Machine Learning from scratch

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Abstract

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4.1 Linear Regression

In statistical modeling, regression analysis consists of a set of statistical processes and techniques for estimating the relationships between a dependent variable (often called the outcome or response variable, or a label in machine learning) and one or more error-free independent variables (often called regressors, predictors, covariates, explanatory variables or features).

The goal of regression is then to predict the value of one or more continuous response variables y given the values of a m-dimensional vector \vec{x} of input (features) variables. Usually we are given a training data set which contains a fixed number N of observations $\{\vec{x_n}\}_{n=1,2,\dots,N}$ together with their corresponding outcome values $\{y_n\}$. The goal is then to predict the value of y for a new given value of $\vec{x} \in \mathbb{R}^m$. For instance, suppose we have a dataset giving the living areas, number of rooms, the district number location and prices of 50 departments in Paris, France:

Living Area (m ²)	Number of Rooms	District	Price (€)
57.45	3	9	319000
47.93	2	15	220000
59.72	3	13	324000
72.85	4	1	411000
46.49	2	7	282000
46.49	2	9	242000
73.69	4	1	392000
61.51	3	12	328000
42.96	2	8	253000
58.14	3	11	327000
43.05	2	19	225000
43.01	2	17	245000
53.63	3	8	300000
21.30	1	3	136000
24.13	1	3	117000
:	:	:	:

We can plot this data to observe the Price according to the Living Area:



Given data like this, we would like to learn to predict the prices of other apartments in Paris, as a function of the input features (Living Area, Rooms, District). Let us first introduce some notation: We set $\vec{x}^{(i)} = (x_1^{(i)}, x_2^{(i)}, x_3^{(i)})$ to be the vector in \mathbb{R}^3 whose entries correspond to the features (Living Area, Rooms, District) of the *i*-th apartment in the training set (see Table). For instance, according to the Table

$$\vec{x}^{(1)} = (57.45, 3, 9)$$

Let $y^{(i)}$ be the output or *target* variable that we are trying to predict; in our case it represents the Price. For instance $y^{(1)} = 319000$. The dataset that we will use to learn consist then of the list of N = 50 training examples:

$$\{(\vec{x}^{(i)}, y^{(i)}) : i = 1, 2, ..., N\}$$

This last list will be then called *training set*.

Remark 4.1. In general when constructing a learning problem we have freedom over which features to use so one may decide to collect more data and include other features. We will say more about feature selection later.

To achieve the goal of predicting the value of y for new given vectors of $\vec{x} \in \mathbb{R}^m$, we will formulate a function $y(\vec{x}, \vec{w})$ whose values are the predictions for new inputs \vec{x} , and where \vec{w} represents a vector of parameters that will be learned from the training set.

The simplest model for the function $y(\vec{x}, \vec{w})$ is the one that consist of a linear combination of the inputs:

(4.2) {{Eq.Sec.Lin.Reg.1}}
$$y(\vec{x}, \vec{w}) = w_0 + w_1 x_1 + \dots + w_m x_m$$

Remark 4.3. The parameter w_0 is usually called a *bias parameter*. The key property of this model is that it is a linear function of the parameters $w_0, ..., w_m$ and also of the input variables $x_1, ..., x_m$.

4.1.1 Non linear models

The class of linear models defined by $\ref{eq:can}$ can be easily extended by considering linear combinations of non-linear functions of the input features \vec{x} . More precisely, suppose $\phi_0, ..., \phi_m : \mathbb{R}^m \longrightarrow \mathbb{R}$ are a collection of *basis* functions. Then, we can consider models of the form:

$$y(\vec{x}, \vec{w}) = w_0 + \sum_{i=1}^{m} w_i \phi_i(\vec{x})$$

For convenience we will set ϕ_0 to be the constant function 1 so that the last equation becomes:

$$(4.4) \quad \{ \text{Eq.Sec.Lin.Reg.2} \} \qquad \qquad y(\vec{x}, \vec{w}) = \sum_{i=0}^m w_i \phi_i(\vec{x}) = \vec{w}^T \phi(x)$$

where
$$\vec{w} = (w_0, ..., w_m)^T$$
 and $\phi = (\phi_0, ..., \phi_m)^T$.

Remark 4.5. Models like **??** are called *linear models* because they are linear in the parameters \vec{w} , nevertheless the model $y(\vec{w}, \vec{x})$ can be a non-linear function of the inputs \vec{x} (it suffices to consider non-linear basis functions). Moreover, notice that if each ϕ_j is defined as the projection onto the j-th coordinate then **??** recovers **??**.

Before deep learning it was common practice in machine learning to use some form of pre-processing of the input variables x, also known as *feature extraction*, expressed in terms of a set of basis functions. The goal was to find a suitable set of basis functions in a way that the resulting learning task could be solved using a simple network model. This is very difficult task. Deep Learning avoids this problem by learning the required nonlinear transformations of the data from the data set itself. To put it in more simpler terms, Deep Learning can help in the task on finding new suitable features that will help to build a more robust and accurate model. This turns out to be very useful for problems about image recognition. We will see more about this in ??. In the following table we provide a list of some common useful choices of the basis functions (assuming a single feature x):

Basis functions $\phi_j(x)$	Model equation $y(x, \vec{w})$	Model nature	
x^{j}	$w_0 + w_1 x + w_2 x^2 + \dots + w_m x^m$	Polynomial	
$\exp\left(-\frac{(x-\mu_j)^2}{2s^2}\right)$	$\sum w_i \exp\left(-\frac{(x-\mu_j)^2}{2s^2}\right)$	Gaussian basis functions	
$\sigma\left(\frac{x-\mu_j}{s}\right)$	$\sum w_i \sigma\left(\frac{x-\mu_i}{s}\right)$	Sigmoidal basis functions	

where σ denotes the sigmoid function defined by

$$\sigma(z) = \frac{1}{1 + \exp(-z)}$$

4.1.2 Maximum likelihood

There are different ways to estimate the parameters \vec{w} for a given model $y(\vec{x}, \vec{w})$ using the dataset $\{(\vec{x}^{(i)}, y^{(i)}) : i = 1, 2, ..., l\}$. Perhaps the most common method consist of finding the parameters \vec{w} that maximize a certain function

called *likelihood function*. To derive such function we must make some assumptions about the relation between the target variables $y^{(i)}$ and the inputs $\vec{x}^{(i)}$.

Let us then assume that the target variable y is given by a deterministic function as in $?? y(\vec{x}, \vec{w})$ with some additive Gaussian noise. This means that the target values and the inputs are related by an equation

(4.6) {{Eq.Sec.Lin.Reg.3}}
$$v = v(\vec{x}, \vec{w}) + \epsilon$$

where ϵ is a random variable that captures either unmodeled effects (*e.g.*, missing important features) or random noise. We assume further that ϵ is a zero-mean Gaussian with variance σ^2 . We can write this last assumption as $\epsilon \sim \mathcal{N}(0, \sigma^2)$. Therefore, the probability density of ϵ is given by

$$p(\epsilon) = \frac{1}{\sqrt{2\pi}\sigma} \exp\left(-\frac{\epsilon^2}{2\sigma^2}\right)$$

By substituting in **??** we then get $p(y|\vec{x}; \vec{w}, \sigma) \simeq \mathcal{N}(y(\vec{x}, \vec{w}, \sigma))$. In other words,

(4.7)
$$p(y|\vec{x}; \vec{w}, \sigma) = \frac{1}{\sqrt{2\pi}\sigma} \exp\left(-\frac{(y - \vec{w}^T \phi(\vec{x}))^2}{2\sigma^2}\right)$$

Finally we can then consider the *design matrix* whose rows correspond to the vectors $\vec{x}^{(i)}$ in the dataset together with the vector or target values \vec{y} , *i. e.*,

There's an error here in this matri one has to includ phi

$$X = \begin{bmatrix} x_1^{(1)} & x_2^{(1)} & x_3^{(1)} & \dots & x_m^{(1)} \\ x_1^{(2)} & x_2^{(2)} & x_3^{(2)} & \dots & x_m^{(2)} \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ x_1^{(N)} & x_2^{(N)} & x_3^{(N)} & \dots & x_m^{(N)} \end{bmatrix}_{N_{YM}} \quad \vec{y} = \begin{bmatrix} y^{(1)} \\ y^{(2)} \\ \ddots \\ y^{(N)} \end{bmatrix}$$

Remark 4.8. Once we start using our dataset we are working under the assumption that each $y^{(i)}$ satisfies a relation as in $\ref{eq:condition}$ and moreover these of the $e^{(i)}$ are themselves independent (and hence also the $y^{(i)}$'s given the $\vec{x}^{(i)}$'s).

From the last assumption we obtain an expression for the likelihood function:

(4.9)
$$L(\vec{w}) = L(\vec{w}; X, \vec{y}) = p(\vec{y}|X, \vec{w}, \sigma^2) = \prod_{i=1}^{N} p(y^{(i)}|\vec{x}^{(i)}, \vec{w}, \sigma)$$

The principle of *maximum likelihood* implies that we should choose the parameters \vec{w} so that the data has the highest possible probability. In other words, we should choose \vec{w} so that the maximum likelihood function $L(\vec{w})$ is maximum. To simplify calculations we can, equivalently, maximize the *log likelihood* $\ell(\vec{w}) = \ln(L(\vec{w}))$. A simple calculation shows that

(4.10)
$$\ell(\vec{w}) = \sum_{i=1}^{N} \ln \left(\frac{1}{\sqrt{2\pi}\sigma} \exp\left(-\frac{(y^{(i)} - \vec{w}^T \phi(\vec{x}^{(i)}))^2}{2\sigma^2} \right) \right)$$

(4.11)
$$= -\frac{N}{2}\ln(2\pi) - \frac{N}{2}\ln(\sigma^2) - \frac{1}{\sigma^2} \cdot \underbrace{\frac{1}{2}\sum_{i=1}^{N}(y^{(i)} - \vec{w}^T\phi(\vec{x}^{(i)}))^2}_{J(\vec{w})}$$

The first two quantities in the last equality are constant and therefore, maximizing the log likelihood $\ell(\vec{w})$ is equivalent to minimizing the *sum-of-squares error function*, also called the *cost function* $J(\vec{w})$:

(4.12) {{Eq.Sec.Lin.Reg.4}}
$$J(\vec{w}) = \frac{1}{2} \sum_{i=1}^{N} (y^{(i)} - \vec{w}^T \phi(\vec{x}^{(i)}))^2$$

We will observe two ways of minimizing the cost function *J*.

4.1.3 Gradient Descent and LMS algorithm

Recall we want to find the parameters \vec{w} so that the cost function $J(\vec{w})$ is minimized. To do this we can use the *gradient descent* algorithm. The idea is to start with an initial guess, say, $\vec{w} = (w_0, w_1, ..., w_m)$ and then iteratively update the parameters using the following rule:

(4.13) {{Eq.Sec.Lin.Reg.5}}
$$\vec{w} \leftarrow \vec{w} - \eta \nabla J(\vec{w})$$

where η is a (usually small) positive number called the *learning rate* and $\nabla J(\vec{w})$ is the gradient of the cost function $J(\vec{w})$. The gradient is a vector whose components are given by the partial derivatives of J with respect to each parameter w_i :

$$\text{(4.14)} \quad \text{\{Eq.Sec.Lin.Reg.6\}\}} \; \nabla J(\vec{w}) = \begin{bmatrix} \frac{\partial J}{\partial w_0} \\ \frac{\partial J}{\partial w_1} \\ \vdots \\ \frac{\partial J}{\partial w_m} \end{bmatrix} = - \begin{bmatrix} \sum_{i=1}^N (y^{(i)} - \vec{w}^T \phi(\vec{x}^{(i)})) \phi_0(\vec{x}^{(i)}) \\ \sum_{i=1}^N (y^{(i)} - \vec{w}^T \phi(\vec{x}^{(i)})) \phi_1(\vec{x}^{(i)}) \\ \vdots \\ \sum_{i=1}^N (y^{(i)} - \vec{w}^T \phi(\vec{x}^{(i)})) \phi_m(\vec{x}^{(i)}) \end{bmatrix}$$

By replacing in **??** and grouping all coordinates we obtain the following update vector rule for the parameters:

(4.15)
$$\vec{w} \leftarrow \vec{w} + \eta \cdot \sum_{i=1}^{N} (y^{(i)} - \vec{w}^T \phi(\vec{x}^{(i)})) \phi(\vec{x}^{(i)})$$

This rule is called the *Least Mean Squares update rule*, sometimes also referred as (LMS) algorithm. The LMS algorithm is a stochastic gradient descent algorithm that updates the parameters \vec{w} using the gradient of the cost function $J(\vec{w})$ at each iteration. The learning rate η controls the step size of the update and can be adjusted to improve convergence. The full pseudo code for the LMS algorithm reads as follows:

Algorithm 1 Least Mean Squares (LMS) Algorithm

Require: Training set $\{(\vec{x}^{(i)}, y^{(i)})\}_{i=1}^{N}$, learning rate η **Require:** Feature functions ϕ_i , j = 0, 1, ..., m

- 1: Initialize $\vec{w} \leftarrow \vec{0}$ or random values
- 2: repeat
- 3: $\vec{w} \leftarrow \vec{w} + \eta \sum_{i=1}^{N} (y^{(i)} \vec{w}^T \phi(\vec{x}^{(i)})) \phi(\vec{x}^{(i)})$
- 4: **until** convergence or maximum iterations **return** \vec{w}

Remark 4.16. This method is called *batch gradient descent* because it uses the entire training set to compute the gradient at each iteration. In practice, we can also use *stochastic gradient descent* (SGD) which updates the parameters using only one training example at a time. This is done by replacing the sum in ?? by a single term $(y^{(i)} - \vec{w}^T \phi(\vec{x}^{(i)}))\phi(\vec{x}^{(i)})$ and iterate over each *i*. The SGD algorithm is usually faster than batch gradient descent and can be used for large datasets.

Finally notice also that the LMS algorithm is a special case of the more general *gradient descent* algorithm. The main difference is that in the LMS algorithm we use the gradient of the cost function $J(\vec{w})$ to update the parameters \vec{w} , while in gradient descent we can use any function to update the parameters. In general gradient descent is susceptible to capture local minima of a function, while the LMS algorithm will capture one global minimum. This is because the cost function $J(\vec{w})$ is a convex function, which means that it has only one global minimum. In other words the LMS will always converge to the same solution regardless of the initial guess \vec{w} .

4.1.4 The Normal Equations

The LMS algorithm is a very powerful method to minimize the cost function $J(\vec{w})$ but it can be slow to converge. In some cases, we can find the optimal solution for the parameters \vec{w} in closed form. This is done by setting the gradient of the cost function $J(\vec{w})$ to zero and solving for \vec{w} . This gives us the so-called normal equations:

4.1.5 Regularization

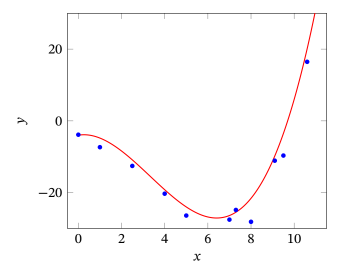
Let us consider a linear model (of polynomial nature) of the form:

$$======>>>>>> bb4ffa3884f86279089af08ae9896dc9ccffb965y(x,\vec{w}) = \sum_{j=0}^{m} w_j \phi_j(x) = w_0 + w_1 x + w_2 x^2 + \dots + w_n x^n + w_n x^$$

where $\phi_j(x) = x^j$. Notice that we are considering a single feature x and in this case m is no longer the number of features but the degree of the polynomial $y(x, \vec{w})$. Sometimes this kind of model is also referred as a model having polynomial features $x, x^2, x^3, ..., x^m$. We will stick to the first terminology for simplicity.

In this situation we want to use the polynomial model given by \ref{star} to fit or approximate a dataset $\{(x^{(i)},y^{(i)}): i=1,...,N\}$. Notice that we are not using the notation $\vec{x}^{(i)}$ because we are only using one feature x. The degree of the polynomial model now becomes a hyperparameter of the model. That is, our goal now is to find the parameters \vec{w} together with the degree m such that the polynomial $y(\vec{w},x)$ is a good approximation of the dataset and such that the model will give suitable predictions for new data. The choice of the degree of the polynomial the becomes important. A high degree polynomial may fit well the dataset but it may not generalize well to new data. This situation is usually know as *overfitting*. Similarly, a low degree polynomial may not fit well the dataset but it may generalize well to new data. This is known as *underfitting*.

In the following figure we show four examples of the results of fitting polynomials on some given data points. The polynomials have degrees 1,2,3,4,5 and 7. The data points are shown in blue and the fitted polynomials are shown in red. The first two polynomials (degree 1 and 2) are underfitting the data, while the last two polynomials (degree 5 and 7) are overfitting the data. The polynomial of degree 3 is a good fit for the data.



4.2 Python Implementation

This section is dedicated to the implementation in python of the LMS algorithm via some problems and exercises taken from Andrew Ng CS229 course at Stanford University. We will use the numpy library which is a powerful library for numerical computing in Python, and the pandas library, a powerful library for data manipulation and analysis. We will also use the matplotlib library to plot the data and the results. In some other problems we will use the scikit-learn library which is a powerful library for machine learning in Python. These notes will contain the main functions and routines in Python. Additional code used to generate the figures and plots will be provided in the Implementations folder of the course repository.

4.2.1 Cost, Gradient and Gradient Descent

Cost function

Let us first implement a function that computes the cost function. Remember that the cost function is given by:

(4.18) {{Eq.Sec.Lin.Reg.8}}
$$J(\vec{w}) = \frac{1}{2} \sum_{i=1}^{N} (y^{(i)} - \vec{w}^T \phi(\vec{x}^{(i)}))^2$$

We will assume that the basis functions ϕ_J are given by projections onto the *j*-th coordinate, *i. e.* $\phi_j(\vec{x}) = x_j$. So that

$$\phi(\vec{x}^{(i)}) = \begin{bmatrix} 1 & x_1^{(i)} & x_2^{(i)} & \cdots & x_m^{(i)} \end{bmatrix}^T$$

In this case we can write the cost function as in ?? as follows:

(4.19) {{Eq.Sec.Lin.Reg.9}}
$$J(\vec{w}) = \frac{1}{2} \sum_{i=1}^{N} (y^{(i)} - \vec{w}^T \vec{x}^{(i)})^2$$

where $\vec{x}_0^{(i)} = 1$ for all i = 1, ..., N. The code to compute the cost function for a single training example \vec{x} and target value y is then given by:

{code:compute_

```
1def compute_cost(X,y,w):
   Compute the cost function for logistic regression.
   Parameters:
  X: numpy array (N,m). Data matrix where N is the number of samples and m is the
     number of features.
  y: numpy array (N,). Vector or target values.
  w: numpy array (M+1,). Weights vector including the bias term w 0.
10
   Cost value : float.
11
12
  N,m = X.shape
  X = np.hstack((np.ones((N, 1)), X)) # Add dummy feature 1 term to X
  w = w.reshape(-1, 1) # Ensure w is a column vector
  y = y.reshape(-1, 1) # Ensure y is a column vector
   cost = (1/2)*np.sum((y - X@w)**2)
  return cost
```

Listing 1: Compute cost

Along all these notes the common conventions and practices will be to use vectorized or Linear algebra notation. In the last code snippet, we start by append to the left a column of ones to the data matrix X. This is done to include the bias term w_0 in the model. Next we ensure that both the weights vector \vec{w} and the outputs vector \vec{y} are both seen as column vectors. The compute_cost function takes \vec{w} and \vec{y} as 1D numpy arrays which are not the same as $N \times 1$ and $(m+1) \times 1$ vectors. Next, we can compute the cost function by simply computing the vector difference y - X@w:

$$\begin{bmatrix} y^{(1)} \\ y^{(2)} \\ \vdots \\ y^{(N)} \end{bmatrix} - \begin{bmatrix} 1 & x_1^{(1)} & x_2^{(1)} & x_3^{(1)} & \dots & x_m^{(1)} \\ 1 & x_1^{(2)} & x_2^{(2)} & x_3^{(2)} & \dots & x_m^{(2)} \\ \vdots & \vdots & \vdots & \vdots & \ddots & \vdots \\ 1 & x_1^{(N)} & x_2^{(N)} & x_3^{(N)} & \dots & x_m^{(N)} \end{bmatrix} \cdot \begin{bmatrix} w_0 \\ w_1 \\ \vdots \\ w_m \end{bmatrix}$$

Finally, the function simply uses the np.sum function to compute the sum of the squares of the differences. The np.sum function is a vectorized function that computes the sum of all the elements of an array much faster than a for loop.

Gradient

The next step is to implement the gradient descent algorithm. Recall that we have to update the parameters \vec{w} using the gradient of the cost function $J(\vec{w})$. First, we create a function to compute the gradient:

{code:compute_

```
Returns:
Gradient: numpy array (m+1,) containing at each entry the gradient with respect to the parameters w.

"""
N,m = X.shape
X = np.hstack((np.ones((N, 1)), X)) # Add dummy feature 1 term to X
w = w.reshape(-1, 1) # Ensure w is a column vector y = y.reshape(-1, 1) # Ensure y is a column vector gradient = - (y-X@w).T @ X

return gradient.flatten() # Return as a flat array for consistency with w shape

Listing 2: Compute Gradient
```

Remark 4.20. The expression in line 17 of the code snippet is a vectorized version of the expression in **??.** Formally we have a matrix product of $(\vec{y} - X \cdot \vec{w})^T$ with X. The former expression is a 1xN vector and the latter is a Nx(m+1) matrix. The result is a 1x(m+1) vector containing the partial derivatives of the cost function with respect to each parameter w_i or in other words, the gradient of the cost function.

LMS algorithm

Finally we proceed to implement the LMS algorithm using gradient descent to minimize the sum-of-squares error function. We will use the compute_cost and compute_gradient functions to compute the cost and the gradient at each iteration.

{code:gradient_

```
idef gradient_descent(X, y, w_in, cost_function, gradient_function,eta=0.01,
    num_iter=1000):
   Batch gradient descent algorithm for linear regression. Updates the weights w by
     taking num_iters gradients steps with learning rate eta
   Parameters:
5
6 X (ndarray (N,m))
                       :Data matrix with N samples and m features.
y (ndarray (N,)) : Vector of target values.
  w_in (ndarray (m+1,)) : Initial weights vector including the bias term w_0.
  cost_function : Function to compute the cost.
  gradient_function : Function to compute the gradient.
n eta (float) : Learning rate.
  num_iter (int)
                     : Maximum number of iterations.
  tol (float)
                    :Tolerance for convergence.
  Returns:
15
   w (ndarray (m+1,)) : Final weights vector.
    cost_history (list): Cost history during training.
  N,m = X.shape
19
   X = np.hstack((np.ones((N, 1)), X)) # Add dummy feature 1 term to X
20
   w = copy.deepcopy(w_in).reshape(-1, 1) # Ensure w is a column vector
   y = y.reshape(-1, 1) # Ensure y is a column vector
   cost_history = [] # To store the cost at each iteration for plotting or analysis
24
  for i in range(num_iter):
26
   # Compute gradient using
```

```
gradient = gradient_function(X,y,w)
w -= eta * gradient

# Save the cost at each iteration
cost_history.append(cost_function(X,y,w))

# Print cost and gradient information every 10 times or as many as num_iters
if less than 10

if i% math.ceil(num_iter / 10) == 0:
    print(f"Iteration {i:4d}: Cost = {cost_history[-1]:8.4f}, Gradient Norm = {
    np.linalg.norm(gradient):8.4f}")
return w, cost_history
```

Listing 3: Gradient Descent

Observation 4.21. It is common practice to consider an average cost function, i. e., we can divide the cost function by N in $\ref{eq:second}$? and obtain:

(4.22) {{Eq.Sec.Lin.Reg.10}}
$$J(\vec{w}) = \frac{1}{2N} \sum_{i=1}^{N} (y^{(i)} - \vec{w}^T \vec{x}^{(i)})^2$$

This is done to make the cost function independent of the number of training examples. Of course this has no effect in finding the minimum as N is a constant. However dividing by N makes our calculations more consistent, especially when working with big data sets. In this case we will have to divide the gradient by N as well. The gradient then turns out to be an "average gradient" which again, makes updates more consistent regardless of the dataset size. We have not done this in the code snippet for simplicity but the reader is encouraged to do so.

4.2.2 House price prediction

Let us go back to our initial motivating example and use our previous implementation of the LMS algorithm. We will use a dataset of house prices stored in a txt file called houses.txt. The dataset contains prices of 100 houses with 4 features (size, number of bedrooms, floors and age). This dataset was derived from the *Ames Housing dataset* which is a popular dataset for regression tasks. The dataset is available at https://www.kagqle.com/datasets/shashanknecrothapa/ames-housing-dataset.

Remark 4.23. Each of the problems or practical exercises will have a separate Jupyter notebook where the reader can observe and play around with the code. The reader is encouraged to use the Jupyter notebooks to run the code and play around with the parameters. The Jupyter notebooks also are available in the Implementations folder of the course repository.

The first four rows of the dataset are shown below:

{tab:house_data

Table 1: House Data Sample

Index	Size (sqft)	Number of bedrooms	Number of floors	Age	Price (in 1000's)
0	952.0	2.0	1.0	65.0	271.5
1	1244.0	3.0	1.0	64.0	300.0
2	1947.0	3.0	2.0	17.0	509.8
3	1725.0	3.0	2.0	42.0	394.0

We can have a better understanding of our dataset by plotting each of the features against the target variable (price):

5 Classification and Logistic Regression

- 5.1 Logistic Regression
- 5.1.1 The perceptron algorithm
- 5.2 Python Implementation
- 6 Generalized Linear Models

Part III

Deep Learning

- 7 Neural Networks
- 8 Convolutional Neural Networks
- 9 Generative Adversarial Networks (GANs)
- 10 Transformers

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10.1 Natural language processing (NLP): Word Embeddings

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10.2 Natural language processing (NLP): word embeddings

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10.3 Attention mechanism

The fundamental idea behind a transformer model is the *attention* mechanism, which allows the model to focus on different parts of the input sequence when making predictions. This mechanism arose from the need to improve the performance of recurrent neural networks (RNNs) for machine translation tasks [BCB16]. Later on, performance was improved considerably by eliminating the RNN architecture altogether and using a fully attention-based architecture, which is the basis of the transformer model [Vas+17].

«««< HEAD Let us consider the following three sentences as an example:

I need to **run** to catch the bus! Paul decided to **run** for president. We had a **run** of bad luck.

In each case, the word *run* has a different meaning depending on the context. The attention mechanism allows the model to focus on the surrounding words to determine the meaning of *run* in each case. For example, in the first sentence, the model can pay more attention to the words *catch* and *bus*, while in the second sentence, it can focus on *Paul* and *president*.

10.3.1 Processing

The input data to a transformer is a collection of vectors $\{\vec{x}^{(i)}\}$ in \mathbb{R}^m where i=1,...,N. As it is usual in these notes, each element of the i-th vector $\vec{x}^{(i)}_j$ is called a *feature* and the data vectors are called *tokens*. These tokens may correspond to words withing a corpus of text, to a patch of pixels within an image, or to any other type of data sensible to be represented (embedded) as a vector. We will associate a matrix $X \in \mathbb{R}^{N \times m}$ to the collection of vectors $\{\vec{x}^{(i)}\}$ as follows:

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(10.1)
$$X = \begin{bmatrix} - & (\vec{x}^{(1)})^T - - \\ - & (\vec{x}^{(2)})^T - - \\ \vdots \\ - & (\vec{x}^{(N)})^T - - \end{bmatrix}$$

The fundamental block of the Transformer will take the matrix X as input and create a new matrix, \hat{X} of the same size:

$$\widetilde{X} = \operatorname{TransformerLayer}(X)$$

The idea is to create a new matrix \widetilde{X} that contains the same information as X, but with the features of each token enhanced by the attention mechanism. We can of course stack (compose) several of these layers in sequence to create a deeper model capable of learning more complex relationships between the tokens. This single transformer layer has two stages: the one acting on columns (features) corresponding to the attention mechanism, and the one acting on rows (tokens) which corresponds to the effect of transforming the features within each token.

Attention

Let us denote by $\vec{y}^{(1)}$, ..., $\vec{y}^{(N)}$ the rows (tokens) of the matrix \widetilde{X} . Each of these tokens should live in an embedding space with a richer semantic structure than the tokens of X. Since each token in X correspond to some data type (say words) and we want to capture some semantic relation between them, each vector $\vec{y}^{(i)}$ should depend on all the tokens from X, i. e, $\vec{x}^{(1)}$, $\vec{x}^{(2)}$, ..., $\vec{x}^{(N)}$. The simplest thing to do is to assume that each $\vec{y}^{(i)}$ depends linearly, or it is linear combination of the tokens in X:

$$\vec{y}^{(i)} = \sum_{i=1}^{N} a_{ij} \vec{x}^{(j)}$$

where α_{ij} are the coefficients of the linear combination. These coefficients will be called *attention* weights. We expect these coefficients to be close to zero whenever the input tokens are not relevant to the output token. For instance, in our previous example: "I need to run to catch the bus", the attention weights for the output token associated with the word catch should be high for the words catch, to (second), bus and run for we need to focus on the object of the action (the bus) and the preceding action (run) to understand the meaning of catch. On the other hand, the attention weights should be low for the words I and need and the first to.

In the following table we present some linguistic intuition about how the weights should be distributed. The first column contains the words of the sentence, the second and third columns contain notation for the input and output tokens, respectively. The fourth column contains the words that should receive low attention weights, while the fifth column contains the words that should receive high attention weights.

We will then impose the following constraints on the attention weights:

- ➤ The attention weights are non-negative: $a_{ij} \ge 0$, as we want to avoid situations in which one coefficient can become large and positive while another one compensated by being large and negative. This is not desirable in our case, as we want to focus on the most relevant tokens.
- ▶ The attention weights sum to one: $\sum_{j=1}^{N} a_{ij} = 1$. This is a normalization condition that ensures that if an output token pays more attention to one input token, it pays less attention to the others.

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{tab:attention ir

Word	Input Token	Output Token	Low Attention	High Attention
I	x^1	y^1	the, bus, catch	I, need, run
need	x^2	y^2	the, bus	need, I, to (first), run
to	x^3	y^3	the, bus, I, catch	to (first), need, run
run	x^4	y^4	I, need	run, to (first), to (second), catch, bus
to	x^5	y ⁵	I, need, to (first)	to (second), run, catch, bus
catch	x^6	y ⁶	I, need, to (first)	catch, to (second), run, the, bus
the	x^7	y^7	I, need, to (first), run	the, bus, catch
bus	x^8	y^8	I, need	bus, the, catch, run

Remark 10.2. Notice that if we assume that the outputs are instead linear combinations of basis functions of the input tokens $\phi_1, ..., \phi_N : \mathbb{R}^m \to \mathbb{R}$, the two conditions above ensure that the basis functions form a partition of unity.

Self-attention

Let us discuss how to determine the attention weights a_{ij} . The idea first is to use an approach similar to the one used in problems related with information retrieval. The following image shows the basic idea used to find the attention weights:

The main idea is to see each of the input vectors $\vec{x}^{(i)}$ as a value vector that will be used to create the output tokens. We will also use $\vec{x}^{(i)}$ as the key vector for the ith input token. Finally we consider each $\vec{x}^{(j)}$ as query vector for the output $\vec{y}^{(j)}$. To achieve the constraints on the attention weights, we will use a softmax function (with no probabilistic interpretation) so that:

$$a_{ij} = \frac{\exp((\vec{x}^{(i)})^T \cdot \vec{x}^{(j)})}{\sum_{k=1}^{N} \exp((\vec{x}^{(i)})^T \cdot \vec{x}^{(k)})}$$

By grouping all the output tokens in a single $N \times m$ matrix Y we obtain a nice matrix formula for the output tokens:

(10.3) {{Eq.subsec.self_attention.1}}
$$Y = Softmax(XX^T)X$$

where the softmax function applied on a $N \times N$ matrix $C = [c_{ij}]$ is a new matrix whose entries given by:

$$Softmax(C)_{ij} = \frac{\exp(c_{ij})}{\sum_{k=1}^{N} \exp(c_{ik})}$$

This process is called *self-attention* because the same input tokens are used as queries, keys and values. The attention weights are computed by taking the dot product of the query vector with the key vectors, and then applying the softmax function. We will see some variations of this later on.

Maybe we have to add some more details here. So far it is not that important, just intuition we get from this

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Network parameters

So far, the transformation we have described to find the output tokens is fixed in the sense that there are no adjustable parameters and therefore this has no learning capacity from data. We would like to build a network that has some flexibility to choose features to focus on when determining the output tokens. For this, we can start by defining modified feature vectors via a linear transformation to the input tokens through a matrix *U* of learnable parameters:

$$\hat{X} = XU$$

where U is an $m \times m$ matrix of learnable weight parameters. Notice that this is analogous to a linear layer in a neural network. By replacing X by \hat{X} in (??) we obtain the following expression for the output tokens:

$$Y = \operatorname{Softmax}(\widehat{X}\widehat{X}^{\mathsf{T}})\widehat{X}$$
 (10.4) {{Eq.subsec.self_attention.2}}
$$= \operatorname{Softmax}(XUU^{\mathsf{T}}X^{\mathsf{T}})XU$$

Remark 10.5. This approach has one remarkable characteristic. The matrix $\widehat{X}\widehat{X}^{\mathsf{T}}$ is symmetric which will in turn imply a symmetric behavior in the attention mechanism. We need much more flexibility, for instance, many tasks in NLP require tokens (words) to be strongly associated with other tokens but not in the opposite sense: The word *hardware* is strongly associated with the word *computer* but the latter may be associated with many other words and its association with the word *hardware* may not be that strong.

To overcome this limitation we still use the same idea but with independent learnable weight matrices for the query, key and value vectors. We will denote these matrices by W_q , W_k and W_v respectively. We then consider:

$$Q = XW_q$$
 $\dim W_q = m \times m_k$
 $K = XW_k$ $\dim W_k = m \times m_k$
 $V = XW_v$ $\dim W_v = m \times m_v$

where m_k and m_v are the dimensions of the key and value vectors respectively. The dimensions are chosen so that we can perform dot products between the query and key vectors (a typycal choice is to take $m_k = m$). Additionally, m_v will govern the dimension of the output tokens. Finally we obtain an expression for the output tokens as follows:

(10.6) {{Eq.subsec.self_attention.3}}
$$Y = Softmax(QK^T)V$$

Remark 10.7. To avoid

10.4 Transformer models

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10.5 Transformer models

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References

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