Gneiting, T. (2008). Editorial: Probabilistic forecasting. *Journal of the Royal Statistical Society: Series A (Statistics in Society)*, 171(2):319–321. doi:10.1111/j.1467-985x.2007.00522.x.
- Gneiting, T. and Raftery, A. E. (2007). Strictly proper scoring rules, prediction, and estimation. *Journal of the American Statistical Association*, 102(477):359–378. doi:10.1198/016214506000001437.
- Hoaglin, D. C. and Andrews, D. F. (1975). The reporting of computation-based results in statistics. *The American Statistician*, 29(3):122–126. doi:10.1080/00031305.1975.10477393.
- Hoffmann, S., Schönbrodt, F., Elsas, R., Wilson, R., Strasser, U., and Boulesteix, A.-L. (2021). The multiplicity of analysis strategies jeopardizes replicability: lessons learned across disciplines. *Royal Society Open Science*, 8(4). doi:10.1098/rsos.201925.
- Holford, N. H. G., Kimko, H. C., Monteleone, J. P. R., and Peck, C. C. (2000). Simulation of clinical trials. *Annual Review of Pharmacology and Toxicology*, 40(1):209–234. doi:10.1146/annurev.pharmtox.40.1.209.
- Hothorn, T., Bretz, F., and Westfall, P. (2008). Simultaneous inference in general parametric models. *Biometrical Journal*, 50(3):346–363. doi:10.1002/bimj.200810425.
- Jelizarow, M., Guillemot, V., Tenenhaus, A., Strimmer, K., and Boulesteix, A.-L. (2010). Over-optimism in bioinformatics: an illustration. *Bioinformatics*, 26(16):1990–1998. doi:10.1093/bioinformatics/btq323.
- Kidwell, M. C., Lazarević, L. B., Baranski, E., Hardwicke, T. E., Piechowski, S., Falkenberg, L.-S., Kennett, C., Slowik, A., Sonnleitner, C., Hess-Holden, C., Errington, T. M., Fiedler, S., and Nosek, B. A. (2016). Badges to acknowledge open practices: A simple, low-cost, effective method for increasing transparency. *PLOS Biology*, 14(5):e1002456. doi:10.1371/journal.pbio.1002456.
- Klein, J. R. and Roodman, A. (2005). Blind analysis in nuclear and particle physics. *Annual Review of Nuclear and Particle Science*, 55(1):141–163. doi:10.1146/annurev.nucl.55.090704.151521.

- Kreutz, C., Can, N. S., Bruening, R. S., Meyberg, R., Mérai, Z., Fernandez-Pozo, N., and Rensing, S. A. (2020). A blind and independent benchmark study for detecting differentially methylated regions in plants. *Bioinformatics*, 36(11):3314–3321. doi:10.1093/bioinformatics/btaa191.
- Kreuzberger, N., Damen, J., Trivella, M., Estcourt, L. J., Aldin, A., Umlauff, L., Vazquez-Montes, M., Wolff, R., Moons, K., Monsef, I., Foroutan, F., Kreuzer, K., and Skoetz, N. (2020). Prognostic models for newly-diagnosed chronic lymphocytic leukaemia in adults: a systematic review and meta-analysis. *Cochrane Database of Systematic Reviews*, 7:CD012022. doi:10.1002/14651858.CD012022.pub2.
- Lawlor, D. A. (2007). Quality in epidemiological research: should we be submitting papers before we have the results and submitting more hypothesis-generating research? *International Journal of Epidemiology*, 36(5):940–943. doi:10.1093/ije/dym168. URL https://doi.org/10.1093/ije/dym168.
- Loder, E., Groves, T., and MacAuley, D. (2010). Registration of observational studies. *BMJ*, 340(feb18 2):c950–c950. doi:10.1136/bmj.c950.
- McCullagh, P. and Nelder, J. A. (2019). Generalized Linear Models. Routledge.
- Monks, T., Currie, C. S. M., Onggo, B. S., Robinson, S., Kunc, M., and Taylor, S. J. E. (2018). Strengthening the reporting of empirical simulation studies: Introducing the STRESS guidelines. *Journal of Simulation*, 13(1):55–67. doi:10.1080/17477778.2018.1442155.
- Morris, T. P., White, I. R., and Crowther, M. J. (2019). Using simulation studies to evaluate statistical methods. *Statistics in Medicine*, 38(11):2074–2102. doi:10.1002/sim.8086.
- Nießl, C., Herrmann, M., Wiedemann, C., Casalicchio, G., and Boulesteix, A.-L. (2021). Over-optimism in benchmark studies and the multiplicity of design and analysis options when interpreting their results. *WIREs Data Mining and Knowledge Discovery*. doi:10.1002/widm.1441.
- Nosek, B. A., Ebersole, C. R., DeHaven, A. C., and Mellor, D. T. (2018). The preregistration revolution. *Proceedings of the National Academy of Sciences*, 115(11):2600–2606. doi:10.1073/pnas.1708274114.
- O'Kelly, M., Anisimov, V., Campbell, C., and Hamilton, S. (2016). Proposed best practice for projects that involve modelling and simulation. *Pharmaceutical Statistics*, 16(2):107–113. doi:10.1002/pst.1789.
- R Core Team (2020). *R: A Language and Environment for Statistical Computing*. R Foundation for Statistical Computing, Vienna, Austria. URL https://www.R-project.org/.
- Riley, R. D., Snell, K. I., Ensor, J., Burke, D. L., Jr, F. E. H., Moons, K. G., and Collins, G. S. (2018). Minimum sample size for developing a multivariable prediction model: PART II binary and time-to-event outcomes. *Statistics in Medicine*, 38(7):1276–1296. doi:10.1002/sim.7992.
- Robin, X., Turck, N., Hainard, A., Tiberti, N., Lisacek, F., Sanchez, J.-C., and Müller, M. (2011). pROC: an open-source package for R and S+ to analyze and compare ROC curves. *BMC Bioinformatics*, 12:77. doi:10.1186/1471-2105-12-77.

- Schmid, C. H. and Griffith, J. L. (2005). Multivariate classification rules: Calibration and discrimination. In Armitage, P. and Colton, T., editors, *Encyclopedia of Biostatistics*, volume 5, pages 3491–3497. Wiley, 2nd edition.
- Schwab, S. and Held, L. (2021). Statistical programming: Small mistakes, big impacts. *Significance*, 18(3):6–7. doi:10.1111/1740-9713.01522.
- Seker, B. O., Reeve, K., Havla, J., Burns, J., Gosteli, M., Lutterotti, A., Schippling, S., Mansmann, U., and Held, U. (2020). Prognostic models for predicting clinical disease progression, worsening and activity in people with multiple sclerosis. *Cochrane Database of Systematic Reviews*, (5). doi:10.1002/14651858.CD013606.
- Simmons, J. P., Nelson, L. D., and Simonsohn, U. (2011). False-positive psychology: Undisclosed flexibility in data collection and analysis allows presenting anything as significant. *Psychological Science*, 22(11):1359–1366. doi:10.1177/0956797611417632.
- Simon, N., Friedman, J., Hastie, T., and Tibshirani, R. (2011). Regularization Paths for Cox's Proportional Hazards Model via Coordinate Descent. *Journal of Statistical Software*, 39(5):1–13. doi:10.18637/jss.v039.i05.
- Skrondal, A. (2000). Design and analysis of monte carlo experiments: Attacking the conventional wisdom. *Multivariate Behavioral Research*, 35(2):137–167. doi:10.1207/s15327906mbr3502_1.
- Smith, M. K. and Marshall, A. (2010). Importance of protocols for simulation studies in clinical drug development. *Statistical Methods in Medical Research*, 20(6):613–622. doi:10.1177/0962280210378949.
- Steverberg, E. W. et al. (2019). Clinical Prediction Models. Springer.
- Tukey, J. W. (1980). We need both exploratory and confirmatory. *The American Statistician*, 34(1):23–25. doi:10.1080/00031305.1980.10482706.
- Ullmann, T., Beer, A., Hünemörder, M., Seidl, T., and Boulesteix, A.-L. (2022). Over-optimistic evaluation and reporting of novel cluster algorithms: an illustrative study. *Advances in Data Analysis and Classification*. doi:10.1007/s11634-022-00496-5.
- Van der Bles, A. M., Van Der Linden, S., Freeman, A. L., Mitchell, J., Galvao, A. B., Zaval, L., and Spiegelhalter, D. J. (2019). Communicating uncertainty about facts, numbers and science. *Royal Society Open Science*, 6(5):181870. doi:10.1098/rsos.181870.
- van Smeden, M., de Groot, J. A. H., Moons, K. G. M., Collins, G. S., Altman, D. G., Eijkemans, M. J. C., and Reitsma, J. B. (2016). No rationale for 1 variable per 10 events criterion for binary logistic regression analysis. *BMC Medical Research Methodology*, 16(1). doi:10.1186/s12874-016-0267-3.
- van Smeden, M., Moons, K. G., de Groot, J. A., Collins, G. S., Altman, D. G., Eijkemans, M. J., and Reitsma, J. B. (2018). Sample size for binary logistic prediction models: Beyond events per variable criteria. *Statistical Methods in Medical Research*, 28(8):2455–2474. doi:10.1177/0962280218784726.
- van Zwet, E. W. and Cator, E. A. (2021). The significance filter, the winner's curse and the need to shrink. *Statistica Neerlandica*, 75(4):437–452. doi:10.1111/stan.12241.

- Vidaurre, D., Bielza, C., and Larrañaga, P. (2013). A survey of L1 regression. *International Statistical Review*, 81(3):361–387. doi:10.1111/insr.12023.
- White, I. R. (2010). Simsum: Analyses of simulation studies including monte carlo error. *The Stata Journal: Promoting communications on statistics and Stata*, 10(3):369–385. doi:10.1177/1536867x1001000305.
- Wicherts, J. M., Veldkamp, C. L. S., Augusteijn, H. E. M., Bakker, M., van Aert, R. C. M., and van Assen, M. A. L. M. (2016). Degrees of freedom in planning, running, analyzing, and reporting psychological studies: A checklist to avoid *p*-hacking. *Frontiers in Psychology*, 7. doi:10.3389/fpsyg.2016.01832.
- Wright, M. N. and Ziegler, A. (2017). ranger: A fast implementation of random forests for high dimensional data in C++ and R. *Journal of Statistical Software*, 77(1):1–17. doi:10.18637/jss.v077.i01.
- Wynants, L., Calster, B. V., Collins, G. S., Riley, R. D., Heinze, G., Schuit, E., Bonten, M. M. J., Dahly, D. L., Damen, J. A., Debray, T. P. A., de Jong, V. M. T., Vos, M. D., Dhiman, P., Haller, M. C., Harhay, M. O., Henckaerts, L., Heus, P., Kammer, M., et al. (2020). Prediction models for diagnosis and prognosis of covid-19: systematic review and critical appraisal. *BMJ*, 369:m1328. doi:10.1136/bmj.m1328.
- Zou, H. (2006). The adaptive lasso and its oracle properties. *Journal of the American Statistical Association*, 101(476):1418–1429. doi:10.1198/016214506000000735.
- Zou, H. and Hastie, T. (2005). Regularization and variable selection via the elastic net. *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*, 67(2):301–320. doi:10.1111/j.1467-9868.2005.00503.x.