# 12: Support Vector Machines (SVMs)

**Previous Next Index** 

# Support Vector Machine (SVM) - Optimization objective

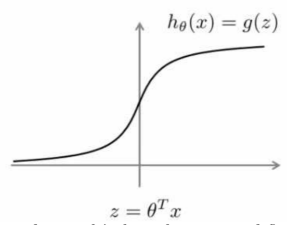
- So far, we've seen a range of different algorithms
  - o With supervised learning algorithms performance is pretty similar
    - What matters more often is;
      - The amount of training data
      - Skill of applying algorithms
- One final supervised learning algorithm that is widely used support vector machine (SVM)
  - Compared to both logistic regression and neural networks, a SVM sometimes gives a cleaner way of learning non-linear functions
  - o Later in the course we'll do a survey of different supervised learning algorithms

### An alternative view of logistic regression

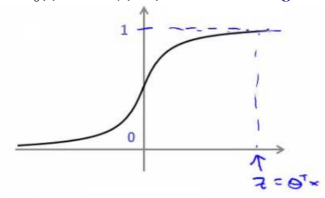
- Start with logistic regression, see how we can modify it to get the SVM
  - As before, the logistic regression hypothesis is as follows

$$h_{\theta}(x) = \frac{1}{1 + e^{-\theta^T x}}$$

o And the sigmoid activation function looks like this



- o In order to explain the math, we use z as defined above
- What do we want logistic regression to do?
  - $\circ$  We have an example where y = 1
    - Then we hope  $h_{\theta}(x)$  is close to 1
    - With  $h_{\theta}(x)$  close to 1,  $(\theta^T x)$  must be **much larger** than o



- $\circ$  Similarly, when y = 0
  - Then we hope  $h_{\theta}(x)$  is close to o

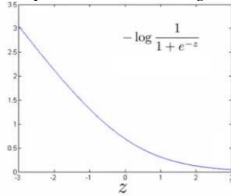
- 12\_Support\_Vector\_Machines
  - With  $h_{\theta}(x)$  close to o,  $(\theta^T x)$  must be **much less** than o
  - o This is our classic view of logistic regression
    - Let's consider another way of thinking about the problem
  - Alternative view of logistic regression
    - If you look at cost function, each example contributes a term like the one below to the overall cost function

$$-(y \log h_{\theta}(x) + (1-y) \log(1 - h_{\theta}(x)))$$

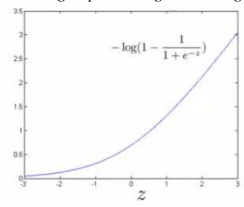
- For the overall cost function, we sum over all the training examples using the above function, and have a 1/m term
- If you then plug in the hypothesis definition  $(h_{\theta}(x))$ , you get an expanded cost function equation;

$$= -y \log \frac{1}{1 + e^{-\theta^T x}} - (1 - y) \log(1 - \frac{1}{1 + e^{-\theta^T x}})$$

- o So each training example contributes that term to the cost function for logistic regression
- If y = 1 then only the first term in the objective matters
  - o If we plot the functions vs. z we get the following graph



- This plot shows the cost contribution of an example when y = 1 given z
  - So if z is big, the cost is low this is good!
  - But if z is o or negative the cost contribution is high
  - This is why, when logistic regression sees a positive example, it tries to set  $\theta^T$  x to be a very large term
- If y = 0 then only the second term matters
  - We can again plot it and get a similar graph

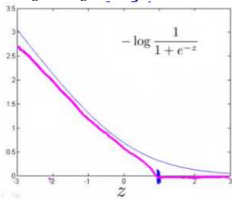


- Same deal, if z is small then the cost is low
  - But if s is large then the cost is massive

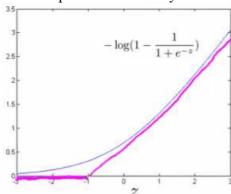
### SVM cost functions from logistic regression cost functions

- To build a SVM we must redefine our cost functions
  - $\circ$  When y = 1
    - Take the y = 1 function and create a new cost function
    - Instead of a curved line create two straight lines (magenta) which acts as an approximation to

the logistic regression y = 1 function



- Take point (1) on the z axis
  - Flat from 1 onwards
  - Grows when we reach 1 or a lower number
- This means we have two straight lines
  - Flat when cost is o
  - Straight growing line after 1
- So this is the new y=1 cost function
  - Gives the SVM a computational advantage and an easier optimization problem
  - We call this function **cost**<sub>1</sub>(**z**)
- Similarly
  - $\circ$  When y = 0
    - Do the equivalent with the y=o function plot



- We call this function **cost**<sub>0</sub>(**z**)
- So here we define the two cost function terms for our SVM graphically
  - How do we implement this?

### The complete SVM cost function

• As a comparison/reminder we have logistic regression below

$$\min_{\theta} \frac{1}{m} \left[ \sum_{i=1}^{m} y^{(i)} \left( -\log h_{\theta}(x^{(i)}) \right) + (1 - y^{(i)}) \left( (-\log(1 - h_{\theta}(x^{(i)}))) \right) \right] + \frac{\lambda}{2m} \sum_{j=1}^{n} \theta_{j}^{2}$$

- o If this looks unfamiliar its because we previously had the sign outside the expression
- For the SVM we take our two logistic regression y=1 and y=0 terms described previously and replace with
   ocost<sub>1</sub>(θ<sup>T</sup> x)
  - $\circ \operatorname{cost}_{\mathbf{O}}(\mathbf{\theta}^T \mathbf{x})$
- So we get

 $\min_{\theta} \frac{1}{m} \sum_{i=1}^{m} \left[ y^{(i)} cost_1(\theta^T x^{(i)}) + (1 - y^{(i)}) cost_0(\theta^T x^{(i)}) \right] + \frac{\lambda}{2m} \sum_{i=1}^{n} \theta_j^2$ 

### SVM notation is slightly different

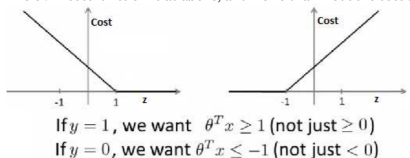
- In convention with SVM notation we rename a few things here
- 1) Get rid of the 1/m terms
  - This is just a slightly different convention
  - $\circ$  By removing 1/m we should get the same optimal values for
    - 1/m is a constant, so should get same optimization
    - e.g. say you have a minimization problem which minimizes to u = 5
      - If your cost function \* by a constant, you still generates the minimal value
      - That minimal value is different, but that's irrelevant
- 2) For logistic regression we had two terms;
  - Training data set term (i.e. that we sum over m) = A
  - Regularization term (i.e. that we sum over n) = **B** 
    - So we could describe it as  $A + \lambda B$
    - Need some way to deal with the trade-off between regularization and data set terms
    - Set different values for  $\lambda$  to parametrize this trade-off
  - $\circ$  Instead of parameterization this as A +  $\lambda$ B
    - For SVMs the convention is to use a different parameter called C
    - So do CA + B
    - If C were equal to  $1/\lambda$  then the two functions (CA + B and A +  $\lambda$ B) would give the same value
- So, our overall equation is

$$\min_{\theta} C \sum_{i=1}^{m} \left[ y^{(i)} cost_1(\theta^T x^{(i)}) + (1 - y^{(i)}) cost_0(\theta^T x^{(i)}) \right] + \frac{1}{2} \sum_{i=1}^{n} \theta_j^2$$

- Unlike logistic,  $h_{\theta}(x)$  doesn't give us a probability, but instead we get a direct prediction of 1 or 0
  - So if  $\theta^T$  x is equal to or greater than  $\theta^T$  = 1
  - $\circ$  Else -->  $h_{\Theta}(x) = o$

# Large margin intuition

- Sometimes people refer to SVM as large margin classifiers
  - We'll consider what that means and what an SVM hypothesis looks like
  - o The SVM cost function is as above, and we've drawn out the cost terms below

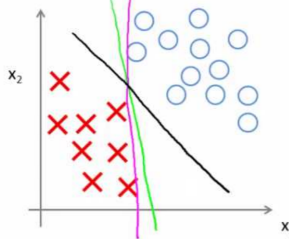


- ∘ Left is cost₁ and right is cost₀
- What does it take to make terms small
  - If y = 1
    - $cost_1(z) = 0$  only when  $z \ge 1$
  - If y = 0
    - $cost_O(z) = o$  only when  $z \le -1$
- Interesting property of SVM
  - If you have a positive example, you only really *need* z to be greater or equal to 0
    - If this is the case then you predict 1
  - SVM wants a bit more than that doesn't want to \*just\* get it right, but have the value be quite a bit bigger than zero
    - Throws in an extra safety margin factor
- Logistic regression does something similar
- What are the consequences of this?
  - o Consider a case where we set C to be huge
    - C = 100,000
    - So considering we're minimizing CA + B

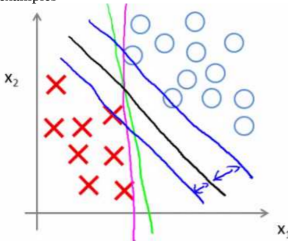
- 12\_Support\_Vector\_Machines
- If C is huge we're going to pick an A value so that A is equal to zero
- What is the optimization problem here how do we make A = o?
- Making A = 0
  - $\blacksquare \text{ If } y = 1$ 
    - Then to make our "A" term o need to find a value of  $\theta$  so  $(\theta^T x)$  is greater than or equal to 1
  - Similarly, if y = 0
    - Then we want to make "A" = 0 then we need to find a value of  $\theta$  so  $(\theta^T x)$  is equal to or less than -1
- So if we think of our optimization problem a way to ensure that this first "A" term is equal to o, we re-factor our optimization problem into just minimizing the "B" (regularization) term, because
  - When A = o --> A\*C = o
- So we're minimizing B, under the constraints shown below

$$Min = \frac{1}{2} \sum_{i=1}^{n} \Theta_{i}^{2}$$
  
S.t.  $\Theta^{T} \times (i) \ge 1$  if  $g^{(i)} = 1$   
 $\Theta^{T} \times (i) \le -1$  if  $g^{(i)} = 0$ 

o Turns out when you solve this problem you get interesting decision boundaries

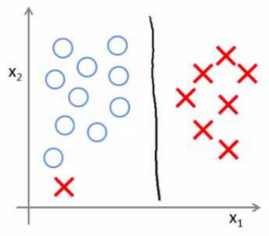


- $\circ\,$  The green and magenta lines are functional decision boundaries which could be chosen by logistic regression
  - But they probably don't generalize too well
- The black line, by contrast is the the chosen by the SVM because of this safety net imposed by the optimization graph
  - More robust separator
- Mathematically, that black line has a larger minimum distance (margin) from any of the training examples

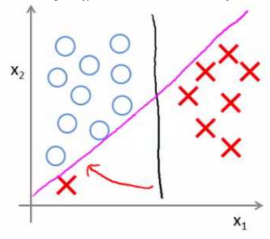


- By separating with the largest margin you incorporate robustness into your decision making process
- We looked at this at when C is very large

- 12\_Support\_Vector\_Machines
  - o SVM is more sophisticated than the large margin might look
    - If you were just using large margin then SVM would be very sensitive to outliers



- You would risk making a ridiculous hugely impact your classification boundary
  - A single example might not represent a good reason to change an algorithm
  - If C is very large then we do use this quite naive maximize the margin approach



- So we'd change the black to the magenta
- But if C is reasonably small, or a not too large, then you stick with the black decision boundary
- What about non-linearly separable data?
  - Then SVM still does the right thing if you use a normal size C
  - So the idea of SVM being a large margin classifier is only really relevant when you have no outliers and you can easily linearly separable data
- o Means we ignore a few outliers

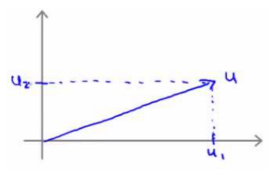
# Large margin classification mathematics (optional)

## **Vector inner products**

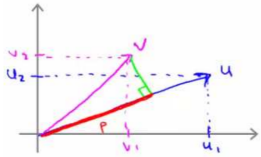
• Have two (2D) vectors u and v - what is the inner product  $(u^T v)$ ?

$$u = \begin{bmatrix} u_1 \\ u_2 \end{bmatrix} v = \begin{bmatrix} v_1 \\ v_2 \end{bmatrix}$$

- Plot *u* on graph
  - i.e  $u_1$  vs.  $u_2$

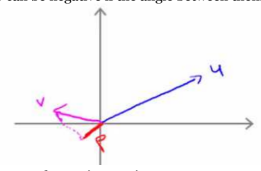


- One property which is good to have is the **norm** of a vector
  - Written as ||u||
    - This is the euclidean length of vector u
  - So  $||\mathbf{u}|| = \mathrm{SQRT}(u_1^2 + u_2^2) = \mathrm{real}$  number
    - i.e. length of the arrow above
    - Can show via Pythagoras
- For the inner product, take v and orthogonally project down onto u
  - First we can plot v on the same axis in the same way  $(v_1 \text{ vs } v_1)$
  - Measure the length/magnitude of the projection



- So here, the green line is the projection
  - p = length along u to the intersection
  - p is the magnitude of the projection of vector v onto vector u
- o Possible to show that

  - u<sup>T</sup> v = p \* ||u||
     So this is one way to compute the inner product
  - $u^T v = u_1 v_1 + u_2 v_2$
  - So therefore
    - $p * ||u|| = u_1v_1 + u_2v_2$
    - This is an important rule in linear algebra
  - We can reverse this too
    - So we could do
      - $\bullet \ v^T u = v_1 u_1 + \mathbf{v}_2 u_2$
      - Which would obviously give you the same number
- o p can be negative if the angle between them is 90 degrees or more



- So here p is negative
- Use the vector inner product theory to try and understand SVMs a little better

### **SVM decision boundary**

13.07.2015 07:17

$$\begin{aligned} & \min_{\theta} \frac{1}{2} \sum_{j=1}^{n} \theta_{j}^{2} \\ & \text{s.t.} \quad \theta^{T} x^{(i)} \geq 1 & \text{if } y^{(i)} = 1 \\ & \theta^{T} x^{(i)} \leq -1 & \text{if } y^{(i)} = 0 \end{aligned}$$

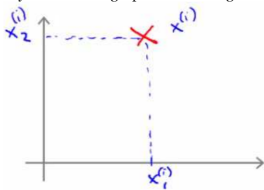
- For the following explanation two simplification
  - $\circ$  Set  $\theta_0$ = o (i.e. ignore intercept terms)
  - $\circ$  Set n = 2  $(x_1, x_2)$ 
    - i.e. each example has only 2 features
- Given we only have two parameters we can simplify our function to

• And, can be re-written as

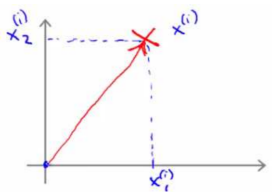
- o Should give same thing
- We may notice that

- $\circ$  The term in red is the norm of  $\theta$ 
  - If we take  $\theta$  as a 2x1 vector
  - If we assume  $\theta_0$  = 0 its still true
- So, finally, this means our optimization function can be re-defined as

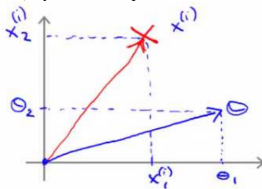
- So the SVM is minimizing the squared norm
- Given this, what are the  $(\theta^T x)$  parameters doing?
  - $\circ$  Given  $\theta$  and given example x what is this equal to
    - We can look at this in a comparable manner to how we just looked at u and v
  - o Say we have a single positive training example (red cross below)



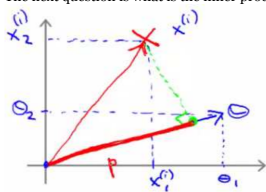
• Although we haven't been thinking about examples as vectors it can be described as such



 $\circ$  Now, say we have our parameter vector  $\theta$  and we plot that on the same axis



• The next question is what is the inner product of these two vectors



- p, is in fact p<sup>i</sup>, because it's the length of p for example i
   Given our previous discussion we know

$$(\theta^T \mathbf{x}^i) = \mathbf{p}^i * ||\theta||$$
$$= \theta_1 \mathbf{x}^i_1 + \theta_2 \mathbf{x}^i_2$$

- So these are both equally valid ways of computing  $\theta^T x^i$
- What does this mean?
  - The constraints we defined earlier

■ 
$$(\theta^T x) >= 1 \text{ if } y = 1$$

$$\blacksquare$$
  $(\Theta^T \mathbf{v}) < = -1$  if  $\mathbf{v} = \Theta$ 

•  $(\theta^T x) <= -1$  if y = 0• Can be replaced/substituted with the constraints

$$p^{i} * ||\theta|| >= 1 \text{ if } y = 1$$

$$p^{i} * ||\theta|| <= -1 \text{ if } y = 0$$

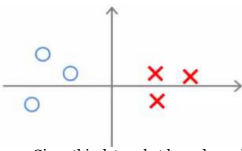
• Writing that into our optimization objective

$$\min_{\theta} \frac{1}{2} \sum_{j=1}^{n} \theta_{j}^{2} = \frac{1}{2} \|\theta\|^{2}$$
s.t.  $p^{(i)} \cdot \|\theta\| \ge 1$  if  $y^{(i)} = 1$ 

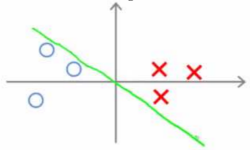
$$p^{(i)} \cdot \|\theta\| \le -1$$
 if  $y^{(i)} = 1$ 

• So, given we've redefined these functions let us now consider the training example below

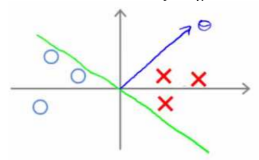
13.07.2015 07:17 9 von 18



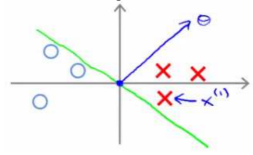
- $\circ$  Given this data, what boundary will the SVM choose? Note that we're still assuming  $\theta_0 = 0$ , which means the boundary has to pass through the origin (0,0)
  - Green line small margins



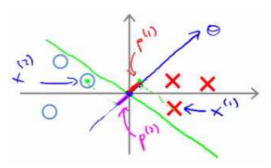
- SVM would not chose this line
  - Decision boundary comes very close to examples
  - Lets discuss why the SVM would **not** chose this decision boundary
- o Looking at this line
  - We can show that  $\theta$  is at 90 degrees to the decision boundary



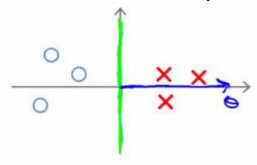
- 0 is always at 90 degrees to the decision boundary (can show with linear algebra, although we're not going to!)
- So now lets look at what this implies for the optimization objective
  - Look at first example (x1)



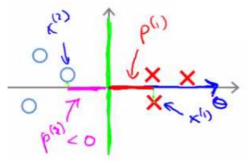
- $\circ$  Project a line from  $x^1$  on to to the  $\theta$  vector (so it hits at 90 degrees)
  - The distance between the intersection and the origin is (p¹)
- Similarly, look at second example (x<sup>2</sup>)
  - Project a line from  $x^2$  into to the  $\theta$  vector
  - This is the magenta line, which will be **negative** (p<sup>2</sup>)
- o If we overview these two lines below we see a graphical representation of what's going on;



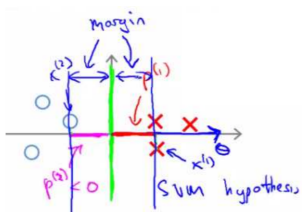
- We find that both these p values are going to be pretty small
- If we look back at our optimization objective
  - We know we need  $p^{1} * ||\theta||$  to be bigger than or equal to 1 for positive examples
    - If p is small
      - Means that  $\|\theta\|$  must be pretty large
  - Similarly, for negative examples we need  $p^2 * ||\theta||$  to be smaller than or equal to -1
    - We saw in this example p<sup>2</sup> is a small negative number
      - So  $\|\theta\|$  must be a large number
- Why is this a problem?
  - The optimization objective is trying to find a set of parameters where the norm of theta is small
    - So this doesn't seem like a good direction for the parameter vector (because as p values get smaller  $||\theta||$  must get larger to compensate)
      - So we should make p values larger which allows  $|\theta|$  to become smaller
- So lets chose a different boundary



- $\circ\,$  Now if you look at the projection of the examples to  $\theta$  we find that  $p^1$  becomes large and  $||\theta||$  can become small
- So with some values drawn in



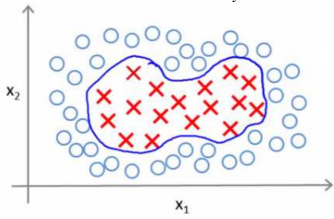
- $\circ$  This means that by choosing this second decision boundary we can make  $|\theta|$  smaller
  - Which is why the SVM choses this hypothesis as better
  - This is how we generate the large margin effect



- The magnitude of this margin is a function of the p values
  - So by maximizing these p values we minimize  $||\theta||$
- Finally, we did this derivation assuming  $\theta_0 = 0$ ,
  - If this is the case we're entertaining only decision boundaries which pass through (0,0)
  - $\circ$  If you allow  $\theta_0$  to be other values then this simply means you can have decision boundaries which cross through the x and y values at points other than (0,0)
  - $\circ$  Can show with basically same logic that this works, and even when  $\theta_0$  is non-zero when you have optimization objective described above (when C is very large) that the SVM is looking for a large margin separator between the classes

# **Kernels - 1: Adapting SVM to non-linear classifiers**

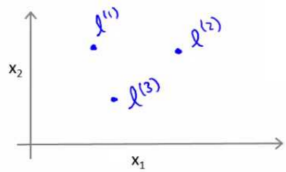
- What are kernels and how do we use them
  - We have a training set
  - We want to find a non-linear boundary



- Come up with a complex set of polynomial features to fit the data
  - Have  $h_{\theta}(x)$  which
    - Returns 1 if the combined weighted sum of vectors (weighted by the parameter vector) is less than or equal to 0
    - Else return o
  - Another way of writing this (new notation) is
    - That a hypothesis computes a decision boundary by taking the sum of the parameter vector multiplied by a **new feature vector f**, which simply contains the various high order x terms
    - e.g.
      - $h_{\theta}(x) = \theta_0 + \theta_1 f_1 + \theta_2 f_2 + \theta_3 f_3$
      - Where
        - $\bullet$   $f_1 = x_1$
        - $f_2 = x_1 x_2$
        - $\bullet$  f<sub>2</sub> = ...
        - i.e. not specific values, but each of the terms from your complex polynomial function
  - Is there a better choice of feature f than the high order polynomials?
    - As we saw with computer imaging, high order polynomials become computationally expensive

New features

- $\circ$  Define three features in this example (ignore  $x_0$ )
- Have a graph of x<sub>1</sub> vs. x<sub>2</sub> (don't plot the values, just define the space)
- o Pick three points in that space



- These points l<sup>1</sup>, l<sup>2</sup>, and l<sup>3</sup>, were chosen manually and are called **landmarks** 
  - Given x, define f1 as the similarity between (x, l1)

$$= \exp(-(||x-l^1||^2)/2\sigma^2)$$

$$= \exp\left(-\frac{||x-l^{(1)}||^2}{2\sigma^2}\right)$$

$$= \exp\left(-\frac{||x-l^{(1)}||^2}{2\sigma^2}\right)$$

- $|| \mathbf{x} \mathbf{l}^{1} ||$  is the euclidean distance between the point x and the landmark  $\mathbf{l}^{1}$  squared
  - Disussed more later
- If we remember our statistics, we know that
  - $\bullet$   $\sigma$  is the standard deviation
  - $\sigma^2$  is commonly called the **variance**
- Remember, that as discussed

$$||x-l^{(1)}||^2 = \sum_{j=1}^n (x_j - l_j^{(1)})^2$$

- o So, f2 is defined as
  - $f_2 = similarity(x, l^1) = exp(-(||x l^2||^2) / 2\sigma^2)$
- And similarly
  - $f_3 = similarity(x, l^2) = exp(-(||x l^1||^2) / 2\sigma^2)$
- This similarity function is called a **kernel** 
  - This function is a **Gaussian Kernel**
- o So, instead of writing similarity between x and l we might write
  - $f_1 = k(x, l^1)$

### Diving deeper into the kernel

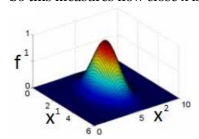
- So lets see what these kernels do and why the functions defined make sense
  - Say x is close to a landmark
    - Then the squared distance will be ~o
      - Sc

- Which is basically e<sup>-0</sup>
  - Which is close to 1
- Say x is far from a landmark
  - Then the squared distance is big
    - Gives e<sup>-large</sup> number
      - Which is close to zero
- Each landmark defines a new features
- If we plot f1 vs the kernel function we get a plot like this
  - $\circ$  Notice that when x = [3,5] then  $f_1 = 1$
  - $\circ$  As x moves away from [3,5] then the feature takes on values close to zero

13 von 18

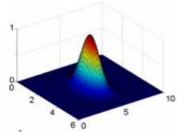
13.07.2015 07:17

o So this measures how close x is to this landmark

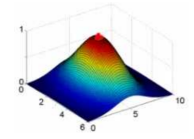


#### What does σ do?

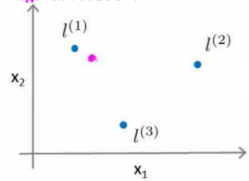
- $\sigma^2$  is a parameter of the Gaussian kernel
  - o Defines the steepness of the rise around the landmark
- Above example  $\sigma^2 = 1$
- Below  $\sigma^2 = 0.5$



- We see here that as you move away from 3,5 the feature f1 falls to zero much more rapidly
- The inverse can be seen if  $\sigma^2 = 3$

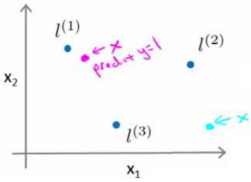


- Given this definition, what kinds of hypotheses can we learn?
  - With training examples x we predict "1" when
  - $\theta_0 + \theta_1 f_1 + \theta_2 f_2 + \theta_3 f_3 >= 0$ 
    - For our example, lets say we've already run an algorithm and got the
      - $\theta_0 = -0.5$
      - $\theta_1 = 1$
      - $\theta_2 = 1$
      - $\theta_2 = 0$
    - Given our placement of three examples, what happens if we evaluate an example at the magenta dot below?

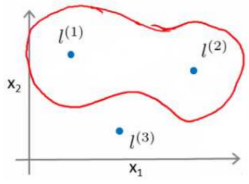


- Looking at our formula, we know f1 will be close to 1, but f2 and f3 will be close to 0
  - So if we look at the formula we have

- 12\_Support\_Vector\_Machines
- $\theta_0 + \theta_1 f_1 + \theta_2 f_2 + \theta_3 f_3 >= 0$
- -0.5 + 1 + 0 + 0 = 0.5
  - 0.5 is greater than 1
- If we had another point far away from all three



- This equates to -0.5
  - So we predict o
- $\circ$  Considering our parameter, for points near  $l^1$  and  $l^2$  you predict 1, but for points near  $l^3$  you predict 0
- Which means we create a non-linear decision boundary that goes a lil' something like this;



- Inside we predict y = 1
- Outside we predict y = o
- So this show how we can create a non-linear boundary with landmarks and the kernel function in the support vector machine
  - o But
    - How do we get/chose the landmarks
    - What other kernels can we use (other than the Gaussian kernel)

# **Kernels II**

- Filling in missing detail and practical implications regarding kernels
- Spoke about picking landmarks manually, defining the kernel, and building a hypothesis function
  - Where do we get the landmarks from?
  - For complex problems we probably want lots of them

### Choosing the landmarks

- Take the training data
- For each example place a landmark at exactly the same location
- So end up with m landmarks
  - One landmark per location per training example
  - Means our features measure how close to a training set example something is
- Given a new example, compute all the f values
  - Gives you a feature vector f (fo to fm)
    - $f_0 = 1$  always
- A more detailed look at generating the f vector
  - $\circ$  If we had a training example features we compute would be using  $(x^i, y^i)$ 
    - So we just cycle through each landmark, calculating how close to that landmark actually x<sup>i</sup> is

- 12\_Support\_Vector\_Machines
  - $f_1^i$ , =  $k(x^i, l^1)$
  - $f_2^i$ , = k(x<sup>i</sup>, l<sup>2</sup>)
  - **.**..
  - $f_m^i$ , =  $k(x^i, l^m)$
  - Somewhere in the list we compare x to itself... (i.e. when we're at f<sub>1</sub><sup>i</sup>)
    - So because we're using the Gaussian Kernel this evalues to 1
  - Take these m features (f<sub>1</sub>, f<sub>2</sub> ... f<sub>m</sub>) group them into an [m +1 x 1] dimensional vector called f
    - f<sup>i</sup> is the f feature vector for the ith example
    - And add a oth term = 1
  - Given these kernels, how do we use a support vector machine

### SVM hypothesis prediction with kernels

- Predict y = 1 if  $(\theta^T f) >= 0$ 
  - $\circ$  Because  $\theta = [m+1 \times 1]$
  - $\circ$  And f = [m +1 x 1]
- $\bullet$  So, this is how you make a prediction assuming you already have  $\theta$ 
  - How do you get θ?

### **SVM** training with kernels

• Use the SVM learning algorithm

$$\min_{\theta} C \sum_{i=1}^{m} y^{(i)} cost_1(\theta^T f^{(i)}) + (1 - y^{(i)}) cost_0(\theta^T f^{(i)}) + \frac{1}{2} \sum_{j=1}^{n} \theta_j^2$$

- o Now, we minimize using f as the feature vector instead of x
- o By solving this minimization problem you get the parameters for your SVM
- In this setup, m = n
  - Because number of features is the number of training data examples we have
- One final mathematic detail (not crucial to understand)
  - $\circ$  If we ignore  $\theta_0$  then the following is true

$$\sum_{j=1}^{n} \theta_j^2 = \theta^T \theta$$

• What many implementations do is

$$\theta^T M \theta$$

- Where the matrix M depends on the kernel you use
- Gives a slightly different minimization means we determine a rescaled version of  $\theta$
- Allows more efficient computation, and scale to much bigger training sets
- If you have a training set with 10 000 values, means you get 10 000 features
  - Solving for all these parameters can become expensive
  - So by adding this in we avoid a for loop and use a matrix multiplication algorithm instead
- You can apply kernels to other algorithms
  - But they tend to be very computationally expensive
  - $\circ\,$  But the SVM is far more efficient so more practical
- Lots of good off the shelf software to minimize this function

### • SVM parameters (C)

- o Bias and variance trade off
- o Must chose C
  - C plays a role similar to 1/LAMBDA (where LAMBDA is the regularization parameter)
- Large C gives a hypothesis of **low bias high variance** --> overfitting
- Small C gives a hypothesis of **high bias low variance** --> underfitting

### • SVM parameters ( $\sigma^2$ )

- Parameter for calculating f values
  - $\blacksquare$  Large  $\sigma^2$  f features vary more smoothly higher bias, lower variance

• Small  $\sigma^2$  - f features vary abruptly - low bias, high variance

# **SVM** - implementation and use

- So far spoken about SVM in a very abstract manner
- What do you need to do this
  - $\circ\,$  Use SVM software packages (e.g. liblinear, libsvm) to solve parameters  $\theta$
  - o Need to specify
    - Choice of parameter C
    - Choice of kernel

### Choosing a kernel

- We've looked at the Gaussian kernel
  - $\circ$  Need to define  $\sigma(\sigma^2)$ 
    - Discussed σ<sup>2</sup>
  - o When would you chose a Gaussian?
    - If n is small and/or m is large
      - e.g. 2D training set that's large
  - o If you're using a Gaussian kernel then you may need to implement the kernel function
    - e.g. a function
      - fi = kernel(x1,x2)
        - Returns a real number
    - Some SVM packages will expect you to define kernel
    - Although, some SVM implementations include the Gaussian and a few others
      - Gaussian is probably most popular kernel
  - NB make sure you perform **feature scaling** before using a Gaussian kernel
    - If you don't features with a large value will dominate the f value
- Could use no kernel linear kernel
  - $\circ$  Predict y = 1 if  $(\theta^T x) >= 0$ 
    - So no f vector
    - Get a standard linear classifier
  - Why do this?
    - If n is large and m is small then
      - Lots of features, few examples
      - Not enough data risk overfitting in a high dimensional feature-space
- Other choice of kernel
  - o Linear and Gaussian are most common
  - o Not all similarity functions you develop are valid kernels
    - Must satisfy Merecer's Theorem
    - SVM use numerical optimization tricks
      - Mean certain optimizations can be made, but they must follow the theorem
  - o Polynomial Kernel
    - We measure the similarity of x and l by doing one of
      - $\blacksquare (\mathbf{x}^T \mathbf{l})^2$
      - $\mathbf{x}^T \mathbf{1} \mathbf{3}$
      - $(x^T l+1)^3$
    - General form is
      - $\bullet$  (x<sup>T</sup>l+Con)<sup>D</sup>
    - If they're similar then the inner product tends to be large
    - Not used that often
    - Two parameters
      - Degree of polynomial (D)
      - Number you add to l (Con)
    - Usually performs worse than the Gaussian kernel
    - Used when x and l are both non-negative
  - String kernel
    - Used if input is text strings
    - Use for text classification
  - o Chi-squared kernel

### • Histogram intersection kernel

#### **Multi-class classification for SVM**

- Many packages have built in multi-class classification packages
- Otherwise use one-vs all method
- Not a big issue

### Logistic regression vs. SVM

- When should you use SVM and when is logistic regression more applicable
- If n (features) is large vs. m (training set)
  - $\circ$  e.g. text classification problem
    - Feature vector dimension is 10 000
    - Training set is 10 1000
    - Then use logistic regression or SVM with a linear kernel
- If n is small and m is intermediate
  - $\circ$  n = 1 1000
  - o m = 10 10 000
  - Gaussian kernel is good
- If n is small and m is large
  - $\circ$  n = 1 1000
  - o m = 50 000+
    - SVM will be slow to run with Gaussian kernel
  - In that case
    - Manually create or add more features
    - Use logistic regression of SVM with a linear kernel
- Logistic regression and SVM with a linear kernel are pretty similar
  - Do similar things
  - Get similar performance
- A lot of SVM's power is using different kernels to learn complex non-linear functions
- For all these regimes a well designed NN should work
  - o But, for some of these problems a NN might be slower SVM well implemented would be faster
- SVM has a convex optimization problem so you get a global minimum
- It's not always clear how to chose an algorithm
  - o Often more important to get enough data
  - o Designing new features
  - o Debugging the algorithm
- SVM is widely perceived a very powerful learning algorithm