

# Survival Mixture Density Networks

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## Abstract

Survival analysis, the art of time-to-event modeling, plays an important role in clinical treatment decisions. Recently, continuous time models built from neural ODEs have been proposed for survival analysis. However, the training of neural ODEs is slow due to the high computational complexity of neural ODE solvers. Here, we propose an efficient alternative for flexible continuous time models, called Survival Mixture Density Networks (Survival MDNs). Survival MDN applies an invertible positive function to the output of Mixture Density Networks (MDNs). While MDNs produce flexible real-valued distributions, the invertible positive function maps the model into the time-domain while preserving a tractable density. Using four datasets, we show that Survival MDN performs better than, or similarly to continuous and discrete time baselines on concordance, integrated Brier score and integrated binomial log-likelihood. Meanwhile, Survival MDNs are also faster than ODE-based models and circumvent binning issues in discrete models.

## 1. Introduction

Survival analysis serves as an important tool in healthcare to assess the risk of events, such as onset of disease (Wilson et al., 1998) or death (Pocock et al., 1982), rehospitalization (Patterson and Lee, 1998) and discharge from hospital (Wang et al., 2020). Survival modeling has been widely used in clinical applications, including improving the prognosis of cancer (Faradmal et al., 2012; Goldstraw et al., 2016; Wang et al., 2019; Lin and Anisa, 2021; Wang et al., 2021), predicting the onset of septic shock (Henry et al., 2015), assessing the survival time of heart failure patients (Ahmad et al., 2017; Kojoria et al., 2004; Jones et al., 2019; Yin et al., 2022) and estimating the graft survival rate of kidney transplant patients (Lee et al., 2019; Rodrigues et al., 2019).

Given patients’ electronic health records including lab tests, vitals, radiology results and clinical notes, doctors need to determine the level of treatment based on the level of risk. For example, WHO guidelines suggest more aggressive treatments for higher risk cardiovascular disease patients (WHO et al., 2007). Therefore, an accurate model of risk is necessary.

Risk in survival analysis is characterized by the conditional distribution of the event time given a patient’s healthcare records. What distinguishes survival analysis from traditional regression problems is that event times can be censored, i.e., only known to lie within a certain range. For example, patients may remain healthy throughout a 10-year

coronary artery disease study (Wilson et al., 1998) so it is only known that such patients survive at least 10 years. Discarding censored times may introduce bias into estimates by underestimating the time until an event, because later times are more likely to be censored and thus thrown away.

Likelihood-based methods are used to estimate survival models (Kalbfleisch and Prentice, 2011). In addition to the usual mass or density computed in maximum likelihood problems, the survival likelihood for censored data includes the survival function, i.e., one minus the cumulative distribution function (CDF) of the distribution. For many distributions, CDF evaluations require explicitly integrating the density. Recent advances in deep learning provide opportunities for flexible survival modeling (LeCun et al., 2015; Ranganath et al., 2016). However, flexible distributions utilizing deep learning, such as those modeled by GANs (Goodfellow et al., 2014; Chapfuwa et al., 2018), may not yield efficient CDF computation.

Traditional survival analysis techniques make distributional assumptions, e.g. log-normal density or proportional hazards, to keep estimation tractable (Kalbfleisch and Prentice, 2011; Cox, 1972). But this limits the flexibility of the model. To move beyond this, discrete time models divide continuous times into a sequence of bins (Miscouridou et al., 2018; Lee et al., 2018; Kvamme and Borgan, 2019) and can approximate arbitrary continuous distributions increasing well as the number of bins increases. However, the choice of bin boundaries is troublesome: it is unclear how best to set the time intervals for each bin, and the survival function for times within a bin is ill-defined. ODE-based continuous time models (Tang et al., 2020, 2022) specify the time-to-event distribution through ODEs. However, the training of ODE-based models is slow due to expensive numerical integration requiring many neural network evaluations for each forward pass (Kelly et al., 2020).

In this work, we propose Survival Mixture Density Networks (Survival MDN). Survival MDN builds off mixture density networks (Bishop, 1994) to allow flexible modeling. Since the time is positive in survival modeling, we apply an invertible positive function to the samples from MDNs. The CDF of Survival MDNs can be obtained easily through the evaluation of the CDF of the mixture components of the MDN, which is simple for mixture components like Gaussians. We evaluate Survival MDN and baselines on four clinical datasets: SUPPORT, METABRIC, GBSG, and MIMIC. On all datasets, Survival MDN performs better than, or as well as, the baselines on concordance, integrated Brier Score and integrated binomial log-likelihood. We also show that training Survival MDNs can be 100 times faster than the ODE-based model SODEN (Tang et al., 2020).<sup>1</sup>

### Generalizable Insights about Machine Learning in the Context of Healthcare

The majority of flexible survival modeling relies on training with the Cox partial likelihood, discrete time modeling, or ordinary differential equations. Training with partial likelihood disables the stochastic gradient descent algorithm and is not scalable for large datasets. Discrete time models have issues with choosing bin boundaries and determining the survival probability for a particular time. ODE-based models use likelihood for training but are slow to train. Our proposed model Survival MDN have several advantages 1) It is a continuous time model 2) It makes fewer distributional assumptions 3) It can be trained with stochastic

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1. The code is available at <https://github.com/XintianHan/Survival-MDN>

gradient descent 4) It is easier to use than discrete models and faster than ODE-based models.

## 2. Background

In this section, we introduce the mathematical foundation of survival analysis and summarize related works. We then describe how our work is distinguished from previous works.

### 2.1. Foundation of Survival Analysis

Survival analysis studies the distribution of event time  $T$  given covariates  $X$ . For example, we would like to know when a patient may die after the admission to ICU. The event time is called the failure time or survival time. We consider the common scenario of *right-censoring* in this work, where only a lower bound of the survival time is observed for some of the patients. We call the lower bound the *censored time*  $C$ . When  $T > C$ , only the censored time  $C$  is observed; when  $T \leq C$ , the failure time  $T$  is observed. We use  $\Delta = I\{T \leq C\}$  to indicate whether the event time is observed and  $U = \min\{T, C\}$  to denote the observed time.

A central quantity that appears in the estimation and use of survival models is the survival function  $S(t|X) = P(T > t|X)$ , i.e., the probability a patient with covariates  $X$  will survival until time  $t$ . By definition,  $S(t|X) = 1 - \text{CDF}(t|X)$ .

Assume we observe i.i.d. datapoints  $\{u_i, \Delta_i, x_i\}_{i=1}^N$  and censoring is random  $T \perp C|X$ . Under these assumptions, and with  $p(t|X)$  denoting the mass or probability density function (PDF) evaluated at  $t$ , the survival likelihood function with a parameter  $\theta$  is proportional to (Kalbfleisch and Prentice, 2011):

$$\prod_{i=1}^N p_{\theta}(u_i|x_i)^{\Delta_i} S_{\theta}(u_i|x_i)^{1-\Delta_i}.$$

In this work, we use the log-likelihood as a training objective function.

### 2.2. Related Work

**Traditional Survival Analysis** Traditionally, survival analysis makes distributional assumptions. The Cox model (Cox, 1972) makes the proportional hazard assumption. The accelerated failure time (AFT) model (Buckley and James, 1979; Wei, 1992) assumes that  $\log(T) = X^T\theta + \epsilon$ , where  $\epsilon$  is a log-logistic distribution. Multiple variants of Cox and AFT models (Aalen, 1980; Bennett, 1983; Cheng et al., 1995; Lin and Ying, 1995; Kalbfleisch and Prentice, 2011; Wu and Witten, 2019) have been proposed to introduce time-varying functions or different distributions. However, these extensions only use linear or simple nonlinear models which may not be flexible enough to model the complex data distribution. Avati et al. (2020) use deep networks to produce the parameters of a lognormal. Though this can capture nonlinear dependence of the lognormal’s parameters on the input, the lognormal assumption may not be appropriate, e.g., if the true distribution has more than one mode.

**Deep Cox Models** The Cox model has been extended with deep networks in several ways. DeepSurv (Katzman et al., 2018) uses a neural network to model the relative risk  $g(X; \theta)$ . Cox-Time (Kvamme et al., 2019) further allows the relative risk to depend on time  $t$ . Kvamme and Borgan (2019) assume the hazard is constant in predefined time intervals. Nagpal et al. (2021) uses a mixture of Cox models parameterized by neural networks. These models optimize the partial likelihood function which does not require the access to survival functions. The partial likelihood is defined by

$$\prod_{i:\Delta_i=1} \frac{\exp(g(u_i, x_i; \theta))}{\sum_{j \in R_i} \exp(g(u_i, x_j; \theta))},$$

where  $R_i = \{j : y_j \geq y_i\}$  denotes the set of patients who survive longer than the  $i$ -th patient. The goal of maximizing the partial likelihood is to make the patient  $i$ 's relative risk at  $u_i$  greater than the other patients who survive longer. However, this requires the whole dataset to evaluate since the risk set involves all the patients. This disadvantage disables the stochastic gradient descent algorithm for training. Though we can use the mini-batches of patients to approximate the risk set  $R_i$ , there are no theoretical guarantees for convergence. When there are thousands of datapoints, stochastic gradient descent is more efficient than gradient descent.

**Deep Discrete Models** Deep categorical survival models (Miscouridou et al., 2018; Fotso, 2018; Goldstein et al., 2020) divide the time axis into a sequence of bins and turn survival analysis into predicting a time's bin. The models use  $K$  bins where the last bin includes all times greater than some value. DeepHit (Lee et al., 2018) adds a rank-based loss and uses discrete models for competing risks. Nnet-survival (Biganzoli et al., 1998; Gensheimer and Narasimhan, 2019) models the survival function by multiplications of conditional probabilities in previous time intervals. These discrete models can approximate arbitrary smooth distributions with increasing fidelity as  $K$  increases (Miscouridou et al., 2018).

However, discrete models have their own problems. These models do not define what happens to the survival function estimation within a bin, at least without additional assumptions e.g. linearly interpolating the CDF. Next, it is challenging to choose the bin boundaries. It is unclear whether to set bin boundaries by population percentiles or by regular intervals (Kvamme and Borgan, 2019; Tang et al., 2020; Craig et al., 2021). Using regular intervals may lead times to concentrate into a small subset of bins. For percentiles, it is unclear whether we should include the censored times into the population. Percentiles of the observed failure times may not equal the percentiles of true failure times. Finally, deep discrete models are based on classification architectures, meaning that they may be overconfident and suffer the same poor calibration observed for classifiers in (Guo et al., 2017).

**ODE-based Models** Recently, continuous time models with neural ODEs (Chen et al., 2018) have been proposed. SODEN (Tang et al., 2020) considers the evolution of cumulative hazard functions as an ODE while Danks and Yau (2022) model the CDF by an ODE. Groha et al. (2020) use the ODE for multi-state survival analysis. ODE-based models have tractable PDF's and CDF's. However, training neural ODEs is slow (Kelly et al., 2020) because of the expensive numerical integration inside ODE solvers. ODE-based models

also require extra hyperparameters related to ODE-solvers, including the solver type, and tolerance level.

**Other Deep Models** Chapfuwa et al. (2018) use GANs for survival distribution modeling. But they do not use the likelihood as an objective for training since the PDF and CDF of a GAN model are intractable. Minimax training of GANs is known to be unstable (Kodali et al., 2017; Bottou et al., 2018). Ranganath et al. (2016) use deep exponential families (Ranganath et al., 2015) with Weibull likelihoods. This approach necessitated the use of black-box variational inference with Monte Carlo gradients (Ranganath et al., 2014; Mohamed et al., 2020), which typically yields both a lower bound on the likelihood and noisier, slower optimizations. Survival stacking (Craig et al., 2021) casts the survival analysis as a classification task by predicting whether one patient is in other patients’ risk sets. But for  $N$  datapoints, survival stacking creates  $O(N^2)$  classification problems which is not tractable for large datasets.

**Our Model** In this work, we propose a new flexible survival model named Survival Mixture Density Networks. Survival MDNs utilize mixture density networks (Bishop, 1994) to allow flexible modeling. With Gaussians as the base distributions, computing CDF and PDF of the model requires the evaluation of standard functions and the error function. The error function can be obtained efficiently via some approximations (Abramowitz et al., 1988). Our simple approach can be trained through stochastic gradient descent and much faster than the ODE-based models. We compare our model with the previous approaches in table 1.

Model	Flexible	Continuous-time	SGD	Without ODE-Solver
Cox	✗	✓	✗	✓
DeepSurv	✗	✓	✗	✓
DeepHit	✓	✗	✓	✓
Nnet-survival	✓	✗	✓	✓
Cox-Time	✓	✓	✗	✓
SODEN	✓	✓	✓	✗
Survival MDN	✓	✓	✓	✓

**Table 1:** Comparison of Different Models

In summary, we propose a continuous-time model that can be trained with stochastic gradients, without numerical ODE solving, and that moves beyond common modeling restrictions (e.g. that the density is log-normal or Cox).

### 3. Survival Mixture Density Networks

Our purpose is to build a survival model that has the following properties:

1. It has a differentiable PDF which can be evaluated efficiently.
2. It has a differentiable CDF which can be evaluated efficiently.

3. It is flexible enough to approximate a wide class of conditional time-to-event distributions  $p(t|x)$  with support over  $\mathbb{R}^+$ .

The first two properties enable efficient training using maximum likelihood and using stochastic gradients. Examples of the last property are models that do not make assumptions like lognormality or proportional hazards.

### 3.1. Mixture Density Networks

Mixture Density Networks (MDNs) (Bishop, 1994) form the key part of Survival MDN. For a given  $x$ , MDNs model the conditional distribution  $p(y|x)$  by mapping  $x$  through a neural network to produce the weights and parameters of a mixture model. Mixture density networks are flexible approximators. For any given  $x$ , with enough components, MDNs can approximate any conditional density  $p(y|x)$  as closely as desired (Bishop, 1994).

In this work, we use Gaussian mixtures (Reynolds et al., 2000; Reynolds, 2009). A discussion on different base distributions can be found in appendix C. Assume we have  $K$  components with weights  $\{w_i\}_{i=1}^K$ , means  $\{\mu_i\}_{i=1}^K$  and standard deviations  $\{\sigma_i\}_{i=1}^K$  such that  $\sum_{i=1}^K w_i = 1$ . The PDF of the Gaussian Mixture Density Network is given by

$$p(y|\{w_i, \mu_i, \sigma_i\}_{i=1}^K) = \sum_{i=1}^K w_i \mathcal{N}(y|\mu_i, \sigma_i^2),$$

where we denote  $\mathcal{N}(y|\mu_i, \sigma_i^2)$  as the density of a Gaussian distributed random variable with mean  $\mu_i$  and variance  $\sigma_i^2$ .

In mixture density networks, we build the conditional distribution by mapping the covariates  $x$  to parameters of the Gaussian Mixture Model through deep neural networks:

$$\{w_i(x), \mu_i(x), \sigma_i(x)\}_{i=1}^K = f_\theta(x),$$

where  $f_\theta$  is a trainable neural network with parameters  $\theta$ .

### 3.2. Survival Mixture Density Networks

We propose Survival Mixture Density Networks (Survival MDN) to satisfy the properties we want for a survival model.

The sampling process for Survival MDN on a given input  $x$  is

1. Calculate  $\{w_i(x), \mu_i(x), \sigma_i(x)\}_{i=1}^K = f_\theta(x)$ .
2. Sample  $y$  according to the PDF  $\sum_{i=1}^K w_i \mathcal{N}(y|\mu_i, \sigma_i^2)$ . To do so, first sample a component  $i$  with probability equal to  $w_i$  and then sample from  $\mathcal{N}(\mu_i, \sigma_i^2)$ .
3. Map  $y$  to the event  $t$  using  $t = g(y) = \log(1 + \exp(y))$ .

The invertible `softplus` function  $g(y) = \log(1 + \exp(y))$  maps the sample from the mixture density network to the positive domain. Another common choice to map the input from  $\mathbb{R}$  to  $\mathbb{R}^+$  is `exp`. We choose `softplus` over `exp` for the reason that `exp` may place high density on very large times.

Next, we show that the PDF and CDF of the Survival MDN is easy to compute. By the change of variables, the Survival MDN PDF at time  $t$  for input  $x$  is:

$$p(t|x) = \left| \frac{dg^{-1}(t)}{dt} \right| \left( \sum_{i=1}^K w_i(x) \mathcal{N}(g^{-1}(t) | \mu_i(x), \sigma_i^2(x)) \right).$$

For the simple choice of the `softplus`, the absolute value term does not depend on the parameters of neural network  $f_\theta$  so this term does not contribute to the log-likelihood training. The Survival MDN CDF at time  $t$  can be computed easily as well. Denote  $F(\cdot | \mu_i, \sigma_i^2)$  as the CDF of the  $i$ -th component in a Gaussian mixture models. Denote  $F(t|x)$  as the CDF of the Survival MDN and  $F_{\text{MDN}}(y|x)$  as the CDF of the underlying MDN. Since `softplus` is an increasing invertible function, we show that the CDF of the Survival MDN at time  $t$  only requires evaluations of the underlying Gaussian CDFs:

$$\begin{aligned} F(t|x) &= F_{\text{MDN}}(g^{-1}(t)|x) \\ &= \int_{-\infty}^{g^{-1}(t)} \sum_{i=1}^K w_i(x) N(y | \mu_i(x), \sigma_i^2(x)) dy \\ &= \sum_{i=1}^K w_i(x) \int_{-\infty}^{g^{-1}(t)} N(y | \mu_i(x), \sigma_i^2(x)) dy \\ &= \sum_{i=1}^K w_i(x) F(g^{-1}(t) | \mu_i(x), \sigma_i^2(x)) \end{aligned}$$

The evaluation of Gaussian CDF's can be done efficiently through the error function `erf`( $\cdot$ ) which is the CDF of the standard normal distribution:

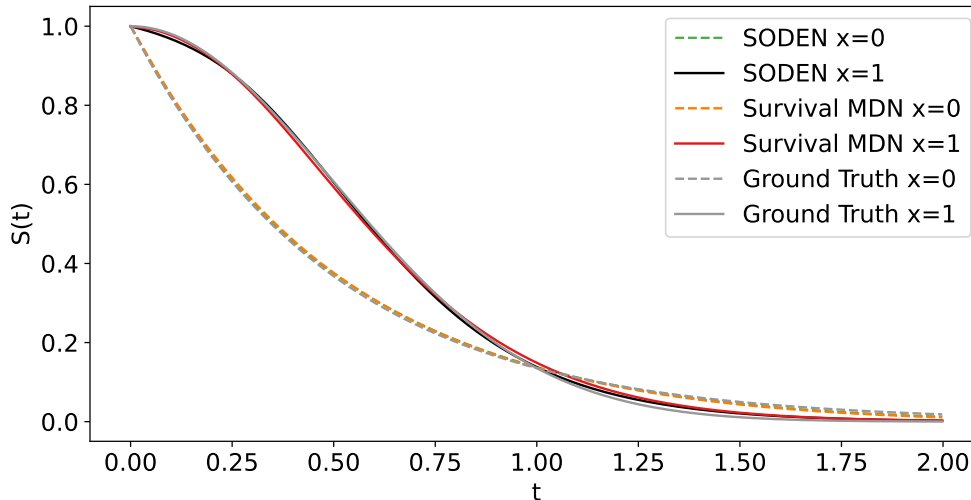
$$F(g^{-1}(t) | \mu_i(x), \sigma_i^2(x)) = \text{erf}((g^{-1}(t) - \mu_i(x)) / \sigma_i(x)).$$

The `erf` function can be computed efficiently via common approximations ([Abramowitz et al., 1988](#)). Now we have satisfied the first two desired properties (PDF and CDF). The last property, flexibility, follows since the Survival MDN maps time-to-event densities to densities over the reals via  $y = g^{-1}(t)$  and a mixture density network with enough components can approximate any smooth density  $p(y|x)$  as closely as desired ([Bishop, 1994](#)).

#### 4. Simulation Study

In this simulation experiment, we test Survival MDN and SODEN on a dataset that the proportional hazard assumption does not hold. We follow the simple simulation setting in SODEN ([Tang et al., 2020](#)). There are two group of  $x$ 's,  $x = 0$  and  $x = 1$ . And the ground truth survival function is:

$$S(t|x) = \exp(-2t) \cdot I\{x = 0\} + \exp(-2t^2) \cdot I\{x = 1\},$$



**Figure 1:** The survival functions of two groups. We show the survival function of two trained model SODEN and Survival MDN together with the ground truth on  $x = 0$  and  $x = 1$  separately. For  $x = 0$ , three survival functions are so close to each other that the green curve for SODEN is covered by the blue curve for Survival MDN and the gray curve for the ground truth.

where  $I$  is the indicator function. This survival distribution does not obey the proportional hazard assumption. Therefore, models that require PH assumption could not fit this dataset well. We generate  $x$  from a Bernoulli distribution with probability 0.5 and then we generate  $t$  using the inverse CDF method. Then we sample the censored time uniformly from  $[0, 2]$ . Instead of simulating a fixed dataset, we use an “online” training method. In each iteration, we generate a new set of 1024 datapoints. We use the likelihood function for training. We train 10,000 iterations for both SODEN and Survival MDN.

We show the resulting survival functions and ground truth in fig. 1. Both Survival MDN and SODEN’s survival functions are close to the ground truth at both  $x = 0$  and  $x = 1$ .

## 5. Real World Experiments

In this section, we compare Survival MDN with baselines Cox, DeepSurv, Cox-Time, Nnet-survival, DeepHit and SODEN. We use four different datasets: SUPPORT, METABRIC, GBSG and MIMIC. We evaluate all models on three different metrics: concordance, integrated binomial log-likelihood and integrated brier score.

### 5.1. Datasets

We choose four different datasets: SUPPORT, METABRIC, GBSG and MIMIC. SUPPORT, METABRIC and GBSG are commonly used datasets for survival analysis, which can be found in `pycox` package. MIMIC is a dataset we preprocessed from MIMIC-iv ([John-](#)



son et al., 2020) in PhysioNet (Goldberger et al., 2000). We describe the details of the datasets here:

- SUPPORT: the Study to Understand Prognoses Preferences Outcomes and Risks of Treatment. It has 14 features. There are 8,873 datapoints, 32% of which are censored. We use the train/valid/test splits from SODEN (Tang et al., 2020)<sup>2</sup>.
- METABRIC: the Molecular Taxonomy of Breast Cancer International Consortium. It has 9 features. There are 1,904 datapoints, 42% of which are censored. We use the train/valid/test splits from the SODEN repository.
- GBSG: The Rotterdam & German Breast Cancer Study Group. It has 7 features. There are 2,232 datapoints, 43% of which are censored.
- MIMIC: The Medical Information Mart for Intensive Care. The SODEN repository does not provide the data files for MIMIC. We choose patients that are alive 24 hours after admission to ICU. We define the event as mortality after admission. We define the censored time as the ICU discharged time. We collect time series features within the 24-hour window after the admission together with the static features. For time series features, we use the minimum, mean and maximum within the window. We remove the features that are missing more than half of the datapoints. Finally, we extract 65 features after preprocessing including common labs and vitals. There are 53,612 datapoints, 82% of which are censored. The SQL code that preprocesses the data from MIMIC-iv is attached in appendix A.

## 5.2. Baselines

We consider baseline models:

- Cox (Cox, 1972): A linear model with the proportional hazards assumption.
- DeepSurv (Katzman et al., 2018): A deep learning model with the linear function in Cox replaced by neural networks.
- Cox-Time (Katzman et al., 2018): A continuous time model that allows the relative risk in Cox to depend on time.
- Nnet-Survival (Gensheimer and Narasimhan, 2019): A discrete time model that models the conditional hazard in each time interval.
- DeepHit (Lee et al., 2018): A deep discrete time model that further adds a rank-based loss to the likelihood as the training objective.
- SODEN (Tang et al., 2020): An ODE-based continuous time model.

For Cox, we use the implementation in the Python package `lifelines`. For DeepSurv, Cox-Time, Nnet-Survival and DeepHit, we use the implementations in the Python package `pycox`. For SODEN, we use the implementation from the SODEN repository.

2. Available at <https://github.com/jiaqima/SODEN>

$P(C > \tau)$	Model	$C_{\tau}^{td}(\uparrow)$	IBLL $_{\tau}(\uparrow)$	IBS $_{\tau}(\downarrow)$
$10^{-8}$	Cox	0.596 ± .002	-0.568 ± .001	0.194 ± .001
	DeepSurv	0.609 ± .003	<b>-0.559</b> ± .002	<b>0.190</b> ± .001
	Cox-Time	0.607 ± .004	0.565 ± .002	0.191 ± .001
	Nnet-Survival	0.624 ± .003	-0.570 ± .004	0.193 ± .001
	DeepHit	<b>0.631</b> ± .003	-0.583 ± .006	0.197 ± .001
	SODEN	0.627 ± .003	-0.563 ± .002	0.191 ± .001
	Survival MDN	0.628 ± .003	<b>-0.559</b> ± .002	<b>0.190</b> ± .002
0.2	Cox	0.596 ± .002	-0.585 ± .001	0.201 ± .000
	DeepSurv	0.609 ± .003	-0.577 ± .002	0.197 ± .001
	Cox-Time	0.606 ± .004	-0.583 ± .002	0.199 ± .001
	Nnet-Survival	0.623 ± .003	-0.586 ± .003	0.201 ± .001
	DeepHit	<b>0.630</b> ± .003	-0.601 ± .006	0.205 ± .002
	SODEN	<b>0.630</b> ± .003	-0.601 ± .006	0.205 ± .002
	Survival MDN	0.628 ± .003	<b>-0.575</b> ± .002	<b>0.196</b> ± .001
0.4	Cox	0.595 ± .002	-0.602 ± .001	0.208 ± .001
	DeepSurv	0.608 ± .002	-0.595 ± .002	0.205 ± .001
	Cox-Time	0.605 ± .004	-0.601 ± .002	0.207 ± .001
	Nnet-Survival	0.623 ± .003	-0.602 ± .003	0.208 ± .001
	DeepHit	<b>0.630</b> ± .003	-0.619 ± .007	0.212 ± .002
	SODEN	0.626 ± .003	-0.597 ± .002	0.205 ± .001
	Survival MDN	0.628 ± .003	<b>-0.593</b> ± .001	<b>0.204</b> ± .001

**Table 2:** Evaluation of all models on SUPPORT with concordance ( $C_{\tau}^{td}$ ), integrated binomial log-likelihood (IBLL $_{\tau}$ ) and integrated Brier score (IBS $_{\tau}$ ). The **bold** number indicates the best performance. We report the mean and the standard error of the mean on all the metrics (mean ± standard error).

### 5.3. Evaluation Metrics

We use the same evaluation metrics as SODEN (Tang et al., 2020). They are concordance, integrated binomial log-likelihood and Brier score. The implementations can be found in the SODEN repository. We briefly describe the three metrics here and refer more detailed descriptions to Tang et al. (2020).

**Concordance** The concordance index is originally proposed by Harrell Jr et al. (1984). It measures the probability that the relative order of the event time of two observations matches the predicted survival probabilities. Antolini et al. (2005) further relaxes the proportional hazard assumption in Harrell’s concordance to create time dependent concordance. Building off the inverse-weighting method in Cheng et al. (1995), Uno et al. (2011) introduces inverse probability weighted concordance to remove the dependence on the censoring distribution. They use the survival distribution of the censoring time  $G(t) = P(C > t)$  as the weight and the Kaplan-Meier estimator for  $G(t)$ . Under the completely random censoring assumption  $C \perp\!\!\!\perp (T, X)$ , the inverse probability weighted estimator is consistent.

$P(C > \tau)$	Model	$C_\tau^{td}(\uparrow)$	IBLL $_\tau(\uparrow)$	IBS $_\tau(\downarrow)$
$10^{-8}$	Cox	0.644 ± .006	-0.508 ± .009	0.169 ± .002
	DeepSurv	0.635 ± .007	-0.517 ± .011	0.171 ± .003
	Cox-Time	0.648 ± .007	-0.511 ± .009	0.172 ± .003
	Nnet-Survival	0.666 ± .005	-0.510 ± .007	0.171 ± .002
	DeepHit	<b>0.674</b> ± .006	-0.514 ± .004	0.174 ± .002
	SODEN	0.661 ± .005	-0.498 ± .008	0.167 ± .003
	Survival MDN	0.667 ± .004	<b>-0.489</b> ± .005	<b>0.165</b> ± .002
0.2	Cox	0.639 ± .006	-0.521 ± .006	0.176 ± .002
	DeepSurv	0.635 ± .006	-0.530 ± .005	0.179 ± .002
	Cox-Time	0.647 ± .005	-0.531 ± .007	0.179 ± .002
	Nnet-Survival	0.662 ± .004	-0.523 ± .003	0.177 ± .001
	DeepHit	<b>0.671</b> ± .004	-0.533 ± .003	0.182 ± .001
	SODEN	0.659 ± .003	-0.516 ± .006	0.174 ± .002
	Survival MDN	0.662 ± .004	<b>-0.510</b> ± .003	<b>0.172</b> ± .001
0.4	Cox	0.637 ± .006	-0.521 ± .006	0.175 ± .002
	DeepSurv	0.635 ± .006	-0.526 ± .005	0.178 ± .002
	Cox-Time	0.644 ± .005	-0.526 ± .006	0.178 ± .002
	Nnet-Survival	0.660 ± .003	-0.519 ± .003	0.176 ± .001
	DeepHit	<b>0.668</b> ± .003	-0.528 ± .003	0.180 ± .001
	SODEN	0.658 ± .004	-0.528 ± .003	0.180 ± .001
	Survival MDN	0.660 ± .002	<b>-0.508</b> ± .003	<b>0.172</b> ± .001

**Table 3:** Evaluation of all models on METABRIC with concordance ( $C_\tau^{td}$ ), integrated binomial log-likelihood (IBLL $_\tau$ ) and integrated Brier score (IBS $_\tau$ ). We report truncated metrics for  $\tau$ 's satisfying  $P(C > \tau) = 10^{-8}, 0.2, 0.4$ . The **bold** number indicates the best performance. We report the mean and the standard error of the mean on all the metrics (mean ± standard error).

This assumption is routinely made for evaluation, e.g. in [Kvamme et al. \(2019\)](#); [Tang et al. \(2020\)](#); [Han et al. \(2021\)](#). Due to the limited number of observations, the estimator of the inverse weight  $1/\hat{G}(t)$  may be very large for some large-enough  $t$ . So [Uno et al. \(2011\)](#) introduce a truncated version of the concordance estimator within a pre-specified time interval  $[0, \tau]$ :

$$C_\tau^{td} = \frac{\sum_{i:\Delta_i=1, u_i < \tau} \sum_{j:u_i < u_j} I(\hat{S}(u_i|x_i) < \hat{S}(u_i|x_j)) / \hat{G}^2(u_i)}{\sum_{i:\Delta_i=1, u_i < \tau} \sum_{j:u_i < u_j} 1 / \hat{G}^2(u_i)},$$

where  $I(\cdot)$  is the indicator function. Here  $\tau$  is used to truncate the large times that have very small  $\hat{G}(t)$ . We choose three  $\tau$ 's that satisfy  $\hat{G}(\tau) = 10^{-8}, 0.2, 0.4$ . When  $\tau = 10^{-8}$ , the truncated concordance is almost equal to the non-truncated version.

**Integrated Brier Score** The Brier score (BS) measures the mean square error between the ground-truth label and the predicted probability for a binary classifier. It measures

both the calibration and discriminative performance. In survival analysis, we evaluate the Brier score at a given time  $t$ . The label is whether the patient can survive after time  $t$  and the predicted probability is the survival function. We also consider an inverse probability weighted estimator (Graf et al., 1999; Gerds and Schumacher, 2006) for the Brier score at time  $t$

$$\text{BS}(t) = \frac{1}{N} \sum_{i=1}^N \left\{ \frac{\hat{S}^2(t|x_i)I(u_i \leq t, \Delta_i = 1)}{\hat{G}(u_i)} + \frac{(1 - \hat{S}(t|u_i))^2 I(u_i > t)}{\hat{G}(t)} \right\},$$

where  $I(\cdot)$  is the indicator function. To consider all times, we use an integrated BS (IBS) over a time interval  $[0, \tau]$ :

$$\text{IBS}_\tau = \frac{1}{\tau} \int_0^\tau \text{BS}(t) dt.$$

To avoid extreme inverse weights, we also report results for  $\tau$ 's that satisfy  $\hat{G}(\tau) = 10^{-8}, 0.2, 0.4$ . When  $\hat{G}(\tau) = 10^{-8}$ ,  $\tau$  is almost equal to the maximum time in the data.

**Integrated Binomial Log-Likelihood** Another common metric for survival analysis is the integrated binomial log-likelihood (IBLL). Different from IBS, IBLL uses binomial log-likelihood at each time step  $t$

$$\text{BLL}(t) = \frac{1}{N} \sum_{i=1}^N \left\{ \frac{\log(1 - \hat{S}(t|x_i)I(u_i \leq t, \Delta_i = 1))}{\hat{G}(u_i)} + \frac{\log(\hat{S}(t|x_i)I(u_i > t))}{\hat{G}(t)} \right\},$$

where  $I(\cdot)$  is the indicator function. And IBLL is defined by

$$\text{IBLL}_\tau = \frac{1}{\tau} \int_0^\tau \text{BLL}(t) dt.$$

We also report results for  $\tau$ 's satisfying  $\hat{G}(\tau) = 10^{-8}, 0.2, 0.4$ .

#### 5.4. Experimental Setup

We randomly split datasets into training, validation, and testing sets. We use the validation set to choose the best epoch from training and hyperparameters and report the results on the test set. For SUPPORT/METABRIC/GBSG, we use 10 splits (8 for training, 1 for validation and 1 for test). For MIMIC, we use 5 splits (3 for training, 1 for validation, and 1 for test) since the size of MIMIC is a bit large. We use random search to create 100 independent trials for different hyperparameters. We use the optimizer RMSProp (Tieleman et al., 2012).

For Survival MDN, following Sudarshan et al. (2020), we use a three-layer neural network that maps the features to a latent representation, and then from the latent representation we use three linear layers to output  $w$ 's,  $\mu$ 's,  $\sigma$ 's separately. We use a `softmax` layer to make sure that the sum of  $w$ 's equals one and use an `exp` function to ensure the standard deviations  $\sigma$ 's are positive. We vary the number of components from 5 to 20.

For other models, we vary the number of layers. Other hyperparameters include the hidden sizes, learning rate, batch normalization, momentum, dropout, and batch size. For DeepHit and Nnet-Survival, we vary the number of time intervals in addition. For other hyperparameters, we use the same tuning ranges as in (Tang et al., 2020). We show the tuning ranges in appendix B.

$P(C > \tau)$	Model	$C_{\tau}^{td}(\uparrow)$	IBLL $_{\tau}(\uparrow)$	IBS $_{\tau}(\downarrow)$
$10^{-8}$	Cox	0.645 ± .009	-0.523 ± .009	0.177 ± .004
	DeepSurv	0.663 ± .007	-0.509 ± .010	<b>0.172</b> ± .004
	Cox-Time	0.654 ± .007	-0.521 ± .009	0.176 ± .003
	Nnet-Survival	0.661 ± .006	-0.516 ± .008	0.174 ± .005
	DeepHit	0.665 ± .008	<b>-0.504</b> ± .017	0.176 ± .005
	SODEN	0.661 ± .012	-0.514 ± .017	0.173 ± .004
	Survival MDN	<b>0.668</b> ± .007	<b>-0.504</b> ± .006	<b>0.172</b> ± .003
0.2	Cox	0.645 ± .009	-0.519 ± .007	0.176 ± .003
	DeepSurv	0.663 ± .007	-0.505 ± .008	0.170 ± .002
	Cox-Time	0.654 ± .007	-0.517 ± .006	0.175 ± .002
	Nnet-Survival	0.661 ± .006	-0.509 ± .006	0.170 ± .003
	DeepHit	0.665 ± .008	-0.510 ± .008	0.172 ± .004
	SODEN	0.661 ± .012	-0.510 ± .009	0.172 ± .004
	Survival MDN	<b>0.668</b> ± .007	<b>-0.501</b> ± .006	<b>0.168</b> ± .002
0.4	Cox	0.645 ± .009	-0.519 ± .007	0.176 ± .003
	DeepSurv	0.663 ± .007	-0.505 ± .008	0.170 ± .002
	Cox-Time	0.654 ± .007	-0.517 ± .006	0.175 ± .002
	Nnet-Survival	0.661 ± .006	-0.509 ± .007	0.170 ± .003
	DeepHit	0.665 ± .008	-0.510 ± .008	0.172 ± .004
	SODEN	0.661 ± .012	-0.510 ± .009	0.172 ± .004
	Survival MDN	<b>0.668</b> ± .007	<b>-0.500</b> ± .006	<b>0.168</b> ± .002

**Table 4:** Evaluation of all models on GBSG with concordance ( $C_{\tau}^{td}$ ), integrated binomial log-likelihood (IBLL $_{\tau}$ ) and integrated Brier score (IBS $_{\tau}$ ). We report truncated metrics for  $\tau$ 's satisfying  $P(C > \tau) = 10^{-8}, 0.2, 0.4$ . The **bold** number indicates the best performance. We report the mean and the standard error of the mean on all the metrics (mean ± standard error).

## 5.5. Results

We report the results on four datasets in table 2 (SUPPORT), table 3 (METABRIC), table 4 (GBSG), and table 5 (MIMIC). For SUPPORT and METABRIC, we use the exact same splits as the SODEN repository so we use their results for the baselines.

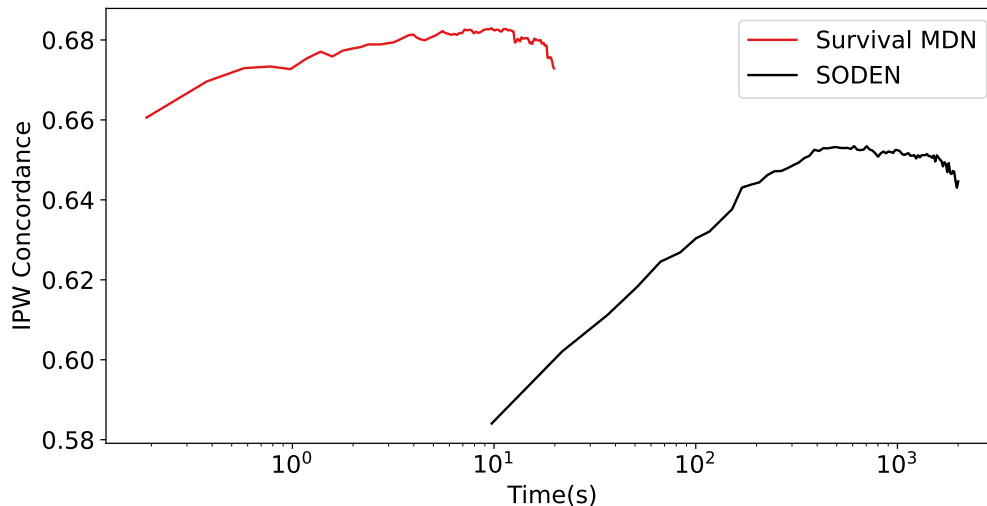
For concordance, DeepHit have the best concordance on SUPPORT and METABRIC while the continuous time model Survival MDN have the best ones on GBSG and MIMIC. For IBLL and IBS, Survival MDN has the best performance across all datasets. The IBLL and IBS care more about the exact survival probability prediction at each time. The discrete time model DeepHit may not give an accurate estimation for survival probability for a particular time since it does not distinguish the times inside one time interval. The discrete time models also have trouble for choosing the bin boundaries (Kvamme and Borgan, 2019; Tang et al., 2020; Craig et al., 2021). Their concordance on MIMIC is worse than SODEN and Survival MDN. Continuous time models Survival MDN and SODEN have similar performance on concordance on four datasets since they are both flexible continuous

time models. There is little difference among  $\hat{G}(\tau) = 10^{-8}, 0.2, 0.4$  for the concordance, IBLL and IBS on small datasets SUPPORT, METABRIC and GBSG, which is the same observation in SODEN (Tang et al., 2020).

$P(C > \tau)$	Model	$C_{\tau}^{td}(\uparrow)$	IBLL $_{\tau}(\uparrow)$	IBS $_{\tau}(\downarrow)$
$10^{-8}$	Cox	0.642 ± .002	-0.211 ± .001	0.061 ± .001
	DeepSurv	<b>0.663</b> ± .001	-0.212 ± .003	0.061 ± .001
	Cox-Time	0.653 ± .001	-0.210 ± .003	0.061 ± .001
	Nnet-Survival	0.649 ± .002	-0.206 ± .000	0.061 ± .001
	DeepHit	0.647 ± .002	-0.206 ± .001	0.061 ± .001
	SODEN	0.659 ± .002	<b>-0.204</b> ± .002	0.060 ± .001
	Survival MDN	0.660 ± .002	<b>-0.204</b> ± .002	<b>0.059</b> ± .001
0.2	Cox	0.711 ± .004	-0.473 ± .133	0.091 ± .014
	DeepSurv	0.734 ± .003	-0.462 ± .150	0.089 ± .015
	Cox-Time	0.726 ± .002	-0.443 ± .126	0.061 ± .001
	Nnet-Survival	0.722 ± .004	-0.229 ± .004	0.066 ± .001
	DeepHit	0.719 ± .004	-0.233 ± .004	0.066 ± .001
	SODEN	0.733 ± .002	-0.229 ± .004	<b>0.065</b> ± .001
	Survival MDN	<b>0.736</b> ± .003	<b>-0.228</b> ± .004	<b>0.065</b> ± .001
0.4	Cox	0.780 ± .002	-0.588 ± .136	0.071 ± .031
	DeepSurv	0.797 ± .001	-0.423 ± .202	0.045 ± .018
	Cox-Time	0.790 ± .002	-0.501 ± .267	0.037 ± .010
	Nnet-Survival	0.784 ± .003	-0.082 ± .003	<b>0.018</b> ± .001
	DeepHit	0.787 ± .003	-0.083 ± .002	0.019 ± .001
	SODEN	<b>0.805</b> ± .005	-0.084 ± .002	0.019 ± .001
	Survival MDN	<b>0.805</b> ± .001	<b>-0.078</b> ± .002	<b>0.018</b> ± .001

**Table 5:** Evaluation of all models on MIMIC with concordance ( $C_{\tau}^{td}$ ), integrated binomial log-likelihood (IBLL $_{\tau}$ ) and integrated Brier score (IBS $_{\tau}$ ). We report truncated metrics for  $\tau$ 's satisfying  $P(C > \tau) = 10^{-8}, 0.2, 0.4$ . The **bold** number indicates the best performance. We report the mean and the standard error of the mean on all the metrics (mean ± standard error).

The training time of SODEN is much longer than Survival MDN. We collect the training time of two models with the same hidden size 32 and number of layers 4 on METABRIC. We use the maximum number of components in the tuning range 20 for Survival MDN. We show the test concordance versus the training time for Survival MDN and SODEN on GeForce RTX 2080 Ti in fig. 2. We can see that Survival MDN reached the peak of the test concordance much faster than SODEN. On average, each epoch of Survival MDN costs 0.20 seconds while each epoch of SODEN costs 23.82 seconds. Training Survival MDN is more than 100 time faster than SODEN.



**Figure 2:** Comparison between Survival MDN and SODEN on IPW Concordance versus training time. The time is shown in log scale.

## 6. Discussion

We propose Survival MDN, a simple flexible continuous time survival modeling. Survival modeling plays an important role in risk estimation and clinical decision making. The proposed model can speed up the training and evaluation of survival modeling which may also accelerate the clinical decision process.

In this work, we combine two simple yet elegant tools—mixture densities and change of variables—to produce flexible survival models. While recent approaches achieve similar flexibility, it is achieved at the expense of training time, complexity, and inconvenient hyper-parameters. Without introducing such complexity, Survival MDNs achieve better or similar performance.

**Limitations** Currently, the proposed model survival MDN only considers Gaussian Mixtures. Though Gaussian Mixtures have a universal approximation power, a combination of different base distributions, e.g. generalized logistics, in mixture density networks may improve the performance. Regarding experimental evaluation, the marginal censoring assumption used in the reweighting estimators is common practice in the literature, but may not be appropriate. Evaluation with censored data is impossible without assumptions, but it could be possible to improve evaluation by making conditional censoring assumptions.

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## Appendix A. MIMIC SQL code

```

select
-- ids
pat.subject_id as subject_id, adm.hadm_id as hadm_id, icu.stay_id as stay_id,
-- demographics
CASE WHEN pat.gender="M" THEN 1 ELSE 0 END as is_male,
CASE WHEN adm.ethnicity="WHITE" THEN 1 ELSE 0 END as is_white,
icu_detail.admission_age as age,
-- weight height
fdw.weight ,
fdh.height ,
-- LOS
icu.los as los_icu_days,
icu_detail.los_hospital as los_hosp_days,
-- death
--icu_detail.icu_intime as icu_intime,
--icu_detail.dod as dod,
TIMESTAMP_DIFF(icu_detail.dod, icu_detail.icu_intime, HOUR) / 24 as time_to_death,
case
    when icu_detail.dod is null then 0
    else 1
end
as death,
-- vitals labs min max mean
vitals.*,
labs.*,
sofa.*
from 'physionet-data.mimic_core.patients' pat
inner join
    'physionet-data.mimic_core.admissions' adm
    on pat.subject_id=adm.subject_id
inner join
    'physionet-data.mimic_icu.icustays' icu
    on adm.subject_id=icu.subject_id
    and
    adm.hadm_id=icu.hadm_id
inner join
    'physionet-data.mimic_derived.first_day_height' fdh
    on
    adm.subject_id = fdh.subject_id and icu.stay_id = fdh.stay_id
inner join
    'physionet-data.mimic_derived.first_day_weight' fdw
    on
    adm.subject_id = fdw.subject_id and icu.stay_id = fdw.stay_id

```

```

inner join
  'physionet-data.mimic_derived.icustay_detail' icu_detail
  on
    adm.subject_id=icu_detail.subject_id
  and
    adm.hadm_id=icu_detail.hadm_id
  and
    icu.stay_id=icu_detail.stay_id
inner join
  'physionet-data.mimic_derived.first_day_sofa' sofa
  on
    adm.subject_id=sofa.subject_id
  and
    adm.hadm_id=sofa.hadm_id
  and
    icu.stay_id=sofa.stay_id

inner join
  'physionet-data.mimic_derived.first_day_vitalsign' vitals
  on
    adm.subject_id=vitals.subject_id
  and
    icu.stay_id=vitals.stay_id
inner join
  'physionet-data.mimic_derived.first_day_lab' labs
  on
    adm.subject_id=labs.subject_id
  and
    icu.stay_id=labs.stay_id
where icu_detail.los_icu > 1
  and pat.gender is not null
  and adm.ethnicity is not null
  and adm.ethnicity != "UNABLE TO OBTAIN"
  and adm.ethnicity != "UNKNOWN"

```

## Appendix B. Tuning Ranges of Hyperparameters

We show the search range of hyperparameters in table 6.

---

Batch size	{32, 64, 128, 256} for METABRIC, GBSG {128, 256, 512} for SUPPORT {512, 1024} for MIMIC
Number of layers	{1, 2, 4}
Hidden size	[2 <sup>2</sup> , 2 <sup>7</sup> ]
Learning rate	[10 <sup>-4.5</sup> , 10 <sup>-1.5</sup> ]
Weight decay	[10 <sup>-9</sup> , 10 <sup>-4</sup> ]
Momentum	[0.85, 0.99]
Dropout	{0, 0.1, 0.5}
Batch normalization	{True, False}
$\alpha$ (Surrogate ranking loss in DeepHit)	[0, 1]
$\sigma$ (Surrogate ranking loss in DeepHit)	{0.25, 1, 5}
Number of intervals (DeepHit, Nnet-survival)	{10, 50, 100, 200, 400} for SUPPORT, METABRIC, GBSG {50, 100, 200, 400, 800} for MIMIC

---

**Table 6:** Tuning ranges of hyperparameters

## Appendix C. Discussion on Different Base Distributions

Here we compare Gaussian base with an alternative base, the generalized logistic distribution, on marginal data generations. We use the following form of the generalized logistic distribution:

$$F(x; \alpha) = 1 - \frac{e^{-\alpha x}}{(1 + e^{-x})^\alpha}.$$

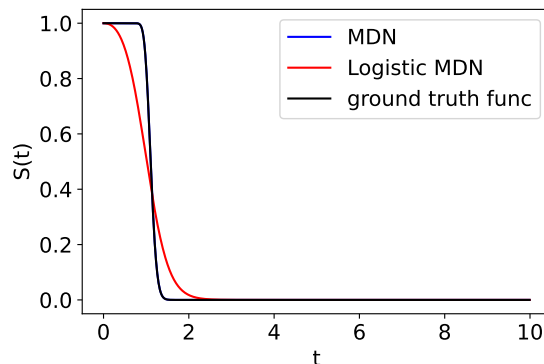
We also shift the generalized logistic distribution using scale and location. In this generalized logistic distribution, we have one more parameter  $\alpha$  which can control the magnitude of the power.

We consider three different marginal data generation cases:

- LogNormal distribution with  $\mu = 0.1$  and  $\sigma = 0.1$ . LogNormal distribution is a common one researchers use in survival analysis. The variance is small in this data generation distribution.
- Student T distribution with degree of freedom one and transformed to positive values through softplus. Student T distribution has a heavy tail.
- Gamma distribution with shape 0.1 and scale 1. When shape is smaller than one, the Gamma distribution put a lot of mass on values close to zero. This may be hard for a mixture model to fit.

We sample the censored time uniformly from  $[0, 10]$ . We still use an online training which generates a whole new batch data in every update step.

The results of fitting LogNormal data is shown in fig. 3. The Gaussian base has survival functions overlapping with the ground truth but the generalized logistic base cannot fit it well.



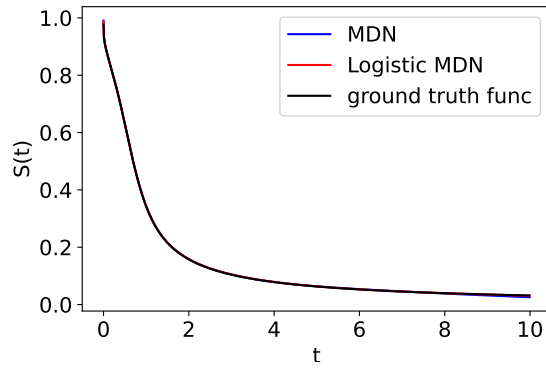
**Figure 3:** LogNormal Data.

The result of fitting student T data is shown in fig. 4. For heavy tailed student T, both Gaussian base and generalized logistic base can also fit it well with survival functions overlapping the ground truth.

The result of fitting Gamma data is shown in fig. 5. The generalized logistic base can fit the Gamma data well while there is some gap between the ground truth and the Gaussian

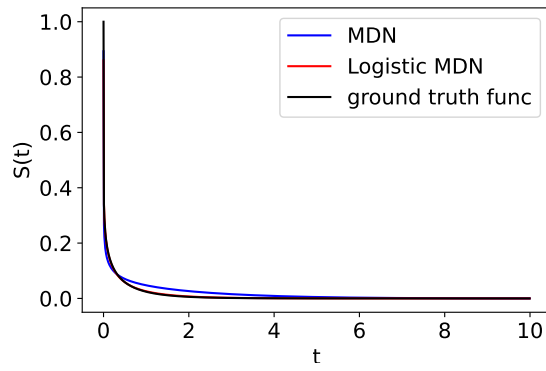


# SURVIVAL MDN



**Figure 4:** Student T + Softplus Data.

base survival function. In Gamma data with a small shape, the generalized logistic base is a better choice.



**Figure 5:** Student T + Softplus Data.