# Bundle Methods for Regularized Risk Minimization

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### Abstract

A wide variety of machine learning problems can be described as minimizing a regularized risk functional, with different algorithms using different notions of risk and different regularizers. Examples include linear Support Vector Machines (SVMs), Gaussian Processes, Logistic Regression, Conditional Random Fields (CRFs), and Lasso amongst others. This paper describes the theory and implementation of a scalable and modular convex solver which solves all these estimation problems. It can be parallelized on a cluster of workstations, allows for data-locality, and can deal with regularizers such as  $L_1$  and  $L_2$  penalties. In addition to the unified framework we present tight convergence bounds, which show that our algorithm converges in  $O(1/\epsilon)$  steps to  $\epsilon$  precision for general convex problems and in  $O(\log(1/\epsilon))$  steps for continuously differentiable problems. We demonstrate the performance of our general purpose solver on a variety of publicly available datasets.

**Keywords:** Optimization, Subgradient methods, Cutting plane method, Bundle methods, regularized risk minimization, parallel optimization

### 1. Introduction

At the heart of many machine learning algorithms is the problem of minimizing a regularized risk functional. That is, one would like to solve

$$\min_{w} J(w) := \lambda \Omega(w) + R_{\text{emp}}(w), \tag{1a}$$

where 
$$R_{\text{emp}}(w) := \frac{1}{m} \sum_{i=1}^{m} l(x_i, y_i, w)$$
 (1b)

is the empirical risk. Moreover,  $x_i \in \mathcal{X} \subseteq \mathbb{R}^d$  are referred to as training instances and  $y_i \in \mathcal{Y}$  are the corresponding labels. l is a (surrogate) convex loss function measuring the discrepancy between y and the predictions arising from using w. For instance, w might enter our model via  $l(x, y, w) = (\langle w, x \rangle - y)^2$ , where  $\langle \cdot, \cdot \rangle$  denotes the standard Euclidean dot product. Finally,  $\Omega(w)$  is a convex function serving the role of a regularizer with regularization constant  $\lambda > 0$ . Typically  $\Omega$  is differentiable and cheap to compute. In contrast, the empirical risk term  $R_{\text{emp}}(w)$  is often non-differentiable, and almost always computationally expensive to deal with.

For instance, if we consider the problem of predicting binary valued labels  $y \in \{\pm 1\}$ , we can set  $\Omega(w) = \frac{1}{2} \|w\|_2^2$  (i.e.,  $L_2$  regularization), and the loss l(x, y, w) to be the binary hinge loss,  $\max(0, 1 - y \langle w, x \rangle)$ , thus recovering linear Support Vector Machines (SVMs) (Joachims, 2006). Using the same regularizer but changing the loss function to  $l(x, y, w) = \log(1 + \exp(-y \langle w, x \rangle))$ , yields logistic regression. Extensions of these loss functions allow us to handle structure in the output space (Bakir et al., 2007) (also see Appendix A for a comprehensive exposition of many common loss functions). On the other hand, changing the regularizer  $\Omega(w)$  to the sparsity inducing  $\|w\|_1$  (i.e.,  $L_1$  regularization) leads to Lasso-type estimation algorithms (Mangasarian, 1965; Tibshirani, 1996; Candes and Tao, 2005).

If the objective function J is differentiable, for instance in the case of logistic regression, we can use smooth optimization techniques such as the standard quasi-Newtons methods like BFGS or its limited memory variant LBFGS (Nocedal and Wright, 1999). These methods are effective and efficient even when m and d are large (Sha and Pereira, 2003; Minka, 2007). However, it is not straightforward to extend these algorithms to optimize a non-differentiable objective, for instance, when dealing with the binary hinge loss (see e.g. Yu et al., 2008).

When J is non-differentiable, one can use nonsmooth convex optimization techniques such as the cutting plane method (Kelly, 1960) or its stabilized version the bundle method (Hiriart-Urruty and Lemaréchal, 1993). The amount of external stabilization that needs to be added is a parameter that requires careful tuning. In this paper, we bypass this problem by noticing that the regularized risk minimization problem (1) already contains a stabilization term, that is, the regularizer  $\Omega(w)$ . Therefore there is no need to add an external stabilizer or to tune it. The resultant algorithm – Bundle Methods for Regularized Risk Minimization (BMRM) – has certain desirable properties: a) it has no parameters to tune, and b) it is applicable to a wide variety of regularized risk minimization problems. Furthermore, we show that BMRM has an  $O(1/\epsilon)$  rate of convergence for nonsmooth problems and  $O(\log(1/\epsilon))$  for smooth problems. This is significantly tighter than the  $O(1/\epsilon^2)$  rates provable for standard bundle methods (Lemaréchal et al., 1995) and a closely related method SVM<sup>struct</sup> (Tsochantaridis et al., 2005). Very briefly, we highlight the two major advantages of our implementation. First, it is completely modular; new loss functions, regularizers, and solvers can be added with relative ease. Second, our architecture allows the empirical risk computation (1b) to be easily parallelized. This makes our solver amenable to large datasets which cannot fit into the memory of a single computer. Our open source C/C++ implementation is freely available for download from http://users.rsise.anu.edu.au/~chteo/BMRM.html.

The outline of our paper is as follows. In Section 2 we describe BMRM and contrast it with standard bundle methods. We also prove rates of convergence. In section 3 we discuss implementation issues and present principled techniques to control memory usage, as well

as to speed up computation via parallelization. Section 4 puts our work in perspective, and discusses related work. Section 5 is devoted to extensive experimental evaluation, which shows that our implementation is comparable to or better than specialized state-of-the-art solvers on a number of publicly available datasets. Finally, we conclude our work and discuss related issues in section 6. In Appendix A we describe various *classes* of loss functions organized according to their common traits in computation. Long proofs are relegated to Appendix B. Before we proceed a brief note about our notation:

**Notation** The indices of elements of a sequence or a set appear in subscript e.g.  $u_1, u_2$ . The *i*-th component of a vector u is denoted by  $u^{(i)}$ . [k] is the shorthand for the set  $\{1, 2, \ldots, k\}$ . The  $L_p$  norm is defined as  $||u||_p = (\sum_{i=1}^d |u^{(i)}|^p)^{1/p}$ , for  $p \ge 1$ , and we use  $||\cdot||$  to denote  $||\cdot||_2$  whenever the context is clear.  $\mathbf{1}_d$  and  $\mathbf{0}_d$  denote the d-dimensional vectors of all ones and zeros respectively.

### 2. Bundle Methods

The precursor to the bundle methods is the cutting plane method (CPM) (Kelly, 1960). CPM uses subgradients, which are a generalization of gradients appropriate for convex functions, including those which are not necessarily smooth. Suppose w' is a point where a convex function J is finite. Then a subgradient is the normal vector of any tangential supporting hyperplane of J at w' (see Figure 1 for geometric intuition). Formally s' is called a subgradient of J at w' if, and only if,

$$J(w) \ge J(w') + \langle w - w', s' \rangle \quad \forall w. \tag{2}$$

The set of all subgradients at w' is called the subdifferential, and is denoted by  $\partial_w J(w')$ . If this set is not empty then J is said to be *subdifferentiable at* w'. On the other hand, if this set is a singleton then the function is said to be *differentiable* at w'. Convex functions are subdifferentiable everywhere in their domain (Hiriart-Urruty and Lemaréchal, 1993).

As implied by (2), J is bounded from below by its linearization (*i.e.*, first order Taylor approximation) at w'. Given subgradients  $s_1, s_2, \ldots, s_t$  evaluated at locations  $w_0, w_1, \ldots, w_{t-1}$ , we can state a tighter (piecewise linear) lower bound for J as follows

$$J(w) \ge J_t^{\text{CP}}(w) := \max_{1 \le i \le t} \{ J(w_{i-1}) + \langle w - w_{i-1}, s_i \rangle \}.$$
 (3)

This lower bound forms the basis of the CPM, where at iteration t the set  $\{w_i\}_{i=0}^{t-1}$  is augmented by

$$w_t := \underset{w}{\operatorname{argmin}} J_t^{\operatorname{CP}}(w). \tag{4}$$

This iteratively refines the piecewise linear lower bound  $J^{\text{CP}}$  and allows us to get close to the minimum of J (see Figure 2 for an illustration).

If  $w^*$  denotes the minimizer of J, then clearly each  $J(w_i) \geq J(w^*)$  and hence  $\min_{0 \leq i \leq t} J(w_i) \geq J(w^*)$ . On the other hand, since  $J \geq J_t^{\text{CP}}$  it follows that  $J(w^*) \geq J_t^{\text{CP}}(w_t)$ . In other words,  $J(w^*)$  is sandwiched between  $\min_{0 \leq i \leq t} J(w_i)$  and  $J_t^{\text{CP}}(w_t)$  (see Figure 3 for an illustration). The CPM monitors the monotonically decreasing quantity

$$\epsilon_t := \min_{0 \le i \le t} J(w_i) - J_t^{\text{CP}}(w_t), \tag{5}$$

and terminates whenever  $\epsilon_t$  falls below a predefined threshold  $\epsilon$ . This ensures that the solution  $J(w_t)$  satisfies  $J(w_t) \leq J(w^*) + \epsilon$ .

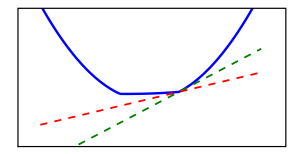


Figure 1: Geometric intuition of a subgradient. The nonsmooth convex function (solid blue) is only subdifferentiable at the "kink" points. We illustrate two of its subgradients (dashed green and red lines) at a "kink" point which are tangential to the function. The normal vectors to these lines are subgradients.

### 2.1 Standard Bundle Methods

Although CPM was shown to be convergent (Kelly, 1960), it is well known (see e.q.Lemaréchal et al., 1995; Belloni, 2005) that CPM can be very slow when new iterates move too far away from the previous ones (i.e., causing unstable "zig-zag" behavior in the iterates).

Bundle methods stabilize CPM by augmenting the piecewise linear lower (e.g.  $J_t^{\text{CP}}(w)$ as in (3)) with a prox-function (i.e., proximity control function) which prevents overly large steps in the iterates (Kiwiel, 1990). Roughly speaking, there are 3 popular types of bundle methods, namely, proximal (Kiwiel, 1990), trust region (Schramm and Zowe, 1992), and level set (Lemaréchal et al., 1995). All three versions use  $\frac{1}{2} \| \cdot \|^2$  as their prox-function, but differ in the way they compute the new iterate:

proximal: 
$$w_t := \underset{w}{\operatorname{argmin}} \{ \frac{\zeta_t}{2} \| w - \hat{w}_{t-1} \|^2 + J_t^{CP}(w) \},$$
 (6)

proximal: 
$$w_t := \underset{w}{\operatorname{argmin}} \{ \frac{\zeta_t}{2} \| w - \hat{w}_{t-1} \|^2 + J_t^{\operatorname{CP}}(w) \},$$
 (6)  
trust region:  $w_t := \underset{w}{\operatorname{argmin}} \{ J_t^{\operatorname{CP}}(w) \mid \frac{1}{2} \| w - \hat{w}_{t-1} \|^2 \le \kappa_t \},$  (7)

level set: 
$$w_t := \underset{w}{\operatorname{argmin}} \{ \frac{1}{2} \| w - \hat{w}_{t-1} \|^2 \mid J_t^{\operatorname{CP}}(w) \le \tau_t \},$$
 (8)

where  $\hat{w}_{t-1}$  is the current prox-center, and  $\zeta_t, \kappa_t$ , and  $\tau_t$  are positive trade-off parameters of the stabilization. Although (6) can be shown to be equivalent to (7) for appropriately chosen  $\zeta_t$  and  $\kappa_t$ , tuning  $\zeta_t$  is rather difficult while a trust region approach can be used for automatically tuning  $\kappa_t$ . Consequently the trust region algorithm BT of Schramm and Zowe (1992) is widely used in practice.

<sup>1.</sup> For brevity we will only describe "first-order" bundle methods, and omit discussion about "second-order" variants such as the bundle-Newton method of Lukšan and Vlček (1998).

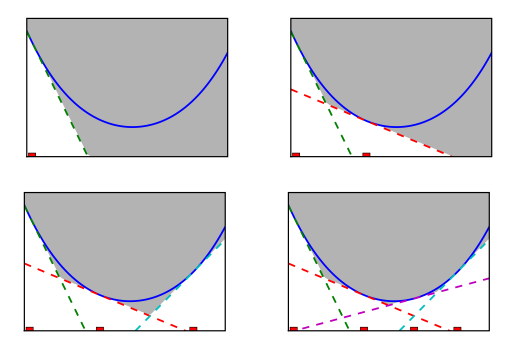


Figure 2: A convex function (blue solid curve) is bounded from below by its linearizations (dashed lines). The gray area indicates the piecewise linear lower bound obtained by using the linearizations. We depict a few iterations of the cutting plane method. At each iteration the piecewise linear lower bound is minimized and a new linearization is added at the minimizer (red rectangle). As can be seen, adding more linearizations improves the lower bound.

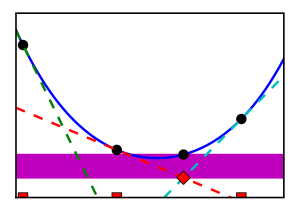


Figure 3: A convex function (blue solid curve) with three linearizations (dashed lines) evaluated at three different locations (red squares). The approximation gap  $\epsilon_3$  at the end of third iteration is indicated by the height of the magenta horizontal band *i.e.*, difference between lowest value of J(w) evaluated so far (lowest black circle) and the minimum of  $J_3^{\text{CP}}(w)$  (red diamond).

Since our methods (see Section 2.2) are closely related to the proximal bundle method, we will now describe them in detail. Similar to the CPM the proximal bundle method also builds a piecewise linear lower bound  $J_t^{\text{CP}}$  (see (3)). In contrast to the CPM, the piecewise linear lower bound augmented with a stabilization term  $\frac{\zeta_t}{2} ||w - \hat{w}_{t-1}||^2$ , is minimized to produce the intermediate iterate  $\bar{w}_t$ . The approximation gap in this case includes the proxfunction:

$$\epsilon_t := J(\hat{w}_{t-1}) - \left[ J_t^{\text{CP}}(\bar{w}_t) + \frac{\zeta_t}{2} \|\bar{w}_t - \hat{w}_{t-1}\|^2 \right].$$
(9)

If  $\epsilon_t$  is less than the pre-defined threshold  $\epsilon$  the algorithm exits. Otherwise, a line search is performed along the line joining  $\hat{w}_{t-1}$  and  $\bar{w}_t$  to produce the new iterate  $w_t$ . If  $w_t$  results in a sufficient decrease of the objective function then it is accepted as the new prox-center  $\hat{w}_t$ ; this is called a serious step. Otherwise, the prox-center remains the same; this is called a null step. Detailed pseudocode can be found in Algorithm 1.

If the approximation gap  $\epsilon_t$  (9) is smaller than  $\epsilon$ , then this ensures that the solution  $J(\hat{w}_{t-1})$  satisfies  $J(\hat{w}_{t-1}) \leq J(w) + \frac{\zeta_t}{2} \|w - \hat{w}_{t-1}\|^2 + \epsilon$  for all w. In particular, if  $J(w^*)$  denotes the optimum as before, then  $J(\hat{w}_{t-1}) \leq J(w^*) + \frac{\zeta_t}{2} \|w^* - \hat{w}_{t-1}\|^2 + \epsilon$ . Contrast this with the approximation guarantee of the CPM, which does not involve the  $\frac{\zeta_t}{2} \|w^* - \hat{w}_{t-1}\|^2$  term.

Although the positive coefficient  $\zeta_t$  is assumed fixed throughout the algorithm, in practice it must be updated after every iteration to achieve faster convergence, and to guarantee a good quality solution (Kiwiel, 1990). Same is the case for  $\kappa_t$  and  $\tau_t$  in trust region and level set bundle methods, respectively. Although the update is not difficult, the procedure relies on other parameters which require careful tuning (Kiwiel, 1990; Schramm and Zowe, 1992;

Lemaréchal et al., 1995). We now describe our method (BMRM) which avoids this problem by exploiting the fact that the objective function of interest to us (1) already contains a regularizer  $\Omega(w)$  which can be used to stabilize the CPM. As we will see, not only is the implementation straightforward, but the rates of convergence also improve from  $O(1/\epsilon^3)$  or  $O(1/\epsilon^2)$  to  $O(1/\epsilon)$ .

# Algorithm 1 Proximal Bundle Method

```
1: input & initialization: \epsilon \geq 0, \ \rho \in (0,1), \ w_1, \ t \leftarrow 0, \ \hat{w}_0 \leftarrow w_0
  2: loop
            t \leftarrow t + 1
  3:
            Compute J(w_{t-1}) and s_t \in \partial_w J(w_{t-1})
  4:
           Update model J_t^{\text{CP}}(w) := \max_{1 \le i \le t} \{J(w_{i-1}) + \langle w - w_{i-1}, s_i \rangle\}

\bar{w}_t \leftarrow \operatorname{argmin}_w J_t^{\text{CP}}(w) + \frac{\zeta_t}{2} \|w - \hat{w}_{t-1}\|^2
           \epsilon_t \leftarrow J(\hat{w}_{t-1}) - \left[J_t^{\text{CP}}(\bar{w}_t) + \frac{\zeta_t}{2} \|\bar{w}_t - \hat{w}_{t-1}\|^2\right]
if \epsilon_t < \epsilon then return \bar{w}_t
  7:
            Linesearch: \eta_t \leftarrow \operatorname{argmin}_{\eta \in \mathbb{R}} J(\hat{w}_{t-1} + \eta(\bar{w}_t - \hat{w}_{t-1}))
                                                                                                                              (if expensive, set \eta_t = 1)
 9:
            w_t \leftarrow \hat{w}_{t-1} + \eta_t(\bar{w}_t - \hat{w}_{t-1})
10:
11:
            if J(\hat{w}_{t-1}) - J(w_t) \ge \rho \epsilon_t then
                SERIOUS STEP: \hat{w}_t \leftarrow w_t
12:
13:
            else
                 NULL STEP: \hat{w}_t \leftarrow \hat{w}_{t-1}
14:
            end if
15:
16: end loop
```

#### 2.2 Bundle Methods for Regularized Risk Minimization (BMRM)

Define:

(subgradient of 
$$R_{\text{emp}}$$
)  $a_t \in \partial_w R_{\text{emp}}(w_{t-1})$  (10)

(offset) 
$$b_t := R_{\text{emp}}(w_{t-1}) - \langle w_{t-1}, a_t \rangle \tag{11}$$

(piecewise linear lower bound of 
$$R_{\text{emp}}$$
)  $R_t^{\text{CP}}(w) := \max_{1 \le i \le t} \langle w, a_i \rangle + b_i$  (12)

(piecewise convex lower bound of 
$$J$$
)  $J_t(w) := \lambda \Omega(w) + R_t^{CP}(w)$  (13)

(iterate) 
$$w_t := \min_{w} J_t(w)$$
 (14)

(approximation gap) 
$$\epsilon_t := \min_{0 \le i \le t} J(w_i) - J_t(w_t)$$
 (15)

We now describe BMRM (Algorithm 2), and contrast it with the proximal bundle method. At iteration t the algorithm builds the lower bound  $R_t^{\text{CP}}$  to the empirical risk  $R_{\text{emp}}$ . The new iterate  $w_t$  is then produced by minimizing  $J_t$  which is  $R_t^{\text{CP}}$  augmented with the regularizer  $\Omega$ ; this is the key difference from the proximal bundle method which uses the  $\frac{\zeta_t}{2} ||w - \hat{w}_{t-1}||^2$  prox-function for stabilization. The algorithm repeats until the approximation gap  $\epsilon_t$  is less than the pre-defined threshold  $\epsilon$ . Unlike standard bundle methods there is no notion of a serious or null step in our algorithm. In fact, our algorithm does not even maintain a prox-center. It can be viewed as a special case of standard bundle methods where the prox-center

is always the origin and never updated (hence every step is a null step). Furthermore, unlike the proximal bundle method, the approximation guarantees of our algorithm do not involve the  $\frac{\zeta_t}{2} \|w^* - w_t\|^2$  term.

# Algorithm 2 BMRM

```
1: input & initialization: \epsilon \geq 0, w_0, t \leftarrow 0

2: repeat

3: t \leftarrow t + 1

4: Compute a_t \in \partial_w R_{\text{emp}}(w_{t-1}) and b_t \leftarrow R_{\text{emp}}(w_{t-1}) - \langle w_{t-1}, a_t \rangle

5: Update model: R_t^{\text{CP}}(w) := \max_{1 \leq i \leq t} \langle w, a_i \rangle + b_i

6: w_t \leftarrow \operatorname{argmin}_w J_t(w) := \lambda \Omega(w) + R_t^{\text{CP}}(w)

7: \epsilon_t \leftarrow \min_{0 \leq i \leq t} J(w_i) - J_t(w_t)

8: until \epsilon_t \leq \epsilon

9: return w_t
```

Algorithm 2 is simple and easy to implement as it does not involve a line search. In fact, whenever efficient (exact) line search is available, it can be used to achieve faster convergence as observed by Franc and Sonnenburg (2008) in the case of linear SVMs with binary hinge loss.<sup>2</sup> We now turn to a variant of BMRM which uses a line search (Algorithm 3); this is a generalization of the optimized cutting plane algorithm for support vector machines (OCAS) of Franc and Sonnenburg (2008). This variant first builds  $R_t^{\text{CP}}$  and minimizes  $J_t$  to obtain an intermediate iterate  $w_t$ . Then, it performs a line search along the line joining  $w_{t-1}^b$  and  $w_t$  to produce  $w_t^b$  which acts like the new prox-center. Note that  $w_t - w_{t-1}^b$  is not necessarily a direction of descent; therefore the line search might return a zero step. Instead of using  $w_t^b$  as the new iterate the algorithm uses the pre-set parameter  $\theta$  to generate  $w_t^c$  on the line segment joining  $w_t^b$  and  $w_t$ . Franc and Sonnenburg (2008) report that setting  $\theta = 0.9$  works well in practice. It is easy to see that Algorithm 3 reduces to Algorithm 2 if we set  $\eta_t = 1$  for all t, and use the same termination criterion.

# Algorithm 3 BMRM with Line Search

```
1: input & initialization: \epsilon \geq 0, \ \theta \in (0,1], \ w_0^b, \ w_0^c \leftarrow w_0^b, \ t \leftarrow 0

2: repeat

3: t \leftarrow t+1

4: Compute a_t \leftarrow \partial_w R_{\text{emp}}(w_{t-1}^c), and b_t \leftarrow R_{\text{emp}}(w_{t-1}^c) - \left\langle w_{t-1}^c, a_t \right\rangle

5: Update model: R_t^{\text{CP}}(w) := \max_{1 \leq i \leq t} \left\langle w, a_i \right\rangle + b_i

6: w_t \leftarrow \operatorname{argmin}_w J_t(w) := \lambda \Omega(w) + R_t^{\text{CP}}(w)

7: Linesearch: \eta_t \leftarrow \operatorname{argmin}_{\eta \in \mathbb{R}} J(w_{t-1}^b + \eta(w_t - w_{t-1}^b))

8: w_t^b \leftarrow w_{t-1}^b + \eta_t(w_t - w_{t-1}^b)

9: w_t^c \leftarrow (1 - \theta)w_t^b + \theta w_t

10: \epsilon_t \leftarrow J(w_t^b) - J_t(w_t)

11: until \epsilon_t \leq \epsilon
```

<sup>2.</sup> A different optimization method but with identical efficient line search procedure is described in Yu et al. (2008).

### 2.3 Dual Problems

In this section, we describe how the sub-problem

$$w_t = \underset{w}{\operatorname{argmin}} J_t(w) := \lambda \Omega(w) + \max_{1 \le i \le t} \langle w, a_i \rangle + b_i$$
 (16)

in Algorithms 2 and 3 is solved via a dual formulation. In fact, we will show that we need not know  $\Omega(w)$  at all, instead it is sufficient to work with its Fenchel dual (Hiriart-Urruty and Lemaréchal, 1993):

**Definition 1 (Fenchel Dual)** Denote by  $\Omega : \mathcal{W} \to \mathbb{R}$  a convex function on a convex set  $\mathcal{W}$ . Then the dual  $\Omega^*$  of  $\Omega$  is defined as

$$\Omega^*(\mu) := \sup_{w \in \mathcal{W}} \langle w, \mu \rangle - \Omega(w). \tag{17}$$

Several choices of regularizers are common. For  $\mathcal{W} = \mathbb{R}^d$  the squared norm regularizer yields

$$\Omega(w) = \frac{1}{2} \|w\|_2^2$$
 and  $\Omega^*(\mu) = \frac{1}{2} \|\mu\|_2^2.$  (18)

More generally, for  $L_p$  norms one obtains (Boyd and Vandenberghe, 2004; Shalev-Shwartz and Singer, 2006):

$$\Omega(w) = \frac{1}{2} \|w\|_p^2$$
 and  $\Omega^*(\mu) = \frac{1}{2} \|\mu\|_q^2$  where  $\frac{1}{p} + \frac{1}{q} = 1$ . (19)

For any positive definite matrix B, we can construct a quadratic form regularizer which allows non-uniform penalization of the weight vector as:

$$\Omega(w) = \frac{1}{2} w^{\top} B w \qquad \text{and} \qquad \Omega^*(\mu) = \frac{1}{2} \mu^{\top} B^{-1} \mu. \tag{20}$$

For the unnormalized negative entropy, where  $\mathcal{W} = \mathbb{R}^d_+$ , we have

$$\Omega(w) = \sum_{i} w^{(i)} \log w^{(i)} - w^{(i)} \quad \text{and} \quad \Omega^*(\mu) = \sum_{i} \exp \mu^{(i)}.$$
(21)

For the normalized negative entropy, where  $W = \{w \mid w \geq 0 \text{ and } ||w||_1 = 1\}$  is the probability simplex, we have

$$\Omega(w) = \sum_{i} w^{(i)} \log w^{(i)} - w^{(i)}$$
 and  $\Omega^*(\mu) = \log \sum_{i} \exp \mu^{(i)}.$  (22)

If  $\Omega$  is differentiable the w at which the supremum of (17) is attained can be written as  $w = \partial_{\mu}\Omega^{*}(\mu)$  (Boyd and Vandenberghe, 2004). In the sequel we will always assume that  $\Omega^{*}$  is twice differentiable. Note that all the regularizers we discussed above are twice differentiable. The following theorem states the dual problem of (16).

**Theorem 2** Denote by  $A = [a_1, ..., a_t]$  the matrix whose columns are the (sub)gradients, and let  $b = [b_1, ..., b_t]$ . The dual problem of

$$w_t = \underset{w \in \mathbb{R}^d}{\operatorname{argmin}} \{ J_t(w) := \underset{1 \le i \le t}{\operatorname{max}} \langle w, a_i \rangle + b_i + \lambda \Omega(w) \} \quad is$$
 (23)

$$\alpha_t = \underset{\alpha \in \mathbb{R}^t}{\operatorname{argmax}} \{ J_t^*(\alpha) := -\lambda \Omega^*(-\lambda^{-1} A \alpha) + \alpha^\top b \mid \alpha \ge 0, \ \|\alpha\|_1 = 1 \}.$$
 (24)

Furthermore,  $w_t$  and  $\alpha_t$  are related by the dual connection  $w_t = \partial \Omega^*(-\lambda^{-1}A\alpha_t)$ .

**Proof** We rewrite (23) as a constrained optimization problem:  $\min_{w,\xi} \lambda \Omega(w) + \xi$  subject to  $\xi \geq \langle w, a_i \rangle + b_i$  for i = 1, ..., t. The corresponding Lagrange function can be written as

$$L(w,\xi,\alpha) = \lambda\Omega(w) + \xi - \alpha^{\top} \left(\xi \mathbf{1}_t - A^{\top}w - b\right) \text{ with } \alpha \ge 0,$$
 (25)

where  $\alpha \geq 0$  denotes that each component of  $\alpha$  is non-negative. Taking derivatives with respect to  $\xi$  yields  $1 - \alpha^{\top} \mathbf{1}_t = 0$ . Moreover, minimization of L with respect to w implies solving  $\max_w \left\langle w, -\lambda^{-1} A \alpha \right\rangle - \Omega(w) = \Omega^*(-\lambda^{-1} A \alpha)$ . Plugging both terms back into (25) allows us to eliminate the primal variables  $\xi$  and w.

Since  $\Omega^*$  is assumed to be twice differentiable and the constraints of (24) are simple, one can easily solve (24) with standard smooth optimization methods such as the *penalty/barrier* methods (Nocedal and Wright, 1999). Recall that for the square norm regularizer  $\Omega(w) = \frac{1}{2} \|w\|_2^2$ , commonly used in SVMs and Gaussian Processes, the Fenchel dual is given by  $\Omega^*(\mu) = \frac{1}{2} \|\mu\|_2^2$ . The following corollary is immediate:

Corollary 3 For quadratic regularization, i.e.,  $\Omega(w) = \frac{1}{2} \|w\|_2^2$ , (24) becomes

$$\alpha_t = \underset{\alpha \in \mathbb{R}^t}{\operatorname{argmax}} \{ -\frac{1}{2\lambda} \alpha^\top A^\top A \alpha + \alpha^\top b \mid \alpha \ge 0, \ \|\alpha\|_1 = 1 \}.$$
 (26)

This means that for quadratic regularization the dual optimization problem is a quadratic program (QP) where the number of constraints equals the number of (sub)gradients computed previously. Since t is typically in the order of 10s to 100s, the resulting QP is very cheap to solve. In fact, we do not even need to know the (sub)gradients explicitly. All that is required to define the QP are the inner products between (sub)gradients  $\langle a_i, a_i \rangle$ .

### 2.4 Convergence Analysis

While the variants of bundle methods we proposed are intuitively plausible, it remains to be shown that they have good rates of convergence. In fact, past results, such as those by Tsochantaridis et al. (2005) suggest a slow  $O(1/\epsilon^2)$  rate of convergence. In this section we tighten their results and show an  $O(1/\epsilon)$  rate of convergence for nonsmooth loss functions and  $O(\log(1/\epsilon))$  rates for smooth loss functions under mild assumptions. More concretely we prove the following two convergence results:

(a) Assume that  $\|\partial_w R_{\text{emp}}(w)\| \leq G$ . For regularizers  $\Omega(w)$  for which  $\|\partial_\mu^2 \Omega^*(\mu)\| \leq H^*$  we prove  $O(1/\epsilon)$  rate of convergence, that is, we show that our algorithm converges to within  $\epsilon$  of the optimal solution in  $O(1/\epsilon)$  iterations.

(b) Under the above conditions, if furthermore  $\|\partial_w^2 J(w)\| \leq H$ , *i.e.*, the Hessian of J is bounded, we can show  $O(\log(1/\epsilon))$  rate of convergence.

For our convergence proofs we use a duality argument similar to those put forward in (Shalev-Shwartz and Singer, 2006; Tsochantaridis et al., 2005), both of which share key techniques with (Zhang, 2003). The crux of our proof argument lies in showing that for nonsmooth loss functions the recurrence  $\epsilon_t - \epsilon_{t+1} \geq c \cdot \epsilon_t^2$  holds for some appropriately chosen constant c. The rates follow by invoking a lemma from Abe et al. (2001). In the case of the smooth losses we show that  $\epsilon_t - \epsilon_{t+1} \geq c' \cdot \epsilon_t$  thus implying an  $O(\log(1/\epsilon))$  rate of convergence.

In order to show the required recurrence, we first observe that by strong duality the values of the primal and dual problems (23) and (24) are equal at optimality. Hence, any progress in  $J_{t+1}$  can be computed in the dual. Next, we observe that the solution of the dual problem (24) at iteration t, denoted by  $\alpha_t$ , forms a feasible set of parameters for the dual problem (24) at iteration t+1 by means of the parameterization  $(\alpha_t, 0)$ , *i.e.*, by padding  $\alpha_t$  with a 0. The value of the objective function in this case equals  $J_t(w_t)$ .

To obtain a lower bound on the improvement due to  $J_{t+1}(w_{t+1})$  we perform a 1-d optimization along  $((1-\eta)\alpha_t, \eta)$  in (24). The constraint  $\eta \in (0,1)$  ensures dual feasibility. We will then bound this improvement in terms of  $\epsilon_t$ . Note that, in general, solving the dual problem (24) results in a increase which is larger than that obtained via the line search. The 1-d minimization is used only for analytic tractability. We now state our key theorem and prove it in Appendix B.

**Theorem 4** Assume that  $\|\partial_w R_{\text{emp}}(w)\| \leq G$  for all  $w \in \text{dom } J$ . Also assume that  $\Omega^*$  has bounded curvature, i.e.,  $\|\partial_{\mu}^2 \Omega^*(\mu)\| \leq H^*$  for all  $\mu \in \{-\lambda^{-1} \sum_{i=1}^{t+1} \alpha_i a_i \text{ where } \alpha_i \geq 0, \ \forall i \ and \ \sum_{i=1}^{t+1} \alpha_i = 1\}$ . In this case we have

$$\epsilon_t - \epsilon_{t+1} \ge \frac{\epsilon_t}{2} \min(1, \lambda \epsilon_t / 4G^2 H^*).$$
 (27)

Furthermore, if  $\|\partial_w^2 J(w)\| \le H$ , then we have

$$\epsilon_{t} - \epsilon_{t+1} \ge \begin{cases} \epsilon_{t}/2 & \text{if } \epsilon_{t} > 4G^{2}H^{*}/\lambda \\ \lambda/8H^{*} & \text{if } 4G^{2}H^{*}/\lambda \ge \epsilon_{t} \ge H/2 \\ \lambda \epsilon_{t}/4HH^{*} & \text{otherwise} \end{cases}$$
(28)

Note that the error keeps on halving initially and settles for a somewhat slower rate of convergence after that, whenever the Hessian of the overall risk is bounded from above. The reason for the difference in the convergence bound for differentiable and non-differentiable losses is that in the former case the gradient of the risk converges to 0 as we approach optimality, whereas in the former case, no such guarantees hold (e.g. when minimizing |x| the (sub)gradient does not vanish at the optimum). The dual of many regularizers, e.g. squares norm, squared  $L_p$  norm, and the entropic regularizer have bounded second derivative. See e.g. (Shalev-Shwartz and Singer, 2006) for a discussion and details. Thus our condition  $\|\partial_{\mu}^2 \Omega^*(\mu)\| \leq H^*$  is not unreasonable. We are now in a position to state our convergence results. The proof is in Appendix B.

**Theorem 5** Assume that  $J(w) \geq 0$  for all w. Under the assumptions of Theorem 4 we can give the following convergence guarantee for Algorithm 2. For any  $\epsilon < 4G^2H^*/\lambda$  the algorithm converges to the desired precision after

$$n \le \log_2 \frac{\lambda J(0)}{G^2 H^*} + \frac{8G^2 H^*}{\lambda \epsilon} - 1$$
 (29)

steps. Furthermore if the Hessian of J(w) is bounded, convergence to any  $\epsilon \leq H/2$  takes at most the following number of steps:

$$n \le \log_2 \frac{\lambda J(0)}{4G^2 H^*} + \frac{4H^*}{\lambda} \max\left[0, H - 8G^2 H^*/\lambda\right] + \frac{4HH^*}{\lambda} \log(H/2\epsilon)$$
 (30)

Several observations are in order: First, note that the number of iterations only depends logarithmically on how far the initial value J(0) is away from the optimal solution. Compare this to the result of Tsochantaridis et al. (2005), where the number of iterations is linear in J(0).

Second, we have an  $O(1/\epsilon)$  dependence in the number of iterations in the non-differentiable case, as opposed to the  $O(1/\epsilon^2)$  rates of Tsochantaridis et al. (2005). In addition to that, the convergence is  $O(\log(1/\epsilon))$  for continuously differentiable problems.

Note that whenever  $R_{\text{emp}}$  is the average over many piecewise linear functions,  $R_{\text{emp}}$  behaves essentially like a function with bounded Hessian as long as we are taking large enough steps not to "notice" the fact that the term is actually nonsmooth.

**Remark 6** For  $\Omega(w) = \frac{1}{2} \|w\|^2$  the dual Hessian is exactly  $H^* = 1$ . Moreover we know that  $H \ge \lambda$  since  $\|\partial_w^2 J(w)\| = \lambda + \|\partial_w^2 R_{\text{emp}}(w)\|$ .

Effectively the rate of convergence of the algorithm is governed by upper bounds on the primal and dual curvature of the objective function. This acts like a condition number of the problem — for  $\Omega(w) = \frac{1}{2} w^{\top} Q w$  the dual is  $\Omega^*(z) = \frac{1}{2} z^{\top} Q^{-1} z$ , hence the largest eigenvalues of Q and  $Q^{-1}$  would have a significant influence on the convergence.

In terms of  $\lambda$  the number of iterations needed for convergence is  $O(\lambda^{-1})$ . In practice the iteration count *does* increase with  $\lambda$ , albeit not as badly as predicted. This is likely due to the fact that the empirical risk  $R_{\rm emp}$  is typically rather smooth and has a certain inherent curvature which acts as a natural regularizer in addition to the regularization afforded by  $\lambda\Omega(w)$ .

For completeness we also state the convergence guarantees for Algorithm 3 and provide a proof in Appendix B.1.

**Theorem 7** Under the assumptions of Theorem 4 Algorithm 3 converges to the desired precision  $\epsilon$  after

$$n \le \frac{8G^2H^*}{\lambda\epsilon}$$

steps for any  $\epsilon < 4G^2H^*/\lambda$ .

## 3. Implementation Issues

### 3.1 Solving the BMRM Subproblem (16) with Limited Memory Space

In Section 2.3 we mentioned the dual of subproblem (16) (i.e., (24)) which is usually easier to solve when the dimensionality d of the problem is larger than the number of iterations t required by BMRM to reach desired precision  $\epsilon$ . Although t is usually in the order of  $10^2$ , a problem with d in the order of  $10^6$  or higher may use up all memory of a typical machine to store the bundle, i.e., linearizations  $\{(a_i, b_i)\}$ , before the convergence is achieved. Here we describe a principled technique which controls the memory usage while maintaining convergence guarantees.

Note that at iteration t, before the computation for new iterate  $w_t$ , Algorithm 2 maintains a bundle of t (sub)gradients  $\{a_i\}_{i=1}^t$  of  $R_{\text{emp}}$  computed at the locations  $\{w_i\}_{i=0}^{t-1}$ . Furthermore, the Lagrange multipliers  $\alpha_{t-1}$  obtained in iteration t-1 satisfy  $\alpha_{t-1} \geq 0$  and  $\sum_{i=1}^{t-1} \alpha_{t-1}^{(i)} = 1$  by the constraints of (24). We define the aggregated (sub)gradient  $\hat{a}_I$ , offset  $\hat{b}_I$  and Lagrange multiplier  $\hat{\alpha}_{t-1}^{(I)}$  as

$$\hat{a}_I := \frac{1}{\hat{\alpha}_{t-1}^{(I)}} \sum_{i \in I} \alpha_{t-1}^{(i)} a_i, \quad \hat{b}_I := \frac{1}{\hat{\alpha}_{t-1}^{(I)}} \sum_{i \in I} \alpha_{t-1}^{(i)} b_i, \quad \text{and} \quad \hat{\alpha}_{t-1}^{(I)} := \sum_{i \in I} \alpha_{t-1}^{(i)}, \tag{31}$$

respectively, where  $I \subseteq [t-1]$  is an index set (Kiwiel, 1983). Clearly, the optimality of (24) at the end of iteration t-1 is maintained when a subset  $\left\{(a_i, b_i, \alpha_{t-1}^{(i)})\right\}_{i \in I}$  is replaced by the aggregate  $(\hat{a}_I, \hat{b}_I, \hat{\alpha}_{t-1}^{(I)})$  for any  $I \subseteq [t-1]$ .

To obtain a new iterate  $w_t$  via (24) with memory space for at most k linearizations, we can, for example, replace  $\{(a_i,b_i)\}_{i\in I}$  with  $(\hat{a}_I,\hat{b}_I)$  where I=[t-k+1] and  $1\leq k\leq t$ . Then, we solve a k-dimensional variant of (24) with  $1\leq [a_1,a_{t-k+2},\ldots,a_t]$ ,  $1\leq [a_1,a_{t-k+2},\ldots,a_t]$ , and  $1\leq k\leq t$ . The optimum of this variant will be lower than or equal to that of (24) as the latter has higher degree of freedom than the former. Nevertheless, solving this variant with  $1\leq k\leq t$  will still guarantee convergence (recall that our convergence proof only uses  $1\leq k\leq t$  will still guarantee the aforementioned number  $1\leq t\leq t$  where  $1\leq t\leq t$  is the "bundle size" since it indicates the number of linearizations the algorithm keeps.

For concreteness, we provide here a memory efficient BMRM variant for the cases where  $\Omega(w) = \frac{1}{2} \|w\|_2^2$  and k = 2. We first see that the dual of subproblem (16) now reads:

$$\eta = \underset{0 \le \eta \le 1}{\operatorname{argmax}} - \frac{1}{2\lambda} \left\| \hat{a}_{[t-1]} + \eta (a_t - \hat{a}_{[t-1]}) \right\|_2^2 + \hat{b}_{[t-1]} + \eta (b_t - \hat{b}_{[t-1]})$$
(32)

$$\equiv \underset{0 \le \eta \le 1}{\operatorname{argmax}} - \frac{\eta}{\lambda} \hat{a}_{[t-1]}^{\top} (a_t - \hat{a}_{[t-1]}) - \frac{\eta^2}{2\lambda} \left\| a_t - \hat{a}_{[t-1]}^{\top} \right\|^2 + \eta (b_t - \hat{b}_{[t-1]}), \tag{33}$$

<sup>3.</sup> In practice, we can remove those linearizations  $\{(a_i,b_i)\}$  whose Lagrange multipliers  $\alpha_i$  are 0 after solving (24). Although this heuristic works well and does not affect the convergence guarantee, there is no bound on the minimum number of linearizations with non-zero Lagrange multipliers needed to achieve convergence.

Since (33) is quadratic in  $\eta$ , we can obtain the optimal  $\eta$  by setting the derivative of the objective in (33) to zero and clipping  $\eta$  in the range [0, 1]:

$$\eta = \min\left(\max\left(0, \frac{b_t - \hat{b}_{[t-1]} + w_{t-1}^{\top} a_t + \lambda \|w_{t-1}\|^2}{\frac{1}{\lambda} \|a_t + \lambda w_{t-1}\|^2}\right), 1\right)$$
(34)

where  $w_{t-1} = -\frac{1}{\lambda}\hat{a}_{[t-1]}$  by the dual connection. With the optimal  $\eta$ , we obtain the new primal iterate  $w_t = (1 - \eta)w_{t-1} - (\eta/\lambda)a_t$ . Algorithm 4 lists the details. Note that this variant is simple to implement and does not require a QP solver.

# **Algorithm 4** BMRM with Aggregation of Previous Linearizations

```
1: input & initialization: \epsilon \geq 0, w_0, t \leftarrow 1

2: Compute a_1 \in \partial_w R_{\text{emp}}(w_0), and b_1 \leftarrow R_{\text{emp}}(w_0) - \langle w_0, a_1 \rangle

3: w_1 \leftarrow -\frac{1}{\lambda}a_1

4: \hat{b}_{[1]} \leftarrow b_1

5: repeat

6: t \leftarrow t + 1

7: Compute a_t \in \partial_w R_{\text{emp}}(w_{t-1}) and b_t \leftarrow R_{\text{emp}}(w_{t-1}) - \langle w_{t-1}, a_t \rangle

8: Compute \eta using Eq. (34)

9: w_t \leftarrow (1 - \eta)w_{t-1} - (\eta/\lambda)a_{t+1}

10: \hat{b}_{[t]} \leftarrow (1 - \eta)\hat{b}_{[t-1]} + \eta b_t

11: \epsilon_t \leftarrow \min_{0 \leq i \leq t} \frac{\lambda}{2} ||w_i||^2 + R_{\text{emp}}(w_i) + \frac{\lambda}{2} ||w_t||^2 - \hat{b}_{[t]}

12: until \epsilon_t \leq \epsilon
```

## 3.2 Parallelization

Algorithms 2, 3, and 4 the evaluation of  $R_{\text{emp}}(w)$  (and  $\partial_w R_{\text{emp}}(w)$ ) is cleanly separated from the computation of new iterate and the choice of regularizer. If  $R_{\text{emp}}$  is additively decomposable over the examples  $(x_i, y_i)$ , i.e., can be expressed as a sum of some independent loss terms  $l(x_i, y_i, w)$ , then we can parallelize these algorithms easily by splitting the datasets and the computation  $R_{\text{emp}}$  over multiple machines. This parallelization scheme not only reduces the computation time but also allows us to handle dataset with size exceeding the memory available on a single machine.

Without loss of generality, we describe a parallelized version of Algorithm 2 here. Assume there are p slave machines and 1 master machine available. At the beginning, we partition a given dataset  $D = \{(x_i, y_i)\}_{i=1}^m$  into p disjoint sub-datasets  $\{D_i\}_{i=1}^p$  and assign one sub-dataset to each slave machine. At iteration t, the master first broadcasts the current iterate  $w_{t-1}$  to all p slaves (e.g. using MPI function MPI::Broadcast (Gropp et al., 1999)). The slaves then compute the losses and (sub)gradients on their local sub-datasets in parallel. As soon as the losses and (sub)gradients computation finished, the master combines the results (e.g. using MPI::AllReduce). With the combined (sub)gradient and offset, the master computes the new iterate  $w_t$  as in Algorithms 2 and 3. This process repeats until convergence is achieved. Detailed pseudocode can be found in Algorithm 5.

### Algorithm 5 Parallel BMRM

```
1: input: \epsilon \geq 0, w_0, dataset D, number of slave machines p
 2: initialization: t \leftarrow 0, assign sub-dataset D_i to slave i, i = 1, \ldots, p
 3: repeat
 4:
          t \leftarrow t + 1
          Master: Broadcast w_{t-1} to all slaves
 5:
         Slaves: Computes R_{\text{emp}}^i(w_{t-1}) := \sum_{(x,y) \in D_i} l(x,y,w_{t-1}) and a_t^i \in \partial_w R_{\text{emp}}^i(w_{t-1})
Master: Aggregate a_t := \frac{1}{|D|} \sum_{i=1}^p a_t^i and b_t := \frac{1}{|D|} \sum_{i=1}^p R_{\text{emp}}^i(w_{t-1}) - \langle w_{t-1}, a_t \rangle
 6:
 7:
          Master: Update model R_t^{CP}(w) := \max_{1 \le j \le t} \langle w, a_j \rangle + b_j
 8:
 9:
          Master: w_t \leftarrow \operatorname{argmin}_w J_t(w) := \lambda \Omega(w) + R_t^{CP}(w)
          Master: \epsilon_t \leftarrow \min_{0 \le i \le t} J(w_i) - J_t(w_t)
11: until \epsilon_t \leq \epsilon
12: return w_t
```

# 4. Related Research

The kernel trick is widely used to transform many existing machine learning algorithms into ones operating on a Reproducing Kernel Hilbert Space (RKHS). One lifts w into an RKHS and replaces all inner product computations with a positive definite kernel function  $k(x, x') \leftarrow \langle x, x' \rangle$ . Examples of algorithms which employ the kernel trick (but essentially still solve (1a)) include Support Vector regression (Vapnik et al., 1997), novelty detection (Schölkopf et al., 2001), Huber's robust regression, quantile regression (Takeuchi et al., 2006), ordinal regression (Herbrich et al., 2000), ranking (Crammer and Singer, 2005), maximization of multivariate performance measures (Joachims, 2005), structured estimation (Taskar et al., 2004; Tsochantaridis et al., 2005), Gaussian Process regression (Williams, 1998), conditional random fields (Lafferty et al., 2001), graphical models (Cowell et al., 1999), exponential families (Barndorff-Nielsen, 1978), and generalized linear models (Fahrmeir and Tutz, 1994).

Traditionally, specialized solvers have been developed for solving the kernel version of (1a) in the dual, e.g. (Chang and Lin, 2001; Joachims, 1999). These algorithms construct the Lagrange dual, and solve for the Lagrange multipliers efficiently. Only recently, research focus has shifted back to solving (1a) in the primal, e.g. (Chapelle, 2007; Joachims, 2006; Sindhwani and Keerthi, 2006). This spurt in research interest is due to three main reasons: First, many interesting problems in diverse areas such as text classification, word-sense disambiguation, and drug design already employ rich high dimensional data which does not necessarily benefit from the kernel trick. All these domains are characterized by large datasets (with m in the order of a million) and very sparse features (e.g. the bag of words representation of a document). Second, efficient factorization methods (e.g. (Fine and Scheinberg, 2001)) can be used for a low rank representation of the kernel matrix thereby effectively rendering the problem linear. Third, approximation methods such as the Random Feature Map proposed by Rahimi and Recht (2008) can efficiently approximate a infinite dimensional nonlinear feature map associated to a kernel by a finite dimensional one. Therefore our focus on the primal optimization problem is not only pertinent but also timely.

The widely used SVM<sup>struct</sup> optimizer of Thorsten Joachims (http://svmlight.joachims.org/svm\_struct.html) is closely related to BMRM. While BMRM can handle many different regularizers and loss functions, SVM<sup>struct</sup> is mainly geared towards square norm regularizers and non-differentiable soft-margin type loss functions. On the other hand, SVM<sup>struct</sup> can handle kernels while BMRM mainly focuses on the primal problem.

Our convergence analysis is closely related to Shalev-Shwartz and Singer (2006) who prove mistake bounds for online algorithms by lower bounding the progress in the dual. Although not stated explicitly, essentially the same technique of lower bounding the dual improvement was used by Tsochantaridis et al. (2005) to show polynomial time convergence of the SVM<sup>struct</sup> algorithm. The main difference however is that Tsochantaridis et al. (2005) only work with a quadratic objective function while the framework proposed by Shalev-Shwartz and Singer (2006) can handle arbitrary convex functions. In both cases, a weaker analysis led to  $O(1/\epsilon^2)$  rates of convergence for nonsmooth loss functions. On the other hand, our results establish a  $O(1/\epsilon)$  rate for nonsmooth loss functions and  $O(\log(1/\epsilon))$  rates for smooth loss functions under mild technical assumptions.

Another related work is SVM<sup>perf</sup> (Joachims, 2006) which solves the SVM with linear kernel in linear time. SVM<sup>perf</sup> finds a solution with accuracy  $\epsilon$  in  $O(md/(\lambda\epsilon^2))$  time, where the m training patterns  $x_i \in \mathbb{R}^d$ . This bound was improved by Shalev-Shwartz et al. (2007) to  $\tilde{O}(1/\lambda\delta\epsilon)$  for obtaining an accuracy of  $\epsilon$  with confidence  $1-\delta$ . Their algorithm, Pegasos, essentially performs stochastic (sub)gradient descent but projects the solution back onto the  $L_2$  ball of radius  $1/\sqrt{\lambda}$ . Note that Pegasos also can be used in an online setting. This, however, only applies whenever the empirical risk decomposes into individual loss terms (e.g. it is not applicable to multivariate performance scores (Joachims, 2005)).

The third related strand of research considers gradient descent in the primal with a line search to choose the optimal step size, see e.g. (Boyd and Vandenberghe, 2004, Section 9.3.1). Under assumptions of smoothness and strong convexity—that is, the objective function can be upper and lower bounded by quadratic functions—it can be shown that gradient descent with line search will converge to an accuracy of  $\epsilon$  in  $O(\log(1/\epsilon))$  steps. Our solver achieves the same rate guarantees for smooth functions, under essentially similar technical assumptions.

We would also like to point out connections to subgradient methods (Nedich and Bertsekas, 2000). These algorithms are designed for nonsmooth functions, and essentially choose an arbitrary element of the subgradient set to perform a gradient descent like update. Let  $||J_w(w)|| \leq G$ , and  $B(w^*, r)$  denote a ball of radius r centered around the minimizer of J(w). By applying the analysis of Nedich and Bertsekas (2000) to the regularized risk minimization problem with  $\Omega(w) := \frac{\lambda}{2} ||w||^2$ , Ratliff et al. (2007) showed that subgradient descent with a fixed, but sufficiently small, stepsize will converge linearly to  $B(w^*, G/\lambda)$ .

Finally, several papers (Keerthi and DeCoste, 2005; Chapelle, 2007) advocate the use of Newton-like methods to solve Support Vector Machines in the "primal". However, they need to take precautions when dealing with the fact that the soft-margin type of loss functions such as the hinge loss is only piecewise differentiable. Instead, our method only requires subdifferentials, which always exist for convex functions, in order to make progress. The large number of and variety of implemented problems shows the flexibility of our approach.

### 5. Experiments

In this section, we examine the convergence behavior of BMRM and show that it is versatile enough to solve a variety of machine learning problems. All our experiments were carried out on a cluster of 24 machines each with a 2.4GHz AMD Dual Core processor and 4GB of RAM. Details of the loss functions, datasets, competing solvers and experimental objectives are described in the following subsections.

### 5.1 Convergence Behavior

We investigated the convergence rate of our method (Algorithm 2) empirically with respect to regularization constant  $\lambda$ , approximation gap  $\epsilon$ , and bundle size k. In addition, we investigated the speedup gained by parallelizing the empirical risk computation. Finally, we examined empirically how generalization performance is related to approximation gap. For simplicity, we focused on the training of a linear SVM with binary hinge loss: <sup>4</sup>

$$\min_{w} J(w) := \frac{\lambda}{2} ||w||^2 + \frac{1}{m} \sum_{i=1}^{m} \max(0, 1 - y_i \langle w, x_i \rangle).$$
 (35)

The experiments were conducted on 6 datasets commonly used in binary classification studies, namely, adult9, astro-ph, news20-b<sup>5</sup>, rcv1, real-sim, and worm. adult9, news20-b, rcv1, and real-sim are available at <a href="http://www.csie.ntu.edu.tw/~cjlin/libsvmtools/datasets/binary.html">http://www.csie.ntu.edu.tw/~cjlin/libsvmtools/datasets/binary.html</a>. astro-ph (Joachims, 2006) and worm (Franc and Sonnenburg, 2008) are available upon request from Thorsten Joachims and Soeren Sonnenburg, respectively. Table 1 summarizes the properties of the datasets.

Dataset	#examples $m$	dimension $d$	density %
adult9	48,842	123	11.27
astro-ph	94,856	99,757	0.08
news20-b	19,954	1,355,191	0.03
rcv1	677,399	47,236	0.15
real-sim	72,201	20,958	0.25
worm	1,026,036	804	25.00

Table 1: Properties of the binary classification datasets used in our experiments.

### 5.1.1 Regularization constant $\lambda$ and approximation gap $\epsilon$

As suggested by the convergence analysis, the linear SVM with the nonsmooth binary hinge loss should converge in  $O(\frac{1}{\lambda\epsilon})$  iterations, where  $\lambda$  and  $\epsilon$  are two parameters which one normally tunes during the model selection phase. Therefore, we investigated the scaling behavior of our method w.r.t. these two parameters. We performed the experiments

<sup>4.</sup> Similar behavior was observed with other loss functions.

<sup>5.</sup> The dataset is originally named news20; we renamed it to avoid confusion with the multiclass version of the dataset.

with unlimited bundle size and with a heuristic that removes subgradients which remained inactive (i.e., Lagrange multiplier = 0) for 10 or more consecutive iterations.<sup>6</sup>

Figure 4 shows the approximation gap  $\epsilon_t$  as a function of number of iterations t. As predicted by our convergence analysis, BMRM converges faster for larger values of  $\lambda$ . Furthermore, the empirical convergence curves exhibit a  $O(\log(1/\epsilon))$  rate instead of the (pessimistic) theoretical rate of  $O(\frac{1}{\epsilon})$ , especially for large values of  $\lambda$ . Interestingly, BMRM converges faster on high-dimensional text datasets (*i.e.*, astro-ph, news20-b, rcv1, and real-sim) than on lower dimensional datasets (*i.e.*, adult9 and worm).

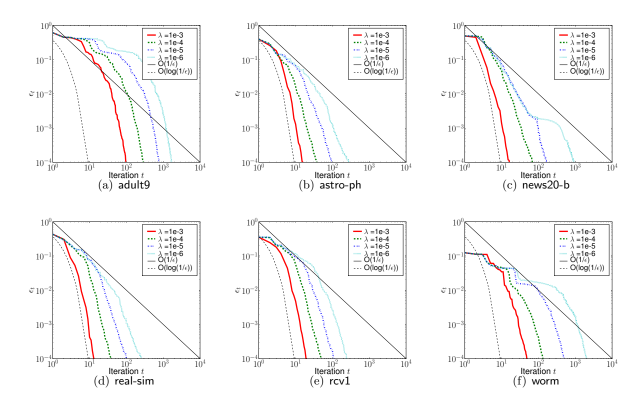


Figure 4: Approximation gap  $\epsilon_t$  as a function of number of iterations t; for different regularization constants  $\lambda$  (and unlimited bundle size).

### 5.1.2 Bundle size

The dual of our method (24) is a concave problem which has dimensionality equal to the number of iterations executed. In the case of linear SVM, (24) is a QP. Hence, as described in Section 3.1, we can trade potentially greater bundle improvement for memory efficiency.

Figure 5 shows the approximation gap  $\epsilon_t$  during the training of linear SVM as a function of the number of iterations t, for different bundle sizes  $k \in \{2, 10, 50, \infty\}$ . In the case of  $k = \infty$ , we employed the same heuristics which remove inactive linearizations as those

<sup>6.</sup> Note that this heuristic does not have any implication in the convergence analysis.

mentioned in section 5.1.1. As expected, the larger k is, the faster the algorithm converges. Although the case k=2 is the slowest, its convergence rate is still faster than the theoretical bound  $\frac{1}{\lambda \epsilon}$ .

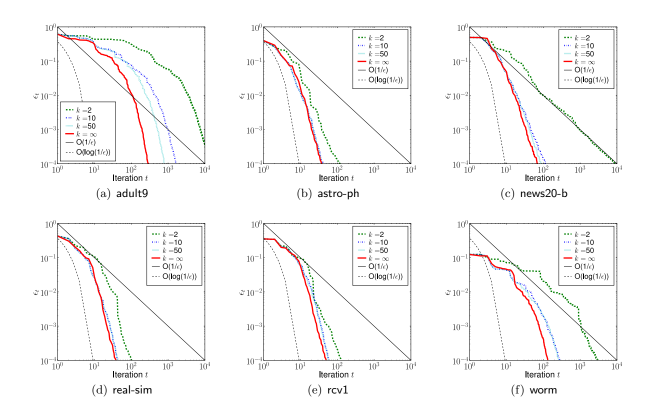


Figure 5: Approximation gap  $\epsilon_t$  as a function of number of iterations t; for different bundle sizes k (and fixed regularization constant  $\lambda = 10^{-4}$ ).

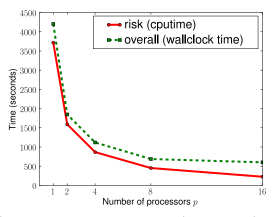
### 5.1.3 Parallelization

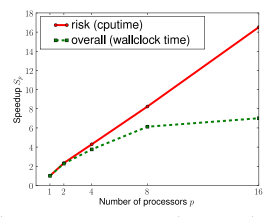
When the empirical risk  $R_{\rm emp}$  is additively decomposable, the loss and subgradient computation can be executed concurrently on multiple processors for different subsets of data points.<sup>7</sup>

We performed experiments for linear SVMs training with parallelized risk computation on the worm dataset. Figure 6(a) shows the wallclock time for the overall training phase (e.g. data loading, risk computation, and solving the QP) and CPU time for just the risk computation as a function of number of processors p. Note that the gap between the two curves essentially tells the runtime upper bound of the sequential part of the algorithm. As expected, both overall and risk computation time decrease as the number of processors p

<sup>7.</sup> This requires only slight modification to the data loading process and the addition of some parallelization related code before and after the code segment for empirical risk computation.

increases. However, in Figure 6(b), we see two different speedups.<sup>8</sup> The speedup for the risk computation is roughly linear as there is no sequential part in it; the speedup of overall computation is approaching a limit<sup>9</sup> as well-explained by Amdahl's law (Amdahl, 1967).





- (a) Risk computation in CPU time (red solid line) and overall computation (*i.e.*, data loading + risk computation + solving the QP) in wallclock time (green dashed line) as a function of number of processors.
- (b) Speedup in risk computation (in CPU time) and overall computation (in wallclock time) as a function of number of processors.

Figure 6: CPU and wallclock time for training linear SVM using parallel BMRM on worm dataset with varying number of processors  $p \in \{1, 2, 4, 8, 16\}$ . In these experiments, regularization constant  $\lambda = 10^{-6}$ , and termination criterion  $\epsilon = 10^{-4}$ .

#### 5.1.4 Generalization versus approximation gap

Since the problems we are considering are convex, all properly convergent optimizers will converge to the same solution. Therefore, comparing generalization performance of the final solution in meaningless. But, in real life one is often interested in the speed with which the algorithm achieves good generalization performance. In this section we study this question. We focus on the generalization (in terms of accuracy) as a function of approximation gap during training. For this experiment, we randomly split each of the datasets into training (60%), validation (20%) and testing (20%) sets.

We first obtained the best  $\lambda \in \{2^{-20}, \dots, 2^0\}$  for each of the datasets using their corresponding validation sets. With these best  $\lambda$ 's, we (re)trained linear SVMs and recorded the testing accuracy as well as the approximation gap at every iteration, with termination criterion  $\epsilon = 10^{-4}$ . Figure 7 shows the difference between the testing accuracy evaluated at every iteration and that after training, as a function of approximation gap at each iteration.

From the figure, it is clear that the testing accuracies on low dimensional datasets (*i.e.*, adult9 and worm) are less stable in general and the approximation gap must be reduced

<sup>8.</sup> Speedup  $S_p = \frac{T_1}{T_p}$  where p is the number of processors and  $T_q$  is the runtime of the parallelized algorithm on q processors.

<sup>9.</sup> The limit of speedup is the inverse of the sequential fraction of the algorithm such as the QP.

to at least  $10^{-3}$  to reach the 0.5% regime of the final testing accuracies; whereas on high dimensional text datasets (*i.e.*, astro-ph, news20-b, rcv1 and real-sim), the testing accuracies arrived at the same regime with approximation gap of  $10^{-2}$  or lower.

In general, the generalization improved as the approximation gap decreased. The improvement in generalization became rather insignificant (say, the maximum of changes in testing accuracies is less than 0.1%) when the approximation gap was further reduced to below some effective threshold  $\epsilon_{\rm eff}$ ; that said, it is not necessary to continue the optimization when  $\epsilon_t \leq \epsilon_{\rm eff}$ . Since  $\epsilon_{\rm eff}$  (or its scale) is not known a priori and the asymptotic analysis in (Shalev-Schwartz and Srebro, 2008) does not reveal the actual scale of  $\epsilon_{\rm eff}$  directly applicable in our case, we carried out another set of experiments to investigate if  $\epsilon_{\rm eff}$  could be estimated with as little effort as possible: For each dataset, we randomly subsampled  $10\%, \ldots, 50\%$  of the training set as sub-datasets and performed the same experiment on all sub-datasets. We then determined the largest  $\epsilon_{\rm eff}$  such that the maximum changes in testing accuracies is less than 0.1%.

Table 5.1.4 shows the (base 10 logarithm of)  $\epsilon_{\rm eff}$  for all sub-datasets as well as the full datasets. It seems that the  $\epsilon_{\rm eff}$  estimated on a smaller sub-dataset is at most 1 order of magnitude larger than the actual  $\epsilon_{\rm eff}$  required on full dataset. In addition, we show in the table that the necessary threshold  $\epsilon_{10\%}$  required by the sub-datasets and the full datasets to attain the final testing accuracies attained by the 10% sub-datasets. The observations obey the analysis in (Shalev-Schwartz and Srebro, 2008) that for a fixed testing accuracy, approximation gap (i.e., optimization error) can be relaxed when more data is given.

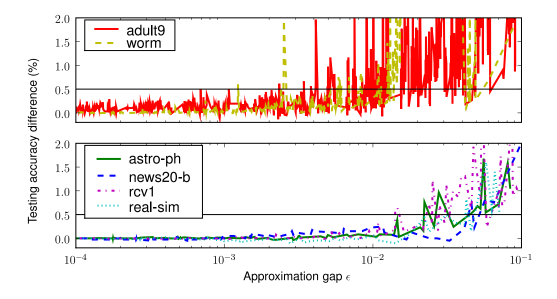


Figure 7: Difference between testing accuracies of intermediate and final models.

<sup>10.</sup> Heuristically, we could terminate the training phase following the *early stopping* strategy by monitoring the changes in accuracies on validation set evaluated in some most recent iterations.

		10%	20%	30%	40%	50%	100%
adult9	Acc. (%)	84.3	84.7	84.9	85.1	85.1	85.2
	$\log_{10} \epsilon_{ ext{eff}}$	-3.90	-3.72	-3.77	-3.88	-3.64	-4.00
	$\log_{10} \epsilon_{10\%}$	-4.01	-1.18	-1.07	-1.16	-1.27	-1.04
astro-ph	Acc. (%)	96.1	96.6	96.4	96.6	96.8	97.4
	$\log_{10} \epsilon_{ ext{eff}}$	-1.48	-1.70	-1.57	-1.49	-1.68	-1.84
	$\log_{10} \epsilon_{10\%}$	-4.00	-1.15	-1.06	-0.98	-1.02	-0.87
news20-b	Acc. (%)	89.9	92.9	94.3	94.5	95.4	96.6
	$\log_{10} \epsilon_{ ext{eff}}$	-2.00	-2.48	-3.87	-1.65	-3.71	-2.84
	$\log_{10} \epsilon_{10\%}$	-4.02	-0.92	-0.70	-0.80	-0.80	-0.67
	Acc. (%)	96.9	97.2	97.4	97.2	97.5	97.6
rcv1	$\log_{10} \epsilon_{ ext{eff}}$	-2.02	-2.40	-1.99	-2.16	-2.34	-2.28
	$\log_{10} \epsilon_{10\%}$	-4.07	-1.19	-1.30	-1.29	-1.13	-1.11
real-sim	Acc. (%)	95.0	95.9	96.3	96.6	96.6	97.2
	$\log_{10} \epsilon_{ ext{eff}}$	-1.74	-1.84	-1.71	-1.99	-1.74	-1.75
	$\log_{10} \epsilon_{10\%}$	-4.02	-1.04	-0.88	-0.87	-0.85	-0.82
worm	Acc. (%)	98.2	98.2	98.2	98.3	98.3	98.4
	$\log_{10} \epsilon_{ ext{eff}}$	-2.43	-2.47	-2.48	-3.62	-2.81	-3.55
	$\log_{10} \epsilon_{10\%}$	-4.00	-1.38	-1.28	-1.37	-1.28	-1.31

Table 2: The first sub-row in each dataset row indicates the testing accuracies of models trained on the corresponding proportions of the dataset. The second sub-row indicates the (base 10 logarithm of) effective threshold such that the maximum difference in testing accuracies of models with approximation gap smaller than that is less than 0.1%. The third sub-row indicates the (base 10 logarithm of) threshold necessary for models to attain the testing accuracy attained by the model trained on the 10% sub-dataset.

### 5.2 Comparison with Existing Bundle Methods

In this section we compare BMRM with a BT implementation obtained from Schramm and Zowe (1992).<sup>11</sup> We also compare the performance of BMRM (Algorithm 2) and LSBMRM (Algorithm 3). The multiclass line search used in LSBMRM can be found in Yu et al. (2008).

For binary classification, we solve the linear SVM (35) on the datasets: adult9, astro-ph, news20-b, rcv1, real-sim, and worm as mentioned in Section 5.1. For multiclass classification, we solve (Crammer and Singer, 2003):

$$\min_{w} J(w) := \frac{\lambda}{2} \|w\|^2 + \frac{1}{m} \sum_{i=1}^{m} \max_{y'_i \in [c]} \left\langle w, e_{y'_i} \otimes x_i - e_{y_i} \otimes x_i \right\rangle + \mathbf{I}(y_i \neq y'_i), \tag{36}$$

where c is the number of classes in the problem,  $e_i$  is the i-th standard basis for  $\mathbb{R}^c$ ,  $\otimes$  denotes Kronecker product; and  $\mathbf{I}(\cdot)$  is an indicator function that has value 1 if its argument is evaluated true, and 0 otherwise. The datasets used in multiclass classification experiments were inex, letter, mnist, news20-m<sup>12</sup>, protein, and usps. inex is available at http://webia.lip6.fr/~bordes/datasets/multiclass/inex.tar.gz and the rest can be found at http://www.csie.ntu.edu.tw/~cjlin/libsvmtools/datasets/multiclass.html. Table 3 summarizes the properties of these datasets.

Dataset	#examples $m$	#classes $c$	dimension $d$	density %
inex	12,107	18	167,295	0.48
letter	20,000	26	16	100.00
mnist	70,000	10	780	19.24
news20-m	19,928	20	62,061	0.13
protein	21,516	3	357	28.31
usps	9,298	10	256	96.70

Table 3: Properties of the multiclass classification datasets used in the experiments.

In each of the experiments, BT was terminated when the norm of the (sub)gradient of J is less than  $10^{-4}$ , whereas BMRM and LSBMRM were terminated when the approximation gap  $\epsilon_t < 10^{-4}$ . In addition to these termination criteria, the maximum number of iterations was set to 2000. Let  $\bar{w}$  denote the "worst" of the three minimizers *i.e.*, the minimizer amongst the three competing methods that has the highest objective function value. To compare these methods, we consider the smallest number of iterations t required to reach 1% of the "worst" minimum  $J(\bar{w})$ :

$$\frac{J(w_t) - J(\bar{w})}{J(\bar{w})} \le 0.01. \tag{37}$$

Figure 8 shows the number of iterations t required by the three methods on each dataset to satisfy (37) as a function of regularization constant  $\lambda \in \{10^{-3}, 10^{-4}, 10^{-5}, 10^{-6}\}$ . As expected, LSBMRM, which uses an exact (but efficient) line search, outperformed both

<sup>11.</sup> The original FORTRAN implementation was automatically converted into C for use in our library

<sup>12.</sup> The dataset is originally named news20; we renamed it to avoid confusion with the binary version of the dataset.

BMRM and BT on all datasets. BMRM performed better than BT on all high dimensional datasets except news20-m but worse on the rest. Although BT tunes the stabilization trade-off parameter  $\kappa_t$  automatically, it still does not guarantee superiority over BMRM which is considerably simpler.

### 5.3 Versatility

In the following subsections, we will illustrate some of the applications of BMRM to various machine learning problems with smooth and non-differentiable loss functions, and with different regularizers. Our aim is to show that BMRM is versatile enough to be used in a variety of seemingly different problems. Readers not interested in this aspect of BMRM can safely skip this subsection.

### 5.3.1 Binary classification

In this section, we evaluate the performance of our method BMRM in the training of binary classifier using linear SVMs (35) and logistic loss:

$$\min_{w} J(w) := \frac{\lambda}{2} ||w||^2 + \frac{1}{m} \sum_{i=1}^{m} \log(1 + \exp(-y_i \langle w, x_i \rangle)), \tag{38}$$

on the binary classification datasets mentioned in Section 5.1 with split similar to that in Section 5.1.4. Since we will compare BMRM with other solvers which use different termination criteria, we consider the CPU time used in reducing the relative difference between the current smallest objective function value and the optimum:

$$\frac{\min_{i \le t} J(w_i) - J(w^*)}{J(w^*)},\tag{39}$$

where  $w_i$  is the weight vector at time/iteration i, and  $w^*$  is the minimizer obtained by running BMRM until the approximation gap  $\epsilon_t < 10^{-4}$ . The best  $\lambda \in \{2^{-20}, \dots, 2^0\}$  for each of the datasets was determined by evaluating the training performance on the corresponding validation set.<sup>13</sup>

In the case of linear SVMs, we compared BMRM to three publicly available state of the art *batch learning* solvers:

- 1. OCAS (Franc and Sonnenburg, 2008) version 0.01 (since this method is equivalent to LSBMRM with binary hinge loss, we refer to this software by LSBMRM for naming consistency),
- 2. LIBLINEAR (Fan et al., 2008) version 1.33 with option "-s 3",
- 3. SVM<sup>perf</sup> (Joachims, 2006) version 2.5 with option "-w 3" and with double precision floating point numbers.

LIBLINEAR solves the dual problem of linear SVM using a coordinate descent method (Hsieh et al., 2008). SVM<sup>perf</sup> was chosen for comparison as it is algorithmically identical to BMRM in this case. Both LIBLINEAR and SVM<sup>perf</sup> provide a "shrinking" technique to speed up

<sup>13.</sup> The corresponding penalty parameter C for LIBLINEAR and OCAS is  $1/(m\lambda)$ , and for SVM<sup>perf</sup> is  $1/(100\lambda)$ .

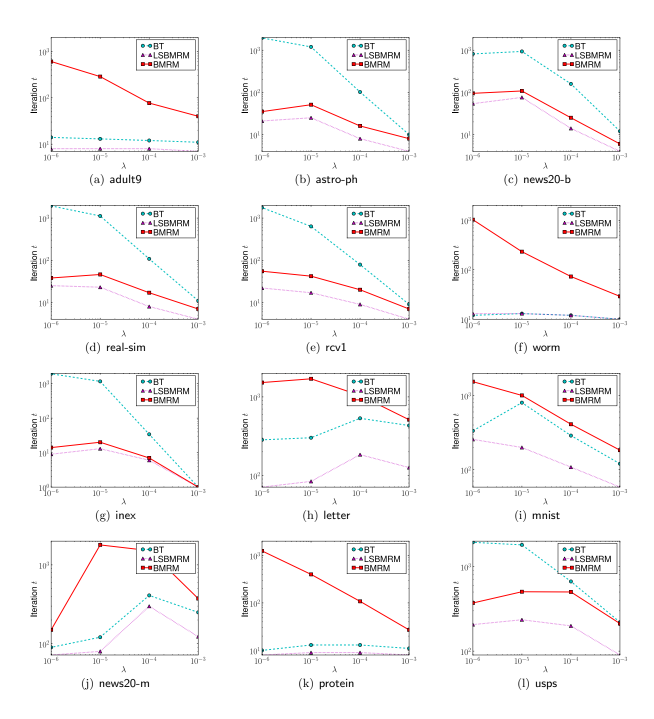


Figure 8: Smallest number of iterations required to satisfy the termination criterion (37) for each dataset and various regularization constants. Note that the points touching the top of the frame (i.e., t = 2000) does not necessarily satisfy (37).

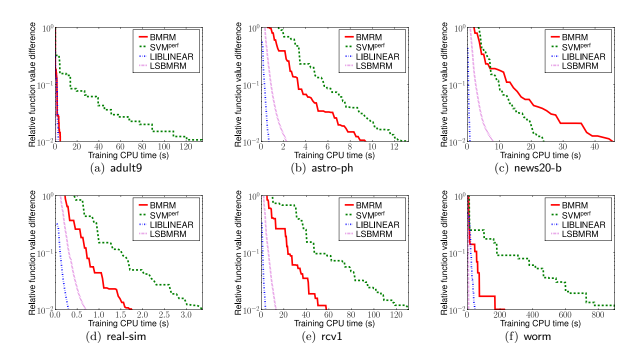


Figure 9: Linear SVMs. Relative primal objective value difference during training.

the algorithms by ignoring some data points which are not likely to affect the objective. Since BMRM does not provide such shrinking technique, we excluded this option in both LIBLINEAR and SVM<sup>perf</sup> for a fair comparison.

Figure 9 shows the relative difference in objective value as a function of training time (CPU seconds) for three methods on various datasets. BMRM is faster than SVM<sup>perf</sup> on all datasets except news20-b. The performance difference observed here is largely due to the differences in the implementations (e.g. feature vector representation, QP solver, etc.). Nevertheless, both BMRM and SVM<sup>perf</sup> are significantly outperformed by LSBMRM and LIBLINEAR on all datasets, and LIBLINEAR is almost always faster than LSBMRM. It is clear from the figure that LSBMRM and LIBLINEAR enjoy progression with "strictly" decreasing objective values; whereas the progress of both BMRM and SVM<sup>perf</sup> are hindered by the "stalling" steps (i.e., the flat line segments in the plots). The fact that LSBMRM is different from BMRM and SVM<sup>perf</sup> by one additional line search step implies that the "stalling" steps is the time that BMRM and SVM<sup>perf</sup> improve the approximation at the regions which do not help reducing the primal objective function value.

In the case of logistic regression, we compare BMRM to the state of the art trust region Newton method for logistic regression (Lin et al., 2008) which is also available in the LIBLINEAR package (option "-s 0"). From Figure 10, we see that LIBLINEAR outperforms BMRM on all datasets and that BMRM suffers from the same "stalling" phenomenon as observed in the linear SVMs case.

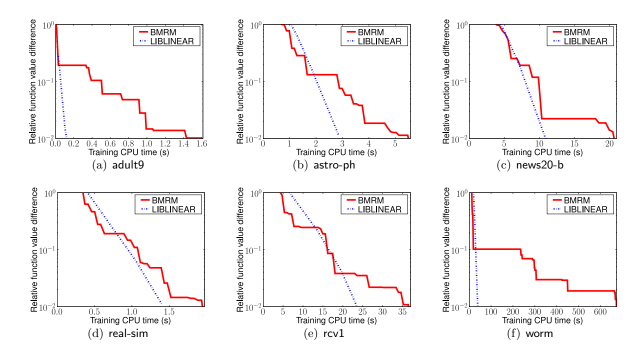


Figure 10: Logistic regression. Relative primal objective value difference during training.

### 5.3.2 Learning the cost matrix for graph matching

In computer vision, there are problems which require matching the objects of interest in a pair of images. These problems are often modeled as attributed graph matching problems where the (extracted) landmark points  $x_i$  in the first image x must be matched to the corresponding points  $x'_{i'}$  in the second image x'. Note that we represent the point  $x_i$  or  $x'_{i'}$  as d-dimensional feature vectors. The attributed graph matching problem is then cast as a Linear Assignment Problem (LAP) which can be solved in worst case  $O(n^3)$  time where n is the number of landmark points (Kuhn, 1955). <sup>14</sup> Formally, the LAP reads

$$\max_{y \in \mathcal{Y}} \sum_{i=1}^{n} \sum_{i'=1}^{n} y_{ii'} C_{ii'}, \tag{40}$$

where  $\mathcal{Y}$  is the set of all  $n \times n$  permutation matrices, and  $C_{ii'}$  is the cost of matching point  $x_i$  to point  $x'_{i'}$ . In the standard setting of graph matching, one way to determine the cost matrix C is as

$$C_{ii'} := -\sum_{k=1}^{d} \left| x_i^{(k)} - x_{i'}^{\prime(k)} \right|^2. \tag{41}$$

<sup>14.</sup> To achieve better matching results, one could further enforce edge-to-edge matching where edge refers to pair of landmark points. This additional matching requirement renders the problem as a *Quadratic Assignment Problem*.

Instead of finding more features to describe the points  $x_i$  and  $x'_{i'}$  that might improve the matching results, Caetano et al. (2007) proposed to learn a weighting to a given set of features that actually improved the matching results in many cases (Caetano et al., 2008).

In (Caetano et al., 2007, 2008) the problem of learning the cost matrix for graph matching is formulated as a  $L_2$  regularized risk minimization with loss function

$$l(x, x', y, w) = \max_{\bar{y} \in \mathcal{Y}} \langle w, \phi(x, x', \bar{y}) - \phi(x, x', y) \rangle + \Delta(\bar{y}, y), \tag{42}$$

where the feature map  $\phi$  is defined as

$$\phi(x, x', y) = -\sum_{i=1}^{n} \sum_{i'=1}^{n} y_{ii'} (|x_i^{(1)} - x_{i'}^{\prime(1)}|^2, \dots, |x_i^{(d)} - x_{i'}^{\prime(d)}|^2), \tag{43}$$

and the label loss  $\Delta$  is defined as the normalized Hamming loss

$$\Delta(\bar{y}, y) = 1 - \frac{1}{n} \sum_{i=1}^{n} \sum_{i'=1}^{n} \bar{y}_{ii'} y_{ii'}.$$
(44)

By (43) and (44), the argument of (42) becomes

$$\langle w, \phi(x, x', \bar{y}) - \phi(x, x', y) \rangle + \Delta(\bar{y}, y) = \sum_{i=1}^{n} \sum_{i'=1}^{n} \bar{y}_{ii'} \tilde{C}_{ii'} + \text{constant}, \tag{45}$$

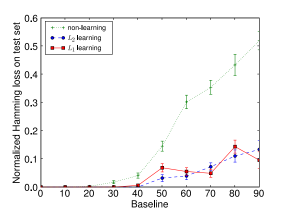
where  $\tilde{C}_{ii'} = -\sum_{k=1}^d w_k |x_i^{(k)} - x_{i'}^{\prime(k)}|^2 - y_{ii'}/n$ . Therefore, (42) is exactly a LAP. We refer interested readers to (Caetano et al., 2007, 2008) for more detailed exposition especially on the use of edge matching (in addition to point matching) which leads to much better performance.

We reproduced the experiment in (Caetano et al., 2008) that used BMRM with  $L_2$  regularization on the CMU house dataset<sup>15</sup>. For this dataset, there are 30 hand-labeled landmark points in each image and the features for those points are the 60-dimensional Shape Context features (Belongie et al., 2001; Caetano et al., 2008). The experiments evaluated the performance of the method for training/validation/testing pairs fixed at baselines (separation of frames)  $0, 10, \ldots, 90$ . Additionally, we ran the same set of experiments with  $L_1$  regularization i.e.,  $\Omega(w) = ||w||_1$ . The matching performance of the cost matrices augmented with learned weight vectors w's are compared with the original non-learning cost matrix i.e., with uniform weight vector  $w = (1, \ldots, 1)$ .

Figure 11 shows the results of the experiments. On the left, we see that the matching performance with learned cost matrices are getting more superior to that of non-learning as the baseline increases. The performance of  $L_1$  and  $L_2$  regularized learning are quite similar on average. On the right are the best learned weights for the features using  $L_1$  regularization (top) and  $L_2$  regularization (bottom) for baseline 50. The weights due to  $L_1$  regularization is considerably sparser (i.e., 42 non-zeros) than that due to  $L_2$  regularization (i.e., 52 non-zeros).

<sup>15.</sup> This dataset consist of a sequence of 111 images of a toy house. Available at http://vasc.ri.cmu.edu/idb/html/motion/house/index.html

<sup>16.</sup> Further description on  $L_1$  regularized BMRM can be found in Appendix C



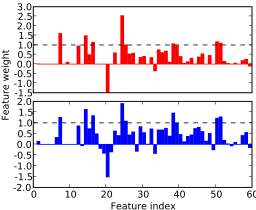


Figure 11: **Left:** Performance on the house dataset as the baseline varies. For each baseline, the minimizer of validation loss is evaluated on all testing examples. The corresponding mean normalized Hamming losses (as points) and its standard errors (as error bars) are reported. **Right:** Feature weights for best models trained with  $L_1$  regularization (top) and  $L_2$  regularization (bottom) for baseline 50. Dashed lines indicate the feature weight value 1.

### 5.3.3 Human action segmentation and recognition

In this section, we consider the problem of joint segmentation and recognition of human action from a video sequence using the discriminative Semi-Markov Models (SMM) proposed by Shi et al. (2008). Denote by  $x = \{x_i\}_{i=1}^n \in \mathcal{X}$  a sequence of n video frames, and by  $y = \{(s_i, c_i)\}_{i=1}^{\bar{n}} \in \mathcal{Y}$  the corresponding segment labeling where  $s_i$  is the starting location of the i-th segment which ends at  $s_{i+1} - 1$ ,  $c_i$  is the frame label for all frames in the segment, and  $\bar{n} \leq n$  the number of segments. For ease of presentation, we append a dummy video frame  $x_{n+1}$  to x and a dummy segment label  $(s_{\bar{n}+1}, c_{\bar{n}+1})$  to y to mark  $x_{n+1}$  as the last segment.

In SMM, there exists a segment variable for each possible segment (i.e., multiple frames) of x that model the frame label and the boundaries (or length) of a segment jointly; these segment variables form a Markov Chain. On the contrary, the Hidden Markov Model (HMM) for the same video x has one frame label variable  $y_i$  for each video frame  $x_i$ . The fact that SMM models multiple frames as one variable allows one to exploit the structure and information in the problem more efficiently than in HMM. The structure exploitation is due to the fact that one human action usually spans several consecutive frames, and the information exploitation is due to the possibility to extract features which only become apparent within a segment of several frames.

The discriminative SMM in (Shi et al., 2008) is formulated as a regularized risk minimization problem where the loss function is

$$l(x, y, w) = \max_{\bar{y} \in \mathcal{Y}} \langle w, \phi(x, \bar{y}) - \phi(x, y) \rangle + \Delta(\bar{y}, y). \tag{46}$$

The feature map  $\phi$  is defined as

$$\phi(x,y) = \left(\sum_{i=1}^{\bar{n}} \phi_1(x,s_i,c_i), \sum_{i=1}^{\bar{n}} \phi_2(x,s_i,s_{i+1},c_i), \sum_{i=1}^{\bar{n}} \phi_3(x,s_i,s_{i+1},c_i,c_{i+1})\right), \tag{47}$$

where  $\phi_1, \phi_2$ , and  $\phi_3$  are some feature functions for the segment boundaries, segments, and adjacent segments, respectively. Let  $y_i$  be the frame label for  $x_i$  according to segment labeling y, the label loss function  $\Delta$  is defined as

$$\Delta(\bar{y}, y) = \sum_{i=1}^{n} \mathbf{I}(\bar{y}_i \neq y_i), \tag{48}$$

where  $I(\cdot)$  is an indicator function as defined in (36). We refer interested readers to (Shi et al., 2008) for more details on the features and the dynamic programming to compute (46) and its subgradient.

We followed the experimental setup in (Shi et al., 2008) by running BMRM for this problem with  $L_2$  (i.e.,  $\Omega(w) = \frac{1}{2} ||w||^2$ ) and  $L_1$  (i.e.,  $\Omega(w) = ||w||_1$ ) regularization, on the Walk-Bend-Draw (WBD) datasets (Shi et al., 2008) which consists of 18 video sequences with 3 human action classes (i.e., waking, bending, drawing). For this dataset, the dimensionality of the image of the feature map  $\phi$  is d = 9917.

Table 4 shows the 6 fold cross validation results for our methods ( $L_1$  and  $L_2$  SMM)<sup>17</sup>, SVMs and SVM-HMM (Tsochantaridis et al., 2005). The latter two are adopted from (Shi et al., 2008). SMM outperforms SVM-HMM and SVM as reported in (Shi et al., 2008). Amongst  $L_1$  and  $L_2$  SMMs, the latter performs the best and converged to optimal. Although  $L_1$  SMM failed to satisfy the termination criterion, the performance is comparable to that of  $L_2$  SMM even with a 40 times *sparser* weight vector (see Figure 12 for the feature weights distributions of  $L_1$  and  $L_2$  SMMs).

Methods	CV mean (std. err.)	#iter.	CPU seconds	nnz(w)
$L_2 \text{ SMM}$	0.931 (0.009)	231	1129	3690
$L_1$ SMM	$0.925 \ (0.009)$	500	2659	84
SVM-HMM	0.870 (0.020)	_	_	_
SVM	$0.840 \ (0.030)$	_	_	_

Table 4: Experimental results on WBD dataset. The second column indicates the mean and standard error of the test accuracy (48). The third and fourth columns indicate the number of iterations and CPU seconds for the training of the final model with the best parameter, and the last column indicates the number of nonzero in the final weight vector w.

### 6. Discussion and Conclusion

The experiments presented in the paper indicate that BMRM is suitable for a wide variety of machine learning problems. In fact, the *modularity* of BMRM not only brings the benefits of

<sup>17.</sup> We set termination criterion  $\epsilon = 10^{-4}$  and limited the maximum number of iteration to 500.

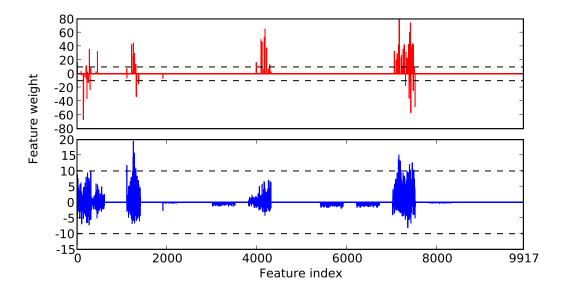


Figure 12: Feature weights for best models trained with  $L_1$  regularization (top) and  $L_2$  regularization (bottom). Dashed lines indicate the feature weight range [ $\pm 10$ ].

parallel and distributed computation but also makes BMRM a natural test bed for trying out new models/ideas on any particular problem with less effort i.e., the user is only required to implement the loss functions and/or regularizers corresponding to different models/ideas.

Nevertheless, we saw in the experiments that BMRM does not guarantee strict improvement in the primal when the dual is solved instead. This phenomenon could significantly hinder the performance of BMRM as seen in some of the experiments. Since efficient line search procedure may not exist for general structured prediction tasks, the *trust region* philosophy used in BT could be a potential strategy to alleviate this problem; we leave this to the future work. We also note that for computationally expensive nonsmooth loss functions, one way to make fuller use of each loss function evaluation is by updating the model  $R_t^{\rm CP}$  with two or more linearizations at a non-diffferentiable location (Frangioni, 1997).

In conclusion, we have presented a variant of standard bundle methods *i.e.*, BMRM, which is algorithmically simpler and, in some senses, more straightforward for regularized risk minimization problems than the standard bundle methods. We also showed a  $O(1/\epsilon)$  rate of convergence for nonsmooth objective functions and  $O(\log(1/\epsilon))$  rates for smooth objective functions.

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### Appendix A. Loss Functions

A multitude of loss functions are commonly used to derive seemingly different algorithms. This often blurs the similarities as well as subtle differences between them, often for historic reasons: Each new loss is typically accompanied by at least one publication dedicated to it. In many cases, the loss is not spelled out explicitly either but instead, it is only given by means of a constrained optimization problem. A case in point are the papers introducing (binary) hinge loss (Bennett and Mangasarian, 1992; Cortes and Vapnik, 1995) and structured loss (Taskar et al., 2004; Tsochantaridis et al., 2005). Likewise, a geometric description obscures the underlying loss function, as in novelty detection (Schölkopf et al., 2001).

Table 5: Scalar loss functions and their derivatives, depending on  $f := \langle w, x \rangle$ , and y.

	Loss $\bar{l}(f,y)$	Derivative $\bar{l}'(f,y)$
Hinge (Bennett and Mangasarian, 1992)	$\max(0, 1 - yf)$	0 if $yf \ge 1$ and $-y$ otherwise
Squared Hinge (Keerthi and DeCoste, 2005)	$\frac{1}{2}\max(0, 1 - yf)^2$	0 if $yf \ge 1$ and $f - y$ otherwise
Exponential (Cowell et al., 1999)	$\exp(-yf)$	$-y\exp(-yf)$
Logistic (Collins et al., 2000)	$\log(1 + \exp(-yf))$	$-y/(1+\exp(yf))$
Novelty (Schölkopf et al., 2001)	$\max(0, 1-f)$	0 if $f \ge 1$ and $-1$ otherwise
Least mean squares (Williams, 1998)	$\frac{1}{2}(f-y)^2$	$\mid f-y \mid$
Least absolute deviation	f-y	$\mid \operatorname{sgn}(f-y)$
Quantile regression (Koenker, 2005)	$\max(\tau(f-y), (1-\tau)(y-f))$	$\tau$ if $f > y$ and $\tau - 1$ otherwise
$\epsilon$ -insensitive (Vapnik et al., 1997)	$\max(0, f-y -\epsilon)$	0 if $ f - y  \le \epsilon$ , else $\operatorname{sgn}(f - y)$
Huber's robust loss (Müller et al., 1997)	$\frac{1}{2}(f-y)^2$ if $ f-y  < 1$ , else $ f-y  - \frac{1}{2}$	$ f-y \text{ if }  f-y  \le 1, \text{ else } \operatorname{sgn}(f-y)$
Poisson regression (Cressie, 1993)	$\exp(f) - yf$	$\mid \exp(f) - y$

Table 6: Vectorial loss functions and their derivatives, depending on the vector f := Wx and on y.

M(f-y)	$\frac{1}{2}(f-y)^{\top}M(f-y)$ where $M \succeq 0$ $\tilde{M}(f-y)$	Multivariate Regression
$\left[ \left[ \sum_{y'} e_{y'} \exp(f'_y) \right] / \sum_{y'} \exp(f'_y) - e_y \right]$	$\log \sum_{y'} \exp(f_{y'}) - f_y$	Softmax Multiclass (Cowell et al., 1999)
where $y^*$ is the argmax of the loss		(Tsochantaridis et al., 2005)
$\Gamma(y,y')(e_{y^*}-e_y)$	$\left  \max_{y'} \Gamma(y, y') (f_{y'} - f_y + \Delta(y, y')) \right  \Gamma(y, y') (e_{y^*} - e_y)$	Scaled Soft-Margin Multiclass
where $y^*$ is the argmax of the loss		(Crammer and Singer, 2003)
$e_{y^*} - e_y$	$\max_{y'}(f_{y'} - f_y + \Delta(y, y'))$	Soft-Margin Multiclass (Taskar et al., 2004)   $\max_{y'}(f_{y'} - f_y + \Delta(y, y'))$
Derivative	Loss	

In this section we give an expository yet unifying presentation of many of those loss functions. Many of them are well known, while others, such as multivariate ranking, hazard regression, or Poisson regression are not commonly used in machine learning. Tables 5 and 6 contain a choice subset of simple scalar and vectorial losses. Our aim is to put the multitude of loss functions in an unified framework, and to show how these losses and their (sub)gradients can be computed efficiently for use in our solver framework.

Note that not all losses, while convex, are continuously differentiable. In this situation we give a subgradient. While this may not be optimal, the convergence rates of our algorithm do not depend on which element of the subdifferential we provide: in all cases the first order Taylor approximation is a lower bound which is tight at the point of expansion.

In this setion, with little abuse of notation,  $v_i$  is understood as the *i*-th component of vector v when v is clearly not an element of a sequence or a set.

#### A.1 Scalar Loss Functions

It is well known (Wahba, 1997) that the convex optimization problem

$$\min_{\xi} \ \xi \text{ subject to } y \langle w, x \rangle \ge 1 - \xi \text{ and } \xi \ge 0$$
 (49)

takes on the value  $\max(0, 1-y\langle w, x\rangle)$ . The latter is a convex function in w and x. Likewise, we may rewrite the  $\epsilon$ -insensitive loss, Huber's robust loss, the quantile regression loss, and the novelty detection loss in terms of loss functions rather than a constrained optimization problem. In all cases,  $\langle w, x \rangle$  will play a key role insofar as the loss is convex in terms of the scalar quantity  $\langle w, x \rangle$ . A large number of loss functions fall into this category, as described in Table 5. Note that not all functions of this type are continuously differentiable. In this case we adopt the convention that

$$\partial_x \max(f(x), g(x)) = \begin{cases} \partial_x f(x) & \text{if } f(x) \ge g(x) \\ \partial_x g(x) & \text{otherwise} \end{cases}$$
 (50)

Since we are only interested in obtaining an arbitrary element of the subdifferential this convention is consistent with our requirements.

Let us discuss the issue of efficient computation. For all scalar losses we may write  $l(x, y, w) = \bar{l}(\langle w, x \rangle, y)$ , as described in Table 5. In this case a simple application of the chain rule yields that  $\partial_w l(x, y, w) = \bar{l}'(\langle w, x \rangle, y) \cdot x$ . For instance, for squared loss we have

$$\bar{l}(\langle w, x \rangle, y) = \frac{1}{2}(\langle w, x \rangle - y)^2$$
 and  $\bar{l}'(\langle w, x \rangle, y) = \langle w, x \rangle - y$ .

Consequently, the derivative of the empirical risk term is given by

$$\partial_w R_{\text{emp}}(w) = \frac{1}{m} \sum_{i=1}^m \bar{l}'(\langle w, x_i \rangle, y_i) \cdot x_i.$$
 (51)

This means that if we want to compute l and  $\partial_w l$  on a large number of observations  $x_i$ , represented as matrix X, we can make use of fast linear algebra routines to pre-compute the vectors

$$f = Xw \text{ and } g^{\top}X \text{ where } g_i = \overline{l}'(f_i, y_i).$$
 (52)

### **Algorithm 6** ScalarLoss(w, X, y)

- 1: **input:** Weight vector w, feature matrix X, and labels y
- 2: Compute f = Xw
- 3: Compute  $r = \sum_{i} \bar{l}(f_i, y_i)$  and  $g = \bar{l}'(f, y)$
- 4:  $g \leftarrow g^{\top} X$
- 5: **return** Risk r and gradient g

This is possible for any of the loss functions listed in Table 5, and many other similar losses. The advantage of this unified representation is that implementation of each individual loss can be done in very little time. The computational infrastructure for computing Xw and  $g^{T}X$  is shared. Evaluating  $\bar{l}(f_{i}, y_{i})$  and  $\bar{l}'(f_{i}, y_{i})$  for all i can be done in O(m) time and it is not time-critical in comparison to the remaining operations. Algorithm 6 describes the details.

An important but often neglected issue is worth mentioning. Computing f requires us to right multiply the matrix X with the vector w while computing f requires the left multiplication of f with the vector f. If f is stored in a row major format then f can be computed rather efficiently while f is expensive. This is particularly true if f cannot fit in main memory. Converse is the case when f is stored in column major format. Similar problems are encountered when f is a sparse matrix and stored in either compressed row format or in compressed column format.

#### A.2 Structured Loss

In recent years structured estimation has gained substantial popularity in machine learning (Tsochantaridis et al., 2005; Taskar et al., 2004; Bakir et al., 2007). At its core it relies on two types of convex loss functions: logistic loss:

$$l(x, y, w) = \log \sum_{y' \in \mathcal{Y}} \exp\left(\left\langle w, \phi(x, y') \right\rangle\right) - \left\langle w, \phi(x, y) \right\rangle, \tag{53}$$

and soft-margin loss:

$$l(x, y, w) = \max_{y' \in \mathcal{Y}} \Gamma(y, y') \left\langle w, \phi(x, y') - \phi(x, y) \right\rangle + \Delta(y, y'). \tag{54}$$

Here  $\phi(x,y)$  is a *joint* feature map,  $\Delta(y,y') \geq 0$  describes the cost of misclassifying y by y', and  $\Gamma(y,y') \geq 0$  is a scaling term which indicates by how much the large margin property should be enforced. For instance, Taskar et al. (2004) choose  $\Gamma(y,y')=1$ . On the other hand Tsochantaridis et al. (2005) suggest  $\Gamma(y,y')=\Delta(y,y')$ , which reportedly yields better performance. Finally, McAllester (2007) recently suggested generic functions  $\Gamma(y,y')$ .

The logistic loss can also be interpreted as the negative log-likelihood of a conditional exponential family model:

$$p(y|x;w) := \exp(\langle w, \phi(x,y) \rangle - g(w|x)), \tag{55}$$

where the normalizing constant g(w|x), often called the log-partition function, reads

$$g(w|x) := \log \sum_{y' \in \mathcal{Y}} \exp\left(\left\langle w, \phi(x, y') \right\rangle\right). \tag{56}$$

As a consequence of the Hammersley-Clifford theorem (Jordan, 2002) every exponential family distribution corresponds to a undirected graphical model. In our case this implies that the labels y factorize according to an undirected graphical model. A large number of problems have been addressed by this setting, amongst them named entity tagging (Lafferty et al., 2001), sequence alignment (Tsochantaridis et al., 2005), segmentation (Rätsch et al., 2007) and path planning (Ratliff et al., 2006). It is clearly impossible to give examples of all settings in this section, nor would a brief summary do this field any justice. We therefore refer the reader to the edited volume Bakir et al. (2007) and the references therein.

If the underlying graphical model is tractable then efficient inference algorithms based on dynamic programming can be used to compute (53) and (54). We discuss intractable graphical models in Section A.2.1, and now turn our attention to the derivatives of the above structured losses.

When it comes to computing derivatives of the logistic loss, (53), we have

$$\partial_w l(x, y, w) = \frac{\sum_{y'} \phi(x, y') \exp \langle w, \phi(x, y') \rangle}{\sum_{y'} \exp \langle w, \phi(x, y') \rangle} - \phi(x, y)$$
 (57)

$$= \mathbf{E}_{y' \sim p(y'|x)} \left[ \phi(x, y') \right] - \phi(x, y). \tag{58}$$

where p(y|x) is the exponential family model (55). In the case of (54) we denote by  $\bar{y}(x)$  the argmax of the RHS, that is

$$\bar{y}(x) := \underset{y'}{\operatorname{argmax}} \Gamma(y, y') \left\langle w, \phi(x, y') - \phi(x, y) \right\rangle + \Delta(y, y'). \tag{59}$$

This allows us to compute the derivative of l(x, y, w) as

$$\partial_w l(x, y, w) = \Gamma(y, \bar{y}(x)) \left[ \phi(x, \bar{y}(x)) - \phi(x, y) \right]. \tag{60}$$

In the case where the loss is maximized for more than one distinct value  $\bar{y}(x)$  we may average over the individual values, since any convex combination of such terms lies in the subdifferential.

Note that (54) majorizes  $\Delta(y, y^*)$ , where  $y^* := \operatorname{argmax}_{y'} \langle w, \phi(x, y') \rangle$  (Tsochantaridis et al., 2005). This can be seen via the following series of inequalities:

$$\Delta(y, y^*) \le \Gamma(y, y^*) \langle w, \phi(x, y^*) - \phi(x, y) \rangle + \Delta(y, y^*) \le l(x, y, w).$$

The first inequality follows because  $\Gamma(y, y^*) \ge 0$  and  $y^*$  maximizes  $\langle w, \phi(x, y') \rangle$  thus implying that  $\Gamma(y, y^*) \langle w, \phi(x, y^*) - \phi(x, y) \rangle \ge 0$ . The second inequality follows by definition of the loss.

We conclude this section with a simple lemma which is at the heart of several derivations of Joachims (2005). While the proof in the original paper is far from trivial, it is straightforward in our setting:

**Lemma 8** Denote by  $\delta(y, y')$  a loss and let  $\phi(x_i, y_i)$  be a feature map for observations  $(x_i, y_i)$  with  $1 \leq i \leq m$ . Moreover, denote by X, Y the set of all m patterns and labels respectively. Finally let

$$\Phi(X,Y) := \sum_{i=1}^{m} \phi(x_i, y_i) \text{ and } \Delta(Y,Y') := \sum_{i=1}^{m} \delta(y_i, y_i').$$
 (61)

Then the following two losses are equivalent:

$$\sum_{i=1}^{m} \max_{y'} \left\langle w, \phi(x_i, y') - \phi(x_i, y_i) \right\rangle + \delta(y_i, y') \text{ and } \max_{Y'} \left\langle w, \Phi(X, Y') - \Phi(X, Y) \right\rangle + \Delta(Y, Y').$$

This is immediately obvious, since both feature map and loss decompose, which allows us to perform maximization over Y' by maximizing each of its m components. In doing so, we showed that aggregating all data and labels into a single feature map and loss yields results identical to minimizing the sum over all individual losses. This holds, in particular, for the sample error loss of Joachims (2005). Also note that this equivalence does *not* hold whenever  $\Gamma(y, y')$  is not constant.

### A.2.1 Intractable Models

We now discuss cases where computing l(x, y, w) itself is too expensive. For instance, for intractable graphical models, the computation of  $\sum_{y} \exp \langle w, \phi(x, y) \rangle$  cannot be computed efficiently. Wainwright and Jordan (2003) propose the use of a convex majorization of the log-partition function in those cases. In our setting this means that instead of dealing with

$$l(x, y, w) = g(w|x) - \langle w, \phi(x, y) \rangle \text{ where } g(w|x) := \log \sum_{y} \exp \langle w, \phi(x, y) \rangle$$
 (62)

one uses a more easily computable convex upper bound on g via

$$\sup_{\mu \in MARG(x)} \langle w, \mu \rangle + H_{Gauss}(\mu|x). \tag{63}$$

Here MARG(x) is an outer bound on the conditional marginal polytope associated with the map  $\phi(x,y)$ . Moreover,  $H_{Gauss}(\mu|x)$  is an upper bound on the entropy by using a Gaussian with identical variance. More refined tree decompositions exist, too. The key benefit of our approach is that the solution  $\mu$  of the optimization problem (63) can immediately be used as a gradient of the upper bound. This is computationally rather efficient.

Likewise note that Taskar et al. (2004) use relaxations when solving structured estimation problems of the form

$$l(x, y, w) = \max_{y'} \Gamma(y, y') \left\langle w, \phi(x, y') - \phi(x, y) \right\rangle + \Delta(y, y'), \tag{64}$$

by enlarging the domain of maximization with respect to y'. For instance, instead of an integer programming problem we might relax the setting to a linear program which is much cheaper to solve. This, again, provides an upper bound on the original loss function.

In summary, we have demonstrated that convex relaxation strategies are well applicable for bundle methods. In fact, the results of the corresponding optimization procedures can be used directly for further optimization steps.

#### A.3 Scalar Multivariate Performance Scores

We now discuss a series of structured loss functions and how they can be implemented efficiently. For the sake of completeness, we give a concise representation of previous work on multivariate performance scores and ranking methods. All these loss functions rely on having access to  $\langle w, x \rangle$ , which can be computed efficiently by using the same operations as in Section A.1.

### **Algorithm 7** ROCScore(X, y, w)

```
1: input: Feature matrix X, labels y, and weight vector w
2: initialization: s_- = m_- and s_+ = 0 and l = \mathbf{0}_m and c = Xw - \frac{1}{2}y
3: \pi \leftarrow \{1, \ldots, m\} sorted in ascending order of c
4: for i = 1 to m do
5: if y_{\pi_i} = -1 then
6: l_{\pi_i} \leftarrow s_+ and s_- \leftarrow s_- - 1
7: else
8: l_{\pi_i} \leftarrow -s_- and s_+ \leftarrow s_+ + 1
9: end if
10: end for
11: Rescale l \leftarrow l/(m_+m_-) and compute r = \langle l, c \rangle and g = l^\top X.
12: return Risk r and subgradient g
```

## A.3.1 ROC Score

Denote by f = Xw the vector of function values on the training set. It is well known that the area under the ROC curve is given by

$$AUC(x, y, w) = \frac{1}{m_{+}m_{-}} \sum_{y_{i} < y_{j}} \mathbf{I}(\langle w, x_{i} \rangle < \langle w, x_{j} \rangle), \tag{65}$$

where  $m_+$  and  $m_-$  are the numbers of positive and negative observations respectively, and  $\mathbf{I}(\cdot)$  is indicator function. Directly optimizing the cost  $1 - \mathrm{AUC}(x, y, w)$  is difficult as it is not continuous in w. By using  $\max(0, 1 + \langle w, x_i - x_j \rangle)$  as the surrogate loss function for all pairs (i, j) for which  $y_i < y_j$  we have the following convex multivariate empirical risk

$$R_{\text{emp}}(w) = \frac{1}{m_{+}m_{-}} \sum_{y_{i} < y_{j}} \max(0, 1 + \langle w, x_{i} - x_{j} \rangle) = \frac{1}{m_{+}m_{-}} \sum_{y_{i} < y_{j}} \max(0, 1 + f_{i} - f_{j}).$$
 (66)

Obviously, we could compute  $R_{\text{emp}}(w)$  and its derivative by an  $O(m^2)$  operation. However Joachims (2005) showed that both can be computed in  $O(m \log m)$  time using a sorting operation, which we now describe.

Denote by  $c = f - \frac{1}{2}y$  an auxiliary variable and let i and j be indices such that  $y_i = -1$  and  $y_j = 1$ . It follows that  $c_i - c_j = 1 + f_i - f_j$ . The efficient algorithm is due to the observation that there are at most m distinct terms  $c_k$ ,  $k = 1, \ldots, m$ , each with different frequency  $l_k$  and sign, appear in (66). These frequencies  $l_k$  can be determined by first sorting c in ascending order then scanning through the labels according to the sorted order of c and keeping running statistics such as the number  $s_-$  of negative labels yet to encounter, and the number  $s_+$  of positive labels encountered. When visiting  $y_k$ , we know  $c_k$  should appears  $s_+$  (or  $s_-$ ) times with positive (or negative) sign in (66) if  $y_k = -1$  (or  $y_k = 1$ ). Algorithm 7 spells out explicitly how to compute  $R_{\text{emp}}(w)$  and its subgradient.

#### A.3.2 Ordinal Regression

Essentially the same preference relationships need to hold for ordinal regression. The only difference is that  $y_i$  need not take on binary values any more. Instead, we may have an

arbitrary number of different values  $y_i$  (e.g. 1 corresponding to 'strong reject' up to 10 corresponding to 'strong accept', when it comes to ranking papers for a conference). That is, we now have  $y_i \in \{1, ..., n\}$  rather than  $y_i \in \{\pm 1\}$ . Our goal is to find some w such that  $\langle w, x_i - x_j \rangle < 0$  whenever  $y_i < y_j$ . Whenever this relationship is not satisfied, we incur a cost  $C(y_i, y_j)$  for preferring  $x_i$  to  $x_j$ . For examples,  $C(y_i, y_j)$  could be constant i.e.,  $C(y_i, y_j) = 1$  (Joachims, 2006) or linear i.e.,  $C(y_i, y_j) = y_j - y_i$ .

Denote by  $m_i$  the number of  $x_j$  for which  $y_j = i$ . In this case, there are  $\bar{M} = m^2 - \sum_{i=1}^n m_i^2$  pairs  $(y_i, y_j)$  for which  $y_i \neq y_j$ ; this implies that there are  $M = \bar{M}/2$  pairs  $(y_i, y_j)$  such that  $y_i < y_j$ . Normalizing by the total number of comparisons we may write the overall cost of the estimator as

$$\frac{1}{M} \sum_{y_i < y_j} C(y_i, y_j) \mathbf{I}(\langle w, x_i \rangle > \langle w, x_j \rangle) \text{ where } M = \frac{1}{2} \left[ m^2 - \sum_i^n m_i^2 \right].$$
 (67)

Using the same convex majorization as above when we were maximizing the ROC score, we obtain an empirical risk of the form

$$R_{\text{emp}}(w) = \frac{1}{M} \sum_{y_i < y_j} C(y_i, y_j) \max(0, 1 + \langle w, x_i - x_j \rangle)$$
 (68)

Now the goal is to find an efficient algorithm for obtaining the number of times when the individual losses are nonzero such as to compute both the value and the gradient of  $R_{\rm emp}(w)$ . The complication arises from the fact that observations  $x_i$  with label  $y_i$  may appear in either side of the inequality depending on whether  $y_j < y_i$  or  $y_j > y_i$ . This problem can be solved as follows: sort f = Xw in ascending order and traverse it while keeping track of how many items with a lower value  $y_j$  are no more than 1 apart in terms of their value of  $f_i$ . This way we may compute the count statistics efficiently. Algorithm 8 describes the details, generalizing the results of Joachims (2006). Again, its runtime is  $O(m \log m)$ , thus allowing for efficient computation.

#### A.3.3 Preference Relations

In general, our loss may be described by means of a set of preference relations  $j \succeq i$  for arbitrary pairs  $(i,j) \in \{1,\ldots m\}^2$  associated with a cost C(i,j) which is incurred whenever i is ranked above j. This set of preferences may or may not form a partial or a total order on the domain of all observations. In these cases efficient computations along the lines of Algorithm 8 exist. In general, this is not the case and we need to rely on the fact that the set P containing all preferences is sufficiently small that it can be enumerated efficiently. The risk is then given by

$$\frac{1}{|P|} \sum_{(i,j)\in P} C(i,j) \mathbf{I}(\langle w, x_i \rangle > \langle w, x_j \rangle) \tag{69}$$

# **Algorithm 8** OrdinalRegression(X, y, w, C)

```
1: input: Feature matrix X, labels y, weight vector w, and score matrix C
 2: initialization: l = \mathbf{0}_n and u_i = m_i \ \forall i \in [n] and r = 0 and g = \mathbf{0}_m
 3: Compute f = Xw and set c = [f - \frac{1}{2}, f + \frac{1}{2}] \in \mathbb{R}^{2m} (concatenate the vectors)
 4: Compute M = (m^2 - \sum_{i=1}^n m_i^2)/2
 5: Rescale C \leftarrow C/M
 6: \pi \leftarrow \{1, \dots, 2m\} sorted in ascending order of c
 7: for i = 1 to 2m do
       j = \pi_i \mod m
       if \pi_i \leq m then
 9:
10:
          for k = 1 to y_i - 1 do
             r \leftarrow r - C(k, y_j) u_k c_j
11:
12:
             g_j \leftarrow g_j - C(k, y_j)u_k
13:
          l_{y_j} \leftarrow l_{y_j} + 1
14:
15:
          for k = y_i + 1 to n do
16:
             r \leftarrow r + C(y_i, k) l_k c_{i+m}
17:
             g_j \leftarrow g_j + C(y_j, k)l_k
18:
          end for
19:
          u_{y_j} \leftarrow u_{y_j} - 1
20:
       end if
21:
22: end for
23: g \leftarrow g^{\top} X
24: return: Risk r and subgradient q
```

Again, the same majorization argument as before allows us to write a convex upper bound

$$R_{\text{emp}}(w) = \frac{1}{|P|} \sum_{(i,j) \in P} C(i,j) \max \left(0, 1 + \langle w, x_i \rangle - \langle w, x_j \rangle\right)$$
 (70)

where 
$$\partial_w R_{\text{emp}}(w) = \frac{1}{|P|} \sum_{(i,j) \in P} C(i,j) \begin{cases} 0 & \text{if } \langle w, x_j - x_i \rangle \ge 1 \\ x_i - x_j & \text{otherwise} \end{cases}$$
 (71)

The implementation is straightforward, as given in Algorithm 9.

#### A.3.4 Ranking

In webpage and document ranking we are often in a situation similar to that described in Section A.3.2, however with the difference that we do not only care about objects  $x_i$  being ranked according to scores  $y_i$  but moreover that different degrees of importance are placed on different documents.

The information retrieval literature is full with a large number of different scoring functions. Examples are criteria such as Normalized Discounted Cumulative Gain (NDCG), Mean Reciprocal Rank (MRR), Precision@n, or Expected Rank Utility (ERU). They are

# **Algorithm 9** Preference(X, w, C, P)

```
1: input: Feature matrix X, weight vector w, score matrix C, and preference set P

2: initialization: r = 0 and g = \mathbf{0}_m

3: Compute f = Xw

4: while (i,j) \in P do

5: if f_j - f_i < 1 then

6: r \leftarrow r + C(i,j)(1 + f_i - f_j)

7: g_i \leftarrow g_i + C(i,j) and g_j \leftarrow g_j - C(i,j)

8: end if

9: end while

10: g \leftarrow g^\top X

11: return Risk r and subgradient g
```

used to address the issue of evaluating rankers, search engines or recommender sytems (Voorhees, 2001; Jarvelin and Kekalainen, 2002; Breese et al., 1998; Basilico and Hofmann, 2004). For instance, in webpage ranking only the first k retrieved documents that matter, since users are unlikely to look beyond the first k, say 10, retrieved webpages in an internet search. Le and Smola (2007) show that these scores can be optimized directly by minimizing the following loss:

$$l(X, y, w) = \max_{\pi} \sum_{i} c_i \langle w, x_{\pi(i)} - x_i \rangle + \langle a - a(\pi), b(y) \rangle.$$
 (72)

Here  $c_i$  is a monotonically decreasing sequence, the documents are assumed to be arranged in order of decreasing relevance,  $\pi$  is a permutation, the vectors a and b(y) depend on the choice of a particular ranking measure, and  $a(\pi)$  denotes the permutation of a according to  $\pi$ . Pre-computing f = Xw we may rewrite (72) as

$$l(f, y) = \max_{\pi} \left[ c^{\mathsf{T}} f(\pi) - a(\pi)^{\mathsf{T}} b(y) \right] - c^{\mathsf{T}} f + a^{\mathsf{T}} b(y)$$
 (73)

and consequently the derivative of l(X, y, w) with respect to w is given by

$$\partial_w l(X, y, w) = (c(\bar{\pi}^{-1}) - c)^\top X \text{ where } \bar{\pi} = \operatorname*{argmax}_{\pi} c^\top f(\pi) - a(\pi)^\top b(y). \tag{74}$$

Here  $\pi^{-1}$  denotes the inverse permutation, such that  $\pi \circ \pi^{-1} = 1$ . Finding the permutation maximizing  $c^{\top} f(\pi) - a(\pi)^{\top} b(y)$  is a linear assignment problem which can be easily solved by the Hungarian Marriage algorithm, that is, the Kuhn-Munkres algorithm.

The original papers by Kuhn (1955) and Munkres (1957) implied an algorithm with  $O(m^3)$  cost in the number of terms. Later, Karp (1980) suggested an algorithm with expected quadratic time in the size of the assignment problem (ignoring log-factors). Finally, Orlin and Lee (1993) propose a linear time algorithm for large problems. Since in our case the number of pages is fairly small (in the order of 50 to 200 per query) the scaling behavior per query is not too important. We used an existing implementation due to Jonker and Volgenant (1987).

Note also that training sets consist of a *collection* of ranking problems, that is, we have several ranking problems of size 50 to 200. By means of parallelization we are able to

### **Algorithm 10** Ranking(X, y, w)

- 1: **input:** Feature matrix X, relevances y, and weight vector w
- 2: Compute vectors a and b(y) according to some ranking measure
- 3: Compute f = Xw
- 4: Compute elements of matrix  $C_{ij} = c_i f_j b_i a_j$
- 5:  $\pi = \text{LinearAssignment}(C)$
- 6:  $r = c^{\top} (f(\pi) f) + (a a(\pi))^{\top} b$
- 7:  $g = c(\pi^{-1}) c$  and  $g \leftarrow g^{\top} X$
- 8: **return** Risk r and subgradient g

distribute the work onto a cluster of workstations, which is able to overcome the issue of the rather costly computation per collection of queries. Algorithm 10 spells out the steps in detail.

## A.3.5 Contingency Table Scores

Joachims (2005) observed that  $F_{\beta}$  scores and related quantities dependent on a contingency table can also be computed efficiently by means of structured estimation. Such scores depend in general on the number of true and false positives and negatives alike. Algorithm 11 shows how a corresponding empirical risk and subgradient can be computed efficiently. As with the previous losses, here again we use convex majorization to obtain a tractable optimization problem.

Given a set of labels y and an estimate y', the numbers of true positives  $(T_+)$ , true negatives  $(T_-)$ , false positives  $(F_+)$ , and false negatives  $(F_-)$  are determined according to a contingency table as follows:

	y > 0	y < 0
y' > 0	$T_{+}$	$F_{+}$
y' < 0	$F_{-}$	$T_{-}$

In the sequel, we denote by  $m_+ = T_+ + F_-$  and  $m_- = T_- + F_+$  the numbers of positives and negative labels in y, respectively. We note that  $F_{\beta}$  score can be computed based on the contingency table (Joachims, 2005) as

$$F_{\beta}(T_{+}, T_{-}) = \frac{(1+\beta^{2})T_{+}}{T_{+} + m_{-} - T_{-} + \beta^{2}m_{+}}.$$
 (75)

If we want to use  $\langle w, x_i \rangle$  to estimate the label of observation  $x_i$ , we may use the following structured loss to "directly" optimize w.r.t.  $F_{\beta}$  score (Joachims, 2005):

$$l(X, y, w) = \max_{y'} \left[ (y' - y)^{\top} f + \Delta(T_+, T_-) \right], \tag{76}$$

where f = Xw,  $\Delta(T_+, T_-) := 1 - F_{\beta}(T_+, T_-)$ , and  $(T_+, T_-)$  is determined by using y and y'. Since  $\Delta$  does not depend on the specific choice of (y, y') but rather just on which sets they disagree, l can be maximized as follows: Enumerating all possible  $m_+m_-$  contingency tables in a way such that given a configuration  $(T_+, T_-)$ ,  $T_+$   $(T_-)$  positive (negative) observations  $x_i$  with largest (lowest) value of  $\langle w, x_i \rangle$  are labeled as positive (negative). This is

# Algorithm 11 $F_{\beta}(X, y, w)$

```
1: input: Feature matrix X, labels y, and weight vector w
  2: Compute f = Xw
  3: \pi^+ \leftarrow \{i: y_i = 1\} sorted in descending order of f
  4: \pi^- \leftarrow \{i : y_i = -1\} sorted in ascending order of f
 5: Let p_0 = 0 and p_i = 2\sum_{k=i}^{m_+} f_{\pi_k^+}, i = 1, \dots, m_+
 6: Let n_0 = 0 and n_i = 2 \sum_{k=i}^{m_-} f_{\pi_k}^-, i = 1, \dots, m_-
  7: y' \leftarrow -y and r \leftarrow -\infty
  8: for i = 0 to m_{+} do
         for j = 0 to m_- do
            r_{\rm tmp} = \Delta(i,j) - p_i + n_j
10:
            if r_{\rm tmp} > r then
11:
                r \leftarrow r_{\text{tmp}}
12:
               T_+ \leftarrow i \text{ and } T_- \leftarrow j
13:
            end if
14:
         end for
15:
16: end for
17: y'_{\pi_{i}^{+}} \leftarrow 1, i = 1, \dots, T_{+}
18: y'_{\pi_{i}^{-}} \leftarrow -1, i = 1, \dots, T_{-}
19: q \leftarrow (y'-y)^{\top} X
20: return Risk r and subgradient q
```

effectively implemented as a nested loop hence run in  $O(m^2)$  time. Algorithm 11 describes the procedure in details.

# A.4 Vector Loss Functions

Next we discuss "vector" loss functions, *i.e.*, functions where w is best described as a matrix (denoted by W) and the loss depends on Wx. Here, we have feature vector  $x \in \mathbb{R}^d$ , label  $y \in \mathbb{R}^k$ , and weight matrix  $W \in \mathbb{R}^{d \times k}$ . We also denote feature matrix  $X \in \mathbb{R}^{m \times d}$  as a matrix of m feature vectors  $x_i$ , and stack up the columns  $W_i$  of W as a vector w.

Some of the most relevant cases are multiclass classification using both the exponential families model and structured estimation, hierarchical models, *i.e.*, ontologies, and multivariate regression. Many of those cases are summarized in Table 6.

#### A.4.1 Unstructured Setting

The simplest loss is multivariate regression, where  $l(x, y, W) = \frac{1}{2}(y - x^{\top}W)^{\top}M(y - x^{\top}W)$ . In this case it is clear that by pre-computing XW subsequent calculations of the loss and its gradient are significantly accelerated.

A second class of important losses is given by plain multiclass classification problems, e.g. recognizing digits of a postal code or categorizing high-level document categories. In this case,  $\phi(x,y)$  is best represented by  $e_y \otimes x$  (using a linear model). Clearly we may view  $\langle w, \phi(x,y) \rangle$  as an operation which chooses a column indexed by y from xW, since all labels

y correspond to a different weight vector  $W_y$ . Formally we set  $\langle w, \phi(x, y) \rangle = [xW]_y$ . In this case, structured estimation losses can be rewritten as

$$l(x, y, W) = \max_{y'} \Gamma(y, y') \left\langle W_{y'} - W_y, x \right\rangle + \Delta(y, y') \tag{77}$$

and 
$$\partial_W l(x, y, W) = \Gamma(y, y^*)(e_{y^*} - e_y) \otimes x.$$
 (78)

Here  $\Gamma$  and  $\Delta$  are defined as in Section A.2 and  $y^*$  denotes the value of y' for which the RHS of (77) is maximized. This means that for unstructured multiclass settings we may simply compute xW. Since this needs to be performed for all observations  $x_i$  we may take advantage of fast linear algebra routines and compute f = XW for efficiency. Likewise note that computing the gradient over m observations is now a matrix-matrix multiplication, too: denote by G the matrix of rows of gradients  $\Gamma(y_i, y_i^*)(e_{y_i^*} - e_{y_i})$ . Then  $\partial_W R_{\text{emp}}(X, y, W) = G^\top X$ . Note that G is very sparse with at most two nonzero entries per row, which makes the computation of  $G^\top X$  essentially as expensive as two matrix vector multiplications. Whenever we have many classes, this may yield significant computational gains.

Log-likelihood scores of exponential families share similar expansions. We have

$$l(x, y, W) = \log \sum_{y'} \exp \langle w, \phi(x, y') \rangle - \langle w, \phi(x, y) \rangle = \log \sum_{y'} \exp \langle W_{y'}, x \rangle - \langle W_y, x \rangle$$
 (79)

$$\partial_W l(x, y, W) = \frac{\sum_{y'} (e_{y'} \otimes x) \exp \langle W_{y'}, x \rangle}{\sum_{y'} \exp \langle W_{y'}, x \rangle} - e_y \otimes x.$$
 (80)

The main difference to the soft-margin setting is that the gradients are *not* sparse in the number of classes. This means that the computation of gradients is slightly more costly.

#### A.4.2 Ontologies

Assume that the labels we want to estimate can be found to belong to a directed acyclic graph. For instance, this may be a gene-ontology graph (Ashburner et al., 2000) a patent hierarchy (Cai and Hofmann, 2004), or a genealogy. In these cases we have a hierarchy of categories to which an element x may belong. Figure 13 gives two examples of such directed acyclic graphs (DAG). The first example is a binary tree, while the second contains nodes with different numbers of children (e.g. node 4 and 12), nodes at different levels having children (e.g. nodes 5 and 12), and nodes which have more than one parent (e.g. node 5). It is a well known fundamental property of trees that they have at most as many internal nodes as they have leaf nodes.

It is now our goal to build a classifier which is able to categorize observations according to which leaf node they belong to (each leaf node is assigned a label y). Denote by k+1 the number of nodes in the DAG including the root node. In this case we may design a feature map  $\phi(y) \in \mathbb{R}^k$  (Cai and Hofmann, 2004) by associating with every label y the vector describing the path from the root node to y, ignoring the root node itself. For instance, for the first DAG in Figure 13 we have

$$\phi(8) = (1,0,1,0,0,0,1,0,0,0,0,0,0,0)$$
 and  $\phi(13) = (0,1,0,0,1,0,0,0,0,0,0,1,0,0)$ 

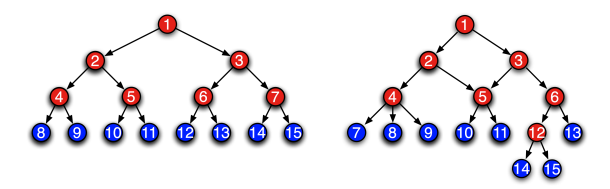


Figure 13: Two ontologies. **Left:** a binary hierarchy with internal nodes  $\{1, \ldots, 7\}$  and labels  $\{8, \ldots 15\}$ . **Right:** a generic directed acyclic graph with internal nodes  $\{1, \ldots, 6, 12\}$  and labels  $\{7, \ldots, 11, 13, \ldots, 15\}$ . Note that node 5 has two parents, namely nodes 2 and 3. Moreover, the labels need not be found at the same level of the tree: nodes 14 and 15 are one level lower than the rest of the nodes.

Whenever several paths are admissible, as in the right DAG of Figure 13 we average over all possible paths. For example, we have

$$\phi(10) = (0.5, 0.5, 0.1, 0.0, 0.0, 0.1, 0.0, 0.0, 0.0)$$
 and  $\phi(15) = (0.1, 0.0, 1, 0.0, 0.0, 0.0, 0.1, 0.0, 0.1)$ .

Also note that the lengths of the paths need not be the same (e.g. to reach 15 it takes a longer path than to reach 13). Likewise, it is natural to assume that  $\Delta(y, y')$ , i.e., the cost for mislabeling y as y' will depend on the similarity of the path. In other words, it is likely that the cost for placing x into the wrong sub-sub-category is less than getting the main category of the object wrong.

To complete the setting, note that for  $\phi(x,y) = \phi(y) \otimes x$  the cost of computing all labels is k inner products, since the value of  $\langle w, \phi(x,y) \rangle$  for a particular y can be obtained by the sum of the contributions for the segments of the path. This means that the values for *all* terms can be computed by a simple breadth first traversal through the graph. As before, we may make use of vectorization in our approach, since we may compute  $xW \in \mathbb{R}^k$  to obtain the contributions on all segments of the DAG before performing the graph traversal. Since we have m patterns  $x_i$  we may vectorize matters by pre-computing XW.

Also note that  $\phi(y) - \phi(y')$  is nonzero only for those edges where the paths for y and y' differ. Hence we only change weights on those parts of the graph where the categorization differs. Algorithm 12 describes the subgradient and loss computation for the soft-margin type of loss function.

The same reasoning applies to estimation when using an exponential families model. The only difference is that we need to compute a *soft-max* over paths rather than exclusively choosing the best path over the ontology. Again, a breadth-first recursion suffices: each of the leaves y of the DAG is associated with a probability p(y|x). To obtain  $\mathbf{E}_{y\sim p(y|x)}[\phi(y)]$  all we need to do is perform a bottom-up traversal of the DAG summing over all proba-

# **Algorithm 12** Ontology(X, y, W)

```
    input: Feature matrix X ∈ ℝ<sup>m×d</sup>, labels y, and weight matrix W ∈ ℝ<sup>d×k</sup>
    initialization: G = 0 ∈ ℝ<sup>m×k</sup> and r = 0
    Compute f = XW and let f<sub>i</sub> = x<sub>i</sub>W
    for i = 1 to m do
    Let D<sub>i</sub> be the DAG with edges annotated with the values of f<sub>i</sub>
    Traverse D<sub>i</sub> to find node y* that maximize sum of f<sub>i</sub> values on the path plus Δ(y<sub>i</sub>, y')
    G<sub>i</sub> = φ(y*) - φ(y<sub>i</sub>)
    r ← r + z<sub>y*</sub> - z<sub>y<sub>i</sub></sub>
    end for
    g = G<sup>T</sup>X
    return Risk r and subgradient g
```

bility weights on the path. Wherever a node has more than one parent, we distribute the probability weight equally over its parents.

### Appendix B. Proofs

#### Proof of Theorem 4

To show Theorem 4 we need several technical intermediate steps. Let  $\gamma_t := J(w_t) - J_t(w_t)$  and recall that  $\epsilon_t := \min_{t' \leq t} J(w_{t'}) - J_t(w_t)$ . The following lemma establishes some useful properties of  $\gamma_t$  and  $\epsilon_t$ .

**Lemma 9** We have  $J_{t'}(w_{t'}) \leq J_t(w_t) \leq J(w^*) \leq J(w_t) = J_{t+1}(w_t)$  for all  $t' \leq t$ . Furthermore,  $\epsilon_t$  is monotonically decreasing with  $\epsilon_t - \epsilon_{t+1} \geq J_{t+1}(w_{t+1}) - J_t(w_t) \geq 0$ . Also,  $\epsilon_t$  upper bounds the distance from optimality via  $\gamma_t \geq \epsilon_t \geq \min_{t' \leq t} J(w_{t'}) - J(w^*)$ .

**Proof** Since  $J_{t'}(w) \leq J_t(w) \leq J(w)$  for all  $t' \leq t$  this property also applies to their respective minima. Moreover, since  $w^*$  minimizes J(w) we have  $J(w^*) \leq J(w_t)$ . Since Taylor expansions are exact at the point of expansion  $J(w_t) = J_{t+1}(w_t)$ . The first inequality follows from the definition of  $\epsilon_t$ , and the fact that  $J_t$  is monotonically increasing. Finally, since  $J_{t'+1}(w_{t'}) = J(w_{t'})$  it is easy to see that  $\gamma_t \geq \epsilon_t = \min_{t' \leq t} J(w_{t'}) - J_t(w_t) \geq \min_{t' \leq t} J(w_{t'}) - J(w^*)$ .

Our second technical lemma allows us to bound the maximum value of a concave function provided that we know its first derivative and a bound on the second derivative.

**Lemma 10** Denote by  $f:[0,1] \to \mathbb{R}$  a concave function with f(0)=0, f'(0)=l, and  $|f''(x)| \le K \ \forall x \in [0,1]$ . Then we have  $\max_{x \in [0,1]} f(x) \ge \frac{l}{2} \min(\frac{l}{K}, 1)$ .

**Proof** We first observe that  $g(x) := lx - \frac{K}{2}x^2 \le f(x) \ \forall x \text{ implies } \max_{x \in [0,1]} f(x) \ge \max_{x \in [0,1]} g(x)$ . g attains the unconstrained maximum  $\frac{l^2}{2K}$  at  $x = \frac{l}{K}$ . Since g is monotonically increasing in  $[0, \frac{l}{K}]$ , if l > K we pick x = 1 which yields constrained maximum  $l - \frac{K}{2} > \frac{l}{2}$ . Taking the minimum over both maxima proves the claim.

To apply the above result, we need to compute the gradient and Hessian of  $J_{t+1}^*(\alpha)$  with respect to the search direction  $((1-\eta)\alpha_t, \eta)$ . The following lemma takes care of the gradient:

**Lemma 11** Denote by  $\alpha_t$  the solution of (23) at time instance t. Moreover, denote by  $\bar{A} = [A, a_{t+1}]$  and  $\bar{b} = [b, b_{t+1}]$  the extended matrices and vectors needed to define the dual problem for step t+1, and let  $\bar{\alpha} \in \mathbb{R}^{t+1}$ . Then the following holds:

$$\partial_{\bar{\alpha}} J_{t+1}^*([\alpha_t, 0]) = \bar{A}^\top w_t + \bar{b} \text{ and}$$
(81)

$$[-\alpha_t, 1]^{\top} \left[ \bar{A}^{\top} w_t + \bar{b} \right] = J_{t+1}(w_t) - J_t(w_t) = \gamma_t$$
(82)

**Proof** By the dual connection  $\partial \Omega^*(-\lambda^{-1}A\alpha_t) = w_t$ . Hence we have that  $\partial_{\bar{\alpha}} - \lambda \Omega^*(-\lambda^{-1}\bar{A}\bar{\alpha}) + \bar{\alpha}^{\top}\bar{b} = \bar{A}^{\top}w_t + \bar{b}$  for  $\bar{\alpha} = [\alpha_t, 0]^{\top}$ . This proves the first claim. To see the second part we eliminate  $\xi$  from of the Lagrangian (25) and write the partial Lagrangian

$$L(w,\alpha) = \lambda \Omega(w) + \alpha^{\top} \left( A^{\top} w + b \right) \text{ with } \alpha \ge 0.$$
 (83)

The result follows by noting that at optimality  $L(w_t, \alpha_t) = J_t(w_t)$  and  $J_{t+1}(w_t) = \lambda \Omega(w_t) + \langle w_t, a_{t+1} \rangle + b_{t+1}$ . Consequently we have

$$J_{t+1}(w_t) - J_t(w_t) = \lambda \Omega(w_t) + \langle w_t, a_{t+1} \rangle + b_{t+1} - \lambda \Omega(w_t) - \alpha_t (A^{\top} w_t + b_t)$$

Rearranging terms proves the claim.

To apply Lemma 10 we also need to bound the second derivative.

**Lemma 12** Under the assumptions of Lemma 11 we have

$$\partial_{\bar{\alpha}}^2 J_{t+1}^*(\bar{\alpha}) = -\lambda^{-1} \bar{A}^\top \partial^2 \Omega^*(-\lambda^{-1} \bar{A}\bar{\alpha}) \bar{A}$$
 (84)

moreover 
$$\bar{A}[-\alpha_t, 1] = \partial_w J(w_t)$$
 (85)

**Proof** The first equality is immediate from the chain rule. Next note that  $\partial_w \Omega(w_t) = -\lambda^{-1} A \alpha_t$  by dual connection. Since  $a_{t+1} = \partial_w R_{\text{emp}}(w_t)$  the claim follows from  $J(w) = R_{\text{emp}}(w) + \lambda \Omega(w)$ .

This result allows us to express the second derivative of the dual objective function (24) in terms of the gradient of the risk functional. The idea is that as we approach optimality, the second derivative will vanish. We will use this fact to argue that for continuously differentiable losses  $R_{\rm emp}(w)$  we enjoy linear convergence throughout.

**Proof** [Theorem 4] We overload the notation for  $J_{t+1}^*$  by defining the following one dimensional concave function

$$J_{t+1}^*(\eta) := J_{t+1}^*([(1-\eta)\alpha_t, \eta]) = -\lambda \Omega^*(-\lambda^{-1}\bar{A}[(1-\eta)\alpha_t^\top, \eta]) + [(1-\eta)\alpha_t^\top, \eta]\bar{b}.$$
 (86)

Clearly,  $J_{t+1}^*(0) = J_t(w_t)$ . Furthermore, by (82), (84), and (85) it follows that

$$\partial_{\eta} J_{t+1}^*(\eta)|_{\eta=0} = [-\alpha_t, 1]^{\top} \partial_{\bar{\alpha}} J_{t+1}^*([\alpha_t, 0]) = \gamma_t \text{ and}$$
 (87)

$$\partial_{\eta}^{2} J_{t+1}^{*}(\eta) = -\lambda^{-1} [-\alpha_{t}, 1]^{\top} \bar{A}^{\top} \partial^{2} \Omega^{*} (-\lambda^{-1} \bar{A} [(1 - \eta)\alpha_{t}, \eta]) \bar{A} [-\alpha_{t}, 1]^{\top}$$
(88)

$$= -\lambda^{-1} [\partial_w J(w_t)]^{\top} \partial^2 \Omega^* (-\lambda^{-1} \bar{A}[(1-\eta)\alpha_t, \eta]) [\partial_w J(w_t)] := r.$$
 (89)

By our assumption on  $\|\partial^2 \Omega^*\| \leq H^*$  we have

$$|r| \le H^* \|\partial_w J(w_t)\|^2 / \lambda.$$

Next we need to bound the gradient of J. For this purpose note that  $\partial_w \lambda \Omega(w_t) = -A^\top \alpha_t$  and moreover that  $\|\alpha_t\|_1 = 1$ . This implies that  $\partial_w \lambda \Omega(w_t)$  lies in the convex hull of the past gradients,  $a_{t'}$ . By our assumption on  $\|\partial_w R_{\text{emp}}(w)\|$  it follows that  $\|\partial_w \lambda \Omega(w_t)\| \leq G$ . We conclude that

$$\|\partial_w J(w_t)\|^2 \le 4G^2 \text{ and } |r| \le 4G^2 H^*/\lambda.$$

Invoking Lemma 10 on  $J_{t+1}^*(\eta) - J_t(w_t)$  shows that

$$J_{t+1}^*(\eta) - J_t(w_t) \ge \frac{\gamma_t}{2} \min(1, \lambda \gamma_t / 4G^2 H^*).$$

We now upper bound the LHS of the above inequality as follows:

$$\epsilon_t - \epsilon_{t+1} \ge J_{t+1}(w_{t+1}) - J_t(w_t) \ge J_{t+1}^*(\eta) - J_t(w_t) \ge \frac{\gamma_t}{2} \min(1, \lambda \gamma_t / 4G^2 H^*).$$
 (90)

The first inequality follows from Lemma 9 while the second follows by observing that  $J_{t+1}(w_{t+1}) = J_{t+1}^*(\alpha_{t+1}) \ge J_{t+1}^*(\eta)$ . The RHS of the third inequality on the other hand can be lower bounded by observing that  $\gamma_t \ge \epsilon_t$ , which follows from Lemma 9. This in turn obtains (27).

For the second part note that (27) already yields the  $\epsilon_t/2$  decrease when  $\epsilon_t \geq 4G^2H^*/\lambda$ . To show the other parts we need to show that the gradient of the regularized risk vanishes as we converge to the optimal solution. Towards this end, we apply Lemma 10 in the primal.<sup>18</sup> This allows us to bound  $\|\partial_w J(w_t)\|$  in terms of  $\gamma_t$ . Plugging in the first and second derivative of  $J(w_t)$  we obtain

$$\gamma_t \ge \frac{1}{2} \left\| \partial_w J(w_t) \right\| \min(1, \left\| \partial_w J(w_t) \right\| / H).$$

If  $\|\partial_w J(w_t)\| > H$ , then  $\gamma_t \ge \frac{1}{2} \|\partial_w J(w_t)\|$  which in turn yields  $|r| \le 4\gamma_t^2 H^*/\lambda$ . Plugging this into Lemma 10 yields a lower bound on the improvement of  $\lambda/8H^*$ .

Finally, for  $\|\partial_w J(w_t)\| \le H$  we have  $\gamma_t \ge \|\partial_w J(w_t)\|^2 / 2H$ , which implies  $|r| \le 2HH^*\gamma_t/\lambda$ . Plugging this into Lemma 10 yields an improvement of  $\lambda \gamma_t / 4HH^* \ge \lambda \epsilon_t / 4HH^*$ .

Since both cases cover the remaining range of convergence, the minimum  $\min(\lambda/8H^*, \lambda\epsilon_t/4HH^*)$  provides a lower bound for the improvement. The crossover point between both terms occurs

<sup>18.</sup> Define  $\bar{J}(\eta) := J(w_t) - J(w_t + \eta p)$  where  $p = -\frac{\partial_w J(w_t)}{\|\partial_w J(w_t)\|}$  is the unit-length gradient. We see that  $\frac{d}{d\eta} \bar{J}(\eta)\Big|_{\eta=0} = [-\partial_w J(w_t + \eta p)^\top p]|_{\eta=0} = \|\partial_w J(w_t)\|$ , and  $\bar{J}(0) = 0$ . Hence Lemma 10 is applicable in this case.

at  $\epsilon_t = H/2$ . Rearranging the conditions leads to the (pessimistic) improvement guarantees of the second claim.

Note that a key step in the above analysis involved bounding  $r := \partial_{\eta}^2 J_{t+1}^*(\eta)$ . For a number of regularizers tighter bounds can be obtained. The following bounds are essentially due to Shalev-Shwartz and Singer (2006):

- For squared norm regularization, i.e.,  $\Omega^*(\mu) = \frac{1}{2} \|\mu\|_2^2$  we have  $r = \|\partial_w J(w_t)\|_2^2$ .
- For  $L_p$  norm regularization, i.e.,  $\Omega^*(\mu) = \frac{1}{2} \|\mu\|_q^2$  we have  $r \leq (q-1) \|\partial_w J(w_t)\|_q^2$ .
- For quadratic form regularization with PD matrix B, i.e.,  $\Omega^*(\mu) = \frac{1}{2}\mu B^{-1}\mu$ , we have  $r = \partial_w J(w_t)^\top B^{-1} \partial_w J(w_t)$ .
- For unnormalized entropic regularization we have  $\partial_{\mu}^{2}\Omega^{*}(\mu) = \operatorname{diag}\left(e^{\mu^{(1)}}, \dots, e^{\mu^{(d)}}\right)$ . Hence we may bound  $r \leq \|\partial_{w}J(w_{t})\|_{2}^{2}\exp\left(\|\mu\|_{\infty}\right)$ . Clearly this bound may be very loose whenever  $\mu$  has only very few large coefficients.
- For normalized entropy regularization, i.e.,  $\Omega^*(\mu) = \log \sum_i \exp \mu^{(i)}$  we have  $r \leq \|\partial_w J(w_t)\|_{\infty}^2$ .

### Proof of Theorem 5

We need the following technical lemma for the proof:

**Lemma 13** Let  $\langle \rho_1, \rho_2, \ldots \rangle$  be a sequence of non-negative numbers satisfying the following recurrence, for  $t \geq 1$ :  $\rho_t - \rho_{t+1} \geq c(\rho_t)^2$ , where c > 0 is a constant. Then for all integers  $t \geq 1$ ,

$$\rho_t \le \frac{1}{c(t-1+\frac{1}{\rho_1 c})}.$$

Furthermore  $\rho_t \leq \rho$  whenever

$$t \le \frac{1}{c\rho} - \frac{1}{\rho_1 c} + 1.$$

This is Sublemma 5.4 of Abe et al. (2001) which is easily proven by induction. Now we can prove the main result.

**Proof** [Theorem 5] For any  $\epsilon_t > 4G^2H^*/\lambda$  it follows from (27) that  $\epsilon_{t+1} \leq \epsilon_t/2$ . Moreover,  $\epsilon_0 \leq J(0)$ , since we know that J is non-negative. Hence we need at most  $\log_2[\lambda J(0)/4G^2H^*]$  to achieve this level of precision. Subsequently we have

$$\epsilon_t - \epsilon_{t+1} \ge \frac{\lambda}{8G^2H^*} \epsilon_t^2. \tag{91}$$

Invoking Lemma 13 by setting  $c = \frac{\lambda}{8G^2H^*}$  and  $\rho_1 = 4G^2H^*/\lambda$  shows that  $\epsilon_t \leq \epsilon$  after at most  $\frac{8G^2H^*}{\lambda\epsilon} - 1$  more steps. This proves the first claim.

To analyze convergence in the second case we need to study two additional phases: for  $\epsilon_t \in [H/2, 4G^2H^*/\lambda]$  we see constant progress. Hence it takes us  $4\lambda^{-2}[8G^2(H^*)^2 - HH^*\lambda]$ 

steps to cover this interval. Finally in the third phase we have  $\epsilon_{t+1} \leq \epsilon_t [1 - \lambda/4HH^*]$ . Starting from  $\epsilon_t = H/2$  we need  $\log_2[2\epsilon/H]/\log_2[1 - \lambda/4HH^*]$  steps to converge. Expanding the logarithm in the denominator close to 1 proves the claim.

### B.1 Proof of Theorem 7

We first note that the termination criterion of Algorithm 3 is slightly different from that of Algorithm 2. In order to apply the convergence results for Algorithm 2 to Algorithm 3 we redefine the following notations:

$$\epsilon_t := J(w_t^b) - J_t(w_t) \tag{92}$$

$$a_{t+1} := \partial_w R_{\text{emp}}(w_t^c), \tag{93}$$

$$b_{t+1} := R_{\text{emp}}(w_t^c) - \langle w_t, a_{t+1} \rangle,$$
 (94)

where

$$w_t^b := \underset{\eta}{\operatorname{argmin}} J(w_{t-1}^b + \eta(w_t - w_{t-1}^b)), \text{ and}$$
 (95)

$$w_t^c := (1 - \theta)w_t^b + \theta w_t. \tag{96}$$

Then we state and prove the following lemma which is crucial to the application of Lemma 11 in the proof.

**Lemma 14** 
$$J_{t+1}(w_t) = \lambda \Omega(w_t) + \langle w_t, a_{t+1} \rangle + b_{t+1}$$

**Proof**  $w_t^b$  is the optimal value of J on the line joining  $w_t$  and  $w_{t-1}^b$  while  $w_t^c$  is a convex combination of  $w_t$  and  $w_t^b$ . Moreover by definition of  $a_{t+1}$  and  $b_{t+1}$  we have  $J(w_t^c) = J_{t+1}(w_t^c)$ . Therefore,

$$J(w_t^c) = J_{t+1}(w_t^c) = \lambda \Omega(w_t^c) + \langle a_{t+1}, w_t^c \rangle + b_{t+1} \ge J(w_t^b). \tag{97}$$

But since  $\Omega$  is convex

$$\Omega(\underbrace{(1-\theta)w_t^b + \theta w_t}_{w_t^c}) \le (1-\theta)\Omega(w_t^b) + \theta\Omega(w_t),$$

which can be rearranged to

$$\theta(\Omega(w_t^b) - \Omega(w_t)) \le \Omega(w_t^b) - \Omega(w_t^c).$$

Multiplying by  $\lambda$  and adding and subtracting  $\theta R_{\text{emp}}(w_t^b)$  and  $\theta R_t(w_t)$  respectively to the above equation

$$\underbrace{\lambda\theta\Omega(w_t^b) + \theta R_{\text{emp}}(w_t^b)}_{\theta J(w_t^b)} - \underbrace{\lambda\theta\Omega(w_t) - \theta R_t(w_t)}_{\theta J_t(w_t)} \leq \underbrace{\lambda\Omega(w_t^b) + R_{\text{emp}}(w_t^b)}_{J(w_t^b)} - \lambda\Omega(w_t^c) - (1 - \theta)R_{\text{emp}}(w_t^b) - \theta R_t(w_t).$$

Plugging in (92) obtains

$$\theta \epsilon_t \le J(w_t^b) - \lambda \Omega(w_t^c) - (1 - \theta) R_{\text{emp}}(w_t^b) - \theta R_t(w_t). \tag{98}$$

Putting (97) and (98) together

$$\langle a_{t+1}, w_t^c \rangle + b_{t+1} \ge J(w_t^b) - \lambda \Omega(w_t^c) \ge (1 - \theta) R_{\text{emp}}(w_t^b) + \theta R_t(w_t) + \theta \epsilon_t. \tag{99}$$

Since  $w_t^c = (1 - \theta)w_t^b + \theta w_t$  it follows that

$$(1-\theta)\left\langle a_{t+1}, w_t^b \right\rangle + \theta \left\langle a_{t+1}, w_t \right\rangle + b_{t+1} \ge (1-\theta)R_{\text{emp}}(w_t^b) + \theta R_t(w_t) + \theta \epsilon_t.$$

Which can be rearranged to

$$(1 - \theta) \left( \left\langle a_{t+1}, w_t^b \right\rangle - R_{\text{emp}}(w_t^b) \right) + \theta \left( \left\langle a_{t+1}, w_t \right\rangle - R_t(w_t) \right) + b_{t+1} \ge \theta \epsilon_t.$$

Since  $\langle w_t^b, a_{t+1} \rangle + b_{t+1}$  is the Taylor approximation of the convex function  $R_{\text{emp}}$  around  $w_t^c$  evaluated at  $w_t^b$  it follows that  $R_{\text{emp}}(w_t^b) \geq \langle w_t^b, a_{t+1} \rangle + b_{t+1}$ . Plugging this into the above equation yields

$$(1-\theta)(-b_{t+1}) + \theta(\langle w_t, a_{t+1} \rangle - R_t(w_t)) + b_{t+1} \ge \theta \epsilon_t.$$

Dividing by  $\theta > 0$  and rearranging yields

$$\langle w_t, a_{t+1} \rangle + b_{t+1} \ge R_t(w_t) + \epsilon_t.$$

The conclusion of the lemma follows from observing that  $R_{t+1}(w_t) = \max(\langle w_t, a_{t+1} \rangle + b_{t+1}, R_t(w_t)) = \langle w_t, a_{t+1} \rangle + b_{t+1}$  and  $J_{t+1}(w_t) = \lambda \Omega(w_t) + R_{t+1}(w_t)$ .

We also need the following two lemmas before we can proceed to the final proof.

**Lemma 15** 
$$\epsilon_t - \epsilon_{t+1} \ge J_{t+1}(w_{t+1}) - J_t(w_t)$$

Proof

$$\epsilon_{t} - \epsilon_{t+1} = J(w_{t}^{b}) - J_{t}(w_{t}) - J(w_{t+1}^{b}) + J_{t+1}(w_{t+1})$$

$$= \underbrace{(J(w_{t}^{b}) - J(w_{t+1}^{b}))}_{\geq 0} + J_{t+1}(w_{t+1}) - J_{t}(w_{t}) \quad \text{(by the definition of } w_{t}^{b})$$

$$\geq J_{t+1}(w_{t+1}) - J_{t}(w_{t})$$

**Lemma 16** Let  $\alpha_t$ ,  $\bar{A} := [a_1, \ldots, a_{t+1}]$ , and  $\bar{b} := [b_1, \ldots, b_{t+1}]$  be as defined in Lemma 11. Then under the assumption of Theorem 4 that  $\|\partial_w R_{\text{emp}}(w)\| \leq G$ , we have

$$[-\alpha_t, 1]^\top \bar{A}^\top \bar{A}[-\alpha_t, 1] \le 4G^2.$$

**Proof** By the dual connection,  $\partial_w \lambda \Omega(w_t) = -A\alpha_t$ . Also,  $\alpha_t \geq 0$ , and  $\|\alpha_t\|_1 = 1$  as it is the optimal solution of (24) at iteration t. It follows that  $\partial_w \lambda \Omega(w_t)$  lies in the convex hull of  $a_{t'} := \partial_w R_{\text{emp}}(w_{t'}^c) \ \forall t' \leq t$ . Therefore  $\|\partial_w \lambda \Omega(w_t)\| \leq G$ . Consequently,

$$[-\alpha_t, 1]^{\top} \bar{A}^{\top} \bar{A}[-\alpha_t, 1] = \|\partial_w \lambda \Omega(w_t) + \partial_w R_{\text{emp}}(w_t^c)\|^2$$
$$= \|\partial_w \lambda \Omega(w_t)\|^2 + 2\partial_w \lambda \Omega(w_t)^{\top} \partial_w R_{\text{emp}}(w_t^c) + \|\partial_w R_{\text{emp}}(w_t^c)\|^2 \le 4G^2,$$

by Cauchy-Schwarz inequality.

Finally, we sketch the proof for Theorem 7.

**Proof** [Theorem 7] (Sketch) Theorem 4 holds for Algorithm 3 by applying Lemmas 14, 15, and 16 into the first part of the proof. Therefore, for  $\epsilon < 4G^2H^*/\lambda$ , (90) reduces to  $\epsilon_t - \epsilon_{t+1} \ge \lambda \epsilon_t / 4G^2H^*$ . Applying Lemma 13 yields  $\epsilon_t \le \frac{1}{c\left(t-1+\frac{1}{\epsilon_1c}\right)}$ , with  $c = \lambda/8G^2H^*$ .

Setting 
$$\frac{1}{c\left(t-1+\frac{1}{\epsilon_1c}\right)}=\epsilon$$
, assuming that  $\epsilon_1>0$ , and solving for  $n$  yields  $n\leq \frac{1}{c\epsilon}=\frac{8G^2H^*}{\lambda\epsilon}$ .

# Appendix C. $L_1$ Regularized BMRM

Following our convention, the  $L_1$  norm regularized BMRM reads

$$\min_{\xi, w} \xi + \lambda \|w\|_1 \text{ subject to } w^{\top} a_i + b_i \le \xi, \ i = 1, \dots, t.$$
 (100)

An equivalent formulation is

$$\min_{\xi, w} \xi \text{ subject to } w^{\top} a_i + b_i \le \xi, \ i = 1, \dots, t \text{ and } \|w\|_1 \le \tau,$$
 (101)

where one can show a monotone correspondence between  $\tau$  and the  $\lambda$  in (100) by comparison of the KKT conditions for the two problems.

Note that our convergence proof does not apply in this case as the Fenchel dual of  $\Omega(w) = ||w||_1$  fails to satisfy the strong convexity assumption. Nevertheless, we see that (101) can be easily solved by CPM where the solution must lie in the  $L_1$  ball of radius  $\tau$ . Finally, we note that the  $L_1$  regularized BMRM can be written in a rather standard linear programming (LP) formulation:

$$\min_{\xi, u, v} \ \xi + \lambda \mathbf{1}_d^{\top} (u + v) \tag{102a}$$

s.t. 
$$a_i^{\top} u - a_i^{\top} v + b_i \le \xi, \ i = 1, \dots, t$$
 (102b)

$$u, v \ge 0, \tag{102c}$$

with the variable of interest w = u - v.