

A simulation approach to calculating minimum sample sizes for prediction modelling

The `pmsims` package for R

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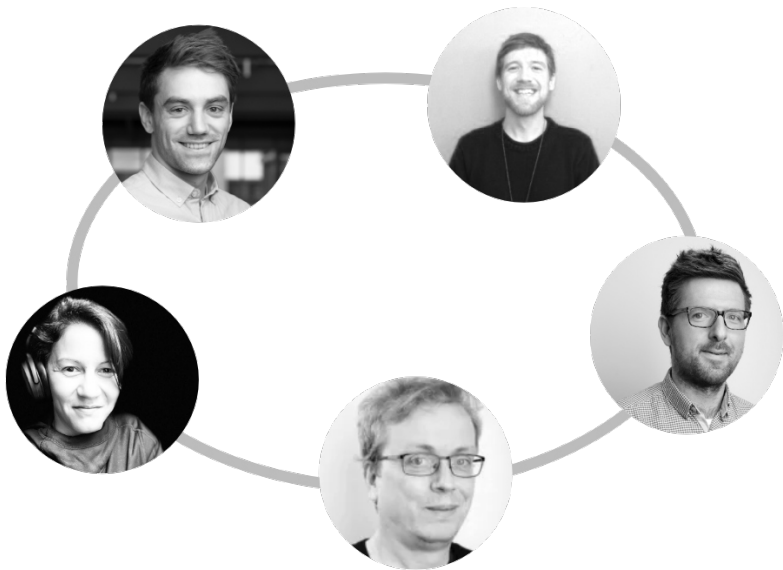
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30-second version

1. Prediction models developed with inadequate samples lead to **overfitting** and **imprecise** estimates.
2. Existing tools use **analytical methods** to derive minimum samples sizes for continuous, binary, and survival outcomes.
3. We've developed a **simulation-based** approach that can be applied to **any** outcome or method.



Overview

1. Background
 - What is the problem?
 - What solutions already exist?
2. Our approach
 - Simulation to identify minimum sample size that satisfies criteria.
 - Flexible, but slower.
 - Gaussian process regression (via `mlpwr`) to speed up.
3. Next steps

What's the problem?

Thousands of prediction models are developed each year. Most are developed with inadequate samples.

- Insufficient sample size was the most common cause of high risk of bias in 731 models for COVID-19.¹ 67% of models were developed on too few patients.
- Sample sizes were inadequate in 56% of models developed using supervised machine learning² and 73% of models in psychiatry.⁴
- 8% of machine learning models published in oncology report a sample size justification.³

Inadequate samples → research waste

- Inadequate samples lead to overfitting and inaccurate estimates of model parameters.
 - Overfitting is where the model captures idiosyncrasies of the development sample, producing inflated estimates of predictive performance that cannot be replicated in the target population.
- Unreliable models may generate inappropriate decisions about patient care or lead to models not being implemented into clinical practice.
- Data collection can be invasive and inconvenient and diverts resources from other activities that benefit patients.

Ensuring sample sizes are sufficient *before model development* would improve patient outcomes by avoiding models developed with inadequate samples and reducing participant burden.

Tools for estimating minimum sample sizes for prediction

Until recently, most studies ignored sample size.

Or they used simple rules-of-thumb (e.g., 10 events per variable).

In 2018, `pmsampsize` was released by Riley et al.

RESEARCH ARTICLE

WILEY Statistics
in Medicine

Minimum sample size for developing a multivariable prediction model: Part I – Continuous outcomes

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In the medical literature, hundreds of prediction models are being developed to predict health outcomes in individuals. For continuous outcomes, typically a linear regression model is developed to predict an individual's outcome value conditional on values of multiple predictors (covariates). To improve model development and reduce the potential for overfitting, a suitable sample size is required in terms of the number of subjects (n) relative to the number of predictor parameters (p) for potential inclusion. We propose that the minimum value of n should meet the following four key criteria: (i) small optimism in predictor effect estimates as defined by a global shrinkage factor of ≥ 0.9 ; (ii) small absolute difference of ≤ 0.05 in the apparent and adjusted R^2 ; (iii) precise estimation (a margin of error $\leq 10\%$ of the true value) of the model's residual standard deviation; and similarly, (iv) precise estimation of the mean predicted outcome value (model intercept). The criteria require prespecification of the user's chosen p and the model's anticipated R^2 as informed by previous studies. The value of n that meets all four criteria provides the minimum sample size required for model development. In an applied example, a new model to predict lung function in African-American women using 25 predictor parameters requires at least 918 subjects to meet all criteria, corresponding to at least 36.7 subjects per predictor parameter. Even larger sample sizes may be needed to additionally ensure precise estimates of key predictor effects, especially when important categorical predictors have low prevalence in certain categories.

KEYWORDS

continuous outcome, linear regression, minimum sample size, multivariable prediction model, R-squared

RESEARCH ARTICLE

WILEY Statistics
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Minimum sample size for developing a multivariable prediction model: PART II – binary and time-to-event outcomes

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When designing a study to develop a new prediction model with binary or time-to-event outcomes, researchers should ensure their sample size is adequate in terms of the number of participants (n) and outcome events (E) relative to the number of predictor parameters (p) considered for inclusion. We propose that the minimum values of n and E (and subsequently the minimum number of events per predictor parameter, EPP) should be calculated to meet the following three criteria: (i) small optimism in predictor effect estimates as defined by a global shrinkage factor of ≥ 0.9 ; (ii) small absolute difference of ≤ 0.05 in the model's apparent and adjusted Nagelkerke's R^2 ; and (iii) precise estimation of the overall risk in the population. Criteria (i) and (ii) aim to reduce overfitting conditional on a chosen p , and require prespecification of the model's anticipated Co-Snell R^2 , which we show can be obtained from previous studies. The values of n and E that meet all three criteria provides the minimum sample size required for model development. Upon application of our approach, a new diagnostic model for Chagas disease requires an EPP of at least 4.8 and a new prognostic model for recurrent venous thromboembolism requires an EPP of at least 23. This reinforces why rules of thumb (eg, 10 EPP) should be avoided. Researchers might additionally ensure the sample size gives precise estimates of key predictor effects; this is especially important when key categorical predictors have few events in some categories, as this may substantially increase the numbers required.

KEYWORDS

binary and time-to-event outcomes, logistic and Cox regression, multivariable prediction model, pseudo R-squared, sample size, shrinkage

The package identifies the minimum sample that results in:

Continuous	Binary
i. Small optimism in predictor effect estimates, indicated by a global shrinkage factor of 0.9.	
ii. Small absolute difference of 0.05 in the apparent and adjusted R^2	
iii. Precise estimation of the model's residual standard deviation.	Precise estimation of the overall risk in the population.
iv. Precise estimation of the model intercept.	

We  pmsampsize, however. . .

pmsampsize has methods for simple continuous, binary, and survival outcome. However, we increasingly need to derive minimum samples for:

Other types of model

e.g., machine learning algorithms such as random forests or gradient boosting.

Other data types

e.g., repeated measures and longitudinal data.

So, we've created a simulation-based framework for sample size estimation for prediction.

A simulation-based framework that derives the minimum sample that achieves:

- Within 10% of the expected large-sample performance;
- Calibration slope of >0.9

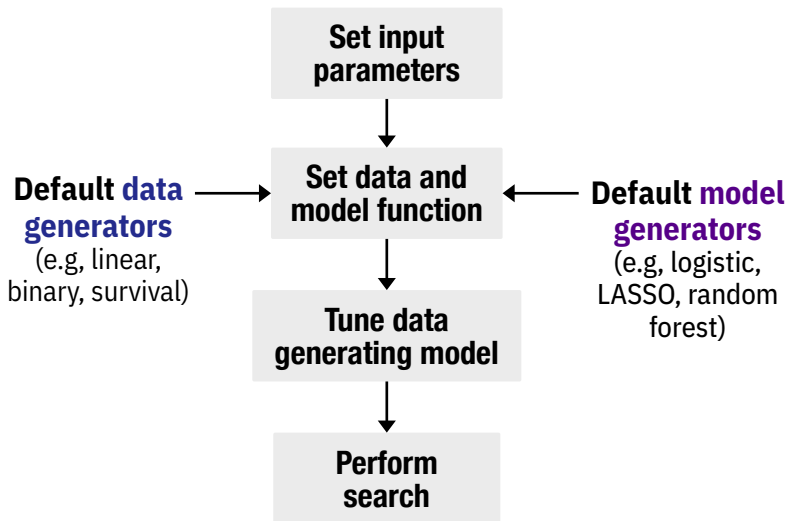
Key features:

- Can be used with any model or data type
- Provides defaults for common model and data types
- Efficient

An R package.

Maybe put slide here about conceptual differences vs. pmsampsize?

Our approach



Slide explaining input parameters

- `simulate_continuous`
- `simulate_binary`
- `simulate_survival`

Which each call:

- `simulate_custom`

Slide explaining default data and model generators

List default data/model/metrics.

Performing the search: `mlpwr`

A simulation-based approach with complex data or models would be too slow.

`mlpwr` is a R package by Felix Zimmer and Rudolf Debelak at the University of Zurich.

“A Power Analysis Toolbox to Find Cost-Efficient Study Designs”

Sample Size Planning for Complex Study Designs: A Tutorial for the `mlpwr` Package

Felix Zimmer, Mirka Henninger, and Rudolf Debelak
University of Zurich

A common challenge in designing empirical studies is determining an appropriate sample size. When more complex models are used, estimates of power can only be obtained using Monte Carlo simulations. In this tutorial, we introduce the R package `mlpwr` to perform simulation-based power analysis based on surrogate modeling. Surrogate modeling is a powerful tool to guide the search for study design parameters that imply a desired power or meet a cost threshold (e.g., in terms of monetary cost). `mlpwr` can be used to search for the optimal allocation when there are multiple design parameters, e.g., when balancing the number of participants and the number of groups in multilevel modeling. At the same time, the approach can take into account the cost of each design parameter, and aims to find a cost-efficient design. We introduce the basic functionality of the package, which can be applied to a wide range of statistical models and study designs. Additionally, we provide two examples based on empirical studies for illustration: one for sample size planning when using an item response theory model, and one for assigning the number of participants and the number of countries for a study using multilevel modeling.

Keywords: simulation, sample size, power analysis, machine learning

Introduction

Reliable testing of scientific hypotheses requires a sufficiently large sample size. A ubiquitous challenge in empirical research is that recruiting large samples is difficult due to resource constraints (e.g., time, money, labor) or ethical constraints (e.g., inconvenience or participation risks). However, if the sample sizes are small, random noise can mask the true effects, e.g., with regard to observed behaviour or cognitive processes. In a

formal hypothesis testing framework, this trade-off between resource constraints and statistical significance is best described by the measure of statistical power. Statistical power describes the probability of finding an effect that is actually present in the population of interest. In general, we want our sample size to be large enough to achieve high statistical power while using as few resources as necessary.

Justifying Sample Sizes

The recent replication crisis has put low statistical power and replicability of scientific research into focus (Button et al., 2013; Open Science Collaboration, 2015). Starting from the observation that most published research results might be wrong (Ioannidis, 2005; Simmons et al., 2011), there have been several developments to improve the replicability of scientific studies (Shrout & Rodgers, 2018). One of these are registered reports, in which research projects are reviewed and conditionally accepted based on sound methodology rather than on the statistical significance of the result. In registered reports, justification of sam-

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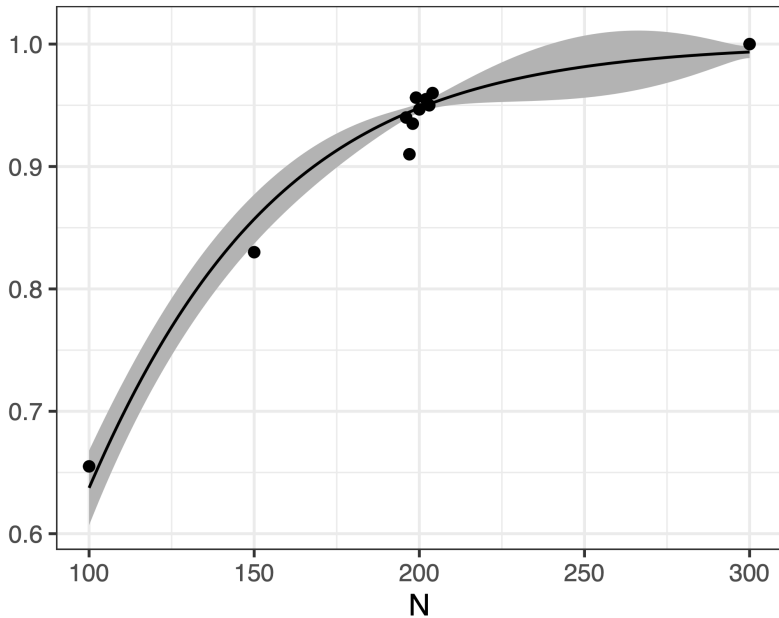
The R syntax for this study is available at the Open Science Framework at <https://osf.io/xebzj/>. All R packages used in this study are available on CRAN.

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Surrogate modeling

- Surrogate modeling aims to approximate a relationship that is costly to investigate with a cheaper function (Bhosekar & Ierapetritou, 2018; Forrester & Keane, 2009).
- We can adopt the idea of surrogate modeling to the functional relationship between study design parameters and power.
- Using this functional relationship, we can predict the power for a sample size that we did not perform a simulation at beforehand.
- Surrogate modeling is more efficient than grid search: In a simple example, our approach required only 20% of the computational effort and performed 50% more simulation runs that used the optimal sample size (Zimmer & Debelak, 2022).

Example



Slide explaining how we calculate the final sample size

1. User specifies input parameters

- The expected large sample performance of the model.
- The range of sample sizes over which to search.
- The number of signal and noise parameters.
- The expected outcome prevalence.

2. Set data, model, and metric functions based on user input

- Use defaults, but can be specified (e.g. `model = "lasso"`).

3. Tune the data generating model

4. Perform search

We then return the minimum sample that is within 10% of expected large sample performance in 80% of replications

Example 1: Binary outcome, logistic regression

including comparison with `pmsampsize`

Example 2: Binary outcome, LASSO regression

Example 3: Custom model function

`simulate_custom`

XGBoost

Maybe put slide with simulations here?

Development status

Package in development; functioning, but more testing needed.

- Follow fediscience.org/@ewan for updates.
- Or enter an email address at tinyurl.com/is-it-ready-yet to get one email when a public release is available.

Please come and talk to us.

Criteria/models/etc. all subject to change.

What's next? (1/2)

1. Machine learning

2. Longitudinal data

What's next? (2/2)

3. Common data types

e.g., clinical, NLP, genetic.

4. Performance

Thank you for listining.

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