## Large-Scale Support Vector Machines: Algorithms and Theory Research Exam

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February 27, 2009

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Large-Scale Support Vector Machines: Algorit

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# Outline

### 1 Introduction: large-scale learning

- 2 Background: Support vector machines
- 3 Primal training methods
  - Stochastic gradient methods
  - Pegasos
- 4 Dual training methods
   Dual coordinate descent
   Bundle method (BMRM)
- 5 Stochastic gradient in learning
- 6 Summary and future directions
- 7 References

# Motivation: the growth of data

- Dataset sizes have been growing at a rapid rate the past few years
- In supervised learning, this growth affects the size of training sets
- Benefit: More information, can make useful predictions
- Concern: Can our methods of analysis scale to such datasets?
  - ► A learning algorithm that scales superlinearly in the size of the training set will be infeasible
  - ► ⇒ Need methods that scale at worst linearly in the number of examples

# Example: ad click data

Google flight tickets Search Advanced Search Preferences			
Web			
Flight Tickets         www.Expedia.com       Book with Expedia for Cheap 2009 Airfares & Ticket Prices Guaranteed         Flight Ticket Deals       www.Travelzoo.com         www.Travelzoo.com       Pricing Your Trip? Let Travelzoo Find the Best Deals. Search Now!         Flight - Cheap Tickets       when You Save Money On Flights With Cheaptickets®, You Win.			
Airline Tickets, Cheap Flights, Cheap Plane Tickets, Hotels, Car <ul> <li>▼</li> <li>Feb 9, 2009 Discount airfares, Cheap Plane Fares, Cheap Air Tickets, Cheap Hotels and Car Rentals Book now and save up to 65% on flight tickets! *</li> <li>www.cheapoair.com/ - 227k - Cached - Similar pages - </li> </ul> <ul> <li>Cheap Travel, Flights, Hotels, Vacations, Car Rentals, Cruise</li> <li>▼</li> <li>Find hotels, airline tickets, vacation packages, travel deals, car rentals and cruises Flight + Hotel; Flight + Car; Hotel + Car</li> <li>Show stock quote for OWW</li> <li>www.orbitz.com/ - Similar pages - </li> </ul>			

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# Defining large-scale learning

- While large-scale can simply mean a large number of examples, a more technical definition follows
- In a learning problem, we want to minimize generalization error subject to two constraints
  - ► There is some maximum number of examples we can pick
  - There is some maximum amount of time available
- Active constraint defines scale of problem
  - ► Number of examples ⇒ small- or medium-scale
  - Time available  $\implies$  large-scale

# This talk

- We focus on methods for large-scale support vector machines (SVMs)
- One of the most popular approaches for binary classification tasks
  - Strong theoretical underpinnings
  - Good performance in practice
- Interested in the training algorithms and some of the theory behind them
  - What are some techniques for large-scale SVMs?
  - Why do they work? (specifically stochastic gradient descent)

# Outline

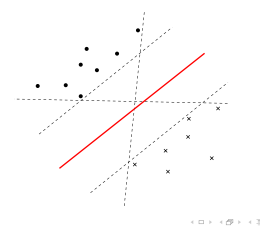
Introduction: large-scale learning

#### Background: Support vector machines

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# Support vector machines

- A support vector machine (SVM) is a binary classifier that finds a maximum margin separating hyperplane
- Intuitively, we expect such a hyperplane to generalize the best



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# Primal problem for SVMs

- Suppose we have a training set  $\mathcal{T} = \{(x_i, y_i)\}_{i=1}^n$
- How do we find the maximum margin separating hyperplane?
- If we allow for misclassifications at the expense of some penalty, the SVM problem is:

### Primal SVM problem

minimize 
$$\frac{\lambda}{2} ||w||^2 + \frac{1}{n} \sum_{i=1}^n [1 - y_i(w \cdot x_i)]_+$$

where  $[\cdot]_+$  denotes the hinge-loss:

$$[x]_+ = \max(0, x)$$

and  $\boldsymbol{w}$  denotes the normal to the hyperplane

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# Interpreting the primal problem

- Can think of the primal problem as  $\ell_2$  regularized hinge-loss minimization
- Falls in the general class of functions  $\hat{g}(w) = \ell_{\rm emp}(w) + r(w)$ 
  - $\ell_{emp}$  measures loss on the training set
  - r is a regularization term
  - Both functions are convex, and hence the optimization is tractable

# Dual formulation

• The dual SVM problem is the following:

Dual SVM problem

maximize 
$$\sum_{i} \alpha_{i} - \frac{1}{2} \sum_{i} \sum_{j} \alpha_{i} \alpha_{j} y_{i} y_{j} (x_{i} \cdot x_{j})$$
subject to  $0 \le \alpha_{i} \le \frac{1}{\lambda n}$ 

• According to the representer theorem, the optimal primal and dual solutions  $w^*$  and  $\alpha^*$  satisfy

$$w^* = \sum_i \alpha_i^* y_i x_i$$

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# Primal vs dual formulation

- Historically, SVMs have been solved using the dual
- Reasons?
  - Naturally extend to kernels
  - Precedent set by "hard margin" case: simple dual optimization constraints
- The primal problem that we stated renders both points largely moot
  - Can also handle kernels
  - Unconstrained!
- So, no a-priori reason to eschew the primal form
  - Opens up some new techniques

# Kernelized dual formulation

### Standard dual problem

$$\text{maximize } \sum_{i} \alpha_i - \frac{1}{2} \sum_{i} \sum_{j} \alpha_i \alpha_j y_i y_j (x_i \cdot x_j) \quad \text{subject to } 0 \le \alpha_i \le \frac{1}{\lambda n}$$

#### ₩

#### Kernelized dual problem

$$\text{maximize } \sum_{i} \alpha_i - \frac{1}{2} \sum_{i} \sum_{j} \alpha_i \alpha_j y_i y_j K(x_i, x_j) \quad \text{subject to } 0 \leq \alpha_i \leq \frac{1}{\lambda n}$$

• Can do the same for the primal form...

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# Kernelized primal formulation

### Standard primal problem

minimize 
$$\frac{\lambda}{2} ||w||^2 + \frac{1}{n} \sum_{i=1}^n [1 - y_i(w \cdot x_i)]_+$$

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### Kernelized primal problem

minimize 
$$\lambda_{f \in H} \frac{\lambda}{2} ||f||_{H}^{2} + \frac{1}{n} \sum_{i=1}^{n} [1 - y_{i}f(x_{i})]_{+}$$

where H is a reproducing kernel Hilbert space (RKHS) with kernel K.

• A Hilbert space is a complete inner product space, and an RKHS can be written  $\{f: f(x) = \sum_i \beta_i K(x, x_i)\}$ 

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# Solving the SVM problem

- Suppose we want to solve the SVM problem with a quadratic programming (QP) solver
- The dual problem requires access to the matrix Q, defined as

$$Q_{ij} = y_i y_j K(x_i, x_j)$$

- Q's size is  $n \times n$ , which for even moderately large training sets is too expensive to store in memory
  - An off-the-shelf solver is insufficient
  - ► We need to design more specialized QP solvers

# Classical SVM solvers: SVM<sup>light</sup> and SMO

- SVM<sup>light</sup>: instead of solving the (large) QP problem, focus on a subset of variables (the working set)
  - Pick a subset of variables that are "likely" to change
  - Now solve this reduced problem using a standard QP solver
- SMO is a special case of SVM<sup>light</sup> where we optimize only two variables at once
  - Advantage: this optimization can be done analytically; does not require an external QP solver
  - ► But we need heuristics for choosing the variables to optimize
- Can implement caching of  ${\boldsymbol{Q}}$  values to improve performance

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## Problems with classical SVM solvers

- **Problem**: These algorithms scale like  $n^2$  in the worst case
  - Infeasible on large datasets
- **Question**: Can we design algorithms that solve the problem more efficiently?
  - We will see some algorithms that scale either linearly or are independent of the number of examples!
  - ► There is a catch, however...

# Linear or kernel SVM?

- Most nascent large-scale solvers looked at the linear SVM case
- Why?
  - ► Simpler!
  - Oft-cited argument is "In some applications, data appear in a rich dimensional feature space, the performances are similar with/without nonlinear mapping", with the canonical example being text classification [HCL<sup>+</sup>08]
  - Most text classification tasks are linearly separable [Joa98]
- Focus on linear SVMs alone is obviously fundamentally limiting: a major weakness of several of these techniques
  - Those that do consider kernels do so in passing; for many, viability for arbitrary kernels is an open question

# Approximate SVM solution

- Newer solvers find approximate solutions to the SVM problem
- If the objective function is f(w), then instead of finding  $w^* = \arg\min f(w)$ , they find  $\tilde{w}$  such that

 $f(w^*) \le f(\tilde{w}) \le f(w^*) + \rho$ 

- The constant  $\rho$  is the (user-controllable) optimization tolerance
  - Runtime is explicitly analyzed in terms of this
  - We call  $\tilde{w}$  a  $\rho$ -optimal solution
- Approximate solutions are meaningful for the SVM problem because the optimization is a surrogate anyway
  - Important point that we discuss later

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 Stochastic gradient methods
 Pegasos

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• Dual coordinate descent
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# Stochastic gradient for SVMs

- Stochastic gradient descent (SGD) underlies at least three SVM training methods: SVM-SGD, NORMA, and Zhang's algorithm
- Idea is simply to apply SGD on the primal SVM problem
  - ► Advantage: Runtime is independent of number of examples
- Seems obvious, so why was it not tried earlier?
  - Historical favouring of the dual over the primal; dual SGA tried in the kernel adatron
  - Association of SGD with backpropagation in multi-layer perceptrons (non-convex problem)
  - Slow convergence rate

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  - ► Slow convergence rate ← will discuss this subsequently

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# Review: stochastic gradient descent (SGD)

- $\bullet\,$  SGD uses a randomized gradient estimate to minimize a function f(w)
  - Instead of  $\nabla f$  , use  $\tilde{\nabla} f$  where  $\mathbb{E}[\tilde{\nabla} f] = \nabla f$
- For empirical loss  $\ell_{emp}(w) = \frac{1}{n} \sum_{i} \ell(x_i, y_i; w)$ :

$$w_{t+1} \leftarrow w_t - \eta \nabla \ell(x_{i(t)}, y_{i(t)}; w_t)$$

where  $\eta$  is the learning rate and  $i(t) \in \{1, \ldots, n\}$  uniformly at random

Pros	Cons	
Fast: "instantaneous" gradient	Have to tune learning rate	
	Slow convergence	

# SGD update for SVMs

Recall the primal problem

$$\min \frac{\lambda}{2} ||w||^2 + \frac{1}{n} \sum_{i=1}^n [1 - y_i(w \cdot x_i)]_+$$

• The SGD update is:

$$w_t \leftarrow (1 - \eta_t \lambda) w_{t-1} - \begin{cases} \eta_t y_{i(t)} x_{i(t)} & \text{ if } y_{i(t)}(w_{t-1} \cdot x_{i(t)}) < 1\\ 0 & \text{ otherwise} \end{cases}$$

where  $i(t) \in \{1, 2, \dots, n\}$  uniformly at random

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# SVM-SGD: the algorithm

for 
$$t = 1 \dots T$$
  
Pick a random example  $(x_i, y_i)$   
 $\eta_t \leftarrow \frac{1}{\lambda(t+t_0)}$   
 $w_t \leftarrow (1 - \eta_t \lambda) w_{t-1}$  // weight decay  
// not correctly classified with confidence  
if  $y_i(w \cdot x_i) < 1$   
 $w_t \leftarrow w_t - \eta_t y_i x_i$   
return  $w_T$ 

Note 1: t<sub>0</sub> is a heuristically chosen constant
 Note 2: Learning rate of η<sub>t</sub> = 1/λ(t+t<sub>0</sub>) initially mysterious...

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# Efficient sparse implementation

- We can represent the weight vector w by a tuple (v,s), where v is a vector and s a scalar
- Then the w update is

$$s \leftarrow (1 - \eta_t \lambda) s$$

$$v \leftarrow v - \eta_t y_i x_i$$

Runtime per iteration is proportional to number of non-zero feature values

# SVM-SGD results

- SVM-SGD can be orders of magnitude faster than methods like SVM<sup>light</sup>, SVM<sup>perf</sup>
- Results on ccat data (781,265 examples with 47,152 features):

Algorithm	Training Time	Primal cost	Test Error
SVM <sup>light</sup>	23642 secs	0.2275	6.02%
SVM <sup>perf</sup>	66 secs	0.2278	6.03%
SVM-SGD	1.4 secs	0.2275	6.01%

Table: Results as reported in [Bot07].

# Handling kernels: NORMA

• To apply SGD with a kernel SVM, we notice that, similar to the representer theorem, our learned weight is always of the form

$$w = \sum \alpha_i y_i \Phi(x_i)$$

- $\implies$  We can implicitly represent w by storing the (non-zero)  $\alpha_i$ 's
- Now the update for each example  $(x_{i(t)}, y_{i(t)})$  becomes

$$\begin{split} (\forall 1 \leq j \leq n) \alpha_j \leftarrow (1 - \eta_t \lambda) \alpha_j \\ \alpha_{i(t)} \leftarrow \alpha_{i(t)} - \begin{cases} \eta_t & \text{ if } y_{i(t)}(w_{t-1} \cdot x_{i(t)}) < 1 \\ 0 & \text{ otherwise} \end{cases} \end{split}$$

# Drawbacks of SGD based methods?

- Both methods converge to  $\rho\text{-optimal solution in }O(1/\rho^2)$  iterations
  - $\blacktriangleright$  Slow: usually expect optimizers to converge in e.g.  $O(\log 1/\rho)$  iterations
- Learning rate tuning
  - Common complaint about gradient methods!
- Question: Fundamental limit to what we can do with SGD?
- Answer: No, simple extensions make it powerful

# Pegasos: extending SGD

• The Pegasos solver extends SGD in two ways:

- Aggressively decrease the learning rate: use  $\eta_t = \frac{1}{\lambda t}$ , no parameter sweep required
- ▶ Project the weight vector onto  $\{x : ||x|| \le 1/\sqrt{\lambda}\}$  (stochastic gradient projection)
- Can prove that these changes allow for convergence in  $\tilde{O}\left(\frac{d}{\lambda\rho}\right)$  time
  - $\blacktriangleright$  Inverse dependence on  $\lambda$  accounts for problem difficulty
- Can also work with kernels, similar to NORMA

## Pegasos: the algorithm

for 
$$t = 1 \dots T$$
  
Pick random  $A_t \subseteq \mathcal{T}$  such that  $|A_t| = k$   
// not correctly classified with confidence  
 $\mathcal{M} := \{(x, y) \in A_t : y(w \cdot x) < 1\}$   
 $\nabla_t := \lambda w_t - \frac{1}{|\mathcal{M}|} \sum_{(x,y) \in \mathcal{M}} yx$ 

Update 
$$w_{t+\frac{1}{2}} \leftarrow w_t - \frac{1}{\lambda t} \cdot \nabla_t //$$
 SGD update  
Let  $w_{t+1} \leftarrow \min\left(1, \frac{1}{\sqrt{\lambda} ||w_{t+\frac{1}{2}}||}\right) w_{t+\frac{1}{2}} //$  Projection step

return  $w_{T+1}$ 

• Note: k does not appear in runtime, and so effectively can be chosen to be 1!

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# Pegasos' convergence

- The projection step makes the learning rate  $\propto \frac{1}{t}$  feasible
- The reason the projection makes sense is the following theorem:

#### Theorem

The optimal SVM solution  $w^*$  satisfies  $||w^*|| \leq \frac{1}{\sqrt{\lambda}}$ 

## Proof of theorem

- By the strong duality theorem, the values of the optimal primal and dual solutions are equal
- Rescaling dual problem so that  $\alpha_i \in [0, 1]$ , we get:

$$\frac{\lambda}{2}||w^*||^2 + \frac{1}{n}\sum_{i=1}^n \ell(x_i, y_i; w^*) = \frac{1}{n}\sum_{i=1}^n \alpha_i^* - \frac{1}{2\lambda n^2}\sum_{i,j} \alpha_i^* \alpha_j^* y_i y_j (x_i \cdot x_j).$$

• But by the representer theorem,  $w^* = \sum \alpha_i^* y_i x_i$ :

$$\frac{\lambda}{2}||w^*||^2 + \frac{1}{n}\sum_{i=1}^n \ell(x_i, y_i; w^*) = \frac{||\alpha^*||_1}{n} - \frac{\lambda}{2}||w^*||^2.$$

Rearranging,

$$\lambda ||w^*||^2 = \frac{||\alpha^*||_1 - \sum_{i=1}^n \ell(x_i, y_i; w^*)}{n} \le 1$$

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# Pegasos results

Runtime comparison of Pegasos (in seconds):

Algorithm	Dataset			
	ccat	covertype	astro-ph	
SVM <sup>light</sup>	20,075	25,514	80	
SVM <sup>perf</sup>	77	85	5	
Pegasos	2	6	2	

Table: Results as reported in [SSSS07].

ccat:  $804414 \times 47236$ , 0.16% dense covertype:  $581012 \times 54$ , 22% dense astro-ph:  $62369 \times 99757$ , 0.08% dense

# Stochastic gradient descent: verdict?

- Observed performance of various methods is good
  - In no small part because individual updates are fast
- ${\, \bullet \,}$  But even with Pegasos, convergence rate is only  $1/\rho$ 
  - Not competitive in terms of optimization
- So why is SGD useful for learning?
- **Answer**: Poor optimization does not necessarily mean poor generalization
  - If SGD can optimize "enough", then we can process more examples and get a good generalization
  - Will discuss this more later
- We now quickly look at a couple of recent SGD-based methods

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# Recent work: FOLOS

- FOLOS is a general solver of convex regularized risk minimization problems i.e.  $\ell_{\rm emp}(w) + r(w)$
- Idea is to do SGD, and an analytic minimization (c.f. projection):

$$\begin{split} w_{t+\frac{1}{2}} &= w_t - \eta_t \tilde{\nabla} \ell_{\mathsf{emp}}(w_t) \\ w_{t+1} &= \operatorname*{argmin}_w \, \left( \frac{1}{2} ||w - w_{t+\frac{1}{2}}||^2 + \eta_{t+\frac{1}{2}} r(w) \right) \end{split}$$

• Can be shown that the update is "forward looking", and implicitly imposes the correct regularization term:

$$w_{t+1} = w_t - \eta_t \tilde{\nabla} \ell_{\mathsf{emp}}(w_t) - \eta_{t+\frac{1}{2}} \nabla r(w_{t+1})$$

• Similar update to Pegasos for  $\ell_2$  regularization, discovers sparsity for  $\ell_1$  regularization

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# Recent work: SGD-QN

- SGD-QN combines stochastic gradient descent and quasi-Newton methods
- Instead of using the inverse Hessian H<sup>-1</sup>, use a diagonal scaling matrix D to approximate it

$$w_{t+1} \leftarrow w_t - \eta_t D \cdot \tilde{\nabla}(\ell_{\mathsf{emp}} + r)(w_t)$$

- No theoretical bound or formal experiments
  - Only briefly described as part of an ICML workshop (where it was the winning method!)
  - Unclear whether projection can be replaced (or augmented) with diagonal scaling

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# Dual training methods Dual coordinate descent Bundle method (BMRM)

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# Dual coordinate descent

- We can use coordinate descent to solve the optimization problem in the dual (DCD)
  - Algorithm underlying LibLinear
- Idea is to simply solve the problem using a univariate minimization
  - Pick some  $\alpha_i$ , and hold  $\alpha_1, \ldots, \alpha_{i-1}, \alpha_{i+1}, \ldots$  constant
  - Find the optimal value of  $\alpha_i$  given the other values
- Fortunately, it turns out that the second step is easy to do for SVMs

## Dual coordinate descent

 $\bullet\,$  If we let  $f(\alpha)$  be the dual objective function, then we want to find

 $\min f(\alpha + de_i)$  subject to  $0 \le \alpha_i + d \le C$ 

where  $C = 1/n\lambda$ . This can be solved with the following:

Solution of the minimization Let  $\nabla_i := (\nabla f(\alpha))_i$ . Then, the solution to the univariate minimization is either  $\alpha_i$  or

$$\alpha_i \leftarrow \min(\max(\alpha_i - \nabla_i / ||x_i||^2, 0), C)$$

Further, by the representer theorem, if we explicitly store w,

$$\nabla_i = y_i(w \cdot x_i) - 1$$

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# DCD algorithm

while  $\alpha$  is not optimal Pick an index  $i \in \{1, ..., n\}$  // potentially stochastic  $\alpha_i^{\text{old}} \leftarrow \alpha_i$   $\nabla_i \leftarrow y_i(w \cdot x_i) - 1$  $\nabla_P \leftarrow \begin{cases} \min(\nabla_i, 0) & \text{if } \alpha_i = 0 \\ \max(\nabla_i, 0) & \text{if } \alpha_i = C \\ \nabla_i & \text{otherwise} \end{cases}$ 

$$\begin{split} \text{if } \nabla_P \neq 0 \text{ // check for non-trivial minimizer} \\ \alpha_i \leftarrow \min(\max(\alpha_i - \nabla_i / ||x_i||^2, 0), C) \\ w \leftarrow w + (\alpha_i - \alpha_i^{\text{old}}) y_i x_i \end{split}$$

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# Analyzing DCD

 ${\ensuremath{\, \bullet }}$  We can interpret the update solely in terms of w

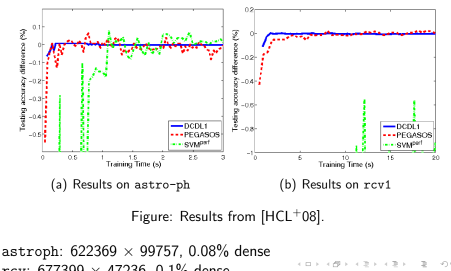
$$w_t = w_{t-1} - (\alpha - \alpha^{\mathsf{old}})y_i x_i$$

- If *i* is chosen randomly, we can think of the algorithm as a form of stochastic gradient descent!
  - Learning rate is chosen via analytic minimization; "optimal" in some sense
  - ► Superior in general to updates for Pegasos? SGD-QN?
- Convergence in batch case in  $O(\log 1/\rho)$  passes over training set
  - Stochastic case is not clear

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# DCD results

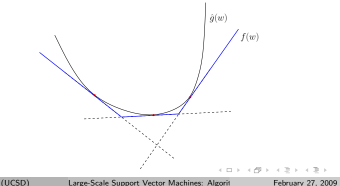
Paper's results indicate it is faster than Pegasos; but some issues about choice of  ${\cal C}$ 



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# Bundle method

- General optimization technique for convex functions: bundle methods
  - Idea is to lower bound a function by an envelope of hyperplanes
  - Regularize solution for stability
- Can apply this idea to SVMs with BMRM (bundle method for risk minimization)
  - Aside: misnomer? Bundle method vs cutting plane



# BMRM update

• Suppose 
$$\hat{g}(w) = \ell_{emp}(w) + r(w)$$

- $f(w) = b_t + \nabla^{(s)} \ell_{emp}(w_t) \cdot w$  defines a hyperplane tangential to  $\hat{g}(w)$  at  $w = w_t$ , where  $\nabla^{(s)}$  denotes a subgradient
- Update the offset  $b_t$  with

$$b_{t+1} = \ell_{\mathsf{emp}}(w_t) - \nabla^{(s)}\ell_{\mathsf{emp}}(w_t) \cdot w_t$$

• The iterates of w are simply taken to be the minimizers of the current approximation:

$$w_{t+1} = \underset{w}{\operatorname{argmin}} \left\{ r(w) + \max_{t' \le t+1} \left[ b_{t'} + (\nabla^{(s)} \ell_{\operatorname{emp}}(w_{t'})) \cdot w \right] \right\}$$

# BMRM update

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$$\hat{g}(w) = \ell_{emp}(w) + r(w)$$

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- Update the offset  $b_t$  with

$$b_{t+1} = \ell_{\mathsf{emp}}(w_t) - \nabla^{(s)} \ell_{\mathsf{emp}}(w_t) \cdot w_t$$

• The iterates of w are simply taken to be the minimizers of the current approximation:

$$w_{t+1} = \underset{w}{\operatorname{argmin}} \left\{ r(w) + \left[ \max_{t' \le t+1} \left[ b_{t'} + (\nabla^{(s)} \ell_{\mathsf{emp}}(w_{t'})) \cdot w \right] \right] \right\}$$

Upper envelope

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# BMRM optimization

- Fortunately, the update admits a simple dual formulation
- Dual bundle problem

The optimization problem for the bundle update is

$$\max_{\alpha} - \frac{1}{2\lambda} \alpha^T Q \alpha + \alpha \cdot b \text{ such that } ||\alpha||_1 = 1, \alpha_i \ge 0,$$

where  $Q_{ij} = \nabla^{(s)} \ell_{emp}(w_i) \cdot \nabla^{(s)} \ell_{emp}(w_j)$ .

- This is a problem whose size at iteration t is  $t \times t$ 
  - Worst case,  $t = O(1/\rho)$ ; and for smooth  $\ell_{emp}$ ,  $t = O(\log 1/\rho)$
  - ► Conjecture that an average of piecewise linear functions (e.g. SVMs) also has roughly t = O(log 1/ρ)!
- Can be solved with a QP solver, or with a line search

# BMRM results

# Results only presented for reducing objective value; not as informative as test error

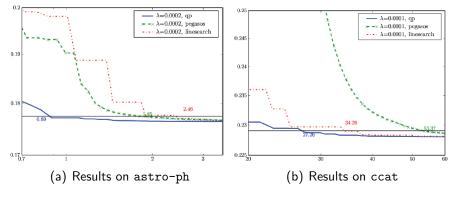


Figure: Objective value vs time results from [SVL07].

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# BMRM and SVM<sup>perf</sup>

- Do the results comparing BMRM and Pegasos contradict those in the Pegasos paper?
  - ► Authors claim SVM<sup>perf</sup> can be seen as a special case of BMRM
- So is SVM<sup>perf</sup> superior to Pegasos?
  - Should compare their generalization ability rather than optimization
  - But ccat results are surprising; Pegasos greatly outperformed till ~ 50 seconds? Hard to imagine Pegasos has comparable generalization!
  - Disappointingly, authors do not discuss this issue at all
- Issue is potentially moot...results in OCAS contradict the ones here!

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# Extension: OCAS

- OCAS is an extension of the BMRM approach
- $\bullet$  Recall that we try to minimize the regularized risk  $\hat{g}(w)$  with a lower envelope f(w)
- The BMRM iterates are guaranteed to satisfy

 $f(w_{t+1}) < f(w_t)$  but **not**  $\hat{g}(w_{t+1}) < \hat{g}(w_t)$ 

- Some iterates are undesirable
- ullet OCAS ensures monotonicity of  $\hat{g}(w_t)$  by a line search
  - Keeps track of best solution, and combines this with current iterate
  - ► Aside: truer bundle method, but calls itself cutting plane!
- Discusses potential for parallelization

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# Why is SGD successful?

- We've seen SVM training algorithms based on SGD
- Advantage: faster processing of each example
- Disadvantage: slower convergence in general
- **Question**: Doesn't the slow convergence rate seriously hamper its viability?

# Why is SGD successful?

- We've seen SVM training algorithms based on SGD
- Advantage: faster processing of each example
- Disadvantage: slower convergence in general
- **Question**: Doesn't the slow convergence rate seriously hamper its viability?
- (Surprising) Answer: Not if we look at the optimization process more closely

# Learning and optimization

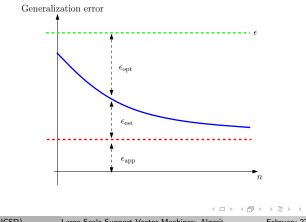
- We have studied SVMs through their optimization problem
  - $\blacktriangleright$  In particular, we looked at runtime as a function of  $\rho$
- But recall that optimization is only a surrogate for generalization
- We hope that minimizing the training error will minimize the generalization error
- Question: Is it necessary to perform strenuous optimization?

# The three components of error

- [BB07] explicitly considers the role of optimization error
- $\bullet\,$  Suppose that we obtain a weight  $\tilde{w}$  using an optimization algorithm run to some finite tolerance
- They decompose the generalization error  $\epsilon(\tilde{w}) = \mathbb{E}[\ell(x,y;\tilde{w})]$  into three components
  - Approximation error  $\epsilon_{app}$ : minimum error due to hypothesis class
  - Estimation error  $\epsilon_{\text{est}}$ : minimum error due to training set
  - Optimization error  $\epsilon_{opt}$ : minimum error due to optimization

# Minimizing generalization error

- If  $n \to \infty$ , then  $\epsilon_{\text{est}} \to 0$
- **Implication**: For a fixed generalization error  $\epsilon$ , as *n* increases we can increase the optimization tolerance  $\rho$



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# The role of estimation error

- We look at the behaviour of estimation error as n gets large
- $\bullet\,$  This lets us connect the behaviour of n and  $\rho$  to the generalization error  $\epsilon\,$

Estimation error bound

Let  $\epsilon^{\ast}$  denote the minimum possible generalization error. Then,

$$0 \le \epsilon - \epsilon^* \le c \cdot \left(\epsilon_{\mathsf{app}} + \frac{d}{n} \log \frac{n}{d} + \rho\right)$$

• Intuition: Estimation error behaves like  $\log n/n$ 

# The role of estimation error

• Now let's bound the individual terms by  $\mathcal{E}$ , the excess error w.r.t. the approximation error:

$$\rho = \Theta(\mathcal{E})$$

$$n = \Theta(d\log(1/\mathcal{E})/\mathcal{E})$$

That implies

$$\epsilon - \epsilon^* \le c \cdot (\epsilon_{\mathsf{app}} + \mathcal{E})$$

• So, we have a way to compare convergence to the generalization optimum

# Estimation error and SGD

• How quickly do GD and SGD get a bound of  $c \cdot (\epsilon_{app} + \mathcal{E})$ ?

Algorithm	Optimization time	Generalization time
GD	$O\left(nd\log\frac{1}{\rho}\right)$	$O\left(\frac{d^2}{\mathcal{E}} \log^2 \frac{1}{\mathcal{E}}\right)$
SGD	$O\left(\frac{d}{\rho}\right)$	$O\left(\frac{d}{\mathcal{E}}\right)$
2GD	$O\left((d^2 + nd)\log\log\frac{1}{\rho}\right)$	$O(\frac{d^2}{\mathcal{E}}\log\frac{1}{\mathcal{E}}\log\log\frac{1}{\mathcal{E}})$
2SGD	$O\left(\frac{d^2}{\rho}\right)$	$O\left(\frac{d^2}{\mathcal{E}}\right)$

- **Conclusion**: SGD generalizes asymptotically faster than GD!
- Key point: SGD's runtime does not depend on the number of examples

# Applying to SVMs

- What implications does this have for SVMs?
  - ► Note: minimizing a regularized loss term
- SGD-based solvers (e.g. Pegasos) should be able to leverage decreased estimation error
  - But can we more accurately characterize this?

# Applying to SVMs

- What implications does this have for SVMs?
  - ► Note: minimizing a regularized loss term
- SGD-based solvers (e.g. Pegasos) should be able to leverage decreased estimation error
  - But can we more accurately characterize this?
- (Very surprising) Implication: The training time should decrease as the number of examples increases

# SVM generalization

- Given more examples, SVM training time should ideally not increase if we want the same generalization error
  - Suppose in time t, we achieve 5% generalization error with 10,000 examples
  - ▶ With 100,000 examples, we can sample to get the same error
- But [SSS08] goes a step further
  - Argues that as number of examples increases, runtime should decrease as a result of decreased estimation error
  - ► That is, it takes us < t time to get 5% generalization error with 100,000 examples

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# SVM generalization bound

• Foundation of the analysis is the following theorem

#### Theorem

Let w be the learned weight vector when SVM optimization is done up to tolerance  $\rho$ , and let  $w_0$  be any other weight vector. Then,

$$g(w) \le g(w_0) + 2\rho + \frac{\lambda}{2} ||w_0||^2 + \tilde{O}(1/\lambda n),$$

where g(w) is the generalization error with weight vector w,

$$g(w) = \mathbb{E}_{(x,y) \sim P(\mathcal{X},\mathcal{Y})}[\ell(x,y;w)]$$

# Proof of generalization bound

- Let  $f(w):=g(w)+\frac{\lambda}{2}||w||^2$  denote the regularized generalization error
- ${\ensuremath{\, \circ }}$  Now decompose g(w) as

$$g(w) = f(w) - \frac{\lambda}{2} ||w||^2$$
  
=  $f(w) - \frac{\lambda}{2} ||w||^2 - \left(f(w_0) - g(w_0) - \frac{\lambda}{2} ||w_0||^2\right)$   
=  $g(w_0) + (f(w) - f(w_0)) + \frac{\lambda}{2} ||w_0||^2 - \frac{\lambda}{2} ||w||^2$   
=  $g(w_0) + (f(w) - f(w^*)) + (f(w^*) - f(w_0))$   
+  $\frac{\lambda}{2} ||w_0||^2 - \frac{\lambda}{2} ||w||^2.$ 

where  $w^* = \operatorname{argmin} f(w)$ .

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# Proof of generalization bound contd.

- Recall that  $\hat{g}(w)$  is the SVM training error
- Second term is a difference of expected losses, can be bounded using the corresponding empirical losses:

$$\begin{split} f(w) - f(w^*) &\leq 2 \max(0, \hat{g}(w) - \hat{g}(w^*)) + O\left(\frac{\log 1/\delta}{\lambda n}\right) \\ &= 2\rho + O\left(\frac{\log 1/\delta}{\lambda n}\right) \text{ by definition of } w \end{split}$$

- $f(w^*) f(w_0) \le 0$  by the optimality of  $w^*$
- Combining these facts,

$$g(w) \le g(w_0) + 2\rho + \frac{\lambda}{2} ||w_0||^2 + \tilde{O}(1/\lambda n)$$

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# Applying the bound

- We can use this bound to find  $T(n, \mathcal{E})$ , the time needed for training with n examples to get excess error  $\mathcal{E}$
- Rewrite  $\rho$  in terms of  $T, \lambda$ : e.g. for Pegasos,

$$g(w) \le g(w_0) + \tilde{O}(d/\lambda T) + \frac{\lambda}{2} ||w_0||^2 + \tilde{O}(1/\lambda n)$$

• Choosing  $\lambda$  to minimize this,

$$g(w) \le g(w_0) + \tilde{O}(||w_0||\sqrt{d/T}) + O(||w_0||/\sqrt{n}) = g(w_0) + \mathcal{E}$$

• Now express T as a function of n and  $\mathcal{E}$ ...

# Pegasos and SVM<sup>perf</sup>

- Easy to get runtime bounds for Pegasos and SVMperf •
- Pegasos:

$$T(n, \mathcal{E}) = \tilde{O}\left(\frac{d}{\left(\mathcal{E}/||w_0|| - O(1/\sqrt{n})\right)^2}\right)$$

• SVM<sup>perf.</sup>

$$T(n,\mathcal{E}) = O\left(\frac{nd}{\left(\mathcal{E}/||w_0|| - O(1/\sqrt{n})\right)^2}\right)$$

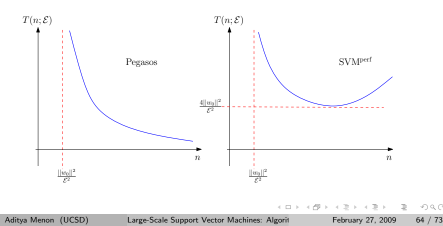
• Implications?

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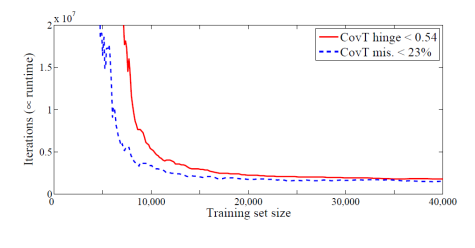
# Pegasos and $\mathsf{SVM}^{\mathsf{perf}}$

- Pegasos' runtime monotonically decreases as a function of n!
- SVM<sup>perf</sup> has a turning point, but after that the runtime increases
  - Turning point where decrease in estimation is offset by increase in iteration cost



# Verifying Pegasos' runtime

Results on covertype:



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# Summary

- We have seen several solvers for SVMs targetting large training sets
  - Primal methods based on SGD
  - Dual solvers based on optimization "tricks"
- Saw why SGD is poor at optimization but good at generalization
  - Runtime agnostic about number of examples
  - Still manages to leverage decreased estimation error
- Runtime for Pegasos decreases with increase in training set size
  - Fundamental limit to dual QP methods?
  - ► SGD for dual c.f. DCD?

# Comparison of methods?

- Experimental comparisons by no means comprehensive
  - ► Pegasos' results directly contradict those in BMRM and OCAS!
  - Insufficient detail on parameter choices
- Motivation for ICML workshop
  - ► SGD-QN performed well, but so did some batch algorithms
  - Optimized interior point method did extremely well!
- Role of loading time
  - If method stores training set in memory, loading time is usually the bottleneck!
  - Online algorithms mix parsing and learning (e.g. Vowpal Wabbit)

# Comment on plausibility

- **Question**: Is supervised learning realistic when the training set is very large?
  - In some domains like bioinformatics, labelling examples is expensive; impossible to completely label a large training set
  - A more realistic setting is semi-supervised learning
- **Answer**: Not always, but there are domains where large training sets are completely labelled
  - Labelling may be a natural byproduct of user actions e.g. Google ad clicks
  - Large user-bases can be leveraged/ "tricked" into doing the labelling e.g. reCAPTCHA
- Nonetheless, (large-scale) semi-supervised SVMs is an important future direction

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# Other research directions

- Other approaches to large-scale SVMs (and learning)
  - ► Parallelization e.g. Cascade SVM, OCAS
  - ► Training set reduction, e.g. active learning, clustering
- Stochastic gradient for kernels?
  - NORMA was more interested in moving target setting; no comparison to other methods
  - Caching? Storing truncated kernel in memory?
- SVMs when data does not fit in memory
  - Completely precludes batch algorithms; incremental SVMs?
- Multi-class SVMs
  - In domains where we might expect large-scale data to arise naturally, classification is usually more complex than binary
  - Need efficient multi-class SVMs
  - ► Some nascent work e.g. LaRank

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#### Questions?

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