Introduction to neural network programming

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SciNet HPC Consortium

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Today's data, code and slides



You can get the slides for today's class at the SciNet Education web page.

https://support.scinet.utoronto.ca/education

Click on the link for the class, and look under "File Storage".

The code and data for today's class are stored on our git repository.

git clone https://gitrepos.scinet.utoronto.ca/public/nn.git

Note that the data may take a moment to download.

About this class



The purpose of this class is to introduce you to the basics of neural network programming in Python. Some notes about the class:

- The material is introductory. If you are already using neural networks it's likely you are already familiar with much of this content.
- We'll be using Python 2.7.X.
- We'll be using the Keras neural-network programming framework, with a *Theano* back end (though it would likely also work using *TensorFlow* too).
- You will need the usual machine learning packages: numpy, matplotlib, scikit-learn.
- You'll obviously also need Theano and Keras installed.

Ask questions!



Neural networks?



Let us begin at the beginning. What are neural networks? Neural networks, also called artificial neural networks,

are a computational model used in machine learning, ..., which is based on a large collection of connected simple units called artificial neurons, loosely analogous to axons in the biological brain. ... Such systems can be trained from examples, rather than explicitly programmed, and excel in areas where the solution or feature detection is difficult to express in a traditional computer program.

Wikipedia

If you're doing pattern recognition of any type, neural networks are worth considering.

Neural networks are commonplace



Neural networks are particularly good at detecting patterns, and for certain problems perform better than any other known class of algorithm. Neural networks are used for

- Image recognition, object detection.
- Natural language processing (voice recognition).
- Novelty detection (detection of outliers).
- Next-word predictions.
- Text sentiment analysis.
- System control (self-driving cars).
- Medical diagnosis.

Neural networks are finding their way into everything.



Neural networks, motivation



Consider the problem of hand-written digit recognition:



How would you go about writing a program which can tell you what digits are displayed?

- All the algorithms you might use to describe what a given number "looks like" are extremely difficult to implement in code. Where do you even start?
- And yet humans can easily tell what these digits are.
- Neural networks are based on a "biologically inspired" approach to solving such classification problems.
- This is one of the classic problems which have been solved using neural networks.

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Neural networks, the approach



Rather than focus on the details of what individual numbers look like, we will instead ignore those details altogether. We will use a different approach:

- Break the dataset of numbers into two or three groups: training, testing, and optionally validation.
- As with other supervised machine-learning algorithms, feed the training data to the neural network and train it to recognize one number from another.
- Rather than focus on details of the numbers, let the neural network figure out the details for itself.

This is the goal of the class. By the end of the class you should be acheiving greater than 98% classification accuracy from your neural network on the hand-written digits data set.

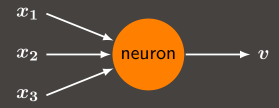
But first, let's start with the basics.



Neurons



Neural networks are built upon "neurons". This is just a fancy way of saying a "function that takes multiple inputs and returns a single output".



The function which the neuron implements is up to the programmer, but it must contain free parameters so that the network can be trained. These functions usually take the form

$$f(x_1,x_2,x_3) = f\left(\sum_{i=1}^3 w_i x_i + b
ight) = f\left(\mathbf{w}\cdot\mathbf{x} + b
ight)$$

Where \mathbf{w} are the 'weights' and \mathbf{b} is the 'bias'. These are the trainable parameters.

Neurons, continued



What might this look like in practise? Consider the data on the right.

| cloudy (c) | $hot\;(h)$ | windy (w) | beach? |
|--------------|------------|-------------|--------|
| 1 | 3 | 2 | 1 |
| 3 | 3 | 2 | 1 |
| 1 | 3 | 3 | 0 |
| 1 | 1 | 2 | 0 |
| 1 | 2 | 2 | 0 |
| 2 | 3 | 1 | 1 |
| 2 | 1 | 2 | 0 |

Our neuron's function might look like:

$$f(c,h,w) = egin{cases} 1 & w_1 imes c + w_2 imes h + w_3 imes w + b \geq 0 \ 0 & w_1 imes c + w_2 imes h + w_3 imes w + b < 0 \end{cases}$$

Where we must now optimize w_1, w_2, w_3, b to match the data.

Training our neuron



How do we optimize? We need to define some sort of "cost function" (sometimes called "loss" or "objective" function):

$$C = rac{1}{2} \sum_i \left(f(\mathbf{w} \cdot \mathbf{x}_i + b) - v_i
ight)^2$$

where v_i are the correct answers, based on the data, associated with each \mathbf{x}_i . Here we are using the "quadratic" cost function.

We then use an optimization algorithm to search for the minimum of C, given ${f x}$ and v.

Neurons, continued more



But there's a difficulty here:

- Having an output being just "1" or "0", as in our previous example, is difficult to deal with, since the function is discontinuous.
- Very small changes in the weights, Δw , can lead to discontinuous changes in the output. This makes using calculus difficult.
- Instead let's change the function of our neuron to use a "sigmoid function" (also called the "logistic function").

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

And so our neuron function becomes

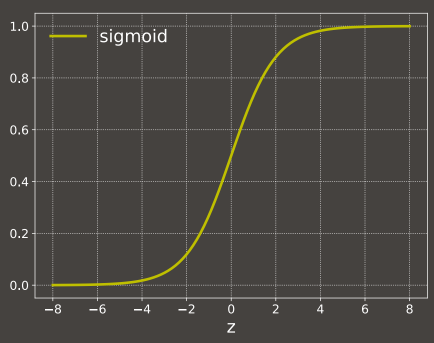
$$f(x_1, x_2, x_3) = f(\mathbf{w} \cdot \mathbf{x} + b) = \frac{1}{1 + e^{-(\mathbf{w} \cdot \mathbf{x} + b)}}$$

Where ${\bf w}$ are again the 'weights' and ${\bf b}$ is the 'bias'.

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Why the sigmoid function?





Because it ranges from 0 to 1 smoothly.

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Using sigmoid neurons



By using the sigmoid neurons, we can approximate the changes in the cost function, C:

$$\Delta Cpprox \sum_{j}rac{\partial C}{\partial w_{j}}\Delta w_{j}+rac{\partial C}{\partial b}\Delta b_{j}$$

meaning that the changes in the cost function are linear in changes to the weights and biases. The cost function changes 'smoothly' with changing parameters.

This allows us to use minimization algorithms to find the optimal values of w and b.

Today's data, code and slides



I'll be using ipython, running from the 'code' directory for the class.

```
ejspence@mycomp ~>
ejspence@mycomp ~> git clone https://gitrepos.scinet.utoronto.ca/public/nn.git
Cloning into 'nn'...
Checking connectivity... done.
ejspence@mycomp ~>
ejspence@mycomp ~> ls
nn
ejspence@mycomp ~> cd nn
ejspence@mycomp nn> ls
code data README.md
ejspence@mycomp nn> cd code
ejspence@mycomp nn/code> ipython --pylab
In [1]:
```

Be careful cutting and pasting code from PDFs!



Our first example



Use the sklearn.datasets.make_blobs command to generate some toy data.

```
In [1]: import sklearn.datasets as skd, sklearn.model_selection as skms

In [2]: import plotting_routines as pr

In [3]:

In [3]:

In [3]: n = 500

In [4]: pos, value = skd.make_blobs(n, centers = 2, center_box = (-3, 3))

In [5]:

In [5]:

In [5]: train_pos, test_pos, train_value, test_value = \
...: skms.train_test_split(pos, value, test_size = 0.2)

In [6]:

In [6]: train_pos.shape, train_value.shape, train_pos[0], train_value[0]

Out[6]: ((400,2), (400,), array([ 2.10552892, 1.31996395]), 1)

In [7]:

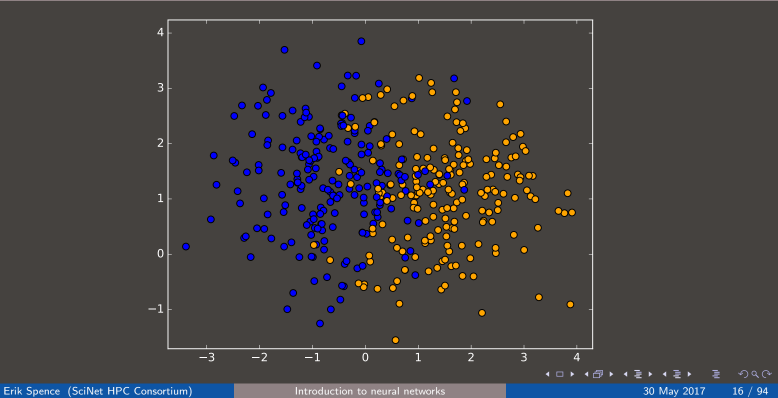
In [7]: pr.plot_dots(train_pos, train_value)

In [8]:
```

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Our data





The goal



What are we trying to accomplish?

- We want to create a neural network which, given a 2D position, can correctly classify the data point (0 or 1).
- To keep things simple, we will begin with a one-neuron network.
- We will use the sigmoid function for our neuron, with the 2 values of the position variable, (x_1, x_2) , as the inputs.
- We will use a technique called "Gradient Descent" to minimize the cost function, and find the best value of w_1 , w_2 and b.

$$f(x_1, x_2) = rac{1}{1 + e^{-(w_1 x_1 + w_2 x_2 + b)}} = rac{1}{1 + e^{-((w_1, w_2) \cdot (x_1, x_2) + b)}}$$

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Gradient Descent



So, again, how do we train our network? We are trying to minimize our cost function:

$$egin{array}{lcl} C = \sum_i C_i &=& rac{1}{2} \sum_i \left(f(\mathrm{w} \cdot \mathrm{x}_i + b) - v_i
ight)^2 \ &=& rac{1}{2} \sum_i \left(rac{1}{1 + e^{-\left((w_1, w_2) \cdot (x_1, x_2)_i + b
ight)}} - v_i
ight)^2 \end{array}$$

Where, again, the v_i are the correct classifications for each $(x_1,x_2)_i$.

The idea behind gradient descent is to calculate the gradient of our function, and then move "downhill".

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Gradient descent, continued



Suppose that our function has only one parameter.

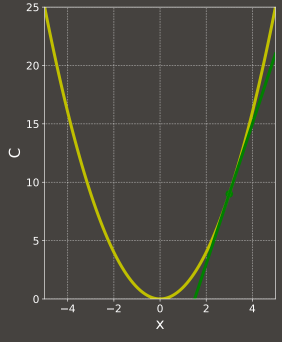
$$C = x^2$$

and we wish to minimize the function. Gradient descent says to move according to the formula:

$$x_{i+1} = x_i - \eta \frac{\partial C}{\partial x_i}$$

where η is called the step size. We then repeat until some stopping criterion is satisfied.

If we have multiple parameters, we step them all.



Gradient Descent, continued more



$$f_i = rac{1}{1 + e^{-ig((w_1, w_2) \cdot (x_1, x_2)_i + big)}}$$

$$C = \sum_i C_i = \sum_i rac{1}{2} \left(f_i - v_i
ight)^2$$

So to find the minimum of our cost function, we need

$$egin{aligned} rac{\partial C}{\partial w_1} &= \sum_i \left(f_i - v_i
ight) f_i \left(1 - f_i
ight) x_{1i} & rac{\partial C}{\partial w_2} &= \sum_i \left(f_i - v_i
ight) f_i \left(1 - f_i
ight) x_{2i} \ & rac{\partial C}{\partial b} &= \sum_i \left(f_i - v_i
ight) f_i \left(1 - f_i
ight) \end{aligned}$$

Gradient Descent, continued even more



So now

$$egin{split} w_1 & o w_1 - \eta \sum_i \left(f_i - v_i
ight) f_i \left(1 - f_i
ight) x_{1i} \ w_2 & o w_2 - \eta \sum_i \left(f_i - v_i
ight) f_i \left(1 - f_i
ight) x_{2i} \ b & o b - \eta \sum_i \left(f_i - v_i
ight) f_i \left(1 - f_i
ight) \ f_i &= rac{1}{1 + e^{-\left((w_1, w_2) \cdot (x_1, x_2)_i + b
ight)}} \end{split}$$

Now we're ready tackle the problem!



Training our neuron, code



```
# first network.py
import numpy as np, numpy.random as npr

def sigma(x, model):
    z = model['w1'] * x[:,0] + \
        model['w2'] * x[:,1] + \
        model['b']
    return 1. / (1. + np.exp(-z))

def build_model(x, v, eta, num_steps = 10000,
    print_best = True):

    model = {'w1': npr.random(), \
        'w2': npr.random(), \
        'b' : npr.random()}
    scale = 100. / float(len(v))
    best = 0.0
    f = sigma(x, model)
```

```
for i in xrange(0, num_steps):
    # Calculate derivatives.
    dCdw1 = sum((f - v) * f * (1 - f) * x[:,0])
    dCdw2 = sum((f - v) * f * (1 - f) * x[:,1])
    dCdb = sum((f - v) * f * (1 - f))

# Update parameters.
model['w1'] -= eta * dCdw1
model['w2'] -= eta * dCdw2
model['b'] -= eta * dCdb

f = sigma(x, model)
score = sum(np.round(f) == v) * scale
if (score > best):
    best, bestmodel = score, model.copy()

# Print out, if requested.
return bestmodel
```

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Our first example, continued



Assume that we've still got our data in memory.

```
In [8]: import first_network as fn

In [9]:

In [9]: model = fn.build_model(train_pos, train_value, eta = 5e-5)

Best by step 0: 48.0 %

Best by step 1000: 87.2 %

Best by step 2000: 87.2 %

Best by step 7000: 87.2 %

Best by step 8000: 87.2 %

Best by step 9000: 87.2 %

Our best model gets 87.2 percent correct!

In [10]:

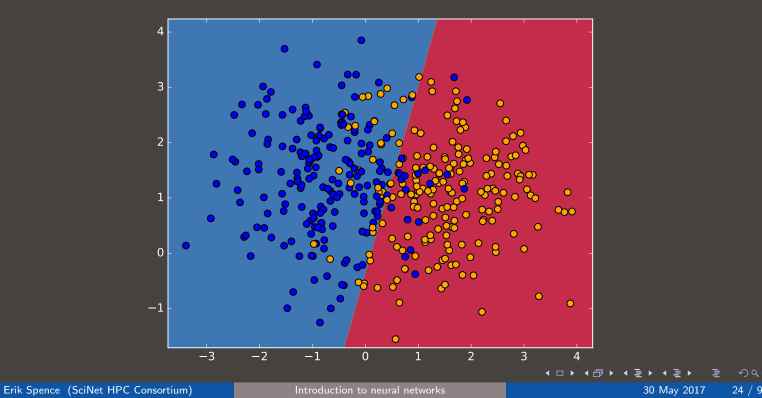
In [10]: pr.plot_decision_boundary(train_pos, train_value, model, fn.predict)

In [11]:
```

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Our fit





Our first example, test data



```
In [11]:
In [11]: import numpy as np
In [12]:
In [12]: f = fn.predict(test_pos, model)
In [13]:
In [13]:
In [13]: sum(np.round(f) == test_value) / float(len(test_value))
Out[13]: 0.88026315789473684
In [14]:
```

88%!

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Some notes on our first example



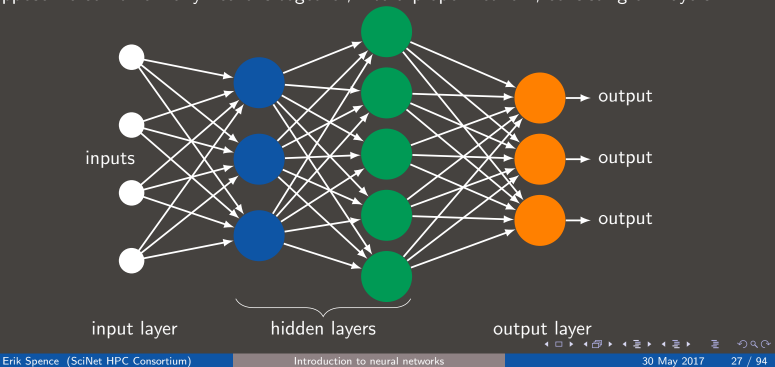
A few observations about our first example.

- Our model only has three free parameters, w_1 , w_2 , b. As such, it is only capable of a linear fit, for data with two independent variables.
- The reason why the fit worked as well as it did is because I separated the data enough that a linear split would be a reasonable thing to do.
- Depending on how well your data were split, your result may not have been as good.
- If we want to be able to categorize more-complex data, such as hand-written digits, we're going to need a more-complex approach.

Neural networks



Suppose we combine many neurons together, into a proper network, consisting of "layers".



Some notes about neural networks



Some details about the graphic on the previous slide:

- The input neurons do not contain any functions. They merely represent the input data being fed into the network.
- ullet Each neuron in the 'hidden' layers and the output layer all contain functions with their own free parameters, ullet and ullet.
- Each neuron outputs a single value. This output is passed to all of the neurons in the subsequent layer. This type of layer is known as a "fully-connected", or "dense", layer.
- The number of free parameters in the neurons in any given layer depends upon the number of neurons in the previous layer.
- The output from the output layer is aggregated into the desired form to calculate the cost function.

Seriously?



You might legitimately wonder why on Earth we would think this would lead anywhere.

- As it happens, this topology is similar to simple biological neural networks.
- Each layer takes the output of the previous layer as its input.
- Each layer makes "decisions" about the information that it receives.
- In this way the later layers are able to make more complex and abstract decisions than the earlier layers.
- A many-layered network can potentially make sophisticated decisions.

However, there are subtleties in training such a network.

Training a neural network



How do we train such a network?

- Suppose that we decide to try to use gradient descent to train the network from three slides ago.
- Each of the neurons has its own set of free parameters, w and b. There are lots of free parameters!
- To update the parameters we need to calculate every $\frac{\partial C}{\partial w_i}$ and $\frac{\partial C}{\partial b}$ for every neuron!
- But how do we calculate those derivatives, especially for the parameters associated with the neurons that are several layers away from the output?

Actually, as it happens, this is a solved problem.



The backpropagation algorithm



To find the gradients of the cost function with respect to the weights and biases we use the "backpropagation algorithm". First let's go over some terminology.

Let the input layer be the zeroth layer. If $x \in \mathbb{R}^{500 \times 2}$ is the input data, then let $a^1 \in \mathbb{R}^{k \times 500}$ be the vector of outputs from the k neurons in the first (hidden) layer:

$$\mathbf{z}^{1} = \mathbf{w}^{1} \mathbf{x}^{T} + \mathbf{b}^{1}$$
 $\mathbf{a}^{1} = \sigma (\mathbf{z}^{1})$

with $\mathbf{w^1} \in \mathbb{R}^{k \times 2}$ and $\mathbf{b^1} \in \mathbb{R}^{k \times 1}$. Similarly,

$$\mathbf{z}^{\ell} = \mathbf{w}^{\ell} \mathbf{a}^{\ell-1} + \mathbf{b}^{\ell} \qquad \mathbf{a}^{\ell} = \sigma\left(\mathbf{z}^{\ell}\right)$$

with $\mathbf{w}^{\ell} \in \mathbb{R}^{m_{\ell} \times m_{(\ell-1)}}$, $\mathbf{b}^{\ell} \in \mathbb{R}^{m_{\ell} \times 1}$, $\mathbf{a}^{\ell} \in \mathbb{R}^{m_{\ell} \times 500}$, where m_{ℓ} is the number of neurons in the ℓ th layer.

The backpropagation algorithm, continued



$$\delta^{M} = rac{\partial C}{\partial z^{M}} =
abla_{\mathbf{a}^{M}} C \circ \sigma' \left(\mathbf{z}^{M}
ight)$$

is the "error" in the last $(M \mathsf{th})$ layer. Recall that

$$C = rac{1}{2} \sum_i \left(a_i^M - v_i
ight)^2$$
 ,

and thus

$$\mathbf{\delta}^{M} = (\mathbf{a}^{M} - \mathbf{v}) \circ \mathbf{\sigma'} (\mathbf{z}^{M}).$$

We now claim that

$$\delta^{\ell} = \left[\left(\mathbf{w}^{\ell+1}\right)^T \delta^{\ell+1} \right] \circ \sigma' \left(z^{\ell} \right)$$

Some algebra reveals that

$$rac{\partial C}{\partial b^\ell} = \delta^\ell$$

and that

$$\frac{\partial C}{\partial w^{\ell}} = \mathbf{a}^{\ell-1} \delta^{\ell}$$

The derivation of these quantities is beyond the scope of this class, but it's not too difficult.

Note that we are now using a_i^{M} as the output of our network.

Our second example

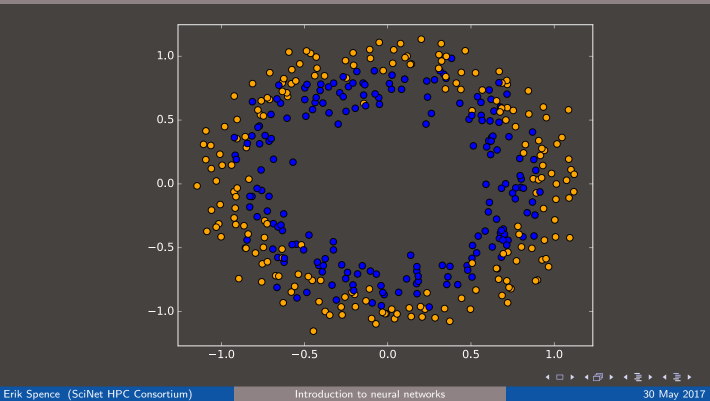


Use the sklearn.datasets.circles command to generate some toy data.

```
In [14]:
In [14]: n = 500
In [15]:
In [15]: pos, value = skd.make_circles(n, noise = 0.1)
In [16]:
In [16]: train_pos, test_pos, train_value, test_value = \
...: skms.train_test_split(pos, value, test_size = 0.2)
In [17]:
In [17]: pr.plot_dots(train_pos, train_value)
In [18]:
```

Our second example, data





The goal



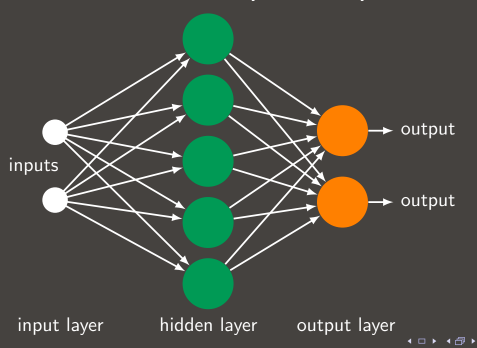
What are we trying to accomplish?

- Just like with our first example, we want to create a network which, given a 2D position, can correctly classify the data point (0 or 1).
- However, obviously a linear fit will not work in this case.
- This time we will create a network with three layers, an input layer, one hidden layer, and an output layer.
- We will use the sigmoid function for all neurons, with the 2 values of the position variable, (x_1, x_2) , as the inputs to the hidden layer, and the outputs of the hidden layer as inputs to the output layer.
- Once again, we'll use gradient descent to minimize the cost function, to find the best values of our weights and biases.

Our neural network



Note that the number of neurons in the hidden layer is arbitrary.



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Training our network, code



```
# second network.py
import numpy as np
import numpy.random as npr

# Sigmoid function.
def sigma(z):
    return 1. / (1. + np.exp(-z))

# Sigmoid prime function.
def sigmaprime(z):
    return sigma(z) * (1. - sigma(z))

# Returns just the predicted values.
def predict(x, model):
    _, _, _, a2 = forward(x, model)
    return np.argmax(a2, axis = 0)
```

```
# second network.py, continued

# Predict the output value, given the model
# and the positions.
def forward(x, model):

# Hidden layer.

z1 = model['w1'].dot(x.T) + model['b1']
a1 = sigma(z1)

# Output layer.
z2 = model['w2'].dot(a1) + model['b2']
a2 = sigma(z2)

return z1, z2, a1, a2
```

Training our network, code, continued



```
def build_model(num_nodes, x, v, eta, output_dim, num_steps = 10000, print_best = True, lam = 0):
 input_dim = np.shape(x)[1]
 model = {'w1': npr.randn(num_nodes, input_dim), 'b1': np.zeros([num_nodes, 1]), \
          'w2': npr.randn(output_dim, num_nodes), 'b2': np.zeros([output_dim, 1])}
 z1, _, a1, a2 = forward(x, model)
 for i in xrange(0, num_steps):
                    delta2[v, range(len(v))] -= 1 # (a^M - v)
   delta2 = a2;
   delta1 = (model['w2'].T).dot(delta2) * sigmaprime(z1)
   dCdb2 = np.sum(delta2, axis = 1, keepdims = True)
   dCdb1 = np.sum(delta1, axis = 1, keepdims = True)
   dCdw2 = delta2.dot(a1.T);
                                 dCdw1 = delta1.dot(x)
   model['w1'] -= eta * (lam * model['w1'] + dCdw1);
                                                         model['b1'] -= eta * dCdb1
   model['w2'] = eta * (lam * model['w2'] + dCdw2);
                                                         model['b2'] -= eta * dCdb2
```

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Our second example, continued



Once again, assume that we've still got our data in memory.

```
In [18]: import second_network as sn

In [19]:

In [19]: model = sn.build_model(10, train_pos, train_value, eta = 0.01, output_dim = 2)

Best by step 0: 51.2 %

Best by step 1000: 83.8 %

Best by step 2000: 84.8 %

:

Best by step 7000: 85.5 %

Best by step 8000: 85.8 %

Best by step 9000: 86.0 %

Our best model gets 86.0 percent correct!

In [20]:

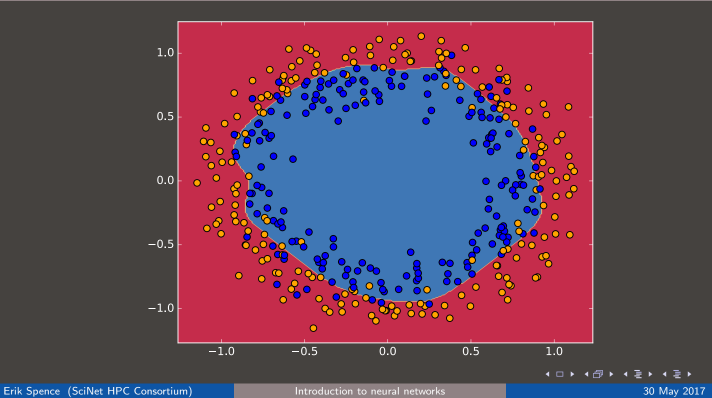
In [20]: pr.plot_decision_boundary(train_pos, train_value, model, sn.predict)

In [21]:
```

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Our fit





Our second example, test data



73%!



Some notes on our second example



A few observations about our second example.

- The choice of η is by trial-and-error.
- Our model has as many free parameters as you like, depending on the number of nodes you use. As such, it is capable of getting an extremely good fit.
- It is not uncommon for the number of parameters in the network to greatly exceed the number of observations. Your machine-learning instincts should be warning you: this situation is ripe for over-fitting.
- Nonetheless, there are techniques that are used to improve the generalization of the model. We'll visit these later in the class.



Handwritten digits



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One of the classic datasets on which to test neural-network techniques is the MNIST dataset.

- A database of handwritten digits, compiled by NIST.
- Contains 60000 training, and 10000 test examples.
- The training digits were written by 250 different people; the test data by 250 different people.
- The digits have been size-normalized and centred.
- Each image is grey scale, 28 x 28 pixels.

We can use our existing code to classify these digits.



Our network



How would we design a network to analyse this data?

- Each image is $28 \times 28 = 784$ pixels. Let the input layer consist of 784 input nodes. Each entry will consist of the grey value for that pixel.
- The output will consist of a one-hot-encoding of the networks analysis of the input data. This means that, if the input image depicts a '7', the output vector should be [0,0,0,0,0,0,0,1,0,0].
- Thus, let there be 10 output nodes, one for each possible digit.
- To start, let's just use a single hidden layer.
- As it happens, the code which we used in the second example can solve this problem.

Hand written digits, continued



The code for reading the data is contained in mnist_loader.py.

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Handwritten digits, test data



82%! Not great. Clearly we have some over-fitting going on. How do we deal with this?

Over-fitting



Over-fitting occurs when a model is excessively fit to noise in the training data, resulting in a model which does not generalize well to the test data.

This can be a serious issue with neural networks. How do we deal with this?

- More data! Either real (original), or artificially created.
- Regularization.
- Dropout.

The first is self-explanatory. We'll go over the later two cases.

Regularization



Regularization is an *ad hoc* technique by which parameters in a fit are penalized to prevent individual parameters from becoming excessively important to the fit.

- This shows up all the time in contexts where over-fitting is to be expected, or the problem is ill-posed (inverse problems).
- Goes by many names: Tikhonov regularization, ridge regression...
- This usually manifests itself as a modification to the cost function, though there are other forms as well.

$$C = rac{1}{2}\sum_i \left(a_i^M - v_i
ight)^2 + rac{\lambda}{2}\sum_j w_j^2$$

where λ is the "regularization parameter", and the sum over j is over all weights in the network.

Regularization, continued



If we call our previous cost function C_0 , then the new cost function is

$$C = C_0 + rac{\lambda}{2} \sum_j w_j^2 \, .$$

The addition of another term in the cost function changes the derivatives used to perform gradient descent.

$$rac{\partial C}{\partial w_k}
ightarrow rac{\partial C_0}{\partial w_k} + \lambda w_k$$

where the derivatives of C_0 are calculated the usual way, using backpropagation. Using gradient descent leads to

$$egin{array}{lll} w_k &
ightarrow & w_k - \eta rac{\partial C_0}{\partial w_k} - \eta \lambda w_k \ &
ightarrow & (1 - \eta \lambda) \, w_k - \eta rac{\partial C_0}{\partial w_k} \end{array}$$

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Training our network, code, continued



```
def build_model(num_nodes, x, v, eta, output_dim, num_steps = 10000, print_best = True, lam = 0):
 input_dim = np.shape(x)[1]
 model = {'w1': npr.randn(num_nodes, input_dim), 'b1': np.zeros([num_nodes, 1]), \
          'w2': npr.randn(output_dim, num_nodes), 'b2': np.zeros([output_dim, 1])}
 z1, _, a1, a2 = forward(x, model)
 for i in xrange(0, num_steps):
                    delta2[v, range(len(v))] -= 1 # (a^M - v)
   delta2 = a2;
   delta1 = (model['w2'].T).dot(delta2) * sigmaprime(z1)
   dCdb2 = np.sum(delta2, axis = 1, keepdims = True)
   dCdb1 = np.sum(delta1, axis = 1, keepdims = True)
   dCdw2 = delta2.dot(a1.T);
                                 dCdw1 = delta1.dot(x)
   model['w1'] -= eta * (lam * model['w1'] + dCdw1);
                                                         model['b1'] -= eta * dCdb1
   model['w2'] = eta * (lam * model['w2'] + dCdw2);
                                                         model['b2'] -= eta * dCdb2
```

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Hand written digits, continued more



The code for the regularization is already part of second_network.py.

Regularization, continued more



Why the improvement in the fitting of the test data?

- The regularization keeps the network from depending on any one particular parameter, or set of parameters, too much.
- This results in the network not focusing too much on any given feature, resulting in better generalization to the test data.
- ullet Trial-and-error is needed to determine a good value of $oldsymbol{\lambda}.$

Regularization isn't used as often as it used to be. Dropout appears to much more popular an approach.

Dropout

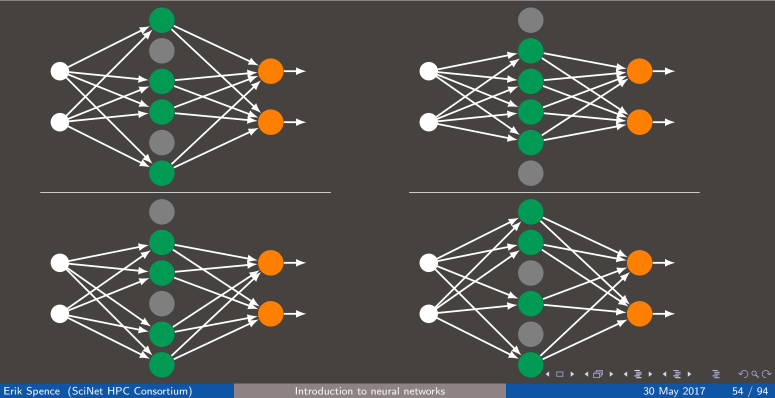


Dropout is a uniquely-neural-network technique to prevent over-fitting.

- The principle is simple: randomly "drop out" neurons from the network, at each iteration of the minimization algorithm.
- Like regularization, this results in the network not putting too much importance on any given weight, since the weights keep randomly disappearing from the network.
- It can be thought of as averaging over several different-but-similar neural networks.
- Different fractions of different layers can be specified for dropout. Again, trial-and-error is needed to determine a good fraction.
- Obviously, the output from the dropout layers must be scaled.
- We're not going to run an example of this.

Dropout, visualized





Neural network frameworks



Now that we have a sense of how neural networks work, we're ready to switch gears and use a 'framework'. Why would you do that?

- Coding your own networks from scratch can be a bit of work. (Though it's easier and cleaner if you use classes).
- Neural network (NN) frameworks have been specifically designed to solve NN problems.
- Python, of course, is not a high-performance language.
- The NN frameworks which have been developed are compiled before being used, thus being much faster than interpreted Python.
- The NN frameworks are also designed to use GPUs, which make things significantly faster than just using CPUs.

The training of neural networks is particularly well suited to solution using GPUs.



Theano



Let's review several of the more-popular NN frameworks out there. First is the Theano programming package.

- Theano is not strictly a NN framework. It was designed to handle multi-dimensional array manipulation.
- Developed by the LISA/MILA lab at the Université de Montréal.
- Written in Python; uses a numpy-like syntax, but generates C code which is compiled before being used.
- Supports symbolic differentiation.
- Is the grand-daddy of NN programming approaches.

Though commonly used in NN programming, it's been falling out of favour due to its long compile times (for some classes of problems) compared to other frameworks.

TensorFlow



TensorFlow is Google's NN framework.

- Released as open source in November 2015.
- The second-generation NN framework developed internally at Google.
- Allegedly capable of running on multiple cores.
- Provides APIs for Python, C++, Java and other languages.
- Can be quite slow.

This framework continues to grow in popularity.



Caffe



Caffe was developed as part of a Ph.D. project at U.C. Berkeley.

- Stands for Convolutional Architecture for Fast Feature Embedding.
- Currently maintained by the Berkeley Vision and Learning Center.
- Written in C++, but has Python and MATLAB interfaces.
- Particularly strong in the image-recognition and analysis area.
- Weak if you're not doing convolution networks (we'll get to this shortly), meaning you're studying text, sound, or time series.
- Only a few native input formats, and only one native output format, HDF5 (though you can get around this using the Python interface).

This is a commonly-encountered framework where ever images are being used.

Torch



Another framework used for neural networks is Torch.

- First released in 2002. Quite mature at this point.
- Written in Lua. Never heard of Lua? Welcome to the club.
- Pytorch was release by Facebook in January.
- Like Theano, Torch is more flexible than just NN. It is more of a generic scientific computing framework.
- Unlike Theano, Torch does not support symbolic differentiation.
- Very strong on GPUs.
- Very fast. Often the fastest depending on the problem being considered.
- Limited visualization capabilities.
- Used and maintained by Facebook, Twitter and other high-profile companies.

Keras



Because Theano is not a native NN framework, we will use Keras on top of Theano.

- Keras is a NN framework.
- It can run on top of either Theano or TensorFlow.
- Because it's a proper framework, all of the NN goodies you need are already built into it.
- Designed for fast development of networks.
- Is compatible with both Python 2 and 3.
- By default it will try to use TensorFlow on the back end; you may need to switch it.
 - Edit your \$HOME/.keras/keras.json file.
 - On Windows, edit your %USERPROFILE%/.keras/keras.json file.
 - You can also set the KERAS_BACKEND environment variable.

Prepping the data



Theano needs the data in a specific format:

- If the input data is 2D, it must have a 'depth' added, to make it 3D, even if the depth is 1.
- If the input data is 1D no depth is needed.
- The labels must be changed to a categorical format (one-hot encoding).

```
In [37]:
In [37]: import keras.utils as ku
Using Theano backend.
In [38]:
In [38]: val_in2 = val_in[0:200,:]
In [39]: val_out2 = val_out[0:200]
In [40]:
In [40]:
In [40]: train_out2 = ku.to_categorical(train_out2, 10)
In [41]: val_out2 = ku.to_categorical(val_out2, 10)
In [42]:
In [42]: train_out = ku.to_categorical(train_out, 10)
In [43]: val_out = ku.to_categorical(val_out, 10)
In [44]:
```

Our network using Keras



Let us re-implement our second network using Keras.

- A "Sequential" model is linear, meaning does not loop back on itself.
- A "Dense" ("fully-connected") layer is the regular layer we've been using.
- Use "input_dim" in the first layer to indicate the shape of the incoming data.
- The "activation" is the output function of the neuron.

```
In [44]: import keras.optimizers as ko
In [45]: import keras.models as km
In [46]: import keras.layers as kl
In [47]:
In [47]: numnodes = 30
In [48]: model = km.Sequential()
In [49]:
In [49]: model.add(kl.Dense(numnodes,
             input_dim = 784,
             activation = 'sigmoid'))
In [50]: model.add(kl.Dense(10,
             activation = 'sigmoid'))
In [51]:
In [51]: model.output_shape
Out[51]: (None, 10)
In [52]:
```

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Our network using Keras, continued



```
In [52]:
In [53]: model.compile(optimizer = ko.SGD(1r = 1), metrics = ['accuracy'],
...: loss = "mean_squared_error")

In [54]:
In [54]:
In [54]: fit = model.fit(train_in2, train_out2, epochs = 100, batch_size = 10,
...: validation_data = (val_in2, val_out2), verbose = 2)

Train on 1000 samples, validate on 200 samples

Epoch 1/100
0s - loss: 0.0742 - acc: 0.4610 - val_loss: 0.0759 - val_acc: 0.4100

Epoch 2/100
0s - loss: 0.0709 - acc: 0.5250 - val_loss: 0.0732 - val_acc: 0.4150

:

Epoch 100/100
0s - loss: 0.0107 - acc: 0.9600 - val_loss: 0.0234 - val_acc: 0.8750
In [55]:
```

Note the over-fitting: training versus validation data.



About that optimization flag



The optimization flag was set to "opt.SGD".

- This stands for "Stochastic Gradient Descent".
- This is similar to regular gradient descent that we used previously.
 - Regular gradient descent is ridiculously slow on large amounts of data.
 - To speed things up, SGD uses a randomly-selected subset of the data (a "batch") to update the coefficients.
 - This is repeated many times, using different batches, until all of the data has been used. This is called an "epoch".
- In practise, regular gradient descent is never used, stochastic gradient descent is used instead, since it's so much faster.
- The only real advantage of regular gradient descent is that it's easier to code, which is why I used it previously.



Our network using Keras, continued



Some notes about the compilation of the model.

- We must specify the loss function with the "loss" argument.
- We must specify the optimization algorithm, using the "optimizer" flag.
- The optimizer can be specified explicitly, as in this example, or just put 'sgd', and get the default values.
- I specified it explicitly so that I could specify the value of η (using 'Ir', the 'learning rate').
- The 'metrics' argument is optional, but is needed if you want the accuracy to be printed.

Our network using Keras, continued more



Now check against the test data.

Again we see the over-fitting rearing its head.

We can do better!

```
In [55]:
In [56]: test_out2 = ku.to_categorical(test_out2, 10)
In [57]:
In [57]: score = model.evaluate(test_in2, test_out2)
In [58]:
In [58]: score
Out[58]: [0.020172787383198738, 0.87]
In [59]:
```

The next steps



We can do better. What's the plan? There are a few simple approaches:

- Use more data.
- Change the activation function.
- Change the cost function.
- Change the optimization algorithm.
- Change the way things are initialized.
- Add regularization, to try to deal with the over-fitting.

We'll try some of these next, but there are some not-so-simple approaches:

• Completely overhaul the network strategy.

We'll take a look at this as well.



The next steps, an aside



There are alot of things we can tweak to make the network do better on the testing data. How do we know what to do?

- In many ways, implementing a network is an art.
- Certain forms and functions and parameters are known to lead to certain types of behaviour, and thus are used in certain situations.
- But choosing the correct values of parameters is often a matter of trial-and-error.
- And choosing the correct activation functions, number of nodes, is also trial-and-error.

Practise is often needed to know how to approach various types of problems. Consult your colleagues, and the literature.



Other activation functions: softplus



Two commonly-used functions:

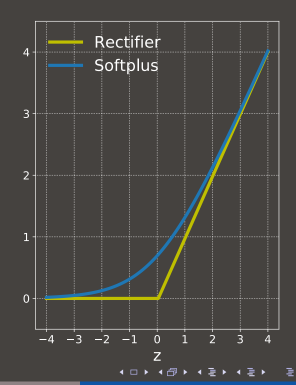
• Rectifier (also called Rectifier Linear Units, or RLUs):

$$f(z) = \max(0, z).$$

• Softplus:

$$f(z) = \ln(1 + e^z).$$

- Good: doesn't suffer from the vanishing-gradient problem.
- Bad: unbounded, could blow up.



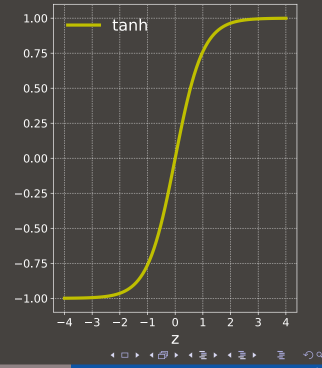
Other activation functions: tanh



Another commonly-used activation function is tanh:

$$f(z) = \tanh(z)$$
.

- Good: stronger gradients than sigmoid, faster learning rate, doesn't suffer from the vanishing-gradient problem.
- Good: because the function is anti-symmetric about zero. This also results in faster learning, at least for deeper networks.



Other activation functions: softmax



One of the more-commonly used output-layer activation functions is the softmax function:

$$s(\mathbf{z}_j) = rac{e^{\mathbf{z}_j}}{\sum\limits_{k=1}^N e^{\mathbf{z}_k}},$$

where N is the number of output neurons. The advantage of this function is that it converts the output to a probability.

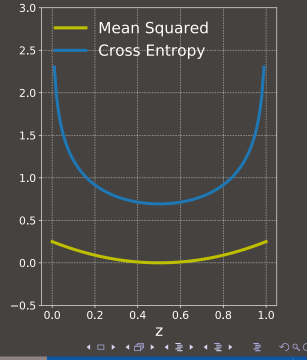
Other cost functions: cross entropy



Probably the most-commonly used cost function is cross entropy:

$$C = -rac{1}{n} \sum_{i}^{n} \left[v_i \log(a_i) + (1-v_i) \log(1-a_i)
ight]$$

- Good: the gradient of cross entropy is directly proportional to the error; learning is faster than with mean squared error.
- Because $0 \le a \le 1$, this is usually used with softmax output.
- ullet v=0.5 in the example on the right.



Other optimization algorithms



There are many algorithms used to minimize the cost function.

- Gradient Descent, and its variations (RMSprop, Adam).
- Newton's method, uses the second derivatives of the cost function. Its variations ("Quasi-Newton") are gaining in popularity, especially DFP, and L-BFGS.
- Conjugate Gradient, which is like a combination of Gradient Descent and Newton's method.
- Levenberg-Marquardt (damped-least-squares) algorithm. Only works on squared cost functions (doesn't work on cross entropy).

If you find that your network won't train on a given optimization algorithm, it may be worth the effort to try a different one.



Our Keras network revisited



```
In [59]: model2 = km.Sequential()
In [60]: model2.add(kl.Dense(numnodes, input_dim = 784, activation = "tanh"))
In [61]: model2.add(kl.Dense(10, activation = "softmax"))
In [62]: model2.compile(loss = "categorical_crossentropy", optimizer = "sgd",
              metrics = ['accuracy'])
In [63]:
In [63]: fit = model2.fit(train_in, train_out, epochs = 100, batch_size = 20,
            validation_data = (val_in, val_out), verbose = 2)
Train on 50000 samples, validate on 10000 samples
Epoch 1/100
2s - loss: 0.6853 - acc: 0.8261 - val_loss: 0.3783 - val_acc: 0.9021
Epoch 2/100
2s - loss: 0.3661 - acc: 0.9002 - val_loss: 0.3049 - val_acc: 0.9163
Epoch 100/100
2s - loss: 0.0543 - acc: 0.9860 - val_loss: 0.1342 - val_acc: 0.9633
In [64]:
```

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Our Keras network revisited, continued



Now check against the test data.

96%! Better!

This is about the best we can do with this approach, at least without using regularization. To push this even further, we need to change our network's architecture.

```
In [64]:
In [64]: test_out = ku.to_categorical(test_out, 10)
In [65]:
In [65]: score = model2.evaluate(test_in, test_out)
In [66]:
In [66]: score
Out[66]: [0.12195164765194058, 0.96540000000000004]
In [67]:
```

Other topics



This workshop is short. There is not time to cover every topic. Some topics you should look into further, if you're going to use NNs in your research:

- preprocessing data: remove unnecessary degrees of freedom, scale and centre the data.
- parameter initialization: how the weights and biases are initialized matters.
- activation functions: there are several activation functions which are used in specific areas of neural networks. Learn which ones apply to your field.
- more cost functions: there are other cost functions which are used in specific fields.
- training failures: the disappearing gradient problem, the exploding gradient problem.

What next?



What we've done so far is pretty good, but it's not going to scale well.

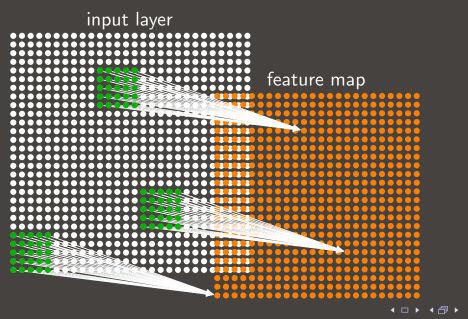
- These are small images, and only black-and-white.
- Imagine we had a more-typical image size (200×200) and 3 colours? Now we're up to 120,000 input parameters.
- We need an approach that is more efficient.
- A good place to start would be an approach that doesn't throw away all of the spatial information.
- The data is 28×28 , not 784×1 .
- We should redesign our network to account for the spatial information. How do we do that?
- The first step called a Convolution Layer.



Convolution layers: feature maps



Create a set of neurons that, instead of using all of the input data, only takes input from a small area of the image. This is called a "feature map".



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Feature maps



Some notes about feature maps.

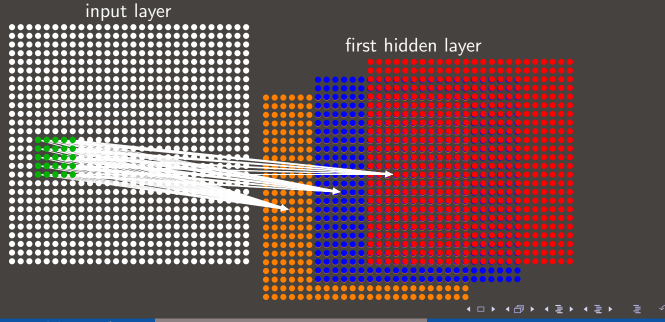
- Notice that the feature map is smaller (24×24) than the input layer (28×28) .
- The size of the feature map is partially set by the 'stride', meaning the number of pixels we shift to use as the input to the next neuron. In this case I've used a stride of 1.
- The weights and biases are shared by all the neurons in the feature map.
- Why? The goal is to train the feature map to recognize a single feature in the input, regardless of its location in the image.
- Consequently, it makes no sense to have a single feature map as the first hidden layer. Rather, multiple feature maps are used as the first layer.
- Feature maps are also called "filters" and "kernels".



Convolution layers, continued more



The first hidden layer, a "convolution layer", consists of multiple feature maps. The same inputs are fed to the neurons in different feature maps.



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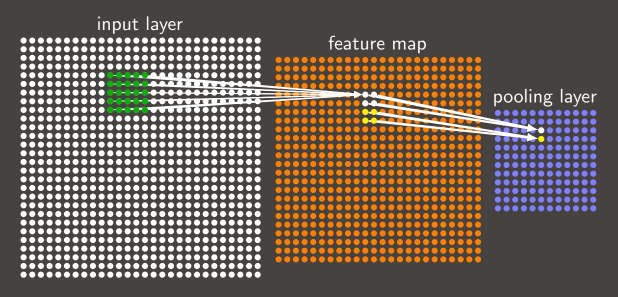
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Pooling layers



Each feature map is often followed by a "pooling layer".



In this case, 2×2 feature map neurons are mapped to a single pooling layer neuron.

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Pooling layers, continued



Some notes about pooling layers.

- The purpose of the layers is reduce the size of the data, and thus the number of parameters.
- The reduction in data also helps with over-fitting.
- Rather than use one of the activation functions we've already discussed, pooling layers use other functions.
- These functions do not have parameters in them which need to be fit.
- The most common function used is 'max', simply taking the maximum input value.
- Other functions are sometimes used, average pooling, L2-norm pooling.



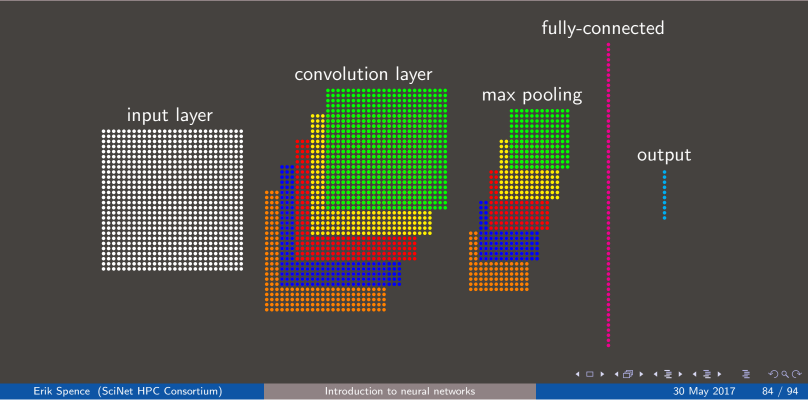
2D data formatting



Generally, 2D images are actually 3D, to deal with colours. Where the third dimension (the "channels") shows up in the dimensionality is indicated with the "image_data_format" function.

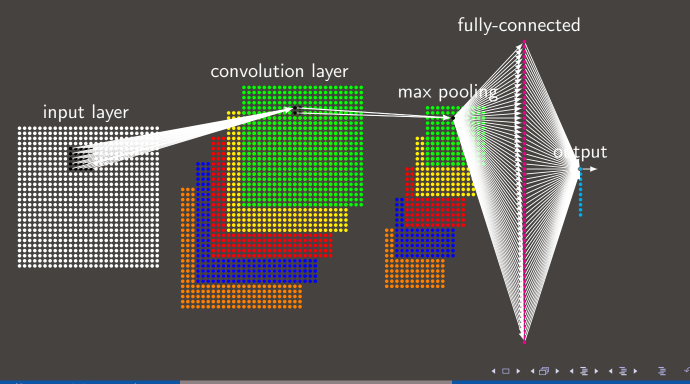
Our network, latest version





Our network, latest version





Our network revisited again



```
In [74]: numfm = 20; numnodes = 100;
In [75]:
In [75]: model3 = km.Sequential()
In [76]:
In [76]: model3.add(kl.Conv2D(numfm, kernel_size = (5, 5), strides = (1, 1),
...: input_shape = (28, 28, 1), activation = "relu"))
In [77]: model3.add(kl.MaxPooling2D(pool_size = (2, 2), strides = (2, 2))
In [78]:
In [78]: model3.add(kl.Flatten())
In [79]:
In [79]: model3.add(kl.Dense(numnodes, activation = "tanh"))
In [80]: model3.add(kl.Dense(10, activation = "softmax"))
In [81]:
```

The "Flatten" layer converts the 2D output to 1D, so that the fully-connected layer can handle it.

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Our network revisited again, continued



| In [81]: model3.summary() | | |
|---|--------------------|---------|
| Layer (type) | Output Shape | Param # |
| conv2d_2 (Conv2D) | (None, 24, 24, 20) | 520 |
| max_pooling2d_1 (MaxPooling2 | | 0 |
| flatten_1 (Flatten) | | 0 |
| dense_1 (Dense) | (None, 100) | 288100 |
| dense_2 (Dense) | (None, 10) | 1010 |
| Total params: 289,630.0 Trainable params: 289,630.0 Non-trainable params: 0.0 | | |
| | | |

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Our network revisited again, more



```
In [82]:
In [82]: model3.compile(loss = "categorical_crossentropy", optimizer = "sgd",
...: metrics = ['accuracy'])

In [83]:
In [83]:
In [83]: fit = model3.fit(train_in, train_out, epochs = 30, batch_size = 20,
...: validation_data = (val_in, val_out), verbose = 2)

Train on 50000 samples, validate on 10000 samples

Epoch 1/30

33s - loss: 0.1219 - acc: 0.9658 - val_loss: 0.1083 - val_acc: 0.9707

Epoch 2/30

33s - loss: 0.0997 - acc: 0.9709 - val_loss: 0.0940 - val_acc: 0.9753

:

Epoch 30/30

17s - loss: 0.0060 - acc: 0.9996 - val_loss: 0.0464 - val_acc: 0.9866

In [84]:
```

Our network revisited again, some more



Now check against the test data.

98.87%! Only 113 / 10000 wrong! Not bad!

You can improve this even more by adding another convolution layer-max pooling layer after the first pair, adding dropout to both layers, and expanding the dataset.

But we'll leave that as an exercise.

```
In [84]:
In [84]: score = model.evaluate(test_in, test_out)
In [85]:
In [85]: score
Out[85]: [0.034107370334264121, 0.9887000000000002]
In [86]:
```

Notes on Convolution Networks



The previous network is called a Convolution Neural Network (CNN), and is quite common in image analysis.

- Often more than a single convolution layer-pooling layer combination will be used.
- This will lead to improved performance, in this case.
- In practice people come up with all manner of combinations of convolution, pooling and fully-connected layers in their networks.
- Trial-and-error is your friend here. Reviewing the literature, you will find themes, but also much art.

Using GPUs



An important note. Graphical Processing Units (GPUs) are particularly good at running NN-training calculations.

| data size | CPU only | | CPU-GPU | |
|-----------|------------|------------|------------|------------|
| | epoch time | total time | epoch time | total time |
| 50000 | 41 s | 21 min 4 s | 4 s | 2 min 43s |
| 250000 | 198 s | 100 min | 26 s | 15 min |

These numbers are for our previous network. These were run on a Power 8 CPU, and a P100 GPU.

Unfortunately, multi-GPU functionality is not yet available in Keras running on Theano (though Theano and TensorFlow both support multiple GPUs).

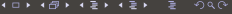


Deep Learning



What is Deep Learning?

- Quite simply: a neural network with many hidden layers.
- Our last network probably qualified as Deep Learning, though barely.
- Up until the mid-2000s neural network research was dominated by "shallow" networks, networks with only 1 or 2 hidden layers.
- The breakthrough came in discovering that it was practical to train networks with a larger number of hidden layers.
- But it only became practical with the advent of sufficient computing power (GPUs) and easily-accessible huge data sets.
- State-of-the-art networks today can contain dozens of layers.



Other neural network types



What we've looked at are called feed-forward networks, since the data only goes in one direction. There are other classes of neural network too, which we will save for an advanced course.

- Recurrent Neural Networks (RNNs). These are networks where output from the network is used as input to the network. These are used for networks that must remember previous information.
- Modular Neural Networks. In this scenario various sub-networks are glued together through an intermediary network.
- Self-organizing maps. Networks which set their own architecture, rather than being specified by the programmer.
- Generative adversarial networks. Two networks work together, one generating data and one judging the data.
- And many others....



Linky goodness



Neural network classes:

- http://neuralnetworksanddeeplearning.com
- http://www.cs.utoronto.ca/~fidler/teaching/2015/CSC2523.html
- http://cs231n.stanford.edu

Backpropagation:

- http://colah.github.io/posts/2015-08-Backprop
- http://cs231n.github.io/optimization-2