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# calibmsm: An R package for calibration plots of the transition probabilities in a multistate model

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#### Abstract

Multistate models, which allow users to answer a wide range of clinical questions, are becoming a more commonly used tool for clinical prediction. It is paramount to evaluate the calibration (as well as other metrics) of a risk prediction model before implementation of the model in practice. Currently no software exists to aid in assessing the calibration of a multistate model. **calibmsm** has been developed to simplify this process for practicing model developers. Calibration of the transition probabilities between given follow up times is made possible through three approaches. The first two utilise calibration techniques for binary and multinomial logistic regression models in combination with inverse probability of censoring weights, whereas the third utilises psuedo-values. All methods are implemented in conjunction with landmarking.

This article details the methodology and provides a comprehensive example on how to assess the calibration of a model developed to predict recovery, adverse events, relapse and survival in patients with blood cancer after a transplantation. This is an illustrative example to showcase how to use the package and motivate a discussion around which of the calibration methods is most appropriate.

Keywords: clinical prediction, calibration, validation, multistate, multi-state, R.

## 1. Introduction

Risk prediction models enable the prediction of clinical events in either diagnostic or prognostic settings (van Smeden et al. 2021) and are used widely to inform clinical practice. A

multistate model (Putter et al. 2007) may be used when there are multiple outcomes of interest, or when a single outcome of interest may be reached via intermediates states. Using a multistate model for prediction is important when the development of an intermediate condition occurring post index date may have an impact on the risk of future outcomes of interest. Multistate models are increasingly being developed given the many medical scenarios where modelling patient movement between clinical states is of intrest (Putter et al. 2006; Le-Rademacher et al. 2018; Lintu et al. 2022; Masia et al. 2017). All risk prediction models developed for use in clinical practice should be validated in a relevant cohort prior to implementation (Steyerberg and Harrell Jr 2016; Sperrin et al. 2022). A key part of the validation process is assessment of the calibration of the model (Van Calster et al. 2019). Ideally calibration curves should be produced which allow evaluation of the calibration over the entire distribution of predicted risk, corresponding to moderate assessment of calibration (Van Calster et al. 2016).

The R package **mstate** (de Wreede *et al.* 2011) provides a comprehensive set of tools to develop a multistate model for a continuously observed multistate survival process. However, currently no software exists to aid researchers in assessing the calibration of a multistate model that has been developed for the purposes of individual risk prediction. **calibmsm** has been developed to enable researchers to estimate calibration curves and scatter plots using three approaches outlined by Pate et al (XXXX REF project 6, may have to put on Arxiv), which focused on assessing the calibration of the transition probabilities out of the starting state. The work in this paper extends the framework to assess the calibration of transition probabilities out of any state j at any time s using landmarking (van Houwelingen 2007; Dafni 2011), provides more details on estimation of the inverse-probability of censoring weights (where relevant), and demonstrates the process for estimating confidence intervals. **calibmsm** is available from the Comprehensive R Archive Network at REF XXXX.

de Wreede et al. (2011) used data from the European Society for Blood and Marrow Transplantation (EBMT 2023) to showcase how to develop a multistate model for clinical prediction of outcomes after bone morrow transplantation in leukemia patients (Figure 1). In this study, we show how to assess the calibration of a model developed on the same EBMT data as a way of illustrating the syntax and workflows of calibmsm. This clinical example also highlights some important differences between the methods in how they deal with informative censoring and computational feasibility, which may impact future uptake of the methods. Details on the methodology are given in section 2. The clinical setting for our example and steps for data preparation and are described in section 3. In section 4 we show how to estimate calibration curves and scatter plots using calibmsm. Section 5 contains a discussion and summary.

# 2. Methodology

#### 2.1. Setup

Let  $X(t) \in \{1, ..., K\}$  be a multistate survival process with K states. We assume a multistate model has already been developed and we want to assess the calibration of the predicted transition probabilities,  $\hat{p}_{j,k}(s,t)$ , in a cohort of interest. The transition probabilities are the probability of being in state k at time t, if in state t at time t, where t at time t, where t at time t

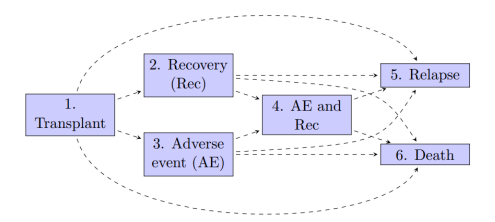


Figure 1: A six-state model for leukemia patients after bone marrow transplantation. Figure taken from (de Wreede *et al.* 2011).

estimate observed event probabilities:

$$o_{i,k}(s,t) = P[X(t) = k | X(s) = j, \hat{p}_{i,k}(s,t)].$$

In the absence of censoring, this can be done using cross sectional calibration techniques in a landmark (van Houwelingen 2007; Dafni 2011) cohort of individuals who are in state j at time s (i.e. methods to assess the calibration of models predicting binary or polytomous outcomes). In the presence of censoring, calibration must be assessed in this landmark cohort of individuals either using these cross sectional techniques in combination with inverse probability of censoring weights, or through pseudo-values. These approaches are detailed in sections 2.2 - 2.6.

#### 2.2. BLR-IPCW calibration curves

The first approach produces calibration curves using a framework for binary logistic regression models with inverse probability of censoring weights (BLR-IPCW). Let  $I_k(t)$  be an indicator for whether an individual is in state k at time t.  $I_k(t)$  is then modeled using a flexible approach with  $\hat{p}_{j,k}(s,t)$  as the sole predictor. This model is fit in the landmark cohort (in state j at time s) of individuals uncensored at time t, weighted using inverse probability of censoring weights (section 2.4). We suggest using a loess smoother:

$$I_k(t) = \operatorname{loess}(\hat{p}_{j,k}(s,t)), \tag{1}$$

or a logistic regression model with restricted cubic splines:

$$logit(I_k(t)) = rcs(logit(\hat{p}_{i,k}(s,t))). \tag{2}$$

Any flexible model for binary outcomes could be used, but these are the most common and are implemented in this package. Observed event probabilities  $\hat{o}_{j,k}(s,t)$  are then estimated as fitted values from these models. The calibration curve is plotted using the set of points

 $\{\hat{p}_{j,k}(s,t),\hat{o}_{j,k}(s,t)\}$ . For unbiased calibration curves, we make the assumption that in the reweighted population, each outcome  $I_k(t)$  is independent from the censoring mechanism.

## 2.3. MLR-IPCW calibration plots

The second approach produces calibration scatter plots using a framework for multinomial logistic regression models with inverse probability of censoring weights (MLR-IPCW). Let  $I_X(t)$  be an polytomous indicator variable taking values  $I_X(t) \in \{1, ..., K\}$  such that  $I_X(t) = k$  if an individual is in state k at time t. The nominal recalibration framework of Van Hoorde  $et\ al.\ (2014,\ 2015)$  is then applied in the landmark cohort of individuals uncensored at time t, weighted using inverse probability of censoring weights (section 2.4). First calculate the log-ratios of the predicted transition probabilities:

$$\hat{LP}_k = ln\left(\frac{\hat{p}_{j,k}(s,t)}{\hat{p}_{j,k_{ref}}(s,t)}\right),\,$$

Then fit the following multinomial logistic regression model:

$$ln\left(\frac{P[I_X(t) = k]}{P[I_X(t) = k_{ref}]}\right) = \alpha_k + \sum_{h=2}^K \beta_{k,h} * s_k(\hat{LP}_h),$$
(3)

where  $k_{ref}$  is an arbitrary reference category which can be reached from state j,  $k \neq k_{ref}$  takes values in the set of states that can be reached from state j, and where s is a vector spline smoother (Yee 2015). Observed event probabilities  $\hat{o}_{j,k}(s,t)$  are then estimated as fitted values from this model. This results in a calibration scatter plot rather than a curve due to all states being modeled simultaneously, as opposed to BLR-IPCW, which is a "one vs all" approach. The scatter occurs because the observed event probabilities for state k vary depending on the predicted transition probabilities of the other states. This is a stronger (Van Calster et al. 2016) form of calibration than that evaluated by BLR-IPCW, and will also result in observed event probabilities which sum to 1. For an unbiased calibration scatter plot, we make the assumption that in the reweighted population, the outcome  $I_X(t)$  is independent from the censoring mechanism.

#### 2.4. Estimation of the inverse probability of censoring weights

The estimand for the weights is  $w_j(s,t)$ , the inverse of the probability of being uncensored at time t if in state j at time s:

$$w_j(s,t) = \frac{1}{P[t_{cens} > t | t > s, X(s) = j, \mathbf{Z}, \mathbf{X(t)}]},$$

where  $\mathbf{X}(\mathbf{t})$  denotes the history of the multistate survival process up to time t, including the transition times. First the estimator  $\hat{P}[t_{cens} > t | t > s, X(s) = j, \mathbf{Z}]$  is calculated by developing an appropriate survival model. The outcome in this model is the time until censoring occurs. Moving into an absorbing state prevents censoring from happening and is treated as a censoring mechanism in this model (i.e. a competing risks approach is not taken when fitting this model).  $\mathbf{X}(\mathbf{t})$  is explicitly conditioned on when defining  $w_j(s,t)$  because the

weights must reflect that censoring can no longer be observed for an individual if they enter an absorbing state at some time  $t_{abs} < t$ . Therefore

$$\hat{P}[t_{cens} > t | t > s, X(s) = j, \mathbf{Z}, \mathbf{X(t)}] = \hat{P}[t_{cens} > min\{t, t_{abs}\} | t > s, X(s) = j, \mathbf{Z}]$$

In calibmsm,  $\hat{P}[t_{cens} > t | t > s, X(s) = j, \mathbf{Z}]$  is estimated using a cox proportional hazards model where all predictors  $\mathbf{Z}$  are assumed to have a linear effect on the log-hazard. However users can also input their own vector of weights. Given the BLR-IPCW and MLR-IPCW approaches are both reliant on correct estimation of the weights, we encourage users to take the time to carefully estimate weights using a well specified model.

Stabilised weights can be estimated by multiplying by the weights  $w_j(s,t)$  by the mean probability of being uncensored:

$$w_j^{stab}(s,t) = \frac{P[t_{cens} > t | t > s, X(s) = j]}{P[t_{cens} > t | t > s, X(s) = j, \mathbf{Z}, \mathbf{X(t)}]}.$$

The numerator can be estimated using an intercept only model, and note there is no dependence on X(t).

Another option is to estimate w(s,t), which is the inverse of the probability of being uncersored at time t if uncensored at time s:

$$w(s,t) = \frac{1}{P[t_{cens} > t | t > s, \mathbf{Z}, \mathbf{X(t)}]}.$$

This can be estimated using the same approach as for  $w_j(s,t)$ , except there is no requirement to be in state j when landmarking at time s. If the censoring mechanism is non-informative after conditioning on the predictors  $\mathbf{Z}$ , then  $w(s,t)=w_j(s,t)$ , and any consistent estimator for w(s,t) will be a consistent estimator of  $w_j(s,t)$ . The advantage is that  $\hat{w}(s,t)$  is calculated by developing a model in the cohort of individuals uncensored at time s, which is a larger cohort than those uncensored and in state j at time s. Therefore  $\hat{w}(s,t)$  will be a more precise estimator than  $\hat{w}_j(s,t)$ . On the contrary, if the assumption of non-informative censoring after conditioning on  $\mathbf{Z}$ , there is a risk of bias.

Finally, we state the importance of using inverse probability of censoring weights, even if the censoring mechanism is believed to be completely non-informative (i.e. happens at random). All multi-state models must have an absorbing state, entry into which prevents censoring from happening. This induces a dependence between the outcome and the censoring mechanism which must be adjusted for using inverse probability of censoring weights.

#### 2.5. Pseudo-value calibration plots

The third approach produces calibration curves using pseudo-values. Pseudo-values can be used to estimate quantities of interest in censored survival data and multistate survival data. For certain estimators  $\hat{\theta}$  (where  $\theta$  must take the form of an expectation), the pseudo-value for individual i is defined as:

$$\hat{\theta}^i = n * \hat{\theta} - (n-1) * \hat{\theta}^{-i}.$$

where  $\hat{\theta}^{-i}$  is equal to  $\hat{\theta}$  estimated in a cohort without individual i. One such estimator for the transition probabilities in a multistate model, P[X(t) = k | X(s) = j], is the Landmark Aalen-Johansen estimator (Putter and Spitoni 2018). Note that this is equivalent to landmarking the cohort and estimating the Aalen-Johansen estimator (Aalen and Johansen 1978), which is what's done in this package. The resulting pseudo-values are vectors with K elements, one for each possible transition. These pseudo-values can replace the outcome  $I_k(t)$  in equations (1) and (2) in order to estimate  $o_{j,k}(s,t)$ .

Pseudo-values are based on the same assumptions as the underlying estimator  $\hat{\theta}$ . The Landmark Aalen-Johansen estimator requires non-informative censoring. The approach to alleviate this is to estimate the pseudo-values within sub-groups of individuals, now making the assumption that censoring is non-informative within the specified subgroups. This can be done by calculating the pseudo-values within subgroups defined by baseline predictors, or subgroups defined by the predicted transition probabilities  $\hat{p}_{j,k}(s,t)$ . Both options are implemented in this package. When pseudo-values are calculated within subgroups, they are still used as the outcome in models (1) and (2) in the same way. Note that the pseudo-values  $\hat{\theta}^i$  are continuous, as opposed to binary  $I_k(t)$ , but the link function in model (2) remains the same to ensure  $\hat{o}_{j,k}(s,t)$  are between zero and one.

#### 2.6. Estimation of confidence intervals

Confidence intervals for both BLR-IPCW and pseudo-value calibration curves can be estimated using bootstrapping. A process for estimating the confidence intervals around the BLR-IPCW calibration curves is as follows:

- 1. Resample validation dataset with replacement
- 2. Landmark the dataset for assessment of calibration
- 3. Calculate inverse probability of censoring weights
- 4. Fit the preferred calibration model in the landmarked dataset (restricted cubic splines or loess smoother)
- 5. Generate observed event probabilities for a fixed vector of predicted transition probabilities (specifically the predicted transition probabilities from the non-bootstrapped landmark validation dataset)

This will produce a number of bootstrapped calibration curves, all plotted over the same vectors of predicted transition probabilities. Taking the  $\frac{\alpha}{2}$  and  $(1-\frac{\alpha}{2})$  percentiles of the observed event probabilities for each predicted transition probability gives the required  $1-\alpha$  confidence interval around the estimated calibration curve. To estimate confidence intervals for the pseudo-value calibration curves using bootstrapping, the same procedure is applied except the third step is replaced with 'calculate the pseudo-values within the landmarked bootstrapped dataset'. For the pseudo-value approach, this will be highly computationally demanding as the pseudo-values must be estimated in every bootstrap dataset.

If using the pseudo-value method, confidence intervals can be calculated using parametric estimates of the standard error when making predictions of the observed event probabilities (i.e. when obtaining fitted values from models (1) or (2)). To obtain a parametric confidence

interval when using the BLR-IPCW approach, firstly, stabilised weights must be used, otherwise the confidence intervals will be too small due to an inflated sample size Hernan and Robins (2020). Furthermore, a robust sandwich-type estimator should be used to estimate the standard error. However, robust sandwich estimators are known to result in conservative confidence intervals (i.e. too large), Hernan and Robins (2020) also highlighted in this simulation by Austin (2016). Furthermore, the size of the confidence interval will also be underestimated as uncertainty in estimation of the weights is not considered, although this is only a real issue at small sample sizes.

Both the parametric and bootstrap confidence intervals have been built into **calibmsm**. However for the reasons outlined above, we recommend using parametric confidence intervals for the pseudo-value calibration curves and bootstrapping for the BLR-IPCW calibration curves. While theoretically feasible using bootstrapping, it is currently unclear how to present confidence intervals for each data point in the calibration scatter plots.

# 3. Clinical setting and data preperation

We utilise data from the European Society for Blood and Marrow Transplantation (EBMT 2023), containing multistate survival data after a transplant for patients with blood cancer. The start of follow up is the day of the transplant and the initial state is alive and in remission. There are three intermediate events (2: recovery, 3: adverse event, or 4: recovery + adverse event), and two absorbing states (5: relapse and 6: death). This data is available from the mstate package (de Wreede et al. 2011).

Four datasets are provided to enable assessment of a multistate model fitted to these data. The first is ebmtcal, which is the same as the ebmt dataset provided in mstate, with two extra variables derived: time until censoring (dtcens) and an indicator for whether censoring was observed (dtcens.s = 1) or an absorbing state was entered (dtcens.s = 0). This dataset contains baseline information on year of transplant (year), age at transplant (age), prophylaxis given (proph), and whether the donor was gender matched (match). The second dataset provided is msebmtcal, which is the ebmt dataset converted into a dataset of class msdata using the process in de Wreede et al. (2011). It contains all transition times, an event indicator for each transition, as well as a trans attribute containing the transition matrix.

```
R> library("calibmsm")
R> data("ebmtcal")
R> head(ebmtcal)
  id
      rec rec.s
                    ae ae.s recae recae.s
                                               rel rel.s
                                                            srv
        22
                   995
                                995
                                               995
                                                            995
                                                                     0
1
   1
                1
                           0
   2
        29
2
                1
                     12
                           1
                                 29
                                               422
                                                            579
                                                                     1
3
   3 1264
                0
                    27
                           1
                               1264
                                           0 1264
                                                        0 1264
                                                                     0
   4
                                                                     1
4
        50
                    42
                           1
                                 50
                                            1
                                                84
                                                        1
                                                            117
                1
5
   5
        22
                1 1133
                           0
                               1133
                                            0
                                               114
                                                        1 1133
                                                                     0
   6
                           1
6
        33
                1
                    27
                                 33
                                            1 1427
                                                        0 1427
                                                                     0
                                          match dtcens dtcens.s
        year agecl proph
1 1995-1998 20-40
                        no no gender mismatch
                                                     995
                                                                 1
2 1995-1998 20-40
                        no no gender mismatch
                                                     422
                                                                 0
```

```
1264
                                                            1
3 1995-1998 20-40
                      no no gender mismatch
4 1995-1998 20-40
                                                            0
                            gender mismatch
                                                  84
                                                            0
5 1995-1998
              >40
                            gender mismatch
                                                114
                      nο
6 1995-1998 20-40
                      no no gender mismatch
                                                1427
```

R> data("msebmtcal")

R> head(msebmtcal)

	id	from	to	trans	Tstart	Tstop	time	$\operatorname{status}$
1	1	1	2	1	0	22	22	1
2	1	1	3	2	0	22	22	0
3	1	1	5	3	0	22	22	0
4	1	1	6	4	0	22	22	0
5	1	2	4	5	22	995	973	0
6	1	2	5	6	22	995	973	0

In the work of de Wreede  $et\ al.\ (2011)$ , the focus is on predicting transition probabilities made at times s=0 and s=100 across a range of follow up times t, and comparing prognosis for patients in different states j. In this study we also focus on assessing the calibration of the transition probabilities made at these times. We assess calibration of the transition probabilities at t=5 years, a common follow up time for cancer prognosis, but calibration of the model may vary for other values of t. We estimate transition probabilities for each individual by developing a model as demonstrated in de Wreede  $et\ al.\ (2011)$ , following the theory of Putter  $et\ al.\ (2007)$ . The predicted transitions probabilities from each state j at times s=0 and s=100 are contained in stacked datasets tps0 and tps100 respectively. A leave-one-out approach was used when estimating these transition probabilities. This means each individual was removed from the development dataset when fitting the multistate model to estimate their transition probabilities. This approach allows validation to be assessed in the same dataset that the model was developed with minimal levels of in-sample optimism. The code for deriving all these datasets is provided in the source code for **calibmsm**.

```
R> data("tps0")
R> head(tps0)
```

```
pstate1
                 pstate2
                            pstate3
                                      pstate4
                                                pstate5
                                                          pstate6
  1 0.1139726 0.2295006 0.08450376 0.2326861 0.1504855 0.1888514
  2 0.1140189 0.2316569 0.08442692 0.2328398 0.1481977 0.1888598
  3 0.1136646 0.2317636 0.08274331 0.2325663 0.1504787 0.1887834
  4 0.1383878 0.1836189 0.07579429 0.2179331 0.1538475 0.2304185
  5 0.1233226 0.1609740 0.05508100 0.1828176 0.1425950 0.3352099
  6 0.1136646 0.2317636 0.08462424 0.2305854 0.1505534 0.1888087
         se1
                    se2
                               se3
                                          se4
                                                      se5
1 0.01291133 0.02369584 0.01257251 0.02323376 0.01648630 0.01601795 1
2 0.01291552 0.02374329 0.01256056 0.02324869 0.01632797 0.01603703 1
3 0.01289444 0.02375770 0.01245752 0.02322375 0.01647890 0.01601525 1
4 0.01857439 0.03004447 0.01462570 0.03018673 0.02124071 0.02416121 1
5 0.01944967 0.03419721 0.01367768 0.03423941 0.02329644 0.03688586 1
6 0.01289444 0.02375770 0.01257276 0.02317348 0.01649531 0.01602438 1
```

```
R> data("tps100")
R> head(tps100)
```

```
id
       pstate1
                  pstate2 pstate3 pstate4
                                             pstate5
                                                        pstate6
  1 0.7013881 0.05239271
                                 0
                                         0 0.1408120 0.1054072
   2 0.7012745 0.05261136
                                 0
                                         0 0.1407625 0.1053516
                                 0
                                         0 0.1407628 0.1053987
   3 0.7011368 0.05270176
                                         0 0.1700565 0.1045183
  4 0.6840325 0.04139266
                                 0
                                         0 0.1500344 0.1264764
  5 0.6804049 0.04308434
                                 0
   6 0.7011368 0.05270176
                                         0 0.1407628 0.1053987
         se1
                    se2 se3 se4
                                        se5
                                                    se6 j
1 0.04691168 0.02077138
                               0 0.03457006 0.03081258 1
                               0 0.03456448 0.03079617 1
2 0.04691218 0.02082871
3 0.04693068 0.02086917
                               0 0.03456101 0.03081033 1
                           0
4 0.05885230 0.02161973
                               0 0.04710517 0.03673242 1
                           0
5 0.06694739 0.02484634
                           0
                               0 0.04905043 0.04628088 1
6 0.04693068 0.02086917
                               0 0.03456101 0.03081033 1
```

# 4. Assessing calibration using calibration curves and scatter plots

The procedure for producing calibration plots requires the use of two functions. The first function, calib\_blr, calib\_pv or calib\_mlr, calculates the data for the calibration plot using the methods described in section 2. The second function, plot.calib\_blr, plot.calib\_pv or plot.calib\_mlr, produces the plots. Separating these processes allows users the flexibility of producing their own Figures once the calibration data has been estimated.

The validation cohort must be provided to functions calib\_blr, calib\_pv and calib\_mlr in two different formats. The data.raw argument requires a data.frame (one observation per individual) and is used to fit the calibration models. For methods BLR-IPCW and MLR-IPCW, data.raw should contain variables dtcens (censoring time) and dtcens.s (censoring indicator, dtcens.s = 1 if the individual is censored at time dtcens, dtcens.s = 0 otherwise), plus any baseline predictors  $\mathbf{Z}$  used to estimate the weights. For the pseudo-value approach, this dataset should contain any baseline predictors  $\mathbf{Z}$  which variables will be grouped by before calculating the pseudo-values. The data.mstate argument requires a dataset of class msdata, and is used to implement the landmarking. A dataset of this class must be produced using the package mstate (de Wreede et al. 2011). Both data.mstate and data.raw should contain corresponding patient ID variables id. The predicted transition probabilities out of state j at time s must then be specified through the tp.pred argument, which must contain a column for each transition k, even if the transition from j to k has zero probability. The rows in tp.pred must be ordered in the same way as those in data.raw. The datasets described in section 3 meet these criteria.

## **4.1.** Plots out of state j = 1 at time s = 0

We start by producing calibration curves for the predicted transition probabilities out of state j = 1 at time s = 0. Given all individuals start in state 1, there is no need to consider the

transition probabilities out of states  $j \neq 1$  at s = 0. Calibration is assessed at follow up time (t = 1826 days). We first evaluate calibration using the BLR-IPCW approach, implemented through the function calib\_blr.

The predicted transition probabilities from state j=1 at time s=0 are extracted from the object tps0. We choose to estimate the calibration curves using restricted cubic splines, and 3 knots are chosen given the reasonably small size of the dataset. Weights are estimated using the internal estimation procedure and the predictor variables year, agecl, proph and match. The w.landmark.type argument assigns whether weights are estimated using all individuals uncensored at time s, or only those uncensored and in state j at time s, as discussed in section 2.4. The maximum weight (w.max = 10) and stabilisation of weights (stabilised = TRUE) are left as default. Weights can also be manually specified using the weights argument. We request 95% confidence intervals for the calibration curves calculated through bootstrapping with 200 bootstrap replicates.

The first element of dat.calib.blr (named plotdata) contains 6 data frames for the calibration curves of the transition probabilities into each of the six states,  $k \in \{1, 2, 3, 4, 5, 6\}$ . Each data frame contains three columns, id: the identifier of each individual; pred: the predicted transition probabilities; obs: the observed event probabilities. The second element (named metadata) is a metadata argument containing information about the data and chosen calibration analysis.

R> str(dat.calib.blr[["plotdata"]])

```
List of 6
$ state1:'data.frame': 1778 obs. of 5 variables:
..$ id : int [1:1778] 2 4 5 7 10 13 14 16 18 19 ...
..$ pred : num [1:1778] 0.114 0.1384 0.1233 0.0974 0.1137 ...
..$ obs : num [1:1778] 0.11 0.104 0.105 0.124 0.11 ...
..$ obs.lower: num [1:1778] 0.0859 0.0789 0.0807 0.0881 0.0861 ...
..$ obs.upper: num [1:1778] 0.133 0.125 0.125 0.164 0.134 ...
```

```
$ state2:'data.frame':
                               1778 obs. of 5 variables:
               : int [1:1778] 2 4 5 7 10 13 14 16 18 19 ...
  ..$ id
               : num [1:1778] 0.232 0.184 0.161 0.212 0.232 ...
  ..$ pred
               : num [1:1778] 0.17 0.186 0.176 0.179 0.17 ...
  ..$ obs.lower: num [1:1778] 0.119 0.156 0.147 0.145 0.119 ...
  ..$ obs.upper: num [1:1778] 0.221 0.218 0.214 0.212 0.221 ...
 $ state3:'data.frame':
                               1778 obs. of 5 variables:
  ..$ id
               : int [1:1778] 2 4 5 7 10 13 14 16 18 19 ...
  ..$ pred
               : num [1:1778] 0.0844 0.0758 0.0551 0.0615 0.0844 ...
               : num [1:1778] 0.1249 0.1167 0.0919 0.1001 0.1248 ...
  ..$ obs.lower: num [1:1778] 0.0971 0.0901 0.0489 0.0614 0.0971 ...
  ..$ obs.upper: num [1:1778] 0.152 0.142 0.135 0.135 0.152 ...
 $ state4:'data.frame':
                               1778 obs. of 5 variables:
  ..$ id
               : int [1:1778] 2 4 5 7 10 13 14 16 18 19 ...
               : num [1:1778] 0.233 0.218 0.183 0.221 0.233 ...
  ..$ pred
               : num [1:1778] 0.243 0.224 0.185 0.228 0.243 ...
  ..$ obs
  ..$ obs.lower: num [1:1778] 0.207 0.195 0.159 0.197 0.207 ...
  ..$ obs.upper: num [1:1778] 0.282 0.255 0.22 0.258 0.282 ...
 $ state5:'data.frame':
                               1778 obs. of 5 variables:
               : int [1:1778] 2 4 5 7 10 13 14 16 18 19 ...
  ..$ id
               : num [1:1778] 0.148 0.154 0.143 0.144 0.149 ...
  ..$ pred
               : num [1:1778] 0.191 0.165 0.222 0.212 0.188 ...
  ..$ obs.lower: num [1:1778] 0.165 0.149 0.178 0.174 0.164 ...
  ..$ obs.upper: num [1:1778] 0.215 0.182 0.26 0.245 0.21 ...
 $ state6:'data.frame':
                               1778 obs. of 5 variables:
  ..$ id
               : int [1:1778] 2 4 5 7 10 13 14 16 18 19 ...
  ..$ pred
               : num [1:1778] 0.189 0.23 0.335 0.264 0.189 ...
               : num [1:1778] 0.207 0.254 0.316 0.28 0.207 ...
  ..$ obs.lower: num [1:1778] 0.177 0.225 0.282 0.252 0.177 ...
  ..$ obs.upper: num [1:1778] 0.225 0.279 0.355 0.303 0.225 ...
R> str(dat.calib.blr[["metadata"]])
List of 8
 $ valid.transitions
                       : num [1:6] 1 2 3 4 5 6
 $ assessed.transitions: num [1:6] 1 2 3 4 5 6
 $ CI
                       : num 95
 $ CI.R.boot
                       : num 200
 $ curve.type
                       : chr "rcs"
 $ ј
                       : num 1
 $ s
                       : num 0
 $ t.eval
                       : num 1826
```

Calibration curves can then be generated using plot.calib\_blr. This S3 generic was written for objects of class calib\_blr and produces the calibration plots using ggplot2 (REF XXXX). The calibration curves (Figure 2) indicate the level of calibration is different for the transition probabilities into each of the different states. The calibration into states 4 and 6 is good, as

R> plot(dat.calib.blr, combine = TRUE, nrow = 2, ncol = 3)

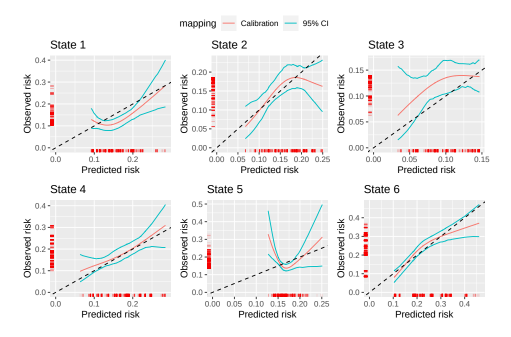


Figure 2: BLR-IPCW calibration curves out of state j = 1 at time s = 0.

both contain the line of prefect calibration across the entire range of predicted risk. State 2 has good calibration over the majority of the predicted risks but over predicts for individuals with the highest predicted risks. Transition probabilities into states 1 and 3 are over and under predicted respectively over most of the range of predicted risks, and large portions of the confidence intervals do not contain the line of perfect calibration. Importantly the calibration of the transition probabilities into state 5 (Relapse), a key clinical outcome in this clinical setting, is extremely poor.

Next we use the pseudo-value approach to assess calibration, implemented through the function calib\_pv. Instead of specifying how the weights are estimated, we now specify variables to define groups within which pseudo-values will be calculated (see section 2.5). We choose to calculate pseudo-values within individuals with the same year of transplant (group.vars), and then split individuals into a further three groups defined by their predicted risk (n.pctls). The number of percentiles should be increased in bigger validation datasets. Year of transplant was identified as a subgrouping variable because a later transplant resulted in a shorter possible follow up, an earlier administrative censoring time, and it was therefore highly predictive of being censored. A parametric confidence interval is estimated as recommended in section 2.6.

```
R> dat.calib.pv <-
+ calib_pv(data.mstate = msebmtcal,
+ data.raw = ebmtcal,
+ j=1,
+ s=0,
+ t.eval = t.eval,</pre>
```

R> plot(dat.calib.pv, combine = TRUE, nrow = 2, ncol = 3)

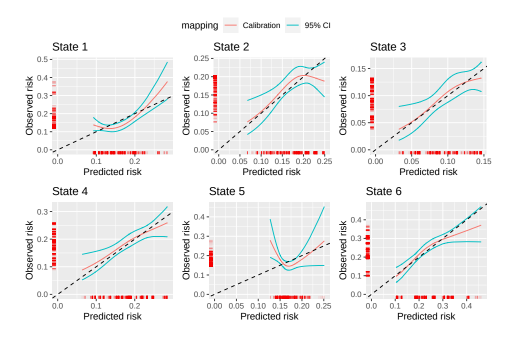


Figure 3: Pseudo-value calibration curves out of state j = 1 at time s = 0.

```
tp.pred = tps0 |>
dplyr::filter(j == 1) |>
dplyr::select(any_of(paste("pstate", 1:6, sep = ""))),

curve.type = "rcs",

rcs.nk = 3,
group.vars = c("year"),

n.pctls = 3,

CI = 95,
CI.type = "parametric")
```

Calibration curves were then generated using plot.calib\_pv. The pseudo-value calibration curves (Figure 3) are largely similar to the BLR-IPCW calibration curves (Figure 2). The agreement in the calibration curves from two completely distinct methods provides reassurance the assessment of calibration is correct. This is with the exception of state k=3, where the pseudo-value calibration plot indicates the transition probabilities are well calibrated, but the BLR-IPCW calibration plot indicates the transition probabilities under predict. If developing this model in a clinical setting, further exploration of this difference would be required. In particular, how the variable year of transplant acts on the censoring mechanism, and for which method the assumptions about independence of the censoring mechanism (section 2) are most likely to hold.

Next we use the MLR-IPCW to evaluate calibration which produces a calibration scatter plot. This is done using the calib\_mlr function, which has the same inputs as calib\_blr.

R> plot(dat.calib.mlr, combine = TRUE)

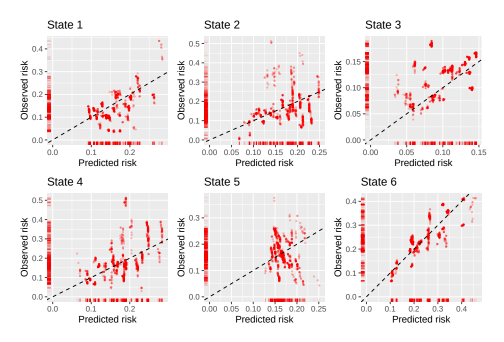


Figure 4: MLR-IPCW calibration curves out of state j = 1 at time s = 0.

The MLR-IPCW calibration plots, produced using plot.calib\_mlr are contained in Figure 4. Within each plot for state k, there is a large amount of variation in calibration of the transition probabilities depending on the predicted transition probabilities into states  $\neq k$ . One valuable insight from these plots is that the variance in the calibration of the transition probabilities into state 6, is considerably smaller than that of state 4, despite these two states both having good calibration according to the BLR-IPCW plots (arguably state 4 looked better calibrated). This means the calibration of the transition probabilities into state 6 remains reasonably consistent, irrespective of the risks of the other states. On the contrary, the calibration of the predicted transition probabilities into state 4 is more highly dependent on the predicted transition probabilities of the other states. This insight can be gained because MLR-IPCW is a stronger (Van Calster et al. 2016) form of calibration assessment than the BLR-IPCW and pseudo-value approaches.

# **4.2.** Plots out of state j = 1 and 3 at time s = 100

In the work of de Wreede et al. (2011) focus then shifts to comparing transition probabilities

when s=100 depending on whether an individual has had an adverse event (state 3) or remains in state 1 (post transplant). Our focus therefore now shifts to assessing the calibration of these transition probabilities. This is done through landmarking as described in section 2. In **calibmsm**, the process remains the same, changing the inputted values j and s, and specifying the appropriate predicted transition probabilities to the argument tp.pred. We start by producing the calibration plots for j=1 and s=100 using the BLR-IPCW (Figure 5) and pseudo-value (Figure 6) methods. Given the small number of data points in this analysis induced by landmarking, we do not produce calibration scatter plots using MLR-IPCW, which may be misleading given the lack of confidence intervals.

```
R> ### Calibration using BLR-IPCW
  dat.calib.blr.j1.s100 <-
     calib_blr(data.mstate = msebmtcal,
                    data.raw = ebmtcal,
+
                    j=1,
                     s = 100,
                     t.eval = t.eval,
                     tp.pred = tps100 |>
                      dplyr::filter(j == 1) |>
                      dplyr::select(any_of(paste("pstate", 1:6, sep = ""))),
                     curve.type = "rcs",
                    rcs.nk = 3,
                     w.covs = c("year", "agecl", "proph", "match"),
                     CI = 95,
                    CI.R.boot = 200)
R> ### Calibration using pseudo-values
  dat.calib.pv.j1.s100 <-
R>
     calib_pv(data.mstate = msebmtcal,
                   data.raw = ebmtcal,
                   j=1,
                   s=100,
+
                   t.eval = t.eval,
                    tp.pred = tps100 |>
                    dplyr::filter(j == 1) |>
                    dplyr::select(any_of(paste("pstate", 1:6, sep = ""))),
                    curve.type = "rcs",
                   rcs.nk = 3,
                   group.vars = c("year"),
                   CI = 95,
                   CI.type = "parametric")
```

There are only four calibration plots because no individuals in state j=1 at time s=100 are in states k=3 (adverse event) or k=4 (recovery + adverse event) after t=1826 days. The calibration of the predicted transition probabilities is very poor. Only for state k=6 is the observed risk a monotonically increasing function of the predicted transition probabilities. We follow this up with the pseudo-value calibration plots (Figure 6) which leads to similar conclusions, as again only state k=6 has a monotonically increasing calibration curve. The

R> plot(dat.calib.blr.j1.s100, combine = TRUE, nrow = 2, ncol = 2)

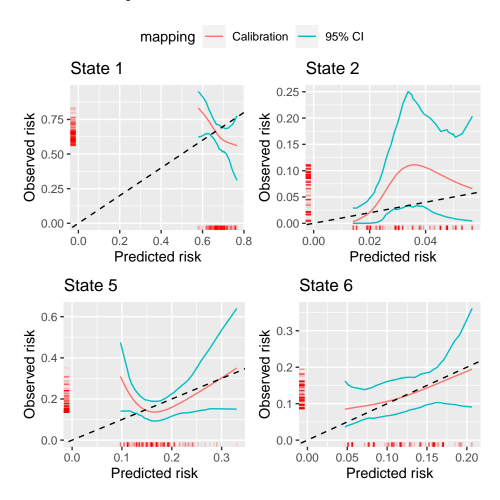


Figure 5: BLR-IPCW calibration curves out of state j = 1 at time s = 100.

confidence intervals are very large. For states k=2 and k=5, we cannot rule out that the poor calibration is a result of sampling variation as opposed to a poorly performing prediction model. A larger validation dataset would be required to get to the bottom of this. For state k=1, the calibration is extremely poor and is unlikely driven by sampling variation.

Next we produce calibration plots for j=3 and s=100 using the BLR-IPCW (Figure 7) and pseudo-value (Figure 8) methods.

R > plot(dat.calib.pv.j1.s100, combine = TRUE, nrow = 2, ncol = 2)

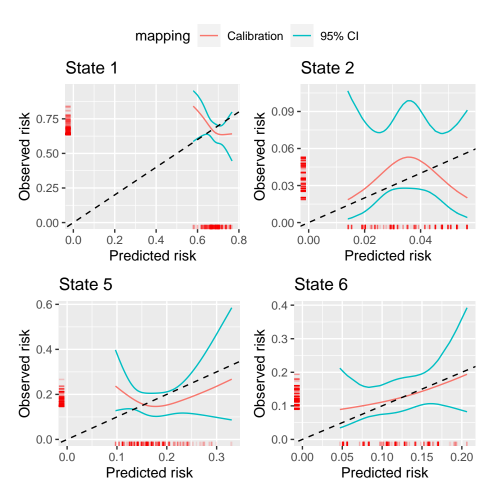


Figure 6: Pseudo-value calibration curves out of state j = 1 at time s = 100.

```
dplyr::select(any of(paste("pstate", 1:6, sep = ""))),
                    curve.type = "rcs",
                    rcs.nk = 3,
                    w.covs = c("year", "agecl", "proph", "match"),
                    CI = 95,
                    CI.R.boot = 200)
  ### Calibration using pseudo-values
R>
  dat.calib.pv.j3.s100 <-
R>
     calib_pv(data.mstate = msebmtcal,
                   data.raw = ebmtcal,
                   j=3,
                   s=100,
                   t.eval = t.eval,
                    tp.pred = tps100 |>
                    dplyr::filter(j == 3) />
                    dplyr::select(any_of(paste("pstate", 1:6, sep = ""))),
                    curve.type = "rcs",
                   rcs.nk = 3,
                   group.vars = c("year"),
                   CI = 95,
                   CI.type = "parametric")
```

Again there are only four possible states that an individual may transition into, although this includes states 3 (adverse event) and 4 (recovery + adverse event), instead of 1 (post transplant) and 2 (recovery). The calibration plots are better than for j=1. For transitions into states k=3,4 and 6, the calibration curves are monotonically increasing and the confidence interval contains the entire line of perfect calibration. This is true when calibration is assessed using BLR-IPCW or pseudo-values. Again the calibration of state 5 is very poor. This makes it difficult to base any clinical decisions on the predicted transition probabilities for relapse out of states j=1 or 3 at time s=100, whereas making clinical decisions based on the risk of death (k=6) after survival for 100 days is more viable, as this was well calibrated for both j=1 and j=3. With the exception of the transition probabilities from j=1 into state k=3 made at time s=0, there has been broad agreement between the calibration curves estimated using the BLR-IPCW and pseudo-value approaches. This provides some reassurance about the assessment of calibration, and that the assumptions on which each method is based are satisfied.

# 5. Discussion

Multistate models are a unique tool for prediction, allowing risks to be updated after key events happen post baseline. Development of multistate models for prediction is becoming more common, yet validation of such models is still very uncommon. A major barrier to implementation of statistical techniques is often the availability of software (Pullenayegum et al. 2016). calibmsm has been developed to aid in the implementation of techniques to assess the calibration of the transition probabilities from a multistate model. This paper has extended previously proposed methods for assessing the calibration of the transition probabilities out

R > plot(dat.calib.blr.j3.s100, combine = TRUE, nrow = 2, ncol = 2)

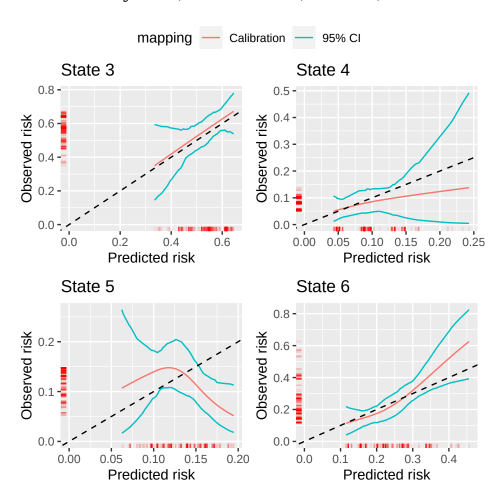
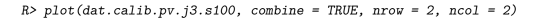


Figure 7: BLR-IPCW calibration curves out of state j=3 at time s=100.



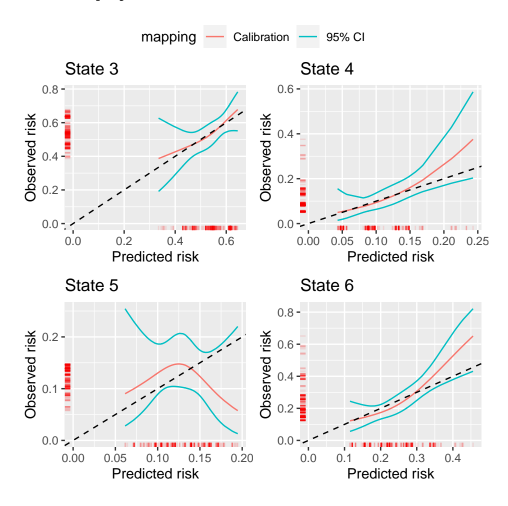


Figure 8: Pseudo-value calibration curves out of state j=3 at time s=100.

of the initial state (XXXX REF PROJECT 6), to the transition probabilities out of any state j at any time s.

All three methods (BLR-IPCW, MLR-IPCW and pseudo-value) have been shown to give an unbiased assessment of calibration under non-informative censoring mechanisms, and a predominately unbiased assessment of calibration under strongly informative censoring **XXXX REF PROJECT 6**. This paper found broadly similar evaluation of calibration when using the BLR-IPCW and pseudo-value methods, however there were discrepancies in the evaluation of calibration of the transition probabilities into state k = 3. Further research evaluating these methods performance in a wider range of simulation scenarios, and by a different research group (Boulesteix *et al.* 2013), would be highly valuable (Heinze *et al.* 2022).

It is also possible to use the calibmsm to validate competing risks models, which are a special case of a multistate models. However, graphical calibration curves already exist for assessing the calibration of a competing risks model (Austin et al. 2022). When fitting graphical calibration curves, it is assumed that censoring is independent of the outcome after conditioning on the complementary log-log transformation of the predicted risk, and the assumption of proportional hazards is made. When implementing BLR-IPCW, the assumption of independence between the censoring mechanism and the outcome in the reweighted population is made. For the pseudo-value approach, censoring is assumed to be non-informative within the specified subgroups. These are different approaches to dealing with the issue of informative censoring in the validation dataset when assessing the calibration of the model. We carried out a sensitivity analyses, comarping BLR-IPCW, pseudo-values and graphical calibration curves for assessing the calibration of a competing risks model out of the first state (post transplant) using the EBMT data (EBMT 2023). This is contained in a vignette 'comparison with graphical calibration curves' provided with the package. All methods resulted in similar calibration curves, but the relative performance of each methods in a wider range of scenarios is still unknown. A comparison of these methods in a simulation when the aforementioned assumptions do and do not hold, and under a range of sample sizes and multistate model structures, would be therefore valuable. Both of these recommendations for future work would constitute phase 3 and 4 studies in the evidence base for new methods (Heinze et al. 2022).

The BLR-IPCW and pseudo-value approaches have different computational burdens. A calibration curve can be obtained reasonably quickly using the BLR-IPCW approach, however estimation of confidence intervals using bootstrapping (the recommend method in section 2.6) will result in a high computational burden in large validation datasets. On the contrary, obtaining the calibration curve itself using the pseudo-value approach has a high computational burden due to estimation of the pseudo-values. Once these have been calculated, a calibration curve and confidence interval can be estimated quickly using parametric techniques, meaning estimation of the confidence interval adds minimal computational burden. We plan to extend the package to allow users to estimate the pseudo-values for each individual seperately before estimating the calibration curve. This will allow the first part of the process to be parallelised and makes estimation of calibration curves using the pseudo-value approach more feasible in large datasets.

Estimation of the weights is clearly of high importance for the BLR-IPCW and MLR-IPCW approaches. If the model to do so is misspecified, this could lead to incorrect evaluation of the calibration. It is possible this is what is causing the difference between the BLR-IPCW and pseudo-value approaches for the calibration of transition probabilities from state

j=1 at time s=0 into state k=3. This package is focused on creation of calibration curves, but is not a dedicated package for estimating inverse probability of censoring weights. We encourage users to create a well specified model for the weights. Custom functions for estimating the weights can be specified through the w.function argument in both calib\_blr and calib\_mlr. Alternatively, weights can be estimated externally and then specified though the weights argument. In this latter case, the internal bootstrapping procedure will not work, as the weights need to be re-estimated in each bootstrap dataset. We have provided a more detailed vignette about how to estimate calibration curves and confidence intervals using bootstrapping when defining your own function to estimate the weights (BLR-IPCW-manual-bootstrap-with-custom-function-for-estimating-the-weights).

In summary, **calibmsm** provides tools to assess the calibration of the transition probabilities of a multistate model or competing risks model using three approaches (BLR-IPCW, MLR-IPCW and pseudo-values). Further comparison of these approaches in targeted simulations to establish their performance under different censoring mechanisms and assumptions would be valuable.

# Computational details

# WILL UPDATE THIS SECTION ONCE PACKAGE HAS BEEN FINALISED AND UPLOADED TO CRAN

The results in this paper were obtained using R 4.3.0 with the **dplyr** 1.1.2, **tidyr** 1.3.0, **gg-plot2** 3.4.2, **ggpubr** 0.6.0, **Hmisc** 5.0.1, **rms** 6.6.0, **VGAM** 1.1.8, **boot** 1.3.28.1, **survival** 3.5.5, **stats** 4.3.0, **magrittr** 2.0.3. R itself and all packages used are available from the Comprehensive R Archive Network (CRAN) at <a href="https://cran.r-project.org/">https://cran.r-project.org/</a>.

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