# Linear models for regression and classification

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### Setup and examples

Loss functions Building a loss function

# (Informal) definition of supervised learning

A (supervised) dataset is a set of n examples:

$$S = \{(x_1, y_1), \dots, (x_n, y_n)\}.$$
 (1)

Informally, given a 'new' pair (x,y) not contained in  $\mathcal{S}$ , we want a function  $f(\cdot)$  such that:

$$f(x) \approx y$$
. (2)

More generally, we can test the model on a separate dataset  $\mathcal{T}$  never seen during training, i.e.,  $\mathcal{S} \cap \mathcal{T} = \emptyset$ .

#### Constraints on the dataset

We always assume implicitly that the elements in S and the elements in T are taken from the same i.i.d, unknown distribution p(x,y).

- **Identically distributed**: the data-generating process is stable (e.g., in recognizing cats, the distribution of species do not change).
- **Independently distributed**: there is no bias in the data collection (e.g., we mostly collect siamese cats).

If the distribution between S and T varies, we talk about domain shift.

# Some motivating examples

- **3 Spam identification**:  $x_i$  is an email, and  $y_i$  describes its probability of being spam.
- **2 Robot navigation**:  $x_i$  is a sensory representation of the environment, and  $y_i$  is a motor command.
- **3 Text translation**:  $x_i$  is a text and  $y_i$  its corresponding translation.
- **Product recommendation**:  $x_i$  is a user, and  $y_i$  its affinity w.r.t. a certain catalogue of products.

Note: ensuring the i.i.d. property sometimes is far from trivial!

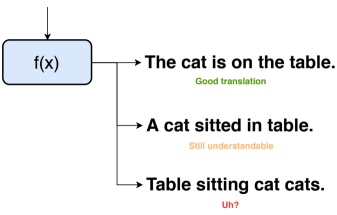


Setup and examples

Loss functions

Building a loss functior

### "Il gatto è sul tavolo."





### Introducing loss functions

Given a point x, a desired value y, and a prediction  $\hat{y} = f(x)$ , we formalize its quality with a loss function  $l(y, \hat{y})$ , such that:

- 1 Low value of loss: good approximation;
- 2 High value of loss: poor approximation.

In this way, learning becomes a problem of minimizing a certain loss quantity that we designed. Importantly, the loss function should be a scalar, vary gradually, and (as we will see) be differentiable.

# Expected risk and empirical risk

The expected loss (risk) of a function f is:

$$f^*(x) = \arg\min\left\{\mathbb{E}_{p(x,y)}\left[l(y,f(x))\right]\right\}. \tag{3}$$

The expected risk is uncomputable, but can be approximated via empirical risk minimization:

$$f^*(x) = \arg\min_{f} \left\{ \frac{1}{n} \sum_{i=1}^{n} l(y_i, f(x_i)) \right\}.$$
 (4)

The gap between the two approaches is called generalization gap.

# Overfitting

Consider the following algorithm (a variation of 1-NN):

$$f(x) = \begin{cases} y & \text{if } (x, y) \in \mathcal{S}, \\ 0 & \text{otherwise}. \end{cases}$$
 (5)

This has 0 training error by construction, but a very large test error: this is an example of overfitting. We will see that (large) neural networks have many counter-intuitive properties when it comes to generalization.

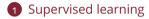
Simone Scardapane Neural Networks 2022/2023 8 / 46

### Some simplifications

To begin our exploration of supervised learning, we will make a few simplifying assumptions:

- The input  ${\bf x}$  is a vector of shape d.
- The output *y* is a single real number.

There are three interesting cases: when  $y \in \mathbb{R}$ , it is a regression task; when  $y \in \{0, \dots, c-1\}$ , it is a classification task; when c=2, it is a binary classification task.



Setup and examples

Loss functions

Building a loss function

#### How to build a loss function?

Building a loss function can be done empirically: considering regression, for example, the prediction error  $(y - \hat{y})$  is a reasonable quantity to penalize.

Since most times we do not care about the sign of the error, these are all valid choices:

Is there a principled way to make the choice?

### Probabilistic formulation

Let us assume that f(x) parameterizes a *probability distribution*  $\hat{p}(y \mid f(x))$  over the possible outputs y. We are now choosing  $\hat{p}(\cdot \mid \cdot)$  instead of  $l(\cdot, \cdot)$ , which can feel more natural. For example, for regression we can assume a Gaussian shape:

$$\hat{p}(y \mid f(x)) = \mathcal{N}(y \mid f(x), \sigma^2), \tag{7}$$

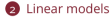
where the model predicts the center of a Gaussian distribution with fixed variance (hyper-parameter).

### Maximum likelihood

The probabilistic formulation also provides a principled way to interpret training by maximizing the likelihood (or log-likelihood) of the model (assuming the elements of the dataset are i.i.d.):

$$f^*(x) = \arg\max_{(x_i, y_i)} \hat{p}(y_i \mid f(x_i)) = \arg\min_{(x_i, y_i)} -\log \hat{p}(y_i \mid f(x_i)).$$
 (8)

Under the Gaussian assumption in (7), we obtain  $-\log \hat{p}(y_i \mid f(x_i)) \propto (y_i - f(x_i))^2$ , i.e., we should optimize the squared loss!



#### Linear models for regression

Solving the LS problem

Overparameterized models and convergence (optional)

Linear models for classification

Calibration and a probabilistic formulation

### What is a linear model?

A linear model f is defined as:

$$f(\mathbf{x}) = \langle \mathbf{w}, \mathbf{x} \rangle = \mathbf{w}^{\top} \mathbf{x} = \sum_{j} w_{j} x_{j},$$
 (9)

where w is a vector of adaptable parameters.

This model is fundamental in many disciplines, ranging from econometrics to statistics.

#### Linear models with bias

A more general formulation considers the inclusion of an offset (bias)  $b \in \mathbb{R}$ :

$$f(\mathbf{x}) = \mathbf{w}^{\top} \mathbf{x} + b. \tag{10}$$

Because we can always rewrite this as  $\mathbf{w}^{\top}\bar{\mathbf{x}}$ , with  $\bar{\mathbf{x}} = [\mathbf{x}; \ 1]$ , we can avoid writing the bias explicitly to simplify the notation.

**Hint**: Everytime we write a linear model, mentally add an offset term whenever needed.

# Graphical representation

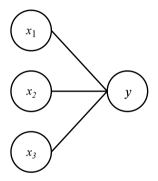


Figure 1: Each arrow represents a *linear* influence on the destination, which sums the results.

### Least-squares cost function

Combining the squared loss with a linear model results in the least-squares optimization problem:

$$LS(\mathbf{w}) = \frac{1}{n} \sum_{i=1}^{n} (y_i - \mathbf{w}^{\top} \mathbf{x}_i)^2.$$
(11)

We can vectorize LS as:

$$LS(\mathbf{w}) = \frac{1}{n} \left\| \mathbf{y} - \mathbf{X} \mathbf{w} \right\|^2, \tag{12}$$

where  $[\mathbf{X}]_i = \mathbf{x}_i$  and  $[\mathbf{y}]_i = y_i$ .



Linear models for regression

#### Solving the LS problem

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### Solving the LS problem

LS is a convex problem, with a simple gradient (normal equations):

$$\nabla \mathsf{LS}(\mathbf{w}) = \frac{2}{n} \mathbf{X}^{\top} (\mathbf{X} \mathbf{w} - \mathbf{y}). \tag{13}$$

LS is special in the sense that  $\nabla LS(\mathbf{w}) = 0$  is a linear equation that can be solved explicitly:

$$\mathbf{w}^* = \left(\mathbf{X}^\top \mathbf{X}\right)^{-1} \mathbf{X}^\top \mathbf{y} = \mathbf{X}^\top \left(\mathbf{X} \mathbf{X}^\top\right)^{-1} \mathbf{y}.$$

$$(14)$$

The terms  $(\mathbf{X}^{\top}\mathbf{X})^{-1}\mathbf{X}^{\top}$  and  $\mathbf{X}^{\top}(\mathbf{X}\mathbf{X}^{\top})^{-1}$  are called the pseudoinverses of  $\mathbf{X}$ , and they require the corresponding matrices to be invertible (full rank).

Simone Scardapane Neural Networks 2022/2023 17/46

### Regularizing the LS problem

Numerical problems in the inversion of  $(\mathbf{X}^{\top}\mathbf{X})$  can be solved by adding a small amount of  $\ell_2$  regularization (ridge regression):

$$LS-REG(\mathbf{w}) = LS(\mathbf{w}) + \frac{\lambda}{2} \|\mathbf{w}\|^2, \tag{15}$$

for some  $\lambda > 0$ . This makes the problem *strictly* convex and forces the solution to be contained in a ball of given radius, modifying the gradient and the explicit solution as:

$$\nabla LS-REG(\mathbf{w}) = \nabla LS(\mathbf{w}) + \lambda \mathbf{w}.$$
 (16)

$$\mathbf{w}^* = \left(\mathbf{X}^{\top} \mathbf{X} + \lambda \mathbf{I}\right)^{-1} \mathbf{X}^{\top} \mathbf{y}. \tag{17}$$

where I is the identity matrix of appropriate shape.

### Show me some code!

#### Generating some data:

```
# Linear model with unknown coefficients
X = torch.randn((10, 5))
y = X @ torch.randn((5, 1))
```

#### Computing a linear model:

```
w = torch.randn((5, 1))
yhat = X @ w # (10, 1)
```

### Computing the objective function:

```
mse = ((y - X @ w)**2).mean()
```

### Show me some code (2)!

#### Explicit solution (numerically unstable):

```
wopt = torch.linalg.inv(X.T @ X) @ X.T @ y
```

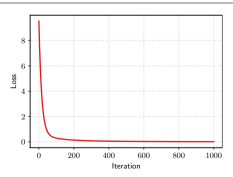
#### Explicit solution (better numerical conditioning):

```
wopt = torch.linalg.solve(X.T @ X, X.T @ y)
```

### Show me some code (3)!

### Simple implementation of gradient descent:

```
1 loss = []
2 for i in range(1000):
3 loss.append(((y - X @ w)**2).mean().item())
4 # Note the sign: the derivative has a minus.
5 w = w + 0.001 * X.T @ (y - X @ w)
```

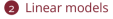


### Computational complexity of LS

Matrix multiplication  $\mathbf{A}$   $\mathbf{B}$  has complexity  $\mathcal{O}(abc)$ , while matrix inversion has complexity

(roughly) cubic.

For large models and datasets, we want algorithms that can scale linearly in both n and d: we will see that gradient descent can satisfy this property.



Linear models for regression

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Calibration and a probabilistic formulation

### Analyzing the convergence

Rewriting the GD step for the LS problem (ignoring the constant factor 2/n):

$$\mathbf{w}_t = \mathbf{w}_{t-1} - \eta \mathbf{X}^{\top} (\mathbf{X} \mathbf{w}_{t-1} - \mathbf{y}). \tag{18}$$

We can use this to write out how the *predictions* over the training set evolve:

$$\hat{\mathbf{y}}_t = \hat{\mathbf{y}}_{t-1} - \eta \mathbf{X} \mathbf{X}^\top \left( \hat{\mathbf{y}}_{t-1} - \mathbf{y} \right) , \tag{19}$$

or, after some manipulation:

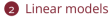
$$(\hat{\mathbf{y}}_t - \mathbf{y}) = \left(\mathbf{I} - \eta \mathbf{X} \mathbf{X}^\top\right) (\hat{\mathbf{y}}_{t-1} - \mathbf{y}) = \left(\mathbf{I} - \eta \mathbf{X} \mathbf{X}^\top\right)^t (\hat{\mathbf{y}}_0 - \mathbf{y}) . \tag{20}$$

See PPA, Chapter 5 for the full analysis.

### Overparameterized models

Eq. (20) is an interesting dynamical system: it will diverge in general, unless (a) all eigenvalues  $\lambda_0, \dots, \lambda_n$  of  $\mathbf{X}\mathbf{X}^{\top}$  are non-negative, and (b)  $\eta \in [0, 1/\lambda_0]$ . In this case, it will decrease to 0. If all eigenvalues are positive, it will do so *exponentially*.

For the latter condition, we need d > n, otherwise the matrix  $\mathbf{X}\mathbf{X}^{\top}$  will have low-rank (hence, some zero eigenvalues). We call this the over-parameterized regime: it is possible in this case for LS to perfectly interpolate all training data exponentially fast!



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### Multi-class classification

In classification, y is an integer  $\{0, \dots, c-1\}$ , such that  $y_i = j$  means that  $\mathbf{x}_i$  is of class j. For example, with c = 3 we might have:

- y = 0: the email is spam;
- y = 1: the email is legit;
- y = 2: the email is dubious.

Solving these as regression tasks is generally not an optimal choice: among other things, it is not guaranteed that classes have a definite ordering.

### Probability distributions

A common solution is to predict a probability distribution over the classes.

A vector  ${\bf a}$  belongs to the probability simplex  $\Delta_c$  if:

(c

$$\sum_{i} [\mathbf{a}]_{i} = 1, \quad [\mathbf{a}]_{i} \ge 0.$$
 (21)

If  $f(\mathbf{x}) = \hat{\mathbf{y}} \in \Delta_c$ , we can interpret it as a categorical probability distribution, e.g., we can select the class with highest probability as:

$$\mathsf{class} = \arg\max_{i} \left[ \hat{\mathbf{y}} \right]_{i}. \tag{22}$$

Simone Scardapane Neural Networks 2022/2023 26 / 46

### A comment on differentiability

Note that we *cannot* easily predict an integer with our models, because it would require some form of threshold operation which is not compatible with gradient descent (gradient zero almost everywhere).

Predicting a probability distribution can be seen as a *soft* approximation to this problem.

#### The softmax function

The softmax function maps any vector to the probability simplex:

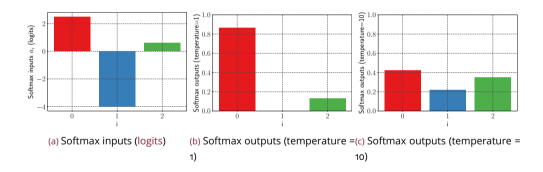
$$[\operatorname{softmax}(\mathbf{a})]_i = \frac{\exp(a_i)}{\sum_j \exp(a_j)}$$
 (23)

The numerator ensures that all outputs are positive, while the denominator ensures that the final vector sums to 1. It can be seen as a soft approximation to the argmax (a better name is in fact *softargmax*).

If desired, we can control the approximation with an additional hyper-parameter  $\tau$  called temperature:

$$[\operatorname{softmax}(\mathbf{a})]_i = \frac{\exp(a_i/\tau)}{\sum_j \exp(a_j/\tau)}$$
 (24)

# Visualizing the softmax function



#### Final model

Our linear model for classification becomes:

$$f(\mathbf{x}) = \operatorname{softmax}(\mathbf{W} \cdot \mathbf{x})$$

$$(c) \qquad (c,d) \qquad (d)$$
(25)

The pre-softmax values  $\mathbf{W}\mathbf{x}$  are called the logits of the model.

### One-hot encoding

In order to compare the predictions with the ground truth, we encode our targets using a one-hot encoding. Given a pair (x, y):

$$y_i = \begin{cases} 1 & \text{if } \mathbf{x} \text{ is of class } i, \\ 0 & \text{otherwise}. \end{cases}$$
 (26)

For example, with 3 classes {cat, dog, other}:

$$cat = [1, 0, 0] \quad dog = [0, 1, 0] \quad other = [0, 0, 1].$$
 (27)

This is a probability distribution putting all the **mass** on a single class.

#### Cross-entropy loss

Finally, we need a loss function l to compare two probability distributions.

The cross-entropy loss is defined for two vectors  $\mathbf{y}, \hat{\mathbf{y}} \in \Delta_c$  as:

$$\mathsf{CE}(\mathbf{y}, \hat{\mathbf{y}}) = -\sum_{i} y_{i} \log \left(\hat{y}_{i}\right) . \tag{28}$$

The CE loss can be derived from the maximum likelihood principle under the assumption that  $f(\mathbf{x})$  encodes a categorical distribution (try it!).

### Logistic regression

A logistic regression is a linear model  $f(\mathbf{x}) = \mathsf{softmax}(\mathbf{W}\mathbf{x})$  trained by optimizing the cross-entropy:

$$LR(\mathbf{W}) = \frac{1}{n} \sum_{i=1}^{n} CE(\mathbf{y}_i, f(\mathbf{x}_i)).$$
 (29)

It is not possible to solve the logistic regression problem explicitly. A linear model for classification has dc parameters.

#### Binary classification

A special case is binary classification, where c=2. In this case, we can predict a single scalar value  $f(\mathbf{x}) \in [0,1]$  since:

$$f(\mathbf{x})$$
 probability of class 1, (30)

$$1-f(\mathbf{x})$$
 probability of class 2 . (31)

The softmax function simplifies to the sigmoid function:

The sigmoid  $\sigma(s) \in [0,1]$  is defined as:

$$\sigma(s) = \frac{1}{1 + \exp(-s)}.\tag{32}$$

# Visualizing the sigmoid function

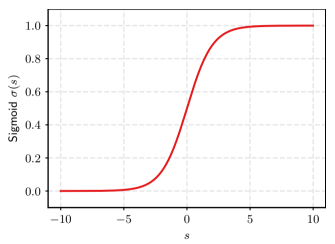


Figure 2: A visualization of the sigmoid function. Note that 0 and 1 are only approached asymptotically.

### Binary logistic regression model

Combining everything, we obtain a binary version of the logistic regression algorithm:

$$BIN-LR(\mathbf{w}) = \frac{1}{n} \sum_{i=1}^{n} \left[ -\underbrace{y_i \log \left( \sigma(\mathbf{w}^{\top} \mathbf{x}) \right)}_{\text{Class 1}} - \underbrace{\left( 1 - y_i \right) \log \left( 1 - \sigma(\mathbf{w}^{\top} \mathbf{x}) \right)}_{\text{Class 2}} \right]$$
(33)

In this case, we can obtain the most probable class from the model as:

$$\mathsf{class} = \begin{cases} 0 & \mathsf{if} \ \sigma(\mathbf{w}^{\top} \mathbf{x}) > 0.5 \,, \\ 1 & \mathsf{otherwise} \,. \end{cases}$$
 (34)

### Gradient of the logistic regression

By manually differentiating we obtain:

$$\sigma'(s) = \sigma(s)(1 - \sigma(s)). \tag{35}$$

Plugging this into the gradient computation we obtain:

$$\nabla \text{BIN-LR}(\mathbf{w}) = \frac{1}{n} \sum_{i=1}^{n} (\sigma(\mathbf{w}^{\top} \mathbf{x}_i) - y_i) \mathbf{x}_i,$$
 (36)

showing its similarity to the regression case.

```
from torch import nn
# Basic one: requires logits in input and indices or one-hot encodings for targets
torch.nn.CrossEntropyLoss()(vtrue, vhat)
# For this one, inputs should be the log of the softmax
torch.nn.NLLLoss()(vtrue, vhat)
# Specialized variants for binary cross-entropy
torch.nn.BCELoss()(ytrue, yhat)
torch.nn.BCEWithLogitsLoss()(ytrue, yhat)
# There are also functional variants
torch.nn.functional.cross_entropy(ytrue, yhat)
# ...
```

### The logsumexp function

Why a variant with logits in input? Note that the ith term of the cross-entropy wrt the logits  $\mathbf{p}$  is given by:

$$-\log\left(\frac{\exp p_i}{\sum_j \exp p_j}\right). \tag{37}$$

In practice, this can be *highly* unstable. However, it can be rewritten as:

$$-p_i + \mathsf{logsumexp}(\mathbf{p})$$
, (38)

where logsumexp is defined as logsumexp( $\mathbf{p}$ ) = log ( $\sum_{i} \exp p_{i}$ ).

https://gregorygundersen.com/blog/2020/02/09/log-sum-exp/

Simone Scardapane Neural Networks 2022/2023 39 / 46

## The logsumexp trick

The reason this is important is that the logsumexp function is invariant in the following sense:

$$\operatorname{logsumexp}(\mathbf{p}) = \log \left( \sum_{i} \exp(p_i - c) \right) + c, \tag{39}$$

where c is an arbitrary constant. By setting  $c = \max{(\mathbf{p})}$ , we can ensure numerical instabilities never occur.

In this sense, softmax can be interpreted as part of the model or as part of the loss; this is not an issue, since  $\arg\max$  only cares about the relative ranking of the values, which is not changed by the softmax.



Linear models for regression

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#### Calibration

A common misconception when doing classification is that  $[f(x)]_i$  can be immediately interpreted as the probability of pattern x being of class i.

However, this is only true whenever the trained model satisfies:

$$p(y = i \mid x) = [f(x)]_i$$
 (40)

We say the model is well calibrated, but this must be checked manually.

Guo, C., et al.. On calibration of modern neural networks. ICML 2017.

Simone Scardapane Neural Networks 2022/2023 41/46

### Measuring calibration

To measure the calibration of a model, we keep a separate validation set, and we split the interval [0,1] into m equispaced bins (each of size 1/m). Define:

- $B_m$  the number of samples from the validation set, whose predicted confidence falls in bin m.
- $p_m$  the average confidence of the network for that bin.
- $a_m$  the average accuracy of the network for these elements.

Then, the expected calibration error (ECE) is given by:

$$\mathsf{ECE} = \sum_{m} \frac{B_m}{n} |a_m - p_m| \,. \tag{41}$$

## Calibration plots

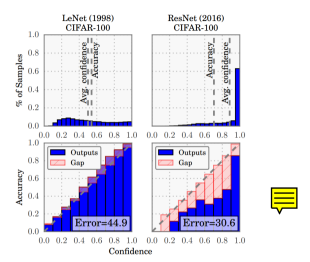


Figure 3: Plotting  $a_m$  against  $p_m$  for every bin gives us a reliability plot (from Guo et al., 2017).

#### **Focal loss**

This topic is important because more complex networks may be highly over (or under) confident, with many methods to improve it (temperature scaling, logit normalization, ...).

A simple (and nonular) ontion is to decrease the weight given to 'easy' samples using a

A simple (and popular) option is to decrease the weight given to 'easy' samples using a variant of cross-entropy call the focal loss:

$$\mathsf{FL}_{\alpha}(\mathbf{y}, \hat{\mathbf{y}}) = -(1 - \hat{\mathbf{y}}_c)^{\alpha} \log \hat{\mathbf{y}}_c, \tag{42}$$

where  $c = \arg \max \mathbf{y}$ .

Simone Scardapane Neural Networks 2022/2023 44/46

Mukhoti, J., et al., 2020. **Calibrating deep neural networks using focal loss**. *Advances in Neural Information Processing Systems*, 33, pp. 15288-15299.

### Visualizing the focal loss

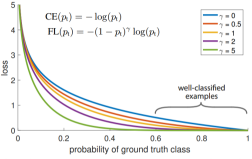


Figure 1. We propose a novel loss we term the *Focal Loss* that adds a factor  $(1-p_t)^{\gamma}$  to the standard cross entropy criterion. Setting  $\gamma>0$  reduces the relative loss for well-classified examples  $(p_t>.5)$ , putting more focus on hard, misclassified examples. As



Lin, T.Y., et al., 2017. Focal loss for dense object detection. In IEEE ICCV (pp. 2980-2988).

# Reading material

• Dive into Deep Learning: Chapters 3 and 4.