

# *Machine Learning in Survival Analysis*



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## Getting Started



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Machine Learning in Survival Analysis

Raphael Sonabend, Andreas Bender

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## Preface

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“Everything happens to everybody sooner or later if there is time enough” - George Bernard Shaw

“...but in this world nothing can be said to be certain, except death and taxes.” - Benjamin Franklin

A logical consequence of Bernard Shaw’s quote is that if there is time enough, then everybody will have experienced a given event at some point. This is one of the central assumptions to survival analysis (specifically to single-event analysis, but we’ll get to that later). As nothing can be certain (except death and taxes), machine learning can be used to predict the probability people will experience the event and *when*. This is exactly the problem that this book tackles.

With immortality only being a theoretical concept, there is never ‘time enough’, hence survival analysis assumes that the event of interest is guaranteed to occur within an object’s lifetime. This event could be a patient entering remission after a cancer diagnosis, the lifetime of a lightbulb after manufacturing, the time taken to finish a race, or any other event that is observed over time. Survival analysis differs from other fields of Statistics in that uncertainty is explicit encoded in the survival problem; this uncertainty is known as ‘censoring’. For example, say a model is being built to predict when a marathon runner will finish a race and to learn this information the model is fed data from every marathon over the past five years. Across this period, there will be many runners who never finish their race. Instead, these runners are said to be ‘censored’ and the model uses all information up until the point of censoring (dropping out of the race), and learns that they ran for at least as long as their censoring time (the time they dropped out). Censoring is unique to survival analysis and without the presence of censoring, survival analysis is mathematically equivalent to regression.

This book covers survival analysis in the most common right-censoring setting for independent censoring, as well as discussing competing risk frameworks for dependent censoring - these terms will all be covered in the introduction of the book.

**A note from Raphael:** I wrote my PhD thesis about machine learning applications to survival analysis as I was interested in understanding why more researchers were not using machine learning models for survival analysis. Since then I’ve had the pleasure to work with, and advise, researchers across different sectors, including pharmaceutical companies, governmental agencies, funding organisations, and research institutions. I hope that this book continues to help researchers discover machine learning survival analysis and to navigate the nuances and complexities it presents.

**A note from Andreas:** FIXME.

## Overview

This textbook is intended to fill a gap in the literature by providing a comprehensive introduction to machine learning in the survival setting. If you are interested in machine learning or survival analysis separately then you might consider James et al. (2013), Hastie, Tibshirani, and Friedman (2001), Bishop (2006) for machine learning and Collett (2014), J. D. Kalbfleisch and Prentice (1973) for survival analysis. This book serves as a complement to the above examples and introduces common machine learning terminology from simpler settings such as regression and classification, but without diving into the detail found in other sources, instead focusing on extension to the survival analysis setting.

This book may be useful for Masters or PhD students who are specialising in machine learning in survival analysis, machine learning practitioners looking to work in the survival setting, or statisticians who are familiar with survival analysis but less so with machine learning. The book could be read cover-to-cover, but this is not advised. Instead it may be preferable to dip into sections of the book as required and use the ‘signposts’ that direct the reader to sections of the book that are relevant to each other.

The book is split into five parts:

**Part I: Survival Analysis and Machine Learning** The book begins by introducing the basics of survival analysis and machine learning and unifying terminology between the two to enable meaningful description of ‘machine learning in survival analysis’ (MLSA). In particular, the survival analysis ‘task’ and survival ‘prediction types’ are defined.

**Part II: Evaluation** The second part of the book discusses measures for evaluating survival models. These are presented in different classes that reflect the prediction types identified in Part I. In each chapter, the measure class is introduced, particular metrics are listed, and commentary is provided on how and when to use the measures. The final chapter of Part II briefly discusses when to use a given measure class and provides recommendations for model comparison.

**Part III: Models** Part III is a deep dive into machine learning models for solving survival analysis problems. This begins with ‘classical’ models that may not be considered ‘machine learning’ and then continues by exploring different classes of machine learning models including random forests, support vector machines, gradient boosting machines, neural networks, and other less common classes. Each model class is introduced in the simpler regression setting and then extensions to survival analysis are discussed. Differences between model implementations are not discussed, instead the focus is on understanding how these models are built for survival analysis - in this way readers are well-equipped to independently follow individual papers introducing specific implementations.

**Part IV: Reduction Techniques** The next part of the book introduces reduction techniques in survival analysis, which is the process of solving the survival analysis task by using methods from other fields. In particular, chapters focus on demonstrating how any survival model can be used in the competing risks setting, discrete time modelling, Poisson methods, pseudovalues (reduction to regression), and other advanced modelling methods.

**Part V: Extensions and Outlook** The final part of the book provides some miscellaneous chapters that may be of use to readers. The first chapter lists common practical problems that occur when running survival analysis experiments and solutions that we have found useful. The next lists open-source software at the time of writing for running machine learning survival analysis experiments. The final chapter is our outlook on survival analysis and where the field may be heading.

Exercises are provided at the end of the book so you can test yourself as you go along.

### Citing this book

Whilst this book remains a work in progress you can cite it as

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```

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We hope you enjoy reading this book.

Raphael and Andreas



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## **Authors**

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Raphael Sonabend is the CEO and Co-Founder of OSPO Now, a company providing virtual open-source program offices as a service. They are also a Visiting Researcher at Imperial College London. Raphael holds a PhD in statistics, specializing in machine learning applications for survival analysis. They created the R packages `mlr3proba`, `survivalmodels`, and the Julia package `SurvivalAnalysis.jl`. Raphael co-edited and co-authored *Applied Machine Learning Using mlr3 in R* (Bischl et al. 2024).

Andreas Bender is...



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## *Acknowledgments*

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We would like to gratefully acknowledge our colleagues that reviewed the content of this book, including: Lukas Burk, Cesaire Fouodo.



# Symbols and Notation

⚠ Minor changes expected!

This page is a work in progress and minor changes will be made over time.

The most common symbols and notation used throughout this book are presented below; in rare cases where different meanings are intended within the book, this will be made clear.

## Fonts, matrices, vectors

A lower-case letter in normal font,  $x$ , refers to a single, fixed observation. When in bold font, a lower-case letter,  $\mathbf{x}$ , refers to a vector of fixed observations, and an upper-case letter,  $\mathbf{X}$ , represents a matrix. Calligraphic letters,  $\mathcal{X}$ , are used to denote sets.

A matrix will always be defined with its dimensions using the notation,  $\mathbf{X} \in \mathcal{X}^{n \times p}$ , or if for example  $\mathcal{X}$  is the set of Reals, it may be written as “ $\mathbf{X}$  is a  $n \times p$  Real-valued matrix”, analogously for integer-valued matrices etc. By default, a ‘vector’ will refer to a column vector, which may be thought of as a matrix with  $n$  rows and one column, and may be represented as:

$$\mathbf{x} = \begin{pmatrix} x_1 \\ x_2 \\ \vdots \\ x_n \end{pmatrix}$$

Vectors are usually defined using transpose notation, for example the vector above may instead be written as  $\mathbf{x}^\top = (x_1 \ x_2 \ \cdots \ x_n)$  or  $\mathbf{x} = (x_1 \ x_2 \ \cdots \ x_n)^\top$ . Vectors may also be defined in a shortened format as  $\mathbf{x} \in \mathcal{X}^{n \times 1}$  or more simply  $\mathbf{x} \in \mathcal{X}^n$ , which implies a column vector of length  $n$  with elements as represented above.

A letter in normal font with one subscript refers to a single element from a vector. For example, given  $\mathbf{x} \in \mathcal{X}^n$ , the  $i$ th element is denoted  $x_i$ . Given a matrix  $\mathbf{X} \in \mathcal{X}^{n \times p}$ , a bold-face lower-case letter with a single subscript refers to the row of a matrix, for example the  $i$ th row would be  $\mathbf{x}_i = (x_{i;1} \ x_{i;2} \ \cdots \ x_{i;p})^\top$ . Whereas a column is referenced with a semi-colon before the subscript, for example the  $j$ th column would be  $\mathbf{x}_{:j} = (x_{1;j} \ x_{2;j} \ \cdots \ x_{n;j})^\top$ . Two subscripts can be used to reference a single element of a matrix, for example  $x_{i;j} \in \mathcal{X}$  would be the element in the  $i$ th row and  $j$ th column of  $\mathbf{X}$ .

## Functions

Typically, a ‘hat’,  $\hat{x}$ , will refer to the prediction or estimation of a variable,  $x$ , with bold-face used again to represent vectors. A ‘bar’,  $\bar{x}$ , refers to the sample mean of  $\mathbf{x}$ . Capital letters in normal font,  $X$ , refer to scalar or vector random variables, which will be made clear from context.  $\mathbb{E}(X)$  is the expectation of the random variable  $X$ . We write  $A \perp\!\!\!\perp B$ , to denote that  $A$  and  $B$  are independent, i.e., that  $P(A \cap B) = P(A)P(B)$ .

A function  $f$ , will either be written as a formal map of domain to codomain,  $f : \mathcal{X} \rightarrow \mathcal{Y}; (x, y) \mapsto f(x, y)$  (which is most useful for understanding inputs and outputs), or more simply and commonly as  $f(x, y)$ . Given a random variable,  $X$ , following distribution  $\zeta$  (mathematically written  $X \sim \zeta$ ), then  $f_X$  denotes the probability density function, and analogously for other distribution defining functions such as the cumulative distribution function, survival function, etc. In the survival analysis context (Chapter 3), a subscript “0” refers to a “baseline” function, for example,  $S_0$  is the baseline survival function.

Finally,  $\exp$ , refers to the exponential function,  $f(x) = e^x$ , and  $\log$  refers to the natural logarithm  $\ln(x) = \log_e(x)$ .

## Variables and acronyms

Common variables and acronyms used in the book are given in Table 0.1 and Table 0.2 respectively.

Table 0.1: Common variables used throughout the book.

Variable	Definition
$\mathbb{R}, \mathbb{R}_{>0}, \mathbb{R}_{\geq 0}, \bar{\mathbb{R}}$	Set of Reals, positive Reals (excl. zero), non-negative Reals (incl. zero), and Reals including $\pm\infty$ .
$\mathbb{N}_{>0}$	Set of Naturals excluding zero.
$(\mathbf{X}, \mathbf{t}, \boldsymbol{\delta})$	Survival data where $\mathbf{X} \in \mathbb{R}^{n \times p}$ is a real-valued matrix of $n$ observations (rows) and $p$ features (columns), $\mathbf{t} \in \mathbb{R}^n$ is a vector of observed outcome times, and $\boldsymbol{\delta} \in \mathbb{R}^n$ is a vector of observed outcome indicators.
$\boldsymbol{\beta}$	Vector of model coefficients/weights, $\boldsymbol{\beta} \in \mathbb{R}^p$ .
$\boldsymbol{\eta}$	Vector of linear predictors, $\boldsymbol{\eta} = (\eta_1 \ \eta_2 \ \dots \ \eta_n)^\top$ , where $\boldsymbol{\eta} = \mathbf{X}\boldsymbol{\beta}$ and $\eta_i = \mathbf{x}_i^\top \boldsymbol{\beta}$ .
$\mathcal{D}, \mathcal{D}_{train}, \mathcal{D}_{test}$	Dataset, training data, and testing data.

Table 0.2: Common acronyms used throughout the book.

Acronym	Definition
AFT	Accelerated Failure Time
cdf	Cumulative Distribution Function
chf	Cumulative Hazard Function
CPH	Cox Proportional Hazards

Acronym	Definition
GBM	Gradient Boosting Machine
GLM	Generalised Linear Model
IPC(W)	Inverse Probability of Censoring (Weighted)
ML	Machine Learning
pdf	Probability Density Function
PH	Proportional Hazards
(S)SVM	(Survival) Support Vector Machine
t.v.i.	Taking Values In



# 1

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## Introduction

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! Major changes expected!

This page is a work in progress and major changes will be made over time.

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### TODO

- Mention somewhere that SA can be used to solve T-year prediciton problems (i.e., see if we can get classif users over to SA).
  - Also SA can be used for censoring/truncation
- 

Writing after a global pandemic, applications of survival analysis are more relevant than ever. Predicting the time from onset of COVID-19 symptoms to hospitalisation, or the time from hospitalisation to intubation, or intubation to death, are all time-to-event predictions that are at the centre of survival analysis. As well as morbid applications, survival analysis predictions may be concerned with predicting the time until a customer cancels their gym membership, or the lifetime of a lightbulb; any event that is guaranteed (or at least very likely) to occur can be modelled by a survival analysis prediction. As these predictions can be so sensitive, for example a model predicting when a child should be taken off breathing support (Data Study Group Team 2020), the best possible predictions, evaluated to the highest standard, are a necessity. In other fields of predictive modelling, machine learning has made incredible breakthroughs (such as AlphaFold), therefore applying machine learning to survival analysis is a natural step in the evolution of an important field.

Survival analysis is the field of Statistics focusing on modelling the distribution of an event, which may mean the time until the event takes place, the risk of the event happening, the probability of the event occurring at a single time, or the event's underlying probability distribution. Survival analysis ('survival') is a unique field of study in Statistics as it includes the added difficulty of 'censoring'. Censoring is best described through example: a study is conducted to determine the mortality rate of a group of patients after diagnoses with a particular disease. If a patient dies during this study then their outcome is 'death' and their time of death can be recorded. However if a patient drops-out of the study before they die, then their time of death (though guaranteed to occur) is unknown and the only available information is the time at which they left the study. This patient is now said to be *censored* at the time they drop out. The censoring mechanism allows as much outcome information (time and event) to be captured as possible for all patients (observations).

Machine learning (ML) is the field of Statistics primarily concerned with building models to either predict outputs from inputs or to learn relationships from data (Hastie, Tibshirani, and Friedman 2001; James et al. 2013). This book is limited to the former case, or more

specifically supervised learning, as this is the field in which the vast majority of survival problems live. Relative to other areas of supervised learning, development in survival analysis has been slow – the majority of developments in machine learning for survival analysis have only been in the past decade (see chapters (?@sec-review)-(Chapter 6)). This appears to have resulted in less interest in the development of machine learning survival models (?@sec-review), less rigour in the evaluation of such models (Chapter 6), and fewer off-shelf/open-source implementations (R. Sonabend et al. 2021). This book seeks to set the foundations for clear workflows, good practice, and precise results for ‘machine learning survival analysis’.

---

## 1.1 Why is this book needed?

Firstly, whilst there are many books dedicated to regression and classification as machine learning problems (the ‘bibles’ of machine learning focus entirely on regression and classification only (Bishop 2006; Hastie, Tibshirani, and Friedman 2001; James et al. 2013)), there is a deficit of books covering the survival analysis setting. By writing this book we hope to fill this gap and enable more practitioners to use cutting-edge methods in survival analysis. Survival analysis has important applications in healthcare, finance, engineering and more, all fields that directly impact upon individual lives on a day-to-day basis, and should perhaps be considered as important as classification and regression. The result of this gap in interest, is the erroneous assumption that one field can be directly applied to another. For example there is evidence of researchers treating censoring as a nuisance to be ignored and using regression models instead (Schwarzer, Vach, and Schumacher 2000). Censoring is indeed a challenge and may contribute to making survival analysis less accessible than other fields, but this need not be the case; a clear unification of terminology and presentation of methods may help make ‘machine learning survival analysis’ more accessible. Added accessibility could lead to more academics (and non-academics) engaging with the field and promoting good standards of practice, as well as developing more novel models and measures.

Where survival models have been developed, these have skewed towards ‘ranking models’, which predict the relative risk of an event occurring (?@sec-surv-set-types). In many applications these predictions are sufficient, for example in randomised control trials if assessing the increased/decreased risk of an event after treatment. However, there are many use-cases where predicting an individual’s survival probability distribution is required. Take, for example, an engineer calculating the lifetime of a plane’s engine.<sup>1</sup> There are three important reasons to replace a jet engine at the optimal time:

- financial: jet engines are very expensive and replacing one sooner than required is a waste of money;
- environmental: an engine being replaced too early is a waste of potential usage;
- safety: if the engine is replaced too late then there is a risk to passengers.

Now consider examples for the three possible ‘prediction types’ the engineer can make:

- i. A ‘relative risk prediction’: This engine is twice as likely to fail as another.
- ii. A ‘survival time prediction’: The engine is expected to fail in 30 days.
- iii. A ‘survival distribution prediction’: The lifetime of the engine is distributed according to the probability distribution  $\zeta$ .

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<sup>1</sup>In this engineering context, survival analysis is usually referred to as reliability analysis.

The first prediction type is not useful as the underlying relative risk may be unknown and the engineer is concerned with the individual lifetime. The second prediction type provides a useful quantity for the engineer to work with however there is no uncertainty captured in this prediction. The third prediction type can capture the uncertainty of failure over the entirety of the positive Reals (though usually only a small subset is possible and useful). With this final prediction type, the engineer can create safe decisions: ‘replace the engine at time  $\tau$ , where  $\tau$  is the time when the predicted probability of survival drops below 60%,  $S(\tau) = 0.6$ ’. There are ethical, economic, and environmental reasons for a good survival distribution prediction and this book considers a distribution prediction to be the most important prediction type.

Evaluating predictions from survival models is of the utmost importance. This is especially important as survival models are often deployed in the public domain, particularly in healthcare. Physical products in healthcare, such as new vaccines, undergo rigorous testing and research in randomised control trials before being publically deployed; the same level of rigour should be expected for the evaluation of survival models that are used in life-and-death situations. Evaluation measures for regression and classification are well-understood with important properties, however survival measures have not undergone the same treatment. For example many survival models are still being evaluated solely with concordance indices that have been repeatedly criticised (Gönen and Heller 2005; Rahman et al. 2017; Schmid and Potapov 2012).

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## 1.2 Reproducibility

This book includes simulations and figures generated in R, the code for any figures or experiments in this book are freely available at <https://github.com/mlsa-book/MLSA> under an MIT licence and all content on this website is available under CC BY 4.0.

### Further reading

- (P. Wang, Li, and Reddy 2019) provides a light-touch but comprehensive survey of machine learning models for survival analysis.



## **Part I**

# **Survival Analysis and Machine Learning**



## 2

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# Machine Learning

TODO (150-200 WORDS)

 Minor changes expected!

This page is a work in progress and minor changes will be made over time.

This chapter covers core concepts in machine learning. This is not intended as a comprehensive introduction and does not cover mathematical theory nor how to run machine learning models using software. Instead, the focus is on introducing important concepts and to provide basic intuition for a general machine learning workflow. This includes the concept of a machine learning task, data splitting (resampling), model training and prediction, evaluation, and model comparison. Recommendations for more comprehensive introductions are given at the end of this chapter, including books that cover practical implementation in different programming languages.

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### 2.1 Basic workflow

This book focuses on *supervised learning*, in which predictions are made for outcomes based on data with observed dependent and independent variables. For example, predicting someone's height is a supervised learning problem as data can be collected for features (independent variables) such as age and sex, and an observable outcome (dependent variable), which is height. Alternatives to supervised learning include *unsupervised learning*, *semi-supervised learning*, and *reinforcement learning*. This book is primarily concerned with *predictive survival analysis*, i.e., making future predictions based on (partially) observed survival outcomes, which falls naturally within the supervised learning domain.

The basic machine learning workflow is represented in Figure 2.1. Data is split into training and test datasets. A learner is selected and is trained on the training data, inducing a fitted model. The features from the test data are passed to the model which makes predictions for the unseen outcomes (Box 1). The outcomes from the test data are passed to a chosen measure with the predictions, which evaluates the performance of the model (Box 2). The process of repeating this procedure to test different training and test data is called *resampling* and running multiple resampling experiments with different models is called *benchmarking*. All these concepts will be explained in this chapter.



Figure 2.1: Basic machine learning workflow with data splitting, model training, predicting, and evaluating. Image from Foss and Kotthoff (2024) (CC BY-NC-SA 4.0).

## 2.2 Tasks

A machine learning task is the specification of the mathematical problem that is to be solved by a given algorithm. For example, “predict the height of a male, 13 year old child”, is a machine learning task. Tasks are derived from datasets and one dataset can give rise to many tasks across any machine learning domain. The dataset described by columns: ‘age’, ‘weight’, ‘height’, ‘sex’, ‘diagnosis’, ‘time of death’, ‘clinician notes’, could give rise to any of the following tasks (and more):

- Predict age from weight, height, and sex - supervised regression task
- Predict sex from age and diagnosis - supervised classification task
- Predict time of death from all other features - supervised survival task
- Categorise observations into clusters - unsupervised clustering
- Learn to speak like a clinician depending on client diagnosis - natural language processing, likely with reinforcement learning

As this book is focused on supervised learning, only the first three of these is covered in this chapter and beyond. The specification of a task is vital for interpreting predictions from a model and its subsequent performance. This is particularly true when separating between deterministic and probabilistic predictions, as discussed later in the chapter.

Formally, let  $\mathbf{x} \in \mathcal{X} \subseteq \mathbb{R}^{n \times p}$  be a matrix with  $p$  features for  $n$  observations and let  $y \in \mathcal{Y}$  be a vector of labels (or *outcomes* or *targets*) for all observations. A dataset is then given by  $\mathcal{D} = ((\mathbf{x}_1, y_1), \dots, (\mathbf{x}_n, y_n))$  where it is assumed  $\mathcal{D} \stackrel{i.i.d.}{\sim} (\mathbb{P}_{xy})^n$  for some unknown distribution  $\mathbb{P}$ .

A machine learning task is the problem of learning the unknown function  $f : \mathcal{X} \rightarrow \mathcal{Y}$  where  $\mathcal{Y}$  specifies the nature of the task, for example classification, regression, or survival.

### 2.2.1 Regression

Regression tasks make continuous predictions, for example someone’s height. Regression may be deterministic, in which case a single continuous value is predicted, or probabilistic, where a probability distribution over the Reals is predicted. For example, predicting an individual’s height as 165cm would be a deterministic regression prediction, whereas predicting their

height follows a  $\mathcal{N}(165, 2)$  distribution would be probabilistic.

Formally, a deterministic regression task is specified by  $f_{Rd} : \mathcal{X} \rightarrow \mathcal{Y} \subseteq \mathbb{R}^n$ , and a probabilistic regression task by  $f_{Rp} : \mathcal{X} \rightarrow \mathcal{S}$  where  $\mathcal{S} \subset \text{Distr}(\mathcal{Y})$  and  $\text{Distr}(\mathcal{Y})$  is the space of distributions over  $\mathcal{Y}$ .

In machine learning, deterministic regression is much more common than probabilistic and hence the shorthand ‘regression’ is used to refer to deterministic regression (in contrast to statistical modeling, where regression usually implies probabilistic regression).

### 2.2.2 Classification

Classification tasks make discrete predictions, for example whether it will rain, snow, or be sunny tomorrow. Similarly to regression, predictions may be deterministic or probabilistic. Deterministic classification predicts which category an observation falls into, whereas probabilistic classification predicts the probability of an observation falling into each category. Predicting it will rain tomorrow is a deterministic prediction whereas predicting  $\hat{p}(\text{rain}) = 0.6; \hat{p}(\text{snow}) = 0.1; \hat{p}(\text{sunny}) = 0.3$  is probabilistic.

Formally, a deterministic classification task is given by  $f_{Cd} : \mathcal{X} \rightarrow \mathcal{Y} \subseteq \mathbb{N}_0$ , and a probabilistic classification task as  $f_{Cp} : \mathcal{X} \rightarrow \mathcal{Y} \subseteq [0, 1]^k$  where  $k$  is the number of categories an observation may fall into. Practically this latter prediction is estimation of the probability mass function  $\hat{p}_Y(y) = P(Y = y)$ . If only two categories are possible, these reduce to the *binary classification* tasks:  $f_{Bd} : \mathcal{X} \rightarrow \{0, 1\}$  and  $f_{Bp} : \mathcal{X} \rightarrow [0, 1]$  for deterministic and probabilistic binary classification respectively.

Note that in the probabilistic binary case it is common to write the task as predicting  $[0, 1]$  not  $[0, 1]^2$  as the classes are mutually exclusive. The class for which probabilities are predicted is referred to as the *positive class*, and the other as the *negative class*.

## 2.3 Training and predicting

The terms *algorithm*, *learner*, and *model* are often conflated in machine learning. A *learner* is a description of a learning algorithm, prediction algorithm, parameters, and hyperparameters. The *learning algorithm* is a mathematical strategy to estimate the unknown mapping from features to outcome as represented by a task,  $f : \mathcal{X} \rightarrow \mathcal{Y}$ . During *training*, data,  $\mathcal{D}$ , is fed into the learning algorithm and induces the *model*  $\hat{f}$ . Whereas the learner defines the framework for training and prediction, the model is the specific instantiation of this framework after training on data.

After training the model, new data,  $\mathbf{x}^*$ , can be fed to the *prediction algorithm*, which is a mathematical strategy that uses the model to make predictions  $\hat{\mathbf{y}} = \hat{f}(\mathbf{x}^*)$ . Algorithms can vary from simple linear equations with coefficients to estimate, to complex iterative procedures that differ considerably between training and predicting.

Algorithms usually involve parameters and hyperparameters. Parameters are learned from data whereas hyperparameters are set beforehand to guide the algorithms. Model *parameters* (or *weights*),  $\theta$ , are coefficients to be estimated during model training. Hyperparameters,  $\lambda$ , control how the algorithms are run but are not directly updated by them. Hyperparameters can be mathematical, for example the learning rate in a gradient boosting machine (Chapter 15), or structural, for example the depth of a decision tree (Chapter 13). The number

of hyperparameters usually increases with learner complexity and affects its predictive performance. Often hyperparameters need to be tuned (Section 2.5) instead of manually set. Computationally, storing  $(\hat{\boldsymbol{\theta}}, \lambda)$  is sufficient to recreate any trained model.

**Box 1 (Ridge regression)**

Let  $f : \mathcal{X} \rightarrow \mathcal{Y}$  be the regression task of interest with  $\mathcal{X} \subseteq \mathbb{R}$  and  $\mathcal{Y} \subseteq \mathbb{R}$ . Let  $(\mathbf{x}, \mathbf{y}) = ((x_1, y_1), \dots, (x_n, y_n))$  be data such that  $x_i \in \mathcal{X}$  and  $y_i \in \mathcal{Y}$  for all  $i = 1, \dots, n$ . Say the **learner** of interest is a regularized linear regression model with **learning algorithm**:

$$(\hat{\beta}_0, \hat{\beta}_1) := \arg \min_{\beta_0, \beta_1} \left\{ \sum_{i=1}^n (y_i - (\beta_0 + \beta_1 x_i))^2 + \gamma \beta_1^2 \right\}.$$

and **prediction algorithm**:

$$\hat{f}(\phi) = \hat{\beta}_0 + \hat{\beta}_1 \phi$$

The **hyperparameters** are  $\lambda = (\gamma \in \mathbb{R}_{>0})$  and the **parameters** are  $\boldsymbol{\theta} = (\beta_0, \beta_1)^\top$ . Say that  $\gamma = 2$  is set and the learner is then trained by passing  $(\mathbf{x}, \mathbf{y})$  to the learning algorithm and thus estimating  $\hat{\boldsymbol{\theta}}$  and  $\hat{f}$ . A **prediction**, can then be made by passing new data  $x^* \in \mathcal{X}$  to the fitted model:  $\hat{y} := \hat{f}(x^*) = \hat{\beta}_0 + \hat{\beta}_1 x^*$ .

## 2.4 Evaluating and benchmarking

To understand if a model is ‘good’, its predictions are evaluated with a *loss function*. Loss functions assign a score to the discrepancy between predictions and true values,  $L : \mathcal{Y} \times \mathcal{Y} \rightarrow \bar{\mathbb{R}}$ . Given (unseen) real-world data,  $(\mathbf{X}^*, \mathbf{y}^*)$ , and a trained model,  $\hat{f}$ , the loss is given by  $L(\hat{f}(\mathbf{X}^*), \mathbf{y}^*) = L(\hat{\mathbf{y}}, \mathbf{y}^*)$ . For a model to be useful, it should perform well in general, meaning its generalization error should be low. The *generalization error* refers to the model’s performance on new data, rather than just the data encountered during training and development.

A model should only be used to make predictions if its generalization error was estimated to be acceptable for a given context. If a model were to be trained and evaluated on the same data, the resulting loss, known as the *training error*, would be an overoptimistic estimate of the true generalization error (James et al. 2013). This occurs as the model is making predictions for data it has already ‘seen’ and the loss is therefore not evaluating the model’s ability to generalize to new, unseen data. Estimation of the generalization error requires *data splitting*, which is the process of splitting available data,  $\mathcal{D}$ , into *training data*,  $\mathcal{D}_{train} \subset \mathcal{D}$ , and *testing data*,  $\mathcal{D}_{test} = \mathcal{D} \setminus \mathcal{D}_{train}$ .

The simplest method to estimate the generalization error is to use *holdout resampling*, which is the process of partitioning the data into one training dataset and one testing dataset, with the model trained on the former and predictions made for the latter. Using 2/3 of the data for training and 1/3 for testing is a common splitting ratio (Kohavi 1995). For independent and identically distributed (iid) data, it is generally best practice to partition the data randomly. This ensures that any potential patterns or information encoded in the ordering of the data are removed, as such patterns are unlikely to generalize to new, unseen

data. For example, in clinical datasets, the order in which patients enter a study might inadvertently encode latent information such as which doctor was on duty at the time, which could theoretically influence patient outcomes. As this information is not explicitly captured in measured features, it is unlikely to hold predictive value for future patients. Random splitting breaks any spurious associations between the order of data and the outcomes.

When data is not iid, for example spatially correlated or time-series data, then random splitting may not be advisable, see Hornung et al. (2023) for an overview of evaluation strategies in non-standard settings.

Holdout resampling is a quick method to estimate the generalization error, and is particular useful when very large datasets are available. However, hold-out resampling has a very high variance for small datasets and there is no guarantee that evaluating the model on one hold-out split is indicative of real-world performance.

*k*-fold cross-validation (CV) can be used as a more robust method to better estimate the generalization error (Hastie, Tibshirani, and Friedman 2001). *k*-fold CV partitions the data into *k* subsets, called *folds*. The training data comprises of  $k - 1$  of the folds and the remaining one is used for testing and evaluation. This is repeated *k* times until each of the folds has been used exactly once as the testing data. The performance from each fold is averaged into a final performance estimate (Figure 2.2). It is common to use  $k = 5$  or  $k = 10$  (Leo Breiman and Spector 1992; Kohavi 1995). This process can be repeated multiple times (*repeated k-fold CV*) and/or *k* can even be set to *n*, which is known as *leave-one-out cross-validation*.

Cross-validation can also be stratified, which ensures that a variable of interest will have the same distribution in each fold as in the original data. This is important, and often recommended, in survival analysis to ensure that the proportion of censoring in each fold is representative of the full dataset (Casalicchio and Burk 2024; Herrmann et al. 2021).

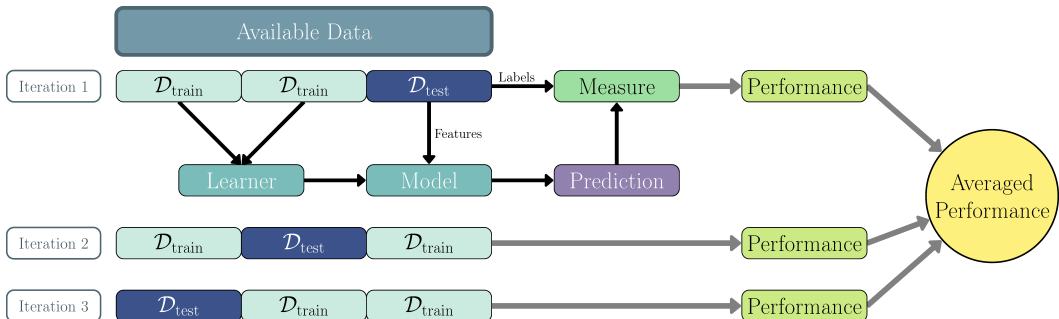


Figure 2.2: Three-fold cross-validation. In each iteration a different dataset is used for predictions and the other two for training. The performance from each iteration is averaged into a final, single metric. Image from Casalicchio and Burk (2024) (CC BY-NC-SA 4.0).

Repeating resampling experiments with multiple models is referred to as a *benchmark experiment*. A benchmark experiment compares models by evaluating their performance on *identical* data, which means the same resampling strategy and folds should be used for all models. Determining if one model is actually better than another is a surprisingly complex topic (Benavoli et al. 2017; Demšar 2006; Dietterich 1998; Nadeau and Bengio 2003) and is out of scope for this book, instead any benchmark experiments performed in this book are purely for illustrative reasons and no results are expected to generalize outside of these experiments.

### Box 2 (Evaluating ridge regression)

Let  $\mathcal{X} \subseteq \mathbb{R}$  and  $\mathcal{Y} \subseteq \mathbb{R}$  and let  $(\mathbf{x}^*, \mathbf{y}^*) = ((x_1^*, y_1^*), \dots, (x_m^*, y_m^*))$  be data previously unseen by the model trained in Box 1 where  $x_i \in \mathcal{X}$  and  $y_i \in \mathcal{Y}$  for all  $i = 1, \dots, m$ . Predictions are made by passing  $\mathbf{x}^*$  to the fitted model yielding  $\hat{\mathbf{y}} = (\hat{y}_1, \dots, \hat{y}_m)$  where  $\hat{y}_i := \hat{f}(x_i^*) = \hat{\beta}_0 + \hat{\beta}_1 x_i^*$ .

Say the mean absolute error is used to evaluate the model, defined by

$$L(\phi, \varphi) = \frac{1}{n} \sum_{i=1}^n |\phi_i - \varphi_i|$$

where  $(\phi, \varphi) = ((\phi_1, \varphi_1), \dots, (\phi_n, \varphi_n))$ .

The model's predictive performance is then calculated as  $L(\hat{\mathbf{y}}, \mathbf{y}^*)$ .

## 2.5 Hyperparameter Optimization

Section 2.3 introduced model hyperparameters, which control how training and prediction algorithms are run. Setting hyperparameters is a critical part of model fitting and can significantly change model performance. *Tuning* is the process of using internal benchmark experiments to automatically select the optimal hyper-parameter configuration. For example, the depth of trees,  $m_r$  in a random forest (Chapter 13) is a potential hyperparameter to tune. This hyperparameter may be tuned over a range of values, say [1, 15] or over a discrete subset, say {1, 5, 15}, for now assume the latter. Three random forests with 1, 5, and 15 tree depth respectively are compared in a benchmark experiment. The depth that results in the model with the optimal performance is then selected for the hyperparameter value going forward. *Nested resampling* is a common method to reduce bias that could occur from using overlapping data for tuning, training, or testing (Simon 2007). Nested resampling is the process of resampling the training set again for tuning and then the optimal model is refit on the entire training data (Figure 2.3).

## 2.6 Conclusion

### Key takeaways

- Machine learning tasks define the predictive problem of interest;
- Regression tasks make predictions for continuous outcomes, such as the amount of rain tomorrow;
- Classification tasks make predictions for discrete outcomes, such as the predicted weather tomorrow;
- Both regression and classification tasks may make deterministic predictions (a single number or category), or probabilistic predictions (the probability of a number or category);
- Models have parameters that are fit during training and hyperparameters that are



Figure 2.3: An illustration of nested resampling. The large blocks represent three-fold CV for the outer resampling for model evaluation and the small blocks represent four-fold CV for the inner resampling for hyperparameter optimization. The light blue blocks are the training sets and the dark blue blocks are the test sets. Image and caption from Becker, Schneider, and Fischer (2024) (CC BY-NC-SA 4.0).

- set or tuned;
- Models should be evaluated on resampled data to estimate the generalization error to understand future performance.

#### Further reading

- *The Elements of Statistical Learning* (Hastie, Tibshirani, and Friedman 2001), *An Introduction to Statistical Learning* (James et al. 2013), and *Pattern Recognition and Machine Learning* (Bishop 2006) for comprehensive introductions and overviews to machine learning.
- *Applied Machine Learning Using mlr3 in R* (Bischl et al. 2024) and *Tidy Modeling* (Kuhn and Silge 2023) for machine learning in R
- *Hands-on Machine Learning with Scikit-Learn, Keras, and TensorFlow* (Géron 2019) for machine learning in Python.
- Bischl et al. (2012) for discussions about more resampling strategies including bootstrapping and subsampling.

# 3

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## Survival Analysis

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TODO (150-200 WORDS)

! Major changes expected!

This page is a work in progress and major changes will be made over time.

*Survival Analysis* is concerned with data where the outcome is the time until an event takes place (a ‘time-to-event’). Because the collection of such data takes place in the temporal domain (it takes time to observe a duration), the event of interest is often unobservable, for example because it did not occur by the end of the data collection period. In survival analysis terminology this is referred to as *censoring*.

This chapter defines basic terminology and mathematical definitions in survival analysis, which are used throughout this book. Building upon this chapter, Chapter 4 introduces event-history analysis, which is a generalisation to settings with multiple, potentially competing or recurrent events, including multi-state outcomes. Concluding this part of the book, Chapter 5 defines different prediction tasks in survival analysis that are used by models and measures to implement machine learning methods.

While these definitions and concepts are not new to survival analysis, it is imperative they are understood to build successful models. Evaluation functions (Part II) can identify if one model is better suited than another to minimize a given objective function, however they cannot identify if the objective function itself was specified correctly, which depends on the assumptions about the data generating process. Evaluating models with the wrong objective function yields meaningless results. Hence, it is of utmost importance for machine learning practitioners to be able identify and specify the survival problem present in their data correctly to ensure models are correctly fit and evaluated.

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### 3.1 Quantifying the Distribution of Event Times

This section introduces functions that can be used to fully characterise a probability distribution, particular focus is given to functions that are important in survival analysis.

Note that we can generally distinguish between event taking place in discrete time or continuous time. For example, consider the time a politician serves in parliament. If we consider the number of election cycles they stay in parliament, it would constitute discrete time, as time can only take values,  $1, 2, 3, \dots$ , that is  $Y \in \mathbb{N}_{>0}$ . On the other hand, the time

an individual stays in hospital is usually determined as the difference between the admission date-time and discharge date-time, which would constitute a continuous time  $Y \in \mathbb{R}_{\geq 0}$ .

In practice the differences are often blurred as time-measurement will naturally be discretized at some level and precision beyond some resolution is often not of interest (hospital length of stay might be interesting up to days or hours, but not minutes and seconds). Also discrete-time methods are often applied to continuous time data and vice versa. It is nevertheless important to make the distinction as it informs mathematical treatment and definition of the different quantities introduced below.

### 3.1.1 Continuous Time

For now, assume a continuous, positive, random variable  $Y$  taking values in (t.v.i.)  $\mathbb{R}_{\geq 0}$ . A standard representation of the distribution of  $Y$  is given by the probability density function (pdf),  $f_Y : \mathbb{R}_{\geq 0} \rightarrow \mathbb{R}_{\geq 0}$ , and cumulative distribution function (cdf),  $F_Y : \mathbb{R}_{\geq 0} \rightarrow [0, 1]; (\tau) \mapsto P(Y \leq \tau)$ .

In survival analysis, it is most common to describe the distribution of event times  $Y$  via the *survival function* and *hazard function* (or *hazard rate*) rather than the pdf or cdf. The survival function is defined as

$$S_Y(\tau) = P(Y > \tau) = \int_{\tau}^{\infty} f_Y(u) \, du, \quad (3.1)$$

which is the probability that an event has not occurred by  $\tau \geq 0$  and thus the complement of the cdf:  $S_Y(\tau) = 1 - F_Y(\tau)$ . By definition,  $S_Y(0) = 1$  and  $S(\tau) \rightarrow 0$  for  $\tau \rightarrow \infty$ .

The hazard function is given by

$$\begin{aligned} h_Y(\tau) &= \lim_{d\tau \searrow 0} \frac{P(\tau \leq Y < \tau + d\tau | Y \geq \tau)}{d\tau} \\ &= \lim_{d\tau \searrow 0} \frac{P(Y \in [\tau, \tau + d\tau] | Y \geq \tau)}{d\tau} \\ &= \frac{f_Y(\tau)}{S_Y(\tau)} \end{aligned} \quad (3.2)$$

where  $d\tau$  denotes a time-interval. The hazard rate is often interpreted as the instantaneous risk of observing an event at  $\tau$ , given that the event has not been observed before  $\tau$ . This is not a probability and  $h_Y$  can be greater than one.

The cumulative hazard function (chf) can be derived from the hazard function by

$$H_Y(\tau) = \int_0^{\tau} h_Y(u) \, du, \quad (3.3)$$

and relates to the survival function via

$$H_Y(\tau) = \int_0^{\tau} h_Y(u) \, du = \int_0^{\tau} \frac{f_Y(u)}{S_Y(u)} \, du = \int_0^{\tau} -\frac{S'_Y(u)}{S_Y(u)} \, du = -\log(S_Y(\tau))$$

These last relationships are particularly important, as many methods estimate the hazard rate, which is then used to calculate the cumulative hazard and survival probability

$$S_Y(\tau) = \exp(-H_Y(\tau)) = \exp\left(-\int_0^{\tau} h_Y(u) \, du\right) \quad (3.4)$$

Unless necessary to avoid confusion, subscripts are dropped from  $S_Y, h_Y$  etc. going forward and instead these functions are referred to as  $S, h$  (and so on).

Usual regression techniques cannot be used to estimate these quantities as  $Y$  is only partially observed, due to different types of censoring and truncation, which are now described.

### 3.1.2 Discrete Time

Now consider a discrete, positive random variable  $\tilde{Y}$  taking value in  $\mathbb{N}_{>0}$  and  $\tau \in \mathbb{N}_{>0}$  some time point in discrete time.

The discrete-time hazard rate

$$h_{\tilde{Y}}^d(\tau) = P(\tilde{Y} = \tau | \tilde{Y} \geq \tau). \quad (3.5)$$

Thus, in contrast to the continuous time hazard Equation 3.2, the discrete time hazard is an actual (conditional) probability, rather than a rate and therefore might be easier to interpret.

The cumulative discrete time hazard is given by

$$H_{\tilde{Y}}^d(\tau) = \sum_{k=1}^{\tau} h_{\tilde{Y}}^d(k) \quad (3.6)$$

We then define the inverse probability

$$s_{\tilde{Y}}^d(\tau) := 1 - h_{\tilde{Y}}^d(\tau) = P(\tilde{Y} > \tau | \tilde{Y} \geq \tau).$$

It follows that the probability to survive beyond time point  $\tau$  is given by

$$S_{\tilde{Y}}^d = P(\tilde{Y} > \tau) = \prod_{k \leq \tau} s_{\tilde{Y}}^d(\tau) = \prod_{k \leq \tau} (1 - h_{\tilde{Y}}^d(\tau)), \quad (3.7)$$

and the unconditional probability for an event at time  $\tau$  is

$$P(\tilde{Y} = \tau) = S_{\tilde{Y}}^d(\tau - 1) h_{\tilde{Y}}^d(\tau). \quad (3.8)$$

When applied to continuous time  $Y$ , the follow-up is divided in  $J$  disjunct intervals  $(a_0, a_1], \dots, (a_{j-1}, a_j], \dots, (a_{J-1}, a_J]$ ,  $j = 1, \dots, J$  such that

$$Y \in (a_{j-1}, a_j] \Leftrightarrow \tilde{Y} = j$$

Thus,

$$h_{\tilde{Y}}^d(j) = P(Y \in (a_{j-1}, a_j] | Y > a_{j-1})$$

and

$$S_{\tilde{Y}}^d(j) = P(Y > a_j) = S_Y(a_j)$$

For more details on discrete time-to-event analysis consider (Tutz and Schmid 2016).

### 3.2 Single-event, right-censored data

The complexity of Survival Analysis compared to other fields arises from the fact that the outcome of interest is often only observed partially. In particular, the time-to-event is often unknown at the end of the observation period, as the event has not occurred yet.

Let,

- $X$  taking values in  $\mathbb{R}^p$  be the generative random variable representing the data *features/covariates/independent variables*.
- $Y$  taking values in  $\mathbb{R}_{\geq 0}$  be the (partially unobservable) *true survival time*.
- $C$  taking values in  $\mathbb{R}_{\geq 0}$  be the (partially unobservable) *true censoring time*.

In the presence of censoring  $C$ , it is impossible to fully observe the true outcome of interest,  $Y$ . Instead, the observable variables are defined by

- $T := \min\{Y, C\}$ , the *outcome time* (realisations are referred to as the *observed outcome time*); and
- $\Delta := \mathbb{I}(Y = T) = \mathbb{I}(Y \leq C)$ , the *event indicator* (also known as the *censoring or status indicator*).

Together  $(T, \Delta)$  is referred to as the *survival outcome* or *survival tuple* and they form the dependent variables. The survival outcome provides a concise mechanism for representing the outcome time and indicating which outcome (event or censoring) took place.

A *survival dataset* is a  $n \times p$  real-valued matrix defined by  $\mathcal{D} = ((\mathbf{x}_1, t_1, \delta_1) \cdots (\mathbf{x}_n, t_n, \delta_n))^{\top}$ , where  $(t_i, \delta_i)$  are realisations of the respective random variables  $(T_i, \Delta_i)$  and  $\mathbf{x}_i$  is a  $p$ -dimensional vector,  $\mathbf{x}_i = (x_{i;1} \ x_{i;2} \ \cdots \ x_{i;p})^{\top}$  of features.

Finally, the following quantities are used frequently throughout this book and survival analysis literature more generally. Let  $(t_i, \delta_i) \stackrel{i.i.d.}{\sim} (T, \Delta)$ ,  $i = 1, \dots, n$ , be observed survival outcomes. Then,

The **set of unique outcome times** is the set of time-points in which at least one observation experiences the event or is censored:

$$\mathcal{U}_O \subseteq \{t_i\}_{i \in \{1, \dots, n\}}$$

The **set of unique event times** is the set of time-points in which at least one observation experiences the event (but not censored):

$$\mathcal{U}_D \subseteq \{t_i : \delta_i = 1\}_{i \in \{1, \dots, n\}}$$

The **ordered, unique events times** may also be denoted by

$$t_{(k)}, \quad k = 1, \dots, m \quad t_{(1)} < t_{(2)} < \cdots < t_{(m)}, \quad m \leq n$$

The **risk set at  $\tau$** , is the index-set of observation units at risk for the event just before  $\tau$

$$\mathcal{R}_{\tau} := \{i : t_i \geq \tau\}$$

where  $i$  is the index of an observation in the data. For right-censored data,  $\mathcal{R}_0 = \{1, \dots, n\}$  and  $\mathcal{R}_\tau \subseteq \mathcal{R}_{\tau'}, \forall \tau > \tau'$ . Note that in a continuous setting, ‘just before’ refers to an infinitesimally smaller time than  $\tau$ , in practice as this is unobservable the risk set is defined at  $\tau$ , hence an observation may both be at risk, and experience an event (or be censored) at  $\tau$ .

The **number of observations at risk at  $\tau$**  is the cardinality of the risk set at  $\tau$ ,

$$n_\tau := \sum_i \mathbb{I}(t_i \geq \tau) = |\mathcal{R}_\tau|$$

Finally, the **number of events at  $\tau$**  is defined by,

$$d_\tau := \sum_i \mathbb{I}(t_i = \tau, \delta_i = 1)$$

For truly continuous variables, one might expect only one event to occur at each observed event time:  $d_{t_i} = 1, \forall i$ . In practice, ties are often observed due to finite measurement precision, such that  $d_\tau > 1$  occurs frequently in real-world datasets.

The quantities  $\mathcal{R}_\tau$ ,  $n_\tau$ , and  $d_\tau$  underlie many models and measures in survival analysis. Several non-parametric and semi-parametric methods (Chapter 12) like the Kaplan-Meier estimator (see Section 3.3) are based on the ratio  $d_\tau/n_\tau$ .

Table 3.1 exemplifies an observed survival dataset, a subset of the `tumor` data (Bender and Scheipl 2018), which contains the time until death in days after operation ( $\delta_i = 1$  if death occurred at the outcome time  $t_i$  and  $\delta_i = 0$  otherwise).

In this example, the above quantities would be:

- $\mathcal{U}_0 = \{268, 397, 519, 1217, 2414\}$ : with 1217 included only once
- $\mathcal{U}_D = \{268, 397, 1217\}$ : with the inclusion of 1217 due to the event at  $t_1$ , not censoring at  $t_5$
- $\mathcal{R}_{\tau=1217} = \{1, 3, 5\}$  (these subjects’ outcome times are greater or equal to  $\tau = 1217$  so they are at risk for the event at this time)
- $n_{\tau=1217} = |\mathcal{R}_{1217}| = 3$
- $d_{\tau=1217} = 1$ : As only  $i = 1$  experienced the event (and not censoring) at this time.

Table 3.1: Subset of the `tumor` (Bender and Scheipl 2018) time-to-event dataset. Rows are individual observations (ID),  $\mathbf{x}_{:j}$  columns are features,  $t$  is observed time-to-event,  $\delta$  is the event indicator.

<b>id (i)</b>	<b>age (<math>\mathbf{x}_{:1}</math>)</b>	<b>sex (x<sub>:2</sub>)</b>	<b>complications (x<sub>:3</sub>)</b>	<b>days (t)</b>	<b>status (<math>\delta</math>)</b>
1	71	female	no	1217	1
2	70	male	no	519	0
3	67	female	yes	2414	0
4	58	male	no	397	1
5	39	female	yes	1217	0
6	59	female	no	268	1

### 3.3 Kaplan-Meier estimator

Before we go further, we introduce a simple, non-parametric estimator for the survival function (Equation 3.1), the Kaplan-Meier estimator (Kaplan and Meier 1958). The estimator is useful for visualising survival data and is a popular baseline model to compare to the predictive performance of more complex methods. In machine learning terms it can be viewed as a “featureless” learner for survival analysis.

Using the quantities introduced in Section 3.2, the estimator is defined by:

$$\hat{S}_{KM}(\tau) = \prod_{k:t_{(k)} \leq \tau} \left(1 - \frac{d_{t_{(k)}}}{n_{t_{(k)}}}\right) \quad (3.9)$$

which is a step-function at the observed ordered event times  $t_{(k)}$ ,  $k = 1, \dots, m$  with  $\hat{S}_{KM}(\tau) = 1 \forall \tau < t_{(1)}$ . It is usually estimated for all unique event times.

For illustration, Figure 3.1 shows the estimated survival probability obtained by applying the Kaplan-Meier estimator to the full `tumor` data, containing observations of  $n = 776$  subjects. By definition, the survival function starts at  $S(t) = 1$  at  $t = 0$  and monotonically decreases towards  $S(t) = 0$  for  $t \rightarrow \infty$ . Dotted lines indicated the median survival, defined as the time at which the survival function reaches  $S(t) = 0.5$ . In this example, the median survival time is approximately 1500 days, as indicated by the dashed lines. This means that 50% of the subjects are expected die within 1500 days after operation.

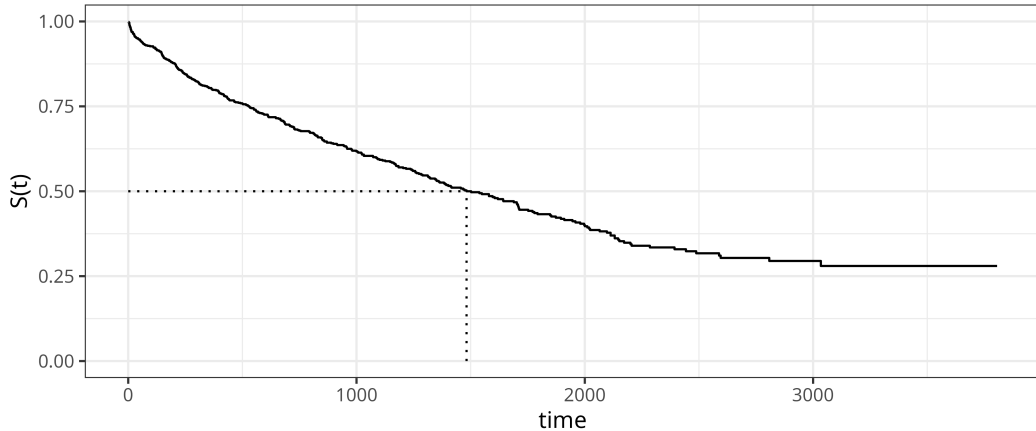


Figure 3.1: Kaplan-Meier estimate for the `tumor` data (Bender and Scheipl 2018). The estimated survival probabilities are given by the solid step-function. Dotted lines indicate the median survival time).

While the Kaplan-Meier estimator does not directly support estimation of covariate effects on the estimated survival probabilities, it is often used for descriptive analysis by applying the estimator to different subgroups (referred to as stratification in survival analysis). For example, Figure 3.2 shows  $\hat{S}_{KM}$  separately for subjects aged 50 years or older and subjects younger than 50 years, respectively. Dashed lines again illustrate median survival times.

However, note that the median survival time does not exist for the younger age group, as their estimated survival function does not cross 0.5.

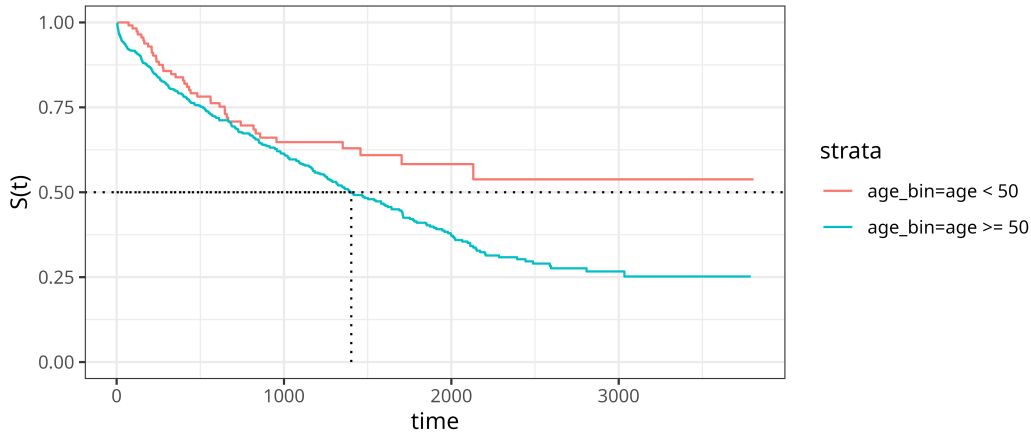


Figure 3.2: Kaplan-Meier estimate for the `tumor` data (Bender and Scheipl 2018) applied to subgroups of subjects of 50 or more years old and less than 50 years old, respectively. The estimated survival probabilities are given by the solid step-function. Dotted lines indicate the median survival time).

Sometimes the KM Estimator is also used to estimate the distribution of censoring times, for example to calculate inverse probability of censoring weights (Section 7.1.1). Throughout this book, we denote the Kaplan-Meier estimator applied to the observed censoring times  $(t_i, 1 - \delta_i)$  as  $\hat{G}_{KM}$ .

### 3.4 Types of Censoring

Three types of censoring are commonly defined in survival analysis: right-censoring, left-censoring, and interval-censoring. The latter can be viewed as the most general case. Multiple types of censoring and/or truncation (Section 3.5) can occur in any given data set and it is vital to identify which types are present in order to correctly select and specify models and measures for the data.

#### Right-censoring

Right-censoring is the most commonly assumed form of censoring in survival data. It occurs when the event of interest was not experienced during the observation period, which may happen because it was no longer observable (for example, due to withdrawal from the study) or because the event did not happen until study end. The exact event time is unknown but it is known that the event is after the observed censoring time, hence *right-censoring* (imagine a timeline from left to right as in Figure 3.3).

Right-censoring can be further divided into Type-I, Type-II and random censoring. Type-I, or *administrative*, censoring occurs at the fixed, pre-defined end of an observation period  $\tau_u$ , in which case the outcome is given by  $(T_i = \min(Y_i, \tau_u), \Delta_i = \mathbb{I}(Y_i \leq \tau_u))$ . Censored

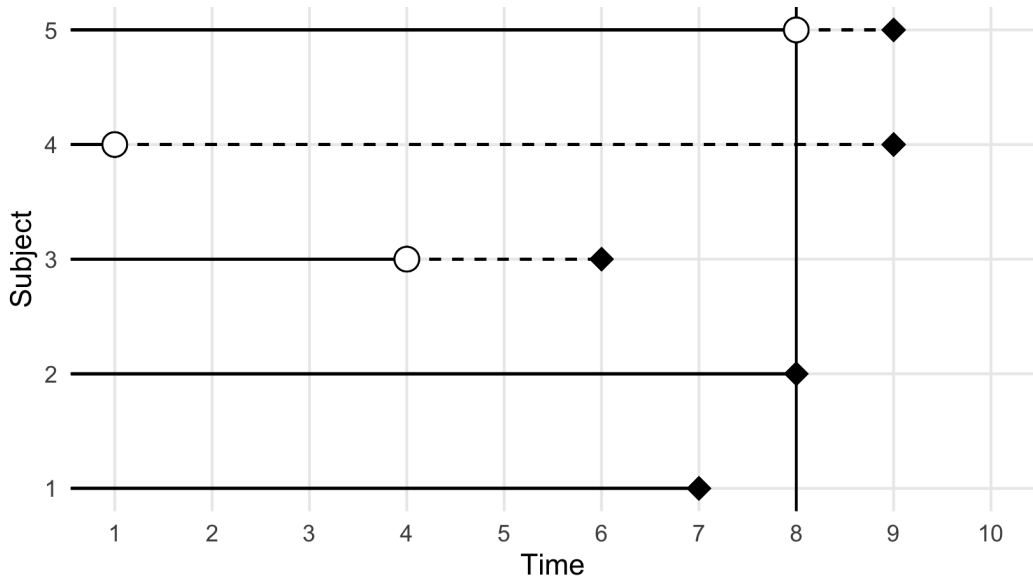


Figure 3.3: Dead and censored subjects (y-axis) over time (x-axis). Black diamonds indicate true death times and white circles indicate censoring times. Diamonds at the end of dashed lines are hypothetical (unknown in reality). Vertical line is the study end time. Subjects 1 and 2 die in the study time. Subject 3 is censored in the study and (unknown) dies within the study time. Subject 4 is censored in the study and (unknown) dies after the study. Subject 5 is censored at study end and (unknown) dies after the end of the study.

observations are therefore represented as  $(\tau_u, 0)$ . Type-II censoring also occurs when the observation period ends. However, in this case the study ends when a pre-defined number of subjects experienced the event of interest and hence  $\tau_u$  is random.

Random censoring occurs when censoring times *randomly* follow an unknown distribution and one observes  $(T_i = \min(Y_i, C_i), \Delta_i = \mathbb{I}(Y_i \leq C_i))$ . Different types of right-censoring can, and sometimes do, co-occur in any given data set.

In practice, these different types of right-censoring are usually handled the same during modeling and evaluation and so this book refers to ‘right-censoring’ generally, which could occur from a combination of the above types.

### Left- and Interval-censoring

Left-censoring occurs when the event is known to have happened at some unknown time before observation time, interval-censoring occurs when the event is known to have happened within some time span, but not the exact time.

Consider a survey about phone use where participants are asked: “How old were you when you used a smart phone for the first time?”. The possible answers are:

- exact age of first use
- didn’t use a smart phone yet
- did or does use a phone but doesn’t remember age of first time use
- did or does use a phone, remembers a specific age range

The first case represents an exactly observed event time, the second case the familiar right-censoring, as the event may occur later in life, but it is unknown when. The third case is referred to as left-censoring, we know the event occurred before the interview, but don’t know when. The fourth case is an example of interval-censoring, as we know the event occurred within some age span, but not the exact age.

Interval- and left-censoring also often occurs in medical contexts. For example, some guidelines suggest annual screenings for skin cancer (starting from a certain age). However, the initial age at which individuals do screenings and regularity of check ups varies widely. If cancer was detected between two screenings, the observation is interval-censored. If cancer is detected at first screening, the observation is left-censored (unless the ‘age’ of the cancer can be narrowed down based on size and other characteristics, in which case it would become interval-censored).

### Censoring Notation

In the presence of left- or interval-censoring the usual representation of survival outcomes as  $(t_i, \delta_i)$  is not sufficient to denote the different types of observations in the data set. Instead, we represent the data as intervals in which the event occurs. Formally, let  $Y_i$  the random variable for time until the event of interest and  $L_i, R_i$  random variables that define an interval  $(L_i, R_i]$  with realisations  $l_i, r_i$ . Let further  $t_i$  the time of observation (for example age at interview in the phone use example) or last-follow up time for subject  $i$ . Then the event time of subject  $i$  is

- left-censored if  $Y_i \in (L_i = 0, R_i = t_i]$ ;
- right-censored at  $t_i$  if  $Y_i \in (L_i = t_i, R_i = \infty)$ ;
- interval-censored  $Y_i \in (L_i = l_i, R_i = r_i], l_i < r_i \leq t_i$
- exactly observed if  $Y_i \in (L_i = t_i, R_i = t_i]$

In the cancer screening example from above it holds that  $r_i = t_i$  and  $l_i$  the last check-up before  $r_i$ . In the phone use example, the participants might specify any age range between 0 and  $t_i$ .

To make this more concrete, consider phone use example data, where  $t_i$  is the age at interview. In practice, such data is often stored by creating two variables representing the left ( $l_i$ ) and right border ( $r_i$ ) of the respective intervals ( $t_i$  is not really needed here to define the outcome, but included for illustration).

id	$t_i$	$l_i$	$r_i$
1	13	13	$\infty$
2	17	15	15
3	16	14	16
4	16	13	15
5	18	0	18

Here, the first subject is right-censored at 13 years ( $l_i = t_i = 13, r_i = \infty$ ), the second subject remembered exactly ( $l_i = r_i = 15 < t_i = 17$ ), the third subject remembers that it was after 14, but not exact age ( $l_i = 14 < r_i = 16 = t_i$ ), fourth subject remembers use after 13 years and latest at 15 years of age ( $l_i = 13 < r_i = 15 < t_i$ ), and the fifth subject uses a smart phone currently at age 18, but cannot specify further ( $l_i = 0 < r_i = 18 = t_i$ ).

From the example above, it is clear, that right- and left-censoring are special cases of interval-censoring. However, if only right- or left-censoring is present, the likelihood and estimation simplifies (see Section 3.6). Also note that in case of left- and interval-censoring the event is known to have occurred, while for right-censoring the event didn't occur during time under observation. For left- and right-censoring one might be tempted to consider it an event  $\delta_i = 1$  without exact time, while right-censoring would be consider a non-event  $\delta_i = 0$ . However, technically it is assumed that the event will always occur, if we wait long enough (for right-censored data in the interval  $(t_i, \infty)$ ). Censoring therefore means having imprecise information about the time of event rather than information about the event occurring or not occurring.

## Dependent vs. Informative Censoring

Censoring may be defined as *uninformative* if  $Y \perp\!\!\!\perp C$  and *informative* otherwise. However, these definitions can be misleading as the term ‘uninformative’ could imply that  $C$  is independent of both  $X$  and  $Y$ , and not just  $Y$ . To avoid misinterpretation, the following definitions are used in this book:

- If  $C \perp\!\!\!\perp X$ , censoring is *feature-independent*, otherwise censoring is *feature-dependent*.
- If  $C \perp\!\!\!\perp Y$ , censoring is *event-independent*, otherwise censoring is *event-dependent*.
- If  $(C \perp\!\!\!\perp Y)|X$ , censoring is conditionally independent of the event given covariates, or *conditionally event-independent*.
- If  $C \perp\!\!\!\perp (X, Y)$ , censoring is *uninformative*, otherwise censoring is *informative*.

Uninformative censoring can generally be well-handled by models as the true underlying distribution of survival times is not affected by censoring. In fact, in this case one could

even use regression models after removing censored observations (if they do not form a high proportion of the data).

In reality, censoring is rarely non-informative as reasons for drop-out or missingness in outcomes tend to be related to the study of interest. Event-dependent censoring is a tricky case that, if not handled appropriately (by a competing-risks framework), can easily lead to poor model development. Imagine a study interested in predicting the time between relapses of stroke but a patient suffers a brain aneurysm due to some separate neurological condition. There is a high possibility that a stroke may have occurred if the aneurysm had not. A survival model is unlikely to distinguish the censoring event (aneurysm) from the event of interest (stroke) and will confuse predictions.

In practice, the majority of models and measures assume that censoring is conditionally event-independent and hence censoring patterns can be predicted/estimated based on the covariates. For example, if studying the survival time of ill pregnant patients in hospital, then dropping out of the study due to pregnancy is clearly dependent on how many weeks pregnant the patient is when the study starts (for the sake of argument assume no early/late pregnancy due to illness).

### 3.5 Censoring vs. Truncation

A common confusion is to conflate censoring and truncation, which is problematic as the methods to handle them differ substantially. Outside of time-to-event settings, truncation usually refers to truncating (or removing) an entire subject from a dataset. As discussed in Chapter 1, truncation in survival analysis refers to partially truncating a period of time and is quite common in the more general event history setting (Chapter 4).

While censored observations have incomplete information about the time-to-event, they are still part of the data set. Whereas truncation leads to observations not entering the data set (at least not at time 0). This will usually introduce bias that needs to be accounted for.

#### Left-truncation

Left-truncation often occurs when inclusion into a study is conditional on the occurrence of another event. Left-truncation plays an important role when modeling recurrent events or multi-state data (Chapter 4), thus the concept will be introduced in more detail.

By example, consider a study from the 18th century (Broström 1987), when childhood and maternal mortality were relatively high. The goal of the study was to establish the effect of a mother's death on the survival of the infant. Since each death was reported to the authorities, an infant was added to the study if and when their mother died. To create a matched cohort, two other infants, whose mothers were alive, were matched into the study based on their age and other relevant features. Thus, groups of three infants within the study had identical features except for the status of the mother (alive or dead). Because of the study design, infants who died before their mothers could never enter into the study. A mother's death is thus referred to as left-truncation event and the infant's age at time of inclusion into the study is referred to as left-truncation time.

More formally, let  $t_i^L$  the subject-specific left-truncation time. Then we only observe subjects with  $y_i > t_i^L$  and subjects with  $y_i < t_i^L$  never enter the data.

This is illustrated in Figure 3.4. Continuing with the example above, say Infant 1 dies at  $t_1$ , while the mother dies at some later time point  $t_1^L$ , therefore infant 1 never enters the study. The mother of Infant 2 dies at  $t_2^L$ , at which point the infant is included in the study and experiences an event at  $t_2$ . Finally, say Infant 3 enters the study at  $t_3^L$  and is censored at 365 days when the study ends.

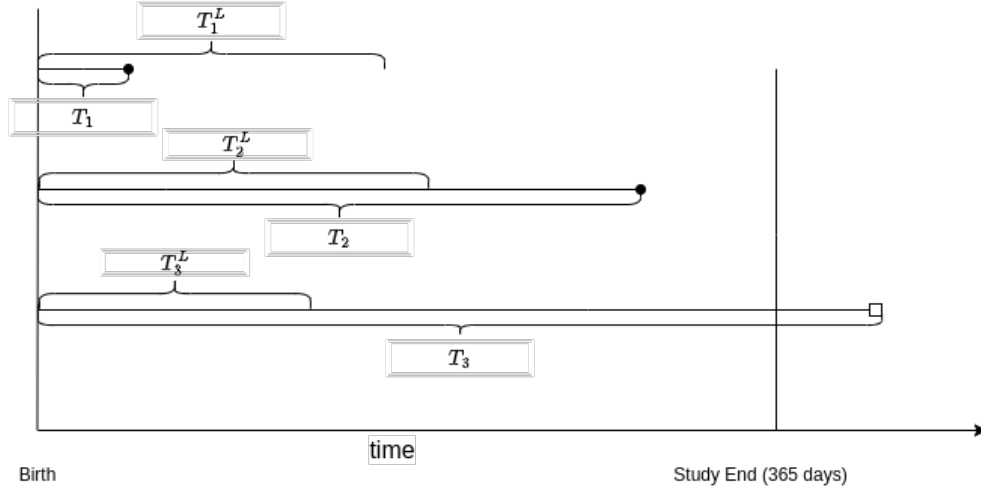


Figure 3.4: Illustration of left-truncation data. Subjects 2 and 3 enter the data set as the study entry condition (mother's death) occurs before the event of interest occurs after the left-truncation time (left-truncation time shorter than event time). Subject 1 on the other hand never enters the data as the event occurs before the study entry condition (left-truncation time longer than event time).

In this example, left-truncation biases the sample towards healthier or more robust infants, as frail infants on die earlier and thus on average before their mothers. This would bias the estimates if not properly taken into account.

Continuing the above example, Figure 3.5 shows the difference between estimated survival probabilities when left-truncation is ignored (left panel) and taken into account (right panel), respectively. It is clear that the survival probabilities were underestimated in both groups, but more so in the group of infants whose mother died (thereby underestimating the effect of the mothers' death on infant survival).

To understand this, consider an excerpt from the infants data in Table 3.3.

Table 3.3: Excerpt of the `infants` (Broström 2024) time-to-event dataset. Rows are individual observations (`id`), `group` indicates matched infants,  $t^L$  is the left-truncation time (time of inclusion into the study),  $t$  is the observed time,  $\delta$  is the event indicator.

group	id	$t^L$	$t$	$\delta$	mother
1	1	55	365	0	dead
1	2	55	365	0	alive
1	3	55	365	0	alive
2	4	13	76	1	dead
2	5	13	365	0	alive

group	id	$t^L$	$t$	$\delta$	mother
2	6	13	365	0	alive
4	7	2	16	1	dead
4	8	2	365	0	alive
4	9	2	365	0	alive

Now recall the calculation of the Kaplan-Meier estimate from Section 3.3. In Table 3.3, without stratifying according to mother's status, the first observed event time is  $t_{(1)} = t_7 = 16$ . Then ignoring left-truncation,

- $\mathcal{R}_{t_{(1)}} = \{1, 2, 3, 4, 5, 6, 7, 8, 9\}$
- $d_{t_{(1)}} = 1$
- $n_{t_{(1)}} = 9$
- $S(t_{(1)}) = 1 - \frac{1}{9} \approx 0.9$

In contrast, when we take left-truncation into account, subjects only enter the risk set for the event after their left-truncation time (we already know they survived until  $t_i^L$  so they are not at risk for the event before that time), thus

- $\mathcal{R}_{t_{(1)}} = \{4, 5, 6, 7, 8, 9\}$
- $d_{t_{(1)}} = 1$
- $n_{t_{(1)}} = 6$
- $S(t_{(1)}) = 1 - \frac{1}{6} \approx 0.8$

Thus, the definition of the KM estimator in Equation 12.1 can still be used in case of left-truncation using the more general risk set definition

$$\mathcal{R}_\tau = \{i : t_i^L \leq \tau \leq t_i\} \quad (3.10)$$

## Right-truncation

Right-truncation often occurs in retrospective sampling based on registry data, when data is queried for cases reported by a certain cut-off time (see for example Vakulenko-Lagun, Mandel, and Betensky (2020)). A common example is the estimation of the incubation period of an infectious disease, which is the time from infection to the disease onset. Only known, symptomatic (and/or tested) cases are entered into the database. At a time  $\tau$ , one can only observe the subset of the infected population that has already experienced the disease, and not the population that is still incubating the disease, hence biasing the data to shorter incubation periods.

Formally, let  $t_i^r$  be the right-truncation time (here time from infection until the time at which the database is queried), then subjects only enter the data set when  $t_i < t_i^r$ . This is illustrated in Figure 3.6 using three subjects. All three subjects were infected during the observation period, however, the right-truncation time  $t_2^r$  of subject 2 is shorter than the incubation period  $t_2$  for this subject, thus at the time of querying the data base, this subject will not be included in the sample, as  $t_2 > t_2^r$ .

Note the difference to right-censoring. If subject 2 was right-censored, the subject would be in our sample and the time of infection would be known - the time of disease onset would be censored. In case of right-truncation on the other hand, the subject is not included in the sample at time of data extraction, as subjects are only included in the registry after disease

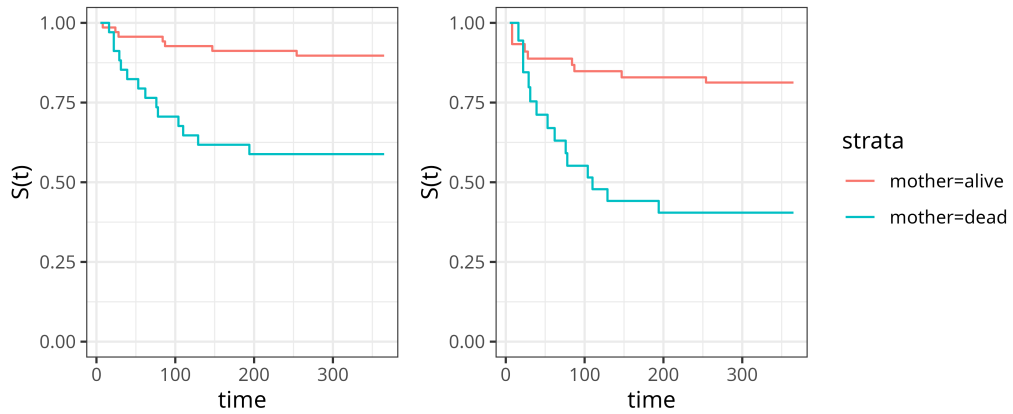


Figure 3.5: Left: Kaplan-Meier estimate of survival probabilities of infants depending on status of the mother ignoring left-truncation. Right: Kaplan-Meier estimate of survival probabilities adjusting for left-truncation.

onset. Overall this leads to a bias towards shorter incubation times and potentially feature values that lead to shorter incubation times.

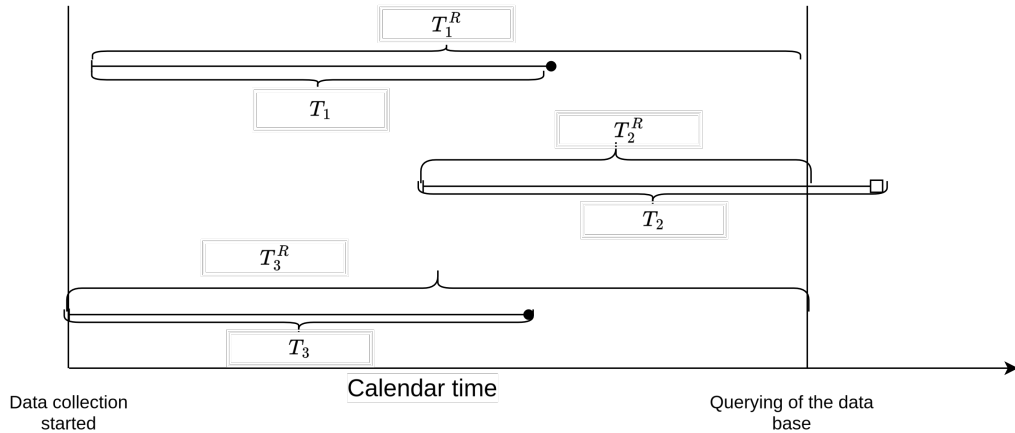


Figure 3.6: Illustration of right-truncation based on registry data. For subjects 1 and 3 the right-truncation time is longer than the incubation period, therefore they are included in the sample when the registry is queried. For subject 2 on the other hand the right-truncation time is shorter, therefore it's excluded from the sample.

As for left-truncation, ignoring right-truncation will lead to biased estimations. While for left-truncated data simple adjustments of the risk set work for non- and semi-parametric methods like the Kaplan-Meier and Cox-type estimators, this is not the case for right-truncated data. However, parametric methods can be employed (see Section 3.6) and generalised product limit estimators for right-truncated data exist (Michael G. Akritas and LaValley 2005).

## 3.6 Estimation

While details about estimation will be given later, when different models are introduced in Part III and Part IV of the book, it is worthwhile discussing some general concepts here, namely parametric and non-parametric approaches.

### 3.6.1 Parametric estimation

Consider for now uncensored data  $(t_i, \delta_i = 1), i = 1, \dots, n$ . A standard approach would be to assume a suitable distribution for the event times  $t_i \stackrel{iid}{\sim} F_Y(\boldsymbol{\theta}), \boldsymbol{\theta} = (\theta_1, \theta_2, \dots)^\top$ , and define the likelihood of the data as

$$\mathcal{L}(\boldsymbol{\theta}) = \prod_{i=1}^n f_Y(t_i | \boldsymbol{\theta}) \quad (3.11)$$

where  $f_Y$  is the pdf of  $F_Y$ .

The model parameters can then be obtained by maximizing the likelihood such that

$$\hat{\boldsymbol{\theta}} = \arg \max_{\boldsymbol{\theta}} \mathcal{L}(\boldsymbol{\theta})$$

However, in the presence of censoring Equation 3.11 is incorrect, as the exact event time is only known for some subjects. For example, for right-censored data ( $\delta_i = 0$ ) we only know that the event occurred after observed censoring time  $t_i$ . Thus the likelihood contribution for such data points is  $P(Y_i > t_i) = S_Y(t_i)$ , whereas for observed event times ( $\delta_i = 1$ ) the likelihood contribution is  $f_Y(t_i)$  as before.

Let now  $\mathcal{O} = \{i : \delta_i = 1\}$  the index set of observed event times and  $\mathcal{RC} = \{i : \delta_i = 0\}$  the index set of censored observations. The likelihood of this data can be written as

$$\begin{aligned} \mathcal{L}(\boldsymbol{\theta}) &\propto \prod_{i \in \mathcal{O}} f_Y(t_i | \boldsymbol{\theta}) \prod_{i \in \mathcal{RC}} S_Y(t_i | \boldsymbol{\theta}) \\ &= \prod_{i=1}^n f_y(t_i | \boldsymbol{\theta})^{\delta_i} S_Y(t_i | \boldsymbol{\theta})^{1-\delta_i} \\ &= \prod_{i=1}^n \frac{f_y(t_i | \boldsymbol{\theta})^{\delta_i}}{S_Y(t_i | \boldsymbol{\theta})^{\delta_i}} S_Y(t_i | \boldsymbol{\theta}) \\ &= \prod_{i=1}^n h_Y(t_i | \boldsymbol{\theta})^{\delta_i} S_Y(t_i | \boldsymbol{\theta}), \end{aligned}$$

where the last equality follows from Equation 3.2.

Similar adjustments to the likelihood can be made for other types of censoring and in the presence of truncation. Following Klein and Moeschberger (2003), we can define the individual likelihood contributions for the different types of censoring as

- observed event at  $t_i$ :  $f_Y(t_i | \boldsymbol{\theta})$
- right-censoring:  $P(Y_i > t_i | \boldsymbol{\theta}) = S_Y(t_i | \boldsymbol{\theta})$
- left-censoring:  $P(Y_i < t_i | \boldsymbol{\theta}) = F_Y(t_i | \boldsymbol{\theta}) = 1 - S_Y(t_i | \boldsymbol{\theta})$

- interval-censoring:  $P(l_i < Y_i \leq r_i | \boldsymbol{\theta}) = S_Y(l_i | \boldsymbol{\theta}) - S_Y(r_i | \boldsymbol{\theta})$

Depending on which of the above contributions occur in the data set, we can now construct our likelihood accordingly. Let  $\mathcal{O}, \mathcal{RC}, \mathcal{LC}, \mathcal{IC}$  non-overlapping subsets of the observed data  $\mathcal{D}$  for subjects with observed event times, right-censoring, left-censoring and interval-censoring, respectively. Assuming independence between observations and in absence of truncation, the likelihood for the observed data can be defined as

$$\mathcal{L}(\boldsymbol{\theta}) \propto \prod_{i \in \mathcal{O}} f_Y(t_i | \boldsymbol{\theta}) \prod_{i \in \mathcal{RC}} S_Y(t_i | \boldsymbol{\theta}) \prod_{i \in \mathcal{LC}} (1 - S_Y(t_i | \boldsymbol{\theta})) \prod_{i \in \mathcal{IC}} (S_Y(l_i | \boldsymbol{\theta}) - S_Y(r_i | \boldsymbol{\theta})) \quad (3.12)$$

In case of truncation, the adjustments are made to all observations, as we have to condition on the event occurring after/before the truncation time. The truncation adjusted likelihood contributions (assuming independence of truncation and event/censoring times) would thus be given by

- left-truncation:

- event:  $f(Y_i = t_i | Y_i \geq t_i^L, \boldsymbol{\theta}) = \frac{f_Y(t_i | \boldsymbol{\theta})}{S_Y(t_i^L | \boldsymbol{\theta})}$
- left-censoring:  $P(Y_i < t_i | Y_i \geq t_i^L, \boldsymbol{\theta}) = \frac{S_Y(t_i | \boldsymbol{\theta})}{S_Y(t_i^L | \boldsymbol{\theta})}$
- interval-censoring:  $P(l_i < Y_i \leq r_i | Y_i \geq t_i^L, \boldsymbol{\theta}) = \frac{S_Y(l_i | \boldsymbol{\theta}) - S_Y(r_i | \boldsymbol{\theta})}{S_Y(t_i^L | \boldsymbol{\theta})}$

- right-truncation:

- event:  $f(Y_i = t_i | Y_i \leq t_i^R, \boldsymbol{\theta}) = \frac{f_Y(t_i | \boldsymbol{\theta})}{F_Y(t_i^R | \boldsymbol{\theta})}$
- left-censoring:  $P(Y_i < t_i | Y_i \leq t_i^R, \boldsymbol{\theta}) = \frac{S_Y(t_i | \boldsymbol{\theta})}{F_Y(t_i^R | \boldsymbol{\theta})}$
- interval-censoring:  $P(l_i < Y_i \leq r_i | Y_i \leq t_i^R, \boldsymbol{\theta}) = \frac{S_Y(l_i | \boldsymbol{\theta}) - S_Y(r_i | \boldsymbol{\theta})}{F_Y(t_i^R | \boldsymbol{\theta})}$

Note that in practice, data sets will often not contain all types of censoring or truncation, in which case Equation 3.12 will contain only a subset of the product terms. This has been illustrated in Equation 3.12 under absence of truncation and  $\mathcal{LC} = \mathcal{IC} = \emptyset$ .

### 3.6.2 Non-parametric estimation

As the name suggests, non-parametric estimation (and semi-parametric estimation) techniques do not make (strong) assumptions about the underlying distribution of event times.

A common principle for such techniques is to partition the follow into intervals or to define specific time-points during the follow-up and to estimate the continuous or discrete time hazards (Equation 3.5) for each interval/time-point. Then derive the respective estimates, for example the survival probability, based on the relationship between the hazard and other quantities (Section 3.1 or Section 3.1.2).

The Kaplan-Meier estimator introduced earlier (Section 3.3) is an example for this principle. There, time-points  $t_{(k)}, k = 1, \dots, m$  are used to calculate the discrete time hazards

$$h^d(t_{(k)}) = P(Y \in (t_{(k-1)}, t_{(k)}] | Y > t_{(k-1)}) = \frac{d_{t_{(k)}}}{n_{t_{(k)}}}. \quad (3.13)$$

The survival probability and the definition of the Kaplan-Meier estimator (Equation 12.1) then follow from Equation 3.7.

Similarly, the Nelson-Aalen estimator (Nelson (1972), O. Aalen (1978)) for the cumulative hazard is obtained via Equation 3.6 as

$$H_{NA}(\tau) = \sum_{k:t_{(k)} \leq \tau} h^d(t_{(k)}) = \sum_{k:t_{(k)} \leq \tau} \frac{d_{t_{(k)}}}{n_{t_{(k)}}}. \quad (3.14)$$

Note that we use a continuous time valued  $\tau$  for the definition, which implies that  $H_{NA,e}(\tau) = H_{NA,e}(t_{(k)}) \forall \tau \in [t_{(k)}, t_{(k+1)})$ . In words: For time points between two unique event times we assume the previous value of the estimate. Similar to the Kaplan-Meier estimator,  $H_{NA}$  can deal with left-truncated data based on the general risk-set definition Equation 3.10.

Based on Equation 3.4 we can also define a survival probability estimator based on the Nelson-Aalen estimator of the cumulative hazard Equation 3.14 as

$$S_{NA}(\tau) = \exp(-H_{NA}(\tau)). \quad (3.15)$$

This relationship is also used by Breslow (1972) to obtain survival probability estimates in the context of the Cox model (Chapter 12).

! Major changes expected!

This page is a work in progress and major changes will be made over time.

### 3.7 Conclusion

Key takeaways

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Further reading

- For practical discussion about survival models in the context of right-censoring and left-truncation see McGough et al. (2021)



# 4

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## *Event-history Analysis*

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TODO (150-200 WORDS)

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In this chapter we take a more general view on time-to-event data. So far, we only considered a single potentially censored, outcome of interest. Here we explore more complex settings with multiple, potentially mutually exclusive events and recurrences of events. In this generalization, the observed data is sometimes referred to as *event-history data* and its analysis as *event-history analysis*.

One way to think about event history data is in terms of transitions between different states, as illustrated in Figure 4.1. Usually, a subject starts out in an initial state 0 (for example, ‘healthy’) and from there transitions to different states. States from which further transitions are possible are called *transient* (displayed as circles), otherwise a state is called *terminal* or *absorbing* (displayed as squares).

In the *single event* setting (Figure 4.1, upper left panel), a subject can only transition to one state (the event of interest). This setting was the focus of Chapter 3. There, the censoring event was considered independent of the event of interest. In the *competing risks* setting (Figure 4.1, upper right panel, Section 4.2), a subject could transition to any of the  $q$  mutually exclusive states, thus the subject is initially *at risk* for a transition to multiple states. Once one of them occurs, the process is considered to have concluded (for the modeling purposes).

In the *recurrent events setting* (Figure 4.1 lower left panel), the same event can be observed multiple times on the same subject (for example recurrent respiratory infections during one year). Two different ways to represent recurrent events are shown: (top) reset the status to 0 after occurrence of an event or (bottom) consider the 1st, 2nd, etc. recurrences of the event as separate states. A detail omitted in the graph: Often recurrent event processes also have a competing, absorbing event. In this more complex setting, but also in general, recurrent events are often represented as multi-state process, which we discuss next. Therefore we forgo detailed discussion of this setting in this book and refer to Cook and Lawless (2007) for a detailed account specific to recurrent events analysis.

In the most general case, the *multi-state* setting (Figure 4.1 lower right panel, Section 4.3), there are multiple transient and terminal states with potential back transitions (for example, moving between different stages of an illness with the possibility of (partial) recovery and death as terminal event).

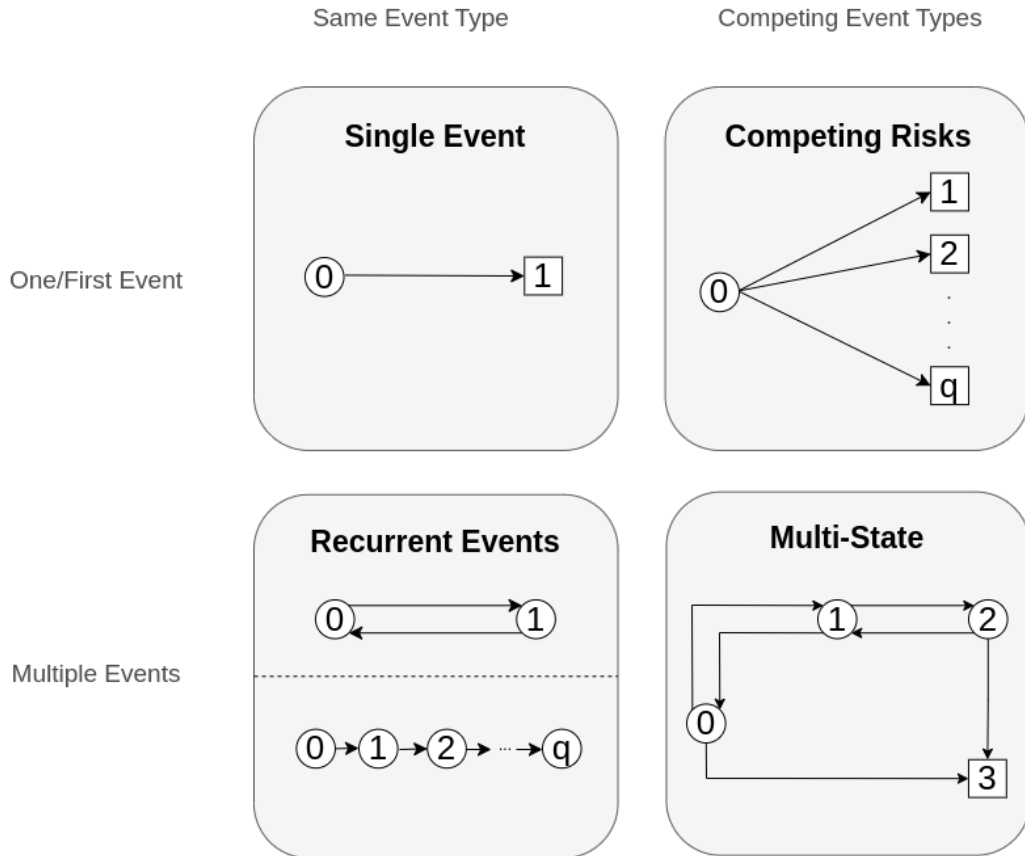


Figure 4.1: Illustration of different types of time-to-event processes. Transient states are displayed as circles, absorbing states are displayed as squares. Top, left: Standard single-event setting with transition from initial state 0 to state 1; Top, right: Competing risks setting with  $q$  competing events. The follow-up ends once one of the  $\{1, \dots, q\}$  events is observed or the study ends; Bottom, left: Recurrent events setting with multiple occurrences of the same event. Bottom, right: Multi-state setting where subjects can transition between multiple transient states with possible back-transitions or to absorbing states.

Note that the concepts discussed in Section 3.4 and Section 3.5 are still relevant here, as, dependent on the specific process, any transition between two states could be subject to different types of censoring and truncation. In particular, remaining in one of the transient states until the end of follow-up constitutes right-censoring with respect to all possible transitions from that state and left-truncation is particularly important as subjects enter the risk sets for a transition at different time points in context of recurrent events and multi-state settings.

---

## 4.1 A process point of view

In order to formalize the different settings more conveniently, we introduce the stochastic process

$$E(\tau) \in \{0, \dots, q\}, \quad \tau \geq 0, \tag{4.1}$$

which indicates the state that is occupied at time  $\tau$ .

Using this notation in the single-event setting we get  $E(\tau) \in \{0, 1\}$  such that the hazard Equation 3.2 could be written as

$$\begin{aligned} h(\tau) &= \lim_{d\tau \searrow 0} \frac{P(Y \in [\tau, \tau + d\tau) | Y \geq \tau)}{d\tau} \\ &= \lim_{d\tau \searrow 0} \frac{P(E(\tau + d\tau) = 1 | E(\tau-) = 0)}{d\tau}, \end{aligned} \tag{4.2}$$

where  $\tau-$  indicates the time point immediately before  $\tau$ .

This notation doesn't yield many advantages in the single-event setting, but will shorten notation later on, particularly in the multi-state setting.

---

## 4.2 Competing Risks

In contrast to single-event survival analysis, competing risks are concerned with the time to the first of multiple, mutually exclusive events.

Table 4.1. shows an excerpt of the `sir.adm` data (Allignol, Beyersmann, and Schumacher 2008) of patients on an intensive care unit (ICU). Time under observation (`time`) could end in one of three outcomes: 1 (discharge alive), 2 (death on ICU) or 0 (neither discharge nor death at the end of follow-up, which constitutes right-censoring at the end of study). The interest was in how pneumonia status (`pneumonia`) at admission to the ICU affects mortality.

Table 4.1: Subset of the `sir.adm` dataset (Allignol, Beyersmann, and Schumacher 2008). Each row represent one subject, `time` is the time under observation, `status` indicates the outcome observed (0: censored at the end of the study, 1: discharged alive from the ICU, 2: death in the ICU). `pneumonia` indicates whether a subject already had pneumonia at ICU admission.

time	status	pneumonia
8	0	no
8	0	no
31	1	yes
5	1	no
9	2	no
5	2	no

Contrast this data to the `tumor` data example in Section 3.3. There, patients were followed even after hospital discharge, thus loss to follow-up could be considered reasonably independent of the event of interest (death). In this study follow-up stopped once patients were discharged. As discharged patients are healthier compared to the ones who remain on ICU, assuming independence between the time until discharge and time until death is unrealistic. Analysis of this data and how the different assumptions (independent censoring vs. competing risks) affects the estimates is discussed in Section 4.2.2 and Section 4.2.4.

#### 4.2.1 Notation and Definitions

In the competing risks setting, everyone starts out in the initial state 0 and can progress to one of the absorbing states  $1, \dots, q$ . The goal is to characterize the process  $E(\tau) \in \{0, \dots, q\}$  in terms of transition hazards and probabilities.

In extension of Equation 4.2, we define *cause-specific* hazards

$$h_e(\tau) = \lim_{d\tau \rightarrow 0} \frac{P(E(\tau + d\tau) = e | E(\tau) = 0)}{d\tau}. \quad (4.3)$$

Analogous to the single-event case, we can also define the cause-specific cumulative hazard

$$H_e(\tau) = \int_0^\tau h_e(u) du \quad (4.4)$$

As competing events are mutually exclusive at any time  $\tau$ , it is possible to define the *all-cause hazard* which is the hazard of any event occurring as the sum of all cause-specific hazards

$$h(\tau) = \sum_{e=1}^q h_e(\tau), \quad (4.5)$$

as well as the *all-cause cumulative hazard*, which can be obtained either via the integral over the all-cause hazard (Equation 4.5) or as sum of cause-specific cumulative hazards (Equation 4.4):

$$H(\tau) = \sum_{e=1}^q H_e(\tau) = \sum_{e=1}^q \int_0^\tau h_e(u) du = \int_0^\tau \sum_{e=1}^q h_e(u) du \int_0^\tau h(u) du \quad (4.6)$$

The *all-cause survival probability* gives the probability that *none* of the events occurred before  $\tau$ . This is usually not estimated directly but calculated from the cause specific hazards instead via Equation 4.6 and Equation 4.7:

$$S(\tau) = P(Y > \tau) = \exp(-H(\tau)) \quad (4.7)$$

Finally, the probability of experiencing an event  $e$  before time  $\tau$ , *which is often referred to as Cumulative Incidence Function* (CIF), is given by

$$\begin{aligned} F_e(\tau) &= P(E(\tau) = e) \\ &= \int_0^\tau f_e(u) \, du = \int_0^\tau S(u-) h_e(u) \, du, \end{aligned} \quad (4.8)$$

where

- $S(u-)$  is the probability of surviving (not experiencing any of the competing events) until the time-point shortly before  $u$
- $f_e(u)du = S(u-)h_e(u)du$  is the probability of experiencing event  $e$  at time point  $u$  (which follows analogously to Equation 3.2).

Note that here we use the notation  $S(u-)$  rather than  $S(u)$  to make explicit that we want the probability to survive until the time point immediately before  $u$ . This doesn't make much difference in continuous time where  $P(T > t) = P(T \geq t)$ , but may be important in (discrete) approximations (as in Section 4.2.2).

$F_e(\tau)$  can be interpreted as the proportion of subjects who experienced event of type  $e$  until time  $\tau$ . Because the events are mutually exclusive, it holds that

$$\sum_{e=1}^q F_e(\tau) + S(\tau) = F(\tau) + S(\tau) = 1,$$

where  $F(\tau)$  is the probability that an event of any type occurring before  $\tau$  and  $S(\tau)$  the probability that no event occurs (Equation 4.7).

Note that all terms of Equation 4.8 can be calculated from the individual hazards (Equation 4.3). Many estimation procedures for the CIF take this approach, consequently referred to as cause-specific hazards approach.

#### 4.2.2 Non-parametric estimators

Non-parametric estimators for the cause-specific (cumulative) hazard (Equation 4.4) in the competing risks setting are derived analogous to the single event case (Section 3.6.2). The CIF then follows from Equation 4.8.

First, recall from Section 3.2 the definitions of the unique ordered event times  $t_{(k)}$ ,  $k = 1, \dots, m$ , the risk-set at time  $t_{(k)}$ ,  $\mathcal{R}_{t_{(k)}}$ , the number of events,  $d_{t_{(k)}}$ , and number of observations at risk  $n_{t_{(k)}}$ . Assume partitioning of the follow-up into  $m$  disjunct intervals  $(t_{(k-1)}, t_{(k)}]$ ,  $k = 1, \dots, m$ , such that  $Y \in (t_{(k-1)}, t_{(k)}] \Leftrightarrow \tilde{Y} = t_{(k)}$ , with  $\tilde{Y}$  defined as in Chapter 20..

An estimate for the *cause-specific* hazard is derived by updating the numerator in Equation 3.13 to  $d_{e,t_{(k)}}$  (the number of events of type  $e$  at time  $t_{(k)}$ ):

$$h_e^d(t_{(k)}) := \frac{d_{e,t_{(k)}}}{n_{t_{(k)}}}, \quad e \in \{1, \dots, q\}. \quad (4.9)$$

The Nelson-Aalen estimator (Equation 3.14) for the *cause-specific* cumulative hazard is then given by

$$H_{NA,e}(\tau) = \sum_{k:t_{(k)} \leq \tau} h_e^d(t_{(k)}) = \sum_{k:t_{(k)} \leq \tau} \frac{d_{e,t_{(k)}}}{n_{t_{(k)}}}, \quad (4.10)$$

which yields a step function for each  $e$ , with jumps at time points  $t_{(k)}$ .

The all-cause survival probability follows from Equation 4.6 and Equation 4.7 as

$$S_{NA}(\tau) = \exp \left( - \sum_{e=1}^q H_{NA,e}(\tau) \right).$$

Finally, the Aalen-Johansen (AJ) estimator (O. O. Aalen and Johansen (1978)) for the CIF follows via Equation 3.8 as

$$F_{AJ,e}(\tau) = \sum_{k:t_{(k)} \leq \tau} S_{NA}(\tau-) h_e^d(\tau) = \sum_{k:t_{(k)} \leq \tau} S_{NA}(t_{(k-1)}) \frac{d_{e,t_{(k)}}}{n_{t_{(k)}}} \quad (4.11)$$

### 4.2.3 Application to mortality of ICU patients

For illustration of the AJ estimator and the interpretation of the CIFs consider the analysis conducted in Beyersmann, Allignol, and Schumacher (2012), based on the data from Table 4.1. Recall that one is interested in estimation of the mortality conditional on pneumonia status at admission, while accounting for discharge from the ICU as competing risk ( $E \in \{"\text{discharge}", "\text{death}"\}$ ). While the AJ estimator cannot naturally incorporate feature information, it can be applied to subgroups of the data (here based on the pneumonia status). Note that this will yield different sets of unique event times in each group, thus the AJ can have jumps at different time-points for the two groups.

TODO: clean up text below

Figure 4.2 shows the AJ estimates of the CIFs for each event type (discharge/death) stratified by pneumonia status. Exemplary, the proportion of subjects with pneumonia being discharged until  $\tau = 120$  days is approximately 75% ( $\hat{F}_{\text{discharge}}(120) = P(Y \leq 120, E = "\text{discharge}") \approx 0.75$ ), while approximately 25% died in the ICU ( $\hat{F}_{\text{death}}(120) = P(Y \leq 120, E = "\text{death}") \approx 0.25$ ). For patients without pneumonia we have  $\hat{F}_{\text{discharge}}(120) \approx 0.91$  and  $\hat{F}_{\text{death}} \approx 0.09$ . In this example,  $\hat{F}_{\text{discharge}} + \hat{F}_{\text{death}} \approx 1$  for both pneumonia groups, as only 14 of 747 patients were censored (neither discharge nor death) at the end of the follow-up.

### 4.2.4 Independent Censoring vs. Competing Risks

It is worth spending some time to consider the difference between independent right-censoring and competing risks. Note that for the estimation of the hazard (Equation 4.9), occurrences of competing events are implicitly assumed right-censored (as  $d_{e,t_{(k)}}$ ) only counts events of type  $e$  and  $n_{t_{(k)}}$  contains the same subjects that would remain if events of type  $\tilde{e} \neq e$  were considered censored before  $t_{(k)}$ . Nevertheless, competing risks are taken into account in the definition of the AJ estimator (Equation 4.11), as the all-cause survival probability (Equation 4.7) depends on all cause-specific hazards.

In contrast, assume that in our analysis of the `sir.adm` data we would consider time of discharge as independent right-censoring. As we only have one other event (death), the data could be treated as single-event, right-censored data as in Chapter 3 and therefore analyzed

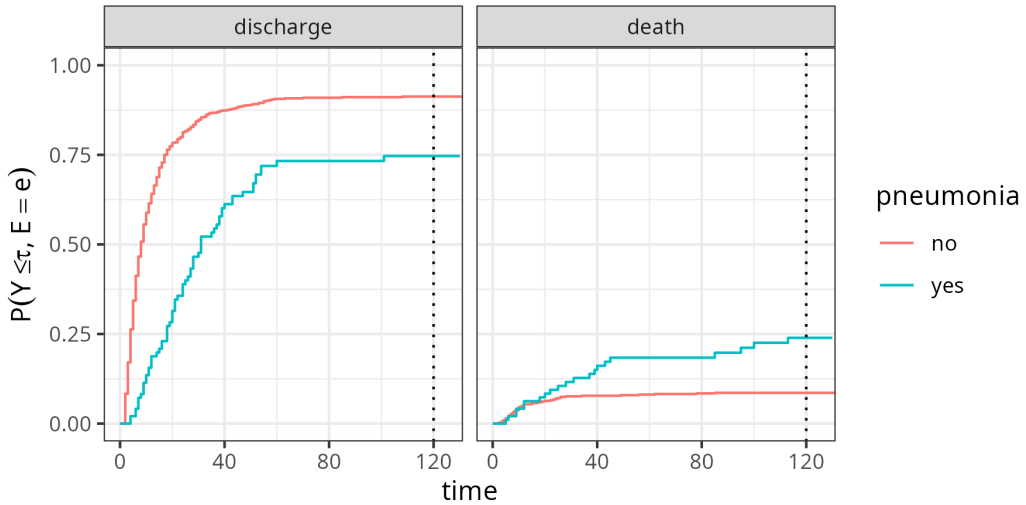


Figure 4.2: Aalen-Johansen estimator for the `sir.adm` data (Allignol, Beyersmann, and Schumacher 2008), stratified by pneumonia status at admission to the ICU. Left panel: Proportion of subjects discharged alive from the ICU. Right panel: Proportion of subjects who died in the ICU.

using the Nelson-Aalen estimator (Equation 3.15). The probability of death before some time-point  $\tau$  could thus be obtained via  $P(Y \leq \tau) = F(\tau) = 1 - S_{NA}(\tau)$ .

Figure 4.3 shows the estimates obtained under the two assumptions. Solid lines indicate the probabilities under the competing risks assumption (identical to the right-hand side of Figure 4.2). Dashed lines are obtained under the independent right-censoring assumption. Clearly, the probabilities of dying at time  $\tau = 120$  are greater when independent censoring is assumed ( $\approx 75\%$  vs.  $\approx 25\%$  in the pneumonia group and  $\approx 62\%$  vs.  $\approx 13\%$  in the no pneumonia group).

### 4.3 Multi-state Models

The multi-state process can be considered the most general type of time-to-event process, as other types (single-event, competing risks, recurrent events) can be viewed as special cases. Multi-state modeling allows realistic depiction of complex processes where subjects can start in different states and transition back and forth between them.

For illustration consider the `prothr` dataset (de Wreede, Fiocco, and Putter 2011) of liver cirrhosis patients from a randomized clinical trial with possible transitions depicted in Figure 4.4. Patients may have normal (state 0) or abnormal (state 1) levels of prothrombin (a protein important for blood clotting, produced by the liver) at the beginning of the trial. Some patients were treated with prednisone (which suppresses immune response and reduces inflammation) and others received a placebo. Death (state 2) constitutes an absorbing state.

The goal of the trial was to investigate if treatment (prednisone) slows down or reverses

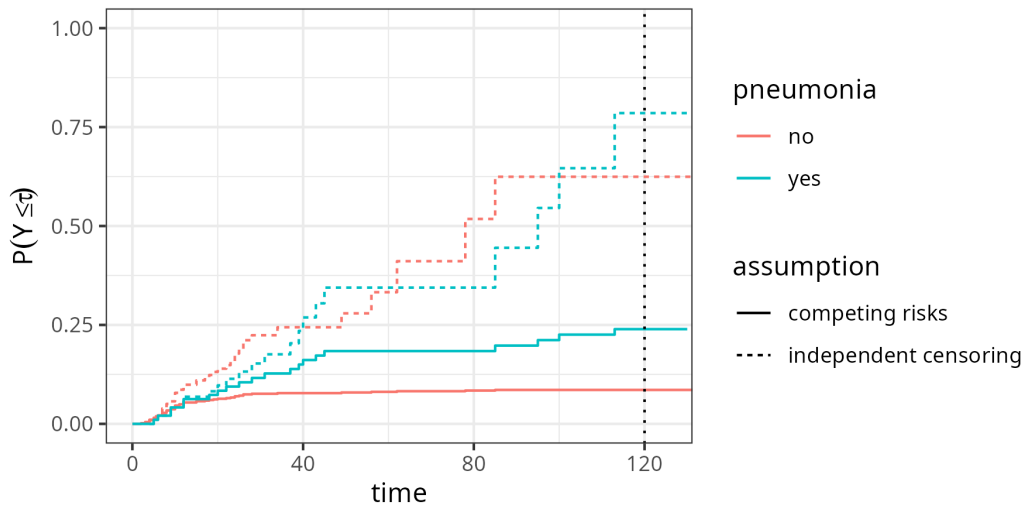


Figure 4.3: Estimation of the probability of dying in the ICU conditional on pneumonia status at admission. Dashed lines give the probabilities under assumption of right-censoring. Solid lines give the probabilities when taking into account discharge as competing risk.

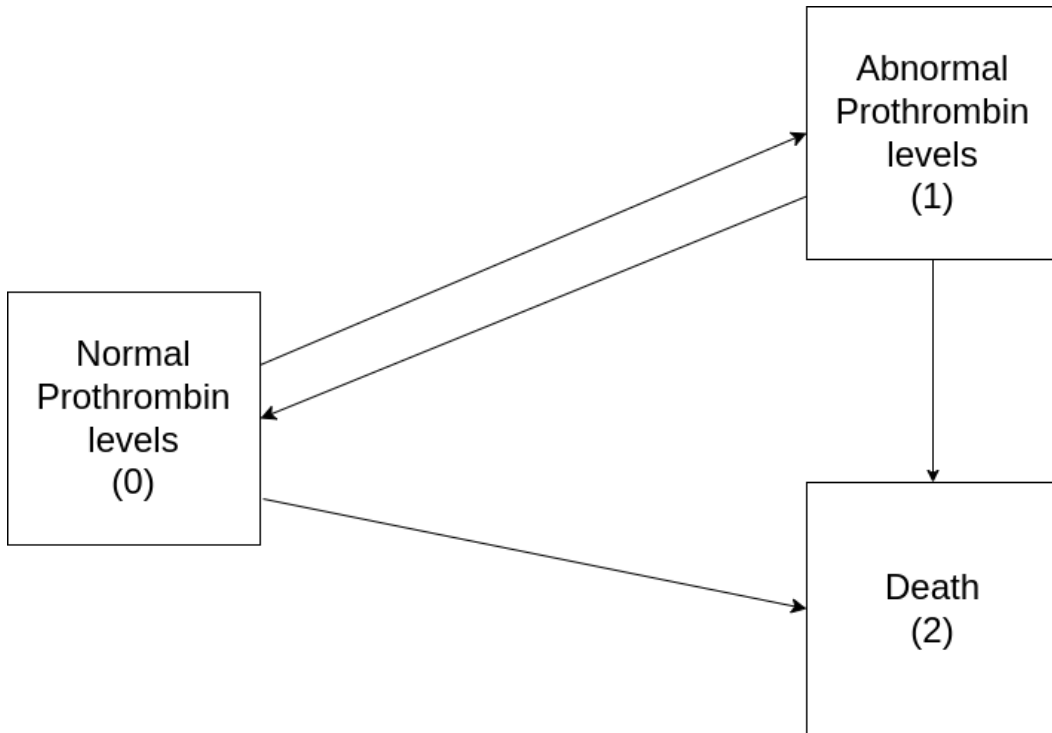


Figure 4.4: Transition graph for the liver cirrhosis patients.

disease progression (transitions  $0 \rightarrow 1$  and  $1 \rightarrow 0$ ) and reduces mortality (transitions  $0 \rightarrow 2$  and  $1 \rightarrow 2$ ).

Table 4.2 shows a subset of the data set and contains for each subject (**id**) one row for each transition for which the subject was at risk for. In this example, this includes transitions that were possible, but didn't happen (counterfactual transitions). The columns **from** and **to** indicate the initial state and the possible end state. **tstart** indicates the time at which the subject entered the risk set for said transitions and **tstop** the time point at which the subjects exited the **from** state (or were censored for any transition). The variable **status** indicates whether the transition was actually made (**status = 1**) or not (**status = 0**). This is necessary, as all possible transitions are listed, so we need an indicator for which transition actually occurred. If **status=0** for all possible transitions, the subject is censored for further transitions. Finally, **treatment** indicates whether a patient was assigned the treatment or placebo group.

Concretely, subject **id=1** already had abnormal prothrombin levels at the beginning of the trial, thus started in state 1 with possible transitions  $1 \rightarrow 0$  and  $1 \rightarrow 2$ . In this case, the patient died, thus transition  $1 \rightarrow 2$  was realized after 151 days, while the transition  $1 \rightarrow 0$  is a ‘counterfactual’ transition that could have happened in the time-span between **tstart=0** and **tend=151**, but didn't. Patient **id=8** also started in state 1, but made a back transition to normal prothrombin levels after 211 days at which time they entered the risk set for transitions  $0 \rightarrow 1$  and  $0 \rightarrow 2$ . Neither of the transitions occurred, as **status=0** for both transitions, which means the subject remained in status 0 until the end of their follow-up at 2770 days (that is was right-censored at 2770 days). Finally, subject **id=46** started in state 0 (normal prothrombin levels), transitioned to state 1 (abnormal levels) after 415 days and then died (transition  $1 \rightarrow 2$ ) two days later. This also illustrates the importance of left-truncation (Section 3.5) in multi-state processes. For example, subjects **id=1** and **id=8** are at risk for the transitions  $1 \rightarrow 0$  and  $1 \rightarrow 2$  from the beginning of the trial (**tstart = 0**). Subject **id=46** on the other hand starts in state 0 and only enters state 1 (and thus the risk set for the transitions  $1 \rightarrow 0$  and  $1 \rightarrow 2$ ) after 415 days (**tstart = 415**). Other subjects in the data may never enter the risk set for these transitions by remaining in state 0 until the end of follow up or by directly transitioning to state 2. The fact that subjects enter the risk sets for different transitions at different time points technically constitutes left-truncation and thus should be taken into account accordingly (Section 4.3.4).

Table 4.2: Subset of the **prothr** dataset (de Wreede, Fiocco, and Putter 2011).

<b>id</b>	<b>from</b>	<b>to</b>	<b>trans</b>	<b>tstart</b>	<b>tstop</b>	<b>status</b>	<b>treatment</b>
1	1	0	3	0	151	0	Placebo
1	1	2	4	0	151	1	Placebo
8	1	0	3	0	211	1	Prednisone
8	1	2	4	0	211	0	Prednisone
8	0	1	1	211	2770	0	Prednisone
8	0	2	2	211	2770	0	Prednisone
46	0	1	1	0	415	1	Prednisone
46	0	2	2	0	415	0	Prednisone
46	1	0	3	415	417	0	Prednisone
46	1	2	4	415	417	1	Prednisone

### 4.3.1 Notation and Definitions

In the competing risks setting, we characterized the data generating process by cause-specific transitions hazards  $h_e(\tau)$  (Equation 4.3). Implicitly these are transition from starting state 0 to end state  $e$ , however, since everyone starts in state 0 this information is ignored. In the multi-state setting on the other hand, subjects can be in different states at different time points. Transitions between different states are therefore characterized by transition-specific hazards, denoted by  $h_{\ell \rightarrow e}(\tau)$  or short  $h_{\ell e}(\tau)$ , where  $\ell$  is the starting state and  $e$  the end state,  $\ell, e \in \{0, \dots, q\}$  (ignoring details such as the starting state must be a transient state and not all states are reachable from each starting state).

Let  $E(\tau) \in \{0, \dots, q\}$  be the state process as before (Equation 4.1). Then, the transition-specific hazard can be defined as

$$h_{\ell e}(\tau) = \lim_{d\tau \rightarrow 0} \frac{P(E(\tau + d\tau) = e | E(\tau-) = \ell)}{d\tau} \quad (4.12)$$

Transition hazards Equation 4.12 indicate the relative risk to enter state  $e$  at time  $\tau$  given occupation of state  $\ell$  at  $\tau-$ , which is the instant before  $\tau$ .

Analogous to the competing risks setting, we can define the transition specific cumulative hazards

$$H_{\ell e}(\tau) = \int_0^\tau h_{\ell e}(u) du \quad (4.13)$$

The probability to transition from state  $\ell$  to  $e$  between two time-points depends on all transitions possible from  $\ell$  and potentially the transitions that have taken place in the past. Thus, other quantities of interest in the multi-state setting are the transition probabilities  $P_{\ell e}(\zeta, \tau) := P(E(\tau) = e | E(\zeta) = \ell)$ , that is the probability to transition from state  $\ell$  to state  $e$  between time points  $\zeta < \tau$ . Implicitly, this notation assumes that the process is Markovian: the transition probability depends only on the state at  $\zeta$  and not any additional past states. Extensions do exist that relax this assumption, for example by including information about the past, but are not relevant for now.

Transition probabilities of a multi-state process are often summarized in a matrix

$$\mathbf{P}(\zeta, \tau) := \begin{pmatrix} P_{00}(\zeta, \tau) & \cdots & P_{0q}(\zeta, \tau) \\ \vdots & \ddots & \vdots \\ P_{q0}(\zeta, \tau) & \cdots & P_{qq}(\zeta, \tau) \end{pmatrix}, \quad (4.14)$$

where rows indicate starting states and columns indicate end states. Some of the elements of  $\mathbf{P}$  might be zero or one depending on the specific process, presence of absorbing states and possible pathways between states. As subjects can only be in one of the  $q+1$  states at  $\tau$ , rows sum to 1:

$$\sum_{e=0}^q P_{\ell e} = 1, \forall \ell \in \{0, \dots, q\}. \quad (4.15)$$

### 4.3.2 Instantaneous transition probabilities

In this section we briefly recap how the transition probabilities can be expressed as the product (integral) of instantaneous transition probabilities

$$p_{\ell e}(\tau) = P(E(\tau + d\tau) = e | E(\tau-) = \ell) \quad (4.16)$$

which can be recognized as the nominator of Equation 4.12, or intuitively as the transition probability between two subsequent time points.

For illustration, consider what is often referred to as an illness-death model depicted in Figure 4.5 (similar to Figure 4.4, but without back-transition), where subjects can transition from healthy state 0 to absorbing state death (2) either directly or via intermediate state 1.

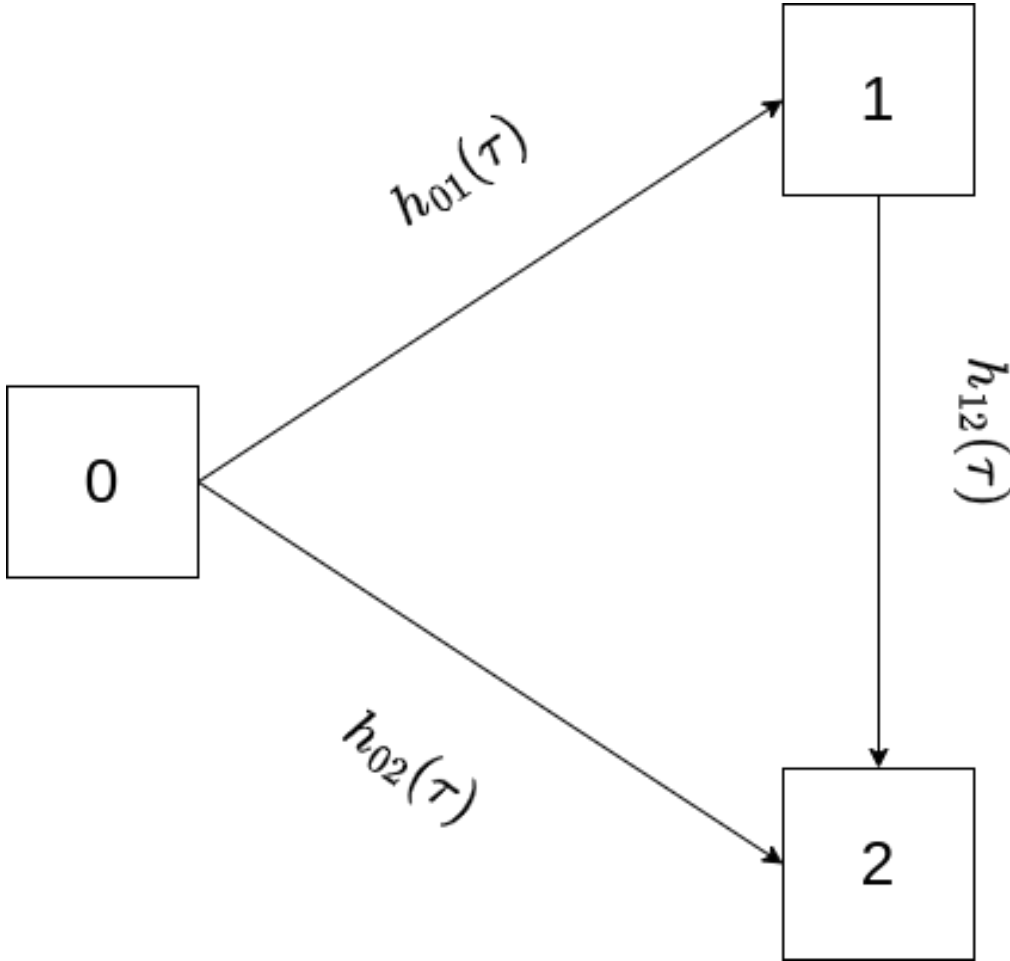


Figure 4.5: An *illness-death model* where subjects can transition from healthy state (0) to death (2) directly or via intermediate illness state (1)

In this example, back-transitions are not possible, therefore the lower triangle of the matrix is filled with zeros and  $P_{22}(\zeta, \tau) = 1, \forall \zeta < \tau$  by virtue of being an absorbing state. The matrix of transition probabilities is thus given as

$$\mathbf{P}(\zeta, \tau) = \begin{pmatrix} P_{00}(\zeta, \tau) & P_{01}(\zeta, \tau) & P_{02}(\zeta, \tau) \\ 0 & P_{11}(\zeta, \tau) & P_{12}(\zeta, \tau) \\ 0 & 0 & 1 \end{pmatrix} \quad (4.17)$$

First, assume that data is collected in discrete time, that is  $\zeta, \tau \in \{0, 1, 2, \dots\}, \zeta < \tau$  and transitions only occur at these discrete time points and not in between. Say we are interested in transition probability  $P_{02}(4, 6)$ , that is the probability to transition from state 0 to state 2

between time points  $\zeta = 4$  and  $\tau = 6$ , given we are in state 0 at time  $\zeta = 4$ . This is possible in the three ways depicted in Figure 4.6.

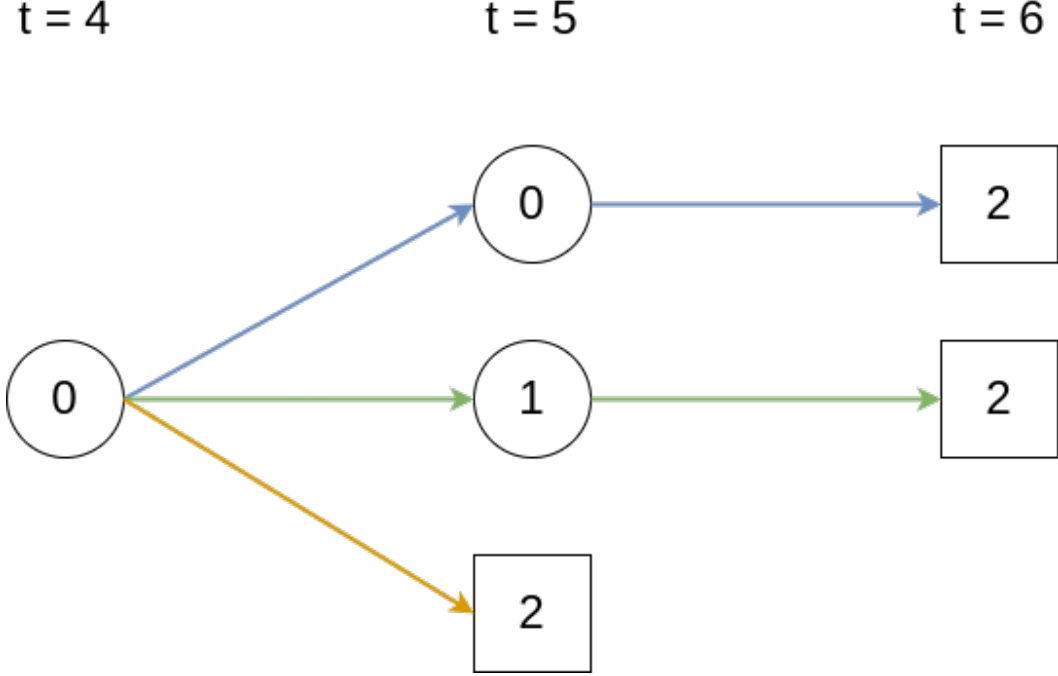


Figure 4.6: Possible paths to transition from state 0 to state 2 between time points 4 and 6

Thus,  $P_{02}(4, 6) = p_{00}(5)p_{02}(6) + p_{01}(6)p_{12}(6) + p_{02}(5)$ , where  $p_{\ell e}(\tau) = P(E(\tau) = e | E(\tau-1) = \ell)$  are the probabilities for transitions  $\ell \rightarrow e$  between two subsequent discrete time points. Thus, in discrete time, the matrix of transition probabilities can be represented as a finite matrix product

$$\mathbf{P}(\zeta, \tau) = \prod_{j=\zeta+1}^{\tau} \begin{pmatrix} p_{00}(j) & p_{01}(j) & p_{02}(j) \\ 0 & p_{11}(j) & p_{12}(j) \\ 0 & 0 & 1 \end{pmatrix}. \quad (4.18)$$

For the concrete example we thus have

$$\begin{aligned} \mathbf{P}(4, 6) &= \begin{pmatrix} P_{00}(4, 6) & P_{01}(4, 6) & P_{02}(4, 6) \\ 0 & P_{11}(4, 6) & P_{12}(4, 6) \\ 0 & 0 & 1 \end{pmatrix} = \prod_{j=5}^6 \begin{pmatrix} p_{00}(j) & p_{01}(j) & p_{02}(j) \\ 0 & p_{11}(j) & p_{12}(j) \\ 0 & 0 & 1 \end{pmatrix} \\ &= \begin{pmatrix} p_{00}(5) & p_{01}(5) & p_{02}(5) \\ 0 & p_{11}(5) & p_{12}(5) \\ 0 & 0 & 1 \end{pmatrix} \begin{pmatrix} p_{00}(6) & p_{01}(6) & p_{02}(6) \\ 0 & p_{11}(6) & p_{12}(6) \\ 0 & 0 & 1 \end{pmatrix} \\ &= \begin{pmatrix} p_{00}(5)p_{00}(6) & p_{00}(5)p_{01}(6) + p_{01}(5)p_{11}(6) & p_{00}(5)p_{02}(6) + p_{01}(5)p_{12}(6) + p_{02}(5) \\ 0 & p_{11}(5)p_{11}(6) & p_{11}(5)p_{12}(6) + p_{12}(5) \\ 0 & 0 & 1 \end{pmatrix}, \end{aligned}$$

where the quantity of interest,  $P_{02}(4, 6)$  is given in the top right corner, but other transition

probabilities are also readily available. For example, the probability to transition from state 1 to 2 between time points  $\zeta = 4$  and  $\tau = 6$  is given as  $P_{12}(4, 6) = p_{11}(5)p_{12}(6) + p_{12}(5)$ .

Returning to the continuous time setting where transitions can occur at any time point, ideas from the discrete time setting still hold. Imagine dividing the interval  $(\zeta, \tau]$  into  $J$  intervals such that  $\zeta = t_0 < t_1 < \dots < t_j < \dots < t_J = \tau$ , assuming that no events occur between time points  $t_j \in \mathbb{R}_+, j = 1, \dots, J$ . Then Equation 4.18 still holds when replacing  $p_{\ell e}(j)$  with  $p_{\ell e}(t_j)$ .

Increasing the number of intervals to infinity or equivalently, reducing the interval size to infinitesimally small intervals  $du$  where only one transition can be observed, leads to a product integral over the instantaneous transition probabilities  $p_{\ell e}(u)$  (Equation 4.16), such that

$$\mathbf{P}(\zeta, \tau) = \lim_{du \rightarrow 0} \prod_{u \in (\zeta, \tau]} \begin{pmatrix} p_{00}(u) & \cdots & p_{0q}(u) \\ \vdots & \ddots & \vdots \\ p_{q0}(u) & \cdots & p_{qq}(u) \end{pmatrix} \quad (4.19)$$

### 4.3.3 Instantaneous transition probabilities and hazards

In context of survival analysis we want to express the transition matrix (Equation 4.19) and thus instantaneous transition probabilities  $p_{\ell e}(u)$  in terms of (cumulative) hazards. To do so, we use, somewhat informally, the following relationships

- From equations Equation 4.12 and Equation 4.13 we can equate the instantaneous transition probabilities to increments of the cumulative hazard (that is the increase in the cumulative hazard within a (fixed, infinitesimally) small interval  $du$ ):  $dH_{\ell e}(u) = h_{\ell e}(u)du = P(E(u + du) = e | E(u-) = \ell) = p_{\ell e}(u)$ ,
- because of relationship Equation 4.15, diagonal elements (transitions into the same state) are set to  $dH_{\ell\ell}(u) := -\sum_{e \neq \ell} dH_{\ell e}(u)$  such that  $1 + dH_{\ell\ell}(u) = 1 - \sum_{e \neq \ell} dH_{\ell e}(u) = 1 - \sum_{e \neq \ell} p_{\ell e}(u) = p_{\ell\ell}(u)$ .

Consequently, Equation 4.19 can be rewritten as

$$\begin{aligned} \mathbf{P}(\zeta, \tau) &= \lim_{du \rightarrow 0} \prod_{u \in (\zeta, \tau]} \begin{pmatrix} p_{00}(u) = 1 + dH_{00}(u) & \cdots & p_{0q}(u) = dH_{0q}(u) \\ \vdots & \ddots & \vdots \\ p_{q0}(u) = dH_{q0}(u) & \cdots & p_{qq}(u) = 1 + dH_{qq}(u) \end{pmatrix} \\ &= \lim_{du \rightarrow 0} \prod_{u \in (\zeta, \tau]} (\mathbf{I} + d\mathbf{H}(u)), \end{aligned}$$

where  $\mathbf{I}$  is a  $(q+1) \times (q+1)$  identity matrix and  $d\mathbf{H}(u)$  is the matrix of increments of the cumulative hazard within infinitesimally small intervals. Thus, similar to the competing risks setting, relationship Equation 4.19 implies that knowledge of the transition specific (cumulative) hazards is sufficient to fully specify the multi-state process.

As analytical solutions of Equation 4.19 only exist for specific models, in practice, the transition probabilities are often once again approximated numerically via a finite matrix product on a discrete time grid  $\zeta = t_0 < t_1 < \dots < t_{J-1} < t_J = \tau$

$$\mathbf{P}(\zeta, \tau) \approx \prod_{j=1}^J (\mathbf{I} + \Delta \mathbf{H}_{\ell e}(t_j)), \quad (4.20)$$

where  $\Delta H_{\ell e}(t_j) = H_{\ell e}(t_j) - H_{\ell e}(t_{j-1})$  is the increment of the cumulative hazards between consecutive time points.

#### 4.3.4 Non-parametric estimation of transition probabilities

From Equation 4.20, it follows that transition probabilities can be estimated by first computing the transition-specific cumulative hazards,  $H_{\ell e}(\tau)$ . Similarly to the competing risks setting (Section 4.2.2), we can first define the transition specific hazards

$$h_{\ell e}^d(t_{(k)}) := \frac{d_{\ell e, t_{(k)}}}{n_{\ell; t_{(k)}}},$$

where

- $d_{\ell e, t_{(k)}}$  is the number of subjects that made the transition  $\ell \rightarrow e$  at time  $t_{(k)}$  and
- $n_{\ell; t_{(k)}}$  is the number of subjects in state  $\ell$  immediately before  $t_{(k)}$ .

The cumulative transition-specific hazards follow as

$$H_{NA, \ell e}(\tau) = \sum_{k: t_{(k)} \leq \tau} h_{\ell e}^d(t_{(k)}) = \sum_{k: t_{(k)} \leq \tau} \frac{d_{\ell e, t_{(k)}}}{n_{\ell; t_{(k)}}},$$

and transition probabilities are obtained via Equation 4.20 as

$$\mathbf{P}(\zeta, \tau) = \prod_{j=1}^J (\mathbf{I} + \Delta \mathbf{H}_{NA, \ell e}(t_{(j)})). \quad (4.21)$$

#### 4.3.5 Application to liver cirrhosis patients

For illustration, consider again the `prothr` data set (Table 4.2), with possible transitions summarized in Figure 4.4. In contrast to the illness-death model in Figure 4.5, back transitions are possible and some subjects already start in the “abnormal” state at the beginning of the study.

Figure 4.7 shows the transition probabilities for the four possible transitions over time for subjects who received treatment and placebo, respectively. In this example back-transitions are possible, therefore, in contrast to the cumulative incidence functions in the competing risks setting, transition probabilities (to transient states) are not monotonously increasing over time. While the probabilities to transition from normal or abnormal state to death ( $0 \rightarrow 2$ ,  $1 \rightarrow 2$ ) increase over time for both groups, probabilities for transitions between the transient states (normal to abnormal and vice versa) increase in the beginning but eventually decreases over time. Overall, prednisone doesn’t appear to have a strong protective effect. Although there appears to be a reduction in  $0 \rightarrow 2$  transitions and an increase in  $1 \rightarrow 0$  transitions between time points 1000 and 3000, this effect doesn’t seem to persist until the end of the follow up.

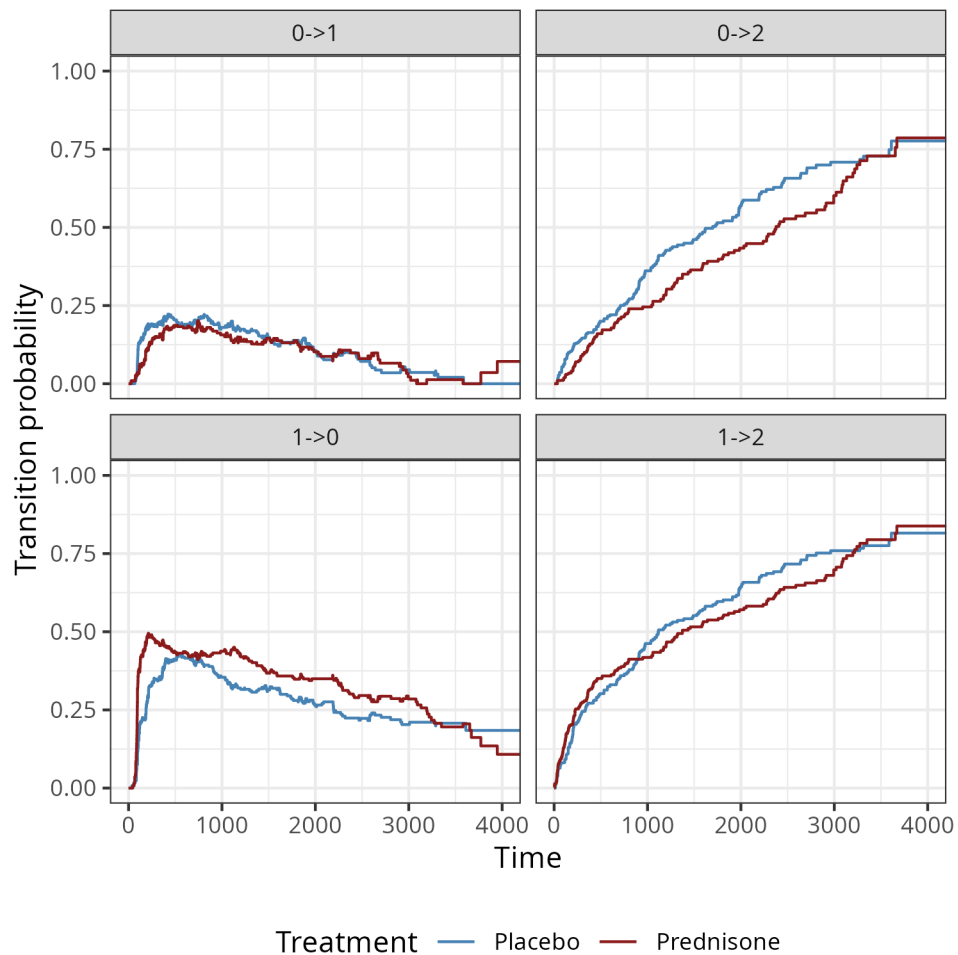


Figure 4.7: Estimated transition probabilities for the different transitions of the prothrombin data example .



# 5

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## Survival Task

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This chapter introduces the different survival problems and formalises them as survival tasks. There are four prediction types in survival analysis: relative risks - predicting the risk of an event, survival times - predicting the time until an event happens, prognostic index - predicting a linear predictor to assess outcomes based on risk factors, and survival distributions - predicting the probability of an event taking place over time. These reduce to three formal survival tasks: deterministic (survival time), ranking (risks and prognostic index), and probabilistic (distribution). This separation of tasks helps create a taxonomy of survival models and losses that is used throughout the book.



Minor changes expected!

This page is a work in progress and minor changes will be made over time.

The preceding chapters have focused on understanding survival data. This final chapter turns to the prediction tasks themselves.

Recall from Section 2.1 that a general survival prediction problem is one in which:

1. a survival dataset,  $\mathcal{D}$ , is split for training,  $\mathcal{D}_{train}$ , and testing,  $\mathcal{D}_{test}$ ;
2. a survival model is fit on  $\mathcal{D}_{train}$ ; and
3. the model predicts a representation of the unknown true survival time,  $Y$ , given  $\mathcal{D}_{test}$ .

The process of model fitting is model-dependent and ranges from non-parametric methods to machine learning approaches; these are discussed in Part III. Deciding what ‘the representation of  $Y$ ’ actually is will be the focus of this chapter.

In general there are four common *prediction types* or *prediction problems* which codify different representations of  $Y$ , these are:

1. Survival distributions;
2. Time-to-event;
3. Relative risks;
4. Prognostic index.

In analogy to classification, the first of these is considered *probabilistic* as it includes uncertainty about the prediction, whereas the second is *deterministic* as the prediction is a single value without associated uncertainty, which may also be known as a point prediction. Technically, the last two prediction types may appear deterministic for the same reason as

(2), however, both implicitly capture uncertainty as neither is a direct prediction of  $Y$ , as will be discussed further below. Whilst these prediction types can be separated in this way, the distribution prediction is the most general case; relative risks and time-to-event can both be derived from a probabilistic prediction and the prognostic index is often used as an interim step before making a probabilistic prediction.

It is vital to clearly separate which prediction problem you are working with, as they are not directly compatible with one another. For example, it is not meaningful to compare a relative risk prediction from one model to a survival distribution prediction of another. Moreover, prediction types may look similar but have very different implications, for example if one erroneously interprets a survival time as a relative risk then they could assume an individual has a worse outcome than they do as longer survival times imply lower risk.

Each prediction type has its own particular advantages and disadvantages and should be used appropriately. For example, survival times are not advised in domains where over-confidence can be detrimental, but can be used when making less-critical predictions, such as a marathon runner's finish time (Figure 5.1, top-left), which is a survival analysis problem as many people do not finish. Whilst clinicians will usually not present a probability distribution to a patient (Figure 5.1, top-right), it is common to relay information such as '5-year survival probability', which is read from a predicted survival distribution. Finally, relative risk predictions are often used for triage and resource allocation, such as assigning patients to particular wards or beds (Figure 5.1, bottom).

Whilst one prediction type is necessary to define a model's output, it is often the case that an algorithm might use another prediction type as an interim prediction. This chaining of predictions is called 'composition' as two components are composed to create a final prediction, this is commonly the case for distribution predictions. It is also possible to convert prediction types between one another. Table 5.1 highlights how distribution and survival time predictions can be transformed to most other forms and how all forms can be transformed to a relative risk prediction. These transformations will be discussed in more detail in the rest of this chapter.

Throughout this chapter let  $\mathcal{X} \subseteq \mathbb{R}^{n \times p}$  be a set representing covariates. Note that the type of censoring in the data is part of the estimation problem and does not affect the prediction types, hence this chapter does not differentiate between right, left, or interval censoring. The focus is primarily on the single-event setting, a brief section on other settings is included at the end of the chapter.

Table 5.1: Converting prediction types between one another. Rows are the original prediction type and columns are the transformed prediction, entries are the transformation methods. 'NA' indicates transformation is not possible between the two prediction types without significant assumptions.  $\phi, \Phi$  are arbitrary functions.

	Distribution	Risk	Time	PI
Distribution, $\hat{S}$	$\hat{S}$	$\sum_t \hat{H}(t)$	RMST( $\tau$ )	NA
Risk, $\hat{r}$	$\Phi(\hat{r})$	$\hat{r}$	NA	NA
Time, $\hat{y}$	Distr( $\hat{y}, \sigma$ )	$-\hat{y}$	$\hat{y}$	NA
PI, $\hat{\eta}$	$\phi(\hat{\eta}, \hat{y})$	$\hat{\eta}$ or $-\hat{\eta}$	NA	$\hat{\eta}$



Figure 5.1: Illustration of prediction types. Survival time predictions (top-left) are limited to domains where uncertainty is less important. Survival probabilities (top-right) are shared with patients as ‘T-year survival probabilities’. Relative risk predictions (bottom) are used for triage and resource allocation.

## 5.1 Predicting Distributions

Predicting a survival distribution means predicting the probability of an individual surviving over time from 0 to  $\infty$ . Ideally one would make predictions over the continuous  $\mathbb{R}_{\geq 0}$ , however, in practice it is more common for predictions to be made over  $\mathbb{N}_0$ . This is due to the majority of models using discrete non-parametric estimators to create a distribution prediction. Distributional prediction can, in theory, target any of the distribution defining functions introduced in Section 3.1, but predicting  $S(t)$  and/or  $h(t)$  is most common. This is a *probabilistic survival task* as uncertainty is explicit in the prediction. Mathematically, the task is defined by  $g : \mathcal{X} \rightarrow \mathcal{S}$  where  $\mathcal{S} \subseteq \text{Distr}(\mathbb{R}_{\geq 0})$  is a set of distributions on  $\mathbb{R}_{\geq 0}$ .

Practically, especially in healthcare, survival distribution predictions are often used to estimate ‘ $\tau$ -year survival probabilities’, which is the probability of survival at a given point in time. Therefore a clinician is not likely to display a survival curve to a patient, but may use their individual features to compute probabilities of survival at key time-points (often, 5, 10 years) – an example of this in use is the PREDICT breast cancer tool (Candido dos Reis et al. 2017). In other contexts, such as engineering, survival distributions might be used to establish thresholds for replacing components. For example, one could use a survival model to estimate the reliability of a plane engine over time, a threshold could be set such that the engine will be replaced when  $\hat{S}(t) < 0.6$ .

Predicting  $\tau$ -year survival probabilities is often confused with a classification problem, which make probabilities for one or more events occurring at a fixed point in time. However, a classification model would be unable to use any observations that were censored before the time of interest and discarding these observations would bias any results.

Another potential confusion is conflating predicting survival distributions with density estimation. Density estimation fits a distribution to the *group*, whereas survival distribution predictions fit distributions to *individuals*. In reality, in the single-event setting, it can be difficult to wrap one’s head around the notion of an individual survival distribution as events occur exactly once at a fixed point in time. In theory, for an observation with event time  $t_i$ , the perfect model would predict guaranteed survival before  $t_i$  followed by guaranteed failure on and after this time, which is the Heaviside function:  $\hat{S}(\tau) = 1$  if  $\tau < t_i$  and  $\hat{S}(\tau) = 0$  otherwise. In practice, this is impossible and would massively overfit the training data and as such all survival models are estimations of this theoretical ideal (Figure 5.2).

## 5.2 Predicting Risks

Predicting risks is perhaps the most common survival problem and is defined as predicting a continuous rank for an individual’s relative risk of experiencing the event. Therefore this might also be known as a ‘ranking’ problem or predicting ‘continuous rankings’. In machine learning terms, this task is the problem of estimating,  $g : \mathcal{X} \rightarrow \mathbb{R}$ .

Interpretation of these rankings is more complex than might be imagined as the meaning differs between models, parametrizations, and even software implementation. To be consistent, in this book a larger risk value *always* corresponds to a higher risk of an event and lower values corresponding to lower risk.

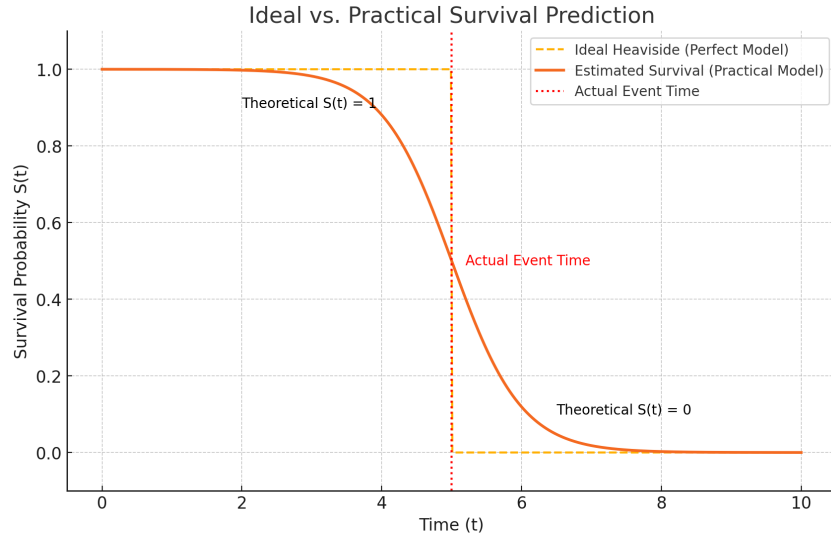


Figure 5.2: Comparing the true event time to theoretical and practical estimates of a survival function. The dashed line represents the theoretical ideal Heaviside function, the solid line represents a realistic estimate and the vertical dotted line is when the event actually takes place.

A potential confusion that should be avoided is conflating this risk prediction with a prediction of absolute risk. Risk predictions are specifically *relative risks*, which means the risks are only comparable to other observations within the same sample. For example, given three subjects,  $\{i, j, k\}$ , a risk prediction may be  $\{0.5, 10, 0.1\}$  respectively. From these predictions, two primary types of conclusion can be drawn.

1) Conclusions comparing subjects:

- The corresponding ranks for  $i, j, k$  are 2, 3, 1;
- $k$  is at the least risk and  $j$  is at the highest risk;
- The risk of  $i$  is slightly higher than that of  $k$  but  $j$ 's risk is considerably higher than both the others.

2) Conclusions comparing risk groups:

- Thresholding risks at 0.4 means  $k$  is at a low-risk but  $i$  and  $j$  are high-risk.
- Thresholding risks at 1.0 means  $i$  and  $k$  are low-risk but  $j$  is high-risk.

Whilst many important conclusions can be drawn from these predictions, the values themselves have no meaning when not compared to other individuals. Similarly, the values have no meanings across research, even if observation  $k$  is at low-risk according to this sample, it may be high risk compared to another.

### 5.2.1 Distributions and risks

In general it is not possible to easily convert a risk prediction to a distribution. However, this may be possible if the risk prediction corresponds to a particular model form. This is discussed further in Section 5.4.

More common is transformation from a distribution to a risk. Theoretically the simplest way to do so would be to take the mean or median of the distribution, however as will be discussed in detail in Section 5.3 this is practically difficult. Instead, a stable approach is to use the ‘ensemble’ or ‘expected mortality’ to create a measure of risk (B. H. Ishwaran et al. 2008). The expected mortality is defined by

$$\sum_t -\log(\hat{S}_i(t)) = \sum_t \hat{H}_i(t)$$

and is closely related to the calibration measure defined in Section 8.1.2. This represents the expected number of events for individuals with similar characteristics to  $i$ . This is a measure of risk as a larger value indicates that among individuals with a similar profile, there is expected to be a larger number of events and therefore have greater risk than those with smaller ensemble mortality. For example, say for an individual,  $i$ , we have:  $(t, \hat{S}_i(t)) = (0, 1), (1, 0.8), (2, 0.4), (3, 0.15)$  then,  $(t, \hat{H}_i(t)) = (0, 0), (1, 0.10), (2, 0.40), (3, 0.82)$  and their relative risk prediction is  $\sum_t \hat{H}_i(t) = 0 + 0.10 + 0.4 + 0.82 = 1.32$ .

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### 5.3 Predicting Survival Times

Predicting a time-to-event is the problem of understanding exactly when an individual will experience an event. Mathematically, the problem is the task of estimating  $g : \mathcal{X} \rightarrow \mathbb{R}_{>0}$ , that is, predicting a single value over  $[0, \infty]$ .

For practical purposes, the expected time-to-event would be the ideal prediction type as it is easy to interpret and communicate. However, such predictions are uncommon as the training data is never fully observed and hence training models is inherently a probabilistic process, with uncertainty being captured from the moment data is censored. In contrast to regression, in which a model could theoretically optimize a performance measure (which can occur when a model is overfit to training data), this is guaranteed not to happen in survival analysis due to the presence of censoring. In the presence of censoring, deterministic predictions may convey a false sense of certainty, as they do not reflect the underlying uncertainty. Hence it is more common to make a probabilistic prediction that can then be reduced to a deterministic one.

#### 5.3.1 Times and risks

Converting a time-to-event prediction to a risk prediction is trivial as the former is a special case of the latter. An individual with a longer survival time will have a lower overall risk: if  $\hat{y}_i, \hat{y}_j$  and  $\hat{r}_i, \hat{r}_j$  are survival time and ranking predictions for subjects  $i$  and  $j$  respectively, then  $\hat{y}_i > \hat{y}_j \Rightarrow \hat{r}_i < \hat{r}_j$ . It is not possible to make the transformation in the opposite direction without making significant assumptions as risk predictions are usually abstract quantities that rarely map to realistic survival times.

#### 5.3.2 Times and distributions

Moving from a survival time to distribution prediction is rare given the reasons outlined above. Theoretically one *could* make a prediction for the expected survival time,  $\hat{y}$  and then assume a particular distributional form. For example, one could assume  $\text{TruncatedNormal}(\hat{y}, \sigma, a =$

$0, b = \infty)$  where  $\hat{y}$  is the predicted expected survival time,  $\sigma$  is a parameter representing variance to be estimated or assumed, and  $\{a, b\}$  is the distribution support. This method clearly has drawbacks given the number of required assumptions and as such is not commonly seen in practice.

In the other direction, it is common to reduce a distribution prediction to a survival time prediction by *attempting* to compute the mean or median of the distribution. When there is no censoring, one can calculate the expectation from the predicted survival function using the ‘Darth Vader rule’ (Muldowney, Ostaszewski, and Wojdowski 2012):

$$\mathbb{E}[Y] = \int_0^\infty S_Y(y) dy \quad (5.1)$$

However, this rule is rarely usable in practice as censoring results in estimated survival distributions being ‘improper’. A valid probability distribution for a random variable  $Y$  satisfies:  $\int f_Y = 1$ ,  $S_Y(0) = 1$  and  $S_Y(\infty) = 0$ . This last condition is often violated in survival distribution predictions, which occurs because many algorithms create predictions using the Kaplan-Meier estimator. Recall from Section 3.3 that the estimator is defined as:

$$\hat{S}_{KM}(\tau) = \prod_{k:t_{(k)} \leq \tau} \left(1 - \frac{d_{t_{(k)}}}{n_{t_{(k)}}}\right)$$

This only reaches zero if every individual at risk at the last observed time-point experiences the event:  $d_{t_{(k)}} = n_{t_{(k)}}$ . In practice, due to administrative censoring, there will almost always be censoring at the final time-point. As a result,  $d_{t_{(k)}} < n_{t_{(k)}}$  and  $\hat{S}(\infty) > 0$ . Heuristics have been proposed to address this including linear extrapolation to zero, or dropping the curve to zero at the final time-point, however these can introduce significant bias in the estimated survival time R. Sonabend, Bender, and Vollmer (2022). Another possibility is to instead report the median survival time, but this is only defined if the survival curves drop below 0.5 within the observed period, which is not guaranteed (Haider et al. 2020).

One alternative is to estimate the *restricted mean survival time* (RMST) (K. Han and Jung 2022; Andersen, Hansen, and Klein 2004). In contrast to Equation 5.1, which integrates the survival curve over the entire time axis, the RMST places an upper-bound on the integral:

$$\text{RMST}(\tau) = \int_0^\tau S_Y(y) dy \quad (5.2)$$

Clearly then  $\text{RMST}(\infty) = \mathbb{E}[Y]$  by Equation 5.1. Whereas Equation 5.1 represents the average survival over  $[0, \infty)$ , Equation 5.2 represents the average survival time up to  $\tau$ . Equivalently, the RMST is the average amount of time each individual spends without experiencing the event up to  $\tau$ . The RMST treats all events happening after  $\tau$  as if they happened at  $\tau$ . Hence, it is a truncated expectation, estimating:  $\mathbb{E}[\min(Y, \tau)]$

The RMST can have several uses when comparing groups but here its relevant for its ability to give a meaningful lower-bound estimate of  $\mathbb{E}[Y]$  in the presence of censoring. For example, say individuals are observed over times  $[0, 100]$  and administrative censoring is present in the data so that  $\hat{S}(\infty) > 0$  and  $\mathbb{E}[Y]$  cannot be reliably computed. However, one can compute  $\text{RMST}(100)$  which gives a lower-bound estimate for the expected survival:  $\text{RMST}(100) \leq \mathbb{E}[Y]$ . This avoids assumptions beyond  $\tau = 100$  and offers an interpretable prediction: “the average survival time is at least  $\text{RMST}(100)$ ” (Figure 5.3). The RMST

avoids the pitfalls of computing the mean and median by remaining valid even when the predicted distribution is improper, provided that  $\tau$  is chosen within the range of observed follow-up times.

As a worked example, say for an individual,  $i$ , we have:  $(t, \hat{S}_i(t)) = (0, 1), (1, 0.8), (2, 0.4), (3, 0.15)$  then (using a Riemann sum approximation for the integral),  $\text{RMST}(4) \approx 1 + 0.8 + 0.4 + 0.15 = 2.35$  whereas  $\text{RMST}(3) \approx 1 + 0.8 + 0.4 = 2.2$ ; the former is a better estimate for  $\mathbb{E}[Y]$ .

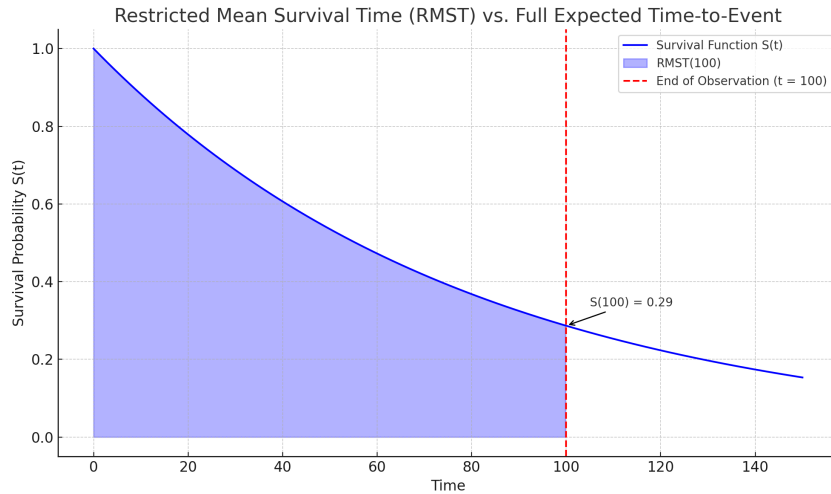


Figure 5.3: Illustrating the RMST at the final observed time-point. The RMST is the area under the survival curve up to the point of interest. The red vertical line represents the final observed time-point, the survival curve (blue) after that point is unknown.

## 5.4 Prognostic Index Predictions

In medical terminology (often used in survival analysis), a prognostic index is a tool that predicts outcomes based on risk factors. Given covariates,  $\mathbf{X} \in \mathbb{R}^{n \times p}$ , and coefficients,  $\boldsymbol{\beta} \in \mathbb{R}^p$ , the *linear predictor* is defined as  $\boldsymbol{\eta} := \mathbf{X}\boldsymbol{\beta}$ . Applying some function  $g$ , which could simply be the identity function  $g(x) = x$ , yields a *prognostic index*,  $g(\boldsymbol{\eta})$ . A prognostic index can serve several purposes, including:

1. Scaling or normalization: simple functions to scale the linear predictor can better support interpretation and visualisation;
2. Capturing non-linear effects: for example the Cox PH model (Chapter 12) applies the transformation  $g(\boldsymbol{\eta}) = \exp(\boldsymbol{\eta})$  to capture more complex relationships between features and outcomes;
3. Aiding in interpretability: in some cases this could simply be  $g(\boldsymbol{\eta}) = -\boldsymbol{\eta}$  to ensure the ‘higher value implies higher risk’ interpretation.

### 5.4.1 Prognostic index, risks, and times

A prognostic index is a special case of the survival ranking task, assuming that there is a one-to-one mapping between the prediction and expected survival times, i.e., as long as there is an intuitive relationship that explains how the prognostic index maps to survival time. For example, one could define  $g(\eta) = 1$  if  $\eta < 5$  or 0 otherwise however this would prevent the prognostic index being used as a ranking method and it would defeat the point of a prognostic index.

As with a risk prediction, there is no general relationship between the prognostic index and survival time. However, there are cases where there may be a clear relationship between the two. For example, one can theoretically estimate a survival time from the linear predictor used in an accelerated failure time model (Section 12.1).

### 5.4.2 Prognostic index and distributions

A general prognostic index cannot be generated from a distribution, however the reverse transformation is very common in survival models. The vast majority of survival models are composed from a group-wise survival time estimate and an estimation of an individual's prognostic index. This will be seen in detail in future chapters (particularly in Chapter 15 and Chapter 16) but in general it is very common for models to assume data follows either proportional hazards or accelerated failure time (Chapter 12). In this case models are formed of two components:

1. A non-parametric estimator (Section 3.3) that estimates the baseline hazard,  $\hat{h}_0$ , or baseline survival function,  $\hat{S}_0$ . Discussed further in Chapter 12, these are group-wise estimates that apply to the entire sample.
2. A sophisticated algorithm that estimates the individual linear prediction,  $\hat{\eta}$ .

These are composed as either:

1.  $\hat{h}_0(t) \exp(\hat{\eta})$  if proportional hazards is assumed; or
2.  $\hat{h}_0(\exp(-\hat{\eta})t) \exp(-\hat{\eta})$  for accelerated failure time models.

## 5.5 Beyond single-event

In Chapter 4, competing risks and multi-state models were introduced with a focus on estimating probability distributions.

In the competing risks setting, ‘survival time’ is ill-defined, it is ambiguous whether this refers to the time until a specific event or until any event takes place. In principle, one could apply the RMSE approach to the all-cause survival function, though this is uncommon in practice. Analogous to the single-event setting, a cause-specific prognostic index can be derived from a cause-specific cumulative incidence function, and vice versa. One might also consider transformations based on all-cause risk, but again such approaches are rarely used in applied settings.

In multi-state models, the primary prediction targets are transition probabilities between states over time. Unlike standard survival settings, there is no single notion of a ‘survival

time'. Instead, one can estimate the sojourn time, which is the expected time spent in a given state, using the estimated transition probabilities. This is particularly well-defined in Markov or semi-Markov models, where sojourn times follow from standard stochastic process theory. These derivations are beyond the scope of this book; for further detail, see for example ([ibe2013markov?](#)).

## 5.6 Conclusion

### Key takeaways

- Survival prediction tasks fall into three categories: probabilistic (predicting survival distributions), deterministic (predicting survival times), and ranking (predicting relative risks or prognostic indices).
- Prediction types are not directly interchangeable; transforming between them requires assumptions and often changes what is being estimated.
- Distribution predictions are the most general, enabling derivation of other forms.
- Survival times are intuitive but prone to overconfidence in censored data.
- Ranking predictions are the most common but their interpretation can be complicated and their utility is limited to within-sample comparisons.

### Further reading

- R. Sonabend, Bender, and Vollmer (2022) reviews methods to transform distribution and ranking/survival time predictions, advantages and disadvantages of methods is discussed in brief in the appendix of Haider et al. (2020).
- For more on the RMST see Uno et al. (2014), K. Han and Jung (2022), and Patrick Royston and Parmar (2011).

## **Part II**

# **Evaluation**



# 6

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## What are Survival Measures?

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TODO (150-200 WORDS)

 Minor changes expected!

This page is a work in progress and minor changes will be made over time.

In this part of the book we discuss one of the most important parts of the machine learning workflow, model evaluation (Foss and Kotthoff 2024). In the next few chapters we will discuss different metrics that can be used to measure a model's performance but before that we will just briefly discuss why model evaluation is so important.

In the simplest case, without evaluation there is no way to know if predictions from a trained machine learning model are any good. Whether one uses a simple Kaplan-Meier estimator, a complex neural network, or anything in between, there is no guarantee any of these methods will actually make useful predictions for a given dataset. This could be because the dataset is inherently difficult for any model to be trained on, perhaps because it is very ‘noisy’, or because a model is simply ill-suited to the task, for example using a Cox Proportional Hazards model when its key assumptions are violated. Evaluation is therefore crucial to trusting any predictions made from a model.

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### 6.1 Survival Measures

Evaluation can be used to assess in-sample and out-of-sample performance.

In-sample evaluation measures the quality of a model's ‘fit’ to data, i.e., whether the model has accurately captured relationships in the training data. However, in-sample measures often cannot be applied to complex machine learning models so this part of the book omits these measures. Readers who are interested in this are directed to Collett (2014) and Hosmer Jr, Lemeshow, and May (2011) for discussion on residuals; Choodari-Oskooei, Royston, and Parmar (2012), Kent and O'Quigley (1988) and Patrick Royston and Sauerbrei (2004) for  $R^2$  type measures; and finally Volinsky and Raftery (2000), HURVICH and TSAI (1989), and Liang and Zou (2008) for information criterion measures.

Out-of-sample measures evaluate the quality of model predictions on new and previously unseen (by the model) data. By following established statistical methods for evaluation, and ensuring that robust resampling methods are used (James et al. 2013), evaluation provides a method for estimating the ‘generalisation error’, which is the expected model performance

on new datasets. This is an important concept as it provides confidence about future model performance without limiting results to the current data. Survival measures are classified into measures of:

- Discrimination (aka ‘separation’) – A model’s discriminatory power refers to how well it separates observations that are at a higher or lower risk of event. For example, a model with good discrimination will predict that (at a given time) a dead patient has a higher probability of being dead than an alive patient.
- Calibration – Calibration is a roughly defined concept (Collins et al. 2014; F. E. Harrell, Lee, and Mark 1996; Rahman et al. 2017; Van Houwelingen 2000) that generally refers to how well a model captures average relationships between predicted and observed values.
- Predictive Performance – A model is said to have good predictive performance (or sometimes ‘predictive accuracy’) if its predictions for new data are ‘close to’ the truth.

These measures could also be categorised into how they evaluate predictions. Discrimination measures compare predictions pairwise where pairs of observations are created and then the predictions for these pairs are compared within and across each other in some way. Calibration measures evaluate predictions holistically by looking at some ‘average’ performance across them to provide an idea of how well suited the model is to the data. Measures of predictive performance evaluate individual predictions and usually take the sample mean of these to estimate the generalisation error.

In the next few chapters we categorise measures by the type of survival prediction they evaluate, which is a more natural taxonomy for selecting measures, but we use the above categories when introducing each measure.

## 6.2 How are Models Evaluated?

As well as using measures to evaluate a model’s performance on a given dataset, evaluation can also be used to measure future performance, to compare and select models, and to tune internal processes. In most cases, models should not be trained/predicted/evaluated on their own, instead a number of simpler reference models should be simultaneously trained and evaluated on the same data, which is known as a ‘benchmark experiment’. This is especially important for survival models, as *all* survival measures depend on the censoring distribution and therefore cannot be interpreted out of context and without comparison to other models. Benchmark experiments are used to empirically compare models across the same data and measures, meaning that if one model outperforms another then that model can be selected for future experiments (though simpler models are preferred if the performance difference is marginal). A model is usually said to ‘outperform’ another if it has a lower generalisation error.

The process of model evaluation is dependent on the measure itself. Measures that are ‘decomposable’ (predictive performance measures) calculate scores for individual predictions and take the sample mean over all scores, on the other hand ‘aggregate’ measures (discrimination and calibration) return a single score over all predictions. The simplest method to estimate the generalisation error is ‘holdout’ resampling, where a dataset  $\mathcal{D}$  is split into non-overlapping subsets for training  $\mathcal{D}_{train}$  and testing  $\mathcal{D}_{test}$ . The model is trained on  $\mathcal{D}_{train}$  and predictions,  $\hat{\mathbf{y}}$  are made based on the features in  $\mathcal{D}_{test}$ . The model is evaluated by using a measure,  $L$ , to compare the predictions to the observed data in the test set,  $L(\mathbf{y}_{test}, \hat{\mathbf{y}})$ .

Where possible, (repeated) k-fold cross-validation (kCV) should be used for more robust estimation of the generalisation error and for model comparison. In kCV, the data is partitioned into  $k$  folds (often  $k$  is 5 or 10), which are non-overlapping subsets. A model is trained on  $k - 1$  folds and evaluated on the  $k$ th fold, this process is repeated until each of the  $k$  folds has acted as the test set exactly once, the computed loss from each iteration is averaged into the final loss, which provides a good estimate of the generalisation error.

For the rest of this part of the book we will introduce different survival measures, discuss their advantages and disadvantages, and in Chapter 11 we will provide some recommendations for choosing measures. We will not discuss the general process of model resampling or evaluation further but recommend Casalicchio and Burk (2024) to readers interested in this topic.



# 7

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## Discrimination Measures

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TODO (150-200 WORDS)

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The next measures discussed are ‘discrimination measures’, which evaluate how well models separate observations into different risk groups. A model is said to have good discrimination if it correctly predicts that one observation is at higher risk of the event of interest than another, where the prediction is ‘correct’ if the observation predicted to be at higher risk does indeed experience the event sooner.

In the survival setting, the ‘risk’ is taken to be the continuous ranking prediction. All discrimination measures are ranking measures, which means that the exact predicted value is irrelevant, only its relative ordering is required. For example given predictions {100, 2, 299.3}, only their rankings, {2, 1, 3}, are used by measures of discrimination.

This chapter begins with time-independent measures (Section 7.1), which measure concordance between pairs of observations at a single observed time point. The next section focuses on time-dependent measures (Section 7.2), which are primarily AUC-type measures that evaluate discrimination over all possible unique time-points and integrate the results for a single metric.

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### 7.1 Time-Independent Measures

The simplest form of discrimination measures are concordance indices, which, in general, measure the proportion of cases in which the model correctly ranks a pair of observations according to their risk. These measures may be best understood in terms of two key definitions: ‘comparable’, and ‘concordant’.

**Definition 7.1** (Concordance). Let  $(i, j)$  be a pair of observations with outcomes  $\{(t_i, \delta_i), (t_j, \delta_j)\}$  and let  $r_i, r_j \in \mathbb{R}$  be their respective risk predictions. Then  $(i, j)$  are called (F. E. J. Harrell et al. 1984; F. E. Harrell, Califff, and Pryor 1982):

- *Comparable* if  $t_i < t_j$  and  $\delta_i = 1$ ;
- *Concordant* if  $r_i > r_j$ .

Note that this book defines risk rankings such that a higher value implies higher risk of event and thus lower expected survival time (`?@sec-surv-set-types`), hence a pair is concordant if  $\mathbb{I}(t_i < t_j, r_i > r_j)$ . Other sources may instead assume that higher values imply lower risk of event and hence a pair would be concordant if  $\mathbb{I}(t_i < t_j, r_i < r_j)$ .

Concordance measures then estimate the probability of a pair being concordant, given that they are comparable:

$$P(r_i > r_j | t_i < t_j \cap \delta_i)$$

These measures are referred to as time *independent* when  $r_i, r_j$  is not a function of time as once the observations are organised into comparable pairs, the observed survival times can be ignored. The time-dependent case is covered in Section 7.2.1.

While various definitions of a ‘Concordance index’ (C-index) exist (discussed in the next section), they all represent a weighted proportion of the number of concordant pairs over the number of comparable pairs. As such, a C-index value will always be between [0, 1] with 1 indicating perfect separation, 0.5 indicating no separation, and 0 being separation in the ‘wrong direction’, i.e. all high risk observations being ranked lower than all low risk observations.

Concordance measures may either be reported as a value in [0, 1], a percentage, or as ‘discriminatory power’, which refers to the percentage improvement of a model’s discrimination above the baseline value of 0.5. For example, if a model has a concordance of 0.8 then its discriminatory power is  $(0.8 - 0.5)/0.5 = 60\%$ . This representation of discrimination provides more information by encoding the model’s improvement over some baseline although is often confused with reporting concordance as a percentage (e.g. reporting a concordance of 0.8 as 80%). In theory this representation could result in a negative value, however this would indicate that  $C < 0.5$ , which would indicate serious problems with the model that should be addressed before proceeding with further analysis. Representing measures as a percentage over a baseline is a common method to improve measure interpretability and closely relates to the ERV representation of scoring rules (Section 9.4).

### 7.1.1 Concordance Indices

Common concordance indices in survival analysis can be expressed as a general measure:

Let  $\hat{\mathbf{r}} = (\hat{r}_1 \ \hat{r}_2 \cdots \hat{r}_m)^\top$  be predicted risks,  $(\mathbf{t}, \boldsymbol{\delta}) = ((t_1, \delta_1) \ (t_2, \delta_2) \cdots (t_m, \delta_m))^\top$  be observed outcomes, let  $W$  be some weighting function, and let  $\tau$  be a cut-off time. Then, the time-independent (‘ind’) *survival concordance index* is defined by,

$$C_{ind}(\hat{\mathbf{r}}, \mathbf{t}, \boldsymbol{\delta} | \tau) = \frac{\sum_{i \neq j} W(t_i) \mathbb{I}(t_i < t_j, \hat{r}_i > \hat{r}_j, t_i < \tau) \delta_i}{\sum_{i \neq j} W(t_i) \mathbb{I}(t_i < t_j, t_i < \tau) \delta_i}$$

The choice of  $W$  specifies a particular variation of the c-index (see below). The use of the cut-off  $\tau$  mitigates against decreased sample size (and therefore high variance) over time due to the removal of censored observations (see Figure 7.1)). For  $\tau$  to be comparable across datasets, a common choice would be to set  $\tau$  as the time at which 80%, or perhaps 90% of the data have been censored or experienced the event.

There are multiple methods for dealing with tied predictions and times. Strictly, tied times are incomparable given the definition of ‘comparable’ given above and hence are usually

ignored in the numerator. On the other hand, ties in the prediction are more problematic but a common method is to set a value of 0.5 for observations when  $r_i = r_j$  (Therneau and Atkinson 2020). Specific concordance indices can be constructed by assigning a weighting scheme for  $W$  which generally depends on the Kaplan-Meier estimate of the survival function of the censoring distribution fit on training data,  $\hat{G}_{KM}$ , or the Kaplan-Meier estimate for the survival function of the survival distribution fit on training data,  $\hat{S}_{KM}$ , or both. Measures that use  $\hat{G}_{KM}$  are referred to as Inverse Probability of Censoring Weighted (IPCW) measures as the estimated censoring distribution is utilised to weight the measure in order to compensate for removed censored observations. This is visualised in Figure 7.1 where  $\hat{G}_{KM}$ ,  $\hat{G}_{KM}^{-2}$ , and  $\hat{S}_{KM}$  are computed based on the `whas` dataset (Hosmer Jr, Lemeshow, and May 2011).

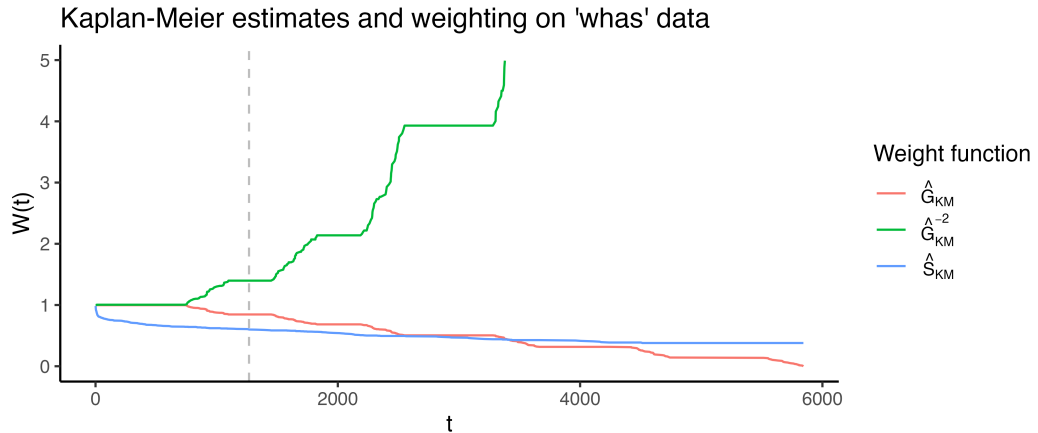


Figure 7.1: Weighting functions obtained on the `whas` dataset. x-axis is follow-up time. y-axis is outputs from one of three weighting functions:  $\hat{G}_{KM}$ , survival function based on the censoring distribution of the `whas` dataset (red), and  $\hat{G}_{KM}^{-2}$  (green),  $\hat{S}_{KM}$ , marginal survival function based on original `whas` dataset (blue). The vertical gray line at  $t = \tau = 1267$  represents the point at which  $\hat{G}(t) < 0.6$ .

The following weights have been proposed for the concordance index:

- $W(t_i) = 1$ : Harrell's concordance index,  $C_H$  (F. E. J. Harrell et al. 1984; F. E. Harrell, Calif, and Pryor 1982), which is widely accepted to be the most common survival measure and imposes no weighting on the definition of concordance. The original measure given by Harrell has no cut-off,  $\tau = \infty$ , however applying a cut-off is now more widely accepted in practice.
- $W(t_i) = [\hat{G}_{KM}(t_i)]^{-2}$ : Uno's C,  $C_U$  (Uno et al. 2011).
- $W(t_i) = [\hat{G}_{KM}(t_i)]^{-1}$
- $W(t_i) = \hat{S}_{KM}(t_i)$
- $W(t_i) = \hat{S}_{KM}(t_i)/\hat{G}_{KM}(t_i)$

All methods assume that censoring is conditionally-independent of the event given the features (`?@sec-surv-set-cens`), otherwise weighting by  $\hat{S}_{KM}$  or  $\hat{G}_{KM}$  would not be applicable. It is assumed here that  $\hat{S}_{KM}$  and  $\hat{G}_{KM}$  are estimated on the training data and not the testing data (though the latter may be seen in some implementations, e.g. Therneau (2015)).

### 7.1.2 Choosing a C-index

With multiple choices of weighting available, choosing a specific measure might seem daunting. Matters are only made worse by debate in the literature, reflecting uncertainty in measure choice and interpretation. In practice, when a suitable cut-off  $\tau$  is chosen, all these weightings perform very similarly (Rahman et al. 2017; Schmid and Potapov 2012). For example, Table 7.1 uses the `whas` data again to compare Harrell's C with measures that include IPCW weighting, when no cutoff is applied (top row) and when a cutoff is applied when  $\hat{G}(t) = 0.6$  (grey line in Figure 7.1). The results are almost identical when the cutoff is applied but still not massively different without the cutoff.

Table 7.1: Comparing C-index measures (calculated on the `whas` dataset using a Cox model with three-fold cross-validation) with no cut-off (top) and a cut-off when  $\hat{G}(t) = 0.6$  (bottom). First column is Harrell's C, second is the weighting  $1/\hat{G}(t)$ , third is Uno's C.

	$W = 1$	$W = G^{-1}$	$W = G^{-2}$
$\tau = \infty$	0.74	0.73	0.71
$\tau = 1267$	0.76	0.75	0.75

On the other hand, if a poor choice is selected for  $\tau$  (cutting off too late) then IPCW measures can be highly unstable (Rahman et al. 2017), for example the variance of Uno's C drastically increases with increased censoring (Schmid and Potapov 2012).

In practice, all C-index metrics provide an intuitive measure of discrimination and as such the choice of C-index is less important than the transparency in reporting. ‘C-hacking’ (R. Sonabend, Bender, and Vollmer 2022) is the deliberate, unethical procedure of calculating multiple C-indices and to selectively report one or more results to promote a particular model or result, whilst ignoring any negative findings. For example, calculating Harrell's C and Uno's C but only reporting the measure that shows a particular model of interest is better than another (even if the other metric shows the reverse effect). To avoid ‘C-hacking’:

- i) the choice of C-index should be made before experiments have begun and the choice of C-index should be clearly reported;
- ii) when ranking predictions are composed (Chapter 18) from distribution predictions, the composition method should be chosen and clearly described before experiments have begun.

As the C-index is highly dependent on censoring within a dataset, C-index values between experiments are not directly comparable, instead comparisons are limited to comparing model rankings, for example conclusions such as “model A outperformed model B with respect to Harrell's C in this experiment”.

## 7.2 Time-Dependent Measures

In the time-dependent case, where the metrics are computed based on specific survival times, the majority of measures are based on the Area Under the Curve, with one exception which is a simpler concordance index.

### 7.2.1 Concordance Indices

In contrast to the measures described above, Antolini's C (Antolini, Boracchi, and Biganzoli 2005) provides a time-dependent ('dep') formula for the concordance index. The definition of 'comparable' is the same for Antolini's C, however, concordance is now determined using the individual predicted survival probabilities calculated at the smaller event time in the pair:

$$P(\hat{S}_i(t_i) < \hat{S}_j(t_i) | t_i < t_j \cap \delta_i)$$

Note that observations are concordant when  $\hat{S}_i(t_i) < \hat{S}_j(t_i)$  as at the time  $t_i$ , observation  $i$  has experienced the event and observation  $j$  has not, hence the expected survival probability for  $\hat{S}_i(t_i)$  should be as close to 0 as possible (noting inherent randomness may prevent the perfect  $\hat{S}_i(t_i) = 0$  prediction) but otherwise should be less than  $\hat{S}_j(t_i)$  as  $j$  is still 'alive'. Once again this probability is estimated with a metric that could include a cut-off and different weighting schemes (though this is not included in Antolini's original definition):

$$C_{dep}(\hat{\mathbf{S}}, \mathbf{t}, \boldsymbol{\delta} | \tau) = \frac{\sum_{i \neq j} W(t_i) \mathbb{I}(t_i < t_j, \hat{S}_i(t_i) < \hat{S}_j(t_i), t_i < \tau) \delta_i}{\sum_{i \neq j} W(t_i) \mathbb{I}(t_i < t_j, t_i < \tau) \delta_i}$$

where  $\hat{\mathbf{S}} = (\hat{S}_1 \ \hat{S}_2 \cdots \hat{S}_m)^\top$ .

Antolini's C provides an intuitive method to evaluate the discrimination of a model based on distribution predictions without depending on compositions to ranking predictions.

### 7.2.2 Area Under the Curve

#### Warning

We are still discussing how to structure and write this section so the contents are all subject to change. The text below is 'correct' but we want to add more detail about estimation of AUC so the book can be more practical, otherwise we may remove the section completely, let us know your thoughts about what you'd like to see here!

AUC, or AUROC, measures calculate the Area Under the Receiver Operating Characteristic (ROC) Curve, which is a plot of the *sensitivity* (or true positive rate (TPR)) against  $1 - specificity$  (or true negative rate (TNR)) at varying thresholds (described below) for the predicted probability (or risk) of event. Figure 7.2 visualises ROC curves for two classification models. The blue line is a featureless baseline that has no discrimination. The red line is a decision tree with better discrimination as it comes closer to the top-left corner.

In a classification setting with no censoring, the AUC has the same interpretation as Harrell's C (Uno et al. 2011). AUC measures for survival analysis were developed to provide a time-dependent measure of discriminatory ability (Patrick J. Heagerty, Lumley, and Pepe 2000). In a survival setting it can reasonably be expected for a model to perform differently over time and therefore time-dependent measures are advantageous. Computation of AUC estimators is complex and as such there are limited accessible metrics available off-shelf. There is limited evidence of these estimators used in the literature, hence discussion of these measures is kept brief.

The AUC, TPR, and TNR are derived from the *confusion matrix* in a binary classification setting. Let  $b_i, \hat{b}_i \in \{0, 1\}$  be the true and predicted binary outcomes respectively for some

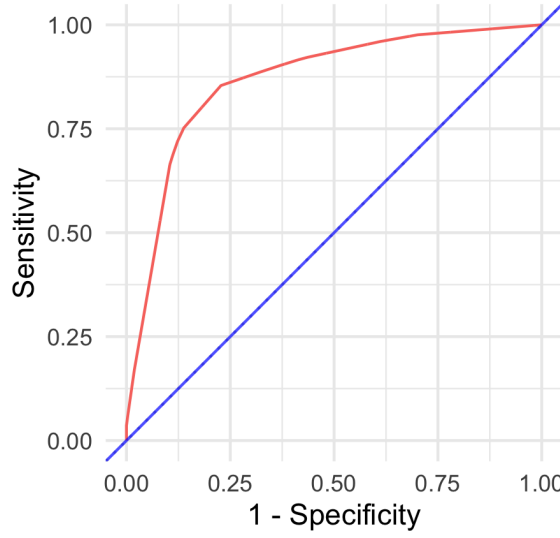


Figure 7.2: ROC Curves for a classification example. Red is a decision tree with good discrimination as it ‘hugs’ the top-left corner. Blue is a featureless baseline with no discrimination as it sits on  $y = x$ .

observation  $i$ . The confusion matrix is then given by:

		$b_i = 1$	$b_i = 0$
$\hat{b}_i = 1$	TP	FP	
	FN	TN	

where  $TN := \sum_i \mathbb{I}(b_i = 0, \hat{b}_i = 0)$  is the number of true negatives,  $TP := \sum_i \mathbb{I}(b_i = 1, \hat{b}_i = 1)$  is the number true positives,  $FP := \sum_i \mathbb{I}(b_i = 0, \hat{b}_i = 1)$  is the number of false positives, and  $FN := \sum_i \mathbb{I}(b_i = 1, \hat{b}_i = 0)$  is the number of false negatives. From these are derived

$$\begin{aligned} TPR &:= \frac{TP}{TP + FN} \\ TNR &:= \frac{TN}{TN + FP} \end{aligned}$$

In classification, a probabilistic prediction of an event can be *thresholded* to obtain a deterministic prediction. For a predicted  $\hat{p} := \hat{P}(b = 1)$ , and threshold  $\alpha$ , the thresholded binary prediction is  $\hat{b} := \mathbb{I}(\hat{p} > \alpha)$ . This is achieved in survival analysis by thresholding the linear predictor at a given time for different values of the threshold and different values of the time. All measures of TPR, TNR and AUC are in the range  $[0, 1]$  with larger values preferred.

Weighting the linear predictor was proposed by Uno *et al.* (2007) (Uno et al. 2007) and provides a method for estimating TPR and TNR via

$$TPR_U(\hat{\eta}, \mathbf{t}, \boldsymbol{\delta} | \tau, \alpha) = \frac{\sum_{i=1}^m \delta_i \mathbb{I}(k(\hat{\eta}_i) > \alpha, t_i \leq \tau) [\hat{G}_{KM}(t_i)]^{-1}}{\sum_{i=1}^m \delta_i \mathbb{I}(t_i \leq \tau) [\hat{G}_{KM}(t_i)]^{-1}}$$

and

$$TNR_U(\hat{\eta}, \mathbf{t} | \tau, \alpha) \mapsto \frac{\sum_{i=1}^m \mathbb{I}(k(\hat{\eta}_i) \leq \alpha, t_i > \tau)}{\sum_{i=1}^m \mathbb{I}(t_i > \tau)}$$

where  $\hat{\eta} = (\hat{\eta}_1 \ \hat{\eta}_2 \ \cdots \ \hat{\eta}_m)^\top$  is a vector of predicted linear predictors,  $\tau$  is the time at which to evaluate the measure,  $\alpha$  is a cut-off for the linear predictor, and  $k$  is a known, strictly increasing, differentiable function.  $k$  is chosen depending on the model choice, for example if the fitted model is PH then  $k(x) = 1 - \exp(-\exp(x))$  (Uno et al. 2007). Similarities can be drawn between these equations and Uno's concordance index, in particular the use of IPCW. Censoring is again assumed to be at least random once conditioned on features. Plotting  $TPR_U$  against  $1 - TNR_U$  for varying values of  $\alpha$  provides the ROC.

The second method, which appears to be more prominent in the literature, is derived from Heagerty and Zheng (2005) (Patrick J. Heagerty and Zheng 2005). They define four distinct classes, in which observations are split into controls and cases.

An observation is a *case* at a given time-point if they are dead, otherwise they are a *control*. These definitions imply that all observations begin as controls and (hypothetically) become cases over time. Cases are then split into *incident* or *cumulative* and controls are split into *static* or *dynamic*. The choice between modelling static or dynamic controls is dependent on the question of interest. Modelling static controls implies that a 'subject does not change disease status' (Patrick J. Heagerty and Zheng 2005), and few methods have been developed for this setting (Kamarudin, Cox, and Kolamunnage-Dona 2017), as such the focus here is on *dynamic* controls. The incident/cumulative cases choice is discussed in more detail below.<sup>1</sup>

The TNR for dynamic cases is defined as

$$TNR_D(\hat{\mathbf{r}}, N | \alpha, \tau) = P(\hat{r}_i \leq \alpha | N_i(\tau) = 0)$$

where  $\hat{\mathbf{r}} = (\hat{r}_1 \ \hat{r}_2 \ \cdots \ \hat{r}_n)^\top$  is some deterministic prediction and  $N(\tau)$  is a count of the number of events in  $[0, \tau]$ . Heagerty and Zheng further specify  $y$  to be the predicted linear predictor  $\hat{\eta}$ . Cumulative/dynamic and incident/dynamic measures are available in software packages 'off-shelf', these are respectively defined by

$$TPR_C(\hat{\mathbf{r}}, N | \alpha, \tau) = P(\hat{r}_i > \alpha | N_i(\tau) = 1)$$

and

$$TPR_I(\hat{\mathbf{r}}, N | \alpha, \tau) = P(\hat{r}_i > \alpha | dN_i(\tau) = 1)$$

where  $dN_i(\tau) = N_i(\tau) - N_i(\tau-)$ . Practical estimation of these quantities is not discussed here.

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<sup>1</sup>All measures discussed in this section evaluate model discrimination from 'markers', which may be a *predictive* marker (model predictions) or a *prognostic* marker (a single covariate). This section always defines a marker as a ranking prediction, which is valid for all measures discussed here with the exception of one given at the end.

The choice between the incident/dynamic (I/D) and cumulative/dynamic (C/D) measures primarily relates to the use-case. The C/D measures are preferred if a specific time-point is of interest (Patrick J. Heagerty and Zheng 2005) and is implemented in several applications for this purpose (Kamarudin, Cox, and Kolamunnage-Dona 2017). The I/D measures are preferred when the true survival time is known and discrimination is desired at the given event time (Patrick J. Heagerty and Zheng 2005).

Defining a time-specific AUC is now possible with

$$AUC(\hat{\mathbf{r}}, N|\tau) = \int_0^1 TPR(\hat{\mathbf{r}}, N|1 - TNR^{-1}(p|\tau), \tau) dp$$

Finally, integrating over all time-points produces a time-dependent AUC and as usual a cut-off is applied for the upper limit,

$$AUC^*(\hat{\mathbf{r}}, N|\tau^*) = \int_0^{\tau^*} AUC(\hat{\mathbf{r}}, N|\tau) \frac{2\hat{p}_{KM}(\tau)\hat{S}_{KM}(\tau)}{1 - \hat{S}_{KM}^2(\tau^*)} d\tau$$

where  $\hat{S}_{KM}, \hat{p}_{KM}$  are survival and mass functions estimated with a Kaplan-Meier model on training data.

Since Heagerty and Zheng's paper, other methods for calculating the time-dependent AUC have been devised, including by Chambliss and Diao (Chambliss and Diao 2006), Song and Zhou (Song and Zhou 2008), and Hung and Chiang (Hung and Chiang 2010). These either stem from the Heagerty and Zheng paper or ignore the case/control distinction and derive the AUC via different estimation methods of TPR and TNR. Blanche *et al.* (2012) (Blanche, Latouche, and Viallon 2012) surveyed these and concluded "regarding the choice of the retained definition for cases and controls, no clear guidance has really emerged in the literature", but agree with Heagerty and Zeng on the use of C/D for clinical trials and I/D for 'pure' evaluation of the marker. Blanche *et al.* (2013) (Blanche, Dartigues, and Jacqmin-Gadda 2013) published a survey of C/D AUC measures with an emphasis on non-parametric estimators with marker-dependent censoring, including their own Conditional IPCW (CIPCW) AUC, which is not discussed further here as it cannot be used for evaluating predictions (R. E. B. Sonabend 2021).

Reviews of AUC measures have produced (sometimes markedly) different results (Blanche, Latouche, and Viallon 2012; Li, Greene, and Hu 2018; Kamarudin, Cox, and Kolamunnage-Dona 2017) with no clear consensus on how and when these measures should be used. The primary advantage of these measures is to extend discrimination metrics to be time-dependent. However, it is unclear how to interpret a threshold of a linear predictor and moreover if this is even the 'correct' quantity to threshold, especially when survival distribution predictions are the more natural object to evaluate over time.

# 8

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## Calibration Measures

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TODO (150-200 WORDS)

 Minor changes expected!

This page is a work in progress and minor changes will be made over time.

Calibration measures evaluate the ‘average’ quality of survival distribution predictions. This chapter is kept relatively short as the literature in this area is scarce (Rahman et al. 2017), this is likely due to the meaning of calibration being unclear in a survival context (Van Houwelingen 2000). However the meaning of calibration is better specified once specific metrics are introduced. As with other measure classes, only measures that can generalise beyond Cox PH models are included here but note that several calibration measures for re-calibrating PH models have been discussed in the literature (Demler, Paynter, and Cook 2015; Van Houwelingen 2000).

Calibration measures can be grouped (Andres et al. 2018) into those that evaluate distributions at a single time-point, ‘1-Calibration’ or ‘Point Calibration’ measures, and those that evaluate distributions at all time-points ‘distributional-calibration’ or ‘probabilistic calibration’ measures. A point-calibration measure will evaluate a function of the predicted distribution at a single time-point whereas a probabilistic measure evaluates the distribution over a range of time-points; in both cases the evaluated quantity is compared to the observed outcome,  $(t, \delta)$ .

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### 8.1 Point Calibration

Point calibration measures can be further divided into metrics that evaluate calibration at a single time-point (by reduction) and measures that evaluate an entire distribution by only considering the event time. The difference may sound subtle but it affects conclusions that can be drawn. In the first case, a calibration measure can only draw conclusions at that one time-point, whereas the second case can draw conclusions about the calibration of the entire distribution. This is the same caveat as using prediction error curves for scoring rules (Section 9.3).

### 8.1.1 Calibration by Reduction

Point calibration measures are implicitly reduction methods as they use classification methods to evaluate a full distribution based on a single point only (Chapter 18). For example, given a predicted survival function  $\hat{S}$ , one could calculate the survival function at a single time point,  $\hat{S}\tau$  and then use probabilistic classification calibration measures. Using this approach one may employ common calibration methods such as the Hosmer–Lemeshow test (Hosmer and Lemeshow 1980). Measuring calibration in this way can have significant drawbacks as a model may be well-calibrated at one time-point but poorly calibrated at all others (Haider et al. 2020). To mitigate this, one could perform the Hosmer–Lemeshow test (or other applicable tests) multiple times with multiple testing correction at many (or all possible) time points, however this would be less efficient and more difficult to interpret than other measures discussed in this chapter.

### 8.1.2 Houwelingen's $\alpha$

As opposed to evaluating distributions at one or more arbitrary time points, one could instead evaluate distribution predictions at meaningful times. van Houwelingen proposed several measures (Van Houwelingen 2000) for calibration but only one generalises to all probabilistic survival models, termed here ‘Houwelingen's  $\alpha$ ’. The measure assesses if the model correctly estimates the theoretical ‘true’ cumulative hazard function of the underlying data generating process,  $H = \hat{H}$ .

The statistic is derived by noting the closely related nature of survival analysis and counting processes, and exploiting the fact that the sum of the cumulative hazard function is an estimate for the number of events in a given time-period (Hosmer Jr, Lemeshow, and May 2011). As this result may seem surprising, below is a short experiment using R that demonstrates how the sum of the cumulative hazard estimated by a Kaplan-Meier estimator is identical to the number of randomly simulated deaths in a dataset:

```
set.seed(42)
library(survival)

event = rbinom(100, 1, 0.7)
times = runif(100)
H = survfit(Surv(times, event) ~ 1)$cumhaz
c("Deaths" = sum(event), "Sum H" = sum(H))

#> Deaths      Sum H
#>    66          66
```

Houwelingen's  $\alpha$  is then defined by substituting  $H$  for the observed total number of deaths and summing over all predictions:

$$H_\alpha(\boldsymbol{\delta}, \hat{\mathbf{H}}, \mathbf{t}) = \frac{\sum_i \delta_i}{\sum_i \hat{H}_i(t_i)}$$

with standard error  $SE(H_\alpha) = \exp(1/\sqrt{\sum_i \delta_i})$ . A model is well-calibrated with respect to  $H_\alpha$  if  $H_\alpha = 1$ .

The next metrics we look at evaluate models across a spectrum of points to assess calibration over time.

## 8.2 Probabilistic Calibration

Calibration over a range of time points may be assessed quantitatively or qualitatively, with graphical methods often favoured. Graphical methods compare the average predicted distribution to the expected distribution, which can be estimated with the Kaplan-Meier curve, discussed next.

### 8.2.1 Kaplan-Meier Comparison

The simplest graphical comparison compares the average predicted survival curve to the Kaplan-Meier curve estimated on the testing data. Let  $\hat{S}_1, \dots, \hat{S}_m$  be predicted survival functions, then the average predicted survival function is the mixture:  $\bar{\hat{S}} = \frac{1}{m} \sum_{i=1}^m \hat{S}_i(\tau)$ . This estimate can be plotted next to the Kaplan-Meier estimate of the survival distribution in a test dataset (i.e., the true data for model evaluation), allowing for visual comparison of how closely these curves align. An example is given in Figure 8.1, a Cox model (CPH), random survival forest, and relative risk tree, are all compared to the Kaplan-Meier estimator. This figure highlights the advantages and disadvantages of this method. The relative risk tree is clearly poorly calibrated as it increasingly diverges from the Kaplan-Meier. In contrast, the Cox model and random forest cannot be directly compared to one another, as both models frequently overlap with each other and the Kaplan-Meier estimator. Hence it is possible to say that the Cox and forests models are better calibrated than the risk tree, however it is not possible to say which of those two is better calibrated and whether their distance from the Kaplan-Meier is significant or not at a given time (when not clearly overlapping).

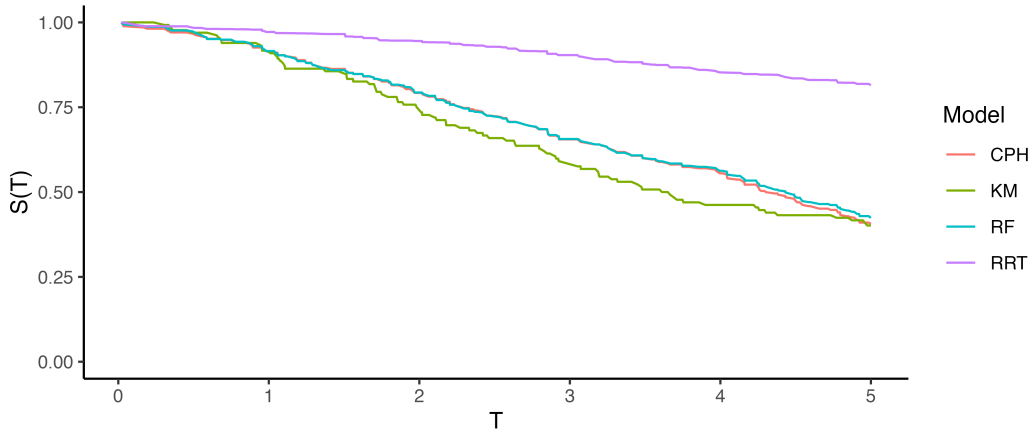


Figure 8.1: Comparing the calibration of a Cox PH (CPH), random forest (RF), and relative risk tree (RRT) to the Kaplan-Meier estimate of the survival function calculated on a test set. The calibration of RRT notably decreases over time whereas RF and CPH are closer to the Kaplan-Meier curve.

This method is useful for making broad statements such as “model X is clearly better calibrated than model Y” or “model X appears to make average predictions close to the Kaplan-Meier estimate”, but that is the limit in terms of useful conclusions. One could refine this method for more fine-grained information by instead using relative risk predictions to create ‘risk groups’ that can be plotted against a stratified Kaplan-Meier, however this

method is harder to interpret and adds even more subjectivity around how many risk groups to create and how to create them (Patrick Royston and Altman 2013). The next measure we consider includes a graphical method as well as a quantitative interpretation.

### 8.2.2 D-Calibration

D-Calibration (Andres et al. 2018; Haider et al. 2020) evaluates a model's calibration by assessing if the predicted survival distributions follow the Uniform distribution as expected, which is motivated by the result that for any random variable  $X$  it follows  $S_X(x) \sim \mathcal{U}(0, 1)$ . This can be tested using a  $\chi^2$  test-statistic:

$$\chi^2 := \sum_{i=1}^n \frac{(O_i - E_i)^2}{E_i}$$

where  $O_1, \dots, O_n$  is the observed number of events in  $n$  groups and  $E_1, \dots, E_n$  is the expected number of events.

To utilise this test, the  $[0, 1]$  codomain of  $S_i$  is cut into  $B$  disjoint contiguous intervals ('bins') over the full range  $[0, 1]$ . Let  $m$  be the total number of observations, then assuming a discrete uniform distribution as the theoretical distribution, the expected number of events in each bin is  $E_i = m/B$  (as the probability of an observation falling into each bin is equal).

The observations in the  $i$ th bin,  $b_i$ , are defined by the set:

$$b_i := \{j = 1, \dots, m : \lceil \hat{S}_i(t_j)B \rceil = i\}$$

where  $j = 1, \dots, m$  are the indices of the observations,  $\hat{S}_i$  are observed (i.e., predicted) survival functions,  $t_i$  are observed (i.e., the ground truth) outcome times, and  $\lceil \cdot \rceil$  is the ceiling function. The observed number of events in  $b_i$  is then the number of observations in that set:  $O_i = |b_i|$ .

The D-Calibration measure, or  $\chi^2$  statistic, is now defined by,

$$D_{\chi^2}(\hat{\mathbf{S}}, \mathbf{t}) := \frac{\sum_{i=1}^B (O_i - \frac{m}{B})^2}{m/B}$$

where  $\hat{\mathbf{S}} = (\hat{S}_1 \ \hat{S}_2 \ \dots \ \hat{S}_m)^\top$  and  $\mathbf{t} = (t_1 \ t_2 \ \dots \ t_m)^\top$ .

This measure has several useful properties. Firstly, one can test the null hypothesis that a model is 'D-calibrated' by deriving a  $p$ -value from comparison to  $\chi^2_{B-1}$ . Secondly,  $D_{\chi^2}$  tends to zero as a model is increasingly well-calibrated, hence the measure can be used for model comparison. Finally, the theory lends itself to an intuitive graphical calibration method as a D-calibrated model implies:

$$p = \frac{\sum_i \mathbb{I}(T_i \leq \hat{F}_i^{-1}(p))}{m}$$

where  $p$  is some value in  $[0, 1]$ ,  $\hat{F}_i^{-1}$  is the  $i$ th predicted inverse cumulative distribution function, and  $m$  is again the number of observations. In words, the number of events occurring at or before each quantile should be equal to the quantile itself, for example 50% of events

should occur before their predicted median survival time. Therefore, one can plot  $p$  on the x-axis and the right hand side of the above equation on the y-axis. A D-calibrated model should result in a straight line on  $x = y$ . This is visualised in Figure 8.2 for the same models as in Figure 8.1. This figure supports the previous findings that the relative risk tree is poorly calibrated in contrast to the Cox model and random forest but again no direct comparison between the latter models is possible.

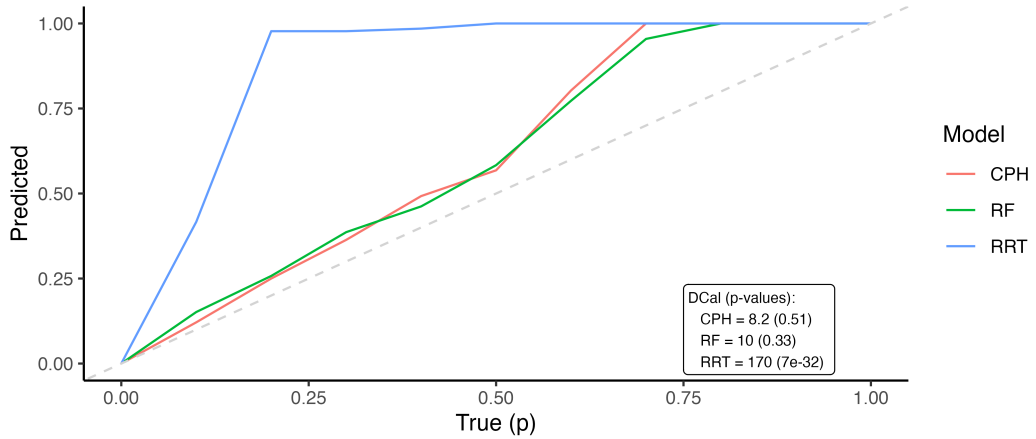


Figure 8.2: Comparing the D-calibration of a Cox PH (CPH), random forest (RF), and relative risk tree (RRT) to the expected distribution on  $y=x$ . As with Figure 8.1, the relative risk tree is clearly not D-calibrated (as supported by the figures in the bottom-right). The CPH and RF are closer to the  $y=x$  however neither follow it perfectly.

Whilst D-calibration has the same problems as the Kaplan-Meier method with respect to visual comparison, at least in this case there are quantities to help draw more concrete solutions. For the models in Figure 8.2, it is clear that the relative risk tree is not D-calibrated with  $p < 0.01$  indicating the null hypothesis of D-calibration, i.e., the predicted quantiles not following a Discrete Uniform distribution, can be comfortably rejected. Whilst the D-calibration for the Cox model is smaller than that of the random forest, the difference is unlikely to be significant, as is seen in the overlapping curves in the figure.

The next chapter will look at scoring rules, which provides a more concrete method to analytically compare the predicted distributions from survival models.



# 9

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## *Evaluating Distributions by Scoring Rules*

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This page is a work in progress and minor changes will be made over time.

Scoring rules evaluate probabilistic predictions and (attempt to) measure the overall predictive ability of a model in terms of both calibration and discrimination (Gneiting and Raftery 2007; Murphy 1973). In contrast to calibration measures, which assess the average performance across all observations on a population level, scoring rules evaluate the sample mean of individual predictions across all observations in a test set. As well as being able to provide information at an individual level, scoring rules are also popular as probabilistic forecasts are widely recognised to be superior to deterministic predictions for capturing uncertainty in predictions (A. P. Dawid 1984; A. Philip Dawid 1986). Formalisation and development of scoring rules has primarily been due to Dawid (A. P. Dawid 1984; A. Philip Dawid 1986; A. Philip Dawid and Musio 2014) and Gneiting and Raftery (Gneiting and Raftery 2007); though the earliest measures promoting “rational” and “honest” decision making date back to the 1950s (Brier 1950; Good 1952). Few scoring rules have been proposed in survival analysis, although the past few years have seen an increase in popularity in these measures. Before delving into these measures, we will first describe scoring rules in the simpler classification setting.

Scoring rules are pointwise losses, which means a loss is calculated for all observations and the sample mean is taken as the final score. To simplify notation, we only discuss scoring rules in the context of a single observation where  $L_i(\hat{S}_i, t_i, \delta_i)$  would be the loss calculated for some observation  $i$  where  $\hat{S}_i$  is the predicted survival function (from which other distribution functions can be derived), and  $(t_i, \delta_i)$  is the observed survival outcome.

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### 9.1 Classification Losses

In the simplest terms, a scoring rule compares two values and assigns them a score (hence ‘scoring rule’), formally we’d write  $L : \mathbb{R} \times \mathbb{R} \mapsto \bar{\mathbb{R}}$ . In machine learning, this usually means comparing a prediction for an observation to the ground truth, so  $L : \mathbb{R} \times \mathcal{P} \mapsto \bar{\mathbb{R}}$  where  $\mathcal{P}$  is a set of distributions. Crucially, scoring rules usually refer to comparisons of true and

predicted *distributions*. As an example, take the Brier score (Brier 1950) defined by:

$$L_{\text{Brier}}(\hat{p}_i, y_i) = (y_i - \hat{p}_i(y_i))^2$$

This scoring rule compares the ground truth to the predicted probability distribution by testing if the difference between the observed event and the truth is minimized. This is intuitive as if the event occurs and  $y_i = 1$ , then  $\hat{p}_i(y_i)$  should be as close to one as possible to minimize the loss. On the other hand, if  $y_i = 0$  then the better prediction would be  $\hat{p}_i(y_i) = 0$ .

This demonstrates an important property of the scoring rule, *properness*. A loss is *proper*, if it is minimized by the correct prediction. In contrast, the loss  $L_{\text{improper}}(\hat{p}_i, y_i) = 1 - L_{\text{Brier}}(\hat{p}_i, y_i)$  is still a scoring rule as it compares the ground truth to the prediction probability distribution, but it is clearly improper as the perfect prediction ( $\hat{p}_i(y_i) = y_i$ ) would result in a score of 1 whereas the worst prediction would result in a score of 0. Proper losses provide a method of model comparison as, by definition, predictions closest to the true distribution will result in lower expected losses.

Another important property is *strict properness*. A loss is *strictly proper* if the loss is uniquely minimized by the ‘correct’ prediction. Consider now the loss  $L_0(\hat{p}_i, y_i) = 0$ . Not only is this a strictly proper scoring rule but it is also proper. The loss can only take the value 0 and is therefore guaranteed to be minimized by the correct prediction. It is clear however that this loss is useless. In contrast, the Brier score is minimized by only one value, which is the optimal prediction (Figure 9.1). Strictly proper losses are particular important for automated model optimisation, as minimization of the loss will result in the ‘optimum score estimator based on the scoring rule’ (Gneiting and Raftery 2007).

Mathematically, a classification loss  $L : \mathcal{P} \times \mathcal{Y} \rightarrow \bar{\mathbb{R}}$  is *proper* if for any distributions  $p_Y, p$  in  $\mathcal{P}$  and for any random variables  $Y \sim p_Y$ , it holds that  $\mathbb{E}[L(p_Y, Y)] \leq \mathbb{E}[L(p, Y)]$ . The loss is *strictly proper* if, in addition,  $p = p_Y$  uniquely minimizes the loss.

As well as the Brier score, which was defined above, another widely used loss is the log loss (Good 1952), defined by

$$L_{\text{logloss}}(\hat{p}_i, y_i) = -\log \hat{p}_i(y_i)$$

These losses are visualised in Figure 9.1, which highlights that both losses are strictly proper (A. Philip Dawid and Musio 2014) as they are minimized when the true prediction is made, and converge to the minimum as predictions are increasingly improved.

## 9.2 Survival Losses

Analogously to classification losses, a survival loss  $L : \mathcal{P} \times \mathbb{R}_{>0} \times \{0, 1\} \rightarrow \bar{\mathbb{R}}$  is *proper* if for any distributions  $p_Y, p$  in  $\mathcal{P}$ , and for any random variables  $Y \sim p_Y$ , and  $C$  t.v.i.  $\mathbb{R}_{>0}$ ; with  $T := \min(Y, C)$  and  $\Delta := \mathbb{I}(T = Y)$ ; it holds that,  $\mathbb{E}[L(p_Y, T, \Delta)] \leq \mathbb{E}[L(p, T, \Delta)]$ . The loss is *strictly proper* if, in addition,  $p = p_Y$  uniquely minimizes the loss. A survival loss is referred to as outcome-independent (strictly) proper if it is only (strictly) proper when  $C$  and  $Y$  are independent.

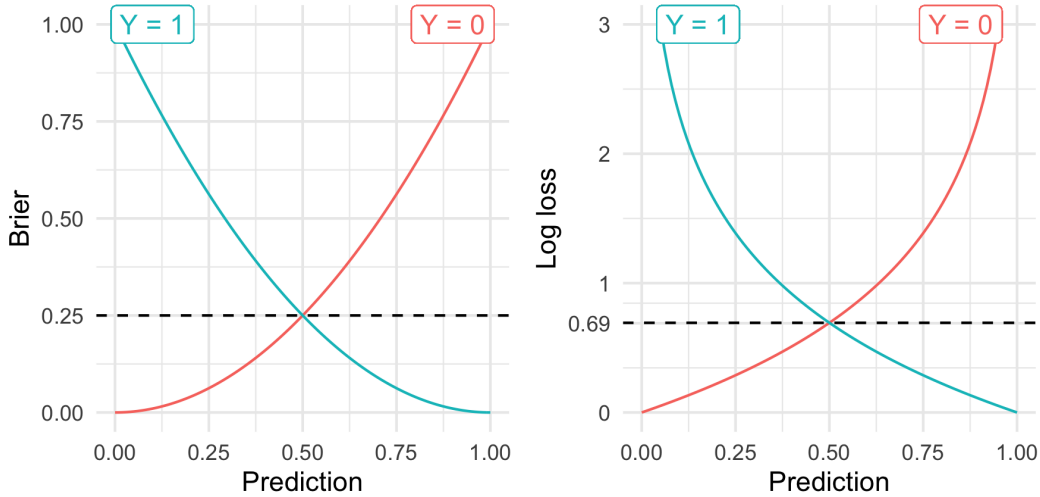


Figure 9.1: Brier and log loss scoring rules for a binary outcome and varying probabilistic predictions. x-axis is a probabilistic prediction in  $[0, 1]$ , y-axis is Brier score (left) and log loss (right). Blue lines are varying Brier score/log loss over different predicted probabilities when the true outcome is 1. Red lines are varying Brier score/log loss over different predicted probabilities when the true outcome is 0. Both losses are minimized when  $\hat{p}_i(y_i) = y_i$ .

With these definitions, the rest of this chapter lists common scoring rules in survival analysis and discusses some of their properties. As with other chapters, this list is likely not exhaustive but will cover commonly used losses.

### 9.2.1 Integrated Graf Score

The Integrated Graf Score (IGS) was introduced by Graf (Graf and Schumacher 1995; Graf et al. 1999) as an analogue to the integrated brier score (IBS) in regression. It is likely the commonly used scoring rule in survival analysis, possibly due to its intuitive interpretation.

The loss is defined by

$$L_{IGS}(\hat{S}_i, t_i, \delta_i | \hat{G}_{KM}) = \int_0^{\tau^*} \frac{\hat{S}_i^2(\tau) \mathbb{I}(t_i \leq \tau, \delta_i = 1)}{\hat{G}_{KM}(t_i)} + \frac{\hat{F}_i^2(\tau) \mathbb{I}(t_i > \tau)}{\hat{G}_{KM}(\tau)} d\tau \quad (9.1)$$

where  $\hat{S}_i^2(\tau) = (\hat{S}_i(\tau))^2$  and  $\hat{F}_i^2(\tau) = (1 - \hat{S}_i(\tau))^2$ , and  $\tau^* \in \mathbb{R}_{\geq 0}$  is an upper threshold to compute the loss up to, and  $\hat{G}_{KM}$  is the Kaplan-Meier trained on the censoring distribution for IPCW (Section 7.1).

At first glance this might seem intimidating but it is worth taking the time to understand the intuition behind the loss. Recall the classification Brier score,  $L(\hat{p}_i, y_i) = (y_i - \hat{p}_i)^2$ , this provides a method to compare and evaluate a probability mass function at one time-point. The *integrated Brier score* (IBS), also known as the *CRPS*, is the integral of the Brier score for all real-valued thresholds (Gneiting and Raftery 2007) and hence allows predictions to be evaluated over multiple points as  $L(\hat{F}_i, y_i) = \int (\hat{F}_i(y_i) - \mathbb{I}(y_i \geq x))^2 dy$  where  $\hat{F}_i$  is the predicted cumulative distribution function and  $x$  is some meaningful threshold. In survival

analysis,  $\hat{F}_i(\tau)$  represents the probability of an observation having experienced the event at or before  $\tau$ , and the ground truth to compare to is therefore whether the observation has actually experienced the event at  $\tau$ , which is the case when  $t_i \leq \tau$ . Hence the IBS becomes  $L(\hat{F}_i, t_i) = \int (\hat{F}_i(\tau) - \mathbb{I}(t_i \leq \tau))^2 d\tau$ . Now for a given time  $\tau$ :

$$L(\hat{F}_i, t) = \begin{cases} (\hat{F}_i(\tau) - 1)^2 = (1 - \hat{F}_i(\tau))^2 = \hat{S}_i^2(\tau), & \text{if } t_i \leq \tau \\ (\hat{F}_i(\tau) - 0)^2 = \hat{F}_i^2(\tau), & \text{if } t_i > \tau \end{cases} \quad (9.2)$$

In words, if an observation has not yet experienced an outcome ( $t_i > \tau$ ) then the loss is minimized when the cumulative distribution function (the probability of having already died) is 0, which is intuitive as the optimal prediction is correctly identifying the observation has yet to experience the event. In contrast, if the observation has experienced the outcome ( $t_i \leq \tau$ ) then the loss is minimized when the survival function (the probability of surviving until  $\tau$ ) is 0, which follows from similar logic.

The final component of the Graf score is accommodating for censoring. At  $\tau$  an observation will either have

1. Not experienced the event:  $I(t_i > \tau)$ ;
2. Experienced the event:  $I(t_i \leq \tau, \delta_i = 1)$ ; or
3. Been censored:  $I(t_i \leq \tau, \delta_i = 0)$

In the Graf score, censored observations are discarded. If they were not then Equation 9.2 would imply their contribution would be treated the same as those who had experienced the event. However this assumption would be entirely wrong as a censored observation is guaranteed not to have experienced the event, hence an ideal prediction for a censored observation is a high survival probability up until the point of censoring, at which time comparison to ground truth is unknown as this is no longer observed.

The act of discarding censored observations means that the sample size decreases over time. To compensate for this, IPCW is used to increasingly weight predictions as  $\tau$  increases. Hence, IPCW weights,  $W_i$  are defined such that

$$W_i = \begin{cases} \hat{G}_{KM}^{-1}(t_i), & \text{if } \mathbb{I}(t_i \leq \tau, \delta_i = 1) \\ \hat{G}_{KM}^{-1}(\tau), & \text{if } \mathbb{I}(t_i > \tau) \end{cases}$$

The weights total 1 when divided over all samples and summed (Graf et al. 1999). They are also intuitive as observations are either weighted by  $G(\tau)$  when they are still alive and therefore still part of the sample, or by  $G(t_i)$  otherwise.

As well as being intuitive, when censoring is uninformative, the Graf score consistently estimates the mean square error  $L(t, S|\tau^*) = \int_0^{\tau^*} [\mathbb{I}(t > \tau) - S(\tau)]^2 d\tau$ , where  $S$  is the correctly specified survival function (Gerds and Schumacher 2006). However, despite these promising properties, the IGS is improper and must therefore be used with care (Rindt et al. 2022; R. Sonabend et al. 2024). In practice, experiments have shown that the effect of impropriety is minimal and therefore this loss should be avoided for automated routines such as model tuning, however it can still be used for model evaluation. In addition, a small adaptation of the loss results in a strictly proper scoring rule simply by altering the weights such that  $W_i = \hat{G}_{KM}^{-1}(t_i)$  for all uncensored observations and 0 otherwise (R. Sonabend et al. 2024), resulting in the reweighted Graf score:

$$L_{RGS}(\hat{S}_i, t_i, \delta_i | \hat{G}_{KM}) = \delta_i \mathbb{I}(t_i \leq \tau^*) \int_0^{\tau^*} \frac{(\mathbb{I}(t_i \leq \tau) - \hat{F}_i(\tau))^2}{\hat{G}_{KM}(t_i)} d\tau$$

The addition of  $\mathbb{I}(t_i \leq \tau^*)$  completely removes observations that experience the event after the cutoff time,  $\tau^*$ , this ensures there are no cases where the  $G(t_i)$  weighting is calculated on time after the cutoff. Including an upper threshold (i.e,  $\tau^* < \infty$ ) effects properness and generalization statements. For example, by evaluating a model using the RGS with a  $\tau^*$  threshold, then the model may be said to only perform well up until  $\tau^*$  with its performance unknown after this time.

The change of weighting slightly alters the interpretation of the contributions at different time-points. By example, let  $(t_i = 4, t_j = 5)$  be two observed survival times, then at  $\tau = 3$ , the Graf score weighting would be  $\hat{G}_{KM}^{-1}(4)$  for both observations, whereas the RGS weights would be  $(KMG^{-1}(4), KMG^{-1}(5))$  respectively, hence there is always more ‘importance’ placed on observations that take longer to experience the event. In practice, the difference between these weights appears to be minimal (R. Sonabend et al. 2024) but as RGS is strictly proper, it is more suitable for automated experiments.

### 9.2.2 Integrated Survival Log Loss

The integrated survival log loss (ISLL) was also proposed by Graf et al. (1999).

$$L_{ISLL}(\hat{S}_i, t_i, \delta_i | \hat{G}_{KM}) = - \int_0^{\tau^*} \frac{\log[\hat{F}_i(\tau)] \mathbb{I}(t_i \leq \tau, \delta_i = 1)}{\hat{G}_{KM}(t_i)} + \frac{\log[\hat{S}_i(\tau)] \mathbb{I}(t_i > \tau)}{\hat{G}_{KM}(\tau)} d\tau$$

where  $\tau^* \in \mathbb{R}_{>0}$  is an upper threshold to compute the loss up to.

Similarly to the IGS, there are three ways to contribute to the loss depending on whether an observation is censored, experienced the event, or alive, at  $\tau$ . Whilst the IGS is routinely used in practice, there is no evidence that ISLL is used, and moreover there are no proofs (or claims) that it is proper.

The reweighted ISLL (RISLL) follows similarly to the RIGS and is also outcome-independent strictly proper (R. Sonabend et al. 2024).

$$L_{RISLL}(\hat{S}_i, t_i, \delta_i | \hat{G}_{KM}) = -\delta_i \mathbb{I}(t_i \leq \tau^*) \int_0^{\tau^*} \frac{\mathbb{I}(t_i \leq \tau) \log[\hat{F}_i(\tau)] + \mathbb{I}(t_i > \tau) \log[\hat{S}_i(\tau)]}{\hat{G}_{KM}(t_i)} d\tau$$

### 9.2.3 Survival density log loss

Another outcome-independent strictly proper scoring rule is the survival density log loss (SDLL) (R. Sonabend et al. 2024), which is given by

$$L_{SDLL}(\hat{f}_i, t_i, \delta_i | \hat{G}_{KM}) = -\frac{\delta_i \log[\hat{f}_i(t_i)]}{\hat{G}_{KM}(t_i)}$$

where  $\hat{f}_i$  is the predicted probability density function. This loss is essentially the classification log loss ( $-\log(\hat{p}_i(t_i))$ ) with added IPCW. Whilst the classification log loss has beneficial

properties such as being differentiable, this is more complex for the SDLL and it is not widely used in practice. A useful alternative to the SDLL which can be readily used in automated procedures is the right-censored log loss.

#### 9.2.4 Right-censored log loss

The right-censored log loss (RCLL) is an outcome-independent strictly proper scoring rule (Avati et al. 2020) that benefits from not depending on IPCW in its construction. The RCLL is defined by

$$L_{RCLL}(\hat{S}_i, t_i, \delta_i) = -\log[\delta_i \hat{f}_i(t_i) + (1 - \delta_i) \hat{S}_i(t_i)]$$

This loss is interpretable when we break it down into its two halves:

1. If an observation is censored at  $t_i$  then all the information we have is that they did not experience the event at the time, so they must be ‘alive’, hence the optimal value is  $\hat{S}_i(t_i) = 1$  (which becomes  $-\log(1) = 0$ ).
2. If an observation experiences the event then the ‘best’ prediction is for the probability of the event at that time to be maximised, as pdfs are not upper-bounded this means  $\hat{f}_i(t_i) = \infty$  (and  $-\log(t_i) \rightarrow \infty$  as  $t_i \rightarrow \infty$ ).

#### 9.2.5 Absolute Survival Loss

The absolute survival loss, developed over time by Schemper and Henderson (2000) and Schmid et al. (2011), is based on the mean absolute error is very similar to the IGS but removes the squared term:

$$L_{ASL}(\hat{S}_i, t_i, \delta_i | \hat{G}_{KM}) = \int_0^{\tau^*} \frac{\hat{S}_i(\tau) \mathbb{I}(t_i \leq \tau, \delta_i = 1)}{\hat{G}_{KM}(t_i)} + \frac{\hat{F}_i(\tau) \mathbb{I}(t_i > \tau)}{\hat{G}_{KM}(\tau)} d\tau$$

where  $\hat{G}_{KM}$  and  $\tau^*$  are as defined above. Analogously to the IGS, the ASL score consistently estimates the mean absolute error when censoring is uninformative (Schmid et al. 2011) but there are also no proofs or claims of properness. The ASL and IGS tend to yield similar results (Schmid et al. 2011) but in practice there is no evidence of the ASL being widely used.

### 9.3 Prediction Error Curves

As well as evaluating probabilistic outcomes with integrated scoring rules, non-integrated scoring rules can be utilised for evaluating distributions at a single point. For example, instead of evaluating a probabilistic prediction with the IGS over  $\mathbb{R}_{\geq 0}$ , instead one could compute the IGS at a single time-point,  $\tau \in \mathbb{R}_{\geq 0}$ , only. Plotting these for varying values of  $\tau$  results in ‘prediction error curves’ (PECs), which provide a simple visualisation for how predictions vary over the outcome. PECs are especially useful for survival predictions as they can visualise the prediction ‘over time’. PECs are mostly used as a graphical guide when comparing few models, rather than as a formal tool for model comparison. An example for

PECs is provided in Figure 9.2 for the IGS where the Cox PH consistently outperforms the SVM.

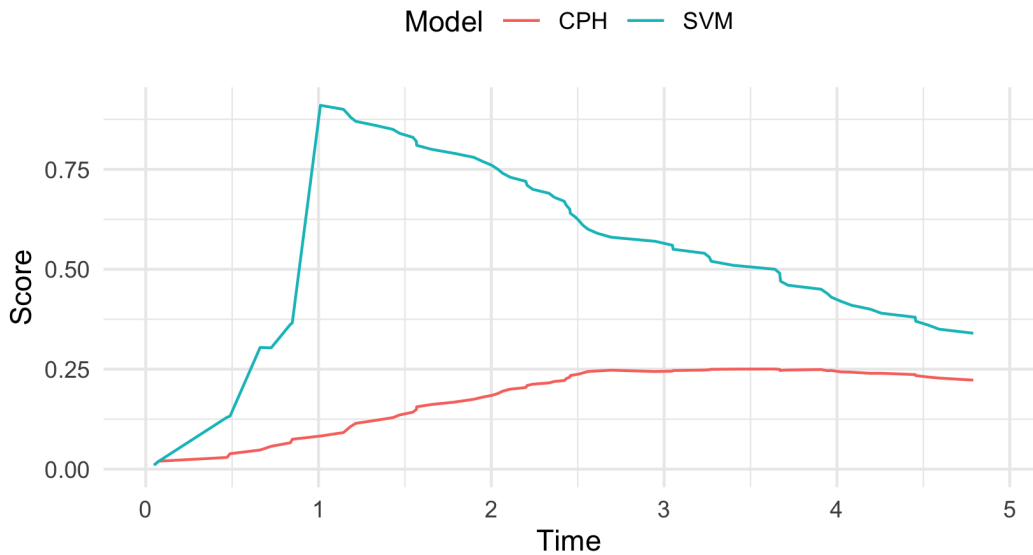


Figure 9.2: Prediction error curves for the CPH and SVM models from Chapter 8. x-axis is time and y-axis is the IGS computed at different time-points. The CPH (red) performs better than the SVM (blue) as it scores consistently lower. Trained and tested on randomly simulated data from `$mlr3proba`.

## 9.4 Baselines and ERV

A common criticism of scoring rules is a lack of interpretability, for example, an IGS of 0.5 or 0.0005 has no meaning by itself, so below we present two methods to help overcome this problem.

The first method, is to make use of baselines for model comparison, which are models or values that can be utilised to provide a reference for a loss, they provide a universal method to judge all models of the same class by (Gressmann et al. 2018). In classification, it is possible to derive analytical baseline values, for example a Brier score is considered ‘good’ if it is below 0.25 or a log loss if it is below 0.693 (Figure 9.1), this is because these are the values obtained if you always predicted probabilities as 0.5, which is a reasonable baseline guess in a binary classification problem. In survival analysis, simple analytical expressions are not possible as losses are dependent on the unknown distributions of both the survival and censoring time. Therefore all experiments in survival analysis must include a baseline model that can produce a reference value in order to derive meaningful results. A suitable baseline model is the Kaplan-Meier estimator (Graf and Schumacher 1995; Lawless and Yuan 2010; Patrick Royston and Altman 2013), which is the simplest model that can consistently estimate the true survival function.

As well as directly comparing losses from a ‘sophisticated’ model to a baseline, one can

also compute the percentage increase in performance between the sophisticated and baseline models, which produces a measure of explained residual variation (ERV) (Edward L. Korn and Simon 1990; Edward L. Korn and Simon 1991). For any survival loss  $L$ , the ERV is,

$$R_L(S, B) = 1 - \frac{L|S}{L|B}$$

where  $L|S$  and  $L|B$  is the loss computed with respect to predictions from the sophisticated and baseline models respectively.

The ERV interpretation makes reporting of scoring rules easier within and between experiments. For example, say in experiment A we have  $L|S = 0.004$  and  $L|B = 0.006$ , and in experiment B we have  $L|S = 4$  and  $L|B = 6$ . The sophisticated model may appear worse at first glance in experiment A (as the losses are very close) but when considering the ERV we see that the performance increase is identical (both  $R_L = 33\%$ ), thus providing a clearer way to compare models.

## 9.5 Conclusion

### Key takeaways

- Scoring rules are a useful tool for measuring a model's overall predictive ability, taking into account calibration and discrimination.
- Strictly proper scoring rules allow models to be compared to one another, which is important when choosing models in a benchmark experiment.
- Many scoring rules for censored data are *not* strictly proper, however experiments suggest that improper rules still provide useful and trustworthy results (R. Sonabend et al. 2024)

### Limitations

- Scoring rules can be difficult to interpret but ERV representations can be a helpful way to overcome this.
- There is no consensus about which scoring rule to use and when so in practice multiple scoring rules may have to be reported in experiments to ensure transparency and fairness of results.
- For non- and semi-parametric survival models that return distribution predictions, estimates of  $f(t)$  are not readily available and require approximations (Rindt et al. 2022), hence measures such as RCLL and SDLL can often not be directly used in practice.

### Further reading

- A. Philip Dawid and Musio (2014) and Gneiting and Raftery (2007) provide a comprehensive summary of scoring rules in regression and classification settings.
- Rindt et al. (2022) and R. Sonabend et al. (2024) review survival scoring rules.

- Rahman et al. (2017) compare measures for external validation including some scoring rules.



# 10

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## Evaluating Survival Time

TODO (150-200 WORDS)

 Minor changes expected!

This page is a work in progress and minor changes will be made over time.

When it comes to evaluating survival time predictions, there are few measures available at our disposal. As a result of survival time predictions being uncommon compared to other prediction types (`?@sec-surv-set-types`), there are limited survival time evaluation measures in the literature. To our knowledge, there are no specialised ‘survival time measures’, instead regression measures are used by ignoring censored observations.

Before presenting these measures, consider what happens when censored observations are discarded. If censoring is truly independent, occurs randomly, and is *very* limited in the data, then there is little harm in discarding observations and treating this as a regression problem. However, if censoring is not independent, then discarding censored observations will lead to missing valuable insights about the model. For example, say the task of interest is to predict the probability of death due to kidney failure and patients are censored if they receive a transplant - this is clearly a competing risk as receiving a transplant greatly reduces the probability of death. If one were to predict the time to death for all patients and to not evaluate the quality of prediction for censored patients, then it would only be possible to conclude about the model’s performance for those who do not receive a transplant. On the surface this may appear to be of value, however, if at the time of prediction it is impossible to know who will receive a transplant (perhaps because the dataset omits relevant information such as time of hospital admission, wait on register, etc.), then for a given prediction for an observation, it would be impossible to know if the prediction is trustworthy - it would be if that patient does not receive a transplant, but would not be if they do not. In short, it is essential that predictions for individuals who end up being censored, are as good as those who are not, simply because there is no method to know which group observations will eventually fall into.

It is interesting to consider if IPCW strategies would compensate for this deficiency, however as we were unable to find research into this method, we have only included measures that we term ‘censoring-ignored regression measures’, which are presented in (P. Wang, Li, and Reddy 2019).

## 10.1 Distance measures

Survival time measures are often referred to as ‘distance’ measures as they measure the distance between the true,  $(t, \delta = 1)$ , and predicted,  $\hat{t}$ , values. These are presented in turn with brief descriptions of their interpretation.

### Censoring-ignored mean absolute error, $MAE_C$

In regression, the mean absolute error (MAE) is a popular measure because it is intuitive to understand as it measures the absolute difference between true and predicted outcomes; hence intuitively one can understand that a model predicting a height of 175cm is clearly better than one predicting a height of 180cm, for a person with true height of 174cm.

$$MAE_C(\hat{\mathbf{t}}, \mathbf{t}, \boldsymbol{\delta}) = \frac{1}{d} \sum_{i=1}^m \delta_i |t_i - \hat{t}_i|$$

Where  $d$  is the number of uncensored observations in the dataset,  $d = \sum_i \delta_i$ .

### Censoring-ignored mean squared error

In comparison to MAE, the mean squared error (MSE), computes the squared differences between true and predicted values. While the MAE provides a smooth, linear, ‘penalty’ for increasingly poor predictions (i.e., the difference between an error of predicting 2 vs. 5 is still 3), but the square in the MSE means that larger errors are quickly magnified (so the difference in the above example is 9). By taking the mean over all predictions, the effect of this inflation is to increase the MSE value as larger mistakes are made.

$$MSE_C(\hat{\mathbf{t}}, \mathbf{t}, \boldsymbol{\delta}) = \frac{1}{d} \sum_{i=1}^m \delta_i (t_i - \hat{t}_i)^2$$

Where  $d$  is again the number of uncensored observations in the dataset,  $d = \sum_i \delta_i$ .

### Censoring-adjusted root mean squared error

Finally, the root mean squared error (RMSE), is simply the square root of the MSE. This allows interpretation on the original scale (as opposed to the squared scale produced by the MSE). Given the inflation effect for the MSE, the RMSE will be larger than the MAE as increasingly poor predictions are made; it is common practice for the MAE and RMSE to be reported together.

$$RMSE_C(\hat{\mathbf{t}}, \mathbf{t}, \boldsymbol{\delta}) = \sqrt{MSE_C(\hat{\mathbf{t}}, \mathbf{t}, \boldsymbol{\delta})}$$

## 10.2 Over- and under-predictions

All of these distance measures assume that the error for an over-prediction ( $\hat{t} > t$ ) should be equal to an under-prediction ( $\hat{t} < t$ ), i.e., that it is ‘as bad’ if a model predicts an outcome time being 10 years longer than the truth compared to being 10 years shorter. In

the survival setting, this assumption is often invalid as it is generally preferred for models to be overly cautious, hence to predict negative events to happen sooner (e.g., predict a life-support machine fails after three years not five if the truth is actually four) and to predict positive events to happen later (e.g., predict a patient recovers after four years not two if the truth is actually three). A simple method to incorporate this imbalance between over- and under-predictions is to add a weighting factor to any of the above measures, for example the  $MAE_C$  might become

$$MAE_C(\hat{\mathbf{t}}, \mathbf{t}, \boldsymbol{\delta}, \lambda, \mu, \phi) = \frac{1}{d} \sum_{i=1}^m \delta_i |(t_i - \hat{t}_i)[\lambda \mathbb{I}(t_i > \hat{t}_i) + \mu \mathbb{I}(t_i < \hat{t}_i) + \phi \mathbb{I}(t_i = \hat{t}_i)]|$$

where  $\lambda, \mu, \phi$  are any Real number to be used to weight over-, under-, and exact-predictions, and  $d$  is as above. The choice of these are highly context dependent and could even be tuned.



# 11

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## *Choosing Measures*

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TODO (150-200 WORDS)

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After reading this part of the book, evaluating survival analysis models may appear more daunting than regression and classification settings, which, in contrast, have fewer (common) measures to choose from. In regression problems, the RMSE and MAE are common choices for evaluating how far predictions are from the truth. In classification, the Brier score or logloss may be used to evaluate probabilistic predictions and the accuracy score or TPR/TNR/FPR/FNR are common for deterministic predictions. In contrast, there are many more measures in survival analysis which are necessarily more complex, due to the need to handle censoring with many possible methods for doing so. Therefore, this final chapter aims to provide some simple to follow guidelines for selecting measures for different types of experiments.

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### 11.1 Defining the experiment

Experiments may be performed to make predictions for new data, compare the performance of multiple models ('benchmark experiments'), investigate patterns in observed data, or some combination of these. Each experiment requires different choices of measures, with different levels of strictness applied to measure assumptions.

#### 11.1.1 Predictive experiments

In the real world, predictive experiments are most common. These are now daily occurrences as machine learning models are routinely deployed on servers to make ongoing predictions. In these cases, the exact task must be precisely stated before any model is deployed and evaluated. Common survival problems to solve include:

1. Identifying low and high risk groups in new data (for resource allocation);
2. Predicting the survival distribution for an individual over time; and
3. Predicting the survival probability for an individual at a specific time.

The first of these is a discrimination problem and it is therefore most important that the model optimises corresponding measures and that measure assumptions are justified. However, even this task may be more complex than it initially seems. For example, while some papers have shown flaws in Harrell's C (Gönen and Heller 2005; Rahman et al. 2017; Schmid and Potapov 2012; Uno et al. 2007), others have demonstrated that common alternatives yield very similar results (Rahman et al. 2017; Therneau and Atkinson 2020) and moreover some prominent alternatives may be harder to interpret due to high variance (Rahman et al. 2017; Schmid and Potapov 2012). In predictive experiment that may require more level of automation, it is important to be careful of C-hacking (Section 7.1.2) and to avoid overoptimistic results. Hence one should not compute a range of concordance indices and report the maximum but instead calculate a single discrimination measure and then establish a pre-defined threshold to determine if the deployed model is optimal, a natural threshold would be 0.5 as anything above this is better than a baseline model. Given Harrell's C to be increasingly over-optimistic with additional censoring (Rahman et al. 2017), it is advisable to use Uno's C instead.

If the task of interest is to predict survival distributions *over time*, then the choice of measure is more limited and only the RCLL and the proper Graf score are recommended. Both these measures can only be interpreted with respect to a baseline so use of the ERV representation is strongly recommended. As with the previous task, establishing a threshold for performance is essential prior to deployment and for ongoing evaluation. It is less clear in these cases what this threshold might be, but the simplest starting point would be to ensure that the model continues to outperform the baseline or a simpler gold-standard model (e.g., the Cox PH).

The final task of interest differs from the previous by only making predictions at a specific time. In this case, prediction error curves, and single-time point calibration measures can be used, as well as scoring rules with shorter cut-offs (i.e., the upper limit of the integral). It is imperative that model performance is never extrapolated outside of the pre-specified time.

### 11.1.2 Benchmark experiments

When conducting benchmark experiments, it is advisable to use a spread of measures so that results can be compared across various properties. In this case, models should be tested against discrimination, calibration, and overall predictive ability (i.e., with scoring rules). As models make different types of predictions, results from these experiments should be limited to metrics that are directly comparable, in other words, two models should only be compared based on the *same* metric. In benchmark experiments, models are compared across the same data and same resampling strategy, hence measure assumptions become less important as they are equally valid or flawed for all models. For example, if one dataset has particularly high amounts of censoring leading to an artificially higher concordance index, then this bias would affect all models equally and the overall experiment would not be affected. Hence, in these experiments it suffices to pick one or two measures for concordance, discrimination, and predictive ability, without having to be overly concerned with the individual metric.

This book recommends using Harrell's C and Uno's C for concordance as these are simplest to compute and including both enables more confidence in model comparison, i.e., if a model outperforms another with respect to both these measures then there can be higher confidence in drawing statements about the model's discriminatory power. For calibration, D-calibration is recommended as it can be meaningfully compared between models, and the RCLL is recommended for a scoring rule (which is proper for outcome-independent censoring). No distance measure is recommended as these do not apply to the vast majority of models. All

these measures can be used for automated tuning, in the case of discrimination tuning to Harrell's C alone should suffice (without also tuning to Uno's C).

### 11.1.3 Investigation

Investigating patterns in observed data is increasingly common as model interpretability methods have become more accessible (Molnar 2019). Before data can be investigated, any model that is trained on the data must first be demonstrated to be a good fit to the data. A model's fit to data can also be evaluated by resampling the data (Chapter 2) and evaluating the predictions. In this case, it is important to choose measures that are interpretable and have justified assumptions. Calibration measures are particularly useful for evaluating if a model is well fit to data, and any of the methods described in Chapter 8 are recommended for this purpose. Discrimination measures *may* be useful, however, given how susceptible they are to censoring, they can be difficult to interpret on their own, and the same is true for scoring rules. One method to resolve ambiguity is to perform a benchmark experiment of multiple models on the same data (ideally with some automated tuning) and then select the best model from this experiment and refit it on the full data (Becker, Schneider, and Fischer 2024) – this is a robust, empirical method that demonstrates a clear trail to selecting a model that outperforms other potential candidates. When investigating a dataset, one may also consider using different measures to assess algorithmic fairness (R. Sonabend et al. 2022), any measure that can be optimised (i.e., where the lowest or highest value is the best) may be used in this case. Finally, there are survival adaptations to the well-known AIC (Liang and Zou 2008) and BIC (Volinsky and Raftery 2000) however as these are generally only applicable to 'classical' models (Chapter 12), they are out of scope for this book and hence have not been discussed.

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## 11.2 Conclusions

This part of the book focused on survival measures. Measures may be used to evaluate model predictions, to tune a model, or to train a model (e.g., in boosting or neural networks). Unlike other settings, there are many different choices of survival measures and it can be hard to determine which to use and when. In practice, like many areas of Statistics, the most important factor is to clearly define any experiment upfront and to be clear about which measures will be used and why. As a rule of thumb, good choices for measures are Harrell's C for evaluating discrimination, with Uno's C supporting findings, D-calibration for calibration, and the RCLL for evaluating overall predictive ability from distribution predictions. Finally, if you are restricted to a single measure choice (e.g., for automated tuning or continuous evaluation of deployed models), then we recommend selecting a scoring rule such as RCLL which captures information about calibration and discrimination simultaneously.



## **Part III**

# **Models**



# 12

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## *Classical Models*

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### 12.1 A Review of Classical Survival Models

This chapter provides a brief review of classical survival models before later chapters move on to machine learning models. ‘Classical’ models are defined with a very narrow scope in this book: low-complexity models that are either non-parametric or have parameters that can be fit with maximum likelihood estimation (or an equivalent method). In contrast, ‘machine learning’ (ML) models require more intensive model fitting procedures such as recursion or iteration. The classical models in this paper are fast to fit and highly interpretable, though can be inflexible and may make unreasonable assumptions. Whereas the ML models are more flexible with hyper-parameters however are computationally more intensive (both in terms of speed and storage), require tuning to produce ‘good’ results, and are often a ‘black-box’ with difficult interpretation.

As classical survival models have been studied extensively for decades, these are only discussed briefly here, primarily these are of interest as many of these models will be seen to influence machine learning extensions. The scope of the models discussed in this chapter is limited to the general book scope (`?@sec-surv-scope`), i.e. single event with right-censoring and no competing-risks, though in some cases these are discussed.

There are several possible taxonomies for categorising statistical models, these include:

- Parametrisation Type: One of non-, semi-, or fully-parametric. \ Non-parametric models assume that the data distribution cannot be specified with a finite set of parameters. In

contrast, fully-parametric models assume the distribution can be specified with a finite set of parameters. Semi-parametric models are a hybrid of the two and are formed of a finite set of parameters *and* an infinite-dimensional ‘nuisance’ parameter.

- Conditionality Type: One of unconditional or conditional. A conditional prediction is one that makes use of covariates in order to condition the prediction on each observation. Unconditional predictors, which are referred to below as ‘estimators’, ignore covariate data and make the same prediction for all individuals.
- Prediction Type: One of ranking, survival time, or distribution (?@sec-surv-set-types).

Table 12.1 summarises the models discussed below into the taxonomies above for reference. Note that the Cox model is listed as predicting a continuous ranking, and not a survival distribution, which may appear inconsistent with other definitions. The reason for this is elaborated upon in Chapter 18. Though the predict-type taxonomy is favoured throughout this book, it is clearer to review classical models in increasing complexity, beginning with unconditional estimators before moving onto semi-parametric continuous ranking predictions, and finally conditional distribution predictors. The review is brief with mathematics limited to the model fundamentals but not including methods for parameter estimation. Also the review is limited to the ‘basic’ model specification and common extensions such as regularization are not discussed though they do exist for many of these models.

All classical models are highly transparent and accessible, with decades of research and many off-shelf implementations. Predictive performance of each model is briefly discussed as part of the review and then again in (R. E. B. Sonabend 2021).

Table 12.1: Table of models discussed in this literature review, classified by parametrisation, prediction type, and conditionality.

Model <sup>1</sup>	Parametrisation <sup>2</sup>	Prediction <sup>3</sup>	Conditionality
Kaplan-Meier	Non	Distr.	Unconditional
Nelson-Aalen	Non	Distr.	Unconditional
Akritis	Non	Distr.	Conditional
Cox PH	Semi	Rank	Conditional
Parametric PH	Fully	Distr.	Conditional
Accelerated Failure Time	Fully	Distr.	Conditional
Proportional Odds	Fully	Distr.	Conditional
Flexible Spline	Fully	Distr.	Conditional

\* 1. All models are implemented in the R package **survival** (Therneau 2015) with the exception of flexible splines, implemented in **flexsurv** (C. Jackson 2016), and the Akritis estimator in **survivalmodels** (R. Sonabend 2020). \* 2. Non = non-parametric, Semi = semi-parametric, Fully = fully-parametric. \* 3. Distr. = distribution, Rank = ranking.

### 12.1.1 Non-Parametric Distribution Estimators

#### Unconditional Estimators

Unconditional non-parametric survival models assume no distribution for survival times and estimate the survival function using simple algorithms based on observed outcomes and no covariate data. The two most common methods are the Kaplan-Meier estimator (**KaplanMeier1958?**), which estimates the average survival function of a training dataset,

and the Nelson-Aalen estimator (O. Aalen 1978; Nelson 1972), which estimates the average cumulative hazard function of a training dataset.

The Kaplan-Meier estimator of the survival function is given by

$$\hat{S}_{KM}(\tau | \mathcal{D}_{train}) = \prod_{t \in \mathcal{U}_O, t \leq \tau} \left(1 - \frac{d_t}{n_t}\right) \quad (12.1)$$

As this estimate is so important in survival models, this book will always use the symbol  $\hat{S}_{KM}$  to refer to the Kaplan-Meier estimate of the average survival function fit on training data  $(T_i, \Delta_i)$ . Another valuable function is the Kaplan-Meier estimate of the average survival function of the *censoring* distribution, which is the same as above but estimated on  $(T_i, 1 - \Delta_i)$ , this will be denoted by  $\hat{G}_{KM}$ .

The Nelson-Aalen estimator for the cumulative hazard function is given by

$$\hat{H}(\tau | \mathcal{D}_{train}) = \sum_{t \in \mathcal{U}_O, t \leq \tau} \frac{d_t}{n_t} \quad (12.2)$$

The primary advantage of these models is that they rely on heuristics from empirical outcomes only and don't require any assumptions about the form of the data. To train the models they only require  $(T_i, \Delta_i)$  and both return a prediction of  $\mathcal{S} \subseteq \text{Distr}(\mathcal{T})$  ((**box-task-surv?**)). In addition, both simply account for censoring and can be utilised in fitting other models or to estimate unknown censoring distributions. The Kaplan-Meier and Nelson-Aalen estimators are both consistent estimators for the survival and cumulative hazard functions respectively.

Utilising the relationships provided in (?@sec-surv-set-types), one could write the Nelson-Aalen estimator in terms of the survival function as  $\hat{S}_{NA} = \exp(-\hat{H}(\tau | \mathcal{D}_{train}))$ . It has been demonstrated that  $\hat{S}_{NA}$  and  $\hat{S}_{KM}$  are asymptotically equivalent, but that  $\hat{S}_{NA}$  will provide larger estimates than  $\hat{S}_{KM}$  in smaller samples (Colosimo et al. 2002). In practice, the Kaplan-Meier is the most widely utilised non-parametric estimator in survival analysis and is the simplest estimator that yields consistent estimation of a survival distribution; it is therefore a natural, and commonly utilised, ‘baseline’ model (Binder and Schumacher 2008; Herrmann et al. 2021; Huang et al. 2020a; P. Wang, Li, and Reddy 2019); estimators that other models should be ‘judged’ against to ascertain their overall performance (Chapter 6).

Not only can these estimators be used for analytical comparison, but they also provide intuitive methods for graphical calibration of models (Section 8.2). These models are never studied for prognosis directly but as baselines, components of complex models (Chapter 18), or graphical tools (Habibi et al. 2018; Jager et al. 2008; Moghimi-dehkordi et al. 2008). The reason for this is due to them having poor predictive performance as a result of omitting explanatory variables in fitting. Moreover, if the data follows a particular distribution, parametric methods will be more efficient (P. Wang, Li, and Reddy 2019).

### Conditional Estimators

The Kaplan-Meier and Nelson-Aalen estimators are simple to compute and provide good estimates for the survival time distribution but in many cases they may be overly-simplistic. Conditional non-parametric estimators include the advantages described above (no assumptions about underlying data distribution) but also allow for conditioning the estimation on the covariates. This is particularly useful when estimating a censoring distribution that may depend on the data (Chapter 6). However predictive performance of conditional non-parametric estimators decreases as the number of covariates increases, and these models are especially poor when censoring is feature-dependent (Gerds and Schumacher 2006).

The most widely used conditional non-parametric estimator for survival analysis is the Akritas estimator (Michael G. Akritas 1994) defined by<sup>1</sup>

$$\hat{S}(\tau|X^*, \mathcal{D}_{train}, \lambda) = \prod_{j:T_j \leq \tau, \Delta_j=1} \left( 1 - \frac{K(X^*, X_j|\lambda)}{\sum_{l=1}^n K(X^*, X_l|\lambda) \mathbb{I}(T_l \geq T_j)} \right)$$

where  $K$  is a kernel function, usually  $K(x, y|\lambda) = \mathbb{I}(|\hat{F}_X(x) - \hat{F}_X(y)| < \lambda), \lambda \in (0, 1]$ ,  $\hat{F}_X$  is the empirical distribution function of the training data,  $X_1, \dots, X_n$ , and  $\lambda$  is a hyper-parameter. The estimator can be interpreted as a conditional Kaplan-Meier estimator which is computed on a neighbourhood of subjects closest to  $X^*$  (Blanche, Dartigues, and Jacqmin-Gadda 2013). To account for tied survival times, the following adaptation of the estimator is utilised (Blanche, Dartigues, and Jacqmin-Gadda 2013)

$$\hat{S}(\tau|X^*, \mathcal{D}_{train}, \lambda) = \prod_{t \in \mathcal{U}_O, t \leq \tau} \left( 1 - \frac{\sum_{j=1}^n K(X^*, X_j|\lambda) \mathbb{I}(T_j = t, \Delta_j = 1)}{\sum_{j=1}^n K(X^*, X_j|\lambda) \mathbb{I}(T_j \geq t)} \right) \quad (12.3)$$

If  $\lambda = 1$  then  $K(\cdot|\lambda) = 1$  and the estimator is identical to the Kaplan-Meier estimator.

The non-parametric nature of the model is highlighted in (Equation 12.3), in which both the fitting and predicting stages are combined into a single equation. A new observation,  $X^*$ , is compared to its nearest neighbours from a training dataset,  $\mathcal{D}_{train}$ , without a separated fitting procedure. One could consider splitting fitting and predicting in order to clearly separate between training and testing data. In this case, the fitting procedure is the estimation of  $\hat{F}_X$  on training data and the prediction is given by (Equation 12.3) with  $\hat{F}_X$  as an argument. This separated fit/predict method is implemented in **survivalmodels** (R. Sonabend 2020). As with other non-parametric estimators, the Akritas estimator can still be considered transparent and accessible. With respect to predictive performance, the Akritas estimator has more explanatory power than non-parametric estimators due to conditioning on covariates, however this is limited to a very small number of variables and therefore this estimator is still best placed as a conditional baseline.

### 12.1.2 Continuous Ranking and Semi-Parametric Models: Cox PH

The Cox Proportional Hazards (CPH) (Cox 1972), or Cox model, is likely the most widely known semi-parametric model and the most studied survival model (Habibi et al. 2018; Moghimi-dehkordi et al. 2008; Reid 1994; P. Wang, Li, and Reddy 2019). The Cox model assumes that the hazard for a subject is proportionally related to their explanatory variables,  $X_1, \dots, X_n$ , via some baseline hazard that all subjects in a given dataset share ('the PH assumption'). The hazard function in the Cox PH model is defined by

$$h(\tau|X_i) = h_0(\tau) \exp(X_i \beta)$$

where  $h_0$  is the non-negative *baseline hazard function* and  $\beta = \beta_1, \dots, \beta_p$  where  $\beta_i \in \mathbb{R}$  are coefficients to be fit. Note the proportional hazards (PH) assumption can be seen as the estimated hazard,  $h(\tau|X_i)$ , is directly proportional to the model covariates  $\exp(X_i \beta)$ . Whilst a form is assumed for the 'risk' component of the model,  $\exp(X_i \beta)$ , no assumptions are made about the distribution of  $h_0$ , hence the model is semi-parametric.

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<sup>1</sup>Arguments and parameters are separated in function signatures by a pipe, '|', where variables to the left are parameters (free variables) and those to the right are arguments (fixed). In this equation,  $\tau$  is a parameter to be set by the user, and  $X^*, \mathcal{D}_{train}, \lambda$  are fixed arguments. This could therefore be simplified to  $\hat{S}(\tau)$  to only include free variables.

The coefficients,  $\beta$ , are estimated by maximum likelihood estimation of the ‘partial likelihood’ (Cox 1975), which only makes use of ordered event times and does not utilise all data available (hence being ‘partial’). The partial likelihood allows study of the informative  $\beta$ -parameters whilst ignoring the nuisance  $h_0$ . The predicted linear predictor,  $\hat{\eta} := X^* \hat{\beta}$ , can be computed from the estimated  $\hat{\beta}$  to provide a ranking prediction.

Inspection of the model is also useful without specifying the full hazard by interpreting the coefficients as ‘hazard ratios’. Let  $p = 1$  and  $\hat{\beta} \in \mathbb{R}$  and let  $X_i, X_j \in \mathbb{R}$  be the covariates of two training observations, then the *hazard ratio* for these observations is the ratio of their hazard functions,

$$\frac{h(\tau|X_i)}{h(\tau|X_j)} = \frac{h_0(\tau) \exp(X_i \hat{\beta})}{h_0(\tau) \exp(X_j \hat{\beta})} = \exp(\hat{\beta})^{X_i - X_j}$$

If  $\exp(\hat{\beta}) = 1$  then  $h(\tau|X_i) = h(\tau|X_j)$  and thus the covariate has no effect on the hazard. If  $\exp(\hat{\beta}) > 1$  then  $X_i > X_j \rightarrow h(\tau|X_i) > h(\tau|X_j)$  and therefore the covariate is positively correlated with the hazard (increases risk of event). Finally if  $\exp(\hat{\beta}) < 1$  then  $X_i > X_j \rightarrow h(\tau|X_i) < h(\tau|X_j)$  and the covariate is negatively correlated with the hazard (decreases risk of event).

Interpreting hazard ratios is known to be a challenge, especially by clinicians who require simple statistics to communicate to patients (Sashegyi and Ferry 2017; Spruance et al. 2004). For example the full interpretation of a hazard ratio of ‘2’ for binary covariate  $X$  would be: ‘assuming that the risk of death is constant at all time-points then the instantaneous risk of death is twice as high in a patient with  $X$  than without’. Simple conclusions are limited to stating if patients are at more or less risk than others in their cohort. Further disadvantages of the model also lie in its lack of real-world interpretability, these include (Reid 1994):

- the PH assumption may not be realistic and the risk of event may not be constant over time;
- the estimated baseline hazard from a non-parametric estimator is a discrete step-function resulting in a discrete survival distribution prediction despite time being continuous; and
- the estimated baseline hazard will be constant after the last observed time-point in the training set (Gelfand et al. 2000).

Despite these disadvantages, the model has been demonstrated to have excellent predictive performance and routinely outperforms (or at least does not underperform) sophisticated ML models (Michael F. Gensheimer and Narasimhan 2018; Luxhoj and Shyur 1997; Van Belle et al. 2011) (and (R. E. B. Sonabend 2021)). It’s simple form and wide popularity mean that it is also highly transparent and accessible.

The next class of models address some of the Cox model disadvantages by making assumptions about the baseline hazard.

### 12.1.3 Conditional Distribution Predictions: Parametric Linear Models

#### Parametric Proportional Hazards

The CPH model can be extended to a fully parametric PH model by substituting the unknown baseline hazard,  $h_0$ , for a particular parameterisation. Common choices for distributions are Exponential, Weibull and Gompertz (John D. Kalbfleisch and Prentice 2011; P. Wang, Li, and Reddy 2019); their hazard functions are summarised in `((tab-survivalists?))` along with the respective parametric PH model. Whilst an Exponential assumption leads to the simplest hazard function, which is constant over time, this is often not realistic in

real-world applications. As such the Weibull or Gompertz distributions are often preferred. Moreover, when the shape parameter,  $\gamma$ , is 1 in the Weibull distribution or 0 in the Gompertz distribution, their hazards reduce to a constant risk ((Figure 12.1)). As this model is fully parametric, the model parameters can be fit with maximum likelihood estimation, with the likelihood dependent on the chosen distribution.

Table 12.2: Exponential, Weibull, and Gompertz hazard functions and PH specification.

Distribution <sup>1</sup>	$h_0(\tau)^2$	$h(\tau X_i)^3$
Exp( $\lambda$ )	$\lambda$	$\lambda \exp(X_i\beta)$
Weibull( $\gamma, \lambda$ )	$\lambda\gamma\tau^{\gamma-1}$	$\lambda\gamma\tau^{\gamma-1} \exp(X_i\beta)$
Gompertz( $\gamma, \lambda$ )	$\lambda \exp(\gamma\tau)$	$\lambda \exp(\gamma\tau) \exp(X_i\beta)$

\* 1. Distribution choices for baseline hazard.  $\gamma, \lambda$  are shape and scale parameters respectively.

\* 2. Baseline hazard function, which is the (unconditional) hazard of the distribution. \* 3. PH hazard function,  $h(\tau|X_i) = h_0(\tau) \exp(X_i\beta)$ .

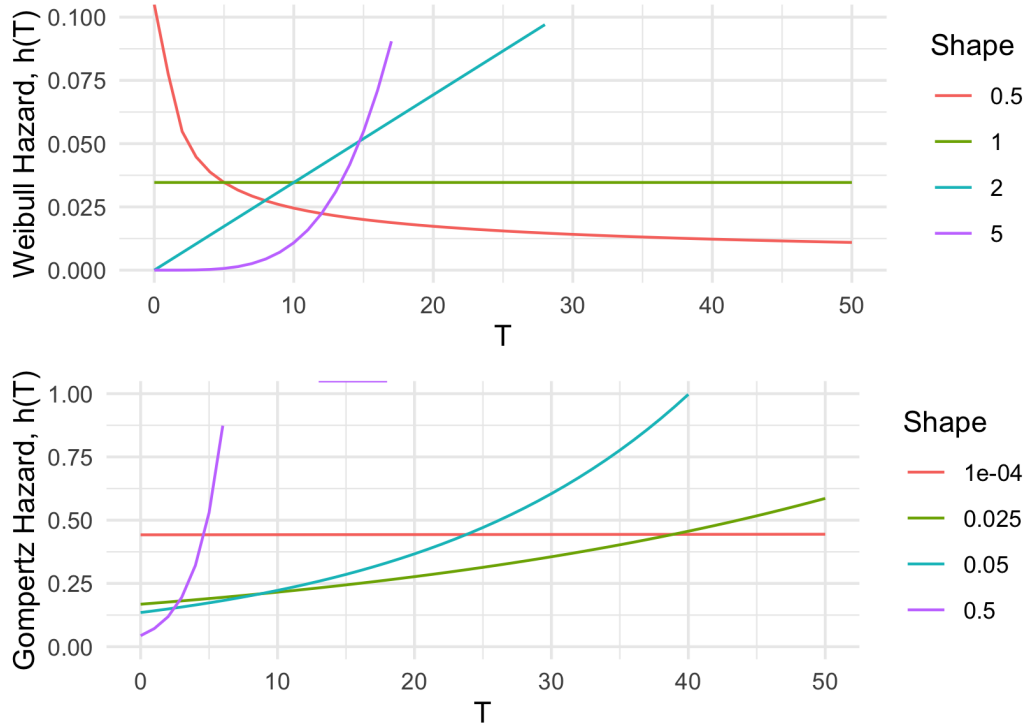


Figure 12.1: Comparing the hazard curves under Weibull and Gompertz distributions for varying values of the shape parameter; scale parameters are set so that each parametrisation has a median of 20. x-axes are time and y-axes are Weibull (top) and Gompertz (bottom) hazards as a function of time.

In the literature, the Weibull distribution tends to be favoured as the initial assumption for the survival distribution (Michael F. Gensheimer and Narasimhan 2018; Habibi et al. 2018; Hielscher et al. 2010; R. and J. 1968; Rahman et al. 2017), though Gompertz is often

tested in death-outcome models for its foundations in modelling human mortality (Gompertz 1825). There exist many tests for checking the goodness-of-model-fit (`?@sec-eval-insample`) and the distribution choice can even be treated as a model hyper-parameter. Moreover it transpires that model inference and predictions are largely insensitive to the choice of distribution (Collett 2014; Reid 1994). In contrast to the Cox model, fully parametric PH models can predict absolutely continuous survival distributions, they do not treat the baseline hazard as a nuisance, and in general will result in more precise and interpretable predictions if the distribution is correctly specified (Reid 1994; Patrick Royston and Parmar 2002).

Whilst misspecification of the distribution tends not to affect predictions too greatly, PH models will generally perform worse when the PH assumption is not valid. PH models can be extended to include time-varying coefficients or model stratification (Cox 1972) but even with these adaptations the model may not reflect reality. For example, the predicted hazard in a PH model will be either monotonically increasing or decreasing but there are many scenarios where this is not realistic, such as when recovering from a major operation where risks tends to increase in the short-term before decreasing. Accelerated failure time models overcome this disadvantage and allow more flexible modelling, discussed next.

### Accelerated Failure Time

In contrast to the PH assumption, where a unit increase in a covariate is a multiplicative increase in the hazard rate, the Accelerated Failure Time (AFT) assumption means that a unit increase in a covariate results in an acceleration or deceleration towards death (expanded on below). The hazard representation of an AFT model demonstrates how the interpretation of covariates differs from PH models,

$$h(\tau|X_i) = h_0(\exp(-X_i\beta)\tau) \exp(-X_i\beta)$$

where  $\beta = (\beta_1, \dots, \beta_p)$  are model coefficients. In contrast to PH models, the ‘risk’ component,  $\exp(-X_i\beta)$ , is the exponential of the *negative* linear predictor and therefore an increase in a covariate value results in a decrease of the predicted hazard. This representation also highlights how AFT models are more flexible than PH as the predicted hazard can be non-monotonic. For example the hazard of the Log-logistic distribution ((Figure 12.2)) is highly flexible depending on chosen parameters. Not only can the AFT model offer a wider range of shapes for the hazard function but it is more interpretable. Whereas covariates in a PH model act on the hazard, in an AFT they act on time, which is most clearly seen in the log-linear representation,

$$\log Y_i = \mu + \alpha_1 X_{i1} + \alpha_2 X_{i2} + \dots + \alpha_p X_{ip} + \sigma \epsilon_i$$

where  $\mu$  and  $\sigma$  are location and scale parameters respectively,  $\alpha_1, \dots, \alpha_p$  are model coefficients, and  $\epsilon_i$  is a random error term. In this case a one unit increase in covariate  $X_{ij}$  means a  $\alpha_j$  increase in the logarithmic survival time. For example if  $\exp(X_i\alpha) = 0.5$  then  $i$  ‘ages’ at double the baseline ‘speed’. Or less abstractly if studying the time until death from cancer then  $\exp(X_i\alpha) = 0.5$  can be interpreted as ‘the entire process from developing tumours to metastasis and eventual death in subject  $i$  is twice as fast than the normal’, where ‘normal’ refers to the baseline when all covariates are 0.

Specifying a particular distribution for  $\epsilon_i$  yields a fully-parametric AFT model. Common distribution choices include Weibull, Exponential, Log-logistic, and Log-Normal (John D. Kalbfleisch and Prentice 2011; P. Wang, Li, and Reddy 2019). The Buckley-James estimator (Buckley and James 1979) is a semi-parametric AFT model that non-parametrically estimates the distribution of the errors however this model has no theoretical justification and is rarely

fit in practice (Wei 1992). The fully-parametric model has theoretical justifications, natural interpretability, and can often provide a better fit than a PH model, especially when the PH assumption is violated (Patel, Kay, and Rowell 2006; Qi 2009; Zare et al. 2015).

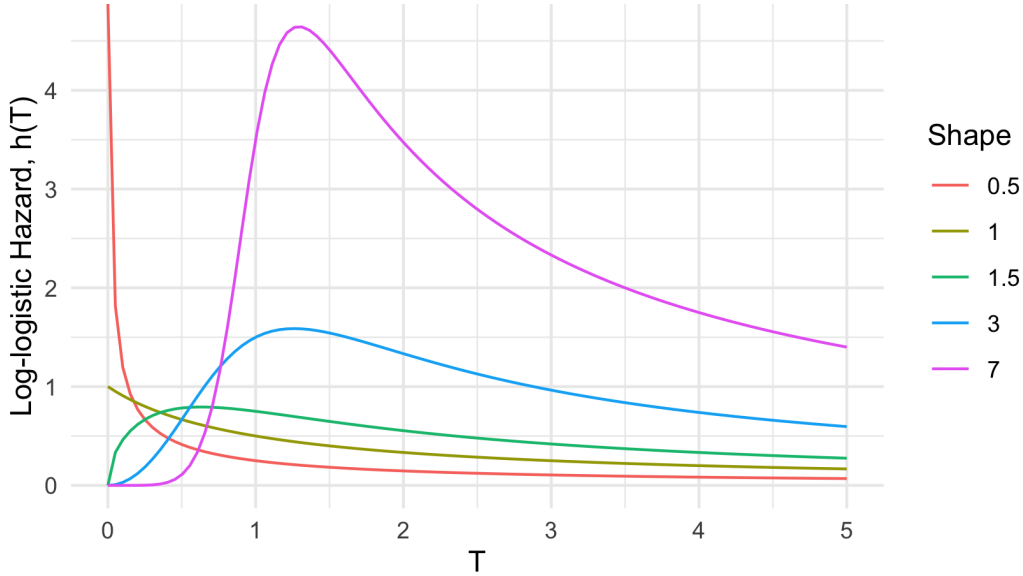


Figure 12.2: Log-logistic hazard curves with a fixed scale parameter of 1 and a changing shape parameter. x-axis is time and y-axis is the log-logistic hazard as a function of time.

### Proportional Odds

Proportional odds (PO) models (Bennett 1983) fit a proportional relationship between covariates and the odds of survival beyond a time  $\tau$ ,

$$O_i(\tau) = \frac{S_i(\tau)}{F_i(\tau)} = O_0(\tau) \exp(X_i\beta)$$

where  $O_0$  is the baseline odds.

In this model, a unit increase in a covariate is a multiplicative increase in the odds of survival after a given time and the model can be interpreted as estimating the log-odds ratio. There is no simple closed form expression for the partial likelihood of the PO model and hence in practice a Log-logistic distribution is usually assumed for the baseline odds and the model is fit by maximum likelihood estimation on the full likelihood (Bennett 1983).

Perhaps the most useful feature of the model is convergence of hazard functions (Kirmani and Gupta 2001), which states  $h_i(\tau)/h_0(\tau) \rightarrow 1$  as  $\tau \rightarrow \infty$ . This property accurately reflects real-world scenarios, for example if comparing chemotherapy treatment on advanced cancer survival rates, then it is expected that after a long period (say 10 years) the difference in risk between groups is likely to be negligible. This is in contrast to the PH model that assumes the hazard ratios are constant over time, which is rarely a reflection of reality.

In practice, the PO model is harder to fit and is less flexible than PH and AFT models, both of which can also produce odds ratios. This may be a reason for the lack of popularity of the PO model, in addition there is limited off-shelf implementations (Collett 2014). Despite

PO models not being commonly utilised, they have formed useful components of neural networks ([?@sec-surv-ml-models-nn](#)) and flexible parametric models (below).

### Flexible Parametric Models – Splines

Royston-Parmar flexible parametric models (Patrick Royston and Parmar 2002) extend PH and PO models by estimating the baseline hazard with natural cubic splines. The model was designed to keep the form of the PH or PO methods but without the semi-parametric problem of estimating a baseline hazard that does not reflect reality (see above), or the parametric problem of misspecifying the survival distribution.

To provide an interpretable, informative and smooth hazard, natural cubic splines are fit in place of the baseline hazard. The crux of the method is to use splines to model time on a log-scale and to either estimate the log cumulative Hazard for PH models,  $\log H(\tau|X_i) = \log H_0(\tau) + X_i\beta$ , or the log Odds for PO models,  $\log O(\tau|X_i) = \log O_0(\tau) + X_i\beta$ , where  $\beta$  are model coefficients to fit,  $H_0$  is the baseline cumulative hazard function and  $O_0$  is the baseline odds function. For the flexible PH model, a Weibull distribution is the basis for the baseline distribution and a Log-logistic distribution for the baseline odds in the flexible PO model.  $\log H_0(\tau)$  and  $\log O_0(\tau)$  are estimated by natural cubic splines with coefficients fit by maximum likelihood estimation. The standard full likelihood is optimised, full details are not provided here. Between one and three internal knots are recommended for the splines and the placement of knots does not greatly impact upon the fitted model (Patrick Royston and Parmar 2002).

Advantages of the model include being: interpretable, flexible, can be fit with time-dependent covariates, and it returns a continuous function. Moreover many of the parameters, including the number and position of knots, are tunable, although Royston and Parmar advised against tuning and suggest often only one internal knot is required (Patrick Royston and Parmar 2002). A recent simulation study demonstrated that even with an increased number of knots (up to seven degrees of freedom), there was little bias in estimation of the survival and hazard functions (Bower et al. 2019). Despite its advantages, a 2018 review (Ng et al. 2018) found only twelve instances of published flexible parametric models since Royston and Parmar's 2002 paper, perhaps because it is more complex to train, has a less intuitive fitting procedure than alternatives, and has limited off-shelf implementations; i.e. is less transparent and accessible than parametric alternatives.

The PH and AFT models are both very transparent and accessible, though require slightly more expert knowledge than the CPH in order to specify the ‘correct’ underlying probability distribution. Interestingly whilst there are many papers comparing PH and AFT models to one another using in-sample metrics ([?@sec-eval-insample](#)) such as AIC (Georgousopoulou et al. 2015; Habibi et al. 2018; Moghimi-dehkordi et al. 2008; Zare et al. 2015), no benchmark experiments could be found for out-of-sample performance. PO and spline models are less transparent than PH and AFT models and are even less accessible, with very few implementations of either. No conclusions can be drawn about the predictive performance of PO or spline models due to a lack of suitable benchmark experiments.



# 13

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## Random Forests

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TODO (150-200 WORDS)

 Minor changes expected!

This page is a work in progress and minor changes will be made over time.

Random forests are a composite (or ensemble) algorithm built by fitting many simpler component models, *decision trees*, and then averaging the results of predictions from these trees. Due to in-built variable importance properties, random forests are commonly used in high-dimensional settings when the number of variables in a dataset far exceeds the number of rows. High-dimensional datasets are very common in survival analysis, especially when considering omics, genetic and financial data. It is therefore no surprise that *random survival forests*, remain a popular and well-performing model in the survival setting.

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### 13.1 Random Forests for Regression

Training of decision trees can include a large number of hyper-parameters and different training steps including ‘growing’ and subsequently ‘pruning’. However, when utilised in random forests, many of these parameters and steps can be safely ignored, hence this section only focuses on the components that primarily influence the resulting random forest. This section will start by discussing decision trees and will then introduce the *bagging* algorithm used to create random forests.

#### 13.1.1 Decision Trees

*Decision trees* are a (relatively) simple machine learning model that are comparatively easy to implement in software and are highly interpretable. The decision tree algorithm takes an input, a dataset, selects a variable that is used to partition the data according to some *splitting rule* into distinct non-overlapping datasets or *nodes* or *leaves*, and repeats this step for the resulting partitions, or *branches*, until some criterion has been reached. The final nodes are referred to as *terminal nodes*.

By example, (Figure 13.1) demonstrates a decision tree predicting the price that a car sells for in India (price in thousands of dollars). The dataset includes as variables the registration year, kilometers driven, fuel type (petrol or automatic), seller type (individual or dealer), transmission type (manual or automatic), and number of owners. The decision tree was

trained with a maximum depth of 2 (the number of rows excluding the top), and it can be seen that with this restriction only the transmission type, registration year, and fuel type were selected variables. During training, the algorithm identified that the first optimal variable to split the data was transmission type, partitioning the data into manual and automatic cars. Manual cars are further subset by registration year whereas automatic cars are split by fuel type. It can also be seen how the average sale price (top value in each leaf) diverges between leaves as the tree splits – the average sale prices in the final leaves are the terminal node predictions.

The graphic highlights several core features of decision trees:

1. They can model non-linear and interaction effects: The hierarchical structure allows for complex interactions between variables with some variables being used to separate all observations (transmission) and others only applied to subsets (year and fuel);
2. They are highly interpretable: it is easy to visualise the tree and see how predictions are made;
3. They perform variable selection: not all variables were used to train the model.

To understand how random forests work, it is worth looking a bit more into the most important components of decision trees: splitting rules, stopping rules, and terminal node predictions.

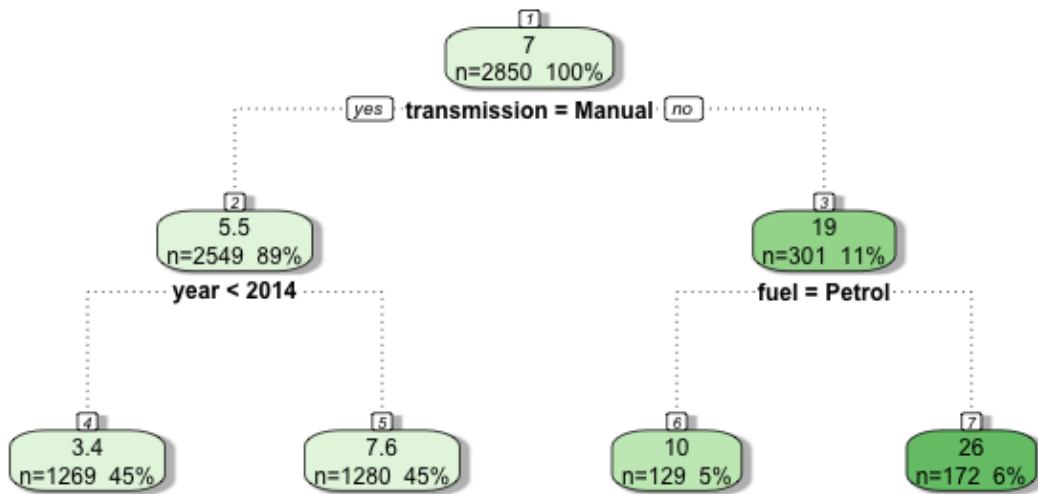


Figure 13.1: Predicting the price a vehicle is sold for in India using a regression tree, dataset from kaggle.com/datasets/nehalbirla/vehicle-dataset-from-cardekho. Rounded rectangles are leaves, which indicate the variable that is being split. Edges are branches, which indicate the cut-off at which the variable is split. Variables are car transmission type (manual or automatic), fuel type (petrol or diesel) and registration year. The number at the top of each leaf is the average selling price in thousands of dollars for all observations in that leaf. The numbers at the bottom of each leaf are the number of observations in the leaf, and the proportion of data contained in the leaf.

### Splitting and Stopping Rules

Precisely how the data partitions/splits are derived and which variables are utilised is determined by the *splitting rule*. The goal in each partition is to find two resulting leaves/nodes that have the greatest difference between them and thus the maximal homogeneity within each leaf, hence with each split, the data in each node should become increasingly similar. The splitting rule provides a way to measure the homogeneity within the resulting nodes. In regression, the most common splitting rule is to select a variable and cut-off (a threshold on the variable at which to separate observations) that minimises the mean squared error in the two potential resulting leaves.

For all decision tree and random forest algorithms going forward, let  $L$  denote some leaf, then let  $L_{xy}, L_x, L_y$  respectively be the set of observations, features, and outcomes in leaf  $L$ . Let  $L_{y;i}$  be the  $i$ th outcome in  $L_y$  and finally let  $\bar{y}_L = \frac{1}{|L_y|} \sum_{i=1}^{|L_y|} L_{y;i}$  be the mean outcome in leaf  $L$ .

Let  $j = 1, \dots, p$  be the index of features and let  $c_j$  be a possible cutoff value for feature  $\mathbf{x}_{;j}$ . Define

$$\begin{aligned} L_{xy}^a(j, c_j) &:= \{(\mathbf{x}_i, y_i) | x_{i;j} < c_j, i = 1, \dots, n\} \\ L_{xy}^b(j, c_j) &:= \{(\mathbf{x}_i, y_i) | x_{i;j} \geq c_j, i = 1, \dots, n\} \end{aligned}$$

as the two leaves containing the set of observations resulting from partitioning variable  $j$  at cutoff  $c_j$ . To simplify equations let  $L^a, L^b$  be shorthands for  $L^a(j, c_j)$  and  $L^b(j, c_j)$ . Then a split is determined by finding the arguments,  $(j^*, c_{j^*}^*)$  that minimise the residual sum of squares across both leaves (James et al. 2013),

$$(j^*, c_{j^*}^*) = \arg \min_{j, c_j} \sum_{y \in L_y^a} (y - \bar{y}_{L^a})^2 + \sum_{y \in L_y^b} (y - \bar{y}_{L^b})^2 \quad (13.1)$$

This method is repeated from the first leaf to the last such that observations are included in a given leaf  $L$  if they satisfy all conditions from all previous branches (splits); features may be considered multiple times in the growing process allowing complex interaction effects to be captured.

Leaves are repeatedly split until a *stopping rule* has been triggered – a criterion that tells the algorithm to stop partitioning data. The stopping rule is usually a condition on the number of observations in each leaf such that leaves will continue to be split until some minimum number of observations has been reached in a leaf. Other conditions may be on the depth of the tree (as in Figure 13.1 which is restricted to a maximum depth of 2), which corresponds to the number of levels of splitting. Stopping rules are often used together, for example by setting a maximum tree depth *and* determining a minimum number of observations per leaf. Deciding the number of minimum observations and/or the maximum depth can be performed with automated hyper-parameter optimisation.

### Terminal Node Predictions

The final major component of decision trees are *terminal node predictions*. As the name suggests, this is the part of the algorithm that determines how to actually make a prediction for a new observation. A prediction is made by ‘dropping’ the new data ‘down’ the tree according to the optimal splits that were found during training. The resulting prediction is then a simple baseline statistic computed from the training data that fell into the corresponding node. In regression, this is commonly the sample mean of the training outcome data.

Returning to Figure 13.1, say a new data point is {transmission = Manual, fuel = Diesel, year = 2015}, then in the first split the left branch is taken as ‘transmission = Manual’, in the second split the right branch is taken as ‘year’ = 2015 ≥ 2014, hence the new data point lands in the second terminal leaf and is predicted to sell for \$7,600. The ‘fuel’ variable is ignored as it is only considered for automatic vehicles. As the final predictions are simple statistics based on training data, all potential predictions can be saved in the original trained model and no complex computations are required in the prediction algorithm.

### 13.1.2 Random Forests

Decision trees often overfit the training data, hence they have high variance, perform poorly on new data, and are not robust to even small changes in the original training data. Moreover, important variables can end up being ignored as only subsets of dominant variables are selected for splitting.

To counter these problems, *random forests* are designed to improve prediction accuracy and decrease variance. Random forests utilise bootstrap aggregation, or *bagging* (Leo Breiman 1996), to aggregate many decision trees. Bagging is a relatively simple algorithm, as follows:

1. **For**  $b = 1, \dots, B$ :
2.  $D_b \leftarrow$  Randomly sample with replacement  $\mathcal{D}_{train}$
3.  $\hat{g}_b \leftarrow$  Train a decision tree on  $D_b$
4. **end For**
5. **return**  $\{\hat{g}_b\}_{b=1}^B$

Step 2 is known as *bootstrapping*, which is the process of sampling a dataset *with* replacement – which is in contrast to more standard subsampling where data is sampled *without* replacement. Commonly, the bootstrapped sample size is the same as the original. However, as the same value may be sampled multiple times, on average the resulting data only contains around 63.2% unique observations (Efron and Tibshirani 1997). Randomness is further injected to decorrelate the trees by randomly subsetting the candidates of features to consider at each split of a tree. Therefore, every split of every tree may consider a different subset of variables. This process is repeated for  $B$  trees, with the final output being a collection of trained decision trees.

Prediction from a random forest for new data  $\mathbf{x}^*$  follows by making predictions from the individual trees and aggregating the results by some function  $\sigma$ , which is usually the sample mean for regression:

$$\hat{g}(\mathbf{x}^*) = \sigma(\hat{g}_1(\mathbf{x}^*), \dots, \hat{g}_B(\mathbf{x}^*)) = \frac{1}{B} \sum_{b=1}^B \hat{g}_b(\mathbf{x}^*)$$

where  $\hat{g}_b(\mathbf{x}^*)$  is the prediction from the  $b$ th tree for some new data  $\mathbf{x}^*$  and  $B$  are the total number of grown trees.

As discussed above, individual decision trees result in predictions with high variance that are not robust to small changes in the underlying data. Random forests decrease this variance by aggregating predictions over a large sample of decorrelated trees, where a high degree of difference between trees is promoted through the use of bootstrapped samples and random candidate feature selection at each split.

Usually many (hundreds or thousands) trees are grown, which makes random forests robust to changes in data and ‘confident’ about individual predictions. Other advantages include having tunable and meaningful hyper-parameters, including: the number of variables to consider for a single tree, the splitting rule, and the stopping rule. Random forests treat trees as *weak learners*, which are not intended to be individually optimized. Instead, each tree captures a small amount of information about the data, which are combined to form a powerful representation of the dataset.

Whilst random forests are considered a ‘black-box’, in that one cannot be reasonably expected to inspect thousands of individual trees, variable importance can still be aggregated across trees, for example by counting the frequency a variable was selected across trees, calculating the minimal depth at which a variable was used for splitting, or via permutation based feature importance. Hence the model remains more interpretable than many alternative methods. Finally, random forests are less prone to overfitting and this can be relatively easily controlled by using *early-stopping* methods, for example by continually growing trees until the performance of the model stops improving.

## 13.2 Random Survival Forests

Unlike other machine learning methods that may require complex changes to underlying algorithms, random forests can be relatively simply adapted to *random survival forests* by updating the splitting rules and terminal node predictions to those that can handle censoring and can make survival predictions. This chapter is therefore focused on outlining different choices of splitting rules and terminal node predictions, which can then be flexibly combined into different models.

### 13.2.1 Splitting Rules

Survival trees and RSFs have been studied for the past four decades and whilst there are many possible splitting rules (Bou-Hamad, Larocque, and Ben-Ameur 2011), only two broad classes are commonly utilised (as judged by number of available implementations, e.g., Pölsterl (2020); Wright and Ziegler (2017); H. Ishwaran et al. (2011)). The first class rely on hypothesis tests, and primarily the log-rank test, to maximise dissimilarity between splits, the second class utilises likelihood-based measures. The first is discussed in more detail as this is common in practice and is relatively straightforward to implement and understand, moreover it has been demonstrated to outperform other splitting rules (Bou-Hamad, Larocque, and Ben-Ameur 2011). Likelihood rules are more complex and require assumptions that may not be realistic, these are discussed briefly.

#### Hypothesis Tests

The log-rank test statistic has been widely utilized as a splitting-rule for survival analysis (Ciampi et al. 1986; B. H. Ishwaran et al. 2008; LeBlanc and Crowley 1993; Segal 1988). The log-rank test compares the survival distributions of two groups under the null-hypothesis that both groups have the same underlying risk of (immediate) events, with the hazard function used to compare underlying risk.

Let  $L^a$  and  $L^b$  be two leaves and let  $h^a, h^b$  be the (theoretical, true) hazard functions in the two leaves respectively and let  $i \in L$  be a shorthand for the indices of the observations in

leaf  $L$  so that  $i = 1, \dots, |L|$ . Define:

- $\mathcal{U}_D$ , the set of unique event times across the data (in both leaves)
- $n_\tau^a$ , the number of observations at risk at  $\tau$  in leaf  $a$

$$n_\tau^a = \sum_{i \in L^a} \mathbb{I}(t_i \geq \tau)$$

- $o_\tau^a$ , the observed number of events in leaf  $a$  at  $\tau$

$$o_\tau^a = \sum_{i \in L^a} \mathbb{I}(t_i = \tau, \delta_i = 1)$$

- $n_\tau = n_\tau^a + n_\tau^b$ , the number of observations at risk at  $\tau$  in both leaves
- $o_\tau = o_\tau^a + o_\tau^b$ , the observed number of events at  $\tau$  in both leaves

Then, the log-rank hypothesis test is given by  $H_0 : h^a = h^b$  with test statistic (Segal 1988),

$$LR(L^a) = \frac{\sum_{\tau \in \mathcal{U}_D} (o_\tau^a - e_\tau^a)}{\sqrt{\sum_{\tau \in \mathcal{U}_D} v_\tau^a}}$$

where:

- $e_\tau^a$  is the expected number of events in leaf  $a$  at  $\tau$

$$e_\tau^a := \frac{n_\tau^a o_\tau}{n_\tau}$$

- $v_\tau^a$  is the variance of the number of events in leaf  $a$  at  $\tau$

$$v_\tau^a := e_\tau^a \left( \frac{n_\tau - o_\tau}{n_\tau} \right) \left( \frac{n_\tau - n_\tau^a}{n_\tau - 1} \right)$$

These results follow as under the assumption of equal hazards in both leafs, the number of events at each  $\tau \in \mathcal{U}_D$  is distributed according to a Hypergeometric distribution. The same statistic results if  $L^b$  is instead considered.

The higher the log-rank statistic, the greater the dissimilarity between the two groups (Figure 13.2), thereby making it a sensible splitting rule for survival, moreover it has been shown that it works well for splitting censored data (LeBlanc and Crowley 1993). Additionally, the log-rank test requires no knowledge about the shape of the survival curves or distribution of the outcomes in either group (Bland and Altman 2004), making it ideal for an automated process that requires no user intervention.

The log-rank *score* rule (Hothorn and Lausen 2003) is a standardized version of the log-rank rule that could be considered as a splitting rule, though simulation studies have demonstrated non-significant improvements in predictive performance when comparing the two (B. H. Ishwaran et al. 2008). Alternative dissimilarity measures and tests have also been suggested as splitting rules, including modified Kolmogorov-Smirnov test and Gehan-Wilcoxon tests (Ciampi et al. 1988). Simulation studies have demonstrated that both of these may have higher power and produce ‘better’ results than the log-rank statistic (Fleming et al. 1980), however neither appears to be commonly used.

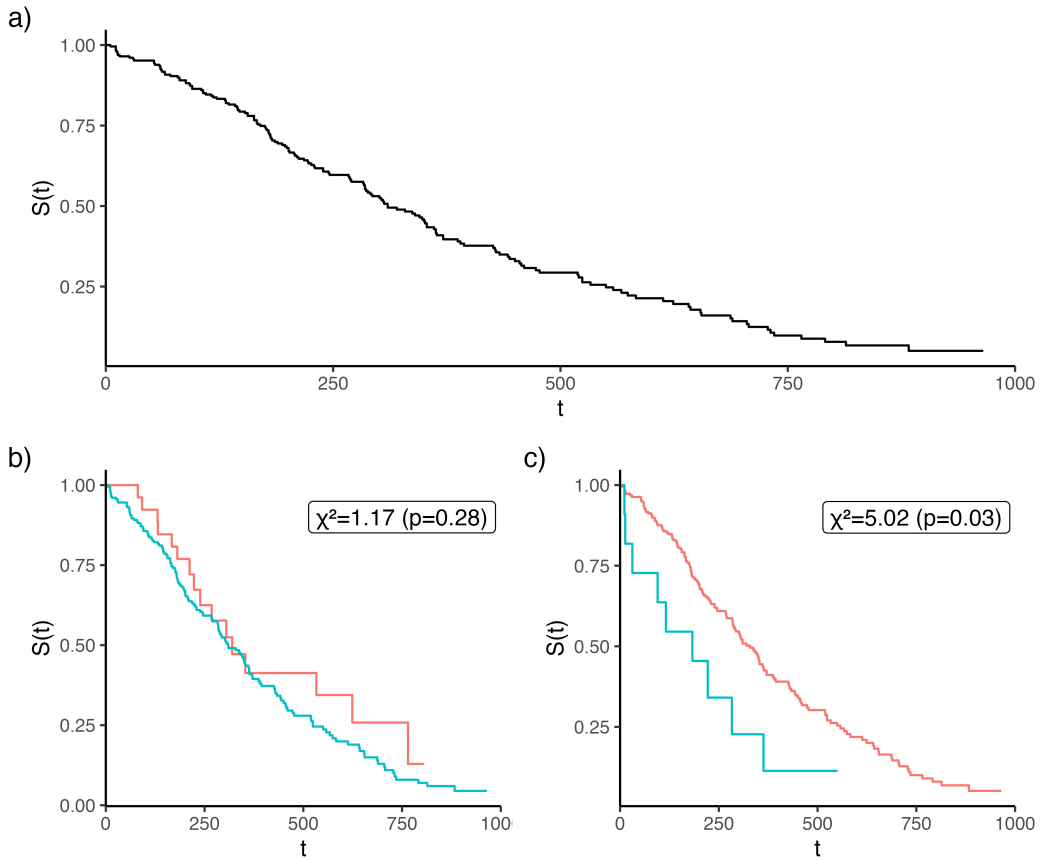


Figure 13.2: Panel (a) is the Kaplan-Meier estimator fit on the complete `lung` dataset from the R package `survival`. (b-c) is the same data stratified according to whether ‘age’ is greater or less than 50 (panel b) or 75 (panel c). The higher  $\chi^2$  statistic (panel c) results in a lower  $p$ -value and a greater difference between the stratified Kaplan-Meier curves. Hence splitting age at 75 results in a greater dissimilarity between the resulting branches and thus makes a better choice for splitting the variable.

In a competing risk setting, Gray's test (Gray 1988) can be used instead of the log-rank test, as it compares cumulative incidence functions rather than all-cause hazards. Similarly to the log-rank test, Gray's test also compares survival distributions using hypothesis tests to determine if there are significant differences between the groups, thus making it a suitable option to build competing risk RSFs.

### Alternative Splitting Rules

A common alternative to the log-rank test is to instead use *likelihood ratio*, or *deviance*, statistics. When building RSFs, the likelihood-ratio statistic can be used to test if the model fit is improved or worsened with each split, thus providing a way to partition data. However, as discussed in `?@sec-surv-obj`, there are many different likelihoods that can be assumed for survival data, and there is no obvious way to determine if one is more sensible than another. Furthermore the choice of likelihood must fit the underlying model assumptions. For example, one could assume the data fits the proportional hazards assumption and in each split one could calculate the likelihood-ratio using the Cox PH partial likelihood. Alternatively, a parametric form could be assumed and a likelihood proposed by LeBlanc and Crowley (1992) may be calculated to test model fit. While potentially useful, these methods are implemented in very few off-shelf software packages, thus empirical comparisons to other splitting rules are lacking.

Other rules have also been studied including comparison of residuals (Therneau, Grambsch, and Fleming 1990), scoring rules (H. Ishwaran and Kogalur 2018), distance metrics (Gordon and Olshen 1985), and concordance metrics (Schmid, Wright, and Ziegler 2016). Experiments have shown different splitting rules may perform better or worse depending on the underlying data (Schmid, Wright, and Ziegler 2016), hence one could even consider treating the splitting rule as a hyper-parameter for tuning. However, if there is a clear goal in prediction, then the choice of splitting rule can be informed by the prediction type. For example, if the goal is to maximise separation, then a log-rank splitting rule to maximise homogeneity in terminal nodes is a natural starting point. Whereas if the goal is to accurately rank observations, then a concordance splitting rule may be optimal.

#### 13.2.2 Terminal Node Prediction

As in the regression setting, the usual strategy for predictions is to create a simple estimate based on the training data that lands in the terminal nodes. However, as seen throughout this book, the choice of estimator in the survival setting depends on the prediction task of interest, which are now considered in turn. First, note that all terminal node predictions can only yield useful results if there are a sufficient number of uncensored observations in each terminal node. Hence, a common RSF stopping rule is the minimum number of *uncensored* observations per leaf, meaning a leaf is not split if that would result in too few uncensored observations in the resulting leaves.

#### Probabilistic Predictions

Starting with the most common survival prediction type, the algorithm requires a simple estimate for the underlying survival distribution in each of the terminal nodes, which can be estimated using the Kaplan-Meier or Nelson-Aalen methods (Hothorn et al. 2004; B. H. Ishwaran et al. 2008; LeBlanc and Crowley 1993; Segal 1988).

Denote  $b$  as a decision tree and  $L^{b(h)}$  as the terminal node  $h$  in tree  $b$ . Then the predicted survival function and cumulative hazard for a new observation  $\mathbf{x}^*$  is,

$$\hat{S}_{b(h)}(\tau | \mathbf{x}^*) = \prod_{i:t_{(i)} \leq \tau} 1 - \frac{d_{t_{(i)}}}{n_{t_{(i)}}}, \quad \{i \in L^{b(h)} : \mathbf{x}^* \in L^{b(h)}\} \quad (13.2)$$

$$\hat{H}_{b(h)}(\tau | \mathbf{x}^*) = \sum_{i:t_{(i)} \leq \tau} \frac{d_{t_{(i)}}}{n_{t_{(i)}}}, \quad \{i \in L^{b(h)} : \mathbf{x}^* \in L^{b(h)}\} \quad (13.3)$$

where  $t_{(i)}$  is the ordered event times and  $d_{t_{(i)}}$  and  $n_{t_{(i)}}$  are the observed number of events, and the number of observations at risk, respectively at  $t_{(i)}$ . See Figure 13.3 for an example using the `lung` dataset (Therneau 2015).

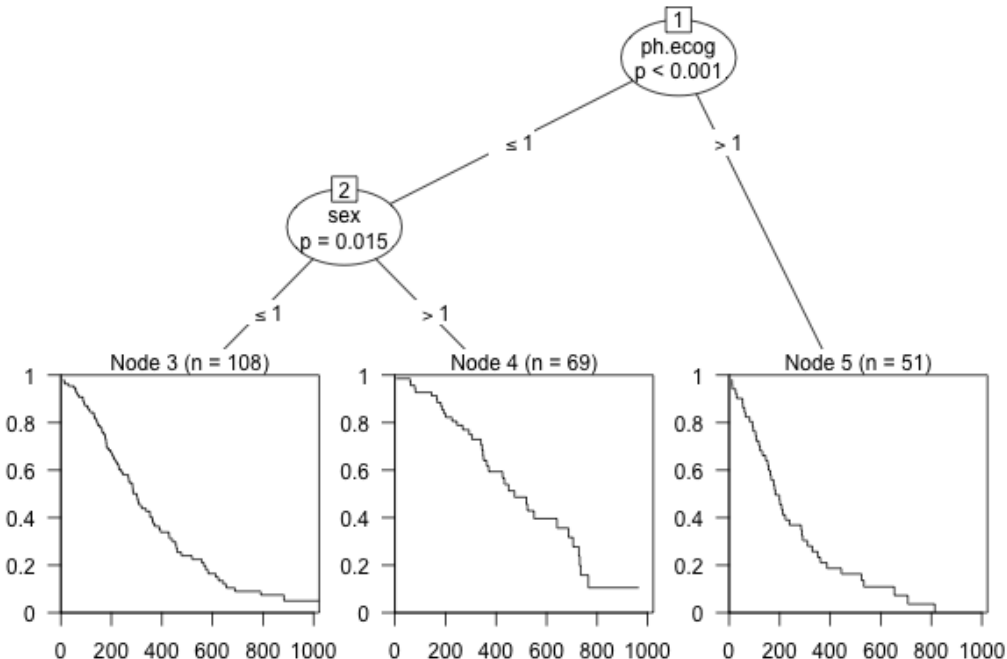


Figure 13.3: Survival tree trained on the `lung` dataset from the R package **survival**. The terminal node predictions are survival curves.

The bootstrapped prediction is the cumulative hazard function or survival function averaged over individual trees. Note that understanding what these bootstrapped functions represents depends on how they are calculated. By definition, a mixture of  $n$  distributions with cumulative distribution functions  $F_i, i = 1, \dots, n$  is given by

$$F(x) = \sum_{i=1}^n w_i F_i(x)$$

Substituting  $F = 1 - S$  and noting  $\sum w_i = 1$  gives the computation  $S(x) = \sum_{i=1}^n w_i S_i(x)$ , allowing the bootstrapped survival function to exactly represent the mixture distribution averaged over all trees:

$$\hat{S}_{Boot}(\tau|\mathbf{x}^*) = \frac{1}{B} \sum_{b=1}^B w_i \hat{S}_b(\tau|\mathbf{x}^*) \quad (13.4)$$

usually with  $w_i = 1/B$  where  $B$  is the number of trees.

In contrast, if one were to instead substitute  $F = 1 - \exp(-H)$ , then the mixture distribution depends on a logarithmic function that can only be approximately computed if predicted survival probabilities are close to one, which is an assumption that deteriorates over time. Therefore, to ensure the bootstrapped prediction accurately represents the underlying mixed probability distribution, the bootstrapped cumulative hazard function should be computed as:

$$\hat{H}_{Boot}(\tau|\mathbf{x}^*) = -\log(\hat{S}_{Boot}(\tau|\mathbf{x}^*)) \quad (13.5)$$

Another practical consideration to take into account is how to average the survival probabilities over the decision trees as each individual Kaplan-Meier estimate may have been trained on different time points. This is overcome by recognising that the Kaplan-Meier estimation results in a piece-wise function that can be linearly interpolated between training data. Figure 13.4 demonstrates this process for three decision trees (panel a), where the survival probability is calculated at all possible time points (panels b-c), and the average is computed with linear interpolation added between time-points (panel d).

Extensions to competing risks follow naturally using bootstrapped cause-specific cumulative incidence functions.

### Deterministic Predictions

As discussed in Chapter 10, predicting and evaluating survival times is a complex and fairly under-researched area. For RSFs, there is an inclination to estimate survival times based on the mean or median survival times of observations in terminal nodes, however this would lead to biased estimations. Therefore, research has tended to focus on relative risk predictions.

As discussed, relative risks are arbitrary values where only the resulting rank matters when comparing observations. In RSFs, each terminal node should be as homogeneous as possible, hence within a terminal node, the risk between observations should be the same. The most common method to estimate average risk appears to be a transformation from the Nelson-Aalen method (B. H. Ishwaran et al. 2008), which exploits results from counting process to provide a measure of expected mortality (Hosmer Jr, Lemeshow, and May 2011) – the same result is used in the van Houwelingen calibration measure discussed in Section 8.1.2. Given new data,  $\mathbf{x}^*$ , falling into terminal node  $b(h)$ , the relative risk prediction is the sum of the predicted cumulative hazard,  $\hat{H}_{b(h)}$ , computed at each observation's observed outcome time:

$$\phi_{b(h)}(\mathbf{x}^*) = \sum_{i \in \mathcal{U}_O} \hat{H}_{b(h)}(t_i|\mathbf{x}^*)$$

where  $\hat{H}_{b(h)}$  is the terminal node prediction as in Equation 13.3. This is interpreted as the number of expected events in  $b(h)$  and the assumption is that a terminal node with more expected events is a higher risk group than a node with less expected events. The bootstrapped risk prediction is the sample mean over all trees:

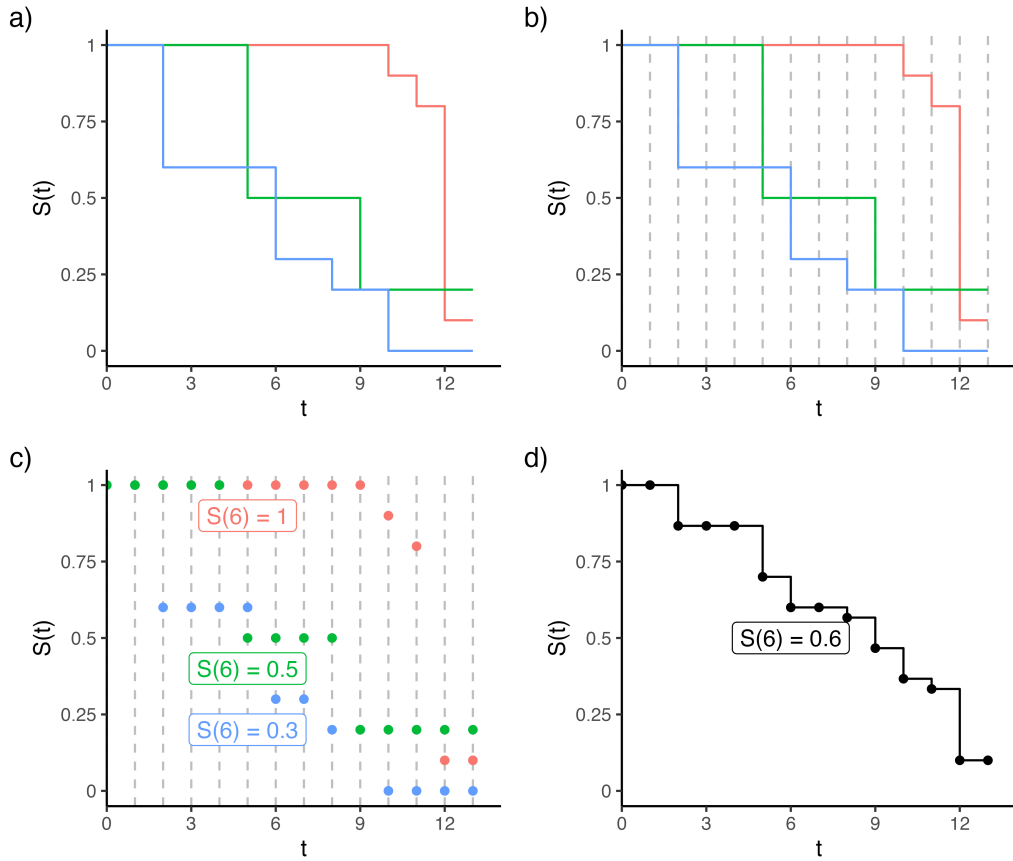


Figure 13.4: Bootstrapping Kaplan-Meier estimators across three decision trees (red, blue, green). Panel a) shows the individual estimates, b) shows the time points to aggregate the trees over, c) is the predicted survival probability from each tree at the desired time points, and d) is the average survival probabilities connected by a step function.

$$\phi_{Boot}(\mathbf{x}^*) = \frac{1}{B} \sum_{b=1}^B \phi_{b(h)}(\mathbf{x}^*)$$

More complex methods have also been proposed that are based on the likelihood-based splitting rule and assume a PH model form (H. Ishwaran et al. 2004; LeBlanc and Crowley 1992). However, these do not appear to be in wide-spread usage.

### 13.3 Conclusion

#### Key takeaways

- Random forests are a highly flexible algorithm that allow the various components to be adapted and altered without major changes to the underlying algorithm. This allows random survival forests (RSFs) to be readily available ‘off-shelf’ in many open-source packages;
- RSFs have in-built variable selection methods that mean they tend to perform well on high-dimensional data, routinely outperforming other models Burk et al. (2024);
- Despite having many potential hyper-parameters to tune, all are intuitive and many can even be ignored as sensible defaults exist in most off-shelf software implementations.

#### Limitations

- Due to the number of trees and the constant bootstrapping procedures, RSFs can be more computationally intensive than other models, though still much less intensive than neural networks and other deep learning methods.
- Despite having some in-built methods for model interpretation, RSFs are still black-boxes that can be difficult to fully interpret.
- With too few trees random forests can have similar limitations to decision trees and with too many random forests can overfit the data. Though most software has sensible defaults to prevent either scenario.

#### Further reading

- A comprehensive review of random survival forests (RSFs) is provided in Bou-Hamad (2011) (Bou-Hamad, Larocque, and Ben-Ameur 2011), which includes extensions to time-varying covariates and different censoring types.
- The discussion of decision trees omitted many methods for growing and pruning trees, if you are interest in those technical details see L. Breiman et al. (1984).
- RSFs have been shown to perform well in benchmark experiments on high-dimensional data, see Herrmann et al. (2021) and Spooner et al. (2020) for examples.
- This chapter considered the most ‘traditional’ forms of RSFs. Conditional inference forests are popular in the regression setting and whilst they are under-researched in survival, see Hothorn et al. (2005) for literature on the topic. A more recent method that seems to perform well is the (accelerated) oblique random survival

forest discussed in (**Jaeger2024?**).



# 14

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## *Support Vector Machines*

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This chapter introduces support vector machines (SVMs) for regression and then describes the extensions to survival analysis. Regression SVMs extend simple linear methods by estimating flexible, non-linear hyperplanes that minimise the difference between predictions and the truth for individual observations. In survival analysis, SVMs may make survival time or ranking predictions, however there is no current formulation for survival distribution predictions. The chapter begins by discussing survival time SVMs and then ranking models before concluding with a hybrid formulation that combines both model forms. This primarily covers the work of Shivaswamy and Van Belle. SVMs are a powerful method for estimating non-linear relationships in data and have proven to be well-performing models in regression and classification. However, SVMs are less developed in survival analysis and have been shown to perform worse than other models in experiments.

 Minor changes expected!

**This page is a work in progress and minor changes will be made over time.**

Support vector machines are a popular class of models in regression and classification settings due to their ability to make accurate predictions for complex high-dimensional, non-linear data. Survival support vector machines (SSVMs) predict continuous responses that can be used as ranking predictions with some formulations that provide survival time interpretations. This chapter starts with SVMs in the regression setting before moving to adaptions for survival analysis.

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### 14.1 SVMs for Regression

In simple linear regression, the aim is to estimate the line  $y = \alpha + x\beta_1$  by estimating the  $\alpha, \beta_1$  coefficients. As the number of coefficients increases, the goal is to instead estimate the *hyperplane*, which divides the higher-dimensional space into two separate parts. To visualize a hyperplane, imagine looking at a room from a birds eye view that has a dividing wall cutting the room into two halves (Figure 14.1). In this view, the room appears to have two dimensions ( $x$ =left-right,  $y$ =top-bottom) and the divider is a simple line of the form  $y = \alpha + x\beta_1$ . In reality, this room is actually three dimensional and has a third dimension ( $z$ =up-down) and the divider is therefore a hyperplane of the form  $y = \alpha + x\beta_1 + z\beta_2$ .

Continuing the linear regression example, consider a simple model where the objective is to find the  $\beta = (\beta_1 \ \beta_2 \cdots \beta_p)^\top$  coefficients that minimize  $\sum_{i=1}^n (g(\mathbf{x}_i) - y_i)^2$  where



Figure 14.1: Visualising a hyperplane by viewing a 3D room in two-dimensions with a wall that is now seen as a simple line. When standing in this room, the wall will clearly exist in three dimensional space.

$g(\mathbf{x}_i) = \alpha + \mathbf{x}_i^\top \boldsymbol{\beta}$  and  $(\mathbf{X}, \mathbf{y})$  is training data such that  $\mathbf{X} \in \mathbb{R}^{n \times p}$  and  $\mathbf{y} \in \mathbb{R}^n$ . In a higher-dimensional space, a penalty term can be added for variable selection to reduce model complexity, commonly of the form

$$\frac{1}{2} \sum_{i=1}^n (g(\mathbf{x}_i) - y_i)^2 + \frac{\lambda}{2} \|\boldsymbol{\beta}\|^2$$

for some penalty term  $\lambda \in \mathbb{R}$ . Minimizing this error function effectively minimizes the *average* difference between all predictions and true outcomes, resulting in a hyperplane that represents the best *linear* relationship between coefficients and outcomes.

Similarly to linear regression, support vector machines (SVMs) (Cortes and Vapnik 1995) also fit a hyperplane,  $g$ , on given training data,  $\mathbf{X}$ . However, in SVMs, the goal is to fit a *flexible* (non-linear) hyperplane that minimizes the difference between predictions and the truth for *individual* observations. A core feature of SVMs is that one does not try to fit a hyperplane that makes perfect predictions as this would overfit the training data and is unlikely to perform well on unseen data. Instead, SVMs use a regularized error function, which allows incorrect predictions (errors) for some observations, with the magnitude of error controlled by an  $\epsilon > 0$  parameter as well as slack parameters,  $\xi' = (\xi'_1 \ \xi'_2 \cdots \xi'_n)^\top$  and  $\xi^* = (\xi^*_1 \ \xi^*_2 \cdots \xi^*_n)^\top$ :

$$\begin{aligned} & \min_{\boldsymbol{\beta}, \alpha, \xi', \xi^*} \frac{1}{2} \|\boldsymbol{\beta}\|^2 + C \sum_{i=1}^n (\xi'_i + \xi^*_i) \\ & \text{subject to } \begin{cases} g(\mathbf{x}_i) \geq y_i - \epsilon - \xi'_i \\ g(\mathbf{x}_i) \leq y_i + \epsilon + \xi^*_i \\ \xi'_i, \xi^*_i \geq 0 \end{cases} \end{aligned} \quad (14.1)$$

$\forall i \in 1, \dots, n$  where  $g(\mathbf{x}_i) = \alpha + \mathbf{x}_i^\top \boldsymbol{\beta}$  for model weights  $\boldsymbol{\beta} \in \mathbb{R}^p$  and  $\alpha \in \mathbb{R}$  and the same training data  $(\mathbf{X}, \mathbf{y})$  as above.

Figure 14.2 visualizes a support vector regression model in two dimensions. The red circles are values within the  $\epsilon$ -tube and are thus considered to have a negligible error. In fact, the red circles do not affect the fitting of the optimal line  $g$  and even if they moved around, as long as they remain within the tube, the shape of  $g$  would not change. In contrast the blue diamonds have an unacceptable margin of error – as an example the top blue diamond will have  $\xi'_i = 0$  but  $\xi^*_i > 0$ , thus influencing the estimation of  $g$ . Points on or outside the epsilon tube are referred to as *support vectors* as they affect the construction of the hyperplane. The  $C \in \mathbb{R}_{>0}$  hyperparameter controls the slack parameters and thus as  $C$  increases, the number of errors (and subsequently support vectors) is allowed to increase resulting in low variance but higher bias, in contrast a lower  $C$  is more likely to introduce overfitting with low bias but high variance (Hastie, Tibshirani, and Friedman 2001).  $C$  should be tuned to control this trade-off.

The other core feature of SVMs is exploiting the *kernel trick*, which uses functions known as *kernels* to allow the model to learn a non-linear hyperplane whilst keeping the computations limited to lower-dimensional settings. Once the model coefficients have been estimated using the optimization above, predictions for a new observation  $\mathbf{x}^*$  can be made using a function of the form

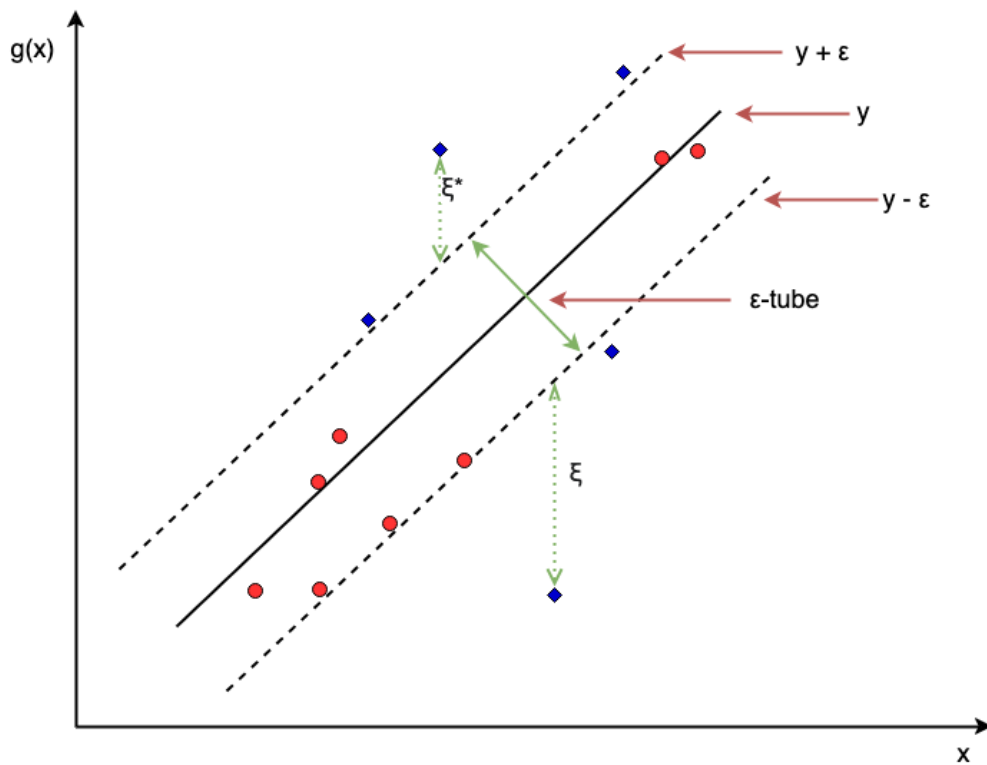


Figure 14.2: Visualising a support vector machine with an  $\varepsilon$ -tube and slack parameters  $\xi'$  and  $\xi^*$ . Red circles are values within the  $\varepsilon$ -tube and blue diamonds are support vectors on and outside the tube. x-axis is single covariate,  $x$ , and y-axis is  $g(x) = x\beta_1 + \alpha$ .

$$\hat{g}(\mathbf{x}^*) = \sum_{i=1}^n \mu_i K(\mathbf{x}^*, \mathbf{x}_i) + \alpha \quad (14.2)$$

Details (including estimation) of the  $\mu_i$  Lagrange multipliers are beyond the scope of this book, references are given at the end of this chapter for the interested reader.  $K$  is a kernel function, with common functions including the linear kernel,  $K(x^*, x_i) = \sum_{j=1}^p x_{ij}x_j^*$ , radial kernel,  $K(x^*, x_i) = \exp(-\omega \sum_{j=1}^p (x_{ij} - x_j^*)^2)$  for some  $\omega \in \mathbb{R}_{>0}$ , and polynomial kernel,  $K(x^*, x_i) = (1 + \sum_{j=1}^p x_{ij}x_j^*)^d$  for some  $d \in \mathbb{N}_{>0}$ .

The choice of kernel and its parameters, the regularization parameter  $C$ , and the acceptable error  $\epsilon$ , are all tunable hyper-parameters, which makes the support vector machine a highly adaptable and often well-performing machine learning method. The parameters  $C$  and  $\epsilon$  often have no clear apriori meaning (especially true in the survival setting predicting abstract rankings) and thus require tuning over a great range of values; no tuning usually results in a poor model fit (Probst, Boulesteix, and Bischl 2019).

## 14.2 SVMs for Survival Analysis

Extending SVMs to the survival domain (SSVMs) is a case of: i) identifying the quantity to predict; and ii) updating the optimization problem (Equation 14.1) and prediction function (Equation 14.2) to accommodate for censoring. In the first case, SSVMs can be used to either make survival time or ranking predictions, which are discussed in turn. The notation above is reused below for SSVMs, with additional notation introduced when required and now using the survival training data  $(\mathbf{X}, \mathbf{t}, \boldsymbol{\delta})$ .

### 14.2.1 Survival time SSVMs

To begin, consider the objective for support vector regression with the  $y$  variable replaced with the usual survival time outcome  $t$ , for all  $i \in 1, \dots, n$ :

$$\begin{aligned} & \min_{\beta, \alpha, \xi', \xi^*} \frac{1}{2} \|\beta\|^2 + C \sum_{i=1}^n (\xi'_i + \xi^*_i) \\ & \text{subject to } \begin{cases} g(\mathbf{x}_i) \geq t_i - \epsilon - \xi'_i \\ g(\mathbf{x}_i) \leq t_i + \epsilon + \xi^*_i \\ \xi'_i, \xi^*_i \geq 0 \end{cases} \end{aligned} \quad (14.3)$$

In survival analysis, this translates to fitting a hyperplane in order to predict the true survival time. However, as with all adaptations from regression to survival analysis, there needs to be a method for incorporating censoring.

Recall the  $(t_l, t_u)$  notation to describe censoring as introduced in Chapter 3 such that the outcome occurs within the range  $(t_l, t_u)$ . Let  $\tau \in \mathbb{R}_{>0}$  be some known time-point, then an observation is:

- left-censored if the survival time is less than  $\tau$ :  $(t_l, t_u) = (-\infty, \tau)$ ;
- right-censored if the true survival time is greater than  $\tau$ :  $(t_l, t_u) = (\tau, \infty)$ ; or

- uncensored if the true survival time is known to be  $\tau$ :  $(t_l, t_u) = (\tau, \tau)$ .

Define  $\mathcal{L} = \{i : t_i > -\infty\}$  as the set of observations with a finite lower-bounded time, which can be seen above to be those that are right-censored or uncensored. Define  $\mathcal{U} = \{i : t_i < \infty\}$  as the analogous set of observations with a finite upper-bounded time, which are those that are left-censored or uncensored.

Consider these definitions in the context of the constraints in Equation 14.3. The first constraint ensures the hyperplane is greater than some lower-bound created by subtracting the slack parameter from the true outcome – given the set definitions above this constraint only has meaning for observations with a finite lower-bound,  $i \in \mathcal{L}$ , otherwise the constraint would include  $g(\mathbf{x}_i) \geq -\infty$ , which is clearly not useful. Similarly the second constraint ensures the hyperplane is less than some upper-bound, which again can only be meaningful for observations  $i \in \mathcal{U}$ . Restricting the constraints in this way leads to the optimization problem (Shivaswamy, Chu, and Jansche 2007) below and visualised in Figure 14.3:

$$\begin{aligned} & \min_{\beta, \alpha, \xi', \xi^*} \frac{1}{2} \|\beta\|^2 + C \left( \sum_{i \in \mathcal{U}} \xi_i + \sum_{i \in \mathcal{L}} \xi_i^* \right) \\ \text{subject to } & \begin{cases} g(\mathbf{x}_i) & \geq t_i - \xi'_i, i \in \mathcal{L} \\ g(\mathbf{x}_i) & \leq t_i + \xi_i^*, i \in \mathcal{U} \\ \xi'_i \geq 0, \forall i \in \mathcal{L}; \xi_i^* \geq 0, \forall i \in \mathcal{U} \end{cases} \end{aligned}$$

If no observations are censored then the optimization becomes the regression optimization in (Equation 14.1). Note that in SSVMs, the  $\epsilon$  parameters are typically removed to better accommodate censoring and to help prevent the same penalization of over- and under-predictions. In contrast to this formulation, one *could* introduce more  $\epsilon$  and  $C$  parameters to separate between under- and over-predictions and to separate right- and left-censoring, however this leads to eight tunable hyperparameters, which is inefficient and may increase overfitting (Fouodo et al. 2018; Land et al. 2011). The algorithm can be simplified to right-censoring only by removing the second constraint completely for anyone censored:

$$\begin{aligned} & \min_{\beta, \alpha, \xi', \xi^*} \frac{1}{2} \|\beta\|^2 + C \sum_{i=1}^n (\xi'_i + \xi_i^*) \\ \text{subject to } & \begin{cases} g(\mathbf{x}_i) & \geq t_i - \xi_i^* \\ g(\mathbf{x}_i) & \leq t_i + \xi'_i, i : \delta_i = 1 \\ \xi'_i, \xi_i^* & \geq 0 \end{cases} \end{aligned}$$

$\forall i \in 1, \dots, n$ . With the prediction for a new observation  $\mathbf{x}^*$  calculated as,

$$\hat{g}(\mathbf{x}^*) = \sum_{i=1}^n \mu_i^* K(\mathbf{x}_i, \mathbf{x}^*) - \delta_i \mu_i' K(\mathbf{x}_i, \mathbf{x}^*) + \alpha$$

Where again  $K$  is a kernel function and the calculation of the Lagrange multipliers is beyond the scope of this book.

### 14.2.2 Ranking SSVMs

Support vector machines can be used to estimate rankings by penalizing predictions that result in disconcordant predictions. Recall the definition of concordance from Chapter 7:

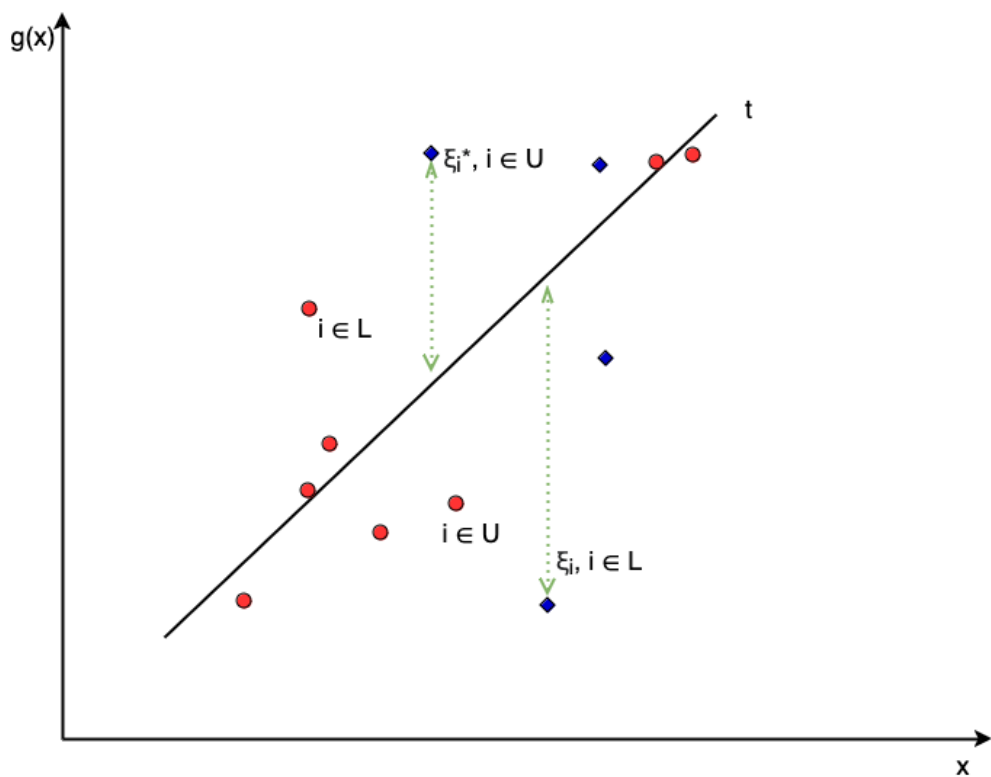


Figure 14.3: Visualising a survival time SVM. Blue diamonds are influential support vectors, which are uncensored or left-censored when  $g(\mathbf{x}) < t$  or uncensored or right-censored when  $g(\mathbf{x}) > t$ . Red circles are non-influential observations.

ranking predictions for a pair of comparable observations  $(i, j)$  where  $t_i < t_j \cap \delta_i = 1$ , are called concordant if  $r_i > r_j$  where  $r_i, r_j$  are the predicted ranks for observations  $i$  and  $j$  respectively and a higher value implies greater risk. Using the prognostic index as a ranking prediction (Section 5.4), a pair of observations is concordant if  $g(\mathbf{x}_i) > g(\mathbf{x}_j)$  when  $t_i < t_j$ , leading to:

$$\begin{aligned} & \min_{\beta, \alpha, \xi} \frac{1}{2} \|\beta\|^2 + \gamma \sum_{i=1}^n \xi_i \\ & \text{subject to } \begin{cases} g(\mathbf{x}_i) - g(\mathbf{x}_j) \geq \xi_i, \forall i, j \in CP \\ \xi_i \geq 0, i = 1, \dots, n \end{cases} \end{aligned}$$

where  $CP$  is the set of comparable pairs defined by  $CP = \{(i, j) : t_i < t_j \wedge \delta_i = 1\}$ . Given the number of pairs, the optimization problem quickly becomes difficult to solve with a very long runtime. To solve this problem Van Belle et al. (2011) found an efficient reduction that sorts observations in order of outcome time and then compares each data point  $i$  with the observation that has the next smallest *survival* time, skipping over censored observations, in maths:  $j(i) := \arg \max_{j \in 1, \dots, n} \{t_j : t_j < t_i\}$ . This is visualized in Figure 14.4 where six observations are sorted by outcome time from smallest (left) to largest (right). Starting from right to left, the first pair is made by matching the observation to the first uncensored outcome to the left, this continues to the end. In order for all observations to be used in the optimisation, the algorithm sets the first outcome to be uncensored hence observation 2 being compared to observation 1.

Using this reduction, the algorithm becomes

$$\begin{aligned} & \min_{\beta, \alpha, \xi} \frac{1}{2} \|\beta\|^2 + \gamma \sum_{i=1}^n \xi_i \\ & \text{subject to } \begin{cases} g(\mathbf{x}_i) - g(\mathbf{x}_{j(i)}) \geq t_i - t_{j(i)} - \xi_i \\ \xi_i \geq 0 \end{cases} \end{aligned}$$

$\forall i = 1, \dots, n$ . Note the updated right hand side of the constraint, which plays a similar role to the  $\epsilon$  parameter by allowing ‘mistakes’ in predictions without penalty.

Predictions for a new observation  $\mathbf{x}^*$  are calculated as,

$$\hat{g}(\mathbf{x}^*) = \sum_{i=1}^n \mu_i (K(\mathbf{x}_i, \mathbf{x}^*) - K(\mathbf{x}_{j(i)}, \mathbf{x}^*)) + \alpha$$

Where  $\mu_i$  are again Lagrange multipliers.

### 14.2.3 Hybrid SSVMs

Finally, Van Belle et al. (2011) noted that the ranking algorithm could be updated to add the constraints of the regression model, thus providing a model that simultaneously optimizes for ranking whilst providing continuous values that can be interpreted as survival time predictions. This results in the hybrid SSVM with constraints  $\forall i = 1, \dots, n$ :

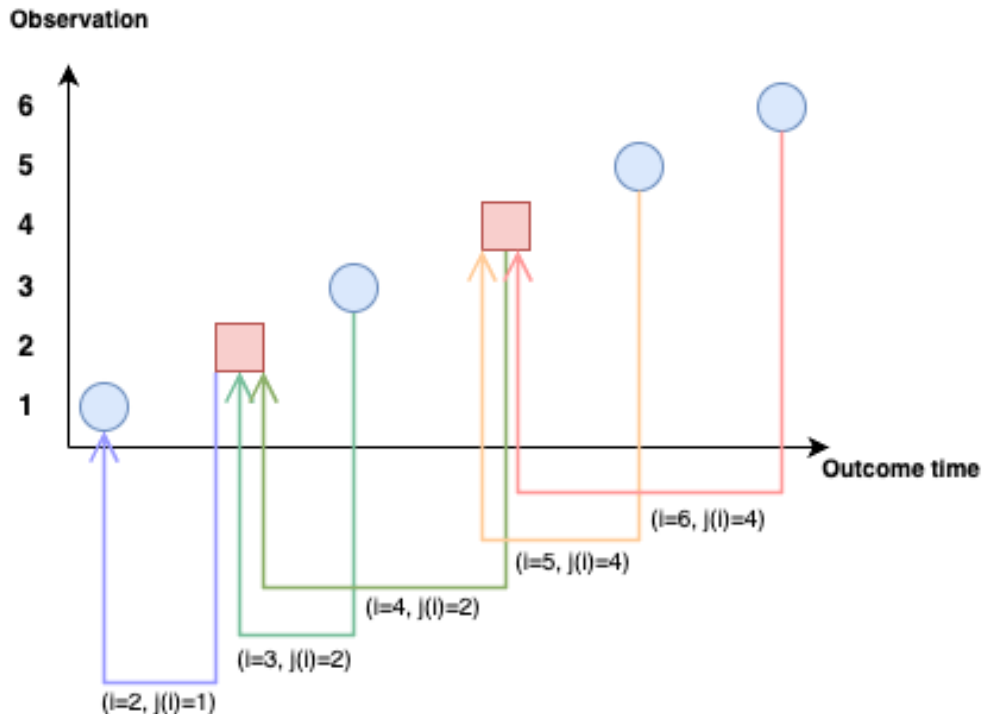


Figure 14.4: Van Belle SVM nearest neighbors reduction. Sorted observations are paired with the nearest uncensored outcome ‘to the left’. Red squares are uncensored observations and blue circles are censored. The observation with the smallest outcome time is always treated as uncensored.

$$\begin{aligned} & \min_{\beta, \alpha, \xi, \xi', \xi^*} \frac{1}{2} \|\beta\|^2 + \gamma \sum_{i=1}^n \xi_i + C \sum_{i=1}^n (\xi'_i + \xi_i^*) \\ \text{subject to } & \begin{cases} g(\mathbf{x}_i) - g(\mathbf{x}_{j(i)}) & \geq t_i - t_{j(i)} - \xi_i \\ g(\mathbf{x}_i) & \leq t_i + \xi_i^*, i : \delta_i = 1 \\ g(\mathbf{x}_i) & \geq t_i - \xi'_i \\ \xi_i, \xi'_i, \xi_i^* & \geq 0 \end{cases} \end{aligned}$$

The blue parts of the equation make up the ranking model and the red parts are the regression model.  $\gamma$  is the penalty associated with the regression method and  $C$  is the penalty associated with the ranking method. Setting  $\gamma = 0$  results in the regression SVM and  $C = 0$  results in the ranking SSVM. Hence, fitting the hybrid model and tuning these parameters is an efficient way to automatically detect which SSVM is best suited to a given task.

Once the model is fit, a prediction from given features  $\mathbf{x}^* \in \mathbb{R}^p$ , can be made using the equation below, again with the ranking and regression contributions highlighted in blue and red respectively.

$$\hat{g}(\mathbf{x}^*) = \sum_{i=1}^n \mu_i (K(\mathbf{x}_i, \mathbf{x}^*) - K(\mathbf{x}_{j(i)}, \mathbf{x}^*)) + \mu_i^* K(\mathbf{x}_i, \mathbf{x}^*) - \delta_i \mu_i' K(\mathbf{x}_i, \mathbf{x}^*) + \alpha$$

where  $\mu_i, \mu_i^*, \mu_i'$  are Lagrange multipliers and  $K$  is a chosen kernel function, which may have further hyper-parameters to select or tune.

### 14.3 Conclusion

#### Key takeaways

- Support vector machines (SVMs) are a highly flexible machine learning method that can use the ‘kernel trick’ to represent infinite dimensional spaces in finite domains;
- Survival SVMs (SSVMs) extend regression SVMs by either making survival time predictions, ranking predictions, or a combination of the two;
- The hybrid SSVM provides an efficient method that encapsulates all the elements of regression and ranking SSVMs and is therefore a good model to include in benchmark experiments to test the potential of SSVMs.

#### Limitations

- SSVMs can only perform well with extensive tuning of hyper-parameters over a wide parameter space. To-date, no papers have experimented with the tuning range for the  $\gamma$  and  $C$  parameters, we note (Fouodo et al. 2018) tune over  $(2^{-5}, 2^5)$ .
- Even using the regression or hybrid model, the authors’ experiments with the SSVM have consistently shown ‘survival time’ estimates tend to be unrealistically large.
- Due to the above limitation, regression estimates cannot be meaningful interpreted and as a consequence there is no sensible composition to create a distribution

prediction from an SSVM. Hence, we are hesitant to suggest usage of SSVMs outside of ranking-based problems.

#### Further reading

- Shivaswamy, Chu, and Jansche (2007), Khan and Bayer Zubek (2008), Land et al. (2011), and Van Belle et al. (2011) to learn more about regression SSVMs.
- Evers and Messow (2008), Van Belle et al. (2007), Van Belle et al. (2008), and Van Belle et al. (2011) for more information about ranking SSVMs.
- Goli, Mahjub, Faradmal, and Soltanian (2016) and Goli, Mahjub, Faradmal, Mashayekhi, et al. (2016) introduce mean residual lifetime optimization SSVMs.
- Fouodo et al. (2018) surveys and benchmarks SSVMs.



# 15

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## *Boosting Methods*

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This page is a work in progress and major changes will be made over time.

Boosting is a machine learning strategy that can be applied to any model class. Similarly to random forests, boosting is an ensemble method that creates a model from a ‘committee’ of learners. The committee is formed of *weak* learners that make poor predictions individually, which creates a *slow learning* approach (as opposed to ‘greedy’) that requires many iterations for a model to be a good fit to the data. Boosting models are similar to random forests in that both make predictions from a large committee of learners. However the two differ in how the members of the committee are correlated and in how they are combined to make a prediction. In random forests, each decision tree is grown independently and their predictions are combined by a simple mean calculation. In contrast, weak learners in a boosting model are fit sequentially with errors from one learner used to train the next, predictions are then made by a linear combination of predictions from each learner (Figure 15.1).

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### 15.1 GBMs for Regression

One of the earliest boosting algorithms is AdaBoost (Freund and Schapire 1996), which is more generally a Forward Stagewise Additive Model (FSAM) with an exponential loss (Hastie, Tibshirani, and Friedman 2001). Today, the most widely used boosting model is the Gradient Boosting Machine (GBM) (J. H. Friedman 2001) or extensions thereof.

Figure 15.1 illustrates the process of training a GBM in a least-squares regression setting:

1. A weak learner,  $f_1$ , often a decision tree of shallow depth is fit on the training data  $(\mathbf{X}, \mathbf{y})$ .
2. Predictions from the learner,  $f_1(\mathbf{X})$ , are compared to the ground truth,  $\mathbf{y}$ , and the residuals are calculated as  $\mathbf{r}_1 = f_1(\mathbf{X}) - \mathbf{y}$ .
3. The next weak learner,  $f_2$ , uses the previous residuals for the target prediction,  $(\mathbf{X}, \mathbf{r}_1)$
4. This is repeated to train  $M$  learners,  $f_1, \dots, f_M$

Predictions are then made as  $\hat{\mathbf{y}} = f_1(\mathbf{X}) + f_2(\mathbf{X}) + \dots + f_M(\mathbf{X})$ .

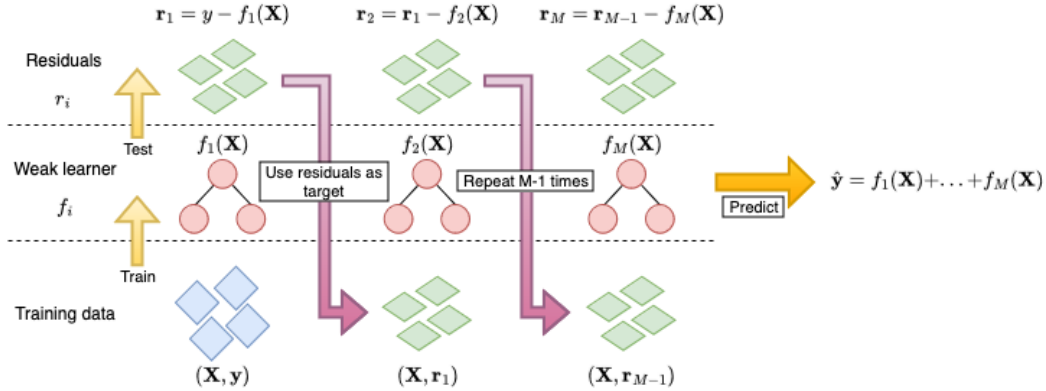


Figure 15.1: Least squares regression Boosting algorithm where the gradient is calculated as the difference between ground truth and predictions.

This is a simplification of the general gradient boosting algorithm, where the residuals are used to train the next model. More generally, a suitable, differentiable loss function relating to the problem of interest is chosen and the negative gradient is computed by comparing the predictions in each iteration with the ground truth. Residuals can be used in the regression case as these are proportional to the negative gradient of the mean squared error.

The algorithm above is also a simplification as no hyper-parameters other than  $M$  were included for controlling the algorithm. In order to reduce overfitting, three common hyper-parameters are utilised:

**Number of iterations,  $M$ :** The number of iterations is often claimed to be the most important hyper-parameter in GBMs and it has been demonstrated that as the number of iterations increases, so too does the model performance (with respect to a given loss on test data) up to a certain point of overfitting (Buhlmann 2006; Hastie, Tibshirani, and Friedman 2001; Schmid and Hothorn 2008a). This makes sense as the foundation of boosting rests on the idea that weak learners can slowly be combined to form a single powerful model. Finding the optimal value of  $M$  is critical as a value too small will result in poor predictions, whilst a value too large will result in model overfitting.

**Subsampling proportion,  $\phi$ :** Sampling a fraction,  $\phi$ , of the training data at each iteration can improve performance and reduce runtime (Hastie, Tibshirani, and Friedman 2001), with  $\phi = 0.5$  often used. Motivated by the success of bagging in random forests, stochastic gradient boosting (J. Friedman 1999) randomly samples the data in each iteration. It appears that subsampling performs best when also combined with shrinkage (Hastie, Tibshirani, and Friedman 2001) and as with the other hyper-parameters, selection of  $\phi$  is usually performed by nested cross-validation.

**Step-size,  $\nu$ :** The step-size parameter is a shrinkage parameter that controls the contribution of each weak learner at each iteration. Several studies have demonstrated that GBMs perform better when shrinkage is applied and a value of  $\nu = 0.1$  is often suggested (Buhlmann and Hothorn 2007; Hastie, Tibshirani, and Friedman 2001; J. H. Friedman 2001; D. K. K. Lee, Chen, and Ishwaran 2019; Schmid and Hothorn 2008a). The optimal values of  $\nu$  and  $M$  depend on each other, such that smaller values of  $\nu$  require larger values of  $M$ , and vice versa. This is intuitive as smaller  $\nu$  results in a slower learning algorithm and therefore more iterations are required to fit the model. Accurately selecting the  $M$  parameter is generally

considered to be of more importance, and therefore a value of  $\nu$  is often chosen heuristically (e.g. the common value of 0.1) and then  $M$  is tuned by cross-validation and/or early-stopping, which is the process of monitoring the model's training performance and stopping when a set performance is reached or when performance stagnates (i.e., no improvement over a set number of rounds).

As well as these parameters, the underlying weak learner hyper-parameters are also commonly tuned. If using a decision tree, then it is usual to restrict the number of terminal nodes in the tree to be between 4 and 8, which corresponds to two or three splits in the tree. Including these hyper-parameters, the general gradient boosting machine algorithm is as follows:

1.  $g_0 \leftarrow$  Initial guess
2. **For**  $m = 1, \dots, M$ :
3.      $\mathcal{D}_{train}^* \leftarrow$  Randomly sample  $\mathcal{D}_{train}$  with probability  $\phi$
4.      $r_{im} \leftarrow -[\frac{\partial L(y_i, g_{m-1}(X_i))}{\partial g_{m-1}(X_i)}], \forall i \in \{i : X_i \in \mathcal{D}_{train}^*\}$
5.     Fit a weak learner,  $h_m$ , to  $(\mathbf{X}, \mathbf{r}_m)$
6.      $g_m \leftarrow g_{m-1} + \nu h_m$
7. **end For**
8. **return**  $\hat{g} = g_M$

Note:

1. The initial guess,  $g_0$ , is often the mean of  $y$  for regression problems but can also simply be 0.
2. Line 4 is the calculation of the negative gradient, which is equivalent to calculating the residuals in a regression problem with the mean squared error loss.
3. Lines 5-6 differ between implementations, with some fitting multiple weak learners and selecting the one that minimizes a simple optimization problem. The version above is simplest to implement and quickest to run, whilst still providing good model performance.

Once the model is trained, predictions are made for new data,  $\mathbf{X}_{test}$  with

$$\hat{Y} = \hat{g}(\mathbf{X}_{test}) = g_0(\mathbf{X}_{test}) + \nu \sum_{i=1}^M g_i(\mathbf{X}_{test})$$

GBMs provide a flexible, modular algorithm, primarily comprised of a differentiable loss to minimise,  $L$ , and the selection of weak learners. This chapter focuses on tree-based weak learners, though other weak learners are possible. Perhaps the most common alternatives are linear least squares (J. H. Friedman 2001) and smoothing splines (Bühlmann and Yu 2003), we will not discuss these further here as decision trees are primarily used for survival analysis, due the flexibility demonstrated in Chapter 13. See references at the end of the chapter for other weak learners. Extension to survival analysis therefore follows by considering alternative losses.

## 15.2 GBMs for Survival Analysis

Unlike other machine learning algorithms that historically ignored survival analysis, early GBM papers considered boosting in a survival context (Ridgeway 1999); though there appears to be a decade gap before further considerations were made in the survival setting. After that period, developments, discussed in this chapter, by Binder, Schmid, and Hothorn, adapted GBMs to a framework suitable for survival analysis.

All survival GBMs make ranking predictions and none are able to directly predict survival distributions. However, depending on the underlying model, the predictions may be indirectly composed into a survival distribution, for example algorithms that assume a proportional hazards (PH) or accelerated failure time (AFT) form. This section starts with those models with simpler underlying forms, then explores more complex alternatives.

### 15.2.1 PH and AFT GBMs

The negative log-likelihood of the semi-parametric PH and fully-parametric AFT models can be derived from the (partial) likelihoods presented in `?@sec-surv-obj`. Given the likelihoods measure the goodness of fit of model parameters, algorithms that use these losses use boosting to train the model coefficients,  $\beta$ , hence at each iteration in the algorithm,  $g_m(\mathbf{x}_i) = \mathbf{x}_i\beta^{(m)}$ , where  $\beta^{(m)}$  are the updated coefficients in iteration  $m$ .

The Cox partial likelihood (Cox 1972, 1975) is given by

$$L^{PH}(\beta) = \prod_{i:\delta_i=1}^n \frac{\exp(\eta_i)}{\sum_{j \in \mathcal{R}_{t_i}} \exp(\eta_j)}$$

with corresponding negative log-likelihood

$$-l^{PH}(\beta) = - \sum_{i=1}^n \delta_i \left[ \eta_i - \log \left( \sum_{j \in \mathcal{R}_{t_i}} \exp(\eta_j) \right) \right] \quad (15.1)$$

where  $\mathcal{R}_{t_i}$  is the set of patients at risk at time  $t_i$  and  $\eta_i = \mathbf{x}_i\beta$ .

The gradient of  $-l^{PH}$  at iteration  $m$  is then

$$r_{im} := \delta_i - \sum_{j=1}^n \delta_j \frac{\mathbb{I}(t_i \geq t_j) \exp(g_{m-1}(\mathbf{x}_i))}{\sum_{k \in \mathcal{R}_{t_j}} \exp(g_{m-1}(\mathbf{x}_k))} \quad (15.2)$$

where  $g_{m-1}(\mathbf{x}_i) = \mathbf{x}_i\beta^{(m-1)}$ .

For non-PH data, boosting an AFT model can outperform boosted PH models (Schmid and Hothorn 2008b). The AFT is defined by

$$\log \mathbf{y} = \boldsymbol{\eta} + \sigma W$$

where  $W$  is a random noise variable independent of  $X$ , and  $\sigma$  is a scale parameter controlling the amount of noise; again  $\boldsymbol{\eta} = \mathbf{X}\beta$ . By assuming a distribution on  $W$ , a distribution is assumed for the full parametric model. The model is boosted by simultaneously estimating

$\sigma$  and  $\beta$ . Assuming a location-scale distribution with location  $g(\mathbf{x}_i)$  and scale  $\sigma$ , one can derive the negative log-likelihood in the  $m$ th iteration as (Klein and Moeschberger 2003)

$$\begin{aligned} -l_m^{AFT}(\beta) = & -\sum_{i=1}^n \delta_i \left[ -\log \sigma + \log f_W \left( \frac{\log(t_i) - \hat{g}_{m-1}(\mathbf{x}_i)}{\hat{\sigma}_{m-1}} \right) \right] + \\ & (1 - \delta_i) \left[ \log S_W \left( \frac{\log(t_i) - \hat{g}_{m-1}(\mathbf{x}_i)}{\hat{\sigma}_{m-1}} \right) \right] \end{aligned}$$

where  $\hat{g}_{m-1}, \hat{\sigma}_{m-1}$  are the location-scale parameters estimated in the previous iteration. Note this key difference to other GBM methods in which two estimates are made in each iteration step. After updating  $\hat{g}_m$ , the scale parameter,  $\hat{\sigma}_m$ , is updated as

$$\hat{\sigma}_m := \arg \min_{\sigma} -l_m^{AFT}(\beta)$$

$\sigma_0$  is commonly initialized as 1 (Schmid and Hothorn 2008b).

As well as boosting fully-parametric AFTs, one could also consider boosting semi-parametric AFTs, for example using the Gehan loss (Johnson and Long 2011) or using Buckley-James imputation (Z. Wang and Wang 2010). However, known problems with semi-parametric AFT models and the Buckley-James procedure (Wei 1992), as well as a lack of off-shelf implementation, mean that these methods are rarely used in practice.

### 15.2.2 Discrimination Boosting

Instead of optimising models based on a given model form, one could instead estimate  $\hat{\eta}$  by optimizing a concordance index, such as Uno's or Harrell's C (Y. Chen et al. 2013; Mayr and Schmid 2014). Consider Uno's C (Section 7.1):

$$C_U(\hat{g}, \mathcal{D}_{train}) = \frac{\sum_{i \neq j} \delta_i \{\hat{G}_{KM}(t_i)\}^{-2} \mathbb{I}(t_i < t_j) \mathbb{I}(\hat{g}(\mathbf{x}_i) > \hat{g}(\mathbf{x}_j))}{\sum_{i \neq j} \delta_i \{\hat{G}_{KM}(t_i)\}^{-2} \mathbb{I}(t_i < t_j)}$$

The GBM algorithm requires that the chosen loss, here  $C_U$ , be differentiable with respect to  $\hat{g}(X)$ , which is not the case here due to the indicator term,  $\mathbb{I}(\hat{g}(X_i) > \hat{g}(X_j))$ , however this term can be replaced with a sigmoid function to create a differentiable loss (Ma and Huang 2006)

$$K(u|\omega) = \frac{1}{1 + \exp(-u/\omega)}$$

where  $\omega$  is a tunable hyper-parameter controlling the smoothness of the approximation. The measure to optimise is then,

$$C_{USmooth}(\beta|\omega) = \sum_{i \neq j} \frac{k_{ij}}{1 + \exp[(\hat{g}(X_j) - \hat{g}(X_i))/\omega]} \quad (15.3)$$

with

$$k_{ij} = \frac{\Delta_i (\hat{G}_{KM}(T_i))^{-2} \mathbb{I}(T_i < T_j)}{\sum_{i \neq j} \Delta_i (\hat{G}_{KM}(T_i))^{-2} \mathbb{I}(T_i < T_j)}$$

The negative gradient at iteration  $m$  for observation  $i$  is then calculated as,

$$r_{im} := - \sum_{j=1}^n k_{ij} \frac{-\exp(\frac{\hat{g}_{m-1}(\mathbf{x}_j) - \hat{g}_{m-1}(\mathbf{x}_i)}{\omega})}{\omega(1 + \exp(\frac{\hat{g}_{m-1}(\mathbf{x}_j) - \hat{g}_{m-1}(\mathbf{x}_i)}{\omega}))} \quad (15.4)$$

The GBM algorithm is then followed as normal with the above loss and gradient. This algorithm may be more insensitive to overfitting than others (Mayr, Hofner, and Schmid 2016), however stability selection (Meinshausen and Bühlmann 2010), which is implemented in off-shelf software packages (Hothorn et al. 2020), can be considered for variable selection.

### 15.2.3 CoxBoost

Finally, ‘CoxBoost’ is an alternative method to boost Cox models and has been demonstrated to perform well in experiments. This algorithm boosts the Cox PH by optimising the penalized partial-log likelihood; additionally the algorithm allows for mandatory (or ‘forced’) covariates (Binder and Schumacher 2008). In medical domains the inclusion of mandatory covariates may be essential, either for model interpretability, or due to prior expert knowledge. CoxBoost deviates from the algorithm presented above by instead using an offset-based approach for generalized linear models (Tutz and Binder 2007).

Let  $\mathcal{I} = \{1, \dots, p\}$  be the indices of the covariates, let  $\mathcal{I}_{mand}$  be the indices of the mandatory covariates that must be included in all iterations, and let  $\mathcal{I}_{opt} = \mathcal{I} \setminus \mathcal{I}_{mand}$  be the indices of the optional covariates that may be included in any iteration. In the  $m$ th iteration, the algorithm fits a weak learner on all mandatory covariates and *one* optional covariate:

$$\mathcal{I}_m = \mathcal{I}_{mand} \cup \{x | x \in \mathcal{I}_{opt}\}$$

In addition, a penalty matrix  $\mathbf{P} \in \mathbb{R}^{p \times p}$  is considered such that  $P_{ii} > 0$  implies that covariate  $i$  is penalized and  $P_{ii} = 0$  means no penalization. In practice, this is usually a diagonal matrix (Binder and Schumacher 2008) and by setting  $P_{ii} = 0, i \in \mathcal{I}_{mand}$  and  $P_{ii} > 0, i \notin \mathcal{I}_{mand}$ , only optional (non-mandatory) covariates are penalized. The penalty matrix can be allowed to vary with each iteration, which allows for a highly flexible approach, however in implementation a simpler approach is to either select a single penalty to be applied in each iteration step or to have a single penalty matrix (Binder 2013).

At the  $m$ th iteration and the  $k$ th set of indices to consider ( $k = 1, \dots, p$ ), the loss to optimize is the penalized partial-log likelihood given by

$$l_{pen}(\gamma_{mk}) = \sum_{i=1}^n \delta_i \left[ \eta_{i,m-1} + \mathbf{x}_{i,\mathcal{I}_{mk}} \gamma_{mk}^\top \right] - \delta_i \log \left( \sum_{j=1}^n \mathbb{I}(t_j \leq t_i) \exp(\eta_{i,m-1} + \mathbf{x}_{i,\mathcal{I}_{mk}} \gamma_{mk}^\top) \right) - \lambda \gamma_{mk} \mathbf{P}_{mk} \gamma_{mk}^\top$$

where  $\eta_{i,m} = \mathbf{x}_i \beta_m$ ,  $\gamma_{mk}$  are the coefficients corresponding to the covariates in  $\mathcal{I}_{mk}$  which is the possible set of candidates for a subset of total candidates  $k = 1, \dots, p$ ;  $\mathbf{P}_{mk}$  is the penalty matrix; and  $\lambda$  is a penalty hyper-parameter to be tuned or selected.<sup>1</sup>

<sup>1</sup>On notation, note that  $\mathbf{P}_{ij}$  refers to the penalty matrix in the  $i$ th iteration for the  $j$ th set of indices, whereas  $P_{ij}$  is the  $(i, j)$ th element in the matrix  $\mathbf{P}$ .

In each iteration, all potential candidate sets (the union of mandatory covariates and one other covariate) are updated by

$$\hat{\gamma}_{mk} = \mathbf{I}_{pen}^{-1}(\hat{\gamma}_{(m-1)k})U(\hat{\gamma}_{(m-1)k})$$

where  $U(\gamma) = \partial l / \partial \gamma(\gamma)$  and  $\mathbf{I}_{pen}^{-1} = \partial^2 l / \partial \gamma \partial \gamma^T (\gamma + \lambda \mathbf{P}_{(m-1)k})$  are the first and second derivatives of the unpenalized partial-log-likelihood. The optimal set is then found as

$$k^* := \arg \max_k l_{pen}(\hat{\gamma}_{mk})$$

and the estimated coefficients are updated with

$$\hat{\beta}_m = \hat{\beta}_{m-1} + \hat{\gamma}_{mk^*}, \quad k^* \in \mathcal{I}_{mk}$$

This deviates from the standard GBM algorithm by directly optimizing  $l_{pen}$  and not its gradient, additionally model coefficients are iteratively updated instead of a more general model form.

### 15.3 Conclusion

#### Key takeaways

- GBMs are a highly flexible and powerful machine learning tool. They have proven particularly useful in survival analysis as minimal adjustments are required to make use of off-shelf software.
- The flexibility of the algorithm allows all the models above to be implemented in relatively few open-source packages.
- There is evidence that boosting models can outperform the Cox PH even in low-dimensional settings (Schmid and Hothorn 2008b), which is not something all ML models can claim.

#### Limitations

- Boosting, especially with tree learners, is viewed as a black-box model that is increasingly difficult to interpret as the number of iterations increase. However, there are several methods for increasing interpretability, such as variable importance and SHAPs (Lundberg and Lee 2017).
- Boosting often relies on intensive computing power, however, dedicated packages such as **xgboost** (T. Chen et al. 2020), exist to push CPU/GPUs to their limits in order to optimise predictive performance.

#### Further reading

- Bühlmann and Yu (2003); Hothorn et al. (2020); Z. Wang and Wang (2010) for more general information and background on componentwise GBMs
- J. H. Friedman (2001); Z. Wang and Wang (2010) for linear least squares weak learners

- Bühlmann and Yu (2003); J. H. Friedman (2001) for decision tree weak learners
- Ridgeway (1999) for early research into GBMs for survival analysis
- Johnson and Long (2011) and Z. Wang and Wang (2010) for semi-parametric AFT boosting

# 16

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## Neural Networks

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### 16.1 Neural Networks

Before starting the survey on neural networks, first a comment about their transparency and accessibility. Neural networks are infamously difficult to interpret and train, with some calling building and training neural networks an ‘art’ (Hastie, Tibshirani, and Friedman 2001). As discussed in the introduction of this book, whilst neural networks are not transparent with respect to their predictions, they are transparent with respect to implementation. In fact the simplest form of neural network, as seen below, is no more complex than a simple linear model. With regard to accessibility, whilst it is true that defining a custom neural network architecture is complex and highly subjective, established models are implemented with a default architecture and are therefore accessible ‘off-shelf’.

#### 16.1.1 Neural Networks for Regression

(Artificial) Neural networks (ANNs) are a class of model that fall within the greater paradigm of *deep learning*. The simplest form of ANN, a feed-forward single-hidden-layer network, is a relatively simple algorithm that relies on linear models, basic activation functions, and simple derivatives. A short introduction to feed-forward regression ANNs is provided to motivate the survival models. This focuses on single-hidden-layer models and increasing this to multiple hidden layers follows relatively simply.

The single hidden-layer network is defined through three equations

$$\begin{aligned} Z_m &= \sigma(\alpha_{0m} + \alpha_m^T X_i), \quad m = 1, \dots, M \\ T &= \beta_{0k} + \beta_k^T Z, \quad k = 1, \dots, K \\ g_k(X_i) &= \phi_k(T) \end{aligned}$$

where  $(X_1, \dots, X_n) \stackrel{i.i.d.}{\sim} X$  are the usual training data,  $\alpha_{0m}, \beta_0$  are bias parameters, and

$\theta = \{\alpha_m, \beta\}$  ( $m = 1, \dots, M$ ) are model weights where  $M$  is the number of hidden units.  $K$  is the number of classes in the output, which for regression is usually  $K = 1$ . The function  $\phi$  is a ‘link’ or ‘activation function’, which transforms the predictions in order to provide an outcome of the correct return type; usually in regression,  $\phi(x) = x$ .  $\sigma$  is the ‘activation function’, which transforms outputs from each layer. The  $\alpha_m$  parameters are often referred to as ‘activations’. Different activation functions may be used in each layer or the same used throughout, the choice is down to expert knowledge. Common activation functions seen in this section include the sigmoid function,

$$\sigma(v) = (1 + \exp(-v))^{-1}$$

tanh function,

$$\sigma(v) = \frac{\exp(v) - \exp(-v)}{\exp(v) + \exp(-v)} \quad (16.1)$$

and ReLU (Nair and Hinton 2010)

$$\sigma(v) = \max(0, v) \quad (16.2)$$

A single-hidden-layer model can also be expressed in a single equation, which highlights the relative simplicity of what may appear a complex algorithm.

$$g_k(X_i) = \sigma_0(\beta_{k0} + \sum_{h=1}^H (\beta_{kh}\sigma_h(\beta_{h0} + \sum_{m=1}^M \beta_{hm}X_{i;m})) \quad (16.3)$$

where  $H$  are the number of hidden units,  $\beta$  are the model weights,  $\sigma_h$  is the activation function in unit  $h$ , also  $\sigma_0$  is the output unit activation, and  $X_{i;m}$  is the  $i$ th observation features in the  $m$ th hidden unit.

An example feed-forward single-hidden-layer regression ANN is displayed in (Figure 16.1). This model has 10 input units, 13 hidden units, and one output unit; two bias parameters are fit. The model is described as ‘feed-forward’ as there are no cycles in the node and information is passed forward from the input nodes (left) to the output node (right).

### Back-Propagation

The model weights,  $\theta$ , in this section are commonly fit by ‘back-propagation’ although this method is often considered inefficient compared to more recent advances. A brief pseudo-algorithm for the process is provided below.

Let  $L$  be a chosen loss function for model fitting, let  $\theta = (\alpha, \beta)$  be model weights, and let  $J \in \mathbb{N}_{>0}$  be the number of iterations to train the model over. Then the back-propagation method is given by,

- **For**  $j = 1, \dots, J$ : [] Forward Pass [i.] Fix weights  $\theta^{(j-1)}$ . [ii.] Compute predictions  $\hat{Y} := \hat{g}_k^{(j)}(X_i|\theta^{(j-1)})$  with (Equation 16.3). [] Backward Pass [iii.] Calculate the gradients of the loss  $L(\hat{Y}|\mathcal{D}_{train})$ . [] Update \*[iv.] Update  $\alpha^{(r)}, \beta^{(r)}$  with gradient descent.
- **End For**

In regression, a common choice for  $L$  is the squared loss,

$$L(\hat{g}, \theta|\mathcal{D}_{train}) = \sum_{i=1}^n (Y_i - \hat{g}(X_i|\theta))^2$$

which may help illustrate how the training outcome,  $(Y_1, \dots, Y_n) \stackrel{i.i.d.}{\sim} Y$ , is utilised for model fitting.

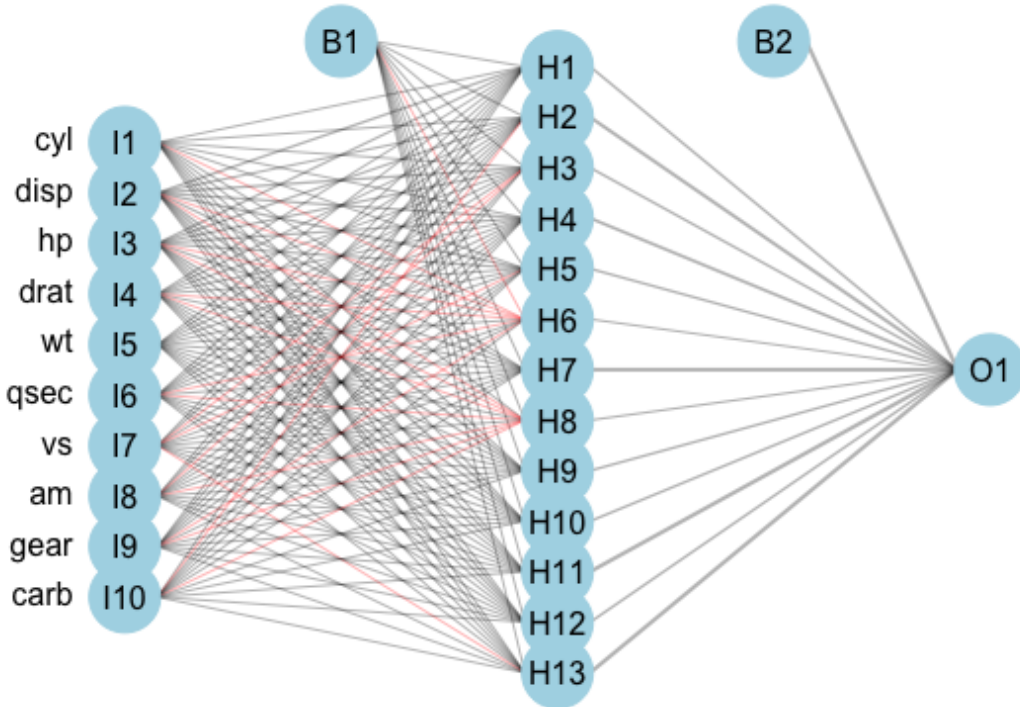


Figure 16.1: Single-hidden-layer artificial neural network with 13 hidden units fit on the `mtcars` (Henderson and Velleman 1981) dataset using the `nnet` (N. Venables and D. Ripley 2002) package, and `gamlss.add` (Stasinopoulos et al. 2020) for plotting. Left column are input variables, I1-I10, second column are 13 hidden units, H1-H13, right column is single output variable, O1. B1 and B2 are bias parameters.

## Making Predictions

Once the model is fitted, predictions for new data follow by passing the testing data as inputs to the model with fitted weights,

$$g_k(X^*) = \sigma_0(\hat{\beta}_{k0} + \sum_{h=1}^H (\hat{\beta}_{kh}\sigma_h(\hat{\beta}_{h0} + \sum_{m=1}^M \hat{\beta}_{hm}X_m^*)))$$

## Hyper-Parameters

In practice, a regularization parameter,  $\lambda$ , is usually added to the loss function in order to help avoid overfitting. This parameter has the effect of shrinking model weights towards zero and hence in the context of ANNs regularization is usually referred to as ‘weight decay’. The value of  $\lambda$  is one of three important hyper-parameters in all ANNs, the other two are: the range of values to simulate initial weights from, and the number of hidden units,  $M$ .

The range of values for initial weights is usually not tuned but instead a consistent range is specified and the neural network is trained multiple times to account for randomness in initialization.

The regularization parameter and number of hidden units,  $M$ , depend on each other and have a similar relationship to the learning rate and number of iterations in the GBMs ([?@sec-surv-ml-models-boost](#)). Like the GBMs, it is simplest to set a high number of hidden units and then tune the regularization parameter (Bishop 2006; Hastie, Tibshirani, and Friedman 2001). Determining how many hidden layers to include, and how to connect them, is informed by expert knowledge and well beyond the scope of this book; decades of research has been required to derive sensible new configurations.

## Training Batches

ANNs can either be trained using complete data, in batches, or online. This decision is usually data-driven and will affect the maximum number of iterations used to train the algorithm; as such this will also often be chosen by expert-knowledge and not empirical methods such as cross-validation.

## Neural Terminology

Neural network terminology often reflects the structures of the brain. Therefore ANN units are referred to as nodes or neurons and sometimes the connections between neurons are referred to as synapses. Neurons are said to be ‘fired’ if they are ‘activated’. The simplest example of activating a neuron is with the Heaviside activation function with a threshold of 0:  $\sigma(v) = \mathbb{I}(v \geq 0)$ . Then a node is activated and passes its output to the next layer if its value is positive, otherwise it contributes no value to the next layer.

### 16.1.2 Neural Networks for Survival Analysis

Surveying neural networks is a non-trivial task as there has been a long history in machine learning of publishing very specific data-driven neural networks with limited applications; this is also true in survival analysis. This does mean however that where limited developments for survival were made in other machine learning classes, ANN survival adaptations have been around for several decades. A review in 2000 by Schwarzer *et al.* surveyed 43 ANNs for diagnosis and prognosis published in the first half of the 90s, however only up to ten

of these are specifically for survival data.<sup>1</sup> Of those, Schwarzer *et al.* deemed three to be ‘na”ive applications to survival data’, and recommended for future research models developed by Liestøl *et al.* (1994) (Liestøl, Andersen, and Andersen 1994), Faraggi and Simon (1995) (Faraggi and Simon 1995), and Biganzoli *et al.* (1998) (E. Biganzoli *et al.* 1998).

This survey will not be as comprehensive as the 2000 survey, and nor has any survey since, although there have been several ANN reviews (B. D. Ripley and Ripley 2001; Huang et al. 2020b; Ohno-Machado 1996; Yang 2010; W. Zhu et al. 2020). ANNs are considered to be a black-box model, with interpretability decreasing steeply as the number of hidden layers and nodes increases. In terms of accessibility there have been relatively few open-source packages developed for survival ANNs; where these are available the focus has historically been in Python, with no R implementations. The new **survivalmodels** (R. Sonabend 2020) package,<sup>2</sup> implements these Python models via **reticulate** (Ushey, Allaire, and Tang 2020). No recurrent neural networks are included in this survey though the survival models SRN (Oh et al. 2018) and RNN-Surv (Giunchiglia, Nemchenko, and Schaar 2018) are acknowledged.

This survey is made slightly more difficult as neural networks are often proposed for many different tasks, which are not necessarily clearly advertised in a paper’s title or abstract. For example, many papers claim to use neural networks for survival analysis and make comparisons to Cox models, whereas the task tends to be death at a particular (usually 5-year) time-point (classification) (I. Han *et al.* 2018; Lundin *et al.* 1999; B. D. Ripley and Ripley 2001; R. M. Ripley, Harris, and Tarassenko 1998; Huseyin Seker *et al.* 2002), which is often not made clear until mid-way through the paper. Reviews and surveys have also conflated these different tasks, for example a very recent review concluded superior performance of ANNs over Cox models, when in fact this is only in classification (Huang et al. 2020a) (RM2) {sec:car\_reduxstrats\_mistakes}. To clarify, this form of classification task does fall into the general *field* of survival analysis, but not the survival *task* ((**box-task-surv?**)). Therefore this is not a comment on the classification task but a reason for omitting these models from this survey.

Using ANNs for feature selection (often in gene expression data) and computer vision is also very common in survival analysis, and indeed it is in this area that most success has been seen (Bello *et al.* 2019; Y.-C. Chen, Ke, and Chiu 2014; Cui *et al.* 2020; Lao *et al.* 2017; McKinney *et al.* 2020; Rietschel, Yoon, and Schaar 2018; H. Seker *et al.* 2002; Zhang *et al.* 2020; X. Zhu, Yao, and Huang 2016), but these are again beyond the scope of this survey.

The key difference between neural networks is in their output layer, required data transformations, the model prediction, and the loss function used to fit the model. Therefore the following are discussed for each of the surveyed models: the loss function for training,  $L$ , the model prediction type,  $\hat{g}$ , and any required data transformation. Notation is continued from the previous surveys with the addition of  $\theta$  denoting model weights (which will be different for each model).

### 16.1.2.1 Probabilistic Survival Models

Unlike other classes of machine learning models, the focus in ANNs has been on probabilistic models. The vast majority make these predictions via reduction to binary classification ???. Whilst almost all of these networks implicitly reduce the problem to classification, most are not transparent in exactly how they do so and none provide clear or detailed interface points in implementation allowing for control over this reduction. Most importantly, the

<sup>1</sup>Schwarzer conflates the prognosis and survival task, therefore it is not clear if all 10 of these are for time-to-event data (at least five definitely are).

<sup>2</sup>Created in order to run the experiments in [@Sonabend2021b].

majority of these models do not detail how valid survival predictions are derived from the binary setting,<sup>3</sup> which is not just a theoretical problem as some implementations, such as the Logistic-Hazard model in **pcox** (Kvamme 2018), have been observed to make survival predictions outside the range [0, 1]. This is not a statement about the performance of models in this section but a remark about the lack of transparency across all probabilistic ANNs.

Many of these algorithms use an approach that formulate the Cox PH as a non-linear model and minimise the partial likelihood. These are referred to as ‘neural-Cox’ models and the earliest appears to have been developed by Faraggi and Simon (Faraggi and Simon 1995). All these models are technically composites that first predict a ranking, however they assume a PH form and in implementation they all appear to return a probabilistic prediction.

**ANN-COX** {#mod-anncox}\ Faraggi and Simon (Faraggi and Simon 1995) proposed a non-linear PH model

$$h(\tau|X_i, \theta) = h_0(\tau) \exp(\phi(X_i\beta)) \quad (16.4)$$

where  $\phi$  is the sigmoid function and  $\theta = \{\beta\}$  are model weights. This model, ‘ANN-COX’, estimates the prediction functional,  $\hat{g}(X^*) = \phi(X^*\hat{\beta})$ . The model is trained with the partial-likelihood function

$$L(\hat{g}, \theta | \mathcal{D}_{train}) = \prod_{i=1}^n \frac{\exp(\sum_{m=1}^M \alpha_m \hat{g}_m(X^*))}{\sum_{j \in \mathcal{R}_{t_i}} \exp(\sum_{m=1}^M \alpha_m \hat{g}_m(X^*))}$$

where  $\mathcal{R}_{t_i}$  is the risk group alive at  $t_i$ ;  $M$  is the number of hidden units;  $\hat{g}_m(X^*) = (1 + \exp(-X^*\hat{\beta}_m))^{-1}$ ; and  $\theta = \{\beta, \alpha\}$  are model weights.

The authors proposed a single hidden layer network, trained using back-propagation and weight optimisation with Newton-Raphson. This architecture did not outperform a Cox PH (Faraggi and Simon 1995). Further adjustments including (now standard) pre-processing and hyper-parameter tuning did not improve the model performance (Mariani et al. 1997). Further independent studies demonstrated worse performance than the Cox model (Faraggi and Simon 1995; Xiang et al. 2000).

**COX-NNET** {#mod-coxnnet}\ COX-NNET (Ching, Zhu, and Garmire 2018) updates the ANN-COX by instead maximising the regularized partial log-likelihood

$$L(\hat{g}, \theta | \mathcal{D}_{train}, \lambda) = \sum_{i=1}^n \Delta_i \left[ \hat{g}(X_i) - \log \left( \sum_{j \in \mathcal{R}_{t_i}} \exp(\hat{g}(X_j)) \right) \right] + \lambda(\|\beta\|_2 + \|w\|_2)$$

with weights  $\theta = (\beta, w)$  and where  $\hat{g}(X_i) = \sigma(wX_i + b)^T\beta$  for bias term  $b$ , and activation function  $\sigma$ ;  $\sigma$  is chosen to be the tanh function ((Equation 16.1)). In addition to weight decay, dropout (Srivastava et al. 2014) is employed to prevent overfitting. Dropout can be thought of as a similar concept to the variable selection in random forests, as each node is randomly deactivated with probability  $p$ , where  $p$  is a hyper-parameter to be tuned.

Independent simulation studies suggest that COX-NNET does not outperform the Cox PH (Michael F. Gensheimer and Narasimhan 2019).

**DeepSurv** {#mod-deepsurv}\ DeepSurv (J. L. Katzman et al. 2018) extends these models to deep learning with multiple hidden layers. The chosen error function is the average

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<sup>3</sup>One could assume they use procedures such as those described in Tutz and Schmid (2016) [@Tutz2016] but there is rarely transparent writing to confirm this.

negative log-partial-likelihood with weight decay

$$L(\hat{g}, \theta | \mathcal{D}_{train}, \lambda) = -\frac{1}{n^*} \sum_{i=1}^n \Delta_i \left[ \left( \hat{g}(X_i) - \log \sum_{j \in \mathcal{R}_{t_i}} \exp(\hat{g}(X_j)) \right) \right] + \lambda \|\theta\|_2^2$$

where  $n^* := \sum_{i=1}^n \mathbb{I}(\Delta_i = 1)$  is the number of uncensored observations and  $\hat{g}(X_i) = \phi(X_i | \theta)$  is the same prediction object as the ANN-COX. State-of-the-art methods are used for data pre-processing and model training. The model architecture uses a combination of fully-connected and dropout layers. Benchmark experiments by the authors indicate that DeepSurv can outperform the Cox PH in ranking tasks (J. Katzman et al. 2016; J. L. Katzman et al. 2018) although independent experiments do not confirm this (Zhao and Feng 2020).

\*\*Cox-Time\*\* {#mod-coxtime}\ Kvamme *et al.* (Kvamme, Borgan, and Scheel 2019) build on these models by allowing time-varying effects. The loss function to minimise, with regularization, is given by

$$L(\hat{g}, \theta | \mathcal{D}_{train}, \lambda) = \frac{1}{n} \sum_{i: \Delta_i=1} \log \left( \sum_{j \in \mathcal{R}_{t_i}} \exp[\hat{g}(X_j, T_i) - \hat{g}(X_i, T_i)] \right) + \lambda \sum_{i: \Delta_i=1} \sum_{j \in \mathcal{R}_{t_i}} |\hat{g}(X_j, T_i)|$$

where  $\hat{g} = \hat{g}_1, \dots, \hat{g}_n$  is the same non-linear predictor but with a time interaction and  $\lambda$  is the regularization parameter. The model is trained with stochastic gradient descent and the risk set,  $\mathcal{R}_{t_i}$ , in the equation above is instead reduced to batches, as opposed to the complete dataset. ReLU activations (Nair and Hinton 2010) and dropout are employed in training. Benchmark experiments indicate good performance of Cox-Time, though no formal statistical comparisons are provided and hence no comment about general performance can be made.

**ANN-CDP** {#mod-anncdp}\ One of the earliest ANNs that was noted by Schwarzer *et al.* (Schwarzer, Vach, and Schumacher 2000) was developed by Liestøl *et al.* (Liestøl, Andersen, and Andersen 1994) and predicts conditional death probabilities (hence ‘ANN-CDP’). The model first partitions the continuous survival times into disjoint intervals  $\mathcal{I}_k$ ,  $k = 1, \dots, m$  such that  $\mathcal{I}_k$  is the interval  $(t_{k-1}, t_k]$ . The model then studies the logistic Cox model (proportional odds) (Cox 1972) given by

$$\frac{p_k(\mathbf{x})}{q_k(\mathbf{x})} = \exp(\eta + \theta_k)$$

where  $p_k = 1 - q_k$ ,  $\theta_k = \log(p_k(0)/q_k(0))$  for some baseline probability of survival,  $q_k(0)$ , to be estimated;  $\eta$  is the usual linear predictor, and  $q_k = P(T \geq T_k | T \geq T_{k-1})$  is the conditional survival probability at time  $T_k$  given survival at time  $T_{k-1}$  for  $k = 1, \dots, K$  total time intervals. A logistic activation function is used to predict  $\hat{g}(X^*) = \phi(\eta + \theta_k)$ , which provides an estimate for  $\hat{p}_k$ .

The model is trained on discrete censoring indicators  $D_{ki}$  such that  $D_{ki} = 1$  if individual  $i$  dies in interval  $\mathcal{I}_k$  and 0 otherwise. Then with  $K$  output nodes and maximum likelihood estimation to find the model parameters,  $\hat{\eta}$ , the final prediction provides an estimate for the conditional death probabilities  $\hat{p}_k$ . The negative log-likelihood to optimise is given by

$$L(\hat{g}, \theta | \mathcal{D}_{train}) = \sum_{i=1}^n \sum_{k=1}^{m_i} [D_{ki} \log(\hat{p}_k(X_i)) + (1 - D_{ki}) \log(\hat{q}_k(X_i))]$$

where  $m_i$  is the number of intervals in which observation  $i$  is not censored.

Liestøl *et al.*{} discuss different weighting options and how they correspond to the PH assumption. In the most generalised case, a weight-decay type regularization is applied to the model weights given by

$$\alpha \sum_l \sum_k (w_{kl} - w_{k-1,l})^2$$

where  $w$  are weights, and  $\alpha$  is a hyper-parameter to be tuned, which can be used alongside standard weight decay. This corresponds to penalizing deviations from proportionality thus creating a model with approximate proportionality. The authors also suggest the possibility of fixing the weights to be equal in some nodes and different in others; equal weights strictly enforces the proportionality assumption. Their simulations found that removing the proportionality assumption completely, or strictly enforcing it, gave inferior results. Comparing their model to a standard Cox PH resulted in a ‘better’ negative log-likelihood, however this is not a precise evaluation metric and an independent simulation would be

preferred. Finally Listøl *et al.* included a warning “The flexibility is, however, obtained at unquestionable costs: many parameters, difficult interpretation of the parameters and a slow numerical procedure’’ (Liestol, Andersen, and Andersen 1994).

**PLANN** {#mod-plann}\ Biganzoli *et al.* (1998) (E. Biganzoli et al. 1998) studied the same proportional-odds model as the ANN-CDP (Liestol, Andersen, and Andersen 1994). Their model utilises partial logistic regression (Efron 1988) with added hidden nodes, hence ‘PLANN’. Unlike ANN-CDP, PLANN predicts a smoothed hazard function by using smoothing splines. The continuous time outcome is again discretised into disjoint intervals  $t_m, m = 1, \dots, M$ . At each time-interval,  $t_m$ , the number of events,  $d_m$ , and number of subjects at risk,  $n_m$ , can be used to calculate the discrete hazard function,<sup>4</sup>

$$\hat{h}_m = \frac{d_m}{n_m}, m = 1, \dots, M \quad (16.5)$$

This quantity is used as the target to train the neural network. The survival function is then estimated by the Kaplan-Meier type estimator,

$$\hat{S}(\tau) = \prod_{m:t_m \leq \tau} (1 - \hat{h}_m) \quad (16.6)$$

The model is fit by employing one of the more ‘usual’ survival reduction strategies in which an observation’s survival time is treated as a covariate in the model (Tutz and Schmid 2016). As this model uses discrete time, the survival time is discretised into one of the  $M$  intervals. This approach removes the proportional odds constraint as interaction effects between time and covariates can be modelled (as time-updated covariates). Again the model makes predictions at a given time  $m$ ,  $\phi(\theta_m + \eta)$ , where  $\eta$  is the usual linear predictor,  $\theta$  is the baseline proportional odds hazard  $\theta_m = \log(h_m(0)/(1 - h_m(0)))$ . The logistic activation provides estimates for the discrete hazard,

$$h_m(X_i) = \frac{\exp(\theta_m + \hat{\eta})}{1 + \exp(\theta_m + \hat{\eta})}$$

which is smoothed with cubic splines (Efron 1988) that require tuning.

A cross-entropy error function is used for training

$$L(\hat{h}, \theta | \mathcal{D}_{train}, a) = - \sum_{m=1}^M \left[ \hat{h}_m \log \left( \frac{h_l(X_i, a_l)}{\hat{h}_m} \right) + (1 - \hat{h}_m) \log \left( \frac{1 - h_l(X_i, a_l)}{1 - \hat{h}_m} \right) \right] n_m$$

where  $h_l(X_i, a_l)$  is the discrete hazard  $h_l$  with smoothing at mid-points  $a_l$ . Weight decay can be applied and the authors suggest  $\lambda \approx 0.01 - 0.1$  (E. Biganzoli et al. 1998), though they make use of an AIC type criterion instead of cross-validation.

This model makes smoothed hazard predictions at a given time-point,  $\tau$ , by including  $\tau$  in the input covariates  $X_i$ . Therefore the model first requires transformation of the input data by replicating all observations and replacing the single survival indicator  $\Delta_i$ , with a time-dependent indicator  $D_{ik}$ , the same approach as in ANN-CDP. Further developments have extended the PLANN to Bayesian modelling, and for competing risks (E. M. Biganzoli, Ambrogi, and Boracchi 2009).

No formal comparison is made to simpler model classes. The authors recommend ANNs primarily for exploration, feature selection, and understanding underlying patterns in the data (E. M. Biganzoli, Ambrogi, and Boracchi 2009).

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<sup>4</sup>Derivation of this as a ‘hazard’ estimator follows trivially by comparison to the Nelson-Aalen estimator.

**Nnet-survival** {#mod-nnetsurvival}\ Aspects of the PLANN algorithm have been generalised into discrete-time survival algorithms in several papers (Michael F. Gensheimer and Narasimhan 2019; **Kvamme2019?**; Mani et al. 1999; Street 1998). Various estimates have been derived for transforming the input data to a discrete hazard or survival function. Though only one is considered here as it is the most modern and has a natural interpretation as the ‘usual’ Kaplan-Meier estimator for the survival function. Others by Street (1998) (Street 1998) and Mani (1999) (Mani et al. 1999) are acknowledged. The discrete hazard estimator (Equation 16.5),  $\hat{h}$ , is estimated and these values are used as the targets for the ANN. For the error function, the mean negative log-likelihood for discrete time (**Kvamme2019?**) is minimised to estimate  $\hat{h}$ ,

$$L(\hat{h}, \theta | \mathcal{D}_{train}) = -\frac{1}{n} \sum_{i=1}^n \sum_{j=1}^{k(T_i)} (\mathbb{I}(T_i = \tau_j, \Delta_i = 1) \log[\hat{h}_i(\tau_j)] + (1 - \mathbb{I}(T_i = \tau_j, \Delta_i = 1)) \log(1 - \hat{h}_i(\tau_j)))$$

where  $k(T_i)$  is the time-interval index in which observation  $i$  dies/is censored,  $\tau_j$  is the  $j$ th discrete time-interval, and the prediction of  $\hat{h}$  is obtained via

$$\hat{h}(\tau_j | \mathcal{D}_{train}) = [1 + \exp(-\hat{g}_j(\mathcal{D}_{train}))]^{-1}$$

where  $\hat{g}_j$  is the  $j$ th output for  $j = 1, \dots, m$  discrete time intervals. The number of units in the output layer for these models corresponds to the number of discrete-time intervals. Deciding the width of the time-intervals is an additional hyper-parameter to consider.

Gensheimer and Narasimhan’s ‘Nnet-survival’ (Michael F. Gensheimer and Narasimhan 2019) has two different implementations. The first assumes a PH form and predicts the linear predictor in the final layer, which can then be composed to a distribution. Their second ‘flexible’ approach instead predicts the log-odds of survival in each node, which are then converted to a conditional probability of survival,  $1 - h_j$ , in a given interval using the sigmoid activation function. The full survival function can be derived with (Equation 16.6). The model has been demonstrated not to outperform the Cox PH with respect to Harrell’s C or the Graf (Brier) score (Michael F. Gensheimer and Narasimhan 2019).

**PC-Hazard** {#mod-pchazard}\ Kvamme and Borgan deviate from nnet-survival in their ‘PC-Hazard’ (**Kvamme2019?**) by first considering a discrete-time approach with a softmax activation function influenced by multi-class classification. They expand upon this by studying a piecewise constant hazard function in continuous time and defining the mean negative log-likelihood as

$$L(\hat{g}, \theta | \mathcal{D}_{train}) = -\frac{1}{n} \sum_{i=1}^n \left( \Delta_i X_i \log \tilde{\eta}_{k(T_i)} - X_i \tilde{\eta}_{k(T_i)} \rho(T_i) - \sum_{j=1}^{k(T_i)-1} \tilde{\eta}_j X_i \right)$$

where  $k(T_i)$  and  $\tau_i$  is the same as defined above,  $\rho(t) = \frac{t - \tau_{k(t)-1}}{\Delta \tau_{k(t)}}$ ,  $\Delta \tau_j = \tau_j - \tau_{j-1}$ , and  $\tilde{\eta}_j := \log(1 + \exp(\hat{g}_j(X_i)))$  where again  $\hat{g}_j$  is the  $j$ th output for  $j = 1, \dots, m$  discrete time intervals. Once the weights have been estimated, the predicted survival function is given by

$$\hat{S}(\tau, X^* | \mathcal{D}_{train}) = \exp(-X^* \tilde{\eta}_{k(\tau)} \rho(\tau)) \prod_{j=1}^{k(\tau)-1} \exp(-\tilde{\eta}_j(X^*))$$

Benchmark experiments indicate similar performance to nnet-survival (**Kvamme2019?**), an unsurprising result given their implementations are identical with the exception of the

loss function (**Kvamme2019?**), which is also similar for both models. A key result found that varying values for interval width lead to significant differences and therefore should be carefully tuned.

**DNNSurv** {#mod-dnnsurv}\ A very recent (pre-print) approach (Zhao and Feng 2020) instead first computes ‘pseudo-survival probabilities’ and uses these to train a regression ANN with sigmoid activation and squared error loss. These pseudo-probabilities are computed using a jackknife-style estimator given by

$$\tilde{S}_{ij}(T_{j+1}, \mathcal{R}_{t_j}) = n_j \hat{S}(T_{j+1} | \mathcal{R}_{t_j}) - (n_j - 1) \hat{S}^{-i}(T_{j+1} | \mathcal{R}_{t_j})$$

where  $\hat{S}$  is the IPCW weighted Kaplan-Meier estimator (defined below) for risk set  $\mathcal{R}_{t_j}$ ,  $\hat{S}^{-i}$  is the Kaplan-Meier estimator for all observations in  $\mathcal{R}_{t_j}$  excluding observation  $i$ , and  $n_j := |\mathcal{R}_{t_j}|$ . The IPCW weighted Kaplan-Meier estimate is found via the IPCW Nelson-Aalen estimator,

$$\hat{H}(\tau | \mathcal{D}_{train}) = \sum_{i=1}^n \int_0^\tau \frac{\mathbb{I}(T_i \leq u, \Delta_i = 1) \hat{W}_i(u)}{\sum_{j=1}^n \mathbb{I}(T_j \geq u) \hat{W}_j(u)} du$$

where  $\hat{W}_i, \hat{W}_j$  are subject specific IPC weights.

In their simulation studies, they found no improvement over other proposed neural networks. Arguably the most interesting outcome of their paper are comparisons of multiple survival ANNs at specific time-points, evaluated with C-index and Brier score. Their results indicate identical performance from all models. They also provide further evidence of neural networks not outperforming a Cox PH when the PH assumption is valid. However, in their non-PH dataset, DNNSurv appears to outperform the Cox model (no formal tests are provided). Data is replicated similarly to previous models except that no special indicator separates censoring and death, this is assumed to be handled by the IPCW pseudo probabilities.

**DeepHit** {#mod-deephit}\ DeepHit (C. Lee et al. 2018) was originally built to accommodate competing risks, but only the non-competing case is discussed here (Kvamme, Borgan, and Scheel 2019). The model builds on previous approaches by discretising the continuous time outcome, and makes use of a composite loss. It has the advantage of making no parametric assumptions and directly predicts the probability of failure in each time-interval (which again correspond to different terminal nodes), i.e.  $\hat{g}(\tau_k | \mathcal{D}_{test}) = \hat{P}(T^* = \tau_k | X^*)$  where again  $\tau_k, k = 1, \dots, K$  are the distinct time intervals. The estimated survival function is found with  $\hat{S}(\tau_K | X^*) = 1 - \sum_{k=1}^K \hat{g}_i(\tau_k | X^*)$ . ReLU activations were used in all fully connected layers and a softmax activation in the final layer. The losses in the composite error function are given by

$$L_1(\hat{g}, \theta | \mathcal{D}_{train}) = - \sum_{i=1}^N [\Delta_i \log(\hat{g}_i(T_i)) + (1 - \Delta_i) \log(\hat{S}_i(T_i))]$$

and

$$L_2(\hat{g}, \theta | \mathcal{D}_{train}, \sigma) = \sum_{i \neq j} \Delta_i \mathbb{I}(T_i < T_j) \sigma(\hat{S}_i(T_i), \hat{S}_j(T_i))$$

for some convex loss function  $\sigma$  and where  $\hat{g}_i(t) = \hat{g}(t | X_i)$ . Again these can be seen to be a cross-entropy loss and a ranking loss. Benchmark experiments demonstrate the model outperforming the Cox PH and RSFs (C. Lee et al. 2018) with respect to separation, and an independent experiment supports these findings (Kvamme, Borgan, and Scheel 2019). However, the same independent study demonstrated worse performance than a Cox PH with respect to the integrated Brier score (Graf et al. 1999).

### 16.1.2.2 Deterministic Survival Models

Whilst the vast majority of survival ANNs have focused on probabilistic predictions (often via ranking), a few have also tackled the deterministic or ‘hybrid’ problem.

**RankDeepSurv** {#mod-rankdeepsurv}\ Jing *et al.* (Jing et al. 2019) observed the past two decades of research in survival ANNs and then published a completely novel solution, RankDeepSurv, which makes predictions for the survival time  $\hat{T} = (\hat{T}_1, \dots, \hat{T}_n)$ . They proposed a composite loss function

$$L(\hat{T}, \theta | \mathcal{D}_{train}, \alpha, \gamma, \lambda) = \alpha L_1(\hat{T}, T, \Delta) + \gamma L_2(\hat{T}, T, \Delta) + \lambda \|\theta\|_2^2$$

where  $\theta$  are the model weights,  $\alpha, \gamma \in \mathbb{R}_{>0}$ ,  $\lambda$  is the shrinkage parameter, by a slight abuse of notation  $T = (T_1, \dots, T_n)$  and  $\Delta = (\Delta_1, \dots, \Delta_n)$ , and

$$L_1(\hat{T}, \theta | \mathcal{D}_{train}) = \frac{1}{n} \sum_{\{i: I(i)=1\}} (\hat{T}_i - T_i)^2; \quad I(i) = \begin{cases} 1, & \Delta_i = 1 \cup (\Delta_i = 0 \cap \hat{T}_i \leq T_i) \\ 0, & \text{otherwise} \end{cases}$$

$$L_2(\hat{T}, \theta | \mathcal{D}_{train}) = \frac{1}{n} \sum_{\{i,j: I(i,j)=1\}} [(T_j - T_i) - (\hat{T}_j - \hat{T}_i)]^2; \quad I(i,j) = \begin{cases} 1, & T_j - T_i > \hat{T}_j - \hat{T}_i \\ 0, & \text{otherwise} \end{cases}$$

where  $\hat{T}_i$  is the predicted survival time for observation  $i$ . A clear contrast can be made between these loss functions and the constraints used in SSVM-Hybrid (Van Belle et al. 2011) (Section 14.2).  $L_1$  is the squared second constraint in 14.2.3 and  $L_2$  is the squared first constraint in 14.2.3. However  $L_1$  in RankDeepSurv discards the squared error difference for all censored observations when the prediction is lower than the observed survival time; which is problematic as if someone is censored at time  $T_i$  then it is guaranteed that their true survival time is greater than  $T_i$  (this constraint may be more sensible if the inequality were reversed). An advantage to this loss is, like the SSVM-Hybrid, it enables a survival time interpretation for a ranking optimised model; however these ‘survival times’ should be interpreted with care.

The authors propose a model architecture with several fully connected layers with the ELU (Clevert, Unterthiner, and Hochreiter 2015) activation function and a single dropout layer. Determining the success of this model is not straightforward. The authors claim superiority of RankDeepSurv over Cox PH, DeepSurv, and RSFs however this is an unclear comparison (RM2) {sec:car\_reduxstrats\_mistakes} that requires independent study.

### 16.1.3 Conclusions

There have been many advances in neural networks for survival analysis. It is not possible to review all proposed survival neural networks without diverting too far from the book scope. This survey of ANNs should demonstrate two points: firstly that the vast majority (if not all) of survival ANNs are reduction models that either find a way around censoring via imputation or discretisation of time-intervals, or by focusing on partial likelihoods only; secondly that no survival ANN is fully accessible or transparent.

Despite ANNs being highly performant in other areas of supervised learning, there is strong evidence that the survival ANNs above are inferior to a Cox PH when the data follows the PH assumption or when variables are linearly related (Michael F. Gensheimer and Narasimhan 2018; Luxhoj and Shyur 1997; Ohno-Machado 1997; Puddu and Menotti 2012; Xiang et al. 2000; Yang 2010; Yasodhara, Bhat, and Goldenberg 2018; Zhao and Feng 2020). There are not enough experiments to make conclusions in the case when the data is non-PH.

Experiments in (R. E. B. Sonabend 2021) support the finding that survival ANNs are not performant.

There is evidence that many papers introducing neural networks do not utilise proper methods of comparison or evaluation (Király, Mateen, and Sonabend 2018) and in conducting this survey, these findings are further supported. Many papers made claims of being ‘superior’ to the Cox model based on unfair comparisons (RM2){sec:car\_reduxstrats\_mistakes} or miscommunicating (or misinterpreting) results (e.g. (Fotso 2018)). At this stage, it does not seem possible to make any conclusions about the effectiveness of neural networks in survival analysis. Moreover, even the authors of these models have pointed out problems with transparency (E. M. Biganzoli, Ambrogi, and Boracchi 2009; Liestol, Andersen, and Andersen 1994), which was further highlighted by Schwarzer *et al.* (Schwarzer, Vach, and Schumacher 2000).

Finally, accessibility of neural networks is also problematic. Many papers do not release their code and instead just state their networks architecture and available packages. In theory, this is enough to build the models however this does not guarantee the reproducibility that is usually expected. For users with a technical background and good coding ability, many of the models above could be implemented in one of the neural network packages in R, such as **nnet** (N. Venables and D. Ripley 2002) and **neuralnet** (Fritsch, Guenther, and N. Wright 2019); though in practice the only package that does contain these models, **survivalmodels**, does not directly implement the models in R (which is much slower than Python) but provides a method for interfacing the Python implementations in **pcox** (Kvamme 2018).

#### Further reading

- Schwarzer, Vach, and Schumacher (2000) provided an early survey of neural networks, focusing on ways in which neural networks have been ‘misused’ in the context of survival analysis. Whilst neural networks have moved on substantially since, their early observations remain valid today.



# 17

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## Choosing Models

TODO (150-200 WORDS)

! Page coming soon!

We are working on this page and it will be available soon!



**Part IV**

**Reduction Techniques**



# 18

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## Reductions

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TODO (150-200 WORDS)

! Major changes expected!

This page is a work in progress and major changes will be made over time.

In this chapter, composition and reduction are formally introduced, defined and demonstrated within survival analysis. Neither of these are novel concepts in general or in survival, with several applications already seen earlier when reviewing models (particularly in neural networks), however a lack of formalisation has led to much repeated work and at times questionable applications ([?@sec-surv-ml-models-nn](#)). The primary purpose of this chapter is to formalise composition and reduction for survival and to unify references and strategies for future use. These strategies are introduced in the context of minimal ‘workflows’ and graphical ‘pipelines’ in order to maximise their generalisability.

A *workflow* is a generic term given to a series of sequential operations. For example a standard ML workflow is fit/predict/evaluate, which means a model is fit, predictions are made, and these are evaluated. In this book, a *pipeline* is the name given to a concrete workflow. Section 18.1 demonstrates how pipelines are represented in this book.

Composition (Section 18.2) is a general process in which an object is built (or composed) from other objects and parameters. Reduction (Section 18.3) is a closely related concept that utilises composition in order to transform one problem into another. Concrete strategies for composition and reduction are detailed in sections Section 18.4 and Section 18.5.

### Notation and Terminology

The notation introduced in Chapter 3 is recapped for use in this chapter: the generative survival template for the survival setting is given by  $(X, T, \Delta, Y, C)$  t.v.i.  $\mathcal{X} \times \mathcal{T} \times \{0, 1\} \times \mathcal{T} \times \mathcal{T}$  where  $\mathcal{X} \subseteq \mathbb{R}^p$  and  $\mathcal{T} \subseteq \mathbb{R}_{\geq 0}$ , where  $C, Y$  are unobservable,  $T := \min\{Y, C\}$ , and  $\Delta = \mathbb{I}(Y = T)$ . Random survival data is given by  $(X_i, T_i, \Delta_i, Y_i, C_i) \stackrel{i.i.d.}{\sim} (X, T, \Delta, Y, C)$ . Usually data will instead be presented as a training dataset,  $\mathcal{D}_{train} = \{(X_1, T_1, \Delta_1), \dots, (X_n, T_n, \Delta_n)\}$  where  $(X_i, T_i, \Delta_i) \stackrel{i.i.d.}{\sim} (X, T, \Delta)$ , and some test data  $\mathcal{D}_{test} = (X^*, T^*, \Delta^*) \sim (X, T, \Delta)$ .

For regression models the generative template is given by  $(X, Y)$  t.v.i.  $\mathcal{X} \subseteq \mathbb{R}^p$  and  $Y \subseteq \mathbb{R}$ . As with the survival setting, a regression training set is given by  $\{(X_1, Y_1), \dots, (X_n, Y_n)\}$  where  $(X_i, Y_i) \stackrel{i.i.d.}{\sim} (X, Y)$  and some test data  $(X^*, Y^*) \sim (X, Y)$ .

## 18.1 Representing Pipelines

Before introducing concrete composition and reduction algorithms, this section briefly demonstrates how these pipelines will be represented in this book.

Pipelines are represented by graphs designed in the following way: all are drawn with operations progressing sequentially from left to right; graphs are comprised of nodes (or ‘vertices’) and arrows (or ‘directed edges’); a rounded rectangular node represents a process such as a function or model fitting/predicting; a (regular) rectangular node represents objects such as data or hyper-parameters. Output from rounded nodes are sometimes explicitly drawn but when omitted the output from the node is the input to the next.

These features are demonstrated in `?@fig-car-example`. Say  $y = 2$  and  $a = 2$ , then: data is provided ( $y = 2$ ) and passed to the shift function ( $f(x) = x + 2$ ), the output of this function ( $y = 4$ ) is passed directly to the next ( $h(x|a) = x^a$ ), this function requires a parameter which is also input ( $a = 2$ ), finally the resulting output is returned ( $y^* = 16$ ). Programmatically,  $a = 2$  would be a hyper-parameter that is stored and passed to the required function when the function is called.

This pipeline is represented as a pseudo-algorithm in `(alg-car-ex?)`, though of course is overly complicated and in practice one would just code  $(y + 2)^a$ .

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## 18.2 Introduction to Composition

This section introduces composition, defines a taxonomy for describing compositors (Section 18.2.1), and provides some motivating examples of composition in survival analysis (Section 18.2.2).

In the simplest definition, a model (be it mathematical, computational, machine learning, etc.) is called a *composite model* if it is built of two or more constituent parts. This can be simplest defined in terms of objects. Just as objects in the real-world can be combined in some way, so can mathematical objects. The exact ‘combining’ process (or ‘compositor’) depends on the specific composition, so too do the inputs and outputs. By example, a wooden table can be thought of as a composite object (Figure 18.1). The inputs are wood and nails, the combining process is hammering (assuming the wood is pre-chopped), and the output is a surface for eating. In mathematics, this process is mirrored. Take the example of a shifted linear regression model. This is defined by a linear regression model,  $f(x) = \beta_0 + x\beta_1$ , a shifting parameter,  $\alpha$ , and a compositor  $g(x|\alpha) = f(x) + \alpha$ . Mathematically this example is overly trivial as this could be directly modelled with  $f(x) = \alpha + \beta_0 + x\beta_1$ , but algorithmically there is a difference. The composite model  $g$ , is defined by first fitting the linear regression model,  $f$ , and then applying a shift,  $\alpha$ ; as opposed to fitting a directly shifted model.

### Why Composition?

Tables tend to be better surfaces for eating your dinner than bundles of wood. Or in modelling terms, it is well-known that ensemble methods (e.g. random forests) will generally outperform their components (e.g. decision trees). All ensemble methods are composite models and this demonstrates one of the key use-cases of composition: improved predictive

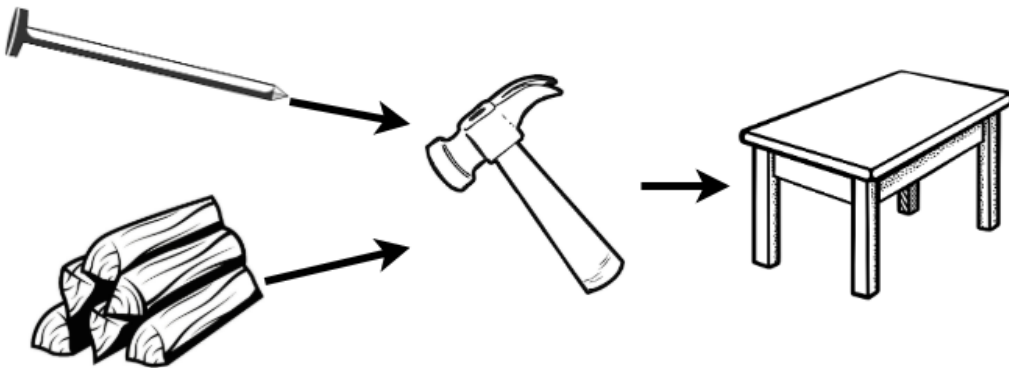


Figure 18.1: Visualising composition in the real-world. A table is a composite object built from nails and wood, which are combined with a hammer ‘compositor’. Figure not to scale.

performance. The second key use-case is reduction, which is fully discussed in Section 18.3. Section 18.2.2 motivates composition in survival analysis by demonstrating how it is already prevalent but requires formalisation to make compositions more transparent and accessible.

### Composite Model vs. Sub-models

A bundle of wood and nails is not a table and 1,000 decision trees are not a random forest, both require a compositor. The compositor in a composite model combines the components into a single model. Considering a composite model as a single model enables the hyper-parameters of the compositor and the component model(s) to be efficiently tuned whilst being evaluated as a single model. This further allows the composite to be compared to other models, including its own components, which is required to justify complexity instead of parsimony in model building (?@sec-eval-why-why).

#### 18.2.1 Taxonomy of Compositors

Just as there are an infinite number of ways to make a table, composition can come in infinite forms. However there are relatively few categories that these can be grouped into. Two primary taxonomies are identified here. The first is the ‘composition type’ and relates to the number of objects composed:

[i]] i. Single-Object Composition (SOC) – This form of composition either makes use of parameters or a transformation to alter a single object. The shifted linear regression model above is one example of this, another is given in Section 18.4.3. i. Multi-Object Composition (MOC) – In contrast, this form of composition combines multiple objects into a single one. Both examples in Section 18.2.2 are multi-object compositions.

The second grouping is the ‘composition level’ and determines at what ‘level’ the composition takes place:

[i]] i. Prediction Composition – This applies at the level of predictions; the component models could be forgotten at this point. Predictions may be combined from multiple models

(MOC) or transformed from a single model (SOC). Both examples in Section 18.2.2 are prediction compositions. i. Task Composition – This occurs when one task (e.g. regression) is transformed to one or more others (e.g. classification), therefore always SOC. This is seen mainly in the context of reduction (Section 18.3). i. Model Composition – This is commonly seen in the context of wrappers (Section 18.5.1.4), in which one model is contained within another. i. Data Composition – This is transformation of training/testing data types, which occurs at the first stage of every pipeline.

### 18.2.2 Motivation for Composition

Two examples are provided below to demonstrate common uses of composition in survival analysis and to motivate the compositions introduced in Section 18.4.

#### Example 1: Cox Proportional Hazards

Common implementations of well-known models can themselves be viewed as composite models, the Cox PH is the most prominent example in survival analysis. Recall the model defined by

$$h(\tau|X_i) = h_0(\tau) \exp(\beta X_i)$$

where  $h_0$  is the baseline hazard and  $\beta$  are the model coefficients.

This can be seen as a composite model as Cox defines the model in two stages (Cox 1972): first fitting the  $\beta$ -coefficients using the partial likelihood and then by suggesting an estimate for the baseline distribution. This first stage produces a linear predictor return type (`?@sec-surv-set-types`) and the second stage returns a survival distribution prediction. Therefore the Cox model for linear predictions is a single (non-composite) model, however when used to make distribution predictions then it is a composite. Cox implicitly describes the model as a composite by writing “alternative simpler procedures would be worth having” (Cox 1972), which implies a decision in fitting (a key feature of composition). This composition is formalised in Section 18.4.1 as a general pipelines. The Cox model utilises the pipeline with a PH form and Kaplan-Meier baseline.

#### Example 2: Random Survival Forests

Fully discussed in Chapter 13, random survival forests are composed from many individual decision trees via a prediction composition algorithm (`((alg-rsf-pred?))`). In general, random forests perform better than their component decision trees, which tends to be true of all ensemble methods. Aggregation of predictions in survival analysis requires slightly more care than other fields due to the multiple prediction types, however this is still possible and is formalised in Section 18.4.4.

## 18.3 Introduction to Reduction

This section introduces reduction, motivates its use in survival analysis (Section 18.3.1), details an abstract reduction pipeline and defines the difference between a complete/incomplete reduction (Section 18.3.2), and outlines some common mistakes that have been observed in the literature when applying reduction (Section 18.3.3).

Reduction is a concept found across disciplines with varying definitions. This report uses the Langford definition: reduction is “a complex problem decomposed into simpler subproblems so that a solution to the subproblems gives a solution to the complex problem” (Langford et al. 2016). Generalisation (or induction) is a common real-world use of reduction, for example sampling a subset of a population in order to estimate population-level results. The true answer (population-level values) may not always be found in this way but very good approximations can be made with simpler sub-problems (sub-sampling).

Reductions are workflows that utilise composition. By including hyper-parameters, even complex reduction strategies can remain relatively flexible. To illustrate reduction by example, recall the table-building example (Section 18.2) in which the task of interest is to acquire a table. The most direct but complex solution is to fell a tree and directly saw it into a table (Figure 18.2, top), clearly this is not a sensible process. Instead the problem can be reduced into simpler sub-problems: saw the tree into bundles of wood, acquire nails, and then use the ‘hammer compositor’ (Figure 18.1) to create a table (Figure 18.2, bottom).

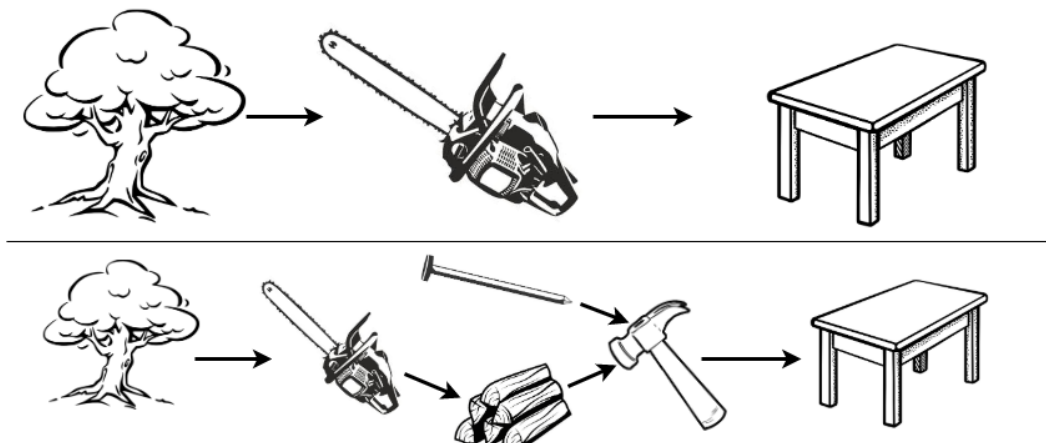


Figure 18.2: Visualising reduction in the real-world. The complex process (top) of directly sawing a tree into a table is inefficient and unnecessarily complex. The reduction (bottom) that involves first creating bundles of wood is simpler, more efficient, and yields the same result, though technically requiring more steps.

In a modelling example, predicting a survival distribution with the Cox model can be viewed as a reduction in which two sub-problems are solved and composed:

- i. predict continuous ranking;
- ii. estimate baseline hazard; and
- iii. compose with (Section 18.4.1).

This is visualised as a reduction strategy in `?@fig-car-cargraph`. The entire process from defining the original problem, to combining the simpler sub-solutions (in green), is the reduction (in red).

### 18.3.1 Reduction Motivation

Formalisation of reduction positively impacts upon accessibility, transparency, and predictive performance. Improvements to predictive performance have already been demonstrated when comparing random forests to decision trees. In addition, a reduction with multiple stages and many hyper-parameters allows for fine tuning for improved transparency and model performance (usual overfitting caveat applies, as does the trade-off described in [?@sec-car-pipelines-trade](#)).

The survey of ANNs ([?@sec-surv-ml-models-nn](#)) demonstrated how reduction is currently utilised without transparency. Many of these ANNs are implicitly reductions to probabilistic classification (Section 18.5.1.6) however none include details about how the reduction is performed. Furthermore in implementation, none provide interface points to the reduction hyper-parameters. Formalisation encourages consistent terminology, methodology and transparent implementation, which can only improve model performance by exposing further hyper-parameters.

Accessibility is improved by formalising specific reduction workflows that previously demanded expert knowledge in deriving, building, and running these pipelines.

Finally there is an economic and efficiency advantage to reduction. A reduction model is relatively ‘cheap’ to explore as they utilise pre-established models and components to solve a new problem. Therefore if a certain degree of predictive ability can be demonstrated from reduction models, it may not be worth the expense of pursuing more novel ideas and hence reduction can help direct future research.

### 18.3.2 Task, Loss, and Data Reduction

Reduction can be categorised into task, loss, and data reduction, often these must be used in conjunction with each other. The direction of the reductions may be one- or two-way; this is visualised in [?@fig-car-reduxdiag](#). This diagram should not be viewed as a strict fit/predict/evaluation workflow but instead as a guidance for which tasks,  $T$ , data,  $D$ , models,  $M$ , and losses,  $L$ , are required for each other. The subscript  $O$  refers to the original object ‘level’ before reduction, whereas the subscript  $R$  is in reference to the reduced object.

The individual task, model, and data compositions in the diagram are listed below, the reduction from survival to classification (Section 18.5.1) is utilised as a running example to help exposition.

- $T_O \rightarrow T_R$ : By definition of a machine learning reduction, task reduction will always be one way. A more complex task,  $T_O$ , is reduced to a simpler one,  $T_R$ , for solving.  $T_R$  could also be multiple simpler tasks. For example, solving a survival task,  $T_O$ , by classification,  $T_R$  (Section 18.5.1).
- $T_R \rightarrow M_R$ : All machine learning tasks have models that are designed to solve them. For example logistic regression,  $M_R$ , for classification tasks,  $T_R$ .
- $M_R \rightarrow M_O$ : The simpler models,  $M_R$ , are used for the express purpose to solve the original task,  $T_O$ , via solving the simpler ones. To solve  $T_O$ , a compositor must be applied, which may transform one (SOC) or multiple models (MOC) at a model- or prediction-level, thus creating  $M_O$ . For example predicting survival probabilities with logistic regression,  $M_R$ , at times  $1, \dots, \tau^*$  for some  $\tau^* \in \mathbb{N}_{>0}$  (Section 18.5.1.4).
- $M_O \rightarrow T_O$ : The original task should be solvable by the composite model. For example predicting a discrete survival distribution by concatenating probabilistic predictions at the times  $1, \dots, \tau^*$  (Section 18.5.1.6).
- $D_O \rightarrow D_R$ : Just as the tasks and models are reduced, the data required to fit these must

likewise be reduced. Similarly to task reduction, data reduction can usually only take place in one direction, to see why this is the case take an example of data reduction by summaries. If presented with 10 data-points  $\{1, 1, 1, 5, 7, 3, 5, 4, 3, 3\}$  then these could be reduced to a single point by calculating the sample mean, 3.3. Clearly given only the number 3.3 there is no strategy to recover the original data. There are very few (if any) data reduction strategies that allow recovery of the original data. Continuing the running example, survival data,  $D_O$ , can be binned (Section 18.5.1.1) to classification data,  $D_R$ .

There is no arrow between  $D_O$  and  $M_O$  as the composite model is never fit directly, only via composition from  $M_R \rightarrow M_O$ . However, the original data,  $D_O$ , is required when evaluating the composite model against the respective loss,  $L_O$ .<sup>1</sup> Reduction should be directly comparable to non-reduction models, hence this diagram does not include loss reduction and instead insists that all models are compared against the same loss  $L_O$ .

A reduction is said to be *complete* if there is a full pipeline from  $T_O \rightarrow M_O$  and the original task is solved, otherwise it is *incomplete*. The simplest complete reduction is comprised of the pipeline  $T_O \rightarrow T_R \rightarrow M_R \rightarrow M_O$ . Usually this is not sufficient on its own as the reduced models are fit on the reduced data,  $D_R \rightarrow M_R$ .

A complete reduction can be specified by detailing:

- i. the original task and the sub-task(s) to be solved,  $T_O \rightarrow T_R$ ;
- ii. the original dataset and the transformation to the reduced one,  $D_O \rightarrow D_R$  (if required); and
- iii. the composition from the simpler model to the complex one,  $M_R \rightarrow M_O$ .

### 18.3.3 Common Mistakes in Implementation of Reduction

In surveying models and measures, several common mistakes in the implementation of reduction and composition were found to be particularly prevalent and problematic throughout the literature. It is assumed that these are indeed mistakes (not deliberate) and result from a lack of prior formalisation. These mistakes were even identified 20 years ago (Schwarzer, Vach, and Schumacher 2000) but are provided in more detail in order to highlight their current prevalence and why they cannot be ignored.

**RM1. Incomplete reduction.** This occurs when a reduction workflow is presented as if it solves the original task but fails to do so and only the reduction strategy is solved. A common example is claiming to solve the survival task by using binary classification, e.g. erroneously claiming that a model predicts survival probabilities (which implies distribution) when it actually predicts a five year probability of death ((**box-task-classif?**)). This is a mistake as it misleads readers into believing that the model solves a survival task ((**box-task-surv?**)) when it does not. This is usually a semantic not mathematical error and results from misuse of terminology. It is important to be clear about model predict types (**?@sec-surv-set-types**) and general terms such as ‘survival predictions’ should be avoided unless they refer to one of the three prediction tasks. **RM2. Inappropriate comparisons.** This is a direct consequence of (RM1) and the two are often seen together. (RM2) occurs when an incomplete reduction is directly compared to a survival model (or complete reduction model) using a measure appropriate for the reduction. This may lead to a reduction model appearing erroneously superior. For example, comparing a logistic regression to a random survival forest (RSF)

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<sup>1</sup>A complete diagram would indicate that  $D_O$  is split into training data, which is subsequently reduced, and test data, which is passed to  $L_O$ . All reductions in this section can be applied to any data splitting process.

(Chapter 13) for predicting survival probabilities at a single time using the accuracy measure is an unfair comparison as the RSF is optimised for distribution predictions. This would be non-problematic if a suitable composition is clearly utilised. For example a regression SSVM predicting survival time cannot be directly compared to a Cox PH. However the SSVM can be compared to a CPH composed with the probabilistic to deterministic compositor, then conclusions can be drawn about comparison to the composite survival time Cox model (and not simply a Cox PH). RM3. Na"ive censoring deletion. This common mistake occurs when trying to reduce survival to regression or classification by simply deleting all censored observations, even if censoring is informative. This is a mistake as it creates bias in the dataset, which can be substantial if the proportion of censoring is high and informative. More robust deletion methods are described in Chapter 22. RM4. Oversampling uncensored observations. This is often seen when trying to reduce survival to regression or classification, and often alongside (RM3). Oversampling is the process of replicating observations to artificially inflate the sample size of the data. Whilst this process does not create any new information, it can help a model detect important features in the data. However, by only oversampling uncensored observations, this creates a source of bias in the data and ignores the potentially informative information provided by the proportion of censoring.

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## 18.4 Composition Strategies for Survival Analysis

Though composition is common practice in survival analysis, with the Cox model being a prominent example, a lack of formalisation means a lack of consensus in simple operations. For example, it is often asked in survival analysis how a model predicting a survival distribution can be used to return a survival time prediction. A common strategy is to define the survival time prediction as the median of the predicted survival curve however there is no clear reason why this should be more sensible than returning the distribution mean, mode, or some random quantile. Formalisation allow these choices to be analytically compared both theoretically and practically as hyper-parameters in a workflow. Four prediction compositions are discussed in this section (**((tab-car-taxredcar?))**), three are utilised to convert prediction types between one another, the fourth is for aggregating multiple predictions. One data composition is discussed for converting survival to regression data. Each is first graphically represented and then the components are discussed in detail. As with losses in the previous chapter, compositions are discussed at an individual observation level but extend trivially to multiple observations.

Table 18.1: Compositions formalised in Section 18.4.

ID <sup>1</sup>	Composition	Type <sup>2</sup>	Level <sup>3</sup>
C1)	Linear predictor to distribution	MOC	Prediction
C2)	Survival time to distribution	MOC	Prediction
C3)	Distribution to survival time	SOC	Prediction
C4)	Survival model averaging	MOC	Prediction
C5)	Survival to regression	SOC	Data

1. ID for reference throughout this book. 2. Composition type. Multi-object composition (MOC) or single-object composition (SOC). 3. Composition level.

### 18.4.1 C1) Linear Predictor → Distribution

This is a prediction-level MOC that composes a survival distribution from a predicted linear predictor and estimated baseline survival distribution. The composition ([?@fig-car-comp-distr](#)) requires:

- $\hat{\eta}$ : Predicted linear predictor.  $\hat{\eta}$  can be tuned by including this composition multiple times in a benchmark experiment with different models predicting  $\hat{\eta}$ . In theory any continuous ranking could be utilised instead of a linear predictor though results may be less sensible ([?@sec-car-pipelines-trade](#)).
- $\hat{S}_0$ : Estimated baseline survival function. This is usually estimated by the Kaplan-Meier estimator fit on training data,  $\hat{S}_{KM}$ . However any model that can predict a survival distribution can estimate the baseline distribution (caveat: see [?@sec-car-pipelines-trade](#)) by taking a uniform mixture of the predicted individual distributions: say  $\xi_1, \dots, \xi_m$  are  $m$  predicted distributions, then  $\hat{S}_0(\tau) = \frac{1}{m} \sum_{i=1}^m \xi_i.S(\tau)$ . The mixture is required as the baseline must be the same for all observations. Alternatively, parametric distributions can be assumed for the baseline, e.g.  $\xi = \text{Exp}(2)$  and  $\xi.S(t) = \exp(-2t)$ . As with  $\hat{\eta}$ , this parameter is also tunable.
- $M$ : Chosen model form, which theoretically can be any non-increasing right-continuous function but is usually one of:
  - Proportional Hazards (PH):  $S_{PH}(\tau|\eta, S_0) = S_0(\tau)^{\exp(\eta)}$
  - Accelerated Failure Time (AFT):  $S_{AFT}(\tau|\eta, S_0) = S_0\left(\frac{\tau}{\exp(\eta)}\right)$
  - Proportional Odds (PO):  $S_{PO}(\tau|\eta, S_0) = \frac{S_0(\tau)}{\exp(-\eta) + (1 - \exp(-\eta))S_0(\tau)}$

Models that predict linear predictors will make assumptions about the model form and therefore dictate sensible choices of  $M$ , for example the Cox model assumes a PH form. This does not mean other choices of  $M$  cannot be specified but that interpretation may be more difficult ([?@sec-car-pipelines-trade](#)). The model form can be treated as a hyper-parameter to tune. \*  $C$ : Compositor returning the composed distribution,  $\zeta := C(M, \hat{\eta}, \hat{S}_0)$  where  $\zeta$  has survival function  $\zeta.S(\tau) = M(\tau|\hat{\eta}, \hat{S}_0)$ .

Pseudo-code for training ([\(\(alg-car-comp-distr-fit?\)\)](#)) and predicting ([\(\(alg-car-comp-distr-pred?\)\)](#)) this composition as a model ‘wrapper’ with sensible parameter choices ([?@sec-car-pipelines-trade](#)) is provided in appendix ([app-car?](#)).

### 18.4.2 C2) Survival Time → Distribution

This is a prediction-level MOC that composes a distribution from a predicted survival time and assumed location-scale distribution. The composition ([?@fig-car-comp-response](#)) requires:

- $\hat{T}$ : A predicted survival time. As with the previous composition, this is tunable. In theory any continuous ranking could replace  $\hat{T}$ , though the resulting distribution may not be sensible ([?@sec-car-pipelines-trade](#)).
- $\xi$ : A specified location-scale distribution,  $\xi(\mu, \sigma)$ , e.g. Normal distribution.
- $\hat{\sigma}$ : Estimated scale parameter for the distribution. This can be treated as a hyper-parameter or predicted by another model.
- $C$ : Compositor returning the composed distribution  $\zeta := C(\xi, \hat{T}, \hat{\sigma}) = \xi(\hat{T}, \hat{\sigma})$ .

Pseudo-code for training ([\(\(alg-car-comp-response-fit?\)\)](#)) and predicting ([\(\(alg-car-comp-response-pred?\)\)](#)) this composition as a model ‘wrapper’ with sensible parameter choices

(?@sec-car-pipelines-trade) is provided in appendix (**app-car?**).

### 18.4.3 C3) Distribution → Survival Time Composition

This is a prediction-level SOC that composes a survival time from a predicted distribution. Any paper that evaluates a distribution on concordance is implicitly using this composition in some manner. Not acknowledging the composition leads to unfair model comparison (Section 18.3.3). The composition (?@fig-car-comp-crank) requires:

- $\zeta$ : A predicted survival distribution, which again is ‘tunable’.
- $\phi$ : A distribution summary method. Common examples include the mean, median and mode. Other alternatives include distribution quantiles,  $\zeta.F^{-1}(\alpha), \alpha \in [0, 1]$ ;  $\alpha$  could be tuned as a hyper-parameter.
- $C$ : Compositor returning composed survival time predictions,  $\hat{T} := C(\phi, \zeta) = \phi(\zeta)$ .

Pseudo-code for training ((**alg-car-comp-crank-fit?**)) and predicting ((**alg-car-comp-crank-pred?**)) this composition as a model ‘wrapper’ with sensible parameter choices (?@sec-car-pipelines-trade) is provided in appendix (**app-car?**).

### 18.4.4 C4) Survival Model Averaging

Ensembling is likely the most common composition in machine learning. In survival it is complicated slightly as multiple prediction types means one of two possible compositions is utilised to average predictions. The (?@fig-car-comp-avg) composition requires:

- $\rho = \rho_1, \dots, \rho_B$ :  $B$  predictions (not necessarily from the same model) of the same type: ranking, survival time or distribution; again ‘tunable’.
- $w = w_1, \dots, w_B$ : Weights that sum to one.
- $C$ : Compositor returning combined predictions,  $\hat{\rho} := C(\rho, w)$  where  $C(\rho, w) = \frac{1}{B} \sum_{i=1}^B w_i \rho_i$ , if  $\rho$  are ranking of survival time predictions; or  $C(\rho, w) = \zeta$  where  $\zeta$  is the distribution defined by the survival function  $\zeta.S(\tau) = \frac{1}{B} \sum_{i=1}^B w_i \rho_i.S(\tau)$ , if  $\rho$  are distribution predictions.

Pseudo-code for training ((**alg-car-comp-avg-fit?**)) and predicting ((**alg-car-comp-avg-pred?**)) this composition as a model ‘wrapper’ with sensible parameter choices (?@sec-car-pipelines-trade) is provided in appendix (**app-car?**).

## 18.5 Novel Survival Reductions

This section collects the various strategies and settings discussed previously into complete reduction workflows. (**tab-car-reduces?**) lists the reductions discussed in this section with IDs for future reference. All strategies are described by visualising a graphical pipeline and then listing the composition steps required in fitting and predicting.

This section only includes novel reduction strategies and does not provide a survey of pre-existing strategies. This limitation is primarily due to time (and page) constraints as every method has very distinct workflows that require complex exposition. Well-established strategies are briefly mentioned below and future research is planned to survey and compare all strategies with respect to empirical performance (i.e. in benchmark experiments).

Two prominent reductions are ‘landmarking’ (Van Houwelingen 2007) and piecewise expo-

nential models (M. Friedman 1982). Both are reductions for time-varying covariates and hence outside the scope of this book. Relevant to this book scope is a large class of strategies that utilise ‘discrete time survival analysis’ (Tutz and Schmid 2016); these strategies include reductions (R7) and (R8). Methodology for discrete time survival analysis has been seen in the literature for the past three decades (Liestol, Andersen, and Andersen 1994). The primary reduction strategy for discrete time survival analysis is implemented in the R package **discSurv**. (Welchowski and Schmid 2019); this is very similar to (R7) except that it enforces stricter constraints in the composition procedures and forces a ‘discrete-hazard’ instead of ‘discrete-survival’ representation (Section 18.5.1.2).

### 18.5.1 R7-R8) Survival → Probabilistic Classification

Two separate reductions are presented in ?@fig-car-R7R8 however as both are reductions to probabilistic classification and are only different in the very last step, both are presented in this section. Steps and compositions of the reduction (?@fig-car-R7R8):

**Fit F1)** A survival dataset,  $\mathcal{D}_{train}$ , is binned,  $B$ , with a continuous to discrete data composition (Section 18.5.1.1). **F2)** A multi-label classification model, with adaptations for censoring,  $g_L(D_B|\theta)$ , is fit on the transformed dataset,  $D_B$ . Optionally,  $g_L$  could be further reduced to binary,  $g_B$ , or multi-class classification,  $g_c$ , (Section 18.5.1.4). **Predict P1)** Testing survival data,  $\mathcal{D}_{test}$ , is passed to the trained classification model,  $\hat{g}$ , to predict pseudo-survival probabilities  $\hat{S}$  (or optionally hazards (Section 18.5.1.2)). **P2a)** Predictions can be composed,  $T_1$ , into a survival distribution prediction,  $\zeta = \zeta_1, \dots, \zeta_m$  (Section 18.5.1.6); or, **P2b)** Predictions can be composed,  $T_2$ , to survival time predictions,  $\hat{T} = \hat{T}_1, \dots, \hat{T}_m$  (Section 18.5.1.7).

Further details for binning, multi-label classification, and transformation of pseudo-survival probabilities are now provided.

#### 18.5.1.1 Composition: Binning Survival Times

An essential part of the reduction is the transformation from a survival dataset to a classification dataset, which requires two separate compositions. The first (discussed here) is to discretise the survival times ( $B(\mathcal{D}_{train}|w)$  in ?@fig-car-R7R8) and the second is to merge the survival time and censoring indicator into a single outcome (Section 18.5.1.2).

Discretising survival times is achieved by the common ‘binning’ composition, in which a continuous outcome is discretised into ‘bins’ according to specified thresholds. These thresholds are usually determined by specifying the width of the bins as a hyper-parameter  $w$ .<sup>2</sup> This is a common transformation and therefore further discussion is not provided here. An example is given below with the original survival data on the left and the binned data on the right ( $w = 1$ ).

X	Time (Cont.)	Died
1	1.56	0
2	2	1
3	3.3	1
4	3.6	0
5	4	0

<sup>2</sup>Binning is described here with equal widths but generalises to unequal widths trivially.

X	Time (Disc.)	Died
1	[1, 2)	0
2	[2, 3)	1
3	[3, 4)	1
4	[3, 4)	0
5	[4, 5)	0

### 18.5.1.2 Composition: Survival to Classification Outcome

The binned dataset still has the unique survival data format of utilising two outcomes for training (time and status) but only making a prediction for one outcome (distribution). In order for this to be compatible with classification, the two outcome variables are composed into a single variable.<sup>3</sup> This is achieved by casting the survival times into a ‘wide’ format and creating a new outcome indicator.<sup>4</sup> Two outcome transformations are possible, the first represents a discrete survival function and the second represents a discrete hazard function.<sup>5</sup>

#### Discrete Survival Function Composition

In this composition, the data in the transformed dataset represents the discrete survival function. The new indicator is defined as follows,

$$Y_{i;\tau} := \begin{cases} 1, & T_i > \tau \\ 0, & T_i \leq \tau \cap \Delta_i = 1 \\ -1, & T_i \leq \tau \cap \Delta_i = 0 \end{cases}$$

At a given discrete time  $\tau$ , an observation,  $i$ , is either alive ( $Y_{i;\tau} = 1$ ), dead ( $Y_{i;\tau} = 0$ ), or censored ( $Y_{i;\tau} = -1$ ). Therefore  $\hat{P}(Y_{i;\tau} = 1) = \hat{S}_i(\tau)$ , motivating this particular choice of representation.

This composition is demonstrated below with the binned data (left) and the composed classification data (right).

X	Time (Disc.)	Died
1	[1, 2)	0
2	[2, 3)	1
3	[3, 4)	1
4	[3, 4)	0
5	[4, 5)	0

X	[1,2)	[2,3)	[3,4)	[4,5)
1	-1	-1	-1	-1
2	1	0	0	0

<sup>3</sup>This is the first key divergence from other discrete-time classification strategies, which use the censoring indicator as the outcome and the time outcome as a feature.

<sup>4</sup>This is the second key divergence from other discrete-time classification strategies, which keep the data in a ‘long’ format.

<sup>5</sup>This is the final key divergence from other discrete-time classification strategies, which enforce the discrete hazard representation.

X	[1,2)	[2,3)	[3,4)	[4,5)
3	1	1	0	0
4	1	1	-1	-1
5	1	1	-1	-1

### Discrete Hazard Function Composition

In this composition, the data in the transformed dataset represents the discrete hazard function. The new indicator is defined as follows,

$$Y_{i;\tau}^* := \begin{cases} 1, & T_i = \tau \cap \Delta_i = 1 \\ -1, & T_i = \tau \cap \Delta_i = 0 \\ 0, & \text{otherwise} \end{cases}$$

At a given discrete time  $\tau$ , an observation,  $i$ , either experiences the event ( $Y_{i;\tau}^* = 1$ ), experiences censoring ( $Y_{i;\tau}^* = -1$ ), or neither ( $Y_{i;\tau}^* = 0$ ). Utilising sequential multi-label classification problem transformation methods (Section 18.5.1.4) results in  $\hat{P}(Y_{i;\tau}^* = 1) = \hat{h}_i(\tau)$ . If methods are utilised that do not ‘look back’ at predictions then  $\hat{P}(Y_{i;\tau}^* = 1) = \hat{p}_i(\tau)$  (Section 18.5.1.4).<sup>6</sup>

This composition is demonstrated below with the binned data (left) and the composed classification data (right).

X	Time (Disc.)	Died
1	[1, 2)	0
2	[2, 3)	1
3	[3, 4)	1
4	[3, 4)	0
5	[4, 5)	0

X	[1,2)	[2,3)	[3,4)	[4,5)
1	-1	0	0	0
2	0	1	0	0
3	0	0	1	0
4	0	0	-1	0
5	0	0	0	-1

### Multi-Label Classification Data

In both compositions, survival data t.v.i.  $\mathbb{R}^p \times \mathbb{R}_{\geq 0} \times \{0, 1\}$  is transformed to multi-label classification data t.v.i.  $\mathbb{R}^p \times \{-1, 0, 1\}^K$  for  $K$  binned time-intervals. The multi-label classification task is defined in Section 18.5.1.4 with possible algorithms.

The discrete survival representation has a slightly more natural interpretation and is ‘easier’ for classifiers to use for training as there are more positive events (i.e. more observations

<sup>6</sup>This important distinction is not required in other discrete-time reduction strategies that automatically condition the prediction by including time as a feature.

alive) to train on, whereas the discrete hazard representation will have relatively few events in each time-point. However the hazard representation leads to more natural predictions (Section 18.5.1.6).

A particular bias that may easily result from the composition of survival to classification data is now discussed.

#### 18.5.1.3 Reduction to Classification Bias

The reduction to classification bias is commonly known (Zhou et al. 2005) but is reiterated briefly here as it must be accounted for in any automated reduction to classification workflow. This bias occurs when making classification predictions about survival at a given time and incorrectly censoring patients who have not been observed long enough, instead of removing them.

By example, say the prediction of interest is five-year survival probabilities after a particular diagnosis, clearly a patient who has only been diagnosed for three years cannot inform this prediction. The bias is introduced if this patient is censored at five-years instead of being removed from the dataset. The result of this bias is to artificially inflate the probability of survival at each time-point as an unknown outcome is treated as censored and therefore alive.

This bias is simply dealt with by removing patients who have not been alive ‘long enough’.<sup>7</sup> Paradoxically, even if a patient is observed to die before the time-point of interest, they should still be removed if they have not been in the dataset ‘long enough’ as failing to do so will result in a bias in the opposite direction, thus over-inflating the proportion of dead observations.

Accounting for this bias is particularly important in the multi-label reduction as the number of observable patients will decrease over time due to censoring.

#### 18.5.1.4 Multi-Label Classification Algorithms

As the work in this section is completely out of the book scope, the full text is in appendix ([app-mlc?](#)). The most important contributions from this section are:

- Reviewing problem transformation methods (Tsoumakas and Katakis 2007) for multi-label classification;
- Identifying that only binary relevance, nested stacking, and classifier chains are appropriate in this reduction; and
- Generalising these methods into a single wrapper for any binary classifier, the ‘LWrapper’.

#### 18.5.1.5 Censoring in Classification

Classification algorithms cannot natively handle the censoring that is included in the survival reduction, but this can be incorporated using one of two approaches.

#### Multi-Class Classification

All multi-label datasets can also handle multi-class data, hence the simplest way in which to handle censoring is to make multi-class predictions in each label for the outcome  $Y_\tau t.v.i.\{-1, 0, 1\}$ . Many off-shelf classification learners can make multi-class predictions natively and simple reductions exist for those that cannot. As a disadvantage to this method,

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<sup>7</sup>Accounting for this bias is only possible if the study start and end dates are known, as well as the date the patient entered the study.

classifiers would then predict if an individual is dead or alive or censored (each mutually exclusive), and not simply alive or dead. Though this could be perceived as an advantage when censoring is informative as this will accurately reflect a real-world competing-risks set-up.

### Subsetting/Hurdle Models

For this approach, the multi-class task is reduced to two binary class tasks: first predict if a subject is censored or not (dead or alive) and only if the prediction for censoring is below some threshold,  $\alpha \in [0, 1]$ , then predict if the subject is alive or not (dead or censored). If the probability of censoring is high in the first task then the probability of being alive is automatically set to zero in the final prediction, otherwise the prediction from the second task is used. Any classifier can utilise this approach and it has a meaningful interpretation, additionally  $\alpha$  is a tunable hyper-parameter. The main disadvantage is increases to storage and run-time requirements as double the number of models may be fit.

Once the datasets have been composed to classification datasets and censoring is suitably incorporated by either approach, then any probabilistic classification model can be fit on the data. Predictions from these models can either be composed to a distribution prediction (R7) or a survival time prediction (R8).

#### 18.5.1.6 R7) Probabilistic Survival → Probabilistic Classification

This final part of the (R7) reduction is described separately for discrete hazard and survival representations of the data (Section 18.5.1.2).

#### Discrete Hazard Representation

In this representation recall that predictions of the positive class,  $P(Y_\tau = 1)$ , are estimating the quantity  $h(\tau)$ . These predictions provide a natural and efficient transformation from predicted hazards to survival probabilities. Let  $\hat{h}_i$  be a predicted hazard function for some observation  $i$ , then the survival function for that observation can be found with a Kaplan-Meier type estimator,

$$\tilde{S}_i(\tau^*) = \prod_{\tau} 1 - \hat{h}_i(\tau)$$

Now predictions are for a pseudo-survival function, which is ‘pseudo’ as it is not right-continuous. Resolving this is discussed below.

#### Discrete Survival Representation

In this representation,  $P(Y_\tau = 1)$  is estimating  $S(\tau)$ , which means that predictions from a classification model result in discrete point predictions and not a right-continuous function. More importantly, there is no guarantee that a non-increasing function will be predicted, i.e. there is no guarantee that  $P(Y_j = 1) < P(Y_i = 1)$ , for time-points  $j > i$ .

Unfortunately there is no optimal way of dealing with predictions of this sort and ‘mistakes’ of this kind have been observed in some software implementation. One point to note is that in practice these are quite rare as the probability of survival will always decrease over time. Therefore the ‘usual’ approach is quite ‘hacky’ and involves imputing increasing predictions with the previous prediction, formally,

$$\tilde{S}(i+1) := \min\{P(Y_{i+1} = 1), P(Y_i = 1)\}, \forall i = \mathbb{R}_{\geq 0}$$

assuming  $\tilde{S}(0) = 1$ . Future research should seek more robust alternatives.

## Right-Continuous Survival Function

From either representation, a \ non-increasing but non-continuous pseudo-survival function,  $\tilde{S}$ , is now predicted. Creating a right-continuous function (' $T_1(\tilde{S})$ ' in `?@fig-car-R7`) from these point predictions (Figure 18.3 (a)) is relatively simple and well-known with accessible off-shelf software. At the very least, one can assume a constant hazard rate between predictions and cast them into a step function (Figure 18.3 (b)). This is a fairly common assumption and is usually valid as bin-width decreases. Alternatively, the point predictions can be smoothed into a continuous function with off-shelf software, for example with polynomial local regression smoothing (Figure 18.3 (c)) or generalised linear smoothing (Figure 18.3 (d)). Whichever method is chosen, the survival function is now non-increasing right-continuous and the (R7) reduction is complete.

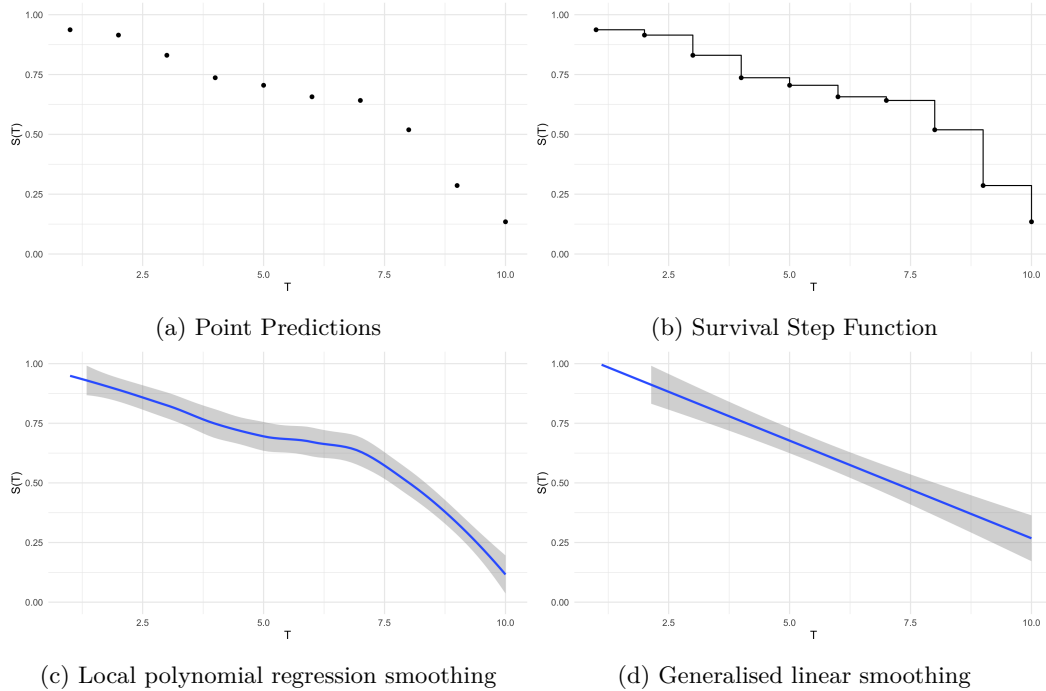


Figure 18.3: Survival function as a: point prediction (a), step function assuming constant risk (b), local polynomial regression smoothing (c), and generalised linear smoothing (d). (c) and (d) computed with **ggplot2** (Wickham 2016).

### 18.5.1.7 R8) Deterministic Survival → Probabilistic Classification

Predicting a deterministic survival time from the multi-label classification predictions is relatively straightforward and can be viewed as a discrete analogue to (C3) (Section 18.4.3). For the discrete hazard representation, one can simply take the predicted time-point for an individual to be time at which the predicted hazard probability is highest however this could easily be problematic as there may be multiple time-points at which the predicted hazard equals 1. Instead it is cleaner to first cast the hazard to a pseudo-survival probability (Section 18.5.1.6) and then treat both representations the same.

Let  $\tilde{S}_i$  be the predicted multi-label survival probabilities for an observation  $i$  such that  $\tilde{S}_i(\tau)$  corresponds with  $\hat{P}(Y_{i;\tau} = 1)$  for label  $\tau \in \mathcal{K}$  where  $Y_{i;\tau}$  is defined in Section 18.5.1.2 and

$\mathcal{K} = \{1, \dots, K\}$  is the set of labels for which to make predictions. Then the survival time transformation is defined by

$$T_2(\tilde{S}_i) = \inf\{\tau \in \mathcal{K} : \tilde{S}_i(\tau) \leq \beta\}$$

for some  $\beta \in [0, 1]$ .

This is interpreted as defining the predicted survival time as the first time-point in which the predicted probability of being alive drops below a certain threshold  $\beta$ . Usually  $\beta = 0.5$ , though this can be treated as a hyper-parameter for tuning. This composition can be utilised even if predictions are not non-increasing, as only the first time the predicted survival probability drops below the threshold is considered. With this composition the (R8) reduction is now complete.

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## 18.6 Conclusions

This chapter introduced composition and reduction to survival analysis and formalised specific strategies. Formalising these concepts allows for better quality of research and most importantly improved transparency. Clear interface points for hyper-parameters and compositions allow for reproducibility that was previously obfuscated by unclear workflows and imprecise documentation for pipelines.

Additionally, composition and reduction improves accessibility. Reduction workflows vastly increase the number of machine learning models that can be utilised in survival analysis, thus opening the field to those whose experience is limited to regression or classification. Formalisation of workflows allows for precise implementation of model-agnostic pipelines as computational objects, as opposed to functions that are built directly into an algorithm without external interface points.

Finally, predictive performance is also increased by these methods, which is most prominently the case for the survival model averaging compositor (as demonstrated by RSFs).



# 19

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## Competing Risks Pipelines

TODO (150-200 WORDS)

! Page coming soon!

We are working on this page and it will be available soon!



# 20

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## *Discrete Time Survival Analysis*

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TODO (150-200 WORDS)

! Page coming soon!

We are working on this page and it will be available soon!



# 21

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## *Connections to Poisson Regression and Processes*

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TODO (150-200 WORDS)

! Page coming soon!

We are working on this page and it will be available soon!



## 22

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# *Connections to Regression and Imputation*

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TODO (150-200 WORDS)

! Major changes expected!

This page is a work in progress and major changes will be made over time.

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TODO

- I think all these sections should have examples in implemented models, e.g., here we can point to SVM models and some neural nets
  - We can also point to neural nets that use reduction to essentially just predict the linear predictor via regression or to use pseudovalues
  - Add pseudovalues
  - Add prediction of the observed outcome (not survival) time
- 

This is a data-level SOC that transforms survival data to regression data by either removing censored observations or ‘imputing’ survival times. This composition is frequently incorrectly utilised (Section 18.3.3) and therefore more detail is provided here than previous compositions. Note that the previous compositions were prediction-level transformations that occur after a survival model makes a prediction, whereas this composition is on a data-level and can take place before model training or predicting.

In Statistics, there are only two methods for removing ‘missing’ values: deletion and imputation; both of these have been attempted for censoring.

Censoring can be beneficial, harmful, or neutral; each will affect the data differently if deleted or imputed. Harmful censoring occurs if the reason for censoring is negative, for example drop-out due to disease progression. Harmful censoring indicates that the true survival time is likely soon after the censoring time. Beneficial censoring occurs if censoring is positive, for example drop-out due to recovery. This indicates that the true survival time is likely far from the censoring time. Finally neutral censoring occurs when no information can be gained about the true survival time from the censoring time. Whilst the first two of these can be considered to be dependent on the outcome, neutral censoring is often the case when censoring is independent of the outcome conditional on the data, which is a standard assumption for the majority of survival models and measures.

### 22.0.0.1 Deletion # {sec-redux-regr-del}

Deletion is the process of removing observations from a dataset. This is usually seen in ‘complete case analysis’ in which observations with ‘missingness’, covariates with missing values, are removed from the dataset. In survival analysis this method is somewhat riskier as the subjects to delete depend on the outcome and not the features. Three methods are considered, the first two are a more brute-force approach whereas the third allows for some flexibility and tuning.

#### Complete Deletion

Deleting all censored observations is simple to implement with no computational overhead. Complete deletion results in a smaller regression dataset, which may be significantly smaller if the proportion of censoring is high. If censoring is uninformative, the dataset is suitably large and the proportion of censoring suitably low, then this method can be applied without further consideration. However if censoring is informative then deletion will add bias to the dataset, although the ‘direction’ of bias cannot be known in advance. If censoring is harmful then censored observations will likely have a similar profile to those that died, thus removing censoring will artificially inflate the proportion of those who survive. Conversely if censoring is beneficial then censored observations may be more similar to those who survive, thus removal will artificially inflate the proportion of those who die.

#### Omission

Omission is the process of omitting the censoring indicator from the dataset, thus resulting in a regression dataset that assumes all observations experienced the event. Complete deletion results in a smaller dataset of dead patients, omission results in no sample size reduction but the outcome may be incorrect. This reduction strategy is likely only justified for harmful censoring. In this case the true survival time is likely close to the censoring time and therefore treating censored observations as dead may be a fair assumption.

#### IPCW

If censoring is conditionally-outcome independent then deletion of censored events is possible by using Inverse Probability of Censoring Weights (IPCW). This method has been seen several times throughout this book in the context of models and measures. It has been formalised as a composition technique by Vock *et al.* (2016) (Vock et al. 2016) although their method is limited to binary classification. Their method weights the survival time of uncensored observations by  $w_i = 1/\hat{G}_{KM}(T_i)$  and deletes censored observations, where  $\hat{G}_{KM}$  is the Kaplan-Meier estimate of the censoring distribution fit on training data. As previously discussed, one could instead consider the Akritas (or any other) estimator for  $\hat{G}_{KM}$ .

Whilst this method does provide a ‘safer’ way to delete censored observations, there is not a necessity to do so. Instead consider the following weights

$$w_i = \frac{\Delta_i + \alpha(1 - \Delta_i)}{\hat{G}_{KM}(T_i)} \quad (22.1)$$

where  $\alpha \in [0, 1]$  is a hyper-parameter to tune. Setting  $\alpha = 1$  equally weights censored and uncensored observations and setting  $\alpha = 0$  recovers the setting in which censored observations are deleted. It is assumed  $\hat{G}_{KM}$  is set to some very small  $\epsilon$  when  $\hat{G}_{KM}(T_i) = 0$ . When  $\alpha \neq 0$  this becomes an imputation method, other imputation methods are now discussed.

### 22.0.0.2 Imputation

Imputation methods estimate the values of missing data conditional on non-missing data and other covariates. Whilst the true value of the missing data can never be known, by carefully conditioning on the ‘correct’ covariates, good estimates for the missing value can be obtained to help prevent a loss of data. Imputing outcome data is more difficult than imputing covariate data as models are then trained on ‘fake’ data. However a poor imputation should still be clear when evaluating a model as testing data remains un-imputed. By imputing censoring times with estimated survival times, the censoring indicator can be removed and the dataset becomes a regression dataset.

#### Gamma Imputation

Gamma imputation (D. Jackson et al. 2014) incorporates information about whether censoring is harmful, beneficial, or neutral. The method imputes survival times by generating times from a shifted proportional hazards model

$$h(\tau) = h_0(\tau) \exp(\eta + \gamma)$$

where  $\eta$  is the usual linear predictor and  $\gamma \in \mathbb{R}$  is a hyper-parameter determining the ‘type’ of censoring such that  $\gamma > 0$  indicates harmful censoring,  $\gamma < 0$  indicates beneficial censoring, and  $\gamma = 0$  is neutral censoring. This imputation method has the benefit of being tunable as  $\gamma$  is a hyper-parameter and there is a choice of variables to condition the imputation. No independent experiments exist studying how well this method performs, nor discussing the theoretical properties of the method.

#### MRL

The Mean Residual Lifetime (MRL) estimator has been previously discussed in the context of SVMs (Section 14.2). Here the estimator is extended to serve as an imputation method. Recall the MRL function,  $MRL(\tau|\hat{S}) = \int_{\tau}^{\infty} \hat{S}(u) du / \hat{S}(\tau)$ , where  $\hat{S}$  is an estimate of the survival function of the underlying survival distribution (e.g.  $\hat{S}_{KM}$ ). The MRL is interpreted as the expected remaining survival time after the time-point  $\tau$ . This serves as a natural imputation strategy where given the survival outcome  $(T_i, \Delta_i)$ , the new imputed time  $T'_i$  is given by

$$T'_i = T_i + (1 - \Delta_i) MRL(T_i | \hat{S})$$

where  $\hat{S}$  would be fit on the training data and could be an unconditional estimator, such as Kaplan-Meier, or conditional, such as Akritas. The resulting survival times are interpreted as the true times for those who died and the expected survival times for those who were censored.

#### Buckley-James

Buckley-James (Buckley and James 1979) is another imputation method discussed earlier (`?@sec-surv-ml-models-boost`). The Buckley-James method uses an iterative procedure to impute censored survival times by the conditional expectation given censoring times and covariates (Z. Wang and Wang 2010). Given the survival tuple for an outcome  $(T_i, \Delta_i)$ , the new imputed time  $T'_i$  is

$$T'_i = \begin{cases} T_i, & \Delta_i = 1 \\ X_i \hat{\beta} + \frac{1}{\hat{S}_{KM}(e_i)} \sum_{e_k < e_i} \hat{p}_{KM}(e_k) e_k & \Delta_i = 0 \end{cases}$$

where  $\hat{S}_{KM}$  is the Kaplan-Meier estimator of the survival distribution estimated on training data and with associated pmf  $\hat{p}_{KM}$  and  $e_i = T_i - X_i\hat{\beta}$  where  $\hat{\beta}$  are estimated coefficients of a linear regression model fit on  $(X_i, T_i)$ . Given the least squares approach, more parametric assumptions are made than other imputation methods and it is more complex to separate model fitting from imputation. Hence, this imputation may only be appropriate on a limited number of data types.

### Alternative Methods

Other methods have been proposed for ‘imputing’ censored survival times though with either less clear discussion or to no benefit. Multiple imputation by chained equations (MICE) has been demonstrated to perform well with covariate data and even outcome data (in a non-survival setting). However no adaptations have been developed to incorporate censoring times into the imputation and therefore is less informative than Gamma imputation.

Re-calibration of censored survival times (Vinzamuri, Li, and Reddy 2017) uses an iterative update procedure to ‘re-calibrate’ censoring times however the motivation behind the method is not sufficiently clear to be of interest in general survival modelling tasks outside of the authors’ specific pipelines.

Finally parametric imputation is defined by making random draws from truncated probability distributions and adding these to the censoring time (P. Royston 2001; Patrick Royston, Parmar, and Altman 2008). Whilst this method is arguably the simplest method and will lead to a sufficiently random sample, i.e. not one skewed by the imputation process, in practice the randomness leads to unrealistic results, with some imputed times being very far from the original censoring times and some being very close.

#### 22.0.0.3 The Decision to Impute or Delete

Deletion methods are simple to implement and fast to compute however they can lead to biasing the data or a significant sample reduction if used incorrectly. Imputation methods can incorporate tuning and have more relaxed assumptions about the censoring mechanism, though they may lead to over-confidence in the resulting outcome and therefore add bias into the dataset. In some cases, the decision to impute or delete is straightforward, for example if censoring is uninformative and only few observations are censored then complete deletion is appropriate. If it is unknown if censoring is informative then this can crudely be estimated by a benchmark experiment. Classification models can be fit on  $\{(X_1, \Delta_1), \dots, (X_n, \Delta_n)\}$  where  $(X_i, \Delta_i) \in \mathcal{D}_{train}$ . Whilst not an exact test, if any model significantly outperforms a baseline, then this may indicate censoring is informative. This is demonstrated in (**tab-car-predcens?**), in which a logistic regression outperforms a featureless baseline in correctly predicting if an observation is censored when censoring is informative, but is no better than the baseline when censoring is uninformative.

Table 22.1: Estimating censoring dependence by prediction. **Sim1** is informative censoring and **Sim7** is uninformative. Logistic regression is compared to a featureless baseline with the Brier score with standard errors. Censoring can be significantly predicted to 95% confidence when informative (**Sim1**) but not when uninformative (**Sim7**).

Data	Baseline	Logistic Regression
<b>Sim1</b>	0.20 (0.14, 0.26)	0.02 (0.01, 0.03)
<b>Sim7</b>	0.19 (0.14, 0.24)	0.16 (0.13, 0.19)

# 23

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## Conclusions

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### 23.1 Common problems in survival analysis

#### 23.1.1 Data cleaning

##### Events at $t=0$

Throughout this book we have defined survival times taking values in the non-negative Reals (zero inclusive)  $\mathbb{R}_{\geq 0}$ . In practice, model implementations assume time is over the positive Reals (zero exclusive). One must therefore consider how to deal with subjects that experience the outcome at 0. There is no established best practice for dealing with this case as the answer may be data-dependent. Possible choices include:

1. Deleting all data where the outcome occurs at  $t = 0$ , this may be appropriate if it only happens in a small number of observations and therefore deletion is unlikely to bias predictions;
2. Update the survival time to the next smallest observed survival time. For example, if the first observation to experience the event after  $t = 0$  happens at  $t = 0.1$ , then set 0.1 as the survival time for any observation experiencing the event at  $t = 0$ . Note this method will not be appropriate when data is over a long period, for example if measuring time over years, then there could be a substantial difference between  $t = 0$  and  $t = 1$ ;
3. Update the survival time to a very small value  $\epsilon$  that makes sense given the context of the data, e.g.,  $\epsilon = 0.0001$ .

##### Continuous v Discrete Time

We defined survival tasks throughout this book assuming continuous time predictions in  $\mathbb{R}_{\geq 0}$ . In practice, many outcomes in survival analysis are recorded on a discrete scale, such as in medical statistics where outcomes are observed on a yearly, daily, monthly, hourly, etc. basis. Whilst discrete-time survival analysis exists for this purpose (Chapter 20), software implementations overwhelming use theory from the 'continuous-time setting. There has not been a lot of research into whether discrete-time methods outperform continuous-time methods when correctly applied to discrete data, however available experiments do not indicate that discrete methods outperform their continuous counterparts (Suresh, Severn,

and Ghosh 2022). Therefore it is recommended to use available software implementations, even when data is recorded on a discrete scale.

### **23.1.2 Evaluation and prediction**

- Which time points to make predictions for?
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## **23.2 What's next for MLSA?**

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## Exercises

TODO (150-200 WORDS)

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