# LxMLS - Lab Guide

July 27, 2016

# Day 0

# **Basic Tutorials**

# 0.1 Today's assignment

In this class we will introduce several fundamental concepts needed further ahead. We start with an introduction to Python, the programming language we will use in the lab sessions, and to Matplotlib and Numpy, two modules for plotting and scientific computing in Python, respectively. Afterwards, we present several notions on probability theory and linear algebra. Finally, we focus on numerical optimization. The goal of this class is to give you the basic knowledge for you to understand the following lectures. We will not enter in too much detail in any of the topics.

# 0.2 Manually Installing the Tools in your own Computer

#### 0.2.1 Desktops vs. Laptops

If you have decided to use one of our provided desktops, all installation procedures have been carried out. You merely need to go to the lxmls-toolkit-student folder inside your home directory and start working! You may go directly to section 0.3. If you wish to use your own laptop, you will need to install Python, the required Python libraries and download the LXMLS code base. It is important that you do this as soon as possible (before the school starts) to avoid unnecessary delays. Please follow the install instructions.

#### 0.2.2 Downloading the labs version from GitHub

The code of LxMLS is available online at GitHub. There are two branches of the code: the master branch contains fully functional code. The student branch contains the same code with some parts deleted, which you must complete in the following exercises. To download the student code, go to

```
https://github.com/LxMLS/lxmls-toolkit
```

and select the student branch in the dropdown menu. This will reload the page to the corresponding branch. Now you just need to click the Clone or download button to obtain the lab tools in a zip format:

```
lxmls-toolkit-student.zip
```

After this you can unzip the file where you want to work and enter the unzipped folder. This will be the place where you will work. **Important:** Be careful to select the student branch and do not install the lxmlstoolkit (using setup.py).

#### 0.2.3 Installing Python 2.7 from Scratch

If you are new to Python the best option right now is the Anaconda platform. You can find installers for Windows, Linux and OSX platforms here

```
https://www.continuum.io/downloads
```

**Important**: Be sure to install the version for Python 2.7. This installer will already provide most of the packages you need to work with the school toolkit. The only package missing will be Theano, you may follow the instructions here to install it

```
http://deeplearning.net/software/theano/install.html#anaconda
```

Note that you just need a normal Theano installation. You do not need the bleeding edge version and it is not recommended to install it.

#### 0.2.4 (Advanced Users) Installing with Pip

If you are familiar with Python you will probably be used to the pip package installer. In this case it might be more easy for you to install the packages yourself using pip

```
pip install -r pip-requirements.txt
```

You may need to pre-append sudo to the system call depending on the platform. this will install the following packages

```
jupyter
numpy
scipy
matplotlib
pyyaml
nltk
theano
```

To check that all the modules we will use work, please run the all\_imports.py file found in the lxmls-toolkit-student folder.

#### 0.2.5 (Advanced Users) Virtual Environments

If you are an experienced Python user and you do not want to add many packages to your existing installation, you can use virtualenv.

```
sudo pip install virtualenv
virtualenv venv
```

This will create a local installation of Python on your work folder. To activate it use

```
source venv/bin/activate
```

then you can install packages locally with no need for root privileges and without touching your original installation.

```
pip install -r pip-requirements.txt
```

Each time you want to work on the school code, you will have to activate the environment. To exit the environment use

```
deactivate
```

If you want to make use of the modules already installed in your main Python installation use

```
virtualenv venv --system-site-packages
```

instead of the conventional call. Bear in mind that using virtual environments can create some problems. For once, the Python binary will be in another location. You will therefore have to be careful when using e.g. shebangs to specify it. The Jupyter-notebook is also known to give problems unless you also install it with the virtual environment. For these reasons, virtual environments are only recommended for students that feel comfortable with Python.

#### 0.2.6 (Advanced Users) Forking and cloning the code on GitHub

It might be the case that you feel very comfortable with scientific Python, know some git/GitHub and want to extend/improve our code base. In that case you can directly clone the project with

```
git clone https://github.com/LxMLS/lxmls-toolkit.git
```

and work from there. Remember to work on the *student* branch!. If you want to contribute to the code-base, you can make pull requests to the *develop* branch or raise issues.

#### 0.2.7 Deciding on the IDE and interactive shell to use

An Integrated Development Environment (IDE) includes a text editor and various tools to debug and interpret complex code. **Important:** As the labs progress you will need an IDE, or at least a good editor and knowledge of pdb/ipdb. This will not be obvious the first days since we will be seeing simpler examples.

As IDE we recommend PyCharm, but feel free to use the software you feel more comfortable with. Py-Charm and other well known IDEs like Spyder are provided with the Anaconda installation.

Aside of an IDE, you will need an interactive command line to run commands. This is very useful to explore variables and functions and quickly debug the exercises. For the most complex exercises you will still need an IDE to modify particular segments of the provided code. As interactive command line we recommend the Jupyter notebook. This also comes installed with Anaconda and is part of the pip-installed packages. The Jupyter notebook is described in the next section. In case you run into problems or you feel uncomfortable with the Jupyter notebook you can use the simpler iPython command line.

## 0.2.8 Jupyter Notebook

Jupyter is good choice when writing Python code. It is an interactive computational environment for data science and scientific computing, where you can combine code execution, rich text, mathematics, plots and rich media. The Jupyter Notebook is a web application that allows you to create and share documents, which contains live code, equations, visualizations and explanatory text. It is very popular in the areas of data cleaning and transformation, numerical simulation, statistical modeling, machine learning and so on. It supports more than 40 programming languages, including all those popular ones used in Data Science such as Python, R, and Scala. It can also produce many different types of output such as images, videos, LaTex and JavaScript. More over with its interactive widgets, you can manipulate and visualize data in real time.

The main features and advantages using the Jupyter Notebook are the following:

- In-browser editing for code, with automatic syntax highlighting, indentation, and tab completion/introspection.
- The ability to execute code from the browser, with the results of computations attached to the code which generated them.
- Displaying the result of computation using rich media representations, such as HTML, LaTeX, PNG, SVG, etc. For example, publication-quality figures rendered by the matplotlib library, can be included inline.
- In-browser editing for rich text using the Markdown markup language, which can provide commentary for the code, is not limited to plain text.
- The ability to easily include mathematical notation within markdown cells using LaTeX, and rendered natively by MathJax.

The basic commands you should know are

Esc	Enter command mode	
Enter	Enter edit mode	
up/down	Change between cells	
Ctrl + Enter	Runs code on selected cell	
Shift + Enter	Runs code on selected cell, jumps to next cell	
restart button	Deletes all variables (useful for troubleshooting)	

Table 1: Basic Jupyter commands

A more detailed user guide can be found here:

http://jupyter-notebook-beginner-guide.readthedocs.io/en/latest/index.html

# 0.3 Solving the Exercises

In the student branch we provide the solve.py script. This can be used to solve the exercises of each day, e.g.

```
python ./solve.py day1
```

You can also undo the solving of an exercise by using

```
python ./solve.py --undo day1
```

Note that this script just downloads the master or student versions of certain files from the GitHub repository. It needs an Internet connection. Since some exercises require you to have the exercises of the previous days completed, the monitors may ask you to use this function. **Important:** Remember to save your own version of the code, otherwise it will be overwritten!.

# 0.4 Python

#### 0.4.1 Python Basics

#### **Pre-requisites**

At this point you should have installed the needed packages. You need also to feel comfortable with an IDE to edit code and an interactive command line. See previous sections for the details. Your work folder will be

```
lxmls-toolkit-student
```

from there, start your interactive command line of choosing e.g.

```
jupyter-notebook
```

and proceed with the following sections

#### Running Python code

We will start by creating and running a dummy program in Python which simply prints the "Hello World!" message to the standard output (this is usually the first program you code when learning a new programming language). There are two main ways in which you can run code in Python:

From a file - Create a file named yourfile.py and write your program in it, using your IDE of choosing e.g. PyCharm:

```
print 'Hello World!'
```

After saving and closing the file, you can run your code by using the run functionality in your IDE. If you wish to run from a command line instead do

```
python yourfile.py
```

This will run the program and display the message "Hello World!". After, that the control will return to the command line or IDE.

In the interactive command line – Start your interactive command line of chosing e.g. Jupyter-notebook. There, you can run Python code by simply writing it and pressing enter (ctr+enter in Jupyter).

```
In[]: print "Hello, World!"
Out[]: Hello, World!
```

but you can also run Python code written into a file.

```
In[]: run ./yourfile.py
Out[]: Hello, World!
```

Keep in mind that you can easily switch between these two modes. You can quickly test commands in the command line directly and e.g. inspect variables. Larger sections of code can be stored and run from files.

#### Help and Documentation

There are several ways to get help on Jupyter:

- Adding a question mark to the end of a function or variable and pressing Enter brings up associated documentation. Unfortunately, not all packages are well documented. Numpy and matplotlib are pleasant exceptions;
- help('if') gets the online documentation for the if keyword;
- help(), enters the help system.
- When at the help system, type q to exit.

#### 0.4.2 Python by Example

#### **Basic Math Operations**

Python supports all basic arithmetic operations, including exponentiation. For example, the following code:

```
print 3 + 5
print 3 - 5
print 3 * 5
print 3 / 5
print 3 ** 5
```

will produce the following output:

```
8
-2
15
0
243
```

Notice that division is always considered as integer division, hence the result being 0 on the example above. To force a floating point division you can force one of the operands to be a floating point number:

```
print 3 / 5.0
0.6
```

Also, notice that the symbol  $\star\star$  is used as exponentation operator, unlike other major languages which use the symbol  $^{\hat{}}$ . In fact, the  $^{\hat{}}$  symbol has a different meaning in Python (bitwise XOR) so, in the beginning, be sure to double-check your code if it uses exponentiation and it is giving unexpected results.

#### **Data Structures**

In Python, you can create lists of items with the following syntax:

```
countries = ['Portugal','Spain','United Kingdom']
```

A string should be surrounded with apostrophes (') or quotes ("). You can access a list with the following:

- len(L), which returns the number of items in L;
- L[i], which returns the item at index *i* (the first item has index 0);
- L[i:j], which returns a new list, containing all the items between indexes i and j-1, inclusive.

**Exercise 0.1** *Use L[i:j] to return the countries in the Iberian Peninsula.* 

#### **Loops and Indentation**

A loop allows a section of code to be repeated a certain number of times, until a stop condition is reached. For instance, when the list you are iterating has reached its end or when a variable has reached a certain value (in this case, you should not forget to update the value of that variable inside the code of the loop). In Python you have while and for loop statements. The following two example programs output exactly the same using both statements: the even numbers from 2 to 8.

```
i = 2
while i < 10:
    print i
    i += 2</pre>
```

```
for i in range(2,10,2):
    print i
```

You can copy and run this Jupyter. Alternatively you can write this into your yourfile.py file an run it as well. Notice something? It is possible that the code did not act as expected or maybe an error message popped up. This brings us to an important aspect of Python: **indentation**. Indentation is the number of blank spaces at the leftmost of each command. This is how Python differentiates between blocks of commands inside and outside a statement, e.g. while, for or other. All commands within a statement have the same number of blank spaces at their leftmost. For instance, consider the following code:

```
a=1
while a <= 3:
    print a
    a += 1</pre>
```

and its output:

```
1
2
3
```

**Exercise 0.2** Can you then predict the output of the following code?:

```
a=1
while a <= 3:
    print a
a += 1</pre>
```

Bear in mind that indentation is often the main source of errors when starting to work with Python. Try to get used to it as quickly as possible. It is also recommendable that you use a text editor that can display all characters e.g. blank space, tabs, since these characters can be visually similar but are considered different by Python. One of the most common mistakes by newcomers to Python is to have their files indented with spaces on some lines and with tabs on other lines. Visually it might appear that all lines have proper indentation, but you will get an IndentationError message if you try it. The recommended way is to use 4 spaces for each indentation level.

#### **Control Flow**

The if statement allows to control the flow of your program. The next program outputs a greeting that depends on the time of the day.

```
hour = 16
if hour < 12:
    print 'Good morning!'
elif hour >= 12 and hour < 20:
    print 'Good afternoon!'
else:
    print 'Good evening!'</pre>
```

#### **Functions**

A function is a block of code that can be reused to perform a similar action. The following is a function in Python.

```
def greet(hour):
    if hour < 12:
        print 'Good morning!'
    elif hour >= 12 and hour < 20:
        print 'Good afternoon!'
    else:
        print 'Good evening!'</pre>
```

You can write this command into Jupyter directly or write them into a file and run the file in Jupyter. Once you do this the function will be available for you to use. Call the function greet with different hours of the day (for example, type greet (16)) and see that the program will greet you accordingly.

**Exercise 0.3** Note that the previous code allows the hour to be less than 0 or more than 24. Change the code in order to indicate that the hour given as input is invalid. Your output should be something like:

<sup>&</sup>lt;sup>1</sup>The PEP8 document (www.python.org/dev/peps/pep-0008) is the official coding style guide for the Python language.

```
greet(50)
Invalid hour: it should be between 0 and 24.
greet(-5)
Invalid hour: it should be between 0 and 24.
```

#### **Profiling**

If you are interested in checking the performance of your program, you can use the command %prun in Jupyter. For example:

```
def myfunction(x):
    ...
%prun myfunction(22)
```

The output of the %prun command will show the following information for each function that was called during the execution of your code:

- ncalls: The number of times this function was called. If this function was used recursively, the output will be two numbers; the first one counts the total function calls with recursions included, the second one excludes recursive calls.
- tottime: Total time spent in this function, excluding the time spent in other functions called from within this function.
- percall: Same as tottime, but divided by the number of calls.
- cumtime: Same as tottime, but including the time spent in other functions called from within this function.
- percall: Same as cumtime, but divided by the number of calls.
- filename:lineno(function): Tells you where this function was defined.

#### **Debugging in Python**

During the lab sessions, there will be situations in which we will use and extend modules that involve elaborated code and statements, like classes and nested functions. Although desirable, it should not be necessary for you to fully understand the whole code to carry out the exercises. It will suffice to understand the algorithm as explained in the theoretical part of the class and the local context of the part of the code where we will be working. For this to be possible is very important that you learn to use and IDE.

An alternative to IDEs, that can also be useful for quick debugging in Jupyter, is the pdb module. This will stop the execution at a given point (called break-point) to get a quick glimpse of the variable structures and to inspect the execution flow of your program. The ipdb is an improved version of pdb that has to be installed by separate. It provides additional functionalities like larger context windows variable auto complete and colors. Unfortunately ipdb has some compatibility problems with Jupyter. We therefore recommend to use ipdb only in spartan configurations such as vim+ipdb as IDE.

In the following example, we use this module to inspect the greet function:

```
def greet(hour):
    if hour < 12:
        print 'Good morning!'
    elif hour >= 12 and hour < 20:
        print 'Good afternoon!'
    else:
        import pdb; pdb.set_trace()
        print 'Good evening!'</pre>
```

Load the new definition of the function by writing this code in a file or a Jupyter cell and running it. Now, if you try greet (50) the code execution should stop at the place where you located the break-point (that is, in the print 'Good evening!' statement). You can now run new commands or inspect variables. For this purpose there are a number of commands you can use<sup>2</sup>, but we provide here a short table with the most useful:

(h)elp	Starts the help menu	
(p)rint	Prints a variable	
(p)retty(p)rint	Prints a variable, with line break (useful for lists)	
(n)ext line	Jumps to next line	
(s)tep	Jumps inside of the function we stopped at	
c(ont(inue))	Continues execution until finding breakpoint or finishing	
(r)eturn	Continues execution until current function returns	
b(reak) n	Sets a breakpoint in in line n	
b(reak) n, condition	Sets a conditional breakpoint in in line n	
l(ist) [n], [m]	Prints 11 lines around current line. Optionally starting in line n or between lines n, m	
w(here)	Shows which function called the function we are in, and upwards (stack)	
u(p)	Goes one level up the stack (frame of the function that called the function we are on)	
d(down)	Goes one level down the stack	
blank	Repeat the last command	
expression	Executes the python expression as if it was in current frame	

Table 2: Basic pdb/ipdb commands, parentheses indicates abbreviation

Getting back to our example, we can type n(ext) once to execute the line we stopped at

Now we can inspect the variable hour using the p(retty)p(rint) option

```
pdb> pp hour
50
```

From here we could keep advancing with the n(ext) option or set a b(reak) point and type c(ontinue) to jump to a new position. We could also execute any python expression which is valid in the current frame (the function we stopped at). This is particularly useful to find out why code crashes, as we can try different alternatives without the need to restart the code again.

#### 0.4.3 Exceptions

Occasionally, a syntactically correct code statement may produce an error when an attempt is made to execute it. These kind of errors are called *exceptions* in Python. For example, try executing the following:

```
10/0
```

A *ZeroDivisionError* exception was raised, and no output was returned. Exceptions can also be forced to occur by the programmer, with customized error messages <sup>3</sup>.

```
raise ValueError("Invalid input value.")
```

 $<sup>^2</sup>$ The complete list can be found at http://docs.python.org/library/pdb.html

 $<sup>^3</sup>$  for a complete list of built-in exceptions, see http://docs.python.org/2/library/exceptions.html

**Exercise 0.4** Rewrite the code in Exercise 0.3 in order to raise a ValueError exception when the hour is less than 0 or more than 24.

Handling of exceptions is made with the *try* statement:

```
while True:
    try:
        x = int(raw_input("Please enter a number: "))
        break
    except ValueError:
        print "Oops! That was no valid number. Try again..."
```

It works by first executing the *try* clause. If no exception occurs, the *except* clause is skipped; if an exception does occur, and if its type matches the exception named in the *except* keyword, the except clause is executed; otherwise, the exception is raised and execution is aborted (if it is not caught by outer *try* statements).

#### **Extending basic Functionalities with Modules**

In Python you can load new functionalities into the language by using the import, from and as keywords. For example we can load the numpy module as

```
import numpy as np
```

then we can run the following on the Jupyter command line

```
np.var?
np.random.normal?
```

The import will make the numpy tools available through the alias np. This shorter alias prevents the code from getting too long if we load lots of modules. The first command will display the help for the method numpy.var using the previously commented symbol? Note that in order to display the help you need the full name of the function including the module name or alias. Modules have also submodules that can be accessed the same way, as shown in the second example.

#### Organizing your Code with your own modules

Creating you own modules is extremely simple. you can for example create the file in your work directory

```
my_tools.py
```

and store there the following code

```
def my_print(input):
    print input
```

From Jupyter you can now import and use this tool as

```
import my_tools
my_tools.my_print("This works!")
```

**Important**: When you modify a module, you need to either re-start the notebook kernel or notify the notebook to reload the module for the changes to take effect. You can use

```
reload(my_tools)
```

for the latter. Other ways of importing one or all the tools from a module are

```
from my_tools import my_print # my_print directly accesible in code
from my_tools import * # will make all functions in my_tools accesible
```

However, this makes reloading the module more complicated. You can also store tools ind different folders. For example, if you store the previous example in the folder

```
day0_tools
and store inside an empty file called
__init__.py
```

then the following import will work

```
import day0_tools.my_tools
```

## 0.4.4 Matplotlib – Plotting in Python

Matplotlib<sup>4</sup> is a plotting library for Python. It supports 2D and 3D plots of various forms. It can show them interactively or save them to a file (several output formats are supported).

```
import numpy as np
import matplotlib.pyplot as plt

X = np.linspace(-4, 4, 1000)

plt.plot(X, X**2*np.cos(X**2))
plt.savefig("simple.pdf")
```

**Exercise 0.5** *Try running the following on Jupyter, which will introduce you to some of the basic numeric and plotting operations.* 

```
# This will import the numpy library
# and give it the np abbreviation
import numpy as np
# This will import the plotting library
import matplotlib.pyplot as plt
# Linspace will return 1000 points,
# evenly spaced between -4 and +4
X = np.linspace(-4, 4, 1000)
# Y[i] = X[i] * *2
Y = X \star \star 2
# Plot using a red line ('r')
plt.plot(X, Y, 'r')
# arange returns integers ranging from -4 to +4
# (the upper argument is excluded!)
Ints = np.arange(-4, 5)
# We plot these on top of the previous plot
# using blue circles (o means a little circle)
plt.plot(Ints, Ints**2, 'bo')
```

<sup>4</sup>http://matplotlib.org/

```
# You may notice that the plot is tight around the line
# Set the display limits to see better
plt.xlim(-4.5,4.5)
plt.ylim(-1,17)
plt.show()
```

### 0.4.5 Numpy – Scientific Computing with Python

Numpy<sup>5</sup> is a library for scientific computing with Python.

#### **Multidimensional Arrays**

The main object of numpy is the multidimensional array. A multidimensional array is a table with all elements of the same type and can have several dimensions. Numpy provides various functions to access and manipulate multidimensional arrays. In one dimensional arrays, you can index, slice, and iterate as you can with lists. In a two dimensional array M, you can use perform these operations along several dimensions.

- M[i,j], to access the item in the  $i^{th}$  row and  $j^{th}$  column;
- M[i:j,:], to get the all the rows between the  $i^{th}$  and  $j-1^{th}$ ;
- M[:,i], to get the  $i^{th}$  column of M.

Again, as it happened with the lists, the first item of every column and every row has index 0.

```
import numpy as np
A = np.array([
    [1,2,3],
    [2,3,4],
    [4,5,6]])

A[0,:] # This is [1,2,3]
A[0] # This is [1,2,3] as well

A[:,0] # this is [1,2,4]

A[1:,0] # This is [2,4]. Why?
    # Because it is the same as A[1:n,0] where n is the size of the array.
```

### **Mathematical Operations**

There are many helpful functions in numpy. For basic mathematical operations, we have np.log, np.exp, np.cos,...with the expected meaning. These operate both on single arguments and on arrays (where they will behave element wise).

```
import matplotlib.pyplot as plt
import numpy as np

X = np.linspace(0, 4 * np.pi, 1000)
C = np.cos(X)
S = np.sin(X)

plt.plot(X, C)
plt.plot(X, S)
```

Other functions take a whole array and compute a single value from it. For example, np.sum, np.mean,... These are available as both free functions and as methods on arrays.

<sup>5</sup>http://www.numpy.org/

```
import numpy as np
A = np.arange(100)
# These two lines do exactly the same thing
print np.mean(A)
print A.mean()
C = np.cos(A)
print C.ptp()
```

**Exercise 0.6** Run the above example and lookup the ptp function/method (use the? functionality in Jupyter).

Exercise 0.7 Consider the following approximation to compute an integral

$$\int_0^1 f(x)dx \approx \sum_{i=0}^{999} \frac{f(i/1000)}{1000}.$$

Use numpy to implement this for  $f(x) = x^2$ . You should not need to use any loops. Note that integer division in Python 2.x returns the floor division (use floats – e.g. 5.0/2.0 – to obtain rationals). The exact value is 1/3. How close is the approximation?

# 0.5 Essential Linear Algebra

Linear Algebra provides a compact way of representing and operating on sets of linear equations.

$$\begin{cases} 4x_1 - 5x_2 = -13 \\ -2x_1 + 3x_2 = 9 \end{cases}$$

This is a system of linear equations in 2 variables. In matrix notation we can write the system more compactly as

$$Ax = b$$

with

$$\mathbf{A}=\left[\begin{array}{cc} 4 & -5 \\ -2 & 3 \end{array}\right]$$
 ,  $\mathbf{b}=\left[\begin{array}{cc} -13 \\ 9 \end{array}\right]$ 

#### 0.5.1 Notation

We use the following notation:

- By  $A \in \mathbb{R}^{m \times n}$ , we denote a **matrix** with m rows and n columns, where the entries of A are real numbers.
- By  $\mathbf{x} \in \mathbb{R}^n$ , we denote a **vector** with n entries. A vector can also be thought of as a matrix with n rows and 1 column, known as a **column vector**. A **row vector** a matrix with 1 row and n columns is denoted as  $\mathbf{x}^T$ , the transpose of  $\mathbf{x}$ .
- The *i*th element of a vector  $\mathbf{x}$  is denoted  $x_i$ :

$$\mathbf{x} = \left[ \begin{array}{c} x_1 \\ x_2 \\ \vdots \\ x_n \end{array} \right].$$

**Exercise 0.8** *In the rest of the school we will represent both matrices and vectors as numpy arrays. You can create arrays in different ways, one possible way is to create an array of zeros.* 

You can check the shape and the data type of your array using the following commands:

```
print a.shape
  (3, 2)
print a.dtype.name
float64
```

This shows you that "a" is an 3\*2 array of type float64. By default, arrays contain 64 bit<sup>6</sup> floating point numbers. You can specify the particular array type by using the keyword dtype.

```
a = np.zeros([m,n],dtype=int)
print a.dtype
int64
```

You can also create arrays from lists of numbers:

```
a = np.array([[2,3],[3,4]])
print a
[[2 3]
[3 4]]
```

There are many more ways to create arrays in numpy and we will get to see them as we progress in the classes.

### 0.5.2 Some Matrix Operations and Properties

• Product of two matrices  $\mathbf{A} \in \mathbb{R}^{m \times n}$  and  $\mathbf{B} \in \mathbb{R}^{n \times p}$  is the matrix  $\mathbf{C} = \mathbf{A}\mathbf{B} \in \mathbb{R}^{m \times p}$ , where

$$C_{ij} = \sum_{k=1}^{n} A_{ik} B_{kj}.$$

Exercise 0.9 You can multiply two matrices by looping over both indexes and multiplying the individual entries.

```
a = np.array([[2,3],[3,4]])
b = np.array([[1,1],[1,1]])
a_dim1, a_dim2 = a.shape
b_dim1, b_dim2 = b.shape
c = np.zeros([a_dim1,b_dim2])
for i in xrange(a_dim1):
    for j in xrange(b_dim2):
        for k in xrange(a_dim2):
        c[i,j] += a[i,k]*b[k,j]
print c
```

This is, however, cumbersome and inefficient. Numpy supports matrix multiplication with the dot function:

 $<sup>^6</sup>$ On your computer, particularly if you have an older computer, int might denote 32 bits integers

```
d = np.dot(a,b)
print d
```

*Important note:* with numpy, you must use dot to get matrix multiplication, the expression a \* b denotes element-wise multiplication.

- Matrix multiplication is associative: (AB)C = A(BC).
- Matrix multiplication is distributive: A(B+C) = AB + AC.
- Matrix multiplication is (generally) not commutative :  $AB \neq BA$ .
- Given two vectors  $\mathbf{x}, \mathbf{y} \in \mathbb{R}^n$  the product  $\mathbf{x}^T \mathbf{y}$ , called **inner product** or **dot product**, is given by

$$\mathbf{x}^T \mathbf{y} \in \mathbb{R} = \begin{bmatrix} x_1 & x_2 & \dots & x_n \end{bmatrix} \begin{bmatrix} y_1 \\ y_2 \\ \vdots \\ y_n \end{bmatrix} = \sum_{i=1}^n x_i y_i.$$

```
a = np.array([1,2])
b = np.array([1,1])
np.dot(a,b)
```

• Given vectors  $\mathbf{x} \in \mathbb{R}^m$  and  $\mathbf{y} \in \mathbb{R}^n$ , the **outer product**  $\mathbf{x}\mathbf{y}^T \in \mathbb{R}^{m \times n}$  is a matrix whose entries are given by  $(xy^T)_{ij} = x_iy_j$ ,

$$\mathbf{x}\mathbf{y}^T \in \mathbb{R}^{m \times n} = \left[ egin{array}{c} x_1 \\ x_2 \\ \vdots \\ x_m \end{array} \right] \left[ egin{array}{cccc} y_1 & y_2 & \dots & y_n \end{array} \right] = \left[ egin{array}{cccc} x_1y_1 & x_1y_2 & \dots & x_1y_n \\ x_2y_1 & x_2y_2 & \dots & x_2y_n \\ \vdots & \vdots & \ddots & \vdots \\ x_my_1 & x_my_2 & \dots & x_my_n \end{array} \right].$$

• The **identity matrix**, denoted  $I \in \mathbb{R}^{n \times n}$ , is a square matrix with ones on the diagonal and zeros everywhere else. That is,

$$I_{ij} = \left\{ \begin{array}{ll} 1 & i = j \\ 0 & i \neq j \end{array} \right.$$

It has the property that for all  $A \in \mathbb{R}^{n \times n}$ , AI = A = IA.

```
I = np.eye(2)
x = np.array([2.3, 3.4])

print I
print np.dot(I,x)

[[ 1.,  0.],
  [ 0.,  1.]]
[2.3, 3.4]
```

• A diagonal matrix is a matrix where all non-diagonal elements are 0.

• The **transpose** of a matrix results from "'flipping" the rows and columns. Given a matrix  $\mathbf{A} \in \mathbb{R}^{m \times n}$ , the transpose  $\mathbf{A}^T \in \mathbb{R}^{n \times m}$  is the  $n \times m$  matrix whose entries are given by  $(A^T)_{ij} = A_{ii}$ .

Also, 
$$(\mathbf{A}^T)^T = \mathbf{A}$$
;  $(\mathbf{A}\mathbf{B})^T = \mathbf{B}^T \mathbf{A}^T$ ;  $(\mathbf{A} + \mathbf{B})^T = \mathbf{A}^T + \mathbf{B}^T$ 

In numpy, you can access the transpose of a matrix as the  $\ensuremath{\mathbb{T}}$  attribute:

```
A = np.array([ [1, 2], [3, 4] ])
print A.T
```

- A square matrix  $\mathbf{A} \in \mathbb{R}^{n \times n}$  is **symmetric** if  $\mathbf{A} = \mathbf{A}^T$ .
- The **trace** of a square matrix  $\mathbf{A} \in \mathbb{R}^{n \times n}$  is the sum of the diagonal elements,  $\operatorname{tr}(\mathbf{A}) = \sum_{i=1}^{n} A_{ii}$

#### 0.5.3 Norms

The **norm** of a vector is informally the measure of the "length" of the vector. The commonly used Euclidean or  $\ell_2$  norm is given by

$$\|\mathbf{x}\|_2 = \sqrt{\sum_{i=1}^n x_i^2}.$$

• More generally, the  $\ell_p$  norm of a vector  $\mathbf{x} \in \mathbb{R}^n$ , where  $p \ge 1$  is defined as

$$\|\mathbf{x}\|_p = \left(\sum_{i=1}^n |x_i|^p\right)^{1/p}.$$

Note:  $\ell_1 \text{ norm} : \|\mathbf{x}\|_1 = \sum_{i=1}^n |x_i| \qquad \ell_{\infty} \text{ norm} : \|\mathbf{x}\|_{\infty} = \max_i |x_i|$ .

## 0.5.4 Linear Independence, Rank, and Orthogonal Matrices

A set of vectors  $\{\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_n\} \subset \mathbb{R}^m$  is said to be (linearly) independent if no vector can be represented as a linear combination of the remaining vectors. Conversely, if one vector belonging to the set can be represented as a linear combination of the remaining vectors, then the vectors are said to be linearly dependent. That is, if

$$\mathbf{x}_j = \sum_{i \neq j} \alpha_i \mathbf{x}_i$$

for some  $j \in \{1, ..., n\}$  and some scalar values  $\alpha_1, ..., \alpha_{i-1}, \alpha_{i+1}, ..., \alpha_n \in \mathbb{R}$ .

- The **rank** of a matrix is the number of linearly independent columns, which is always equal to the number of linearly independent rows.
- For  $A \in \mathbb{R}^{m \times n}$ , rank $(A) \leq \min(m, n)$ . If rank $(A) = \min(m, n)$ , then A is said to be full rank.
- For  $\mathbf{A} \in R^{m \times n}$ , rank $(\mathbf{A}) = \operatorname{rank}(\mathbf{A}^T)$ .
- For  $\mathbf{A} \in \mathbb{R}^{m \times n}$ ,  $\mathbf{B} \in \mathbb{R}^{n \times p}$ , rank $(\mathbf{A}\mathbf{B}) \leq \min(\operatorname{rank}(\mathbf{A}), \operatorname{rank}(\mathbf{B}))$ .
- For  $\mathbf{A}, \mathbf{B} \in \mathbb{R}^{m \times n}$ ,  $rank(\mathbf{A} + \mathbf{B}) \le rank(\mathbf{A}) + rank(\mathbf{B})$ .
- Two vectors  $\mathbf{x}, \mathbf{y} \in \mathbb{R}^n$  are **orthogonal** if  $\mathbf{x}^T \mathbf{y} = 0$ . A square matrix  $\mathbf{U} \in \mathbb{R}^{n \times n}$  is orthogonal if all its columns are orthogonal to each other and are normalized ( $\|\mathbf{x}\|_2 = 1$ ), It follows that

$$\mathbf{U}^T\mathbf{U}=\mathbf{I}=\mathbf{U}\mathbf{U}^T.$$

# 0.6 Probability Theory

Probability is the most used mathematical language for quantifying uncertainty. The **sample space**  $\mathfrak{X}$  is the set of possible outcomes of an experiment. **Events** are subsets of  $\mathfrak{X}$ .

**Example 0.1 (discrete space)** Let H denote "heads" and T denote "tails." If we toss a coin twice, then  $\mathfrak{X} = \{HH, HT, TH, TT\}$ . The event that the first toss is heads is  $A = \{HH, HT\}$ .

A sample space can also be *continuous* (eg.,  $\mathcal{X} = \mathbb{R}$ ). The union of events A and B is defined as  $A \cup B = \{\omega \in \mathcal{X} \mid \omega \in A \lor \omega \in B\}$ . If  $A_1, \ldots, A_n$  is a sequence of sets then  $\bigcup_{i=1}^n A_i = \{\omega \in \mathcal{X} \mid \omega \in A_i \text{ for at least one i}\}$ . We say that  $A_1, \ldots, A_n$  are **disjoint** or **mutually exclusive** if  $A_i \cap A_j = \emptyset$  whenever  $i \neq j$ .

We want to assign a real number P(A) to every event A, called the **probability** of A. We also call P a **probability distribution** or **probability measure**.

**Definition 0.1** A function P that assigns a real number P(A) to each event A is a **probability distribution** or a **probability measure** if it satisfies the three following axioms:

Axiom 1:  $P(A) \ge 0$  for every A

Axiom 2:  $P(\mathfrak{X}) = 1$ 

Axiom 3: If  $A_1, \ldots, A_n$  are disjoint then

$$P\left(\bigcup_{i=1}^{n} A_i\right) = \sum_{i=1}^{n} P(A_i).$$

One can derive many properties of *P* from these axioms:

$$P(\varnothing) = 0$$

$$A \subseteq B \Rightarrow P(A) \le P(B)$$

$$0 \le P(A) \le 1$$

$$P(A') = 1 - P(A) \quad (A' \text{ is the complement of } A)$$

$$P(A \cup B) = P(A) + P(B) - P(A \cap B)$$

$$A \cap B = \phi \Rightarrow P(A \cup B) = P(A) + P(B).$$

An important case is when events are **independent**, this is also a usual approximation which lends several practical advantages for the computation of the joint probability.

**Definition 0.2** Two events A and B are independent if

$$P(AB) = P(A)P(B) \tag{1}$$

often denoted as  $A \perp B$ . A set of events  $\{A_i : i \in I\}$  is independent if

$$P\left(\bigcap_{i\in J}A_i\right)=\prod_{i\in J}P(A_i)$$

for every finite subset I of I.

For events *A* and *B*, where P(B) > 0, the **conditional probability** of *A* given that *B* has occurred is defined as:

$$P(A|B) = \frac{P(AB)}{P(B)}. (2)$$

Events A and B are independent if and only if P(A|B) = P(A). This follows from the definitions of independence and conditional probability.

A preliminary result that forms the basis for the famous Bayes' theorem is the law of total probability which states that if  $A_1, \ldots, A_k$  is a partition of  $\mathcal{X}$ , then for any event B,

$$P(B) = \sum_{i=1}^{k} P(B|A_i)P(A_i).$$
 (3)

Using Equations 2 and 3, one can derive the Bayes' theorem.

**Theorem 0.1 (Bayes' Theorem)** Let  $A_1, \ldots A_k$  be a partition of  $\mathfrak{X}$  such that  $P(A_i) > 0$  for each i. If P(B) > 0 then, for each  $i = 1, \ldots, k$ ,

$$P(A_i|B) = \frac{P(B|A_i)P(A_i)}{\sum_j P(B|A_j)P(A_j)}.$$
(4)

**Remark 0.1**  $P(A_i)$  is called the prior probability of  $A_i$  and  $P(A_i|B)$  is the posterior probability of  $A_i$ .

**Remark 0.2** *In Bayesian Statistical Inference, the Bayes' theorem is used to compute the estimates of distribution parameters from data. Here, prior is the initial* belief about the parameters, likelihood is the distribution function of the parameter (usually trained from data) and posterior is the updated belief about the parameters.

### 0.6.1 Probability distribution functions

A **random variable** is a mapping  $X : \mathcal{X} \to \mathbb{R}$  that assigns a real number  $X(\omega)$  to each outcome  $\omega$ . Given a random variable X, an important function called the **cumulative distributive function** (or **distribution function**) is defined as:

**Definition 0.3** *The* **cumulative distribution function** *CDF*  $F_X : \mathbb{R} \to [0,1]$  *of a random variable* X *is defined by*  $F_X(x) = P(X \le x)$ .

The CDF is important because it captures the complete information about the random variable. The CDF is right-continuous, non-decreasing and is normalized ( $\lim_{x\to-\infty} F(x)=0$  and  $\lim_{x\to\infty} F(x)=1$ ).

**Example 0.2 (discrete CDF)** Flip a fair coin twice and let X be the random variable indicating the number of heads. Then P(X = 0) = P(X = 2) = 1/4 and P(X = 1) = 1/2. The distribution function is

$$F_X(x) = \begin{cases} 0 & x < 0 \\ 1/4 & 0 \le x < 1 \\ 3/4 & 1 \le x < 2 \\ 1 & x \ge 2. \end{cases}$$

**Definition 0.4** X is discrete if it takes countable many values  $\{x_1, x_2, \ldots\}$ . We define the **probability function** or **probability mass function** for X by

$$f_X(x) = P(X = x).$$

**Definition 0.5** A random variable X is **continuous** if there exists a function  $f_X$  such that  $f_X \ge 0$  for all x,  $\int_{-\infty}^{\infty} f_X(x) dx = 1$  and for every  $a \le b$ 

$$P(a < X < b) = \int_{a}^{b} f_X(x) dx. \tag{5}$$

*The function*  $f_X$  *is called the* **probability density function** (PDF). We have that

$$F_X(x) = \int_{-\infty}^{x} f_X(t)dt$$

and  $f_X(x) = F'_X(x)$  at all points x at which  $F_X$  is differentiable.

A discussion of a few important distributions and related properties:

#### 0.6.2 Bernoulli

The **Bernoulli distribution** is a discrete probability distribution that takes value 1 with the success probability p and 0 with the failure probability q = 1 - p. A single Bernoulli trial is parametrized with the success probability p, and the input  $k \in \{0,1\}$  (1=success, 0=failure), and can be expressed as

$$f(k; p) = p^k q^{1-k} = p^k (1-p)^{1-k}$$

#### 0.6.3 Binomial

The probability distribution for the number of successes in n Bernoulli trials is called a **Binomial distribution**, which is also a discrete distribution. The Binomial distribution can be expressed as exactly j successes is

$$f(j,n;p) = \binom{n}{j} p^{j} q^{n-j} = \binom{n}{j} p^{j} (1-p)^{n-j}$$

where n is the number of Bernoulli trials with probability p of success on each trial.

#### 0.6.4 Categorical

The **Categorical distribution** (often conflated with the Multinomial distribution, in fields like Natural Language Processing) is another generalization of the Bernoulli distribution, allowing the definition of a set of possible outcomes, rather than simply the events "success" and "failure" defined in the Bernoulli distribution. Considering a set of outcomes indexed from 1 to n, the distribution takes the form of

$$f(x_i; p_1, ..., p_n) = p_i.$$

where parameters  $p_1, ..., p_n$  is the set with the occurrence probability of each outcome. Note that we must ensure that  $\sum_{i=1}^{n} p_i = 1$ , so we can set  $p_n = 1 - \sum_{i=1}^{n-1} p_i$ .

#### 0.6.5 Multinomial

The **Multinomial distribution** is a generalization of the Binomial distribution and the Categorical distribution, since it considers multiple outcomes, as the Categorial distribution, and multiple trials, as in the Binomial distribution. Considering a set of outcomes indexed from 1 to n, the vector  $[x_1, ..., x_n]$ , where  $x_i$  indicates the number of times the event with index i occurs, follows the Multinomial distribution

$$f(x_1,...,x_n;p_1,...,p_n) = \frac{n!}{x_1!...x_n!}p_1^{x_1}...p_n^{x_n}.$$

Where parameters  $p_1, ..., p_n$  represent the occurrence probability of the respective outcome.

#### 0.6.6 Gaussian Distribution

A very important theorem in probability theory is the **Central Limit Theorem**. The Central Limit Theorem states that, under very general conditions, if we sum a very large number of mutually independent random variables, then the distribution of the sum can be closely approximated by a certain specific continuous density called the normal (or Gaussian) density. The normal density function with parameters  $\mu$  and  $\sigma$  is defined as follows:

$$f_X(x) = \frac{1}{\sqrt{2\pi}\sigma} e^{-(x-\mu)^2/2\sigma^2}, -\infty < x < \infty.$$

Figure 1 compares a plot of normal density for the cases  $\mu = 0$  and  $\sigma = 1$ , and  $\mu = 0$  and  $\sigma = 2$ .

#### 0.6.7 Maximum Likelihood Estimation

Until now we assumed that, for every distribution, the parameters  $\theta$  are known and are used when we calculate  $p(x|\theta)$ . There are some cases where the values of the parameters are easy to infer, such as the probability p of getting a head using a fair coin, used on a Bernoulli or Binomial distribution. However, in many problems, these values are complex to define and it is more viable to estimate the parameters using the data x. For

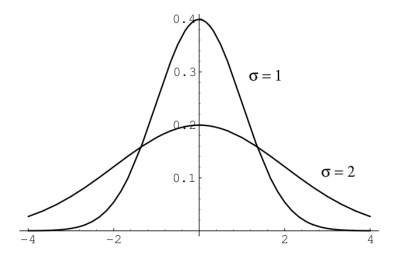


Figure 1: Normal density for two sets of parameter values.

instance, in the example above with the coin toss, if the coin is somehow tampered to have a biased behavior, rather than examining the dynamics or the structure of the coin to infer a parameter for p, a person could simply throw the coin n times, count the number of heads h and set  $p = \frac{h}{n}$ . By doing so, the person is using the data x to estimate  $\theta$ .

With this in mind, we will now generalize this process by defining the probability  $p(\theta|x)$  as the probability of the parameter  $\theta$ , given the data x. This probability is called **likelihood**  $\mathcal{L}(\theta|x)$  and measures how well the parameter  $\theta$  models the data x. The likelihood can be defined in terms of the distribution f as

$$\mathcal{L}(\theta|x_1,...,x_n) = \prod_{i=1}^n f(x_i|\theta)$$

where  $x_1, ..., x_n$  are independently and identically distributed (i.i.d.) samples.

To understand this concept better, we go back to the tampered coin example again. Suppose that we throw the coin 5 times and get the sequence [1,1,1,1,1] (1=head, 0=tail). Using the Bernoulli distribution (see Section 0.6.2) f to model this problem, we get the following likelihood values:

- $\mathcal{L}(0,x) = f(1,0)^5 = 0^5 = 0$
- $\mathcal{L}(0.2, x) = f(1, 0.2)^5 = 0.2^5 = 0.00032$
- $\mathcal{L}(0.4, x) = f(1, 0.4)^5 = 0.4^5 = 0.01024$
- $\mathcal{L}(0.6, x) = f(1, 0.6)^5 = 0.6^5 = 0.07776$
- $\mathcal{L}(0.8, x) = f(1, 0.8)^5 = 0.8^5 = 0.32768$
- $\mathcal{L}(1,x) = f(1,1)^5 = 1^5 = 1$

If we get the sequence [1,0,1,1,0] instead, the likelihood values would be:

- $\mathcal{L}(0,x) = f(1,0)^3 f(0,0)^2 = 0^3 \times 1^2 = 0$
- $\mathcal{L}(0.2, x) = f(1, 0.2)^3 f(0, 0.2)^2 = 0.2^3 \times 0.8^2 = 0.00512$
- $\mathcal{L}(0.4, x) = f(1, 0.4)^3 f(0, 0.4)^2 = 0.4^3 \times 0.6^2 = 0.02304$
- $\mathcal{L}(0.6, x) = f(1, 0.6)^3 f(0, 0.6)^2 = 0.6^3 \times 0.4^2 = 0.03456$
- $\mathcal{L}(0.8, x) = f(1, 0.8)^3 f(0, 0.8)^2 = 0.8^3 \times 0.2^2 = 0.02048$
- $\mathcal{L}(1,x) = f(1,1)^5 = 1^3 \times 0^2 = 0$

We can see that the likelihood is the highest when the distribution f with parameter p is the best fit for the observed samples. Thus, the best estimate for p according to x would be the value for which  $\mathcal{L}(p|x)$  is the highest.

The value of the parameter  $\theta$  with the highest likelihood is called **maximum likelihood estimate (MLE)** and is defined as

$$\hat{\theta}_{mle} = argmax_{\theta} \mathcal{L}(\theta|x)$$

Finding this for our example is relatively easy, since we can simply derivate the likelihood function to find the absolute maximum. For the sequence [1,0,1,1,0], the likelihood would be given as

$$\mathcal{L}(p|x) = f(1,p)^3 f(0,p)^2 = p^3 (1-p)^2$$

And the MLE estimate would be given by:

$$\frac{\delta \mathcal{L}(p|x)}{\delta p} = 0,$$

which resolves to

$$p_{mle} = 0.6$$

**Exercise 0.10** Over the next couple of exercises we will make use of the Galton dataset, a dataset of heights of fathers and sons from the 1877 paper that first discussed the "regression to the mean" phenomenon. This dataset has 928 pairs of numbers.

• *Use the* load() *function in the* galton.py *file to load the dataset. The file is located under the* lxmls/readers *folder*<sup>7</sup>. *Type the following in your Python interpreter:* 

```
import galton as galton
galton_data = galton.load()
```

- What are the mean height and standard deviation of all the people in the sample? What is the mean height of the fathers and of the sons?
- Plot a histogram of all the heights (you might want to use the plt.hist function and the ravel method on arrays).
- Plot the height of the father versus the height of the son.
- You should notice that there are several points that are exactly the same (e.g., there are 21 pairs with the values 68.5 and 70.2). Use the ? command in ipython to read the documentation for the numpy.random.randn function and add random jitter (i.e., move the point a little bit) to the points before displaying them. Does your impression of the data change?

#### 0.6.8 Conjugate Priors

**Definition 0.6** let  $\mathcal{F} = \{f_X(x|s), s \in \mathcal{X}\}$  be a class of likelihood functions; let  $\mathcal{P}$  be a class of probability (density or mass) functions; if, for any x, any  $p_S(s) \in \mathcal{P}$ , and any  $f_X(x|s) \in \mathcal{F}$ , the resulting a posteriori probability function  $p_S(s|x) = f_X(x|s)p_S(s)$  is still in  $\mathcal{P}$ , then  $\mathcal{P}$  is called a conjugate family, or a family of **conjugate priors**, for  $\mathcal{F}$ .

# 0.7 Numerical optimization

Most problems in machine learning require minimization/maximization of functions (likelihoods, risk, energy, entropy, etc.,). Let  $x^*$  be the value of x which minimizes the value of some function f(x). Mathematically, this is written as

$$x^* = \arg\min_{x} f(x)$$

In a few special cases, we can solve this minimization problem analytically in closed form (solving for optimal  $x^*$  in  $\nabla_x f(x^*) = 0$ ), but in most cases it is too cumbersome (or impossible) to solve these equations

 $<sup>^7</sup>$  You might need to inform python about the location of the lxmls labs toolkit. To do so you need to import sys and use the sys.path.append function to add the path to the toolkit readers.

analytically, and they must be tackled numerically. In this section we will cover some basic notions of numerical optimization. The goal is to provide the intuitions behind the methods that will be used in the rest of the school. There are plenty of good textbooks in the subject that you can consult for more information (Nocedal and Wright, 1999; Bertsekas et al., 1995; Boyd and Vandenberghe, 2004).

The most common way to solve the problems when no closed form solution is available is to resort to an iterative algorithm. In this Section, we will see some of these iterative optimization techniques. These iterative algorithms construct a sequence of points  $x^{(0)}, x^{(1)}, \ldots \in \text{domain}(f)$  such that hopefully  $x^t = x^*$  after a number of iterations. Such a sequence is called the **minimizing sequence** for the problem.

#### 0.7.1 Convex Functions

One important property of a function f(x) is whether it is a **convex function** (in the shape of a bowl) or a **non-convex function**. Figures 2 and 3 show an example of a convex and a non-convex function. Convex functions are particularly useful since you can guarantee that the minimizing sequence converges to the true global minimum of the function, while in non-convex functions you can only guarantee that it will reach a local minimum.



Figure 2: Illustration of a convex function. The line segment between any two points on the graph lies entirely above the curve.

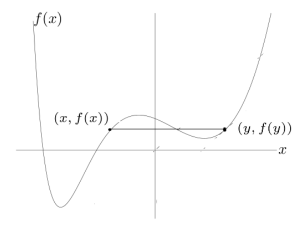


Figure 3: Illustration of a non-convex function. Note the line segment intersecting the curve.

Intuitively, imagine dropping a ball on either side of Figure 2, the ball will roll to the bottom of the bowl independently from where it is dropped. This is the main benefit of a convex function. On the other hand, if you drop a ball from the left side of Figure 3 it will reach a different position than if you drop a ball from its right side. Moreover, dropping it from the left side will lead you to a much better (*i.e.*, lower) place than if you drop the ball from the right side. This is the main problem with non-convex functions: there are no guarantees about the quality of the local minimum you find.

More formally, some concepts to understand about convex functions are: A **line segment** between points  $x_1$  and  $x_2$ : contains all points such that

$$x = \theta x_1 + (1 - \theta)x_2$$

where  $0 \le \theta \le 1$ .

A convex set contains the line segment between any two points in the set

$$x_1, x_2 \in C$$
,  $0 \le \theta \le 1 \implies \theta x_1 + (1 - \theta)x_2 \in C$ .

A function  $f : \mathbb{R}^n \to R$  is a **convex function** if the domain of f is a convex set and

$$f(\theta \mathbf{x} + (1 - \theta)\mathbf{y}) \le \theta f(\mathbf{x}) + (1 - \theta)f(\mathbf{y})$$

for all  $\mathbf{x}, \mathbf{y} \in \mathbb{R}^n$ ,  $0 \le \theta \le 1$ 

#### 0.7.2 Derivative and Gradient

The **derivative** of a function is a measure of how the function varies with its input variables. Given an interval [a,b] one can compute how the function varies within that interval by calculating the average slope of the function in that interval:

$$\frac{f(b) - f(a)}{b - a}. (6)$$

The derivative can be seen as the limit as the interval goes to zero, and it gives us the slope of the function at that point.

$$\frac{\partial f}{\partial x} = \lim_{h=0} \frac{f(x+h) - f(x)}{h} \tag{7}$$

Table 3 shows derivatives of some functions that we will be using during the school.

Function $f(x)$	Derivative $\frac{\partial f}{\partial x}$
$x^2$	2x
$x^n$	$nx^{n-1}$
$\log(x)$	$\frac{1}{x}$
$\exp(x)$	$\exp(x)$
$\frac{1}{x}$	$-\frac{1}{x^2}$

Table 3: Some derivative examples

An important rule of derivation is the chain rule. Consider  $h = f \circ g$ , and u = g(x), then:

$$\frac{\partial h}{\partial x} = \frac{\partial f}{\partial u} \cdot \frac{\partial g}{\partial x} \tag{8}$$

**Example 0.3** Consider the function  $h(x) = \exp(x^2)$ , this can be decomposed as  $h(x) = f(g(x)) = f(u) = \exp(u)$ , where  $u = g(x) = x^2$  and has derivative  $\frac{\partial h}{\partial x} = \frac{\partial f}{\partial u} \cdot \frac{\partial u}{\partial x} = \exp(u) \cdot 2x = \exp(x^2) \cdot 2x$ 

**Exercise 0.11** Consider the function  $f(x) = x^2$  and its derivative  $\frac{\partial f}{\partial x}$ . Look at the derivative of that function at points [-2,0,2], draw the tangent to the graph in that point  $\frac{\partial f}{\partial x}(-2) = -4$ ,  $\frac{\partial f}{\partial x}(0) = 0$ , and  $\frac{\partial f}{\partial x}(2) = 4$ . For example, the tangent equation for x = -2 is y = -4x - b, where b = f(-2). The following code plots the function and the derivatives on those points using matplotlib (See Figure 4).

```
a = np.arange(-5,5,0.01)
f_x = np.power(a,2)
plt.plot(a,f_x)

plt.xlim(-5,5)
plt.ylim(-5,15)

k= np.array([-2,0,2])
plt.plot(k,k**2,"bo")
for i in k:
    plt.plot(a, (2*i)*a - (i**2))
```

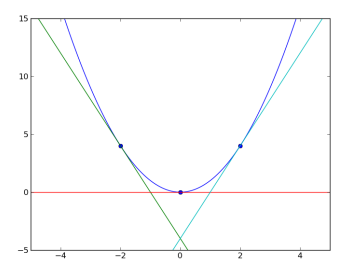


Figure 4: Illustration of the gradient of the function  $f(x^2)$  at three different points x = [-2, 0.2]. Note that at point x = 0 the gradient is zero which corresponds to the minimum of the function.

The **gradient** of a function is a generalization of the derivative concept we just saw before for several dimensions. Let us assume we have a function  $f(\mathbf{x})$  where  $\mathbf{x} \in \mathbb{R}^2$ , so  $\mathbf{x}$  can be seen as a pair  $\mathbf{x} = [x_1, x_2]$ . Then, the gradient measures the slope of the function in both directions:  $\nabla_{\mathbf{x}} f(\mathbf{x}) = [\frac{\partial f}{\partial x_1}, \frac{\partial f}{\partial x_2}]$ .

#### 0.7.3 Gradient Based Methods

Gradient based methods are probably the most common methods used for finding the minimizing sequence for a given function. The methods used in this class will make use of the function value  $f(\mathbf{x}) : \mathbb{R}^n \to R$  as well as the gradient of the function  $\nabla_{\mathbf{x}} f(\mathbf{x})$ . The simplest method is the **Gradient descent** method, an unconstrained first-order optimization algorithm.

The intuition of this method is as follows: You start at a given point  $\mathbf{x}_0$  and compute the gradient at that point  $\nabla_{\mathbf{x}_0} f(\mathbf{x})$ . You then take a step of length  $\eta$  on the direction of the negative gradient to find a new point:  $\mathbf{x}_1 = \mathbf{x}_0 - \eta \nabla_{\mathbf{x}_0} f(\mathbf{x})$ . Then, you compute the gradient at this new point,  $\nabla_{\mathbf{x}_1} f(\mathbf{x})$ , and take a step of length  $\eta$  on the direction of the negative gradient to find a new point:  $\mathbf{x}_2 = \mathbf{x}_1 - \eta \nabla_{\mathbf{x}_1} f(\mathbf{x})$ . You proceed until you have reached a minimum (local or global). Recall from the previous subsection that you can identify the minimum by testing if the norm of the gradient is zero:  $||\nabla f(\mathbf{x})|| = 0$ .

There are several practical concerns even with this basic algorithm to ensure both that the algorithm converges (reaches the minimum) and that it does so in a fast way (by fast we mean the number of function and gradient evaluations).

- Step Size  $\eta$  A first question is how to find the step length  $\eta$ . One condition is that *eta* should guarantee sufficient decrease in the function value. We will not cover these methods here but the most common ones are **Backtracking line search** or the **Wolf Line Search** (Nocedal and Wright, 1999).
- **Descent Direction** A second problem is that using the negative gradient as direction can lead to a very slow convergence. Different methods that change the descent direction by multiplying the gradient by a matrix **B** have been proposed that guarantee a faster convergence. Two notable methods are the Conjugate Gradient (CG) and the Limited Memory Quasi Newton methods (LBFGS) (Nocedal and Wright, 1999).
- Stopping Criteria Finally, it will normally not be possible to reach full convergence either because it will be too slow, or because of numerical issues (computers cannot perform exact arithmetic). So normally we need to define a stopping criteria for the algorithm. Three common criteria (that are normally used together) are: a maximum number of iterations; the gradient norm be smaller than a given threshold  $||\nabla f(\mathbf{x})|| \leq \eta_1$ , or the normalized difference in the function value be smaller than a given threshold  $\frac{|f(\mathbf{x}_t) f(\mathbf{x}_{t-1})|}{\max(|f(\mathbf{x}_t)|,|f(\mathbf{x}_{t-1})|)} \leq \eta_2$

Algorithm 1 shows the general gradient based algorithm. Note that for the simple gradient descent algorithm **B** is the identity matrix and the descent direction is just the negative gradient of the function.

#### Algorithm 1 Gradient Descent

- 1: **given** a starting point  $\mathbf{x}_0$ , i = 0
- 2: repeat
- 3: Compute step size  $\eta$
- 4: Compute descent direction  $-\mathbf{B}\nabla f(x_i)$ .
- 5:  $\mathbf{x}_{i+1} \leftarrow \mathbf{x}_i \eta \mathbf{B} \nabla f(\mathbf{x}_i)$
- 6:  $i \leftarrow i + 1$
- 7: until stopping criterion is satisfied.

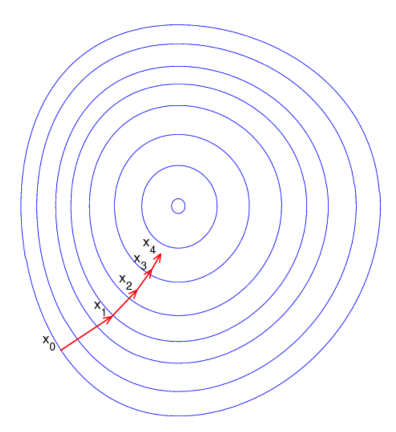


Figure 5: Illustration of gradient descent. The blue circles correspond to contours of the function (each blue circle is a set of points which have the same function value), while the red lines correspond to steps taken in the negative gradient direction.

**Exercise 0.12** Consider the function  $f(x) = (x+2)^2 - 16 \exp(-(x-2)^2)$ . Make a function that computes the function value given x.

```
def get_y(x):
    return (x+2)**2 - 16*np.exp(-((x-2)**2))
```

*Draw a plot around*  $x \in [-8, 8]$ .

```
x = np.arange(-8, 8, 0.001)

y = map(lambda u: get_y(u), x)

plt.plot(x, y)

plt.show()
```

Calculate the derivative of the function f(x), implement the function get\_grad(x).

```
def get_grad(x):
    return (2*x+4)-16*(-2*x + 4)*np.exp(-((x-2)**2))
```

Use the method gradient\_descent to find the minimum of this function. Convince yourself that the code is doing the proper thing. Look at the constants we defined. Note, that we are using a simple approach to pick the step size (always have the value step\_size) which is not necessarily correct.

```
def gradient_descent (start_x, func, grad):
    # Precision of the solution
    prec = 0.0001
    #Use a fixed small step size
    step\_size = 0.1
    #max iterations
    max_iter = 100
    x_new = start_x
    res = []
    for i in xrange(max_iter):
        x\_old = x\_new
        #Use beta egual to -1 for gradient descent
       x_new = x_old - step_size * grad(x_new)
        f_x_{new} = func(x_{new})
       f_x_old = func(x_old)
        res.append([x_new,f_x_new])
        if(abs(f_x_new - f_x_old) < prec):
            print "change in function values too small, leaving"
            return np.array(res)
    print "exceeded maximum number of iterations, leaving"
    return np.array(res)
```

Run the gradient descent algorithm starting from  $x_0 = -8$  and plot the minimizing sequence.

```
x_0 = -8
res = gradient_descent(x_0,get_y,get_grad)
plt.plot(res[:,0],res[:,1],'+')
plt.show()
```

Figure 6 shows the resulting minimizing sequence. Note that the algorithm converged to a minimum, but since the function is not convex it converged only to a local minimum.

Now try the same exercise starting from the initial point  $x_0 = 8$ .

```
x_0 = 8
res = gradient_descent(x_0,get_y,get_grad)
plot(res[:,0],res[:,1],'+')
```

Figure 7 shows the resulting minimizing sequence. Note that now the algorithm converged to the global minimum. However, note that to get to the global minimum the sequence of points jumped from one side of the minimum to the other. This is a consequence of using a wrong step size (in this case too large). Repeat the previous exercise changing both the values of the step-size and the precision. What do you observe?

During this school we will rely on the numerical optimization methods provided by Scipy (scientific computing library in python), which are very efficient implementations.

# 0.8 Python Exercises

#### 0.8.1 Numpy and Matplotlib

**Exercise 0.13** 1. Consider the function  $f(x) = (x+2)^2 - 16 \exp(-(x-2)^2)$ . Draw a plot around the  $x \in [-8, 8]$  region.

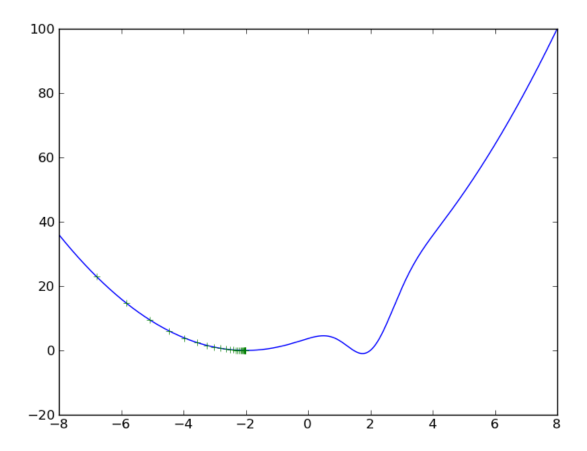


Figure 6: Example of running gradient descent starting on point  $x_0 = -8$  for function  $f(x) = (x+2)^2 - 16 \exp(-(x-2)^2)$ . The function is represented in blue, while the points of the minimizing sequence are displayed as green plus signs.

- 2. What is  $\frac{\partial f}{\partial x}$ ?
- 3. Use gradient descent to find a local minimum starting from  $x_0 = -4$  and  $x_0 = +4$ , with  $\eta = .01$ . Plot all of the intermediate estimates that you obtain in the same plot.

```
import numpy as np
import matplotlib.pyplot as plt
X = np.linspace(-8, 8, 1000)
Y = (X+2)**2 - 16*np.exp(-((X-2)**2))
# derivative of the function f
def get_Y_dev(x):
    return (2*x+4)-16*(-2*x + 4)*np.exp(-((x-2)**2))
def grad_desc(start_x, eps, prec):
    runs the gradient descent algorithm and returns the list of estimates
    example of use grad_desc(start_x=3.9, eps=0.01, prec=0.00001)
    . . .
    x_new = start_x
    x_old = start_x + prec * 2
    res = [x\_new]
    while abs(x_old-x_new) > prec:
        x\_old = x\_new
```

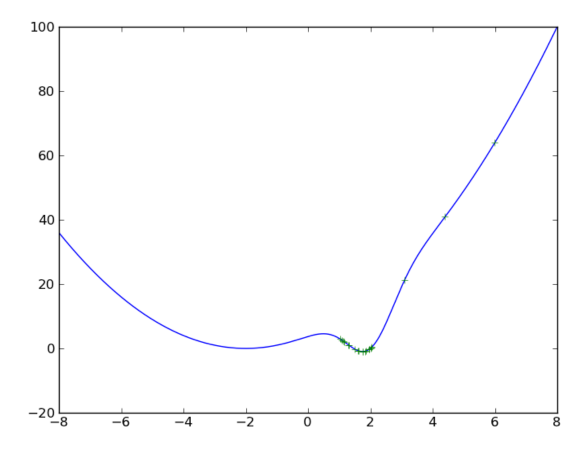


Figure 7: Example of running gradient descent starting on point  $x_0 = 8$  for function  $f(x) = (x+2)^2 - 16 \exp(-(x-2)^2)$ . The function is represented in blue, while the points of the minimizing sequence are displayed as green plus signs.

```
x_new = x_old - eps * get_Y_dev(x_new)
res.append(x_new)
return np.array(res)
```

**Exercise 0.14** Consider the linear regression problem (ordinary least squares) on the Galton dataset, with a single response variable

$$y = x^T w + \varepsilon$$

The linear regression problem is, given a set  $\{y^{(i)}\}_i$  of samples of y and the corresponding  $x^{(i)}$  vectors, estimate w to minimise the sum of the  $\varepsilon$  variables. Traditionally this is solved analytically to obtain a closed form solution (although this is **not the way in which it should be computed** in this exercise, linear algebra packages have an optimised solver, with numpy, use numpy.linalg.lstsg).

Alternatively, we can define the error function for each possible w:

$$e(\boldsymbol{w}) = \sum_{i} \left( \boldsymbol{x^{(i)}}^T \boldsymbol{w} - \boldsymbol{y^{(i)}} \right)^2.$$

- 1. Derive the gradient of the error  $\frac{\partial e}{\partial w_i}$ .
- 2. Implement a solver based on this for two dimensional problems (i.e.,  $w \in \mathbb{R}^2$ ).

3. Use this method on the Galton dataset from the previous exercise to estimate the relationship between father and son's height. Try two formulas

$$s = f w_1 + \varepsilon, \tag{9}$$

where s is the son's height, and f is the father heights; and

$$s = fw_1 + 1w_0 + \varepsilon \tag{10}$$

where the input variable is now two dimensional: (f,1). This allows the intercept to be non-zero.

- 4. Plot the regression line you obtain with the points from the previous exercise.
- 5. Use the np.linalg.lstsqfunction and compare to your solution.

### 0.8.2 Debugging

**Exercise 0.15** Use the debugger to debug the buggy.py script which attempts to repeatedly perform the following computation:

- 1. *Start*  $x_0 = 0$
- 2. Iterate
  - (a)  $x'_{t+1} = x_t + r$ , where r is a random variable.
  - (b) if  $x'_{t+1} >= 1$ ., then stop.
  - (c) if  $x'_{t+1} \le 0$ , then  $x_{t+1} = 0$
  - (d) else  $x_{t+1} = x'_{t+1}$ .
- 3. Return the number of iterations.

Having repeated this computation a number of times, the programme prints the average. Unfortunately, the program has a few bugs, which you need to fix.

# Day 1

# Classification

This day will serve as an introduction to machine learning. We recall some fundamental concepts about decision theory and classification. We also present some widely used models and algorithms and try to provide the main motivation behind them. There are several textbooks that provide a thorough description of some of the concepts introduced here: for example, Mitchell (1997), Duda et al. (2001), Schölkopf and Smola (2002), Joachims (2002), Bishop (2006), Manning et al. (2008), to name just a few. The concepts that we introduce in this chapter will be revisited in later chapters, where the same algorithms and models will be adapted to structured inputs and outputs. For now, we concern only with multi-class classification (with just a few classes).

# Today's assignment

The assignment of today's class is to implement a classifier called Naïve Bayes, and use it to perform sentiment analysis on a corpus of book reviews from Amazon.

#### 1.1 Notation

In what follows, we denote by  $\mathfrak{X}$  our *input set* (also called *observation set*) and by  $\mathfrak{Y}$  our *output set*. We will make no assumptions about the set  $\mathfrak{X}$ , which can be continuous or discrete. In this lecture, we consider *classification* problems, where  $\mathfrak{Y} = \{c_1, \ldots, c_K\}$  is a finite set, consisting of K classes (also called *labels*). For example,  $\mathfrak{X}$  can be a set of documents in natural language, and  $\mathfrak{Y}$  a set of topics, the goal being to assign a topic to each document.

We use upper-case letters for denoting random variables, and lower-case letters for value assignments to those variables: for example,

- X is a random variable taking values on X,
- *Y* is a random variable taking values on *y*,
- $x \in \mathcal{X}$  and  $y \in \mathcal{Y}$  are particular values for X and Y.

We consider *events* such as X = x, Y = y, etc.

For simplicity reasons, throughout this lecture we will use modified notation and let P(y) denote the *probability* associated with the event Y = y (instead of  $P_Y(Y = y)$ ). Also, *joint* and *conditional* probabilities are denoted as  $P(x,y) \triangleq P_{X,Y}(X = x \land Y = y)$  and  $P(x|y) \triangleq P_{X|Y}(X = x \mid Y = y)$ , respectively. From the laws of probabilities:

$$P(x,y) = P(y|x)P(x) = P(x|y)P(y),$$
 (1.1)

for all  $x \in \mathcal{X}$  and  $y \in \mathcal{Y}$ .

Quantities that are predicted or estimated from the data will be appended a hat-symbol: for example, estimations of the probabilities above are denoted as  $\hat{P}(y)$ ,  $\hat{P}(x,y)$  and  $\hat{P}(y|x)$ ; and a prediction of an output will be denoted  $\hat{y}$ .

We assume that a *training dataset*  $\mathcal{D}$  is provided which consists of M input-output pairs (called *examples* or *instances*):

$$\mathcal{D} = \{ (x^1, y^1), \dots, (x^M, y^M) \} \subseteq \mathcal{X} \times \mathcal{Y}.$$
(1.2)

The **goal of (supervised) machine learning** is to use the training dataset  $\mathcal{D}$  to learn a function h (called a *classifier*) that maps from  $\mathcal{X}$  to  $\mathcal{Y}$ : this way, given a new instance  $x \in \mathcal{X}$  (test example), the machine makes a prediction  $\hat{y}$  by evaluating h on x, i.e.,  $\hat{y} = h(x)$ .

# 1.2 Generative Classifiers: Naïve Bayes

If we knew the *true* distribution P(X, Y), the best possible classifier (called Bayes optimal) would be one which predicts according to

$$\hat{y} = \arg \max_{y \in \mathcal{Y}} P(y|x)$$

$$= \arg \max_{y \in \mathcal{Y}} \frac{P(x,y)}{P(x)}$$

$$=^{\dagger} \arg \max_{y \in \mathcal{Y}} P(x,y)$$

$$= \arg \max_{y \in \mathcal{Y}} P(y)P(x|y), \qquad (1.3)$$

where in  $\dagger$  we used the fact that P(x) is constant with respect to y.

Generative classifiers try to estimate the probability distributions P(Y) and P(X|Y), which are respectively called the *class prior* and the *class conditionals*. They assume that the data are generated according to the following generative story (independently for each m = 1, ..., M):

- 1. A class  $y_m \sim P(Y)$  is drawn from the class prior distribution;
- 2. An input  $x_m \sim P(X|Y = y_m)$  is drawn from the corresponding class conditional.

Figure 1.1 shows an example of the Bayes optimal decision boundary for a toy example with K = 2 classes, M = 100 points, class priors  $P(y_1) = P(y_2) = 0.5$ , and class conditionals  $P(x|y_i)$  given by 2-D Gaussian distributions with the same variance but different means.

Simple Data Set -- Mean1= (-1.00,-1.00) Var1 = 0.50 Mean2= (1.00,1.00) Var2= 0.50 Nr. Points=100.00, Balance=0.50 Train-Dev-Test (0.80,.0.00,0.20)

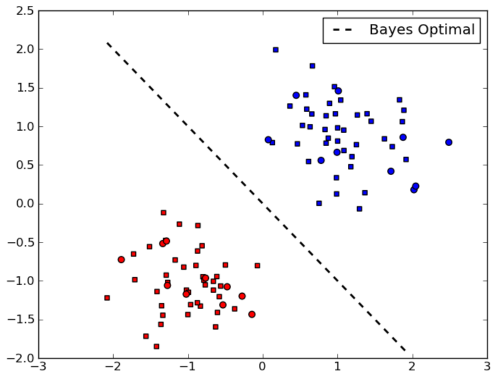


Figure 1.1: Example of a dataset together with the corresponding Bayes optimal decision boundary. The input set consists in points in the real plane,  $\mathcal{X} = \mathcal{R}$ , and the output set consists of two classes (Red and Blue). Training points are represented as squares, while test points are represented as circles.

#### 1.2.1 Training and Inference

Training a generative model amounts to *estimating* the probabilities P(Y) and P(X|Y) using the dataset  $\mathcal{D}$ , yielding estimates  $\hat{P}(y)$  and  $\hat{P}(x|y)$ . This estimation is usually called *training* or *learning*.

After we are done training, we are given a new input  $x \in \mathcal{X}$ , and we want to make a prediction according to

$$\hat{y} = \arg \max_{y \in \mathcal{Y}} \hat{P}(y) \hat{P}(x|y), \tag{1.4}$$

using the estimated probabilities. This is usually called *inference* or *decoding*.

We are left with two important problems:

- 1. How should the distributions  $\hat{P}(Y)$  and  $\hat{P}(X|Y)$  be "defined"? (i.e., what kind of independence assumptions should they state, or how should they factor?)
- 2. How should parameters be estimated from the training data  $\mathfrak{D}$ ?

The first problem strongly depends on the application at hand. Quite often, there is a natural decomposition of the input variable *X* into *J* components,

$$X = (X_1, \dots, X_I). \tag{1.5}$$

The naïve Bayes method makes the following assumption:  $X_1, \ldots, X_J$  are conditionally independent given the class. Mathematically, this means that

$$P(X|Y) = \prod_{j=1}^{J} P(X_j|Y).$$
 (1.6)

Note that this independence assumption greatly reduces the number of parameters to be estimated (degrees of freedom) from  $O(\exp(J))$  to O(J), hence estimation of  $\hat{P}(X|Y)$  becomes much simpler, as we shall see. It also makes the overall computation much more efficient for large J and it decreases the risk of overfitting the data. On the other hand, if the assumption is over-simplistic it may increase the risk of under-fitting.

For the second problem, one of the simplest ways to solve it is using *maximum likelihood estimation*, which aims to maximize the probability of the training sample, assuming that each point was generated independently. This probability (call it  $P(\mathcal{D})$ ) factorizes as

$$P(\mathcal{D}) = \prod_{m=1}^{M} P(x^{m}, y^{m})$$

$$= \prod_{m=1}^{M} P(y^{m}) \prod_{j=1}^{J} P(x_{j}^{m} | y^{m}).$$
(1.7)

#### 1.2.2 Example: Multinomial Naïve Bayes for Document Classification

We now consider a more realistic scenario where the naïve Bayes classifier may be applied. Suppose that the task is *document classification*:  $\mathcal{X}$  is the set of all possible documents, and  $\mathcal{Y} = \{y_1, \dots, y_K\}$  is a set of classes for those documents. Let  $\mathcal{V} = \{w_1, \dots, w_I\}$  be the vocabulary, i.e., the set of words that occur in some document.

A very popular document representation is through a "bag-of-words": each document is seen as a collection of words along with their frequencies; word ordering is ignored. We are going to see that this is equivalent to a naïve Bayes assumption with the *multinomial model*. We associate to each class a multinomial distribution, which ignores word ordering, but takes into consideration the frequency with which each word appears in a document. For simplicity, we assume that all documents have the same length L. Each document x is assumed to have been generated as follows. First, a class y is generated according to P(y). Then, x is generated by sequentially picking words from  $\mathcal V$  with replacement. Each word  $w_j$  is picked with probability  $P(w_j|y)$ . For example, the probability of generating a document  $x = w_j \dots w_{j_L}$  (i.e., a sequence of L words  $w_{j_1}, \dots, w_{j_L}$ ) is

$$P(x|y) = \prod_{l=1}^{L} P(w_{j_l}|y) = \prod_{j=1}^{J} P(w_j|y)^{n_j(x)},$$
(1.8)

<sup>&</sup>lt;sup>1</sup>We can get rid of this assumption by defining a distribution on the document length. Everything stays the same if that distribution is uniform up to a maximum document length.

where  $n_i(x)$  is the number of occurrences of word  $w_i$  in document x.

Hence, the assumption is that word occurrences (*tokens*) are independent given the class. The parameters that need to be estimated are  $\hat{P}(y_1), \ldots, \hat{P}(y_K)$ , and  $\hat{P}(w_j|y_k)$  for  $j=1,\ldots,J$  and  $k=1,\ldots,K$ . Given a training sample  $\mathcal{D}=\{(x^1,y^1),\ldots,(x^M,y^M)\}$ , denote by  $\mathcal{I}_k$  the indices of those instances belonging to the kth class. The maximum likelihood estimates of the quantities above are:

$$\hat{P}(y_k) = \frac{|\mathcal{I}_k|}{M}, \qquad \hat{P}(w_j|y_k) = \frac{\sum_{m \in \mathcal{I}_k} n_j(x^m)}{\sum_{i=1}^{J} \sum_{m \in \mathcal{I}_k} n_i(x^m)}.$$
(1.9)

In words: the class priors' estimates are their relative frequencies (as before), and the class-conditional word probabilities are the relative frequencies of those words across documents with that class.

# 1.3 Assignment

With the previous theoretical background, you will be able to solve today's assignment.

**Exercise 1.1** In this exercise we will use the Amazon sentiment analysis data (Blitzer et al., 2007), where the goal is to classify text documents as expressing a positive or negative sentiment (i.e., a classification problem with two classes). We are going to focus on book reviews. To load the data, type:

```
import lxmls.readers.sentiment_reader as srs
scr = srs.SentimentCorpus("books")
```

This will load the data in a bag-of-words representation where rare words (occurring less than 5 times in the training data) are removed.

- 1. Implement the Naïve Bayes algorithm. Open the file multinomial\_naive\_bayes.py, which is inside the classifiers folder. In the MultinomialNaiveBayes class you will find the train method. We have already placed some code in that file to help you get started.
- 2. After implementing, run Naïve Bayes with the multinomial model on the Amazon dataset (sentiment classification) and report results both for training and testing:

```
import lxmls.classifiers.multinomial_naive_bayes as mnbb

mnb = mnbb.MultinomialNaiveBayes()
params_nb_sc = mnb.train(scr.train_X,scr.train_y)
y_pred_train = mnb.test(scr.train_X,params_nb_sc)
acc_train = mnb.evaluate(scr.train_y, y_pred_train)
y_pred_test = mnb.test(scr.test_X,params_nb_sc)
acc_test = mnb.evaluate(scr.test_y, y_pred_test)
print "Multinomial Naive Bayes Amazon Sentiment Accuracy train: %f test: %f"%(
acc_train,acc_test)
```

3. Observe that words that were not observed at training time cause problems at test time. Why? To solve this problem, apply a simple add-one smoothing technique: replace the expression in Eq. 1.9 for the estimation of the conditional probabilities by

$$\hat{P}(w_j|c_k) = \frac{1 + \sum_{m \in \mathcal{I}_k} n_j(x^m)}{J + \sum_{i=1}^J \sum_{m \in \mathcal{I}_k} n_i(x^m)}.$$

where I is the number of distinct words.

This is a widely used smoothing strategy which has a Bayesian interpretation: it corresponds to choosing a uniform prior for the word distribution on both classes, and to replace the maximum likelihood criterion by a maximum a posteriori approach. This is a form of regularization, preventing the model from overfitting on the training data. See e.g. Manning and Schütze (1999); Manning et al. (2008) for more information. Report the new accuracies.

#### **Discriminative Classifiers** 1.4

In the previous sections we discussed generative classifiers, which require us to model the class prior and class conditional distributions (P(Y)) and P(X|Y), respectively). Recall, however, that a classifier is *any* function which maps objects  $x \in \mathcal{X}$  onto classes  $y \in \mathcal{Y}$ . While it's often useful to model how the data was generated, it's not required. Classifiers that do not model these distributions are called discriminative classifiers.

#### 1.4.1 **Features**

For the purpose of understanding discriminative classifiers, it is useful to think about each  $x \in X$  as an abstract object which is subject to a set of descriptions or measurements, which are called *features*. A feature is simply a real number that describes the value of some property of x. For example, in the previous section, the features of a document were the number of times each word  $w_i$  appeared in it.

Let  $g_1(x), \dots, g_I(x)$  be *J* features of *x*. We call the vector

$$g(x) = (g_1(x), \dots, g_J(x))$$
 (1.10)

a feature vector representation of x. The map  $g: \mathfrak{X} \to \mathbb{R}^J$  is called a feature mapping.

In NLP applications, features are often binary-valued and result from evaluating propositions such as:

$$g_1(x) \triangleq \begin{cases} 1, & \text{if sentence } x \text{ contains the word } Ronaldo \\ 0, & \text{otherwise.} \end{cases}$$
 (1.11)
$$g_2(x) \triangleq \begin{cases} 1, & \text{if all words in sentence } x \text{ are capitalized} \\ 0, & \text{otherwise.} \end{cases}$$
 (1.12)
$$g_3(x) \triangleq \begin{cases} 1, & \text{if } x \text{ contains any of the words } amazing, excellent or :-)} \\ 0, & \text{otherwise.} \end{cases}$$
 (1.13)

$$g_2(x) \triangleq \begin{cases} 1, & \text{if all words in sentence } x \text{ are capitalized} \\ 0, & \text{otherwise.} \end{cases}$$
 (1.12)

$$g_3(x) \triangleq \begin{cases} 1, & \text{if } x \text{ contains any of the words } amazing, excellent \text{ or } :-) \\ 0, & \text{otherwise.} \end{cases}$$
 (1.13)

In this example, the feature vector representation of the sentence "Ronaldo shoots and scores an amazing goal!" would be g(x) = (1, 0, 1).

In multi-class learning problems, rather than associating features only with the input objects, it is useful to consider joint feature mappings  $f: \mathfrak{X} \times \mathfrak{Y} \to \mathbb{R}^D$ . In that case, the joint feature vector f(x,y) can be seen as a collection of joint input-output measurements. For example:

$$f_1(x,y) \triangleq \begin{cases} 1, & \text{if } x \text{ contains } Ronaldo, \text{ and topic } y \text{ is sport} \\ 0, & \text{otherwise.} \end{cases}$$
 (1.14)  
 $f_2(x,y) \triangleq \begin{cases} 1, & \text{if } x \text{ contains } Ronaldo, \text{ and topic } y \text{ is politics} \\ 0, & \text{otherwise.} \end{cases}$ 

$$f_2(x,y) \triangleq \begin{cases} 1, & \text{if } x \text{ contains } Ronaldo, \text{ and topic } y \text{ is politics} \\ 0, & \text{otherwise.} \end{cases}$$
 (1.15)

A very simple form of defining a joint feature mapping which is often employed is via:

$$f(x,y) \triangleq g(x) \otimes e_{y}$$

$$= (0,...,0,\underbrace{g(x)}_{y \text{th slot}},0,...,0)$$
(1.16)

where  $g(x) \in \mathbb{R}^J$  is a input feature vector,  $\otimes$  is the Kronecker product ( $[a \otimes b]_{ij} = a_i b_j$ ) and  $e_y \in \mathbb{R}^K$ , with  $[e_y]_c = 1$  iff y = c, and 0 otherwise. Hence  $f(x, y) \in \mathbb{R}^{J \times K}$ .

#### 1.4.2 Inference

Linear classifiers are very popular in natural language processing applications. They make their decision based on the rule:

$$\hat{y} = \arg\max_{y \in \mathcal{Y}} w \cdot f(x, y). \tag{1.17}$$

where

- $w \in \mathbb{R}^D$  is a weight vector;
- $f(x,y) \in \mathbb{R}^D$  is a feature vector;

•  $w \cdot f(x,y) = \sum_{d=1}^{D} w_d f_d(x,y)$  is the inner product between w and f(x,y).

Hence, each feature  $f_d(x, y)$  has a weight  $w_d$  and, for each class  $y \in \mathcal{Y}$ , a score is computed by linearly combining all the weighted features. All these scores are compared, and a prediction is made by choosing the class with the largest score.

**Remark 1.1** With the design above (Eq. 1.16), and decomposing the weight vector as  $\mathbf{w} = (\mathbf{w}_{c_1}, \dots, \mathbf{w}_{c_K})$ , we have that

$$\boldsymbol{w} \cdot \boldsymbol{f}(\boldsymbol{x}, \boldsymbol{y}) = \boldsymbol{w}_{\boldsymbol{y}} \cdot \boldsymbol{g}(\boldsymbol{x}). \tag{1.18}$$

In words: each class  $y \in \mathcal{Y}$  gets its own weight vector  $\mathbf{w}_y$ , and one defines a input feature vector  $\mathbf{g}(x)$  that only looks at the input  $x \in \mathcal{X}$ . This representation is very useful when features only depend on input x since it allows a more compact representation. Note that the number of features is normally very large.

**Remark 1.2** The multinomial naïve Bayes classifier described in the previous section is an instance of a linear classifier. Recall that the naïve Bayes classifier predicts according to  $\hat{y} = \arg\max_{y \in \mathcal{Y}} \hat{P}(y)\hat{P}(x|y)$ . Taking logs, in the multinomial model for document classification this is equivalent to:

$$\hat{y} = \arg \max_{y \in \mathcal{Y}} \log \hat{P}(y) + \log \hat{P}(x|y)$$

$$= \arg \max_{y \in \mathcal{Y}} \log \hat{P}(y) + \sum_{j=1}^{J} n_j(x) \log \hat{P}(w_j|y)$$

$$= \arg \max_{y \in \mathcal{Y}} w_y \cdot g(x), \tag{1.19}$$

where

$$w_{y} = (b_{y}, \log \hat{P}(w_{1}|y), ..., \log \hat{P}(w_{J}|y))$$

$$b_{y} = \log \hat{P}(y)$$

$$g(x) = (1, n_{1}(x), ..., n_{J}(x)).$$
(1.20)

Hence, the multinomial model yields a prediction rule of the form

$$\hat{y} = \arg \max_{y \in \mathcal{Y}} w_y \cdot g(x). \tag{1.21}$$

#### 1.4.3 Online Discriminative Algorithms

We now discuss two discriminative classification algorithms. These two algorithms are called *online* (or *stochastic*) algorithms because they only process one data point (in our example, one document) at a time. Algorithms which look at the whole dataset at once are called *offline*, or *batch* algorithms, and will be discussed later.

#### Perceptron

The *perceptron* (Rosenblatt, 1958) is perhaps the oldest algorithm used to train a linear classifier. The perceptron works as follows: at each round, it takes an element *x* from the dataset, and uses the current model to make a prediction. If the prediction is correct, nothing happens. Otherwise, the model is corrected by adding the feature vector w.r.t. the correct output and subtracting the feature vector w.r.t. the predicted (wrong) output. Then, it proceeds to the next round.

Alg. 2 shows the pseudo-code of the perceptron algorithm. As it can be seen, it is remarkably simple; yet it often reaches a very good performance, often better than the Naïve Bayes, and usually not much worse than maximum entropy models or SVMs (which will be described in the following section). <sup>2</sup>

A weight vector w defines a *separating hyperplane* if it classifies all the training data correctly, *i.e.*, if  $y^m = \arg\max_{y \in \mathcal{Y}} w \cdot f(x^m, y)$  hold for  $m = 1, \ldots, M$ . A dataset  $\mathcal{D}$  is *separable* if such a weight vector exists (in general, w is not unique). A very important property of the perceptron algorithm is the following: if  $\mathcal{D}$  is separable, then the number of mistakes made by the perceptron algorithm until it finds a separating hyperplane is *finite*. This means that if the data are separable, the perceptron will eventually find a separating hyperplane w.

There are other variants of the perceptron (e.g., with regularization) which we omit for brevity.

<sup>&</sup>lt;sup>2</sup>Actually, we are showing a more robust variant of the perceptron, which averages the weight vector as a post-processing step.

#### Algorithm 2 Averaged perceptron

```
1: input: dataset \mathcal{D}, number of rounds R
 2: initialize t = 0, w^t = 0
 3: for r = 1 to R do
       \mathfrak{D}_s = \operatorname{shuffle}(\mathfrak{D})
       for i = 1 to M do
           m = \mathcal{D}_s(i)
 6:
 7:
           t = t + 1
           take training pair (x^m, y^m) and predict using the current model:
                                                                 \hat{y} \leftarrow \arg\max_{y' \in \mathcal{Y}} w^t \cdot f(x^m, y')
           update the model: w^{t+1} \leftarrow w^t + f(x^m, y^m) - f(x^m, \hat{y})
 9.
        end for
10:
11: end for
12: output: the averaged model \hat{w} \leftarrow \frac{1}{t} \sum_{i=1}^{t} w^{i}
```

**Exercise 1.2** We provide an implementation of the perceptron algorithm in the class Perceptron (file perceptron.py).

1. Run the following commands to generate a simple dataset similar to the one plotted on Figure 1.1:

```
import lxmls.readers.simple_data_set as sds
sd = sds.SimpleDataSet(nr_examples=100, g1 = [[-1,-1],1], g2 = [[1,1],1], balance=0.
5, split=[0.5,0,0.5])
```

2. Run the perceptron algorithm on the simple dataset previously generated and report its train and test set accuracy:

```
import lxmls.classifiers.perceptron as percc

perc = percc.Perceptron()
params_perc_sd = perc.train(sd.train_X,sd.train_y)
y_pred_train = perc.test(sd.train_X,params_perc_sd)
acc_train = perc.evaluate(sd.train_y, y_pred_train)
y_pred_test = perc.test(sd.test_X,params_perc_sd)
acc_test = perc.evaluate(sd.test_Y, y_pred_test)
print "Perceptron Simple Dataset Accuracy train: %f test: %f"%(acc_train,acc_test)
```

3. Plot the decision boundary found:

```
fig,axis = sd.plot_data()
fig,axis = sd.add_line(fig,axis,params_perc_sd,"Perceptron","blue")
```

Change the code to save the intermediate weight vectors, and plot them every five iterations. What do you observe?

4. Run the perceptron algorithm on the Amazon dataset.

#### Margin Infused Relaxed Algorithm (MIRA)

The MIRA algorithm (Crammer and Singer, 2002; Crammer et al., 2006) has achieved very good performance in NLP problems. Recall that the Perceptron takes an input pattern and, if its prediction is wrong, adds the quantity  $[f(x^m, y^m) - f(x^m, \hat{y})]$  to the weight vector. MIRA changes this by adding  $\eta^t[f(x^m, y^m) - f(x^m, \hat{y})]$  to the weight vector. The difference is the step size  $\eta^t$ , which depends on the iteration t.

There is a theoretical basis for this algorithm, which we now briefly explain. At each round t, MIRA updates the weight vector by solving the following optimization problem:

#### Algorithm 3 MIRA

```
1: input: dataset \mathcal{D}, parameter \lambda, number of rounds R
 2: initialize t = 0, w^t = 0
 3: for r = 1 to R do
          \mathfrak{D}_s = \operatorname{shuffle}(\mathfrak{D})
          for i = 1 to M do
 5:
              m = \mathcal{D}_s(i)
 6:
 7:
              t = t + 1
              take training pair (x^m, y^m) and predict using the current model:
 8:
                                                                                  \hat{y} \leftarrow \arg\max_{y' \in \mathcal{Y}} w^t \cdot f(x^m, y')
              compute loss: \ell^t = \boldsymbol{w}^t \cdot f(x^m, \hat{\boldsymbol{y}}) - \boldsymbol{w}^t \cdot f(x^m, y^m) + \rho(\hat{\boldsymbol{y}}, y^m)
 9:
              compute stepsize: \eta^t = \min\left\{\lambda^{-1}, \frac{\ell^t}{\|f(x^m, y^m) - f(x^m, \hat{y})\|^2}\right\} update the model: w^{t+1} \leftarrow w^t + \eta^t(f(x^m, y^m) - f(x^m, \hat{y}))
10:
11:
          end for
12:
13: end for
14: output: the averaged model \hat{w} \leftarrow \frac{1}{t} \sum_{i=1}^{t} w^{i}
```

$$\boldsymbol{w}^{t+1} \leftarrow \operatorname*{arg\,min}_{\boldsymbol{w},\boldsymbol{\xi}} \qquad \qquad \boldsymbol{\xi} + \frac{\lambda}{2} \|\boldsymbol{w} - \boldsymbol{w}^t\|^2 \tag{1.22}$$

s.t. 
$$\mathbf{w} \cdot f(x^m, y^m) \ge \mathbf{w} \cdot f(x^m, \hat{y}) + 1 - \xi$$
 (1.23)

$$\xi \ge 0, \tag{1.24}$$

where  $\hat{y} = \arg\max_{y' \in \mathcal{Y}} \boldsymbol{w}^t \cdot f(x^m, y')$  is the prediction using the model with weight vector  $\boldsymbol{w}^t$ . By inspecting Eq. 1.22 we see that MIRA attempts to achieve a tradeoff between *conservativeness* (penalizing large changes from the previous weight vector via the term  $\frac{\lambda}{2} \| \boldsymbol{w} - \boldsymbol{w}^t \|^2$ ) and *correctness* (by requiring, through the constraints, that the new model  $\boldsymbol{w}^{t+1}$  "separates" the true output from the prediction with a margin (although slack  $\xi \geq 0$  is allowed).<sup>3</sup> Note that, if the prediction is correct  $(\hat{y} = y^m)$  the solution of the problem Eq. 1.22 leaves the weight vector unchanged  $(\boldsymbol{w}^{t+1} = \boldsymbol{w}^t)$ . This quadratic programming problem has a closed form solution:<sup>4</sup>

$$\boldsymbol{w}^{t+1} \leftarrow \boldsymbol{w}^t + \eta^t(f(\boldsymbol{x}^m, \boldsymbol{y}^m) - f(\boldsymbol{x}^m, \hat{\boldsymbol{y}})),$$

with

$$\eta^{t} = \min \left\{ \lambda^{-1}, \frac{w^{t} \cdot f(x^{m}, \hat{y}) - w^{t} \cdot f(x^{m}, y^{m}) + \rho(\hat{y}, y^{m})}{\|f(x^{m}, y^{m}) - f(x^{m}, \hat{y})\|^{2}} \right\},$$

where  $\rho: \mathcal{Y} \times \mathcal{Y} \to \mathbb{R}_+$  is a non-negative cost function, such that  $\rho(\hat{y}, y)$  is the cost incurred by predicting  $\hat{y}$  when the true output is y; we assume  $\rho(y, y) = 0$  for all  $y \in \mathcal{Y}$ . For simplicity, we focus here on the 0/1-cost (but keep in mind that other cost functions are possible):

$$\rho(\hat{y}, y) = \begin{cases} 1 & \text{if } \hat{y} \neq y \\ 0 & \text{otherwise.} \end{cases}$$
 (1.25)

MIRA is depicted in Alg. 3. For other variants of MIRA, see Crammer et al. (2006).

**Exercise 1.3** We provide an implementation of the MIRA algorithm. Compare it with the perceptron for various values of  $\lambda$ 

```
import lxmls.classifiers.mira as mirac

mira = mirac.Mira()
mira.regularizer = 1.0 # This is lambda
params_mira_sd = mira.train(sd.train_X, sd.train_y)
y_pred_train = mira.test(sd.train_X, params_mira_sd)
```

<sup>&</sup>lt;sup>3</sup>The intuition for this large margin separation is the same for support vector machines, which will be discussed in §1.4.4.

<sup>&</sup>lt;sup>4</sup>Note that the perceptron updates are identical, except that we always have  $\eta_t = 1$ .

```
acc_train = mira.evaluate(sd.train_y, y_pred_train)
y_pred_test = mira.test(sd.test_X, params_mira_sd)
acc_test = mira.evaluate(sd.test_y, y_pred_test)
print "Mira Simple Dataset Accuracy train: %f test: %f"%(acc_train,acc_test)
fig,axis = sd.add_line(fig,axis,params_mira_sd,"Mira","green")

params_mira_sc = mira.train(scr.train_X,scr.train_y)
y_pred_train = mira.test(scr.train_X,params_mira_sc)
acc_train = mira.evaluate(scr.train_y, y_pred_train)
y_pred_test = mira.test(scr.test_X,params_mira_sc)
acc_test = mira.evaluate(scr.test_y, y_pred_test)
print "Mira Amazon Sentiment Accuracy train: %f test: %f"%(acc_train,acc_test)
```

Compare the results achieved and separating hiperplanes found.

#### 1.4.4 Batch Discriminative Classifiers

As we have mentioned, the perceptron and MIRA algorithms are called *online* or *stochastic* because they look at one data point at a time. We now describe two discriminative classifiers which look at all points at once; these are called *offline* or *batch* algorithms.

#### **Maximum Entropy Classifiers**

The notion of *entropy* in the context of Information Theory (Shannon, 1948) is one of the most significant advances in mathematics in the twentieth century. The principle of *maximum entropy* (which appears under different names, such as "maximum mutual information" or "minimum Kullback-Leibler divergence") plays a fundamental role in many methods in statistics and machine learning (Jaynes, 1982). <sup>5</sup> The basic rationale is that choosing the model with the highest entropy (subject to constraints that depend on the observed data) corresponds to making the fewest possible assumptions regarding what was unobserved, making uncertainty about the model as large as possible.

For example, if we throw a die and want to estimate the probability of its outcomes, the distribution with the highest entropy would be the uniform distribution (each outcome having of probability a 1/6). Now suppose that we are only told that outcomes  $\{1,2,3\}$  occurred 10 times in total, and  $\{4,5,6\}$  occurred 30 times in total, then the principle of maximum entropy would lead us to estimate P(1) = P(2) = P(3) = 1/12 and P(4) = P(5) = P(6) = 1/4 (i.e., outcomes would be uniform within each of the two groups).

This example could be presented in a more formal way. Suppose that we want to use binary features to represent the outcome of the die throw. We use two features:  $f_{123}(x,y)=1$  if and only if  $y\in\{1,2,3\}$ , and  $f_{456}(x,y)=1$  if and only if  $y\in\{4,5,6\}$ . Our observations state that in 40 throws, we observed  $f_{123}$  10 times (25%) and  $f_{456}$  30 times (75%). The maximum entropy principle states that we want to find the parameters w of our model, and consequently the probability distribution  $P_w(Y|X)$ , which makes  $f_{123}$  have an expected value of 0.25 and  $f_{456}$  have an expected value of 0.75. These constraints,  $E[f_{123}]=0.25$  and  $E[f_{456}]=0.75$ , are known as first moment matching constraints.

An important fundamental result, which we will not prove here, is that the maximum entropy distribution  $P_w(Y|X)$  under first moment matching constraints is a *log-linear model*. <sup>8</sup> It has the following parametric form:

$$P_{w}(y|x) = \frac{\exp(w \cdot f(x,y))}{Z(w,x)}$$
(1.26)

The denominator in Eq. 1.26 is called the *partition function*:

$$Z(\boldsymbol{w}, \boldsymbol{x}) = \sum_{\boldsymbol{y}' \in \boldsymbol{\mathcal{Y}}} \exp(\boldsymbol{w} \cdot \boldsymbol{f}(\boldsymbol{x}, \boldsymbol{y}')). \tag{1.27}$$

<sup>&</sup>lt;sup>5</sup>For an excellent textbook on Information Theory, we recommend Cover et al. (1991).

<sup>&</sup>lt;sup>6</sup>For an introduction of maximum entropy models, along with pointers to the literature, see http://www.cs.cmu.edu/~aberger/maxent.html.

<sup>&</sup>lt;sup>7</sup>In general, these constraints mean that feature expectations under that distribution  $\frac{1}{M} \sum_m E_{Y \sim P_w}[f(x_m, Y)]$  must match the observed relative frequencies  $\frac{1}{M} \sum_m f(x_m, y_m)$ .

<sup>&</sup>lt;sup>8</sup>Also called a a Boltzmann distribution, or an exponential family of distributions.

An important property of the partition function is that the gradient of its logarithm equals the feature expectations:

$$\nabla_{w} \log Z(w, x) = E_{w}[f(x, Y)] = \sum_{y' \in \mathcal{Y}} P_{w}(y'|x) f(x, y').$$
 (1.28)

The average conditional log-likelihood is:

$$\mathcal{L}(w; \mathcal{D}) = \frac{1}{M} \log P_{w}(y^{1}, \dots, y^{M} | x^{1}, \dots, x^{M})$$

$$= \frac{1}{M} \log \prod_{m=1}^{M} P_{w}(y^{m} | x^{m})$$

$$= \frac{1}{M} \sum_{m=1}^{M} \log P_{w}(y^{m} | x^{m})$$

$$= \frac{1}{M} \sum_{m=1}^{M} (w \cdot f(x^{m}, y^{m}) - \log Z(w, x^{m})).$$
(1.29)

We try to find the parameters w that maximize the log-likelihood  $\mathcal{L}(w; \mathcal{D})$ ; to avoid overfitting, we add a regularization term that penalizes values of w that have a high magnitude. The optimization problem becomes:

$$\hat{w} = \underset{w}{\operatorname{arg max}} \mathcal{L}(w; \mathcal{D}) - \frac{\lambda}{2} ||w||^{2}$$

$$= \underset{w}{\operatorname{arg min}} - \mathcal{L}(w; \mathcal{D}) + \frac{\lambda}{2} ||w||^{2}. \tag{1.30}$$

Here we use the squared  $L_2$ -norm as the regularizer, but other norms are possible. The scalar  $\lambda \geq 0$  controls the amount of regularization. Unlike the naïve Bayes examples, this optimization problem does not have a closed form solution in general; hence we need to resort to numerical optimization (see section 0.7). Let  $F_{\lambda}(w; \mathcal{D}) = -\mathcal{L}(w; \mathcal{D}) + \frac{\lambda}{2} ||w||^2$  be the objective function in Eq. 1.30. This function is convex, which implies that a local optimum of Eq. 1.30 is also a global optimum.  $F_{\lambda}(w; \mathcal{D})$  is also differentiable: its gradient is

$$\nabla_{w} F_{\lambda}(w; \mathcal{D}) = \frac{1}{M} \sum_{m=1}^{M} (-f(x^{m}, y^{m}) + \nabla_{w} \log Z(w, x^{m})) + \lambda w$$

$$= \frac{1}{M} \sum_{m=1}^{M} (-f(x^{m}, y^{m}) + E_{w}[f(x^{m}, Y)]) + \lambda w. \tag{1.31}$$

A batch gradient method to optimize Eq. 1.30 is shown in Alg. 4. Essentially, Alg. 4 iterates through the following updates until convergence:

$$w^{t+1} \leftarrow w^{t} - \eta_{t} \nabla_{w} F_{\lambda}(w^{t}; \mathcal{D})$$

$$= (1 - \lambda \eta_{t}) w^{t} + \eta_{t} \frac{1}{M} \sum_{m=1}^{M} (f(x^{m}, y^{m}) - E_{w}[f(x^{m}, Y)]). \qquad (1.32)$$

Convergence is ensured for suitable stepsizes  $\eta_t$ . Monotonic decrease of the objective value can also be ensured if  $\eta_t$  is chosen with a suitable line search method, such as Armijo's rule (Nocedal and Wright, 1999). In practice, more sophisticated methods exist for optimizing Eq. 1.30, such as conjugate gradient or L-BFGS. The latter is an example of a quasi-Newton method, which only requires gradient information, but uses past gradients to try to construct second order (Hessian) approximations.

In large-scale problems (very large M) batch methods are slow. *Online* or *stochastic* optimization are attractive alternative methods. Stochastic gradient methods make "noisy" gradient updates by considering only a single instance at the time. The resulting algorithm, called Stochastic Gradient Descent (SGD) is shown as Alg. 5. At each round t, an instance m(t) is chosen, either randomly (stochastic variant) or by cycling through the dataset (online variant). The stepsize sequence must decrease with t: typically,  $\eta_t = \eta_0 t^{-\alpha}$  for some  $\eta_0 > 0$ 

<sup>&</sup>lt;sup>9</sup>In a Bayesian perspective, this corresponds to choosing independent Gaussian priors  $p(w_d) \sim \mathcal{N}(0; 1/\lambda^2)$  for each dimension of the weight vector.

#### Algorithm 4 Batch Gradient Descent for Maximum Entropy

- input: D, λ, number of rounds T, learning rate sequence (η<sub>t</sub>)<sub>t=1,...,T</sub>
   initialize w<sup>1</sup> = 0
- 3: **for** t = 1 **to** T **do**
- 4: **for** m = 1 **to** M **do**
- take training pair  $(x^m, y^m)$  and compute conditional probabilities using the current model, for each  $y' \in \mathcal{Y}$ :

$$P_{\boldsymbol{w}^t}(\boldsymbol{y}'|\boldsymbol{x}^m) = \frac{\exp(\boldsymbol{w}^t \cdot f(\boldsymbol{x}^m, \boldsymbol{y}'))}{Z(\boldsymbol{w}, \boldsymbol{x}^m)}$$

6: compute the feature vector expectation:

$$E_w[f(x^m,Y)] = \sum_{y' \in \mathcal{Y}} P_{w^t}(y'|x^m) f(x^m,y')$$

- 7: end for
- 8: choose the stepsize  $\eta_t$  using, e.g., Armijo's rule
- 9: update the model:

$$w^{t+1} \leftarrow (1 - \lambda \eta_t) w^t + \eta_t M^{-1} \sum_{m=1}^{M} (f(x^m, y^m) - E_w[f(x^m, Y)])$$

10: **end for** 11: **output:**  $\hat{w} \leftarrow w^{T+1}$ 

and  $\alpha \in [1,2]$ , tuned in a development partition or with cross-validation.

**Exercise 1.4** We provide an implementation of the L-BFGS algorithm for training maximum entropy models in the class MaxEnt\_batch, as well as an implementation of the SGD algorithm in the class MaxEnt\_online.

1. Train a maximum entropy model using L-BFGS on the Simple data set (try different values of  $\lambda$ ). Compare the results with the previous methods. Plot the decision boundary.

```
import lxmls.classifiers.max_ent_batch as mebc

me_lbfgs = mebc.MaxEntBatch()
me_lbfgs.regularizer = 1.0
params_meb_sd = me_lbfgs.train(sd.train_X,sd.train_y)
y_pred_train = me_lbfgs.test(sd.train_X,params_meb_sd)
acc_train = me_lbfgs.evaluate(sd.train_y, y_pred_train)
y_pred_test = me_lbfgs.test(sd.test_X,params_meb_sd)
acc_test = me_lbfgs.evaluate(sd.test_y, y_pred_test)
print "Max-Ent batch Simple Dataset Accuracy train: %f test: %f"%(acc_train,acc_test_)
fig,axis = sd.add_line(fig,axis,params_meb_sd,"Max-Ent-Batch","orange")
```

2. Train a maximum entropy model using L-BFGS, on the Amazon dataset (try different values of  $\lambda$ ) and report training and test set accuracy. What do you observe?

```
params_meb_sc = me_lbfgs.train(scr.train_X,scr.train_y)
y_pred_train = me_lbfgs.test(scr.train_X,params_meb_sc)
acc_train = me_lbfgs.evaluate(scr.train_y, y_pred_train)
y_pred_test = me_lbfgs.test(scr.test_X,params_meb_sc)
acc_test = me_lbfgs.evaluate(scr.test_y, y_pred_test)
print "Max-Ent Batch Amazon Sentiment Accuracy train: %f test: %f"%(acc_train, acc_test)
```

#### Algorithm 5 SGD for Maximum Entropy

```
    input: D, λ, number of rounds T, learning rate sequence (η<sub>t</sub>)<sub>t=1,...,T</sub>
    initialize w<sup>1</sup> = 0
    for t = 1 to T do
```

4: choose m = m(t) randomly

5: take training pair  $(x^m, y^m)$  and compute conditional probabilities using the current model, for each  $y' \in y$ :

$$P_{\boldsymbol{w}^t}(\boldsymbol{y}'|\boldsymbol{x}^m) = \frac{\exp(\boldsymbol{w}^t \cdot f(\boldsymbol{x}^m, \boldsymbol{y}'))}{Z(\boldsymbol{w}, \boldsymbol{x}^m)}$$

6: compute the feature vector expectation:

$$E_{\boldsymbol{w}}[f(\boldsymbol{x}^m,\boldsymbol{Y})] = \sum_{\boldsymbol{y}' \in \boldsymbol{\mathcal{Y}}} P_{\boldsymbol{w}^t}(\boldsymbol{y}'|\boldsymbol{x}^m) f(\boldsymbol{x}^m,\boldsymbol{y}')$$

7: update the model:

$$\boldsymbol{w}^{t+1} \leftarrow (1 - \lambda \eta_t) \boldsymbol{w}^t + \eta_t \left( f(\boldsymbol{x}^m, \boldsymbol{y}^m) - E_{\boldsymbol{w}}[f(\boldsymbol{x}^m, \boldsymbol{Y})] \right)$$

8: end for

9: **output:**  $\hat{\boldsymbol{w}} \leftarrow \boldsymbol{w}^{T+1}$ 

3. Now, fix  $\lambda = 1.0$  and train with SGD (you might try to adjust the initial step). Compare the objective values obtained during training with those obtained with L-BFGS. What do you observe?

```
import lxmls.classifiers.max_ent_online as meoc

me_sgd = meoc.MaxEntOnline()
me_sgd.regularizer = 1.0
params_meo_sc = me_sgd.train(scr.train_X,scr.train_y)
y_pred_train = me_sgd.test(scr.train_X,params_meo_sc)
acc_train = me_sgd.evaluate(scr.train_y, y_pred_train)
y_pred_test = me_sgd.test(scr.test_X,params_meo_sc)
acc_test = me_sgd.evaluate(scr.test_y, y_pred_test)
print "Max-Ent Online Amazon Sentiment Accuracy train: %f test: %f"%(acc_train, acc_test)
```

#### **Support Vector Machines**

Support vector machines are also a discriminative approach, but they are not a probabilistic model at all. The basic idea is that, if the goal is to accurately predict outputs (according to some cost function), we should focus on that goal in the first place, rather than trying to estimate a probability distribution (P(Y|X) or P(X,Y)), which is a more difficult problem. As Vapnik (1995) puts it, "do not solve an estimation problem of interest by solving a more general (harder) problem as an intermediate step."

We next describe the *primal* problem associated with multi-class support vector machines (Crammer and Singer, 2002), which is of primary interest in natural language processing. There is a significant amount of literature about Kernel Methods (Schölkopf and Smola, 2002; Shawe-Taylor and Cristianini, 2004) mostly focused on the *dual* formulation. We will not discuss non-linear kernels or this dual formulation here.<sup>10</sup>

Consider  $\rho(y',y)$  as a non-negative cost function, representing the cost of assigning a label y' when the correct label was y. For simplicity, we focus here on the 0/1-cost defined by Equation 1.25 (but keep in mind that other cost functions are possible). The *hinge*  $loss^{11}$  is the function

$$\ell(\boldsymbol{w}; \boldsymbol{x}, \boldsymbol{y}) = \max_{\boldsymbol{y}' \in \mathcal{Y}} \left[ \boldsymbol{w} \cdot \boldsymbol{f}(\boldsymbol{x}, \boldsymbol{y}') - \boldsymbol{w} \cdot \boldsymbol{f}(\boldsymbol{x}, \boldsymbol{y}) + \rho(\boldsymbol{y}', \boldsymbol{y}) \right]. \tag{1.33}$$

<sup>&</sup>lt;sup>10</sup>The main reason why we prefer to discuss the primal formulation with linear kernels is that the resulting algorithms run in linear time (or less), while known kernel-based methods are quadratic with respect to *M*. In large-scale problems (large *M*) the former are thus more appealing.

appealing. The hinge loss for the 0/1 cost is sometimes defined as  $\ell(w; x, y) = \max\{0, \max_{y' \neq y} w \cdot f(x, y') - w \cdot f(x, y) + 1\}$ . Given our definition of  $\rho(\hat{y}, y)$ , note that the two definitions are equivalent.

#### Algorithm 6 Stochastic Subgradient Descent for SVMs

- 1: **input:**  $\mathfrak{D}$ ,  $\lambda$ , number of rounds T, learning rate sequence  $(\eta_t)_{t=1,\dots,T}$
- 2: initialize  $w^1 = 0$
- 3: **for** t = 1 **to** T **do**
- 4: choose m = m(t) randomly
- 5: take training pair  $(x^m, y^m)$  and compute the "cost-augmented prediction" under the current model:

$$\tilde{y} = \underset{y' \in \mathcal{Y}}{\arg\max} \, w^t \cdot f(x^m, y') - w^t \cdot f(x^m, y^m) + \rho(y', y)$$

6: update the model:

$$\boldsymbol{w}^{t+1} \leftarrow (1 - \lambda \eta_t) \boldsymbol{w}^t + \eta_t \left( f(\boldsymbol{x}^m, \boldsymbol{y}^m) - f(\boldsymbol{x}^m, \tilde{\boldsymbol{y}}) \right)$$

- 7: end for
- 8: **output:**  $\hat{w} \leftarrow w^{T+1}$

Note that the objective of Eq. 1.33 becomes zero when y' = y. Hence, we always have  $\ell(w; x, y) \geq 0$ . Moreover, if  $\rho$  is the 0/1 cost, we have  $\ell(w; x, y) = 0$  if and only if the weight vector is such that the model makes a correct prediction with a *margin* greater than 1: *i.e.*,  $w \cdot f(x, y) \geq w \cdot f(x, y') + 1$  for all  $y' \neq y$ . Otherwise, a positive loss is incurred. The idea behind this formulation is that not only do we want to make a correct prediction, but we want to make a *confident* prediction; this is why we have a loss unless the prediction is correct with some margin.

Support vector machines (SVM) tackle the following optimization problem:

$$\hat{w} = \arg\min_{w} \sum_{m=1}^{M} \ell(w; x^{m}, y^{m}) + \frac{\lambda}{2} ||w||^{2},$$
 (1.34)

where we also use the squared  $L_2$ -norm as the regularizer. For the 0/1-cost, the problem in Eq. 1.34 is equivalent to:

$$\underset{\boldsymbol{w},\boldsymbol{\xi}}{\operatorname{arg\,min}} \qquad \qquad \sum_{m=1}^{M} \xi_m + \frac{\lambda}{2} \|\boldsymbol{w}\|^2 \tag{1.35}$$

s.t. 
$$\mathbf{w} \cdot f(\mathbf{x}^m, \mathbf{y}^m) \ge \mathbf{w} \cdot f(\mathbf{x}^m, \tilde{\mathbf{y}}^m) + 1 - \xi_m, \quad \forall m, \tilde{\mathbf{y}}^m \in \mathcal{Y} \setminus \{\mathbf{y}^m\}.$$
 (1.36)

Geometrically, we are trying to choose the linear classifier that yields the largest possible separation margin, while we allow some violations, penalizing the amount of slack via extra variables  $\xi_1, \ldots, \xi_m$ . There is now a trade-off: increasing the slack variables  $\xi_m$  makes it easier to satisfy the constraints, but it will also increase the value of the cost function.

Problem 1.34 does not have a closed form solution. Moreover, unlike maximum entropy models, the objective function in 1.34 is non-differentiable, hence smooth optimization is not possible. However, it is still convex, which ensures that any local optimum is the global optimum. Despite not being differentiable, we can still define a *subgradient* of the objective function (which generalizes the concept of gradient), which enables us to apply subgradient-based methods. A stochastic subgradient algorithm for solving Eq. 1.34 is illustrated as Alg. 6. The similarity with maximum entropy models (Alg. 5) is striking: the only difference is that, instead of computing the feature vector expectation using the current model, we compute the feature vector associated with the cost-augmented prediction using the current model.

A variant of this algorithm was proposed by Shalev-Shwartz et al. (2007) under the name *Pegasos*, with excellent properties in large-scale settings. Other algorithms and software packages for training SVMs that have become popular are SVMLight (http://svmlight.joachims.org) and LIBSVM (http://www.csie.ntu.edu.tw/~cjlin/libsvm/), which allow non-linear kernels. These will generally be more suitable for smaller datasets, where high accuracy optimization can be obtained without much computational effort.

**Remark 1.3** *Note the similarity between the stochastic (sub-)gradient algorithms (Algs. 5–6) and the online algorithms seen above (perceptron and MIRA).* 

**Exercise 1.5** Implement the SVM primal algorithm (Hint: look at the models implemented earlier, you should only need to change a few lines of code). Do this by creating a file SVM. py and implement class SVM. Then, repeat the MaxEnt exercise now using SVMs, for several values of  $\lambda$ :

```
import lxmls.classifiers.svm as svmc
svm = svmc.SVM()
svm.regularizer = 1.0 # This is lambda
params_svm_sd = svm.train(sd.train_X,sd.train_y)
y_pred_train = svm.test(sd.train_X,params_svm_sd)
acc_train = svm.evaluate(sd.train_y, y_pred_train)
y_pred_test = svm.test(sd.test_X,params_svm_sd)
acc_test = svm.evaluate(sd.test_y, y_pred_test)
print "SVM Online Simple Dataset Accuracy train: %f test: %f"%(acc_train,acc_test)
fig,axis = sd.add_line(fig,axis,params_svm_sd,"SVM","orange")
params_svm_sc = svm.train(scr.train_X, scr.train_y)
y_pred_train = svm.test(scr.train_X,params_svm_sc)
acc_train = svm.evaluate(scr.train_y, y_pred_train)
v_pred_test = svm.test(scr.test_X,params_svm_sc)
acc_test = svm.evaluate(scr.test_y, y_pred_test)
print "SVM Online Amazon Sentiment Accuracy train: %f test: %f"%(acc_train,acc_test)
```

Compare the results achieved and separating hiperplanes found.

## 1.5 Comparison

Table 1.1 provides a high-level comparison among the different algorithms discussed in this chapter.

	Naive Bayes	Perceptron	MIRA	MaxEnt	SVMs
Generative/Discriminative	G	D	D	D	D
Performance if true model	Bad	Fair (may	Good	Good	Good
not in the hipothesis class		not converge)			
Performance if features overlap	Fair	Good	Good	Good	Good
Training	Closed Form	Easy	Easy	Fair	Fair
Hyperparameters to tune	1 (smoothing)	0	1	1	1

Table 1.1: Comparison among different algorithms.

**Exercise 1.6** • Using the simple dataset run the different algorithms varying some characteristics of the data: like the number of points, variance (hence separability), class balance. Use function run\_all\_classifiers in file labs/run\_all\_classifiers.py which receives a dataset and plots all decisions boundaries and accuracies. What can you say about the methods when the amount of data increases? What about when the classes become too unbalanced?

#### 1.6 Final remarks

Some implementations of the discussed algorithms are available on the Web:

- SVMLight: http://svmlight.joachims.org
- LIBSVM: http://www.csie.ntu.edu.tw/~cjlin/libsvm/
- Maximum Entropy: http://homepages.inf.ed.ac.uk/lzhang10/maxent\_toolkit.html
- MALLET: http://mallet.cs.umass.edu/.

## Day 2

# **Sequence Models**

In this class, we relax the assumption that the data points are independently and identically distributed (i.i.d.) by moving to a scenario of *structured prediction*, where the inputs are assumed to have temporal or spacial dependencies. We start by considering sequential models, which correspond to a *chain structure*: for instance, the words in a sentence. In this lecture, we will use part-of-speech tagging as our example task.

We start by defining the notation for this lecture in Section 2.1. Afterwards, in section 2.2, we focus on the well known Hidden Markov Models and in Section 2.3 we describe how to estimate its parameters from labeled data. In Section 2.4 we explain the inference algorithms (Viterbi and Forward-Backward) for sequence models. These inference algorithms will be fundamental for the rest of this lecture, as well as for the next lecture on *discriminative* training of sequence models. In Section 2.6 we describe the task of Part-of-Speech tagging, and how the Hidden Markov Models are suitable for this task. Finally, in Section 2.7 we address unsupervised learning of Hidden Markov Models through the Expectation Maximization algorithm.

## Today's assignment

The assignment of today's class is to implement one inference algorithm for Hidden Markov Models, used to find the most likely hidden state sequence given an observation sequence.

#### 2.1 Notation

In what follows, we assume a finite set of observation labels,  $\Sigma := \{w_1, \dots, w_J\}$ , and a finite set of state labels,  $\Lambda := \{c_1, \dots, c_K\}$ . We denote by  $\Sigma^*$ ,  $\Lambda^*$  the two infinite sets of sequences obtained by grouping the elements of each label set including repetitions and the empty string  $\varepsilon^1$ . Elements of  $\Sigma^*$  and  $\Lambda^*$  are strings of observations and strings of states, respectively. Throughout this class, we assume our input set is  $\mathfrak{X} = \Sigma^*$ , and our output set is  $\mathfrak{Y} = \Lambda^*$ . In other words, our inputs are observation sequences,  $x = x_1 x_2 \dots x_N$ , for some  $N \in \mathbb{N}$ , where each  $x_i \in \Sigma$ ; given such an x, we seek the corresponding state sequence,  $y = y_1 y_2 \dots y_N$ , where each  $y_i \in \Lambda$ . We also consider two special states: the start symbol, which starts the sequence, and the stop symbol, which ends the sequence.

Moreover, in this lecture we will assume two scenarios:

- 1. Supervised learning. We will train models from a sample set of paired observation and state sequences,  $\mathcal{D}_L := \{(x^1, y^1), \dots, (x^M, y^M)\} \subseteq \mathcal{X} \times \mathcal{Y}.$
- 2. *Unsupervised learning*. We will train models from the set of observations only,  $\mathcal{D}_U := \{x^1, \dots, x^M\} \subseteq \mathcal{X}$ .

Our notation is summarized in Table 2.1.

#### 2.2 Hidden Markov Models

Hidden Markov Models (HMMs) are one of the most common sequence probabilistic models, and have been applied to a wide variety of tasks. HMMs are particular instances of directed probabilistic graphical models (or Bayesian networks) which have a chain topology. In a Bayesian network, every random variable is represented

<sup>&</sup>lt;sup>1</sup>More formally, we say  $\Sigma^* := \{\varepsilon\} \cup \Sigma \cup \Sigma^2 \cup ...$  and  $\Lambda^* := \{\varepsilon\} \cup \Lambda \cup \Lambda^2 \cup ...$  is the Kleene closure of each of the two sets above.

Notation		
$\mathcal{D}_L$	training set (including labeled data)	
$\mathcal{D}_{U}$	training set (unlabeled data only)	
M	number of training examples	
$x = x_1 \dots x_N$	observation sequence	
$y = y_1 \dots y_N$	state sequence	
N	length of the sequence	
$x_i$	observation at position $i$ in the sequence, $i \in \{1,, N\}$	
$y_i$	state at position $i$ in the sequence, $i \in \{1,, N\}$	
Σ	observation set	
J	number of distinct observation labels	
$w_{j}$	particular observation, $j \in \{1, \dots, J\}$	
Λ	state set	
K	number of distinct state labels	
$c_k$	particular state, $k \in \{1,, K\}$	

Table 2.1: General notation used in this class

as a node in a graph, and the edges in the graph are directed and represent probabilistic dependencies between the random variables. For an HMM, the random variables are divided into two sets, the *observed variables*,  $X = X_1 \dots X_N$ , and the *hidden variables*  $Y = Y_1 \dots Y_N$ . In the HMM terminology, the observed variables are called *observations*, and the hidden variables are called *states*. The states are generated according to a first order Markov process, in which the  $i^{th}$  state  $Y_i$  depends only of the previous state  $Y_{i-1}$ . Two special states are the start symbol, which starts the sequence, and the stop symbol, which ends the sequence. In addition, states emit observation symbols. In an HMM, it is assumed that all observations are independent given the states that generated them.

As you may find out with today's assignment, implementing the inference routines of the HMM can be challenging. We start with a small and very simple (also very unrealistic!) example. The idea is that you may compute the desired quantities by hand and check if your implementation yields the correct result.

**Example 2.1** Consider a person who is only interested in four activities: walking in the park (walk), shopping (shop), cleaning the apartment (clean) and playing tennis (tennis). Also, consider that the choice of what the person does on a given day is determined exclusively by the weather on that day, which can be either rainy or sunny. Now, supposing that we observe what the person did on a sequence of days, the question is: can we use that information to predict the weather on each of those days? To tackle this problem, we assume that the weather behaves as a discrete Markov chain: the weather on a given day depends only on the weather on the previous day. The entire system can be described as an HMM.

For example, assume we are asked to predict the weather conditions on two different sequences of days. During these two sequences, we observed the person performing the following activities:

- "walk walk shop clean"
- "clean walk tennis walk"

This will be our test set.

Moreover, and in order to train our model, we are given access to three different sequences of days, containing both the activities performed by the person and the weather on those days, namely:

- "walk/rainy walk/sunny shop/sunny clean/sunny"
- "walk/rainy walk/rainy shop/rainy clean/sunny"
- "walk/sunny shop/sunny shop/sunny clean/sunny"

This will be our training set.

Figure 2.2 shows the HMM model for the first sequence of the training set, which already includes the start and stop symbols. The notation is summarized in Table 2.2.

**Exercise 2.1** Load the simple sequence dataset. From the ipython command line create a simple sequence object and look at the training and test set.

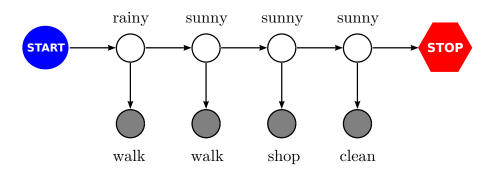


Figure 2.1: Diagram showing the conditional independence relations of the HMM. As an example, the variables are the values of the first sentence of the training set of the simple sequence.

HMM Notation		
x	observed sequence "walk walk shop clean"	
N=4	observation length	
i	position in the sentence: $i \in \{1 N\}$	
$\Sigma = \{  ext{walk, shop, clean, tennis} \}$	observation set	
j	index into the observation set $j \in \{1,, J\}$	
$X_i = w_j$	observation at position $i$ has value $w_j$	
$\Lambda = \{ \text{rainy, sunny} \}$	state set	
k	index into state set $k \in \{1,, K\}$	
$Y_i = c_k$	state at position $i$ has value $c_k$	

Table 2.2: HMM notation for the simple example.

Get in touch with the classes used to store the sequences, you will need this for the next exercise. Note that each label is internally stored as a number. This number can be used as index of an array to store information regarding that label.

```
for sequence in simple.train.seq_list:
    print sequence

walk/rainy walk/sunny shop/sunny clean/sunny
walk/rainy walk/rainy shop/rainy clean/sunny
walk/sunny shop/sunny shop/sunny clean/sunny

for sequence in simple.train.seq_list:
    print sequence.x

[0, 0, 1, 2]
[0, 0, 1, 2]
[0, 1, 1, 2]
for sequence in simple.train.seq_list:
    print sequence.y
[0, 1, 1, 1]
```

The probability distributions  $P(Y_i|Y_{i-1})$  are called *transition probabilities*; the distributions  $P(Y_1|Y_0 = \mathtt{start})$  are the *initial probabilities*, and  $P(Y_{N+1} = \mathtt{stop}|Y_N)$  the *final probabilities*.<sup>2</sup> Finally, the distributions  $P(X_i|Y_i)$  are called *emission probabilities*.

A first order HMM model has the following independence assumptions over the joint distribution P(X = x, Y = y):

• **Independence of previous states.** The probability of being in a given state at position i only depends on the state of the previous position i - 1. Formally,

$$P(Y_i = y_i | Y_{i-1} = y_{i-1}, Y_{i-2} = y_{i-2}, \dots, Y_1 = y_1) = P(Y_i = y_i | Y_{i-1} = y_{i-1})$$

defining a first order Markov chain.<sup>3</sup>

• **Homogeneous transition.** The probability of making a transition from state  $c_l$  to state  $c_k$  is independent of the particular position in the sequence. That is, for all  $i, t \in \{1, ..., N\}$ ,

$$P(Y_i = c_k | Y_{i-1} = c_l) = P(Y_t = c_k | Y_{t-1} = c_l)$$

• **Observation independence.** The probability of observing  $X_i = x_i$  at position i is fully determined by the state  $Y_i$  at that position. Formally,

$$P(X_i = x_i | Y_1 = y_1, \dots, Y_i = y_i, \dots, Y_N = y_N) = P(X_i = x_i | Y_i = y_i)$$

This probability is independent of the particular position so, for every i and t, we can write:

$$P(X_i = w_i | Y_i = c_k) = P(X_t = w_i | Y_t = c_k)$$

These conditional independence assumptions are crucial to allow efficient inference, as it will be described.

The distributions that define the HMM model are summarized in Table 2.3. For each one of them we will use a short notation to simplify the exposition.

HMM distributions			
Name	probability distribution	short notation	array size
initial probability	$P(Y_1 = c_k   Y_0 = \text{start})$	$P_{\text{init}}(c_k \text{start})$	K
transition probability	$P(Y_i = c_k   Y_{i-1} = c_l)$	$P_{\text{trans}}(c_k c_l)$	$K \times K$
final probability	$P(Y_{N+1} = \operatorname{stop} Y_N = c_k)$	$P_{\text{final}}(\text{stop} c_k)$	K
emission probability	$P(X_i = w_j   Y_i = c_k)$	$P_{\text{emiss}}(w_j c_k)$	$J \times K$

Table 2.3: HMM probability distributions.

The joint distribution can be expressed as:

$$P(X_1 = x_1, \dots, X_N = x_N, Y_1 = y_1, \dots, Y_N = y_N) = P_{\text{init}}(y_1|\text{start}) \times \left(\prod_{i=1}^{N-1} P_{\text{trans}}(y_{i+1}|y_i)\right) \times P_{\text{final}}(\text{stop}|y_N) \times \prod_{i=1}^{N} P_{\text{emiss}}(x_i|y_i), \tag{2.1}$$

which for the example of Figure 2.1 is:

$$\begin{split} P(X_1 = x_1, \dots, X_4 = x_4, Y_1 = y_1, \dots, Y_4 = y_4) = \\ P_{\text{init}}(\text{rainy}|\text{start}) \times P_{\text{trans}}(\text{sunny}|\text{rainy}) \times P_{\text{trans}}(\text{sunny}|\text{sunny}) \times P_{\text{trans}}(\text{sunny}|\text{sunny}) \times \\ P_{\text{final}}(\text{stop}|\text{sunny}) \times P_{\text{emiss}}(\text{walk}|\text{rainy}) \times P_{\text{emiss}}(\text{walk}|\text{sunny}) \times P_{\text{emiss}}(\text{shop}|\text{sunny}) \\ \times P_{\text{emiss}}(\text{clean}|\text{sunny}). \end{split}$$

<sup>&</sup>lt;sup>2</sup>Note that the initial and final probabilities are asymmetric.

<sup>&</sup>lt;sup>3</sup>The order of the Markov chain depends on the number of previous positions taken into account. The remainder of the exposition can be easily extended to higher order HMMs, giving the model more generality, but making inference more expensive.

In the next section we turn our attention to estimating the different probability distributions of the model.

## 2.3 Finding the Maximum Likelihood Parameters

One important problem in HMMs is to estimate the model parameters, *i.e.*, the distributions depicted in Table 2.3. We will refer to the set of all these parameters as  $\theta$ . In a supervised setting, the HMM model is trained to maximize the joint log-likelihood of the data. Given a dataset  $\mathfrak{D}_L$ , the objective being optimized is:

$$\arg\max_{\theta} \sum_{m=1}^{M} \log P_{\theta}(X = x^{m}, Y = y^{m}), \tag{2.3}$$

where  $P_{\theta}(X = x^m, Y = y^m)$  is given by Eq. 2.1.

In some applications (*e.g.* speech recognition) the observation variables are continuous, hence the emission distributions are real-valued (*e.g.* mixtures of Gaussians). In our case, both the state set and the observation set are discrete (and finite), therefore we use multinomial distributions for the emission and transition probabilities. Multinomial distributions are attractive for several reasons: first of all, they are easy to implement; secondly, the maximum likelihood estimation of the parameters has a simple closed form. The parameters are just normalized counts of events that occur in the corpus (the same as the Naïve Bayes from previous class).

Given our labeled corpus  $\mathcal{D}_L$ , the estimation process consists of counting how many times each event occurs in the corpus and normalize the counts to form proper probability distributions. Let us define the following quantities, called sufficient statistics, that represent the counts of each event in the corpus:

Initial Counts: 
$$C_{\text{init}}(c_k) = \sum_{m=1}^{M} \mathbf{1}(y_1^m = c_k);$$
 (2.4)

**Transition Counts**: 
$$C_{\text{trans}}(c_k, c_l) = \sum_{m=1}^{M} \sum_{i=2}^{N} \mathbf{1}(y_i^m = c_k \wedge y_{i-1}^m = c_l);$$
 (2.5)

Final Counts: 
$$C_{\text{final}}(c_k) = \sum_{m=1}^{M} \mathbf{1}(y_N^m = c_k);$$
 (2.6)

Emission Counts: 
$$C_{\text{emiss}}(w_j, c_k) = \sum_{m=1}^{M} \sum_{i=1}^{N} \mathbf{1}(x_i^m = w_j \wedge y_i^m = c_k);$$
 (2.7)

Here  $y_i^m$ , the underscript denotes the state index position for a given sequence, and the superscript denotes the sequence index in the dataset, and the same applies for the observations. Note that 1 is an indicator function that has the value 1 when the particular event happens, and zero otherwise. In other words, the previous equations go through the training corpus and count how often each event occurs. For example, Eq. 2.5 counts how many times  $c_k$  follows state  $c_l$ . Therefore,  $C_{\text{trans}}(\text{sunny}, \text{rainy})$  contains the number of times that a sunny day followed a rainy day.

After computing the counts, one can perform some sanity checks to make sure the implementation is correct. Summing over all entries of each count table we should observe the following:

- **Initial Counts** Should sum to the number of sentences:  $\sum_{k=1}^{K} C_{\text{init}}(c_k) = M$
- Transition/Final Counts Should sum to the number of tokens:  $\sum_{k,l=1}^{K} C_{\text{trans}}(c_k, c_l) + \sum_{k=1}^{K} C_{\text{final}}(c_k) = MN$
- Emission Counts Should sum to the number of tokens:  $\sum_{j=1}^{J} \sum_{k=1}^{K} C_{\text{emiss}}(w_j, c_k) = MN$ .

Using the sufficient statistics (counts) the parameter estimates are:

$$P_{\text{init}}(c_k|\text{start}) = \frac{C_{\text{init}}(c_k)}{\sum_{l=1}^{K} C_{\text{init}}(c_l)}$$

$$P_{\text{final}}(\text{stop}|c_l) = \frac{C_{\text{final}}(c_l)}{\sum_{k=1}^{K} C_{\text{trans}}(c_k, c_l) + C_{\text{final}}(c_l)}$$
(2.8)

$$P_{\text{final}}(\text{stop}|c_l) = \frac{C_{\text{final}}(c_l)}{\sum_{k=1}^{K} C_{\text{trans}}(c_k, c_l) + C_{\text{final}}(c_l)}$$
(2.9)

$$P_{\text{trans}}(c_k|c_l) = \frac{C_{\text{trans}}(c_k, c_l)}{\sum_{\nu=1}^{K} C_{\text{trans}}(c_{\nu}, c_l) + C_{\text{final}}(c_l)}$$
(2.10)

$$P_{\text{trans}}(c_{k}|c_{l}) = \frac{C_{\text{trans}}(c_{k},c_{l})}{\sum_{p=1}^{K} C_{\text{trans}}(c_{p},c_{l}) + C_{\text{final}}(c_{l})}$$

$$P_{\text{emiss}}(w_{j}|c_{k}) = \frac{C_{\text{emiss}}(w_{j},c_{k})}{\sum_{q=1}^{J} C_{\text{emiss}}(w_{q},c_{k})}$$
(2.10)

**Exercise 2.2** *The provided function* train\_supervised *from the* hmm.py *file implements the above parameter estimates.* Run this function given the simple dataset above and look at the estimated probabilities. Are they correct? You can also check the variables ending in \_counts instead of \_probs to see the raw counts (for example, typing hmm.initial\_counts will show you the raw counts of initial states). How are the counts related to the probabilities?

```
import lxmls.sequences.hmm as hmmc
hmm = hmmc.HMM(simple.x_dict, simple.y_dict)
hmm.train_supervised(simple.train)
print "Initial Probabilities:", hmm.initial_probs
[ 0.66666667  0.333333333]
print "Transition Probabilities:", hmm.transition_probs
          0. 1
[ 0.5
          0.625]]
print "Final Probabilities:", hmm.final_probs
[ 0.
         0.3751
print "Emission Probabilities", hmm.emission_probs
[[ 0.75
          0.25 ]
          0.375]
[ 0.25
          0.375]
 [ 0.
```

#### 2.4 Decoding a Sequence

Given the learned parameters and a new observation sequence  $x = x_1 \dots x_N$ , we want to find the sequence of hidden states  $y^* = y_1^* \dots y_N^*$  that "best" explains it. This is called the *decoding* problem. There are several ways to define what we mean by the "best"  $y^*$ , depending on our goal: for instance, we may want to minimize the probability of error on each hidden variable  $Y_{ij}$  or we may want to find the best assignment to the sequence  $Y_1 \dots Y_N$  as a whole. Therefore, finding the best sequence can be accomplished through different approaches:

 A first approach, normally called posterior decoding or minimum risk decoding, consists in picking the highest state posterior for each position *i* in the sequence:

$$y_i^* = \underset{y_i \in \Lambda}{\arg \max} P(Y_i = y_i | X_1 = x_1, \dots, X_N = x_N).$$
 (2.12)

Note, however, that this approach does not guarantee that the sequence  $y^* = y_1^* \dots y_N^*$  will be a valid sequence of the model. For instance, there might be a transition between two of the best state posteriors with probability zero.

• A second approach, called Viterbi decoding, consists in picking the best global hidden state sequence:

$$y^* = \underset{y=y_1...y_N}{\arg \max} P(Y_1 = y_1, ..., Y_N = y_N | X_1 = x_1, ..., X_N = x_N)$$

$$= \underset{y=y_1...y_N}{\arg \max} P(Y_1 = y_1, ..., Y_N = y_N, X_1 = x_1, ..., X_N = x_N).$$
(2.13)

Both approaches (which will be explained in Sections 2.4.2 and 2.4.3, respectively) rely on dynamic programming and make use of the independence assumptions of the HMM model. Moreover, they use an alternative representation of the HMM called a *trellis*.

A trellis unfolds all possible states for each position and it makes explicit the independence assumption: each position only depends on the previous position. Here, each column represents a position in the sequence and each row represents a possible state. Figure 2.2 shows the trellis for the particular example in Figure 2.1.

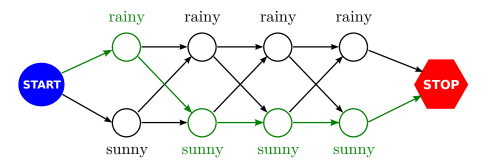


Figure 2.2: Trellis representation of the HMM in Figure 2.1, for the observation sequence "walk walk shop clean", where each hidden variable can take the values rainy or sunny.

Considering the trellis representation, note that we can include the following information:

- an *initial probability* to the arrows that depart from the start symbol;
- a *final probability* to the arrows that reach the stop symbol;
- a transition probability to the remaining arrows; and,
- an *emission probability* to each circle, which is the probability that the observed symbol is emitted by that particular state.

For convenience, we will be working with log-probabilities, rather than probabilities.<sup>4</sup> Therefore, if we associate to each circle and arrow in the trellis a score that corresponds to the log-probabilities above, and if we define the score of a path connecting the start and stop symbols as the sum of the scores of the circles and arrows it traverses, then the goal of finding the most likely sequence of states (Viterbi decoding) corresponds to finding the path with the highest score.

The trellis scores are given by the following expressions:

• For each state  $c_k$ :

$$score_{init}(c_k) = log P_{init}(Y_1 = c_k | start).$$
 (2.14)

• For each position  $i \in 1, ..., N-1$  and each pair of states  $c_k$  and  $c_l$ :

$$score_{trans}(i, c_k, c_l) = log P_{trans}(Y_{i+1} = c_k | Y_i = c_l).$$
(2.15)

• For each state  $c_1$ :

$$score_{final}(c_l) = log P_{final}(stop|Y_N = c_l).$$
 (2.16)

• For each position  $i \in 1, ..., N$  and state  $c_k$ :

$$score_{emiss}(i, c_k) = log P_{emiss}(X_i = x_i | Y_i = c_k).$$
 (2.17)

<sup>&</sup>lt;sup>4</sup>This will be motivated further in Section 2.4.1, where we describe how operations can be performed efficiently in the log-domain.

In the next exercise, you will compute the trellis scores.

**Exercise 2.3** Convince yourself that the score of a path in the trellis (summing over the scores above) is equivalent to the log-probability  $\log P(X = x, Y = y)$ , as defined in Eq. 2.2. Use the given function compute\_scores on the first training sequence and confirm that the values are correct. You should get the same values as presented below.

```
initial_scores, transition_scores, final_scores, emission_scores = hmm.compute_scores(
   simple.train.seq_list[0])
print initial_scores
[-0.40546511 -1.09861229]
print transition_scores
[[[-0.69314718
                -infl
  [-0.69314718 -0.47000363]]
 [[-0.69314718
                     -inf]
  [-0.69314718 -0.47000363]]
 [[-0.69314718
                     -infl
  [-0.69314718 -0.47000363]]]
print final_scores
        -inf -0.980829251
print emission_scores
[[-0.28768207 -1.38629436]
 [-0.28768207 -1.38629436]
 [-1.38629436 -0.98082925]
       -inf -0.98082925]]
```

*Note that scores which are*  $-\infty$  (-inf) *correspond to zero-probability events.* 

#### 2.4.1 Computing in log-domain

We will see that the decoding algorithms will need to multiply twice as many probability terms as the length N of the sequence. This may cause underflowing problems when N is large, since the nested multiplication of numbers smaller than 1 may easily become smaller than the machine precision. To avoid that problem, Rabiner (1989) presents a scaled version of the decoding algorithms that avoids this problem. An alternative, which is widely used, is computing in the log-domain. That is, instead of manipulating probabilities, manipulate log-probabilities (the scores presented above). Every time we need to multiply probabilities, we can sum their log-representations, since:

$$\log(\exp(a) \times \exp(b)) = a + b. \tag{2.18}$$

Sometimes, we need to add probabilities. In the log domain, this requires us to compute

$$\log(\exp(a) + \exp(b)) = a + \log(1 + \exp(b - a)), \tag{2.19}$$

where we assume that a is smaller than b.

Exercise 2.4 Look at the module sequences/log\_domain.py. This module implements a function logsum\_pair (logx, logy) to add two numbers represented in the log-domain; it returns their sum also represented in the log-domain. The function logsum(logv) sums all components of an array represented in the log-domain. This will be used later in our decoding algorithms. To observe why this is important, type the following:

```
import numpy as np
a = np.random.rand(10)
np.log(sum(np.exp(a)))
2.8397172643228661
```

```
np.log(sum(np.exp(10*a)))
10.121099917705818

np.log(sum(np.exp(100*a)))
93.159220940569128

np.log(sum(np.exp(1000*a)))
inf

from lxmls.sequences.log_domain import *
logsum(a)
2.8397172643228665

logsum(10*a)
10.121099917705818

logsum(100*a)
93.159220940569114

logsum(1000*a)
925.88496219586864
```

#### 2.4.2 Posterior Decoding

Posterior decoding consists in picking state with the highest posterior for each position in the sequence independently; for each i = 1, ..., N:

$$y_i^* = \underset{y_i \in \Lambda}{\arg \max} P(Y_i = y_i | X = x).$$
 (2.20)

The **sequence posterior distribution** is the probability of a particular hidden state sequence given that we have observed a particular sequence. Moreover, we will be interested in two other posteriors distributions: the **state posterior distribution**, corresponding to the probability of being in a given state in a certain position given the observed sequence; and the **transition posterior distribution**, which is the probability of making a particular transition, from position i to i + 1, given the observed sequence. They are formally defined as follows:

Sequence Posterior: 
$$P(Y = y | X = x) = \frac{P(X = x, Y = y)}{P(X = x)};$$
 (2.21)

**State Posterior**: 
$$P(Y_i = y_i | X = x);$$
 (2.22)

**Transition Posterior**: 
$$P(Y_{i+1} = y_{i+1}, Y_i = y_i | X = x)$$
. (2.23)

To compute the posteriors, a first step is to be able to compute the likelihood of the sequence P(X = x), which corresponds to summing the probability of all possible hidden state sequences.

**Likelihood**: 
$$P(X = x) = \sum_{y \in \Lambda^N} P(X = x, Y = y).$$
 (2.24)

The number of possible hidden state sequences is exponential in the length of the sequence ( $|\Lambda|^N$ ), which makes the sum over all of them hard. In our simple example, there are  $2^4=16$  paths, which we can actually explicitly enumerate and calculate their probability using Equation 2.1. But this is as far as it goes: for example, for Part-of-Speech tagging with a small tagset of 12 tags and a medium size sentence of length 10, there are  $12^{10}=61917364224$  such paths. Yet, we must be able to compute this sum (sum over  $y\in\Lambda^N$ ) to compute the above likelihood formula; this is called the inference problem. For sequence models, there is a well known dynamic programming algorithm, the **Forward-Backward** (FB) algorithm, which allows the computation to be performed in linear time, 5 by making use of the independence assumptions.

The FB algorithm relies on the independence of previous states assumption, which is illustrated in the trellis view by having arrows only between consecutive states. The FB algorithm defines two auxiliary probabilities, the forward probability and the backward probability.

<sup>&</sup>lt;sup>5</sup>The runtime is linear with respect to the sequence length. More precisely, the runtime is  $O(N|\Lambda|^2)$ . A naive enumeration would cost  $O(|\Lambda|^N)$ .

#### Efficient forward probability computation

The forward probability represents the probability that in position i we are in state  $Y_i = c_k$  and that we have observed  $x_1, \ldots, x_i$  up to that position. Therefore, its mathematical expression is:

Forward Probability: forward
$$(i, c_k) = P(Y_i = c_k, X_1 = x_1, \dots, X_i = x_i)$$
 (2.25)

Using the independence assumptions of the HMM we can compute forward  $(i, c_k)$  using all the forward computations {forward (i-1,c) for  $c \in \Lambda$ }. In order to facilitate the notation of the following argument we will denote by  $x_{i:j}$  the assignment  $X_i = x_i, \ldots, X_j = x_j$ . Therefore we can write forward  $(i, y_i)$  as  $P(y_i, x_{1:i})$  and rewrite the forward expression as follows:

$$P(y_i, x_{1:i}) = \sum_{y_{i-1} \in \Lambda} P(y_i, y_{i-1}, x_{1:i}) = \sum_{y_{i-1} \in \Lambda} P(x_i | y_i, y_{i-1}, x_{1:i-1}) \cdot P(y_i | y_{i-1}, x_{1:i-1}) \cdot P(y_{i-1}, x_{1:i-1})$$
(2.26)

Using the **Observation independence** and the **Independence of previous states** properties of the first order HMM we have  $P(x_i|y_i,y_{i-1},x_{1:i-1}) = P(x_i|y_i)$  and  $P(y_i|y_{i-1},x_{1:i-1}) = P(y_i|y_{i-1})$ . Therefore equation (2.26) can be written, for  $i \in \{2,\ldots,N\}$  (where N is the length of the sequence), as

forward
$$(i, y_i) = \sum_{y_{i-1} \in \Lambda} P(x_i|y_i,) \cdot P(y_i|y_{i-1}) \cdot \text{forward}(i-1, y_{i-1})$$
 (2.27)

Using equation (2.27) we have proved that the forward probability can be defined by the following recurrence rule:

$$forward(1,c_k) = P_{init}(c_k|start) \times P_{emiss}(x_1|c_k)$$
(2.28)

forward
$$(i, c_k) = \left(\sum_{c_l \in \Lambda} P_{\text{trans}}(c_k|c_l) \times \text{forward}(i-1, c_l)\right) \times P_{\text{emiss}}(x_i|c_k)$$
 (2.29)

forward
$$(N+1, \text{stop}) = \sum_{c_l \in \Lambda} P_{\text{final}}(\text{stop}|c_l) \times \text{forward}(N, c_l).$$
 (2.30)

Using the forward trellis one can compute the likelihood simply as:

$$P(X = x) = \text{forward}(N + 1, \text{stop}). \tag{2.31}$$

Although the forward probability is enough to calculate the likelihood of a given sequence, we will also need the backward probability to calculate the state posteriors.

#### Efficient backward probability computation

The backward probability is similar to the forward probability, but operates in the inverse direction. It represents the probability of observing  $x_{i+1}, \ldots, x_N$  from position i + 1 up to N, given that at position i we are at state  $Y_i = c_l$ :

**Backward Probability**: backward(
$$i$$
,  $c_l$ ) =  $P(X_{i+1} = x_{i+1}, \dots, X_N = x_N | Y_i = c_l)$ . (2.32)

Using the independence assumptions of the HMM we can compute backward  $(i, c_k)$  using all the backward computations {backward (i + 1, c) for  $c \in \Lambda$ }. Therefore we can write backward  $(i, y_i)$  as  $P(x_{i+1:N}|y_i)$  and rewrite the forward expression as follows:

$$P(x_{i+1:N}|y_i) = \sum_{y_{i+1} \in \Lambda} P(x_{i+1:N}, y_{i+1}|y_i) = \sum_{y_{i+1} \in \Lambda} P(x_{i+2:N}|y_i, y_{i+1}, x_{i+1}) P(x_{i+1}, |y_{i+1}, y_i) P(y_{i+1}|y_i)$$
(2.33)

Using the previous equation we have proved that the backward probability can be defined by the following recurrence rule:

$$backward(N, c_l) = P_{final}(stop|c_l)$$
 (2.34)

$$backward(i, c_l) = \sum_{c_k \in \Lambda} P_{trans}(c_k|c_l) \times backward(i+1, c_k) \times P_{emiss}(x_{i+1}|c_k)$$
 (2.35)

$$\operatorname{backward}(i, c_l) = \sum_{c_k \in \Lambda} P_{\operatorname{trans}}(c_k | c_l) \times \operatorname{backward}(i + 1, c_k) \times P_{\operatorname{emiss}}(x_{i+1} | c_k)$$

$$\operatorname{backward}(0, \operatorname{start}) = \sum_{c_k \in \Lambda} P_{\operatorname{init}}(c_k | \operatorname{start}) \times \operatorname{backward}(1, c_k) \times P_{\operatorname{emiss}}(x_1 | c_k).$$

$$(2.35)$$

Using the backward trellis one can compute the likelihood simply as:.

$$P(X = x) = \text{backward}(0, \text{start}). \tag{2.37}$$

#### The forward backward algorithm

We have seen how we can compute the probability of a sequence x using the the forward and backward probabilities by computing forward (N+1, stop) and backward (0, start) respectively. Moreover, the probability of a sequence x can be computed with both forward and backward probabilities at a particular position i. The probability of a given sequence x at any position i in the sequence can be computed as follows 6:

$$P(X = x) = \sum_{c_k \in \Lambda} P(X_1 = x_1, \dots, X_N = x_N, Y_i = c_k)$$

$$= \sum_{c_k \in \Lambda} \underbrace{P(X_1 = x_1, \dots, X_i = x_i, Y_i = c_k)}_{\text{forward}(i, c_k)} \times \underbrace{P(X_{i+1} = x_{i+1}, \dots, X_N = x_N | Y_i = c_k)}_{\text{backward}(i, c_k)}$$

$$= \sum_{c_k \in \Lambda} \text{forward}(i, c_k) \times \text{backward}(i, c_k). \tag{2.38}$$

This equation will work for any choice of i. Although redundant, this fact is useful when implementing an HMM as a sanity check that the computations are being performed correctly, since one can compute this expression for several *i*; they should all yield the same value.

Algorithm 7 shows the pseudo code for the forward backward algorithm. The reader can notice that the forward and backward computations in the algorithm make use of  $P_{emiss}$  and  $P_{trans}$ . There are a couple of details that should be taken into account if the reader wants to understand the algorithm using scores instead of probabilities.

- $forward(i,\hat{c})$  is computed using  $P_{emiss}(x_i|\hat{c})$  which does not depend on the sum over all possible states  $c_k \in \Lambda$ . Therefore when taking the logarithm of the sum over all possible states the recurrence of the forward computations can be split as a sum of two logarithms.
- backward(i,  $\hat{c}$ ) is computed using  $P_{\text{trans}}(c_k|\hat{c})$  and  $P_{\text{emiss}}(x_{i+1}|c_k)$  both of which depend on  $c_k$ . Therefore when taking the logarithm of the sum the expression cannot be split as a sum of logarithms.

Exercise 2.5 Run the provided forward-backward algorithm on the first train sequence. Observe that both the forward and the backward passes give the same log-likelihood.

```
log_likelihood, forward = hmm.decoder.run_forward(initial_scores, transition_scores,
   final_scores, emission_scores)
print 'Log-Likelihood =', log_likelihood
Log-Likelihood = -5.06823232601
log_likelihood, backward = hmm.decoder.run_backward(initial_scores, transition_scores,
   final_scores, emission_scores)
print 'Log-Likelihood =', log_likelihood
Log-Likelihood = -5.06823232601
```

<sup>&</sup>lt;sup>6</sup> The second equality in 2.38 can be justified as follows. Since  $Y_i = y_i$  is observed then  $X_{i+1}, \dots, X_N$  is conditionally independent from any  $X_k$  for  $k \leq i$ . Therefore  $P(x_1, ..., x_N, y_i) = P(x_{i+1}, ..., x_N | y_i, x_1, ..., x_i) \cdot P(y_i, x_1, ..., x_i) = P(x_{i+1}, ..., x_N | y_i) \cdot P(y_i, x_1, ..., x_i) = P(x_{i+1}, ..., x_N | y_i) \cdot P(y_i, x_1, ..., x_i) = P(x_{i+1}, ..., x_N | y_i) \cdot P(y_i, x_1, ..., x_i) = P(x_{i+1}, ..., x_N | y_i) \cdot P(y_i, x_1, ..., x_i) = P(x_{i+1}, ..., x_N | y_i) \cdot P(y_i, x_1, ..., x_i) = P(x_{i+1}, ..., x_N | y_i) \cdot P(y_i, x_1, ..., x_i) = P(x_{i+1}, ..., x_N | y_i) \cdot P(y_i, x_1, ..., x_i) = P(x_{i+1}, ..., x_N | y_i) \cdot P(y_i, x_1, ..., x_i) = P(x_{i+1}, ..., x_N | y_i) \cdot P(y_i, x_1, ..., x_i) = P(x_{i+1}, ..., x_N | y_i) \cdot P(y_i, x_1, ..., x_i) = P(x_{i+1}, ..., x_N | y_i) \cdot P(y_i, x_1, ..., x_i) = P(x_{i+1}, ..., x_N | y_i) \cdot P(y_i, x_1, ..., x_i) = P(x_{i+1}, ..., x_N | y_i) \cdot P(y_i, x_1, ..., x_i) = P(x_{i+1}, ..., x_N | y_i) \cdot P(y_i, x_1, ..., x_N | y_i) \cdot P$  $backward(i, y_i) \cdot forward(i, y_i)$ .

#### Algorithm 7 Forward-Backward algorithm

```
1: input: sequence x_1, \ldots, x_N, scores P_{\text{init}}, P_{\text{trans}}, P_{\text{final}}, P_{\text{emiss}}
 2: Forward pass: Compute the forward probabilities
 3: for c_k \in \Lambda do
        forward(1, c_k) = P_{\text{init}}(c_k | \text{start}) \times P_{\text{emiss}}(x_1 | c_k)
 5: end for
 6: for i = 2 to N do
         for \hat{c} \in \Lambda do
 7:
            forward(i, \hat{c}) = \left(\sum_{c_k \in \Lambda} P_{\text{trans}}(\hat{c}|c_k) \times \text{forward}(i-1, c_k)\right) \times P_{\text{emiss}}(x_i|\hat{c})
 8:
 9:
         end for
10: end for
11: Backward pass: Compute the backward probabilities
12: for c_k \in \Lambda do
        backward(N, c_k) = P_{final}(stop|c_k)
13:
14: end for
15: for i = N - 1 to 1 do
        for \hat{c} \in \Lambda do
16:
            \mathsf{backward}(i, \hat{c}) = \sum_{c_k \in \Lambda} P_{\mathsf{trans}}(c_k | \hat{c}) \times \mathsf{backward}(i+1, c_k) \times P_{\mathsf{emiss}}(x_{i+1} | c_k)
17:
18:
         end for
19: end for
20: output: The forward and backward probabilities.
```

Given the forward and backward probabilities, one can compute both the state and transition posteriors (you can hint why by looking at the term inside the sum in Eq. 2.38).

$$\begin{aligned} \textbf{State Posterior:} \quad & P(Y_i = y_i | X = x) = \frac{\text{forward}(i, y_i) \times \text{backward}(i, y_i)}{P(X = x)}; \end{aligned} \end{aligned} \tag{2.39}$$
 
$$\begin{aligned} & \textbf{Transition Posterior:} \quad & P(Y_i = y_i, Y_{i+1} = y_{i+1} | X = x) = \\ & \frac{\text{forward}(i, y_i) \times P_{\text{trans}}(y_{i+1} | y_i) \times P_{\text{emiss}}(x_{i+1} | y_{i+1}) \times \text{backward}(i+1, y_{i+1})}{P(X = x)}. \end{aligned} \end{aligned}$$

A graphical representation of these posteriors is illustrated in Figure 2.3. On the left it is shown that forward  $(i, y_i) \times \text{backward}(i, y_i)$  returns the sum of all paths that contain the state  $y_i$ , weighted by P(X = x); on the right we can see that forward  $(i, y_i) \times P_{\text{trans}}(y_{i+1}|y_i) \times P_{\text{emiss}}(x_{i+1}|y_{i+1}) \times \text{backward}(i+1, y_{i+1})$  returns the same for all paths containing the edge from  $y_i$  to  $y_{i+1}$ .

As a practical example, given that the person performs the sequence of actions "walk walk shop clean", we want to know the probability of having been raining in the second day. The state posterior probability for this event can be seen as the probability that the sequence of actions above was generated by a sequence of weathers and where it was raining in the second day. In this case, the possible sequences would be all the sequences which have rainy in the second position.

Using the state posteriors, we are ready to perform posterior decoding. The strategy is to compute the state posteriors for each position  $i \in \{1, ..., N\}$  and each state  $c_k \in \Lambda$ , and then pick the arg-max at each position:

$$\widehat{y}_i := \underset{y_i \in \Lambda}{\arg \max} P(Y_i = y_i | X = x). \tag{2.41}$$

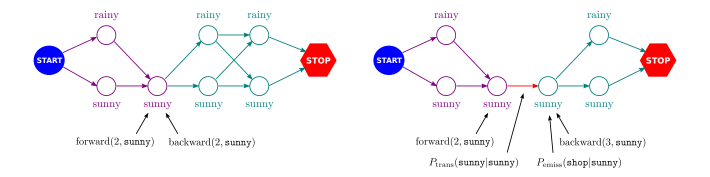


Figure 2.3: A graphical representation of the components in the state and transition posteriors. Recall that the observation sequence is "walk walk shop clean".

**Exercise 2.6** Compute the node posteriors for the first training sequence (use the provided compute\_posteriors function), and look at the output. Note that the state posteriors are a proper probability distribution (the lines of the result sum to 1).

#### **Exercise 2.7** Run the posterior decode on the first test sequence, and evaluate it.

```
y_pred = hmm.posterior_decode(simple.test.seq_list[0])
print "Prediction test 0:", y_pred

walk/rainy walk/rainy shop/sunny clean/sunny

print "Truth test 0:", simple.test.seq_list[0]

walk/rainy walk/sunny shop/sunny clean/sunny
```

#### *Do the same for the second test sequence:*

```
y_pred = hmm.posterior_decode(simple.test.seq_list[1])
print "Prediction test 1:", y_pred

clean/rainy walk/rainy tennis/rainy walk/rainy

print "Truth test 1:", simple.test.seq_list[1]

clean/sunny walk/sunny tennis/sunny walk/sunny
```

What is wrong? Note the observations for the second test sequence: the observation tennis was never seen at training time, so the probability for it will be zero (no matter what state). This will make all possible state sequences have zero probability. As seen in the previous lecture, this is a problem with generative models, which can be corrected using smoothing (among other options).

Change the train\_supervised method to add smoothing:

```
def train_supervised(self, sequence_list, smoothing):
```

Try, for example, adding 0.1 to all the counts, and repeating this exercise with that smoothing. What do you observe?

```
hmm.train_supervised(simple.train, smoothing=0.1)
y_pred = hmm.posterior_decode(simple.test.seq_list[0])
print "Prediction test 0 with smoothing:", y_pred
walk/rainy walk/rainy shop/sunny clean/sunny
print "Truth test 0:", simple.test.seq_list[0]
walk/rainy walk/sunny shop/sunny clean/sunny
y_pred = hmm.posterior_decode(simple.test.seq_list[1])
print "Prediction test 1 with smoothing:", y_pred
clean/sunny walk/sunny tennis/sunny walk/sunny
print "Truth test 1:", simple.test.seq_list[1]
clean/sunny walk/sunny tennis/sunny walk/sunny
```

#### 2.4.3 Viterbi Decoding

**Viterbi decoding** consists in picking the best global hidden state sequence  $\hat{y}$  as follows:

$$\widehat{y} = \underset{y \in \Lambda^N}{\arg\max} P(Y = y | X = x) = \underset{y \in \Lambda^N}{\arg\max} P(X = x, Y = y). \tag{2.42}$$

The Viterbi algorithm is very similar to the forward procedure of the FB algorithm, making use of the same trellis structure to efficiently represent the exponential number of sequences without prohibitive computation costs. In fact, the only difference from the forward-backward algorithm is in the recursion 2.29 where instead of summing over all possible hidden states, we take their maximum.

**Viterbi** viterbi
$$(i, y_i) = \max_{y_1...y_{i-1}} P(Y_1 = y_1, ..., Y_i = y_i, X_1 = x_1, ..., X_i = x_i)$$
 (2.43)

The Viterbi trellis represents the path with maximum probability in position i when we are in state  $Y_i = y_i$ and that we have observed  $x_1, \ldots, x_i$  up to that position. The Viterbi algorithm is defined by the following recurrence:

viterbi(1, 
$$c_k$$
) =  $P_{\text{init}}(c_k|\text{start}) \times P_{\text{emiss}}(x_1|c_k)$  (2.44)

$$viterbi(i, c_k) = \left(\max_{c_l \in \Lambda} P_{trans}(c_k|c_l) \times viterbi(i-1, c_l)\right) \times P_{emiss}(x_i|c_k)$$
(2.45)

$$viterbi(i, c_k) = \left(\max_{c_l \in \Lambda} P_{trans}(c_k|c_l) \times viterbi(i-1, c_l)\right) \times P_{emiss}(x_i|c_k)$$

$$backtrack(i, c_k) = \left(\arg\max_{c_l \in \Lambda} P_{trans}(c_k|c_l) \times viterbi(i-1, c_l)\right)$$

$$viterbi(N+1, stop) = \max_{c_l \in \Lambda} P_{final}(stop|c_l) \times viterbi(N, c_l)$$

$$(2.45)$$

$$viterbi(N+1, stop) = \max_{c_l \in \Lambda} P_{final}(stop|c_l) \times viterbi(N, c_l)$$
(2.47)

$$\operatorname{backtrack}(N+1,\operatorname{stop}) = \operatorname{arg\,max}_{c_l \in \Lambda} P_{\operatorname{final}}(\operatorname{stop}|c_l) \times \operatorname{viterbi}(N,c_l). \tag{2.48}$$

Algorithm 8 shows the pseudo code for the Viterbi algorithm. Note the similarity with the forward algorithm. The only differences are:

- Maximizing instead of summing;
- Keeping the argmax's to backtrack.

#### Algorithm 8 Viterbi algorithm

```
1: input: sequence x_1, \ldots, x_N, scores P_{\text{init}}, P_{\text{trans}}, P_{\text{final}}, P_{\text{emiss}}
 2: Forward pass: Compute the best paths for every end state
 3: Initialization
 4: for c_k \in \Lambda do
        viterbi(1, c_k) = P_{\text{init}}(c_k|\text{start}) \times P_{\text{emiss}}(x_1|c_k)
 6: end for
 7: for i = 2 to N do
         for c_k \in \Lambda do
            viterbi(i, c_k) = \left(\max_{c_l \in \Lambda} P_{\text{trans}}(c_k | c_l) \times \text{viterbi}(i - 1, c_l)\right) \times P_{\text{emiss}}(x_i | c_k)
            backtrack(i, c_k) = \left( arg \max_{c_l \in \Lambda} P_{trans}(c_k | c_l) \times viterbi(i - 1, c_l) \right)
10:
         end for
11:
12: end for
13: \max_{u \in \Lambda^N} P(X = x, Y = y) := \max_{c_l \in \Lambda} P_{\text{final}}(\text{stop}|c_l) \times \text{viterbi}(N, c_l)
15: Backward pass: backtrack to obtain the most likely path
16: \widehat{y}_N = \operatorname{arg\,max}_{c_l \in \Lambda} P_{\text{final}}(\operatorname{stop}|c_l) \times \operatorname{viterbi}(N, c_l)
17: for i = N - 1 to 1 do
18: \hat{y}_i = \text{backtrack}(i+1, \hat{y}_{i+1})
19: end for
20: output: the viterbi path \hat{y}.
```

## 2.5 Assignment

With the previous theoretical background, you have the necessary tools to solve today's assignment.

Exercise 2.8 Implement a method for performing Viterbi decoding in file sequence\_classification\_decoder.py.

```
def run_viterbi(self, initial_scores, transition_scores, final_scores,
  emission_scores):
```

Hint: look at the implementation of run\_forward. Also check the help for the numpy methods max and argmax. This method will be called by

```
def viterbi_decode(self, sequence)
```

in the module sequence\_classifier.py.

Test your method on both test sequences and compare the results with the ones given.

```
hmm.train_supervised(simple.train, smoothing=0.1)
y_pred, score = hmm.viterbi_decode(simple.test.seq_list[0])
print "Viterbi decoding Prediction test 0 with smoothing:", y_pred, score
walk/rainy walk/rainy shop/sunny clean/sunny -6.02050124698

print "Truth test 0:", simple.test.seq_list[0]

walk/rainy walk/sunny shop/sunny clean/sunny

y_pred, score = hmm.viterbi_decode(simple.test.seq_list[1])
print "Viterbi decoding Prediction test 1 with smoothing:", y_pred, score
clean/sunny walk/sunny tennis/sunny walk/sunny -11.713974074

print "Truth test 1:", simple.test.seq_list[1]
```

Note: since we didn't run the train\_supervised method again, we are still using the result of the training using smoothing. Therefore, you should compare these results to the ones of posterior decoding with smoothing.

## 2.6 Part-of-Speech Tagging (POS)

Part-of-Speech (PoS) tagging is one of the most important NLP tasks. The task is to assign each word a grammatical category, or Part-of-Speech, *i.e.* noun, verb, adjective,... Recalling the defined notation,  $\Sigma$  is a vocabulary of word types, and  $\Lambda$  is the set of Part-of-Speech tags.

In English, using the Penn Treebank (PTB) corpus (Marcus et al., 1993), the current state of the art for part of speech tagging is around 97% for a variety of methods.

In the rest of this class we will use a subset of the PTB corpus, but instead of using the original 45 tags we will use a reduced tag set of 12 tags, to make the algorithms faster for the class. In this task, x is a sentence (*i.e.*, a sequence of word tokens) and y is the sequence of possible PoS tags.

The first step is to load the corpus. We will start by loading 1000 sentences for training and 1000 sentences both for development and testing. Then we train the HMM model by maximum likelihood estimation.

Look at the transition probabilities of the trained model (see Figure 2.4), and see if they match your intuition about the English language (e.g. adjectives tend to come before nouns).

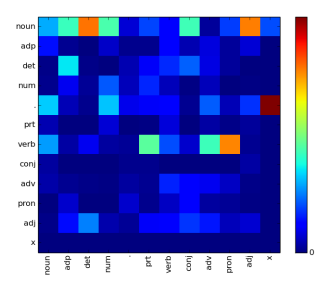


Figure 2.4: Transition probabilities of the trained model. Each column is the previous state and row is the current state. Note the high probability of having Noun after Determinant or Adjective, or of having Verb after Nouns or Pronouns, as expected.

**Exercise 2.9** *Test the model using both posterior decoding and Viterbi decoding on both the train and test set, using the methods in class HMM:* 

```
viterbi_pred_train = hmm.viterbi_decode_corpus(train_seq)
posterior_pred_train = hmm.posterior_decode_corpus(train_seq)
eval_viterbi_train = hmm.evaluate_corpus(train_seq, viterbi_pred_train)
eval_posterior_train = hmm.evaluate_corpus(train_seq, posterior_pred_train)
print "Train Set Accuracy: Posterior Decode %.3f, Viterbi Decode: %.3f"%(
    eval_posterior_train, eval_viterbi_train)
Train Set Accuracy: Posterior Decode 0.985, Viterbi Decode: 0.985

viterbi_pred_test = hmm.viterbi_decode_corpus(test_seq)
posterior_pred_test = hmm.posterior_decode_corpus(test_seq)
eval_viterbi_test = hmm.evaluate_corpus(test_seq, viterbi_pred_test)
eval_posterior_test = hmm.evaluate_corpus(test_seq, posterior_pred_test)
print "Test Set Accuracy: Posterior Decode %.3f, Viterbi Decode: %.3f"%(
    eval_posterior_test, eval_viterbi_test)
Test Set Accuracy: Posterior Decode 0.350, Viterbi Decode: 0.509
```

What do you observe? Remake the previous exercise but now train the HMM using smoothing. Try different values (0,0.1,0.01,1) and report the results on the train and development set. (Use function pick\_best\_smoothing).

```
best_smoothing = hmm.pick_best_smoothing(train_seq, dev_seq, [10,1,0.1,0])
Smoothing 10.000000 -- Train Set Accuracy: Posterior Decode 0.731, Viterbi Decode: 0.691
Smoothing 10.000000 -- Test Set Accuracy: Posterior Decode 0.712, Viterbi Decode: 0.675
Smoothing 1.000000 -- Train Set Accuracy: Posterior Decode 0.887, Viterbi Decode: 0.865
Smoothing 1.000000 -- Test Set Accuracy: Posterior Decode 0.818, Viterbi Decode: 0.792
Smoothing 0.100000 -- Train Set Accuracy: Posterior Decode 0.968, Viterbi Decode: 0.965
Smoothing 0.100000 -- Test Set Accuracy: Posterior Decode 0.851, Viterbi Decode: 0.842
Smoothing 0.000000 -- Train Set Accuracy: Posterior Decode 0.985, Viterbi Decode: 0.985
Smoothing 0.000000 -- Test Set Accuracy: Posterior Decode 0.370, Viterbi Decode: 0.526
hmm.train_supervised(train_seq, smoothing=best_smoothing)
viterbi_pred_test = hmm.viterbi_decode_corpus(test_seq)
posterior_pred_test = hmm.posterior_decode_corpus(test_seq)
eval_viterbi_test = hmm.evaluate_corpus(test_seq, viterbi_pred_test)
eval_posterior_test = hmm.evaluate_corpus(test_seq, posterior_pred_test)
print "Best Smoothing %f -- Test Set Accuracy: Posterior Decode %.3f, Viterbi Decode: %.
    3f"% (best_smoothing, eval_posterior_test, eval_viterbi_test)
Best Smoothing 0.100000 -- Test Set Accuracy: Posterior Decode 0.837, Viterbi Decode: 0.
   827
```

Perform some error analysis to understand were the errors are coming from. You can start by visualizing the confusion matrix (true tags vs predicted tags). You should get something like what is shown in Figure 2.5.

## 2.7 Unsupervised Learning of HMMs

We next address the problem of *unsupervised* learning. In this setting, we are not given any labeled data; all we get to see is a set of natural language sentences. The underlying question is:

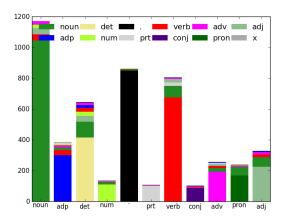


Figure 2.5: Confusion Matrix for the previous example. Predicted tags are columns and the true tags corresponds to the constituents of each column.

Can we learn something from raw text?

This task is challenging, since the process by which linguistic structures are generated is not always clear; and even when it is, it is typically too complex to be formally expressed. Nevertheless, unsupervised learning has been applied to a wide range of NLP tasks, such as: Part-of-Speech Induction (Schütze, 1995; Merialdo, 1994; Clark, 2003), Dependency Grammar Induction (Klein and Manning, 2004; Smith and Eisner, 2006), Constituency Grammar Induction (Klein and Manning, 2004), Statistical Word Alignments (Brown et al., 1993) and Anaphora Resolution (Charniak and Elsner, 2009), just to name a few.

Different motivations have pushed research in this area. From both a linguistic and cognitive point of view, unsupervised learning is useful as a tool to study language acquisition. From a machine learning point of view, unsupervised learning is a fertile ground for testing new learning methods, where significant improvements can yet be made. From a more pragmatic perspective, unsupervised learning is required since annotated corpora is a scarce resource for different reasons. Independently of the reason, unsupervised learning is an increasingly active field of research.

A first problem with unsupervised learning, since we don't observe any labeled data (i.e., the training set is now  $\mathcal{D}_U = \{x^1, \dots, x^M\}$ ), is that most of the methods studied so far (e.g., Perceptron, MIRA, SVMs) cannot be used since we cannot compare the true output with the predicted output. Note also that a direct minimization of the *complete negative log-likelihood* of the data,  $\log P_{\theta}(X^1 = x^1, \dots, X^M = x^m)$ , is very challenging, since it would require marginalizing out (i.e., summing over) all possible hidden variables:

$$\log P_{\theta}(X^{1} = x^{1}, \dots, X^{M} = x^{m}) = \sum_{m=1}^{M} \log \sum_{y \in \Lambda} P_{\theta}(X = x^{m}, Y = y).$$
 (2.49)

Note also that the objective above is *non-convex* even for a linear model: hence, it may have local minima, which makes optimization much more difficult.

The most common optimization method in the presence of hidden (latent) variables is the Expectation Maximization (EM) algorithm, described in the next section. Note that this algorithm is a generic optimization routine that does not depend on a particular model. Later, in Section 2.7.2 we will apply the EM algorithm to the task of part-of-speech induction, where one is given raw text and a number of clusters and the task is to cluster words that behave similarly in a grammatical sense.

#### 2.7.1 Expectation Maximization Algorithm

Given the training corpus  $\mathcal{D}_U := \{x^1, \dots, x^M\}$ , training seeks model parameters  $\theta$  that minimize the negative log-likelihood of the corpus:

Negative Log Likelihood: 
$$\mathcal{L}(\theta) = \widehat{\mathbb{E}}[-\log P_{\theta}(X)] = -\frac{1}{M} \sum_{m=1}^{M} \log \left( \sum_{y^m \in \Lambda} P_{\theta}(X = x^m, Y = y^m) \right),$$
 (2.50)

where  $\widehat{\mathbb{E}}[f(X)] := \frac{1}{M} \sum_{m=1}^{M} f(x^m)$  denotes the empirical average of a function f over the training corpus. Because of the hidden variables  $y^1, \ldots, y^M$ , the likelihood term contains a sum over all possible hidden structures inside of a logarithm, which makes this quantity hard to compute.

The most common minimization algorithm to fit the model parameters in the presence of hidden variables is the Expectation-Maximization (EM) algorithm. The EM procedure can be thought of intuitively in the following way. If we observe the hidden variables' values for all sentences in the corpus, then we could easily compute the maximum likelihood value of the parameters as described in Section 2.3. On the other hand, if we had the model parameters we could label data using the model, and collect the sufficient statistics described in Section 2.3. However, since we are working in an unsupervised setting, we never get to observe the hidden state sequence. Instead, given the training set  $\mathcal{D}_U = \{x^1 \dots x^M\}$ , we will need to collect *expected* sufficient statistics (in this case, *expected* counts) that represent the number of times that each hidden variable is expected to be used with the current parameters setting. These expected sufficient statistics will then be used during learning as "fake" observations of the hidden variables. Using the state and transition posterior distributions described in Equations 2.22 and 2.23, the expected sufficient statistics can be computed by the following formulas:

Initial Counts: 
$$C_{\text{init}}(c_k) = \sum_{m=1}^{M} P_{\theta}(Y_1^m = c_k \mid X^m = x^m);$$
 (2.51)

**Transition Counts**: 
$$C_{\text{trans}}(c_k, c_l) = \sum_{m=1}^{M} \sum_{i=2}^{N} P_{\theta}(Y_i^m = c_k \wedge Y_{i-1}^m = c_l \mid X^m = x^m);$$
 (2.52)

Final Counts: 
$$C_{\text{final}}(c_k) = \sum_{m=1}^{M} P_{\theta}(Y_N^m = c_k \mid X^m = x^m);$$
 (2.53)

Emission Counts: 
$$C_{\text{emiss}}(w_j, c_k) = \sum_{m=1}^{M} \sum_{i=1}^{N} \mathbf{1}(x_i^m = w_j) P_{\theta}(Y_i^m = c_k \mid X^m = x^m);$$
 (2.54)

Compare the previous equations with the ones described in Section 2.3 for the same quantities (Eqs. 2.4–2.7). The main difference is that while in the presence of supervised data you sum the observed events, when you have no label data you sum the posterior probabilities of each event. If these probabilities were such that the probability mass was around single events then both equations will produce the same result.

The EM procedure starts with an initial guess for the parameters  $\theta^{\hat{0}}$  at time t=0. The algorithm keeps iterating until it converges to a local minima of the negative log likelihood. Each iteration is divided into two steps:

- The **E-Step** (Expectation) computes the posteriors for the hidden variables  $P_{\theta^t}(Y|X=x^m)$  given the current parameter values  $\theta^t$  and the observed variables  $X=x^m$  for the m-th sentence. For an HMM, this can be done through the Forward-Backward algorithm described in the previous sections.
- The **M-step** (Maximization) uses those posteriors  $P_{\theta^t}(Y|X=x^m)$  to "softly fill in" the values of the hidden variables  $Y^m$ , and collects the sufficient statistics: initial counts (Eq. 2.51), transition counts (Eq. 2.52), final counts (Eq. 2.53), and emission counts (Eq. 2.54). These counts are then used to estimate maximum likelihood parameters  $\theta^{t+1}$ , as described in Section 2.3.

The EM algorithm is guaranteed to converge to a local minimum of the negative log-likelihood  $\mathcal{L}(\theta)$ , under mild conditions. Note that we are not committing to the best assignment of the hidden variables, but summing the occurrences of each parameter weighed by the posterior probability of all possible assignments. This modular split into two intuitive and straightforward steps accounts for the vast popularity of EM.<sup>7</sup>

Algorithm 9 presents the pseudo code for the EM algorithm.

One important thing to note in Algorithm 9 is that for the HMM model we already have all the model pieces we require. In fact the only method we don't have yet implemented from previous classes is the method to update the counts given the posteriors.

**Exercise 2.10** *Implement the method to update the counts given the state and transition posteriors.* 

def update\_counts(self, sequence, state\_posteriors, transition\_posteriors):

 $<sup>^7</sup>$ More formally, EM minimizes an *upper bound* of  $\mathcal{L}(\theta)$  via block-coordinate descent. See Neal and Hinton (1998) for details. Also, even though we are presenting EM in the context of HMMs, this algorithm can be more broadly applied to any probabilistic model with latent variables.

#### Algorithm 9 EM algorithm.

```
1: input: dataset \mathcal{D}_U, initial model \theta
 2: for t = 1 to T do
 3:
       E-Step:
 4:
       Clear counts: C_{\text{init}}(.) = C_{\text{trans}}(.,.) = C_{\text{final}}(.) = C_{\text{emiss}}(.,.) = 0
       for x^m \in \mathcal{D}_U do
 6:
          Compute posterior expectations P_{\theta}(Y_i|X=x^m) and P_{\theta}(Y_i,Y_{i+1}|X=x^m) using the current model \theta
 7:
          Update counts as shown in Eqs. 2.51–2.54.
 8:
 9:
       end for
10:
       M-Step:
11:
       Update the model parameters \theta based on the counts.
12:
13: end for
```

You now have all the pieces to implement the EM algorithm. Look at the code for EM algorithm in file sequences/hmm.py and check it for yourself.

```
def train_EM(self, dataset, smoothing=0, num_epochs=10, evaluate=True):
   self.initialize_random()
    if evaluate:
        acc = self.evaluate_EM(dataset)
        print "Initial accuracy: %f"%(acc)
    for t in xrange(1, num_epochs):
        #E-Step
        total_log_likelihood = 0.0
        self.clear_counts(smoothing)
        for sequence in dataset.seq_list:
            # Compute scores given the observation sequence.
            initial_scores, transition_scores, final_scores, emission_scores = \
                self.compute_scores(sequence)
            state_posteriors, transition_posteriors, log_likelihood = \
                self.compute_posteriors(initial_scores,
                                        transition_scores,
                                        final_scores,
                                        emission_scores)
            self.update_counts(sequence, state_posteriors, transition_posteriors)
            total_log_likelihood += log_likelihood
        print "Iter: %i Log Likelihood: %f"%(t, total_log_likelihood)
        #M-Step
        self.compute_parameters()
        if evaluate:
             ### Evaluate accuracy at this iteration
            acc = self.evaluate_EM(dataset)
            print "Iter: %i Accuracy: %f"%(t,acc)
```

#### 2.7.2 Part of Speech Induction

In this section we present the Part-of-Speech induction task. Part-of-Speech tags are pre-requisite for many text applications. The task of Part-of-Speech tagging where one is given a labeled training set of words and respective tags is a well studied task with several methods achieving high prediction quality, as we saw in the first part of this chapter.

On the other hand the task of Part-of-Speech induction where one does not have access to a labeled corpus is a much harder task with a huge space for improvement. In this case, we are given only the raw text along with sentence boundaries and a predefined number of clusters we can use. This problem can be seen as a clustering problem. We want to cluster words that behave grammatically in the same way on the same cluster. This is a much harder problem.

Depending on the task at hand we can pick an arbitrary number of clusters. If the goal is to test how well our method can recover the true POS tags then we should use the same number of clusters as POS tags. On the other hand, if the task is to extract features to be used by other methods we can use a much bigger number of clusters (e.g., 200) to capture correlations not captured by POS tags, like lexical affinity.

Note, however that nothing is said about the identity of each cluster. The model has no preference in assigning cluster 1 to nouns vs cluster 2 to nouns. Given this non-identifiability several metrics have been proposed for evaluation (Reichart and Rappoport, 2009; Haghighi and Klein, 2006; Meilă, 2007; Rosenberg and Hirschberg, 2007). In this class we will use a common and simple metric called **1-Many**, which maps each cluster to majority pos tag that it contains (see Figure 2.6 for an example).

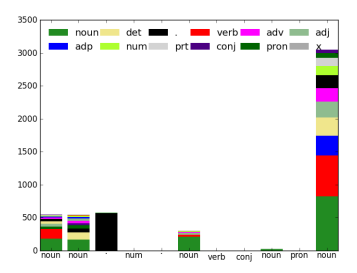


Figure 2.6: Confusion Matrix example. Each cluster is a column. The best tag in each column is represented under the column (1-many) mapping. Each color represents a true Pos Tag.

**Exercise 2.11** Run 20 epochs of the EM algorithm for part of speech induction:

```
hmm.train_EM(train_seq, 0.1, 20, evaluate=True)
viterbi_pred_test = hmm.viterbi_decode_corpus(test_seq)
posterior_pred_test = hmm.posterior_decode_corpus(test_seq)
eval_viterbi_test =
                    hmm.evaluate_corpus(test_seq, viterbi_pred_test)
eval_posterior_test = hmm.evaluate_corpus(test_seq, posterior_pred_test)
Initial accuracy: 0.303638
Iter: 1 Log Likelihood: -101824.763927
Iter: 1 Accuracy: 0.305441
Iter: 2 Log Likelihood: -78057.108346
Iter: 2 Accuracy: 0.321976
Iter: 3 Log Likelihood: -77813.725501
Iter: 3 Accuracy: 0.357451
Iter: 4 Log Likelihood: -77192.947674
Iter: 4 Accuracy: 0.385109
Iter: 5 Log Likelihood: -76191.800849
Iter: 5 Accuracy: 0.392123
Iter: 6 Log Likelihood: -75242.572729
Iter: 6 Accuracy: 0.391121
Iter: 7 Log Likelihood: -74392.892496
Iter: 7 Accuracy: 0.404249
Iter: 8 Log Likelihood: -73357.542833
```

```
Iter: 8 Accuracy: 0.399940
Iter: 9 Log Likelihood: -72135.182778
Iter: 9 Accuracy: 0.399238
Iter: 10 Log Likelihood: -70924.246230
Iter: 10 Accuracy: 0.395430
Iter: 11 Log Likelihood: -69906.561800
Iter: 11 Accuracy: 0.394328
Iter: 12 Log Likelihood: -69140.228623
Iter: 12 Accuracy: 0.390821
Iter: 13 Log Likelihood: -68541.416423
Iter: 13 Accuracy: 0.391522
Iter: 14 Log Likelihood: -68053.456865
Iter: 14 Accuracy: 0.389117
Iter: 15 Log Likelihood: -67667.318961
Iter: 15 Accuracy: 0.386411
Iter: 16 Log Likelihood: -67337.685686
Iter: 16 Accuracy: 0.385409
Iter: 17 Log Likelihood: -67054.571821
Iter: 17 Accuracy: 0.385409
Iter: 18 Log Likelihood: -66769.973881
Iter: 18 Accuracy: 0.385409
Iter: 19 Log Likelihood: -66442.608458
Iter: 19 Accuracy: 0.385409
confusion_matrix = cm.build_confusion_matrix(test_seq.seq_list, viterbi_pred_test,
                                            len(corpus.tag_dict), hmm.get_num_states())
cm.plot_confusion_bar_graph(confusion_matrix, corpus.tag_dict,
                            xrange(hmm.get_num_states()), 'Confusion matrix')
```

Note: your results may not be the same as in this example since we are using a random start, but the trend should be the same, with log-likelihood increasing at every iteration.

In the previous exercise we used an HMM to do Part-of-Speech induction using 12 clusters (by omission the HMM uses as number of hidden states the one provided by the corpus). A first observation is that the log-likelihood is always increasing as expected. Another observation is that the accuracy goes up from 32% to 38%. Note that normally you will run this algorithm for 200 iterations, we stopped earlier for time constraints. Another observation is that the accuracy is not monotonic increasing, this is because the likelihood is not a perfect proxy for the accuracy. In fact all that the likelihood is measuring are co-occurrences of words in the corpus; it has no idea of pos tags. The fact that we are improving derives from the fact that language is not random but follows some specific hidden patterns. In fact this patterns are what true pos-tags try to capture. A final observation is that the performance is really bad compared to the supervised scenario, so there is a lot of space for improvement. The actual state of the art is around 71% for fully unsupervised (Graça, 2010; Berg-Kirkpatrick et al., 2010) and 80% (Das and Petrov, 2011) using parallel data and information from labels in the other language.

Looking at Figure 2.6, it shows the confusion matrix for this particular example. A first observation is that most clusters are mapped to nouns, verbs or punctuation. This is a known fact since there are many more nouns and verbs than any other tags. Since maximum likelihood prefers probabilities to be uniform (imagine two parameters: in one setting both have value 0.5 so the likelihood will be 0.5\*0.5 = 0.25, while in the other case one as 0.1 and 0.9 so the maximum likelihood is 0.09). Several approaches have been proposed to address this problem, moving towards a Bayesian setting (Johnson, 2007), or using Posterior Regularization (Graça et al., 2009).

## Day 3

# **Learning Structured Predictors**

In this class, we will continue to focus on sequence classification, but instead of following a *generative* approach (like in the previous chapter) we move towards *discriminative* approaches. Recall that the difference between these approaches is that generative approaches attempt to model the probability distribution of the data, P(X,Y), whereas discriminative ones only model the conditional probability of the sequence, given the observed data, P(Y|X).

### Today's Assignment

The assignment of today's class is to implement the structured version of the perceptron algorithm.

## 3.1 Classification vs Sequential Classification

Table 3.1 shows how the models for classification have counterparts in *sequential* classification. In fact, in the last chapter we discussed the Hidden Markov model, which can be seen as a generalization of the Naïve Bayes model for sequences. In this chapter, we will see a generalization of the Perceptron algorithm for sequence problems (yielding the Structured Perceptron, Collins 2002) and a generalization of Maximum Entropy model for sequences (yielding Conditional Random Fields, Lafferty et al. 2001). Note that both these generalizations are not specific for sequences and can be applied to a wide range of models (we will see in tomorrow's lecture how these methods can be applied to parsing). Although we will not cover all the methods described in Chapter 1, bear in mind that all of those have a structured counterpart. It should be intuitive after this lecture how those methods could be extended to structured problems, given the perceptron example.

Classification	Sequences	
Generative		
Naïve Bayes (Sec. 1.2)	Hidden Markov Models (Sec. 2.2)	
Discriminative		
Perceptron (Sec. 1.4.3)	Structured Perceptron (Sec. 3.5)	
Maximum Entropy (Sec. 1.4.4)	Conditional Random Fields (Sec. 3.4)	

Table 3.1: Summary of the methods used for classification and sequential classification covered in this guide.

Throughout this chapter, we will be searching for the solution of

$$\underset{y \in \Lambda^{N}}{\arg \max} P(Y = y | X = x) = \underset{y \in \Lambda^{N}}{\arg \max} w \cdot f(x, y). \tag{3.1}$$

where w is a weight vector, and f(x,y) is a feature vector. We will see that this sort of linear models are more flexible than HMMs, in the sense that they may incorporate more general features while being able to reuse the same decoders (based on the Viterbi and forward-backward algorithms). In fact, the exercises in this lab will not touch the decoders that have been developed in the previous lab. Only the training algorithms and the function that compute the scores will change.

As in the previous section,  $y = y_1 \dots y_N$  is a sequence so the maximization is over an exponential number of objects, making it intractable for brute force methods. Again we will make an assumption analogous to the

"first order Markov independence property," so that the features may decompose as a sum over nodes and edges in a trellis. This is done by assuming that expression 3.1 can be written as:

$$\underset{y \in \Lambda^{N}}{\operatorname{arg max}} \sum_{i=1}^{N} \underbrace{w \cdot f_{\text{emiss}}(i, x, y_{i})}_{\text{score}_{\text{emiss}}} + \underbrace{w \cdot f_{\text{init}}(x, y_{1})}_{\text{score}_{\text{init}}} + \sum_{i=1}^{N-1} \underbrace{w \cdot f_{\text{trans}}(i, x, y_{i}, y_{i+1})}_{\text{score}_{\text{trans}}} + \underbrace{w \cdot f_{\text{final}}(x, y_{N})}_{\text{score}_{\text{final}}}$$
(3.2)

In other words, the scores score<sub>emiss</sub>, score<sub>init</sub>, score<sub>trans</sub> and score<sub>final</sub> are now computed as inner products between weight vectors and feature vectors rather than log-probabilities. The feature vectors depend locally on the output variable (*i.e.*, they only look at a single  $y_i$  or a pair  $y_i$ ,  $y_{i+1}$ ); however they may depend globally on the observed input  $x = x_1, ..., x_N$ .

#### 3.2 Features

In this section we will define two simple feature sets.<sup>1</sup> The first feature set will only use identity features, and will mimic the features used by the HMM model from the previous section. This will allow us to directly compare the performance of a generative vs a discriminative approach. Table 3.2 depicts the features that are implicit in the HMM, which was the subject of the previous chapter. These features are indicators of initial, transition, final, and emission events.

Condition	Name
$y_i = c_k \& i = 0$	Initial Features
$y_i = c_k \& y_{i-1} = c_l$	Transition Features
$y_i = c_k \& i = N$	Final Features
$x_i = w_j \& y_i = c_k$	Emission Features

Table 3.2: IDFeatures feature set. This set replicates the features used by the HMM model.

Note that the fact that we were using a generative model has forced us (in some sense) to make strong independence assumptions. However, since we now move to a discriminative approach, where we model P(Y|X) rather than P(X,Y), we are not tied anymore to some of these assumptions. In particular:

- We may use "overlapping" features, *e.g.*, features that fire simultaneously for many instances. For example, we can use a feature for a word, such as a feature which fires for the word "brilliantly", and another for prefixes and suffixes of that word, such as one which fires if the last two letters of the word are "ly". This would lead to an awkward model if we wanted to insist on a generative approach.
- We may use features that depend arbitrarily on the *entire input sequence x*. On the other hand, we still need to resort to "local" features with respect to the *outputs* (*e.g.* looking only at consecutive state pairs), otherwise decoding algorithms will become more expensive.

This leads us to the second feature set, composed of features that are traditionally used in POS tagging with discriminative models. See Table 3.3 for some examples. Of course, we could have much more complex features, looking arbitrarily to the input sequence. We are not going to have them in this exercise only for performance reasons (to have less features and smaller caches). State-of-the-art sequence classifiers can easily reach over one million features!

Our features subdivide into two groups:  $f_{\rm emiss}$ ,  $f_{\rm init}$ , and  $f_{\rm final}$  are all instances of *node features*, depending only on a single position in the state sequence (a node in the trellis);  $f_{\rm trans}$  are *edge features*, depending on two consecutive positions in the state sequence (an edge in the trellis). Similarly as in the previous chapter, we have the following scores (also called *log-potentials* in the literature on CRFs and graphical models):

• Initial scores. These are scores for the initial state. They are given by

$$score_{init}(x, y_1) = w \cdot f_{init}(x, y_1). \tag{3.3}$$

• Transition scores. These are scores for two consecutive states at a particular position. They are given by

$$score_{trans}(i, x, y_i, y_{i+1}) = w \cdot f_{trans}(i, x, y_i, y_{i+1}). \tag{3.4}$$

<sup>&</sup>lt;sup>1</sup>Although not required, all the features we will use are binary features, indicating whether a given condition is satisfied or not.

Condition	Name
$y_i = c_k \& i = 0$	Initial Features
$y_i = c_k \& y_{i-1} = c_l$	Transition Features
$y_i = c_k \& i = N$	Final Features
$x_i = w_i \& y_i = c_k$	Basic Emission Features
$x_i = w_i \& w_i$ is uppercased & $y_i = c_k$	Uppercase Features
$x_i = w_i \& w_i$ contains digit & $y_i = c_k$	Digit Features
$x_i = w_i \& w_i$ contains hyphen & $y_i = c_k$	Hyphen Features
$x_i = w_i \& w_i[0i] \forall i \in [1, 2, 3] \& y_i = c_k$	Prefix Features
$ x_i = w_i \& w_i[ w_i  - i w_i ] \forall i \in [1, 2, 3] \& y_i = c_k$	Suffix Features

Table 3.3: Extended feature set. Some features in this set could not be included in the HMM model. The features included in the bottom row are all considered emission features for the purpose of our implementation, since they all depend on i, x and  $y_i$ .

• Final scores. These are scores for the final state. They are given by

$$score_{final}(x, y_N) = w \cdot f_{final}(x, y_N). \tag{3.5}$$

• Emission scores. These are scores for a state at a particular position. They are given by

$$score_{emiss}(i, x, y_i) = w \cdot f_{emiss}(i, x, y_i). \tag{3.6}$$

### 3.3 Discriminative Sequential Classifiers

Given a weight vector w, the conditional probability  $P_w(Y = y | X = x)$  is then defined as follows:

$$P_{w}(Y = y | X = x) = \frac{1}{Z(w, x)} \exp\left(w \cdot f_{\text{init}}(x, y_{1}) + \sum_{i=1}^{N-1} w \cdot f_{\text{trans}}(i, x, y_{i}, y_{i+1}) + w \cdot f_{\text{final}}(x, y_{N}) + \sum_{i=1}^{N} w \cdot f_{\text{emiss}}(i, x, y_{i})\right) (3.7)$$

where the normalizing factor Z(w, x) is called the *partition function*:

$$Z(w, x) = \sum_{y \in \Lambda^{N}} \exp \left( w \cdot f_{\text{init}}(x, y_{1}) + \sum_{i=1}^{N-1} w \cdot f_{\text{trans}}(i, x, y_{i}, y_{i+1}) + w \cdot f_{\text{final}}(x, y_{N}) + \sum_{i=1}^{N} w \cdot f_{\text{emiss}}(i, x, y_{i}) \right) (3.8)$$

#### 3.3.1 Training

For training, the important problem is that of obtaining the weight vector w that lead to an accurate classifier. We will discuss two possible strategies:

1. Maximizing conditional log-likelihood from a set of labeled data  $\{(x^m, y^m)\}_{m=1}^M$ , yielding **conditional random fields**. This corresponds to the following optimization problem:

$$\hat{w} = \arg\max_{w} \sum_{m=1}^{M} \log P_{w}(Y = y^{m} | X = x^{m}).$$
 (3.9)

To avoid overfitting, it is common to regularize with the Euclidean norm function, which is equivalent to considering a zero-mean Gaussian prior on the weight vector. The problem becomes:

$$\widehat{\boldsymbol{w}} = \arg\max_{\boldsymbol{w}} \sum_{m=1}^{M} \log P_{\boldsymbol{w}}(Y = y^{m} | X = x^{m}) - \frac{\lambda}{2} \| \boldsymbol{w} \|^{2}.$$
 (3.10)

This is precisely the structured variant of the maximum entropy method discussed in Chapter 1. Unlike HMMs, this problem does not have a closed form solution and has to be solved with numerical optimization.

2. Alternatively, running the **structured perceptron** algorithm to obtain a weight vector *w* that accurately classifies the training data. We will see that this simple strategy achieves results which are competitive with conditional log-likelihood maximization.

#### 3.3.2 Decoding

For decoding, there are three important problems that need to be solved:

- 1. Given X = x, compute the most likely output sequence  $\hat{y}$  (the one which maximizes  $P_w(Y = y | X = x)$ ).
- 2. Compute the posterior marginals  $P_w(Y_i = y_i | X = x)$  at each position *i*.
- 3. Evaluate the partition function Z(w, x).

Interestingly, all these problems can be solved by using the very same algorithms that were already implemented for HMMs: the Viterbi algorithm (for 1) and the forward-backward algorithm (for 2–3). All that changes is the way the scores are computed.

#### 3.4 Conditional Random Fields

Conditional Random Fields (CRF) (Lafferty et al., 2001) can be seen as an extension of the Maximum Entropy (ME) models to structured problems.<sup>2</sup> They are *globally* normalized models: the probability of a given sentence is given by Equation 3.7. There are only two differences with respect to the standard ME model described a couple of days ago for multi-class classification:

- Instead of computing the posterior marginals P(Y = y|X = x) for all possible configurations of the output variables (which are exponentially many), it assumes the model decompose into "parts" (in this case, nodes and edges), and it computes only the posterior marginals for those parts,  $P(Y_i = y_i|X = x)$  and  $P(Y_i = y_i, Y_{i+1} = y_{i+1}|X = x)$ . Crucially, we can compute these quantities by using the very same forward-backward algorithm that we have described for HMMs.
- Instead of updating the features for all possible outputs  $y' \in \Lambda^N$ , we again exploit the decomposition into parts above and update only "local features" at the nodes and edges.

Algorithm 10 shows the pseudo code to optimize a CRF with the stochastic gradient descent (SGD) algorithm (our toolkit also includes an implementation of a quasi-Newton method, L-BFGS, which converges faster, but for the purpose of this exercise, we will stick with SGD).

**Exercise 3.1** In this exercise you will train a CRF using different feature sets for Part-of-Speech Tagging. Start with the code below, which uses the ID feature set from table 3.2.

<sup>&</sup>lt;sup>2</sup>An earlier, less successful, attempt to perform such an extension was via Maximum Entropy Markov models (MEMM) (McCallum et al., 2000). There each factor (a node or edge) is a *locally* normalized maximum entropy model. A shortcoming of MEMMs is its so-called *labeling bias* (Bottou, 1991), which makes them biased towards states with few successor states (see Lafferty et al. (2001) for more information).

#### Algorithm 10 SGD for Conditional Random Fields

- 1: **input:**  $\mathfrak{D}$ ,  $\lambda$ , number of rounds T, learning rate sequence  $(\eta_t)_{t=1,\dots,T}$
- 2: initialize  $w^1 = 0$
- 3: **for** t = 1 **to** T **do**
- 4: choose m = m(t) randomly
- 5: take training pair  $(x^m, y^m)$  and compute the probability  $P_w(Y = y^m | X = x^m)$  using the current model w and Eq. 3.7.
- 6: for every i,  $y'_{i'}$  and  $y'_{i+1}$ , compute marginal probabilities  $P(Y_i = y'_i | X = x)$  and  $P(Y_i = y'_i, Y_{i+1} = y'_{i+1} | X = x^m)$  using the current model w, for each node and edge, through the forward-backward algorithm.
- 7: compute the feature vector expectation:

$$E_{w^{t}}[f(x^{m},Y)] = \sum_{y_{1}} P(y_{1}|x^{m}) f_{\text{init}}(x^{m},y_{1}) + \sum_{i=1}^{N-1} \sum_{y_{i},y_{i+1}} P(y_{i},y_{i+1}|x^{m}) f_{\text{trans}}(i,x^{m},y_{i},y_{i+1}) + \sum_{y_{N}} P(y_{N}|x^{m}) f_{\text{final}}(x^{m},y_{N}) + \sum_{i=1}^{N} \sum_{y_{i}} P(y_{i}|x^{m}) f_{\text{emiss}}(i,x^{m},y_{i}).$$
(3.11)

8: update the model:

$$w^{t+1} \leftarrow (1 - \lambda \eta_t) w^t + \eta_t (f(x^m, y^m) - E_{w^t}[f(x^m, Y)])$$

- 9: end for
- 10: **output:**  $\widehat{w} \leftarrow w^{T+1}$

```
crf_online = crfo.CRFOnline(corpus.word_dict, corpus.tag_dict, feature_mapper)
crf_online.num_epochs = 20
crf_online.train_supervised(train_seq)
Epoch: 0 Objective value: -5.779018
Epoch: 1 Objective value: -3.192724
Epoch: 2 Objective value: -2.717537
Epoch: 3 Objective value: -2.436614
Epoch: 4 Objective value: -2.240491
Epoch: 5 Objective value: -2.091833
Epoch: 6 Objective value: -1.973353
Epoch: 7 Objective value: -1.875643
Epoch: 8 Objective value: -1.793034
Epoch: 9 Objective value: -1.721857
Epoch: 10 Objective value: -1.659605
Epoch: 11 Objective value: -1.604499
Epoch: 12 Objective value: -1.555229
Epoch: 13 Objective value: -1.510806
Epoch: 14 Objective value: -1.470468
Epoch: 15 Objective value: -1.433612
Epoch: 16 Objective value: -1.399759
Epoch: 17 Objective value: -1.368518
Epoch: 18 Objective value: -1.339566
Epoch: 19 Objective value: -1.312636
```

You will receive feedback when each epoch is finished, note that running the 20 epochs might take a while. After training is done, evaluate the learned model on the training, development and test sets.

```
pred_train = crf_online.viterbi_decode_corpus(train_seq)
pred_dev = crf_online.viterbi_decode_corpus(dev_seq)
pred_test = crf_online.viterbi_decode_corpus(test_seq)
```

```
eval_train = crf_online.evaluate_corpus(train_seq, pred_train)
eval_dev = crf_online.evaluate_corpus(dev_seq, pred_dev)
eval_test = crf_online.evaluate_corpus(test_seq, pred_test)

print "CRF - ID Features Accuracy Train: %.3f Dev: %.3f Test: %.3f"%(eval_train,eval_dev,
eval_test)
```

You should get values similar to these:

```
Out[]: CRF - ID Features Accuracy Train: 0.949 Dev: 0.846 Test: 0.858
```

Compare with the results achieved with the HMM model (0.837 on the test set, from the previous lecture). Even using a similar feature set a CRF yields better results than the HMM from the previous lecture. Perform some error analysis and figure out what are the main errors the tagger is making. Compare them with the errors made by the HMM model. (Hint: use the methods developed in the previous lecture to help you with the error analysis).

**Exercise 3.2** Repeat the previous exercise using the extended feature set. Compare the results.

```
feature_mapper = exfc.ExtendedFeatures(train_seq)
feature_mapper.build_features()
crf_online = crfo.CRFOnline(corpus.word_dict, corpus.tag_dict, feature_mapper)
crf\_online.num\_epochs = 20
crf_online.train_supervised(train_seq)
Epoch: 0 Objective value: -7.141596
Epoch: 1 Objective value: -1.807511
Epoch: 2 Objective value: -1.218877
Epoch: 3 Objective value: -0.955739
Epoch: 4 Objective value: -0.807821
Epoch: 5 Objective value: -0.712858
Epoch: 6 Objective value: -0.647382
Epoch: 7 Objective value: -0.599442
Epoch: 8 Objective value: -0.562584
Epoch: 9 Objective value: -0.533411
Epoch: 10 Objective value: -0.509885
Epoch: 11 Objective value: -0.490548
Epoch: 12 Objective value: -0.474318
Epoch: 13 Objective value: -0.460438
Epoch: 14 Objective value: -0.448389
Epoch: 15 Objective value: -0.437800
Epoch: 16 Objective value: -0.428402
Epoch: 17 Objective value: -0.419990
Epoch: 18 Objective value: -0.412406
Epoch: 19 Objective value: -0.405524
pred_train = crf_online.viterbi_decode_corpus(train_seq)
pred_dev = crf_online.viterbi_decode_corpus(dev_seq)
pred_test = crf_online.viterbi_decode_corpus(test_seq)
eval_train = crf_online.evaluate_corpus(train_seq, pred_train)
eval_dev = crf_online.evaluate_corpus(dev_seq, pred_dev)
eval_test = crf_online.evaluate_corpus(test_seq, pred_test)
print "CRF - Extended Features Accuracy Train: %.3f Dev: %.3f Test: %.3f"%(eval_train,
   eval_dev, eval_test)
```

You should get values close to the following:

Compare the errors obtained with the two different feature sets. Do some error analysis: what errors were correct by using more features? Can you think of other features to use to solve the errors found?

The main lesson to learn from this exercise is that, usually, if you are not satisfied by the accuracy of your algorithm, you can perform some error analysis and find out which errors your algorithm is making. You can then add more features which attempt to improve those specific errors (this is known as *feature engineering*). This can lead to two problems:

- More features will make training and decoding more expensive. For example, if you add features that depend on the current word and the previous word, the number of new features is the square of the number of different words, which is quite large. For example, the Penn Treebank has around 40000 different words, so you are adding a lot of new features, even though not all pairs of words will ever occur. Features that depend on three words (previous, current, and next) are even more numerous.
- If features are very specific, such as the (previous word, current word, next word) one just mentioned, they might occur very rarely in the training set, which leads to overfit problems. Some of these problems (not all) can be mitigated with techniques such as smoothing, which you already learned about.

## 3.5 Structured Perceptron

The structured perceptron (Collins, 2002), namely its averaged version, is a very simple algorithm that relies on Viterbi decoding and very simple additive updates. In practice this algorithm is very easy to implement and behaves remarkably well in a variety of problems. These two characteristics make the structured perceptron algorithm a natural first choice to try and test a new problem or a new feature set.

Recall what you learned about the perceptron algorithm (Section 1.4.3) and compare it against the structured perceptron (Algorithm 11).

#### Algorithm 11 Averaged Structured perceptron

- 1: **input:**  $\mathfrak{D}$ , number of rounds T
- 2: initialize  $w^1 = 0$
- 3: **for** t = 1 **to** T **do**
- 4: choose m = m(t) randomly
- 5: take training pair  $(x^m, y^m)$  and predict using the current model w, through the Viterbi algorithm:

$$\widehat{y} \leftarrow \arg\max_{y' \in \Lambda^N} w^t \cdot f(x^m, y')$$

- 6: update the model:  $w^{t+1} \leftarrow w^t + f(x^m, y^m) f(x^m, \widehat{y})$
- 7: end for
- 8: **output:** the averaged model  $\hat{w} \leftarrow \frac{1}{T} \sum_{t=1}^{T} w^{t}$

There are only two differences, which mimic the ones already seen for the comparison between CRFs and multi-class ME models:

- Instead of explicitly enumerating all possible output configurations (exponentially many of them) to compute  $\hat{y} := \arg \max_{y' \in \mathcal{Y}} w \cdot f(x^m, y')$ , it finds the best sequence through the Viterbi algorithm.
- Instead of updating the features for the entire  $\hat{y}$ , it updates only the node and edge features at the positions where the labels are different—i.e., where mistakes are made.

## 3.6 Assignment

**Exercise 3.3** *Implement the structured perceptron algorithm.* 

To do this, edit file sequences/structured\_perceptron.py and implement the function

```
def perceptron_update(self, sequence):
    pass
```

This function should apply one round of the perceptron algorithm, updating the weights for a given sequence, and returning the number of predicted labels (which equals the sequence length) and the number of mistaken labels.

Hint: adapt the function

```
def gradient_update(self, sequence, eta):
```

defined in file sequences/crf\_online.py. You will need to replace the computation of posterior marginals by the Viterbi algorithm, and to change the parameter updates according to Algorithm 11. Note the role of the functions

```
self.feature_mapper.get_*_features()
```

in providing the indices for the features obtained for  $f(x^m, y^m)$  or  $f(x^m, \widehat{y})$ 

**Exercise 3.4** Repeat Exercises 3.1–3.2 using the structured perceptron algorithm instead of a CRF. Report the results. Here is the code for the simple feature set:

```
feature_mapper = idfc.IDFeatures(train_seq)
feature_mapper.build_features()
print "Perceptron Exercise"
sp = spc.StructuredPerceptron(corpus.word_dict, corpus.tag_dict, feature_mapper)
sp.num\_epochs = 20
sp.train_supervised(train_seq)
Epoch: 0 Accuracy: 0.656806
Epoch: 1 Accuracy: 0.820898
Epoch: 2 Accuracy: 0.879176
Epoch: 3 Accuracy: 0.907432
Epoch: 4 Accuracy: 0.925239
Epoch: 5 Accuracy: 0.939956
Epoch: 6 Accuracy: 0.946284
Epoch: 7 Accuracy: 0.953790
Epoch: 8 Accuracy: 0.958499
Epoch: 9 Accuracy: 0.955114
Epoch: 10 Accuracy: 0.959235
Epoch: 11 Accuracy: 0.968065
Epoch: 12 Accuracy: 0.968212
Epoch: 13 Accuracy: 0.966740
Epoch: 14 Accuracy: 0.971302
Epoch: 15 Accuracy: 0.968653
Epoch: 16 Accuracy: 0.970419
Epoch: 17 Accuracy: 0.971891
Epoch: 18 Accuracy: 0.971744
Epoch: 19 Accuracy: 0.973510
pred_train = sp.viterbi_decode_corpus(train_seq)
pred_dev = sp.viterbi_decode_corpus(dev_seq)
pred_test = sp.viterbi_decode_corpus(test_seq)
eval_train = sp.evaluate_corpus(train_seq, pred_train)
eval_dev = sp.evaluate_corpus(dev_seq, pred_dev)
eval_test = sp.evaluate_corpus(test_seq, pred_test)
print "Structured Perceptron - ID Features Accuracy Train: %.3f Dev: %.3f Test: %.3f"%(
   eval_train, eval_dev, eval_test)
```

Structured Perceptron - ID Features Accuracy Train: 0.984 Dev: 0.835 Test: 0.840

#### Here is the code for the extended feature set:

```
feature_mapper = exfc.ExtendedFeatures(train_seq)
feature_mapper.build_features()
sp = spc.StructuredPerceptron(corpus.word_dict, corpus.tag_dict, feature_mapper)
sp.num\_epochs = 20
sp.train_supervised(train_seq)
Epoch: 0 Accuracy: 0.764386
Epoch: 1 Accuracy: 0.872701
Epoch: 2 Accuracy: 0.903458
Epoch: 3 Accuracy: 0.927594
Epoch: 4 Accuracy: 0.938484
Epoch: 5 Accuracy: 0.951141
Epoch: 6 Accuracy: 0.949816
Epoch: 7 Accuracy: 0.959529
Epoch: 8 Accuracy: 0.957616
Epoch: 9 Accuracy: 0.962325
Epoch: 10 Accuracy: 0.961148
Epoch: 11 Accuracy: 0.970567
Epoch: 12 Accuracy: 0.968212
Epoch: 13 Accuracy: 0.973216
Epoch: 14 Accuracy: 0.974393
Epoch: 15 Accuracy: 0.973951
Epoch: 16 Accuracy: 0.976600
Epoch: 17 Accuracy: 0.977483
Epoch: 18 Accuracy: 0.974834
Epoch: 19 Accuracy: 0.977042
pred_train = sp.viterbi_decode_corpus(train_seq)
pred_dev = sp.viterbi_decode_corpus(dev_seq)
pred_test = sp.viterbi_decode_corpus(test_seq)
eval_train = sp.evaluate_corpus(train_seq, pred_train)
eval_dev = sp.evaluate_corpus(dev_seq, pred_dev)
eval_test = sp.evaluate_corpus(test_seq, pred_test)
print "Structured Perceptron - Extended Features Accuracy Train: %.3f Dev: %.3f Test: %.
   3f"%(eval_train, eval_dev, eval_test)
```

#### And here are the expected results:

Structured Perceptron - Extended Features Accuracy Train: 0.984 Dev: 0.888 Test: 0.890

## Day 4

# Syntax and Parsing

In this lab we will implement some exercises related with parsing.

## 4.1 Phrase-based Parsing

## 4.1.1 Context Free Grammars

Let T be an *alphabet* (*i.e.*, a finite set of symbols), and denote by  $T^*$  its Kleene closure, *i.e.*, the infinite set of strings produced with those symbols:

$$\mathfrak{I}^* = \emptyset \cup \mathfrak{I} \cup \mathfrak{I}^2 \cup \dots$$

A *language* L is a subset of  $\mathfrak{T}^*$ . The "complexity" of a language L can be loosely defined by how hard it is to construct a machine (an *automaton*) capable of distinguishing the words in L from the elements of  $\mathfrak{T}^*$  which are not in L. If L is finite, a very simple automaton can be built which just memorizes the strings in L. The next simplest case is that of *regular languages*, which are recognizable by *finite state machines*. These are the languages that can be expressed by regular expressions. An example (where  $\mathfrak{T} = \{a,b\}$ ) is the language  $L = \{ab^{n_1}aa^{n_2} \mid n_1, n_2 \in \mathbb{N}\}$ , which corresponds to the regular expression ab\*a+. *Hidden Markov models* (studied in previous lectures) can be seen as a stochastic version of finite state machines.

A step higher in the hierarchy of languages leads to *context-free languages*, which are more complex than regular languages. These are languages that are generated by *context-free grammars*, and recognizable by *push-down automata* (which are slightly more complex than finite state machines). In this section we describe context-free grammars and how they can be made probabilistic. This will yield models that are more powerful than hidden Markov models, and are specially amenable for modeling the syntax of natural languages.<sup>2</sup>

A *context-free grammar* (CFG) is a tuple  $G = \langle \mathcal{N}, \mathcal{T}, \mathcal{R}, \mathcal{S} \rangle$  where:

- 1.  $\mathbb{N}$  is a finite set of *non-terminal* symbols. Elements of  $\mathbb{N}$  are denoted by upper case letters (A, B, C, . . .). Each non-terminal symbol is a syntactic category: it represents a different type of phrase or clause in the sentence.
- 2.  $\mathcal{T}$  is a finite set of *terminal* symbols (disjoint from  $\mathcal{N}$ ). Elements of  $\mathcal{T}$  are denoted by lower case letters  $(a,b,c,\ldots)$ . Each terminal symbol is a surface word: terminal symbols make up the actual content of sentences. The set  $\mathcal{T}$  is called the *alphabet* of the language defined by the grammar G.
- 3.  $\mathcal{R}$  is a set of *production rules*, *i.e.*, a finite relation from  $\mathcal{N}$  to  $(\mathcal{N} \cup \mathcal{T})^*$ . G is said to be in Chomsky normal form (CNF) if production rules in  $\mathcal{R}$  are either of the form  $A \to BC$  or  $A \to a$ .
- 4. S is a *start symbol*, used to represent the whole sentence. It must be an element of  $\mathbb{N}$ .

Any CFG can be transformed to be in CNF without loosing any expressive power in terms of the language it generates. Hence, we henceforth assume that *G* is in CNF without loss of generality.

To see how CFGs can model the syntax of natural languages, consider the following sentence,

<sup>&</sup>lt;sup>1</sup>We recommend the classic book by Hopcroft et al. (1979) for a thorough introduction on the subject of automata theory and formal languages.

<sup>&</sup>lt;sup>2</sup>This does not mean that natural languages are context free. There is an immense body of work on grammar formalisms that relax the "context-free" assumption, and those formalisms have been endowed with a probabilistic framework as well. Examples are: lexical functional grammars, head-driven phrase structured grammars, combinatorial categorial grammars, tree adjoining grammars, etc. Some of these formalisms are *mildly context sensitive*, a relaxation of the "context-free" assumption which still allows polynomial parsing algorithms. There is also equivalence in expressive power among several of these formalisms.

She enjoys the Summer school.

along with a grammar (in CNF) with the following production rules:

S --> NP VP

NP --> Det N

NP --> She

VP --> V NP

V --> enjoys

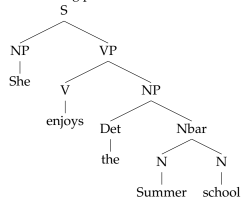
Det --> the

Nbar --> N N

N --> Summer

N --> school

With this grammar, we may derive the following parse tree:



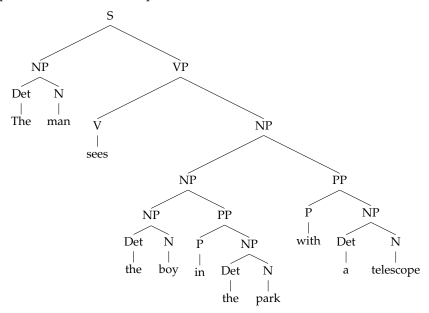
## 4.1.2 Ambiguity

A fundamental characteristic of natural languages is ambiguity. For example, consider the sentence

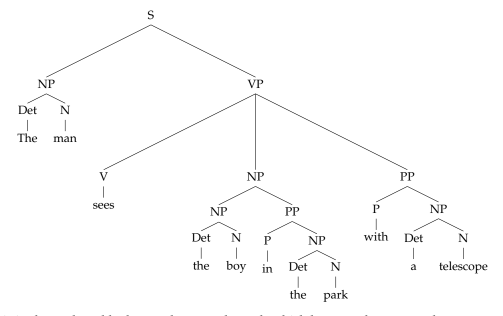
The man sees the boy in the park with a telescope.

for which all the following parse trees are plausible interpretations:

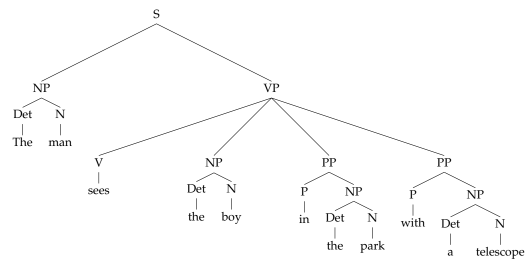
1. The boy is in a park and he has a telescope:



2. The boy is in a park, and the man sees him using a telescope as an instrument:



3. The man is in the park and he has a telescope, through which he sees a boy somewhere:



The ambiguity is caused by the several places to each the prepositional phrase could be attached. This kind of syntactical ambiguity (*PP-attachment*) is one of the most frequent in natural language.

## 4.1.3 Probabilistic Context-Free Grammars

A probabilistic context-free grammar is a tuple  $G_{\theta} = \langle \mathbb{N}, \mathbb{T}, \mathbb{R}, \mathbb{S}, \theta \rangle$ , where  $\langle \mathbb{N}, \mathbb{T}, \mathbb{R}, \mathbb{S} \rangle$  is a CFG and  $\theta$  is a vector of parameters, one per each production rule in  $\mathbb{R}$ . Assuming that the grammar is in CNF, each rule of the kind  $Z \to XY$  is endowed a conditional probability

$$\theta_{Z\to XY} = P_{\theta}(XY|Z),$$

and each unary rule of the kind  $Z \rightarrow w$  is endowed with a conditional probability

$$\theta_{Z\to w} = P_{\theta}(w|Z).$$

For these conditional probabilities to be well defined, the entries of  $\theta$  must be non-negative and need to normalize properly for each  $Z \in \mathbb{N}$ :

$$\sum_{X,Y \in \mathcal{N}} \theta_{Z \to XY} + \sum_{w \in \mathcal{T}} \theta_{Z \to w} = 1.$$

Let s be a string and t a parse tree derivation for s. For each  $r \in \mathcal{R}$ , let  $n_r(t,s)$  be the number of times production rule r appears in the derivation. According to this generative model, the joint probability of t and s factors as

the product of the conditional probabilities above:

$$P(t,s) = \prod_{r \in \mathcal{R}} \theta_r^{n_r(t,s)}.$$

For example, for the sentence above (She enjoys the Summer school) this probability would be

$$\begin{split} P(t,s) &= P(\text{NP VP}|S) \times P(\text{She}|\text{NP}) \times P(\text{V NP}|\text{VP}) \times P(\text{enjoys}|\text{V}) \\ &\times P(\text{Det Nbar}|\text{NP}) \times P(\text{the}|\text{Det}) \times P(\text{N N}|\text{Nbar}) \\ &\times P(\text{Summer}|\text{N}) \times P(\text{school}|\text{N}). \end{split} \tag{4.1}$$

When a sentence is ambiguous, the most likely parse tree can be obtained by maximizing the conditional probability P(t|s); this quantity is proportional to P(t,s) and therefore the latter quantity can be maximized. The number of possible parse trees, however, grows exponentially with the sentence length, rendering a direct maximization intractable. Fortunately, a generalization of the Viterbi algorithm exists which uses dynamic programming to carry out this computation. This is the subject of the next section.

## 4.1.4 The CKY Parsing Algorithm

#### Algorithm 12 CKY algorithm

```
1: input: probabilistic CFG G_{\theta} in CNF, and sentence s = w_1 \dots w_N (each w_i is a word)
   {We'll fill the CKY table bottom-up with partial probabilities \delta and backtrack pointers \psi. This is similar to
    the Viterbi algorithm but it works for parse trees rather than sequences.}
 5: {Initialization}
 6: for i = 1 to N do
       for each production rule r \in \mathbb{R} of the form Z \to w_i do
 7:
 8:
         \delta(i,i,Z) = \theta_{Z \to w_i}
       end for
 9:
10: end for
11:
    {Induction}
12:
    for i = 2 to N do {i is length of span}
       for j = 1 to N - i + 1 do \{j \text{ is start of span}\}\
14:
         for each non-terminal Z \in \mathcal{N} do
15:
            Set partial probability:
16:
                                \delta(j,j+i-1,Z) = \max_{\substack{X,Y\\j < k < j+i}} \delta(j,k-1,X) \times \delta(k,j+i-2,Y) \times \theta_{Z \to XY}
```

17: Store backpointer:

$$\psi(j,j+i-1,Z) = \operatorname*{arg\,max}_{\substack{X,Y\\j < k < j+i}} \delta(j,k-1,X) \times \delta(k,j+i-2,Y) \times \theta_{Z \to XY}$$

```
18: end for
19: end for
20: end for
21:
22: {Termination}
23: P(s,\hat{t}) = \delta(1,N,\mathbb{S})
24: Backtrack through \psi to obtain most likely parse tree \hat{t}
```

One of the most widely known algorithm for parsing natural language sentences is the Cocke-Kasami-Younger (CKY) algorithm. Given a grammar in CNF with  $|\mathcal{R}|$  production rules, its runtime complexity for parsing a sentence of length N is  $O(N^3|\mathcal{R}|)$ . We present here a simple extension of the CKY algorithm that obtains the most likely parse tree of a sentence, along with its probability. Given a sequence of words  $w_i$  in a

sentence  $s = w_1, ..., w_N$ , we want to learn probability for each possible non-terminal rule  $Z \in \mathcal{R}$ , and take the most probable parsing tree that generates the sentence. <sup>3</sup> This is displayed in Alg. 12.

This algorithm assigns a probability  $\delta(i,j,Z)$ , for every possible subtree of the sentence, starting from word j and spanning k words  $(w_j,w_{j+1},\ldots,w_{j+k})$ , which are generated by rule Z. The process goes from spanning sequences of length one to n and for each size partitioning the subtree into two parts X, and Y to compute the probability that each production rule  $Z \to X, Y$  generates the subsequence. Those of length one correspond to the set of terminal rules  $Z \to w_i$ , to which it is assign a conditional probability  $\delta(i,i,Z) = \theta_{Z \to w_i}$ . At each step the most probable rule  $Z \to X, Y$  is stored in  $\psi(i,j,Z)$ . Once the process is completed, the sentence is recognized by the grammar if the subsequence containing the entire sentence is matched by the start symbol, and the most probable parse tree  $\hat{t}$  can be recovered by iterating backwards through the stored rules.

**Exercise 4.1** In this simple exercise, you will see the CKY algorithm in action. There is a Javascript applet that illustrates how CKY works (in its non-probabilistic form). Go to http://lxmls.it.pt/2015/cky.html, and observe carefully the several steps taken by the algorithm. Write down a small grammar in CNF that yields multiple parses for the ambiguous sentence The man saw the boy in the park with a telescope, and run the demo for this particular sentence. What would happen in the probabilistic form of CKY?

## 4.1.5 Learning the Grammar

There is an immense body of work on *grammar induction* using probabilistic models (see *e.g.*, Manning and Schütze (1999) and the references therein, as well as the most recent works of Klein and Manning (2002); Smith and Eisner (2005); Cohen et al. (2008)): this is the problem of learning the parameters of a grammar from plain sentences only. This can be done in an EM fashion (like in sequence models), except that the forward-backward algorithm is replaced by inside-outside. Unfortunately, the performance of unsupervised parsers is far from good, at present days. Much better results have been produced by supervised systems, which, however, require expensive annotation in the form of *treebanks*: this is a corpus of sentences annotated with their corresponding syntactic trees. The following is an example of an annotated sentence in one of the most widely used treebanks, the *Penn Treebank* (http://www.cis.upenn.edu/~treebank/):

Exercise 4.2 This exercise will show you that real-world sentences can have complicated syntactic structures. There is a parse tree visualizer in http://www.ark.cs.cmu.edu/treeviz/. Go to your local data/treebanks folder and open the file PTB\_excerpt.txt. Copy a few trees from the file, one at a time, and examine their parse trees in the visualizer.

A treebank makes possible to learn a parser in a supervised fashion. The simplest way is via a generative approach. Instead of counting transition and observation events of an HMM (as we did for sequence models), we now need to count production rules and symbol occurrences to estimate the parameters of a probabilistic context-free grammar. While performance would be much better than that of unsupervised parsers, it would still be rather poor. The reason is that the model we have described so far is oversimplistic: it makes too strong independence assumptions. In this case, the Markovian assumptions are:

- 1. Each terminal symbol w in some position i is independent of everything else given that it was derived by the rule  $Z \to w$  (i.e., given its parent Z);
- 2. Each pair of non-terminal symbols X and Y spanning positions i to j, with split point k, is independent of everything else given that they were derived by the rule  $Z \to XY$  (i.e., given their parent Z).

<sup>&</sup>lt;sup>3</sup>Similarly, the forward-backward algorithm for computing posterior marginals in sequence models can be extended for context-free parsing. It takes the name *inside-outside algorithm*. See Manning and Schütze (1999) for details.

The next section describes some model refinements that complicate the problem of parameter estimation, but usually allow for a dramatic improvement on the quality of the parser.

#### 4.1.6 Model Refinements

A number of refinements has been made that yield more accurate parsers. We mention just a few:

**Parent annotation.** This strategy splits each non-terminal symbol in the grammar (*e.g.* Z) by annotating it with all its possible parents (*e.g.* creates nodes  $Z^X, Z^Y, \ldots$  every time production rules like  $X \to Z \cdot, X \to \cdot Z$ , or  $Y \to Z \cdot, Y \to \cdot Z$  exist in the original grammar). This increases the vertical Markovian length of the model, hence weakening the independence assumptions. Parent annotation was initiated by Johnson (1998) and carried on in the unlexicalized parsers of Klein and Manning (2003) and follow-up works.

**Lexicalization.** A particular weakness of PCFGs is that they ignore word context for interior nodes in the parse tree. Yet, this context is relevant in determining the production rules that should be invoked in the derivation. A way of overcoming this limitation is by *lexicalizing* parse trees, *i.e.*, annotating each phrase node with the lexical item (word) which governs that phrase: this is called the *head* word of the phrase. Fig. 4.1 shows an example of a lexicalized parse tree. To account for lexicalization, each non-terminal symbol in the grammar (*e.g.* Z) is split into many symbols, each annotated with a word that may govern that phrase (*e.g.*  $Z^{w_1}, Z^{w_2}, \ldots$ ). This greatly increases the size of the grammar, but it has a significant impact in performance. A string of work involving lexicalized PCFGs includes Magerman (1995); Charniak (1997); Collins (1999).

**Discriminative models.** Similarly as in sequence models (where it was shown how to move from an HMM to a CRF), we may abandon our generative model and consider a discriminative one. An advantage of doing that is that it becomes much easier to adopt non-local input features (*i.e.*, features that depend arbitrarily on the surface string), for example the kind of features obtained via lexicalization, and much more. The CKY parsing algorithm can still be used for decoding, provided the feature vector decompose according to *non-terminal symbols* and *production rules*. In this case, productions and non-terminals will have a score which does not correspond to a log-probability; the partition function and the posterior marginals can be computed with the inside-outside algorithm. See Taskar et al. (2004) for an application of structured SVMs to parsing, and Finkel et al. (2008) for an application of CRFs.

**Latent variables.** Splitting the variables in the grammar by introducing latent variables appears as an alternative to lexicalization and parent annotation. There is a string of work concerning latent variable grammars, either for the generative and discriminative cases: Petrov and Klein (2007, 2008a,b). Some related work also considers coarse-to-fine parsing, which iteratively applies more and more refined models: Charniak et al. (2006).

History-based parsers. Finally, there is a totally different line of work which models parsers as a sequence of greedy shift-reduce decisions made by a push-down automaton (Ratnaparkhi, 1999; Henderson, 2003). When discriminative models are used, arbitrary conditioning can be done on past decisions made by the automaton, allowing to include features that are difficult to handle by the other parsers. This comes at the price of greediness in the decisions taken, which implies suboptimality in maximizing the desired objective function.

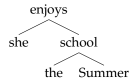
## 4.2 Dependency Parsing

#### 4.2.1 Motivation

Consider again the sentence

She enjoys the Summer school.

along with the lexicalized parse tree displayed in Fig. 4.1. If we drop the phrase constituents and keep only the head words, the parse tree would become:



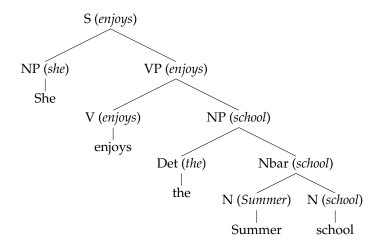


Figure 4.1: A lexicalized parse tree for the sentence She enjoys the Summer school.

This representation is called a *dependency tree*; it can be alternatively represented as shown in Fig. 4.2. Dependency trees retain the lexical relationships involved in lexicalized phrase-based parse trees. However, they drop phrasal constituents, which render non-terminal nodes unnecessary. This has computational advantages (no grammar constant is involved in the complexity of the parsing algorithms) as well as design advantages (no grammar is necessary, and treebank annotations are way simpler, since no internal constituents need to be annotated). It also shifts the focus from internal syntactic structures and generative grammars (Chomsky, 1965) to lexical and transformational grammars (Tesnière, 1959; Hudson, 1984; Melčuk, 1988; Covington, 1990). Arcs connecting words are called *dependency links* or *dependency arcs*. In an arc  $\langle h, m \rangle$ , the source word h is called the *head* and the target word m is called the *modifier*.

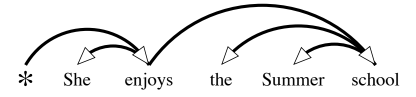


Figure 4.2: A dependency tree for the sentence *She enjoys the Summer school*. Note the additional dummy root symbol (\*) which is included for convenience.

#### 4.2.2 Projective and Non-projective Parsing

Dependency trees constructed using the method just described (*i.e.*, lexicalization of context-free phrase-based trees) always satisfy the following properties:

- 1. Each word (excluding the dummy root symbol) has exactly one parent.
- 2. The dummy root symbol has no parents.
- 3. There are no cycles.
- 4. The dummy root symbol has exactly one child.
- 5. All arcs are *projective*. This means that for any arc  $\langle h, m \rangle$ , all words in its span (*i.e.*, all words lying between h and m) are descendents from h (i.e. there is a directed path of dependency links from h to such word).

Conditions 1–3 ensure that the set of dependency links form a well-formed tree, rooted in the dummy symbol, which spans all the words of the sentence. Condition 4 requires that there is a single link departing from the root. Finally, a tree satisfying condition 5 is said *projective*: it implies that arcs cannot cross (*e.g.*, we cannot have arcs  $\langle h, m \rangle$  and  $\langle h', m' \rangle$  such that the word positions are ordered as h < h' < m < m').

In many languages (*e.g.*, those which have free-order) we would like to relax the assumption that all trees must be projective. Even in languages which have fixed word order (such as English) there are syntactic

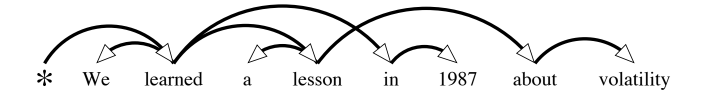


Figure 4.3: A non-projective parse tree.

phenomena which are awkward to characterize using projective trees arising from the context-free assumption. Usually, such phenomena are characterized with additional linguistic artifacts (e.g., traces, Wh-movement, etc.). An example is the sentence (extracted from the Penn Treebank)

We learned a lesson in 1987 about volatility.

There, the prepositional phrase *in 1987* should be attached to the verb phrase headed by *learned* (since this is *when* we learned the lesson), but the other prepositional phrase *about volatility* should be attached to the noun phrase headed by *lesson* (since the *lesson* was about volatility). To explain such phenomenon, context-free grammars need to use additional machinery which allows words to be scrambled (in this case, via a movement transformation and the consequent insertion of a trace). In the dependency-based formalism, we can get rid of all those artifacts altogether by allowing *non-projective* parse trees. These are trees that satisfy conditions 1-3 above, but not necessarily conditions 4 or 5.4 The dependency tree in Fig. 4.3 is non-projective: note that the arc  $\langle lesson, about \rangle$  is not projective.

We end this section by mentioning that dependency trees can have their arcs labeled, to provide more detailed syntactic information. For example, the arc  $\langle enjoys, She \rangle$  could be labeled as SUBJ to denote that the modifier She has a subject function, and the arc  $\langle enjoys, school \rangle$  could be labeled as OBJ to denote that the modifier school has an object function. For simplicity, we resort to unlabeled trees, which just convey the backbone structure. To cope with the labels, one can use either a joint model that infers the backbone and labels altogether, or to have a two-stage approach that first gets the backbone structure, and then the arc labels.

#### 4.2.3 Algorithms for Projective Dependency Parsing

We now turn our attention to *algorithms* for obtaining a dependency parse tree. We start by considering a simple kind of models which are called *arc-factored*. These models assign a score  $s_{\theta}(h, m)$  to each possible arc  $\langle h, m \rangle$  connecting a pair of words; they then score a particular dependency tree t by summing over the individual scores of the arcs that are present in the tree:

$$score_{\theta}(t) = \sum_{\langle h, m \rangle \in t} s_{\theta}(h, m).$$

As usual, from the point of view of the parsing algorithm, it does not matter whether the scores come from a generative or discriminative approach, and which features were used to compute the scores. The three important inference tasks are:

1. Obtain the tree with the largest score,

$$\hat{t} = \underset{t}{\operatorname{arg max score}}_{\theta}(t).$$

2. Compute the partition function (for a log-linear model),

$$Z(\mathbf{s}_{\theta}) = \sum_{t} \exp(\operatorname{score}_{\theta}(t)),$$

where  $\mathbf{s}_{\theta}$  is short-hand notation for the set of all the  $s_{\theta}(h, m)$  coefficients.

<sup>&</sup>lt;sup>4</sup>It is also common to impose conditions 1–4, in which case the tree need not be projective, but it must have a single link departing from the root. The algorithms to be described below can be adapted for this case.

3. Compute the posterior marginals for all the possible arcs (which for a log-linear model is the gradient of the log-partition function),

$$P_{\theta}(\langle h, m \rangle \in t) = \frac{\partial \log Z(\mathbf{s}_{\theta})}{\partial s_{\theta}(h, m)}.$$

As explained in this morning's lecture, in *projective* dependency parsing using arc-factored models, the three tasks above can be solved in time  $O(N^3)$ , through Eisner's algorithm (Eisner, 1996), which is represented as Algorithm 13. Eisner's algorithm is related with the CKY algorithm in the sense that it also parses a sentence in a bottom-up fashion, building larger spans from smaller spans. In fact, the CKY algorithm can be adapted in a naïve manner to perform projective dependency parsing by dropping the constituent symbols and keeping track of the head positions of each span. This naïve strategy, however, would force us to manipulate five indices when combining two spans to form a larger span (the start and end points of the large span, the mid point, and the head positions for the two spans being combined), which would increase the complexity of the algorithm to  $O(N^5)$ . Eisner's algorithm uses the notion of *complete* and *incomplete* spans (in which all the right modifiers and all the left modifiers of a given head are all built separately) to reduce the complexity back to  $O(N^3)$ . By replacing maximizations with summations, the same dynamic programming algorithm can be used to compute the partition function and the posterior marginals.

## Algorithm 13 Eisner's algorithm for first-order projective dependency parsing

```
1: input: Arc scores s_{\theta}(h, m), for h \in \{0, ..., N\}, m \in \{1, ..., N\}, and h \neq m, associated with a sentence
    s = w_1 \dots w_N.
 2: {Initialization}
 3: for i = 0 to N do
       {Initialize incomplete spans.}
 4:
 5:
       incomplete[i, i, \leftarrow] := 0.0
       incomplete[i, i, \rightarrow] := 0.0
 6:
 7:
 8:
       {Initialize complete spans.}
 9:
       complete[i, i, \leftarrow] := 0.0
       complete[i, i, \rightarrow] := 0.0
10:
11: end for
12:
   {Induction}
13:
14: for k = 1 to N do {k is length of span}
       for s = 0 to N - k do \{s \text{ is start of span}\}\
15:
          Set t := s + k \{ t \text{ is end of span} \}
16:
17:
          {First, create incomplete spans.}
18:
          incomplete[s, t, \leftarrow] := \max_{s < r < t} (complete[s, r, \rightarrow] + complete[r + 1, t, \leftarrow] + s_{\theta}(t, s))
19:
20:
          incomplete[s, t, \rightarrow] := \max_{s \le r < t} (complete[s, r, \rightarrow] + complete[r + 1, t, \leftarrow] + s_{\theta}(s, t))
21:
          {Then, create complete spans.}
22:
          complete[s, t, \leftarrow] := \max_{s < r < t}(complete[s, r, \leftarrow] + incomplete[r, t, \leftarrow])
23:
          complete[s, t, \rightarrow] := \max_{s < r < t} (incomplete[s, r + 1, \rightarrow] + complete[r + 1, t, \rightarrow])
24:
25:
       end for
    end for
26:
27:
    {Termination}
29: Backtrack to obtain the actual tree, whose score is complete [0, N, \rightarrow].
```

## 4.2.4 Algorithms for Non-Projective Dependency Parsing

We turn our attention to *non-projective* dependency parsing. In that case, efficient solutions also exist for the three problems above; interestingly, they are based in combinatorial algorithms which are not related at all with dynamic programming:

• The first problem corresponds to finding the maximum weighted directed spanning tree in a directed graph. This problem is well known in combinatorics and can be solved in  $O(N^3)$  using Chu-Liu-Edmonds'

algorithm (Chu and Liu, 1965; Edmonds, 1967).<sup>5</sup> This has first been noted by McDonald et al. (2005).

• The second and third problems can be solved by invoking another important result in combinatorics, the *matrix-tree theorem* (Tutte, 1984). This fact has been noticed independently by Smith and Smith (2007); Koo et al. (2007); McDonald and Satta (2007). The cost is that of computing a determinant and inverting a matrix, which can be done in time  $O(N^3)$ . The procedure is as follows. We first consider the directed weighted graph formed by including all the possible dependency links  $\langle h, m \rangle$  (including the ones departing from the dummy root symbol, for which h = 0 by convention), along with weights given by  $\exp(s_{\theta}(h, m))$ , and compute its (N + 1)-by-(N + 1) Laplacian matrix L whose entries are:

$$L_{hm} = \begin{cases} \sum_{h'=0}^{N} \exp(s_{\theta}(h', m)), & \text{if } h = m, \\ -\exp(s_{\theta}(h, m))), & \text{otherwise.} \end{cases}$$
 (4.2)

Denote by  $\hat{L}$  the (0,0)-minor of L, *i.e.*, the matrix obtained from L by removing the first row and column. Consider its determinant det  $\hat{L}$  and its inverse  $\hat{L}^{-1}$ . Then:

- the partition function is given by

$$Z(\mathbf{s}_{\theta}) = \det \hat{L}; \tag{4.3}$$

- the posterior marginals are given by

$$P_{\theta}(\langle h, m \rangle \in t) = \begin{cases} \exp(s_{\theta}(h, m)) \cdot ([\hat{L}^{-1}]_{mm} - [\hat{L}^{-1}]_{mh}) & \text{if } h \neq 0 \\ \exp(s_{\theta}(0, m)) \cdot [\hat{L}^{-1}]_{mm} & \text{otherwise.} \end{cases}$$
(4.4)

**Exercise 4.3** In this exercise you are going to experiment with arc-factored non-projective dependency parsers.

The CoNLL-X and CoNLL 2008 shared task datasets (Buchholz and Marsi, 2006; Surdeanu et al., 2008) contain dependency treebanks for 14 languages. In this lab, we are going to experiment with the Portuguese and English datasets. We preprocessed those datasets to exclude all sentences with more than 15 words; this yielded the files:

- data/deppars/portuguese\_train.conll,
- data/deppars/portuguese\_test.conll,
- data/deppars/english\_train.conll,
- data/deppars/english\_test.conll.
- 1. After importing all the necessary libraries, load the Portuguese dataset:

```
import sys
sys.path.append('.')

import lxmls.parsing.dependency_parser as depp

dp = depp.DependencyParser()
dp.read_data("portuguese")
```

Observe the statistics which are shown. How many features are there in total?

2. We will now have a close look on the features that can be used in the parser. Examine the file:

```
lxmls/parsing/dependency_features.py.
```

*The following method takes a sentence and computes a vector of features for each possible arc*  $\langle h, m \rangle$ *:* 

```
def create_arc_features(self, instance, h, m, add=False):
    '''Creates features for arc h-->m.'''
```

We grouped the features in several subsets, so that we can conduct some ablation experiments:

There is a asymptotically faster algorithm by Tarjan (1977) which solves the same problem in  $O(N^2)$ .

- Basic features that look only at the parts-of-speech of the words that can be connected by an arc;
- Lexical features that also look at these words themselves;
- Distance features that look at the length and direction of the dependency link (i.e., distance between the two words);
- Contextual features that look at the context (part-of-speech tags) of the words surrounding h and m.

In the default configuration, only the basic features are enabled. The total number of features is the quantity observed in the previous question. With this configuration, train the parser by running 10 epochs of the structured perceptron algorithm:

```
dp.train_perceptron(10)
dp.test()
```

What is the accuracy obtained in the test set? (Note: the shown accuracy is the fraction of words whose parent was correctly predicted.)

3. Repeat the previous exercise by subsequently enabling the lexical, distance and contextual features:

```
dp.features.use_lexical = True
dp.read_data("portuguese")
dp.train_perceptron(10)
dp.test()

dp.features.use_distance = True
dp.read_data("portuguese")
dp.train_perceptron(10)
dp.test()

dp.features.use_contextual = True
dp.read_data("portuguese")
dp.train_perceptron(10)
dp.test()
```

For each configuration, write down the number of features and test set accuracies. Observe the improvements obtained when more features were added.

Feel free to engineer new features!

4. Which of the three important inference tasks discussed above (computing the most likely tree, computing the partition function, and computing the marginals) need to be performed in the structured perceptron algorithm? What about a maximum entropy classifier, with stochastic gradient descent? Check your answers by looking at the following two methods in code/dependency\_parser.py:

```
def train_perceptron(self, n_epochs):
...

def train_crf_sgd(self, n_epochs, sigma, eta0 = 0.001):
...
```

Repeat the last exercise by training a maximum entropy classifier, with stochastic gradient descent, using  $\lambda = 0.01$  and a initial stepsize of  $\eta_0 = 0.1$ :

```
dp.train_crf_sgd(10, 0.01, 0.1)
dp.test()
```

Compare the results with those obtained by the perceptron algorithm.

5. Train a parser for English using your favourite learning algorithm:

```
dp.read_data("english")
dp.train_perceptron(10)
dp.test()
```

The predicted trees are placed in the file data/deppars/english\_test.conll.pred. To get a sense of which errors are being made, you can check the sentences that differ from the gold standard (see the data in data/deppars/english\_test.conll) and visualize those sentences, e.g., in http://www.ark.cs.cmu.edu/treeviz/.

6. (Optional.) Implement Eisner's algorithm for projective dependency parsing. The pseudo-code is shown as Algorithm 13. Implement this algorithm as the function

in file dependency\_decoder.py. The input is a matrix of arc scores, whose dimension is (N+1)-by-(N+1), and whose (h,m) entry contains the score  $s_{\theta}(h,m)$ . In particular, the first row contains the scores for the arcs that depart from the root, and the first column's values, along with the main diagonal, are to be ignored (since no arcs point to the root, and there are no self-pointing arcs). To make your job easier, we provide an implementation of the backtracking part:

so you just need to build complete/incomplete spans and their backtrack pointers and then call

```
heads = -np.ones(N+1, dtype=int)
self.backtrack_eisner(incomplete_backtrack, complete_backtrack, 0, N, 1, 1,
heads)
return heads
```

to obtain the final parse.

To test the algorithm, retrain the parser on the English data (where the trees are actually all projective) by setting the flag dp.projective to True:

```
dp = depp.DependencyParser()
dp.features.use_lexical = True
dp.features.use_distance = True
dp.features.use_contextual = True
dp.read_data("english")
dp.projective = True
dp.train_perceptron(10)
dp.test()
```

You should get the following results:

```
Number of sentences: 8044
Number of tokens: 80504
Number of words: 12202
Number of pos: 48
Number of features: 338014
Epoch 1
Training accuracy: 0.835637168541
Epoch 2
Training accuracy: 0.922426254687
Epoch 3
Training accuracy: 0.947621628947
Epoch 4
Training accuracy: 0.960326602521
Epoch 5
Training accuracy: 0.967689840538
Epoch 6
Training accuracy: 0.97263631025
Training accuracy: 0.97619370285
Epoch 8
Training accuracy: 0.979209016579
Epoch 9
Training accuracy: 0.98127569228
Epoch 10
Training accuracy: 0.981320865519
Test accuracy (509 test instances): 0.886732599366
```

#### 4.2.5 Model Refinements

A number of refinements has been made that yield more accurate dependency parsers. We mention just a few:

**Sibling and grandparent features.** The arc-factored assumption fails to capture correlations between pairs of arcs. The dynamic programming algorithms for the *projective* case can be extended (at some additional cost) to handle features that look at consecutive sibling arcs on the same side of the head (*i.e.*, pairs of arcs of the form  $\langle h, m \rangle$  and  $\langle h, s \rangle$  with h < m < s or h > m > s, such that no arc  $\langle h, r \rangle$  exists with r between m and s. This has been done by Eisner and Satta (1999). Similarly, grandparents can also be accommodated with similar extensions (Carreras, 2007). These are called "second-order models."

For the non-projective case, however, any extension beyond the arc-factored model becomes NP-hard (McDonald and Satta, 2007). Yet, approximate algorithms have been proposed to handle "second-order models" that seem to work well: a greedy method (McDonald et al., 2006), loopy belief propagation (Smith and Eisner, 2008), a linear programming relaxation (Martins et al., 2009), and a dual decomposition method (Koo et al., 2010).

**Third-order models.** For the projective case, third order models have also been considered (Koo and Collins, 2010). This was extended to the non-projective case by Martins et al. (2013).

**Transition-based parsers.** Like in the phrase-based case, there is a totally different line of work which models parsers as a sequence of greedy shift-reduce decisions (Nivre et al., 2006; Huang and Sagae, 2010). These parsers seem to be very fast (expected linear time) and only slightly less accurate than the state-of-the-art. Solutions have been worked out for the non-projective case also (Nivre, 2009).

#### 4.2.6 External Links

If you want to check a demo of a parser that use some of the extensions above, you can do so at

```
http://demo.ark.cs.cmu.edu/parse
```

This demo uses TurboParser (http://www.ark.cs.cmu.edu/TurboParser), a free software implementation of a fast and accurate dependency parser that can parse thousands of tokens per second with state-of-theart accuracies (way faster than the Python implementation that you have used at the labs). Feel free to try it at home!

Other well-known software toolkits that implement dependency parsers are:

- MSTParser(http://www.seas.upenn.edu/~strctlrn/MSTParser/MSTParser.html), a graph-based dependency parser;
- MaltParser (http://www.maltparser.org/), a transition-based parser;
- DPO3 (http://groups.csail.mit.edu/nlp/dpo3/), a third-order projective parser;
- Linear-time dynamic programming parser (http://acl.cs.qc.edu/~lhuang/), a fast transition-based parser.

## Day 5

# Deep Learning I

## 5.1 Today's assignment

Today's class will be divided in two main parts. Firstly, we will learn about computation graphs and the Backpropagation algorithm. We will also code Backpropagation in Numpy for a simple model (a feed forward network). Secondly, we will learn the basics of the Theano module for Python. Theano allows you to implement generic computation graphs, automatically compute gradients with respect to their parameters and make use of Graphical Processing Units (GPUs) for optimal speed.

If you are new to the topic, you should aim to understand the concept of computation graph, finish the Backpropagation exercise and attain a basic understanding of Theano. If you already know Backpropagation well and have experience with normal Python, you should aim to complete the whole day.

## 5.2 Introduction to Deep Learning and Theano

Deep learning is the name behind the latest wave of neural network research. This is a very old topic, dating from the first half of the 20th century, that has attained formidable impact in the machine learning community recently. There is nothing particularly difficult in deep learning. You have already visited all the mathematical principles you need in the first days of the labs of this school. At their core, deep learning models are just functions mapping vector inputs **x** to vector outputs **y**, constructed by composing linear and non-linear functions. This composition can be expressed in the form of a *computation graph*, where each node applies a function to its inputs and passes the result as its output. The parameters of the model are the weights given to the different inputs in nodes applying linear functions. This vaguely resembles synapse strengths in human neural networks, hence the name artificial neural networks.

Due to their compositional nature, gradient methods and the chain rule can be applied learn the parameters of these models regardless of their complexity. See Section for a refresh on the basic concept. We will also refer to the gradient learning methods introduced in Section 1.4.4. Today we will focus on *feed-forward networks*. Tomorrow we will extended today's class to *recurrent neural networks* (RNNs).

Some of the changes that led to the surge of deep learning are not only improvements on the existing neural network algorithms, but also the increase in the amount of data available and computing power. In particular, the use of Graphical Processing Units (GPUs) has allowed neural networks to be applied to very large datasets. Working with GPUs is not trivial as it requires dealing with specialized hardware. Luckily, as it is often the case, we are one Python import away from solving this problem.

For the particular case of deep learning, there is a growing number of python toolboxes available that allow you to design custom computational graphs for GPUs as e.g. Theano<sup>1</sup> or TensorFlow<sup>2</sup>.

In these labs we will be working with Theano. Theano allows us to express computation graphs symbolically in terms of basic algebraic operations. It also automatically computes gradients and produces CUDA-compatible code for GPUs. The exercises are designed to gain a low-level understanding of Theano. If you are only looking forward to utilize pre-designed models, the Keras toolbox<sup>3</sup> provides high-level operations compatible with both Theano and TensorFlow.

<sup>&</sup>lt;sup>1</sup>http://deeplearning.net/software/theano/

<sup>&</sup>lt;sup>2</sup>https://www.tensorflow.org/

<sup>&</sup>lt;sup>3</sup>http://keras.io/

## 5.3 Computation Graphs

## 5.3.1 Example: The computation graph of a log-linear model

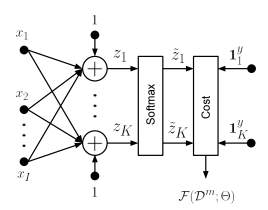


Figure 5.1: Representation of a log-linear model as a computation graph: a composition of linear and non-linear transformations. The classification cost for the m-th training example  $\mathbb{D}^m = \{x, y\}$  is also shown. Note  $\mathbf{1}^y$  is an indicator vector with a one in position y.

A computation graph is just a way of expressing a multi-input multi-output function with a directed acyclic graph. Each node in the graph applies a function over the outputs of its parent nodes and passes the result to its child-nodes, thus attaining very complex functions by *composing* simpler functions. To introduce the concept we will work with a very simple example, a categorical distribution over *K* classes parametrized as a log-linear model. This is given by

$$p(y = k|x) = \frac{1}{Z(\mathbf{W}, \mathbf{b}, \mathbf{x})} \exp\left(\sum_{i=1}^{I} W_{ik} x_i + b_k\right), \tag{5.1}$$

where

$$Z(\mathbf{W}, \mathbf{b}, \mathbf{x}) = \sum_{k'=1}^{K} \exp\left(\sum_{i=1}^{I} W_{ik'} x_i + b_{k'}\right)$$
 (5.2)

is the partition function ensuring that all output values sum to one, so that we can interpret these values as probabilities. The model thus receives a feature vector  $\mathbf{x} \in \mathbb{R}^I$  and assigns a probability over  $y \in 1 \cdots K$  possible class indices. It is parametrized by weights and bias  $\Theta = \{\mathbf{W}, \mathbf{b}\}$ , with  $\mathbf{W} \in \mathbb{R}^{K \times J}$  and  $\mathbf{b} \in \mathbb{R}^K$ . This is a well known classifier bearing resemblance to the models seen in the second day of the school, in particular the maxent model<sup>4</sup>.

Another way of looking at these models is as the composition of the linear transformation

$$z_k = \sum_{i=1}^{I} W_{ik} x_i + b_k, (5.3)$$

with the softmax non-linear transformation

$$p(y=k|x) \equiv \tilde{z}_k = \frac{\exp(z_k)}{\sum_{k'=1}^K \exp(z_{k'})}.$$
(5.4)

This is shown as a computation graph in Fig. 5.1. Note that in the following sections we will also use  $\mathbf{z}$  and  $\tilde{\mathbf{z}}$  to denote the output of linear and non-linear functions respectively.

<sup>&</sup>lt;sup>4</sup>There are some differences with respect to Eq.1.26, like the use of a bias **b**. Also, if we consider the binary joint feature mapping  $f(x,y) = g(x) \otimes e_y$  of Eq.1.16, the maximum entropy classifier in Eq.1.26 becomes a special case of Eq.5.1, in which the feature vector **x** only take binary values and the bias parameters in **b** are all set to zero.

#### 5.3.2 Stochastic Gradient Descent: a refresher

As we saw on day one, the parameters of a log linear model  $\Theta = \{W, b\}$  can be learned with Stochastic Gradient Descent (SGD). To apply SGD we first need to define an error function that measures how good we are doing for any given parameter values. To remain close to the maximum entropy example, we will use as cost function the average minus posterior probability of the correct class, also known as the Cross-Entropy (CE) criterion. Bear in mind, however, that we could pick other non-linear functions and cost functions that do not have a probabilistic interpretation. For example, the same principle could be applied to a regression problem where the cost is the Mean Square Error (MSE).

For a training data-set  $\mathcal{D} = \{(\mathbf{x}^1, y^1), \dots, (\mathbf{x}^M, y^M)\}$  of M examples, the CE cost function is given by

$$\mathcal{F}(\mathcal{D};\Theta) = -\frac{1}{M} \sum_{m=1}^{M} \log p(y^m = k(m) | \mathbf{x}^m), \tag{5.5}$$

where k(m) is the correct class index for the m-th example. To learn the parameters of this model with SGD, all we need to do is compute the gradient of the cost  $\nabla \mathcal{F}$  with respect to the parameters of the model and iteratively update our parameter estimates as

$$\mathbf{W} \leftarrow \mathbf{W} - \eta \nabla_{\mathbf{W}} \mathcal{F} \tag{5.6}$$

and

$$\mathbf{b} \leftarrow \mathbf{b} - \eta \nabla_{\mathbf{b}} \mathcal{F},$$
 (5.7)

where  $\eta$  is the learning rate. Note that in practice we will use a mini-batch of examples as opposed to the whole train set. Very often, more elaborated learning rules as e.g. momentum or Adagrad are used. Bear in mind that, in general, these still require the computation of the gradients as the main step. The reasoning here outlined will also be applicable to these.

## 5.3.3 Deriving Gradients in Computation Graphs using the Chain Rule

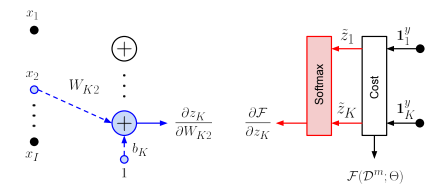


Figure 5.2: Forward-pass (blue) and Backpropagation (red) calculations to estimate the gradient of weight  $W_{K2}$  and bias  $b_K$  of a log-linear model.

The expressions for  $\nabla \mathcal{F}$  are well known in the case of log-linear models. However, for the sake of the introduction to deep learning, we will show how they can be derived by exploiting the decomposition of the cost function into the computational graph seen in the last section (and represented in Fig. 5.1). To simplify notation, and without loss of generality, we will work with the classification cost of an individual example

$$\mathcal{F}(\mathcal{D}^m; \Theta) = -\log p(y^m = k(m)|\mathbf{x}^m), \tag{5.8}$$

where  $\mathcal{D}^m = \{(\mathbf{x}^m, y^m)\}$ . Due to linearity, extending the gradients of Eq. 5.8 to M examples as in Eq. 5.5 simply requires computing the average of the gradients for the individual examples. That is

$$\nabla_{\mathbf{W}} \mathcal{F}(\mathcal{D}; \Theta) = -\frac{1}{M} \sum_{m=1}^{M} \nabla_{\mathbf{W}} \log p(y^m = k(m) | \mathbf{x}^m)$$
 (5.9)

All we need to do to compute  $\nabla_{\mathbf{W}} \mathfrak{F}(\mathfrak{D}; \Theta)$  is to have an expression for  $\nabla_{\mathbf{W}} \log p(y^m = k(m) | \mathbf{x}^m)$ .

Lets start by computing the element (k,i) of the gradient matrix  $\nabla_{\mathbf{W}} \mathcal{F}(\mathcal{D};\Theta)$ , which contains the partial derivative with respect to the weight  $W_{ki}$ . To do this, we invoke the **chain rule** to split the derivative calculation into two terms at variable  $z_{k'}$  (Eq.5.3) with  $k' = 1 \cdots K$ 

$$\frac{\partial \mathcal{F}(\mathcal{D}; \Theta)}{\partial W_{ki}} = \sum_{k'=1}^{K} \frac{\partial \mathcal{F}(\mathcal{D}; \Theta)}{\partial z_{k'}} \frac{\partial z_{k'}}{\partial W_{ki}}.$$
 (5.10)

We have thus transformed the problem of computing the derivative into computing two easier derivatives. Since  $z_{k'}$  only depends on the weight  $W_{ki}$  in a linear way (see the graph in Fig. 5.2), the second derivative in Eq.5.10 is given by

$$\frac{\partial z_{k'}}{\partial W_{ki}} = \frac{\partial}{\partial W_{ki}} \left( \sum_{i'=1}^{I} W_{k'i'} x_{i'}^m + b_{k'} \right) = \begin{cases} x_i^m & \text{if } k = k' \\ 0 & \text{otherwise} \end{cases}, \tag{5.11}$$

which implies that

$$\frac{\partial \mathcal{F}(\mathcal{D}^m; \Theta)}{\partial W_{ki}} = \frac{\partial \mathcal{F}(\mathcal{D}^m; \Theta)}{\partial z_k} x_i^m. \tag{5.12}$$

The first term derivative is given by

$$\frac{\partial \mathcal{F}(\mathcal{D}^m; \Theta)}{\partial z_k} = \frac{\partial}{\partial z_k} \left( z_{k(m)} - \log \left( \sum_{k'=1}^K \exp(z_{k'}) \right) \right) = \begin{cases} 1 - \tilde{z}_k & \text{if } k = k(m) \\ -\tilde{z}_k & \text{otherwise} \end{cases}.$$
 (5.13)

From the formula for each element and a single example, we can now obtain the gradient matrix for a batch of *M* examples by simply averaging and expressing the previous equations in vector form as follows

$$\nabla_{\mathbf{W}} \mathcal{F}(\mathcal{D}; \Theta) = -\frac{1}{M} \sum_{m=1}^{M} \left( \mathbf{1}^{y^m} - \tilde{\mathbf{z}}^m \right) (\mathbf{x}^m)^T.$$
 (5.14)

Here  $\mathbf{1}^{y^m} \in \mathbb{R}^K$  is a vector of zeros with a one in  $y^m = k(m)$ , which is the index of the correct class for the example m.

In order to compute the derivatives of the cost function with respect to the bias parameters  $b_k$ , we only need to compute one additional derivative

$$\frac{\partial z_{k'}}{\partial b_k} = \begin{cases} 1 & \text{if } k = k' \\ 0 & \text{otherwise} \end{cases}$$
 (5.15)

This leads us to the last gradient expression

$$\nabla_{\mathbf{b}} \mathcal{F}(\mathcal{D}; \Theta) = -\frac{1}{M} \sum_{m=1}^{M} \left( \mathbf{1}^{y^m} - \tilde{\mathbf{z}}^m \right). \tag{5.16}$$

An important consequence of the previous derivation is the fact that each gradient of the parameters  $\nabla_{\mathbf{W}} \mathcal{F}(\mathcal{D}; \Theta)$  and  $\nabla_{\mathbf{b}} \mathcal{F}(\mathcal{D}; \Theta)$  can be computed from two terms:

- 1. The derivative of the cost with respect to the linear transformation applying the weight  $\partial \mathcal{F}(\mathcal{D}^m; \Theta) / \partial z_k$ , denoted in red.
- 2. The derivative of the forward-pass up to the linear transformation applying the weight  $\partial z_k/\partial W_{ik}$ , denoted in blue. This is always equal to the variable multiplying that weight and one in the case of the bias.

This is the key to the Backpropagation algorithm as we will see in the next Section 5.4.

## 5.4 Going Deeper than Log-linear by using Composition

## 5.4.1 The Multilayer Perceptron or Feed-Forward Network

We have seen that just by using the chain rule we can easily compute gradients for compositions of two functions (one non-linear and one linear). However, there was nothing in the derivation that would stop us from composing more than two functions. Let us think, for example of the following model

## Algorithm 14 Forward pass of a Multi-Layer Perceptron (MLP) or Feed-Forward (FF) network

- 1: **input:** Initial parameters for an MLP of *N* layers  $\Theta = \{ \mathbf{W}^1, \mathbf{b}^1, \cdots \mathbf{W}^N, \mathbf{b}^N \}$
- 2: **input**: Input data vector  $\tilde{\mathbf{z}}^0 \equiv \mathbf{x}$ .
- 3: **for** n = 1 **to** N 1 **do**
- 4: Apply linear transformation

$$z_j^n = \sum_{i=1}^I W_{ji}^n \tilde{z}_i^{n-1} + b_j^n$$

5: Apply non-linear transformation e.g. sigmoid (hereby denoted  $\sigma()$ )

$$\tilde{z}_j^n = \sigma(z_j^n) = \frac{1}{1 + \exp(-z_j^n)}$$

- 6: end for
- 7: Apply final linear transformation

$$z_{k}^{N} = \sum_{j=1}^{J} W_{kj}^{N} \tilde{z}_{j}^{N-1} + b_{k}^{N}$$

8: Apply final non-linear transformation e.g. softmax

$$p(y = k|x) \equiv \tilde{z}_k^N = \frac{\exp(z_k^N)}{\sum_{k'=1}^K \exp(z_k^N)}$$

In a similar fashion to the log-linear model, the MLP/FF can be expressed as a computation graph. This is displayed in Fig. 5.3

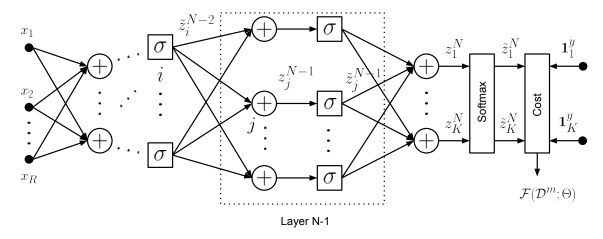


Figure 5.3: Representation of a Multi-Layer Perceptron (MLP) or Feed-Forward (FF) network as a computation graph. The classification cost for the m-th training example  $\mathbb{D}^m = \{x, y\}$  is also shown.

## Some considerations:

- MLPs/FFs are characterized by applying functions in a set of layers subsequently to a single input. This
  characteristic is also shared by convolutional networks, although the latter also have parameter tying
  constraints.
- The non-linearities in the intermediate layers are usually one-to-one transformations. The most typical are the sigmoid, hyperbolic tangent and the rectified linear unit (ReLU).

• The output non-linearity is determined by the output to be estimated. In order to estimate probability distributions the softmax is typically used. For regression problems a last linear layer is used instead.

## 5.4.2 Backpropagation: an overview

For the examples in this chapter we will consider the case in which we are estimating a distribution over classes, thus we will use the CE cost function (Eq. 5.5).

To compute the gradient with respect the parameters of the *n*-th layer, we just need to apply the chain rule as in the previous section, consecutively. Fortunately, we do not need to repeat this procedure for each layer as it is easy to spot a recursive rule (the Backpropagation recursion) that is valid for many neural models, including feed-forward networks (such as MLPs) as well as recurrent neural networks (RNNs) with minor modifications. The Backpropagation method, which is given in Algorithm 15 for the case of an MLP, consists of the following steps:

- The forward pass step, where the input signal is injected though the network in a forward fashion (see Alg. 14)
- The backpropagation step, where the derivative of the cost function (also called error) is injected back through the network and backpropagated according to the derivative rules (see steps 8-17 in Alg. 15)
- Finally, the gradients with respect to the parameters are computed by multiplying the input signal from the forward pass and the backpropagated error signal, at the corresponding places in the network (step 18 in Alg. 15)
- Given the gradients computed in the previous step, the model weights can then be easily update according to a specified learning rule (step 19 in Alg. 15 uses a mini-batch SGD update rule).

The main step of the method is the backpropagation step, where one has to compute the backpropagation recursion rules for a specific network. The next section presents a careful deduction of these recursion rules, for the present MLP model.

## 5.4.3 Backpropagation: deriving the rule

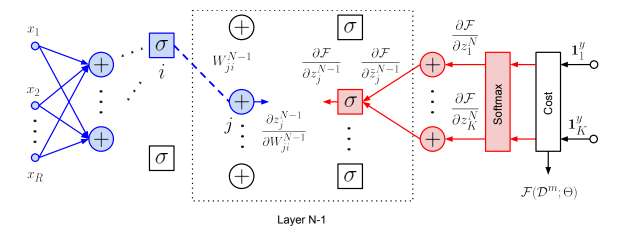


Figure 5.4: Forward-pass (blue) and Backpropagation (red) calculations to estimate the gradient of weight  $W_{ji}$  at layer N-1 of a MLP.

In a generic MLP we would like to compute the values of all parameters  $\Theta = \{\mathbf{W}^1, \mathbf{b}^1, \cdots \mathbf{W}^N, \mathbf{b}^N\}$ . As explained previously, we will thus need to compute the backpropagated error at each node  $\partial \mathcal{F}(\mathcal{D}^m; \Theta) / \partial z_k^n$ , and the corresponding derivative for the forward-pass  $\partial z_k^n / \partial W_{ik}$ , for  $n = 1 \cdots N$ . Fortunately, it is easy to spot a recursion that will allow us to compute these values for each node, given all its child nodes. To spot it, we can start trying to compute the gradients one layer before the output layer (see Fig. 5.4), i.e. layer N-1. We start by splitting at N-1

$$\frac{\partial \mathcal{F}(\mathcal{D}^m; \Theta)}{\partial W_{ji}} = \sum_{j'=1}^{J} \frac{\partial \mathcal{F}(\mathcal{D}^m; \Theta)}{\partial z_{j'}^{N-1}} \frac{\partial z_{j'}^{N-1}}{\partial W_{ji}} = \frac{\partial \mathcal{F}(\mathcal{D}^m; \Theta)}{\partial z_{j}^{N-1}} \frac{\partial z_{j}^{N-1}}{\partial W_{ji}}$$
(5.17)

where, as in the case of the log-linear model, we have used the fact that only  $z_j$  depends on  $W_{ji}$ . We now advance forward from  $z_j^N$  and apply the non-linearity to obtain  $\tilde{z}_j^N$ . As long as the non-linearity is a one-to-one operation (and this is often the case in neural-networks), we will also get rid of the summation leading to

$$\frac{\partial \mathcal{F}(\mathcal{D}^m; \Theta)}{\partial W_{ji}} = \left(\sum_{j'=1}^{J} \frac{\partial \mathcal{F}(\mathcal{D}^m; \Theta)}{\partial \tilde{z}_{j'}^{N-1}} \frac{\partial \tilde{z}_{j'}^{N-1}}{\partial z_{j}^{N-1}}\right) \frac{\partial z_{j}^{N-1}}{\partial W_{ji}} = \frac{\partial \mathcal{F}(\mathcal{D}^m; \Theta)}{\partial \tilde{z}_{j}^{N-1}} \frac{\partial \tilde{z}_{j}^{N-1}}{\partial z_{j}^{N-1}} \frac{\partial z_{j}^{N-1}}{\partial W_{ji}}.$$
 (5.18)

We will not be so lucky in the next application of the chain rule. Looking at Fig. 5.4, it is clear that the derivatives at each node in layer N-1 will depend on all values of layer N. We thus have

$$\frac{\partial \mathcal{F}(\mathcal{D}^m; \Theta)}{\partial W_{ji}} = \left(\sum_{k'=1}^K \frac{\partial \mathcal{F}(\mathcal{D}^m; \Theta)}{\partial z_{k'}^N} \frac{\partial z_{k'}^N}{\partial \tilde{z}_i^{N-1}}\right) \frac{\partial \tilde{z}_j^{N-1}}{\partial z_j^{N-1}} \frac{\partial z_j^{N-1}}{\partial W_{ji}}.$$
 (5.19)

Now, comparing Eq. 5.17 with Eq. 5.19 it is easy to see that, if we denote the derivative of the cost (or error) with respect to the *N* linear layer of the graph as

$$e^{N-1} = \frac{\partial \mathcal{F}(\mathcal{D}^m; \Theta)}{\partial z_{\nu}^{N-1}},\tag{5.20}$$

then we can relate this to the cost derivative with respect to the preceding layer with

$$e^{N-1} = \left(\sum_{k'=1}^{K} e^{N} \frac{\partial z_{k'}^{N}}{\partial \tilde{z}_{i}^{N-1}}\right) \frac{\partial \tilde{z}_{j}^{N-1}}{\partial z_{i}^{N-1}} = \left(\sum_{k'=1}^{K} e_{k'}^{N} W_{k'j}^{N}\right) \cdot \tilde{z}_{j}^{N-1} \cdot (1 - \tilde{z}_{j}^{N-1}), \tag{5.21}$$

where we have used the fact that for sigmoid non-linearities

$$\frac{\partial \tilde{z}_j}{\partial z_i} = \tilde{z}_j \cdot (1 - \tilde{z}_j). \tag{5.22}$$

This formula can also be interpreted as traversing the graph in Fig. 5.3 backwards, multiplying at each node the backpropagated error by the derivative of its inputs and summing the errors propagated from different nodes. A more formal expression of the algorithm can be found in Algorithm 15.

#### Algorithm 15 Mini-batch SGD with Back-Propagation

- 1: **input:** Data  $\mathcal{D} = \{\mathcal{D}_1, \mathcal{D}_2, ..., \mathcal{D}_B\}$  split into B mini-batches of size M', MLP of N layers, with parameters  $\Theta = \{\mathbf{W}^1, \mathbf{b}^1, \cdots \mathbf{W}^N, \mathbf{b}^N\}$ , number of rounds T, learning rate  $\eta$
- 2: initialize parameters  $\Theta$  randomly
- 3: **for** t = 1 **to** T **do**
- 4: **for** b = 1 **to** B **do**
- 5: **for** m = 1 **to** M' **do**
- 6: Compute the forward pass for each of the M' examples in batch b; keep not only  $p(y^m|\mathbf{x}^m) \equiv \tilde{\mathbf{z}}^{m,N}$  but also all the intermediate non-linear outputs  $\tilde{\mathbf{z}}^{m,1} \cdots \tilde{\mathbf{z}}^{m,N}$ .
- 7: end for
- 8: **for** n = N **to** 1 **do**
- 9: **if** n==N **then**
- 10: **for** m = 1 **to** M' **do**
- 11: Initialize the error at last layer, for each example *m*. For the softmax with CE cost this is given by:

$$\mathbf{e}^{m,N} = (\mathbf{1}_{k(m)} - \tilde{\mathbf{z}}^{m,N}).$$

- 12: end for
- 13: else
- 14: **for** m = 1 **to** M' **do**
- 15: Backpropagate the error through the linear layer, for each example *m*:

$$\mathbf{e}^m = \left( (\mathbf{W}^{n+1})^T \mathbf{e}^{m,n+1} \right)$$

16: Backpropagate the error through the non-linearity, for the sigmoid this is:

$$\mathbf{e}^{m,n} = \mathbf{e}^m \odot \tilde{\mathbf{z}}^{m,n} \odot (1 - \tilde{\mathbf{z}}^{m,n}),$$

where  $\odot$  is the element-wise product and the 1 is replicated to match the size of  $\tilde{\mathbf{z}}^n$ .

- 17: end for
- 18: end if
- 19: Compute the gradients using the backpropagated errors and the inputs from the forward pass

$$abla_{\mathbf{W}^n} \mathfrak{F}(\mathfrak{D}; \Theta) = -\frac{1}{M'} \sum_{m=1}^{M'} \mathbf{e}^{m,n} \cdot \left( \tilde{\mathbf{z}}^{m,n-1} \right)^T,$$

$$\nabla_{\mathbf{b}^n} \mathfrak{F}(\mathfrak{D}; \Theta) = -\frac{1}{M'} \sum_{m=1}^{M'} \mathbf{e}^{m,n}.$$

20: Update the parameters

$$\mathbf{W}^n \leftarrow \mathbf{W}^n - \eta \nabla_{\mathbf{W}^n} \mathfrak{F},$$

$$\mathbf{b}^n \leftarrow \mathbf{b}^n - \eta \nabla_{\mathbf{b}^n} \mathfrak{F}.$$

- 21: end for
- 22: end for
- 23: end for

#### **Exercise 5.1** Start by loading Amazon sentiment corpus used in day 1

```
import numpy as np
import lxmls.readers.sentiment_reader as srs
scr = srs.SentimentCorpus("books")
train_x = scr.train_X.T
train_y = scr.train_y[:, 0]
test_x = scr.test_X.T
test_y = scr.test_y[:, 0]
```

Go to lxmls/deep\_learning/mlp.py:class NumpyMLP:def grads() and complete the code of the NumpyMLP class with the Backpropagation recursion that we just saw. Once you are done. Try different network geometries by increasing the number of layers and layer sizes e.g.

```
# Neural network modules
import lxmls.deep_learning.mlp as dl
import lxmls.deep_learning.sgd as sgd
# Model parameters
geometry = [train_x.shape[0], 20, 2]
actvfunc = ['sigmoid', 'softmax']
# Instantiate model
mlp = dl.NumpyMLP(geometry, actvfunc)
```

You can test the different hyperparameters

```
# Model parameters
n_iter = 5
bsize = 5
lrate = 0.01
# Train
sgd.SGD_train(mlp, n_iter, bsize=bsize, lrate=lrate, train_set=(train_x, train_y))
acc_train = sgd.class_acc(mlp.forward(train_x), train_y)[0]
acc_test = sgd.class_acc(mlp.forward(test_x), test_y)[0]
print "MLP (%s) Amazon Sentiment Accuracy train: %f test: %f" % (geometry, acc_train, acc_test)
```

### 5.4.4 Some final reflections on Backpropagation

If you are new to the neural network topic, this is about the most important piece of theory you should learn about deep learning. Here are some reflections that you should keep in mind.

- Thanks to the multi-layer structure and the chain rule, Backpropagation allows models that compose linear and non-linear functions with any depth (in principle<sup>5</sup>).
- The formulas are also valid for other cost functions and output layer non-linearities with minor modifications. It is only necessary to compute the equivalent of Eq. 5.13.
- The formulas are also valid for hidden non-linearities other than the sigmoid. Element-wise non-linear transformations still allow the simplification in Eq. 5.21. With little effort it is also possible to deal with other cases.
- However, there is an important limitation: unlike the log-linear models, the optimization problem is *non convex*. This removes some formal guarantees, most importantly we can get trapped in local minima during training.

<sup>&</sup>lt;sup>5</sup>Not exactly, since it is possible to run into numerical problems.

## 5.5 Deriving gradients and GPU code with Theano

#### 5.5.1 An Introduction to Theano

As you may have observed, the speed of SGD training for MLPs slows down considerably when we increase the number of layers. One reason for this is that the code that we use here is not very optimized. It is thought for you to learn the basic principles. Even if the code was more optimized, it would still be very slow for reasonable network sizes. The cost of computing each linear layer is proportional to the dimensionality of the previous and current layers, which in most cases will be rather large.

For this reason most deep learning applications use Graphics Processing Units (GPU) in their computations. This specialized hardware is normally used to accelerate computer graphics, but can also be used for general computation intensive tasks. However, we need to deal with specific interfaces and operations in order to use a GPU. This is where Theano comes in. Theano is a multidimensional symbolic expression python module with focus on neural networks. It will provide us with the following nice features:

- Symbolic expressions: Express the operations of the MLP (forward pass, cost) symbolically, as mathematical operations rather than explicit code
- Symbolic Differentiation: As a consequence of the previous feature, we can compute gradients of arbitrary mathematical functions automatically.
- GPU integration: The code will be ready to work on a GPU, provided that you have one and it is active within Theano. It will also be faster on normal CPUs since the symbolic operations are compiled to C code.
- Theano is focused on Deep Learning, with an active community and several tutorials easily available.

However, this does not come at a free price. There are a number of limitations

- Symbolic algebra is more difficult to debug, as we can not easily step in at each operation.
- Working with CPU and GPU code implies that we have to be more careful about the types of the variables.
- Theano tends to output long error messages. However, once you get used to it, error messages accurately point the source of the problem.
- Handling recurrent neural networks is much simpler than in Numpy but it still implies working with complicated constructs that are complicated to debug.

**Exercise 5.2** Get in contact with Theano. Learn the difference between a symbolic representation and a function. Start by implementing the first layer of our previous MLP in Numpy

```
# Numpy code
x = test_x # Test set
W1, b1 = mlp.params[:2] # Weights and bias of fist layer
z1 = np.dot(W1, x) + b1 # Linear transformation
tilde_z1 = 1/(1+np.exp(-z1)) # Non-linear transformation
```

Now we will implement this in Theano. We start by creating the variables over which we will produce the operations. For example the symbolic input is defined as

```
# Theano code.
# NOTE: We use undescore to denote symbolic equivalents to Numpy variables.
# This is no Python convention!.
import theano
import theano.tensor as T
_x = T.matrix('x')
```

Note that this variable does not have any particular value, nor a space reserved in memory for it. It contains just a symbolic definition of what the variable can contain. The particular values will be given when we use it to compile a function.

We could actually use the same definition format to define the weights and give their particular values as inputs to the compiled function. However, since we will be using a more complicated format in later exercises, we will use it here as well. The shared class allows to define variables that are shared across functions. They are also given a concrete value so that we do not need to give it for each function call. This format is therefore ideal for the weights of our network.

```
_W1 = theano.shared(value=W1, name='W1', borrow=True)
_b1 = theano.shared(value=b1, name='b1', borrow=True, broadcastable=(False, True))
```

**Important:** One of the main differences between Numpy and theano data is broadcast. In Numpy if we sum an array with shape (N, M) to one with shape (1, M), the second array will be copied N times to form a (N, M) matrix. This is known as broadcasting. In Theano this is not automatic. You need to specify broadcasting explicitly. This is important for example when using a bias, which will be copied to match the number of examples in the batch. In other cases, like when using variables for recurrent neural networks, broadcast has to be set to False. Broadcast is one of the typical source of errors when you start working with Theano. Keep this in mind.

Now lets describe the operations we want to do with the variables. Again only symbolically. This is done by replacing our usual operations by Theano symbolic ones when necessary e. g. the internal product dot() or the sigmoid. Some operations like e.g. + are automatically recognized by Theano (operator overloading).

```
_z1 = T.dot(_W1, _x) + _b1
_tilde_z1 = T.nnet.sigmoid(_z1)

# Keep in mind that naming variables is useful when debugging
_z1.name = 'z1'
_tilde_z1.name = 'tilde_z1'
```

When debugging the code it is often useful to print the graph of computations.

```
# Perceptron computation graph
theano.printing.debugprint(_tilde_z1)

sigmoid [@A] 'tilde_z1'
    |Elemwise{add,no_inplace} [@B] 'z1'
    |dot [@C] ''
    | |W1 [@D]
    | |x [@E]
    |b1 [@F]
```

It is important to keep in mind that, until this point, we do not have a function we can use to produce any practical input. In order to obtain this we have to compile this function by calling

```
layer1 = theano.function([_x], _tilde_z1)
```

Note the use of [] for the input variables, even if we just specify one variable. We can now do a test to compare the Numpy and Theano implementations and see that they give the same outputs.

```
# Check Numpy and Theano match
assert np.allclose(tilde_z1, layer1(x.astype(theano.config.floatX))), \
"Numpy and Theano Perceptrons are different"
```

One of the main sources of errors in Theano is the type of the arrays used. To ensure compatibility with the GPU check every array that you are going to work with and be sure it has the propper type To automatically switch between float64 (CPU) and float32 (GPU) types Theano provides the floatX type. Keep in mind not using these is the number one source of errors when you start with Theano.

```
train_x = train_x.astype(theano.config.floatX)
train_y = train_y.astype('int32')
```

## 5.5.2 Symbolic Forward Pass

In the previous section you have seen how to create symbolic Theano functions with shared parameters. You have thus all you need to implement the whole forward pass of a generic MLP in Theano.

**Exercise 5.3** Complete the method \_forward() inside of the lxmls/deep\_learning/mlp.py:class TheanoMLP. Note that this is called only once at the initialization of the class. To debug your implementation put a breakpoint at the \_\_init\_\_ function call. Hint: Note that this is very similar to NumpyMLP.forward(). You just need to keep track of the symbolic variable representing the output of the network after each layer is applied and compile the function at the end. After you are finished instantiate a Theano class and check that Numpy and Theano forward pass are the same.

```
mlp_a = dl.NumpyMLP(geometry, actvfunc)
mlp_b = dl.TheanoMLP(geometry, actvfunc)
```

To help debugging in Theano is sometimes useful to switch off the optimizer. This helps Theano point out which part of the Python code generated the error

```
theano.config.optimizer='None'
```

In general, the best way to debug Theano is to **compile often**. Try to compile the intermediate variables and perform the algebraic operations in Numpy. Once you are done, the following code should yield no error

```
assert np.allclose(mlp_a.forward(test_x), mlp_b.forward(test_x)), \
"ERROR: Numpy and Theano forward passes differ"
```

#### 5.5.3 Symbolic Differentiation

In the previous section we compiled the forward pass of a MLP. In this section we will do the same with the cost used for training. We will also derive the gradients although this will be trivial once we have the cost function compiled.

**Exercise 5.4** We first see an example that does not use any of the code in TheanoMLP but rather continues from what you wrote in Ex. 5.2. In this exercise you completed a sigmoid layer with Theano. To get some values for the weights we used the first layer of the network you trained in 5.2. Now we are going to use the second layer as well. This is thus assuming that your network in 5.2 has only two layers e.g. the recommended geometry (I, 20, 2). Make sure this is the case before starting this exercise.

For the sake of clarity, lets write here the part of Ex. 5.2 that we had completed

Now we just need to complete this with the second layer, using a softmax non-linearity

With this, we could compile a function to obtain the output of the network symb\_tilde\_z2 for a given input symb\_x. In this exercise we are however interested in obtaining the misclassification cost. This is given in Eq. 5.5. First we are going to need the symbolic variable for the correct output

```
_y = T.ivector('y')
```

The minus posterior probability of the class given the input is the same as selecting the k(m)-th softmax output, were k(m) is the index of the correct class for  $x^m$ . If we want to do this for a vector  $\mathbf{y}$  containing M different examples, we can write this as

```
_F = -T.mean(T.log(_tilde_z2[_y, T.arange(_y.shape[0])]))
```

Now obtaining a function that computes the gradient could not be easier.

```
_nabla_F = T.grad(_F, _W1)
nabla_F = theano.function([_x, _y], _nabla_F)
```

To finish this exercise have a look at the TheanoMLP class. As you may realise it just implements what is shown above for the generic case of N layers.

## 5.5.4 Symbolic mini-batch update

One important aspect to take into account is that to obtain a good speed up, the batch update operation must be carried out inside of Theano. To do this we may use the updates argument of theano.function

**Exercise 5.5** Define the updates list. This is a list where each element is a tuple of a parameter and the update rule to be applied that parameter. In this case we are defining the SGD update rule, but take into account that using more complex update rules like e.g. momentum or adam implies just replacing the last line of the following snippet.

```
mlp_c = dl.TheanoMLP(geometry, actvfunc)
    _x = T.matrix('x')
    _y = T.ivector('y')
    _F = mlp_c._cost(_x, _y)
# SGD update rule
updates = [(par, par - lrate*T.grad(_F, par)) for par in mlp_c.params]
```

This will return the cost of each batch and update the MLP parameters at the indent same time using updates

```
batch_up = theano.function([_x, _y], _F, updates=updates)
n_batch = int(np.ceil(float(train_x.shape[1])/bsize))
```

Once we have defined this, we can compare speed and accuracy of the Numpy and simple gradient versions using

```
import time
# Model
geometry = [train_x.shape[0], 20, 2]
actvfunc = ['sigmoid', 'softmax']
```

```
# Numpy MLP
        = dl.NumpyMLP(geometry, actvfunc)
mlp_a
         = time.clock()
init_t
sgd.SGD_train(mlp_a, n_iter, bsize=bsize, lrate=lrate, train_set=(train_x, train_y))
print "\nNumpy version took %2.2f sec" % (time.clock() - init_t)
acc_train = sgd.class_acc(mlp_a.forward(train_x), train_y)[0]
acc_test = sgd.class_acc(mlp_a.forward(test_x), test_y)[0]
print "Amazon Sentiment Accuracy train: %f test: %f\n" % (acc_train, acc_test)
# Theano grads
mlp_b = dl.TheanoMLP(geometry, actvfunc)
init_t = time.clock()
sgd.SGD_train(mlp_b, n_iter, bsize=bsize, lrate=lrate, train_set=(train_x, train_y))
print "\nCompiled gradient version took %2.2f sec" % (time.clock() - init_t)
acc_train = sgd.class_acc(mlp_b.forward(train_x), train_y)[0]
acc_test = sgd.class_acc(mlp_b.forward(test_x), test_y)[0]
print "Amazon Sentiment Accuracy train: %f test: %f\n" % (acc_train, acc_test)
# Theano batch update
init_t = time.clock()
sgd.SGD_train(mlp_c, n_iter, batch_up=batch_up, n_batch=n_batch, bsize=bsize,
             train_set=(train_x, train_y))
print "\nTheano compiled batch update version took %2.2f" % (time.clock() - init_t)
acc_train = sgd.class_acc(mlp_c.forward(train_x), train_y)[0]
acc_test = sgd.class_acc(mlp_c.forward(test_x), test_y)[0]
print "Amazon Sentiment Accuracy train: %f test: %f\n"%(acc_train,acc_test)
```

As you may observe, just computing the gradients with Theano may not lead to a decrease, but rather an increase in computing time. To maximally exploit the power of Theano, it is necessary to bundle both computations and data together using approaches like the compiled batch update.

## Day 6

# **Deep Learning II**

## 6.1 Today's assignment

Today's class will be focused on advanced deep learning concepts, mainly Recurrent Neural Networks (RNNs). In the first day we saw how the chain-rule allowed us to compute gradients for arbitrary computation graphs. Today we will see that we can still do this for more complex models like Recurrent Neural Networks (RNNs). In these models we will input data in different points of the graph, which will correspond to different time instants. The key factor to consider is that, for a fixed number of time steps, this is still a computation graph and all what we saw on the first day applies with no need for extra math.

If you managed to finish the previous day completely you should aim at finishing this as well. If you still have pending exercises from the first day e.g. the Theano part. It is recommended that you try to solve them first and then continue with this day. If you are not interested in Theano, the first exercise is also a pure Numpy implementation that you can use to learn about RNNs.

## 6.2 Recurrent Neural Networks: Backpropagation Through Time

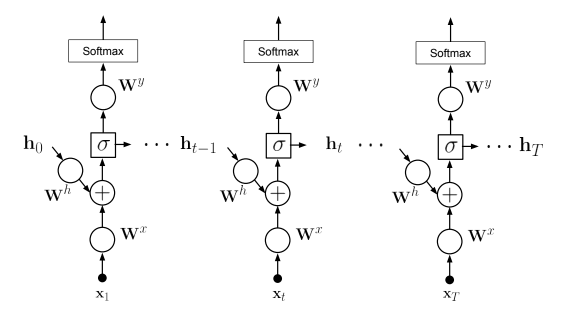


Figure 6.1: The simplest RNN can be seen as replicating a single hidden-layer FF network T times and passing the intermediate hidden variable  $\mathbf{h}_t$  across different steps. Note that all nodes operate over vector inputs e.g.,  $\mathbf{x}_t \in \mathbb{R}^I$ . Circles indicate matrix multiplications.

The previous day focused on Feed Forward (FF) networks. These networks are ill suited to learn variable length patterns since they only accept inputs of a fixed size. In order to learn sequences using neural networks, we need therefore to define some architecture that is able to process variable length inputs. Recurrent Neural Networks (RNNs) solve this problem by unfolding the computation graph in time. In other words, the network

is replicated as many times as it is necessary to cover the sequence to be modeled. In order to model the sequence one or more connections across different time instants are created. This allows the network to have a memory in time and thus capture complex patterns in sequences. In the simplest model, depicted in Fig. 6.1, and detailed in Algorithm 6.2, a RNN is created by replicating a single hidden-layer FF network T times and passing the intermediate hidden variable across different steps.

Since RNNs are computation graphs involving the same operations as FF networks, they can also be trained with backpropagation. However the error is propagated over the length of the entire sequence. This often leads to numerical problems know as *vanishing* and *exploding* gradients. A number of solutions are used to mitigate this issue. One simple, yet inelegant, method is clipping the gradients to a fixed threshold. Another solution is to resort to more complex RNN models that are able to better handle long range dependencies and are less sensitive to this phenomena. It is important to bear in mind, however, that all RNNs still use backpropagation as seen in the previous day, although it is often referred as *Backpropagation through time*.

#### Algorithm 16 Forward pass of a Recurrent Neural Network (RNN)

- 1: **input:** Initial parameters for an RNN Input  $\Theta = \{ \mathbf{W}^x \in \mathbb{R}^{H \times I}, \mathbf{W}^h \in \mathbb{R}^{H \times H}, \mathbf{W}^y \in \mathbb{R}^{K \times H} \}$  input, recurrent and output transformations respectively.
- 2: **input**: Input data matrix  $\mathbf{x} \in \mathbb{R}^{\hat{I} \times T}$  of size T. Initial recurrent variable  $\mathbf{h}_0$ .
- 3: **for** t = 1 **to** T 1 **do**
- 4: Apply linear transformation combining input and recurrent signals

$$z_{jt}^{h} = \sum_{i=1}^{I} W_{ji}^{x} x_{it} + \sum_{i=1}^{J} W_{jj'}^{h} h_{j't-1}$$

5: Apply non-linear transformation e.g. sigmoid (hereby denoted  $\sigma()$ )

$$h_{jt} = \sigma(z_{jt}^h) = \frac{1}{1 + \exp(-z_{jt}^h)}$$

- 6: end for
- 7: Apply final linear transformation to each of the recurrent variables  $\mathbf{h}_1 \cdots \mathbf{h}_T$

$$z_{kt}^y = \sum_{j=1}^J W_{kj}^y h_{jt}$$

8: Apply final non-linear transformation e.g. softmax

$$p(y_t = k | \mathbf{x}_{t:}) = \frac{\exp(z_{kt}^y)}{\sum_{k'=1}^K \exp(z_{k't}^y)}$$

**Exercise 6.1** Convince yourself that a RNN is just an FF unfolded in time. Run the NumpyRNN code. Set break-points and compare with what you learned about backpropagation in the previous day.

To work with RNNs we will use the Part-of-speech data-set seen in the sequence models day.

We will need to redo the indices of the data so that they are consecutive and cast all data to numpy arrays of int32 for compatibility with GPUs. This function will also add reverse indices to recover tag and word from its index word\_dict and tag\_dict

```
# Redo indices
train_seq, test_seq, dev_seq = pcc.compacify(train_seq, test_seq, dev_seq, theano=True)
# Get number of words and tags in the corpus
nr_words = len(train_seq.x_dict)
nr_tags = len(train_seq.y_dict)
```

Load and configure the NumpyRNN. Remember to user reload if you want to modify the code inside the rnns module

```
import lxmls.deep_learning.rnn as rnns
reload(rnns)
# RNN configuration
SEED = 1234  # Random seed to initialize weigths
emb_size = 50  # Size of word embeddings
hidden_size = 20  # size of hidden layer
# RNN
np_rnn = rnns.NumpyRNN(nr_words, emb_size, hidden_size, nr_tags, seed=SEED)
# Example sentence
x0 = train_seq[0].x
y0 = train_seq[0].y
```

Compute forward pass and gradients

```
# Forward pass
p_y, y_rnn, h, z1, x = np_rnn.forward(x0, all_outputs=True)
# Gradients
numpy_rnn_gradients = np_rnn.grads(x0, y0)
```

## 6.3 The Scan operation in Theano

Handling variable length computation graphs in an automatic fashion is not simple. Theano provides the *scan* function for this purpose. The scan function acts as a symbolic "for" loop. Since, unlike for normal python "for" loops, it is not possible to put a breakpoint in the scan loop, the design of graphs with scan has to be handled with care. Toolboxes like Keras conveniently abstract the user from such constructs. However, for complex designs it will be necessary to be able to use scan or equivalent functions.

**Exercise 6.2** Understand the basics of scan with these examples. Scan allows you to build computation graphs with a variable number of nodes. It acts as a python "for" loop but it is symbolic. The following example should help you understand the basic scan functionality. It generates a sequence for a given length. Run it and modify it. Try to arrive at an error and understand what happened.

```
import numpy as np
import theano
import theano.tensor as T
theano.config.optimizer='None'

def square(x):
    return x**2

# Python
def np_square_n_steps(nr_steps):
    out = []
    for n in np.arange(nr_steps):
        out.append(square(n))
    return np.array(out)
```

```
# Theano
nr_steps = T.lscalar('nr_steps')
h, _ = theano.scan(fn=square, sequences=T.arange(nr_steps))
th_square_n_steps = theano.function([nr_steps], h)

# Compare both
print np_square_n_steps(10)
print th_square_n_steps(10)
```

The following example should help you understand about matrix multiplications and passing values from one iteration to the other. At each step, we will multiply the output of the previous step by a matrix A. We start with an initial vector s0. The matrix and vector are random but normalized to result on a Markov chain (this is irrelevant for the use of scan).

```
# Configuration
nr_states = 3
nr_steps = 5

# Transition matrix
A = np.abs(np.random.randn(nr_states, nr_states))
A = A/A.sum(0, keepdims=True)
# Initial state
s0 = np.zeros(nr_states)
s0[0] = 1
```

```
# Numpy version
def np_markov_step(s_tm1):
    s_t = np.dot(s_tm1, A.T)
    return s_t

def np_markov_chain(nr_steps, A, s0):
    # Pre-allocate space
    s = np.zeros((nr_steps+1, nr_states))
    s[0, :] = s0
    for t in np.arange(nr_steps):
        s[t+1, :] = np_markov_step(s[t, :])
    return s

np_markov_chain(nr_steps, A, s0)
```

```
# Theano version
# Store variables as shared variables
th_A = theano.shared(A, name='A', borrow=True)
th_s0 = theano.shared(s0, name='s0', borrow=True)
# Symbolic variable for the number of steps
th_nr_steps = T.lscalar('nr_steps')
def th_markov_step(s_tm1):
    s_t = T.dot(s_t, th_A.T)
    # Remember to name variables
    s_t.name = 's_t'
    return s_t
s, _ = theano.scan(th_markov_step,
                   outputs_info=[dict(initial=th_s0)],
                   n_steps=th_nr_steps)
th\_markov\_chain = theano.function([th\_nr\_steps], T.concatenate((th\_s0[None, :], s), 0))
th_markov_chain(nr_steps)
```

## 6.4 A RNN in Theano for Part-of-Speech Tagging

**Exercise 6.3** Complete the theano code for a RNN inside lxmls/deep\_learning/rnn.py. Use exercise 6.1 for a numpy example and 6.2 to learn how to handle scan. Keep in mind that you only need to implement the forward pass! Theano will handle backpropagation for us.

```
# Instantiate the class
rnn = rnns.RNN(nr_words, emb_size, hidden_size, nr_tags, seed=SEED)
# Compile the forward pass function
x = T.ivector('x')
th_forward = theano.function([x], rnn._forward(x).T)
```

When working with theano, it is more difficult to localize the source of errors. It is therefore important to work step by step and test the code frequently. To debug we suggest to implement and compile the forward pass first. You can use this code for testing. If it raises no error you are good to go.

```
assert np.allclose(th_forward(x0), np_rnn.forward(x0)), \
"Numpy and Theano forward pass differ!"
```

Once you are confident the forward pass is working you can test the gradients

```
# Compile function returning the list of gradients
x = T.ivector('x')  # Input words
y = T.ivector('y')  # gold tags
p_y = rnn._forward(x)
cost = -T.mean(T.log(p_y)[T.arange(y.shape[0]), y])
grads_fun = theano.function([x, y], [T.grad(cost, par) for par in rnn.param])
```

```
# Compare numpy and theano gradients
theano_rnn_gradients = grads_fun(x0, y0)
for n in range(len(theano_rnn_gradients)):
    assert np.allclose(numpy_rnn_gradients[n], theano_rnn_gradients[n]), \
    "Numpy and Theano gradients differ in step n"
```

Finally, it is time to test our network in the task of POS tagging. Lets define the optimization parameters and compile batch update and prediction functions

```
# Parameters
lrate = 0.5  # Learning rate
n_iter = 5  # Number of iterations
# Get list of SGD batch update rule for each parameter
updates = [(par, par - lrate*T.grad(cost, par)) for par in rnn.param]
# compile
rnn_prediction = theano.function([x], T.argmax(p_y, 1))
rnn_batch_update = theano.function([x, y], cost, updates=updates)
```

To train the newtwork with SGD, you can use the following code for this purpose

```
nr_words = sum([len(seq.x) for seq in train_seq])
for i in range(n_iter):
    # Training
    cost = 0
    errors = 0
    for n, seq in enumerate(train_seq):
        cost += rnn_batch_update(seq.x, seq.y)
        errors += sum(rnn_prediction(seq.x) != seq.y)
    acc\_train = 100*(1-errors*1./nr\_words)
   print "Epoch %d: Train cost %2.2f Acc %2.2f %%" % (i+1, cost, acc_train),
    # Evaluation
    errors = 0
    for n, seq in enumerate(dev_seq):
       errors += sum(rnn_prediction(seq.x) != seq.y)
    acc_dev = 100*(1-errors*1./nr_words)
    print " Devel Acc %2.2f %%" % acc_dev
    sys.stdout.flush()
```

## 6.5 The Importance of Pre-training

One of the key insights that has played a role in the rise of deep learning in NLP tasks is the use of neural word-embeddings. These are just numeric representations of words that can be learned from unsupervised data using simple FF networks such as e.g. skip-grams.

Such representations can be plugged into supervised models such as the RNN that we just trained to initialize its initial layer. The use of pre-trained embeddings very often leads to important improvements in performance.

**Exercise 6.4** Test the effect of using pre-trained embeddings. Run the following code to download the embeddings. Reset the layer parameters and initialize the embedding layer with the pre-trained embeddings. Then run the training code from the last exercise.

```
# Embeddings Path
import lxmls.deep_learning.embeddings as emb
import os
reload(emb)
if not os.path.isfile(EMBEDDINGS):
    emb.download_embeddings('senna_50', "data/senna_50")
E = emb.extract_embeddings("data/senna_50", train_seq.x_dict)
# Reset model to remove the effect of training
rnn = rnns.reset_model(rnn, seed=SEED)
# Set the embedding layer to the pre-trained values
rnn.param[0].set_value(E.astype(theano.config.floatX))
# Now re-run SGD training
```

## 6.6 More Complex RNNs: LSTMs

As previously mentioned, the basic RNN that we saw until now suffers from a number of drawbacks like exploding or vanishing gradients, and the inability to model long term dependencies. These deficiencies are solved by more complex RNNs that include more complex internal logics, and are able to selectively forget, thus capturing better long term information. The so called Long Short-Term Memory (LSTM) and Gated Recurrent Unit (GRU) have replaced simple RNNs in most of modern models. However, they only imply more complex architectures. For an excellent visual explanation of the differences between simple RNNs, GRUs and LSTMs please visit

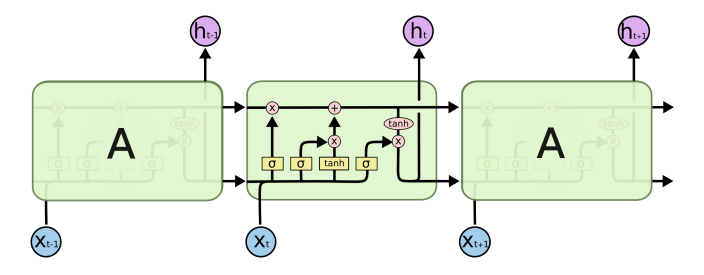


Figure 6.2: Detailed explanation of the LSTM internal logic, see source: http://colah.github.io/posts/2015-08-Understanding-LSTMs/.

http://colah.github.io/posts/2015-08-Understanding-LSTMs/

**Exercise 6.5** Convince yourself that LSTMs are just more complex RNNs. Run them, play around with the hyper parameters and compare the RNN and LSTM classes.

```
# Instantiate LSTM
lstm = rnns.LSTM(nr_words, emb_size, hidden_size, nr_tags)
# Compile prediction and bacth update functions
lstm_prediction = theano.function([x], T.argmax(lstm._forward(x), 1))
lstm_cost = -T.mean(T.log(lstm._forward(x))[T.arange(y.shape[0]), y])
# Get list of SGD batch update rule for each parameter
lstm_updates = [(par, par - lrate*T.grad(lstm_cost, par)) for par in lstm.param]
# compile
lstm_batch_update = theano.function([x, y], lstm_cost, updates=lstm_updates)
```

```
nr_words = sum([len(seq.x) for seq in train_seq])
for i in range(n_iter):
   # Training
   cost = 0
   errors = 0
   for n, seq in enumerate(train_seq):
       cost += lstm_batch_update(seq.x, seq.y)
       errors += sum(lstm_prediction(seq.x) != seq.y)
   acc_train = 100*(1-errors*1./nr_words)
   print "Epoch %d: Train cost %2.2f Acc %2.2f %%" % (i+1, cost, acc_train),
    # Evaluation:
   errors = 0
   for n, seq in enumerate(dev_seq):
        errors += sum(lstm_prediction(seq.x) != seq.y)
   acc_dev = 100*(1-errors*1./nr_words)
   print " Devel Acc %2.2f %%" % acc_dev
    sys.stdout.flush()
```

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