



CZECH TECHNICAL UNIVERSITY IN PRAGUE  
Faculty of Nuclear Sciences and Physical Engineering



# **Generative and discriminative models for set data**

## **Generativní a diskriminativní modely pro množinová data**

Research Project

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*Author's declaration:*

I declare that this Bachelor's Degree Project is entirely my own work and I have listed all the used sources in the bibliography.

Prague, June 1, 2021

Jméno Autora

## Název práce

*Obor:* Celý název oboru (nikoliv zkratka)

*Zaměření:* Celý název zaměření (Pokud obor neobsahuje zaměření, tuto řádku odstranit.)

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[illegible]

**Klíčová slova:** klíčová slova (nebo výrazy) seřazená podle abecedy a oddělená čárkou

### Title of the Work

*Author:* Author's Name

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**Key words:** keywords in alphabetical order separated by commas

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# Introduction

Paragraphs of the Introduction. . .

# Chapter 1

## Multiple Instance Learning

In standard machine learning problems each sample is represented by a fixed vector  $\mathbf{x}$  of values, nevertheless in multiple instance learning (MIL) we deal with samples which are represented by a set of vectors. These vectors are called *instances* and come from an instance space  $\mathcal{X}$ . Sets of these instances are called *bags* and come from bag space  $\mathcal{B} = \mathcal{P}_F(\mathcal{X})$ , where  $\mathcal{P}_F(\mathcal{X})$  denotes all finite subsets of  $\mathcal{X}$ . With this in mind, we can easily write down any bag  $b$  as  $b = \{\mathbf{x} \in \mathcal{X}\}_{\mathbf{x} \in b}$ . Each bag can be arbitrarily large or empty thus the size of bag  $|b| \in \mathbb{N}_0$ . There may exist intrinsic labeling of instances, but we are only interested in labeling at the bag levels. Bag labels come from a finite set  $C$  and what we want in MIL is learning a predictor in the form  $f : \mathcal{B}(\mathcal{X}) \rightarrow C$  which can also be rewritten in the form  $f(\{\mathbf{x}\}_{\mathbf{x} \in b})$ . We consider supervised setting, in which each sample of the dataset is attributed a label. We can denote available data by  $\mathcal{D} = \{(b_i, y_i) \in \mathcal{B} \times C \mid i \in \{1, 2, \dots, |\mathcal{D}|\}\}$ , where  $|\mathcal{D}|$  denotes size of  $\mathcal{D}$ .

### 1.1 Cross-Validation

Probably the simplest and most widely used method for estimating prediction error is *cross-validation*. This method directly estimates the expected extra-sample error

$$\text{Err} = \mathbb{E} \left[ L(Y, \hat{f}(X)) \right], \quad (1.1)$$

the average generalization error when the method  $\hat{f}(X)$  is applied to an independent test sample from the joint distribution of  $X$  and  $Y$ . As mentioned earlier, we might hope that cross-validation estimates the conditional error, with the training set  $\mathcal{T}$  held fixed. But cross-validation typically estimates well only the expected prediction error.

#### K-Fold Cross-Validation

Ideally, if we had enough data, we would set aside a validation set and use it to assess the performance of our prediction model. Since data are often scarce, this is usually not possible. To finesse the problem, K-fold cross-validation uses part of the available data to fit the model, and a different part to test it. We split the data into  $K$  roughly equal-sized parts; for example, when  $K = 5$ , the scenario could look like this

train	train	validation	train	train
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For the  $k$ th part (third above), we fit the model to the other  $K - 1$  parts of the data, and calculate the prediction error of the fitted model when predicting the  $k$ th part of the data. We do this for  $k = \{1, 2, \dots, K\}$  and combine the  $K$  estimates of prediction error.

Here are more details. Let  $\kappa : \{1, \dots, N\} \rightarrow \{1, \dots, K\}$  be an indexing function that indicates the partition to which observation  $i$  is allocated by the randomization. Denote by  $\hat{f}^{-k}(x)$  the fitted function, computed with the  $k$ th part of the data removed. Then the cross-validation estimate of prediction error is

$$\text{CV}(\hat{f}) = \frac{1}{N} \sum_{i=1}^N L(y_i, \hat{f}^{-\kappa(i)}(x_i)). \quad (1.2)$$

Typical choices of  $K$  are 5 or 10 and even case  $K = N$ , which is known as *leave-one-out* cross-validation.

## 1.2 Experiment

Suppose we have two classes 0 and 1 (known as binary classification), which means that bags are labeled either as 0 or 1. What happens, if we have many more bags, for example, labeled as class 1? This situation is very common in anomaly detection, where known anomalies are quite rare.

Let's assume train set is composed of 80% bags labeled as 1 and 5% bags labeled as 0, all randomly chosen. Test set is composed of 20% bags labeled as 1 and 95% labeled as 0, in other words it is complement of train set. Validation set is very similar to train set in terms of ratios, it contains 20% bags labeled as 1 and 2% bags labeled as 0. Train set is used to train our model, after that we evaluate loss function of the model with help of validation a test set, where number of dense layers is our hyperparameter. The purpose of this simulation is to find number of dense layer in which the loss is minimal and compare these 2 values. This experiment was performed 5 times then results were averaged, totally on 6 different datasets.

As we can see in Figure 1.1, the loss evaluated on different sets varies therefore it is likely to make mistakes when choosing our hyperparameter if we don't have enough input data.

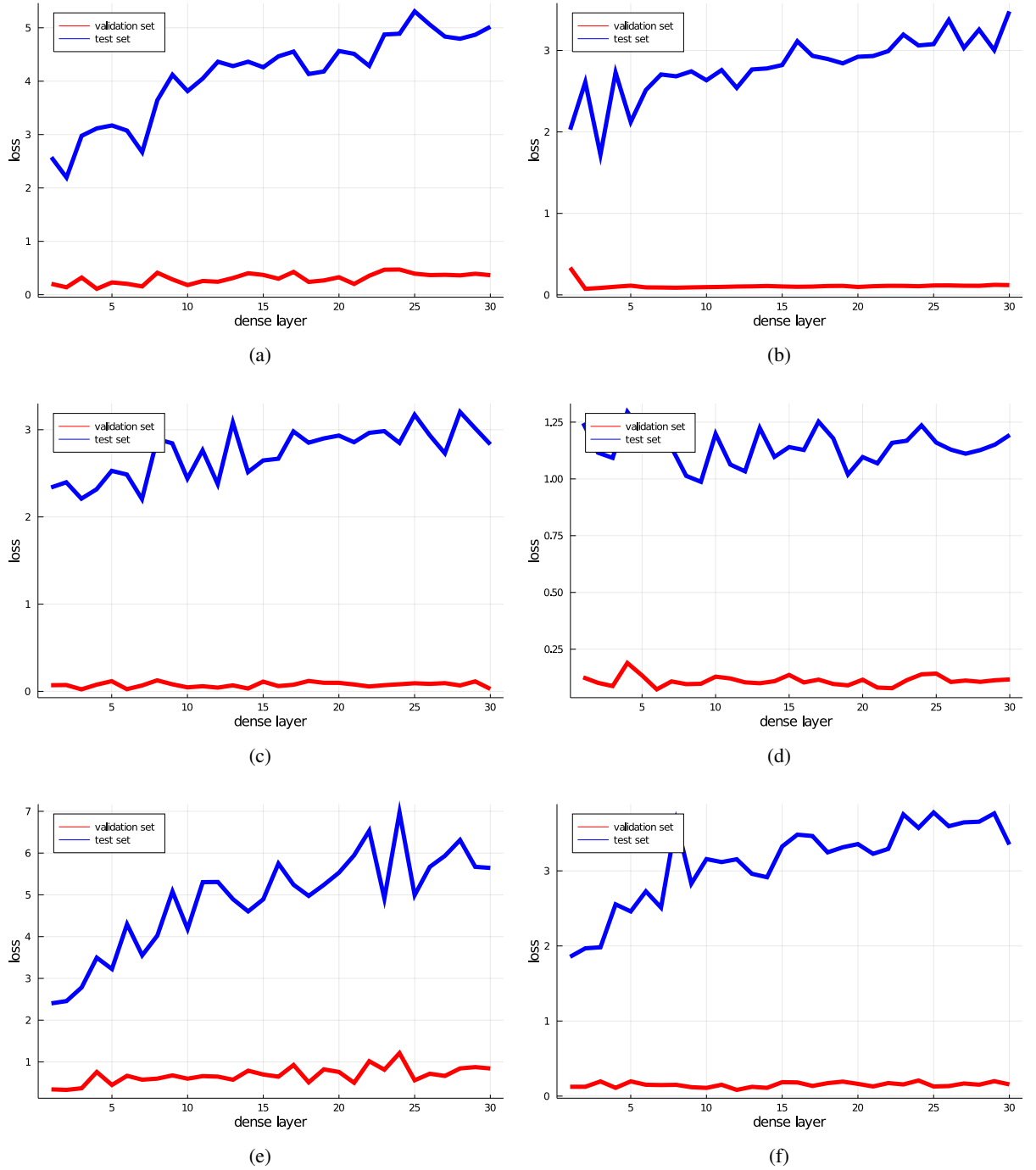


Figure 1.1: Evaluation of loss function with the use of validation and test set on different datasets.



# Conclusion

Text of the conclusion...

# Bibliography

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