ROUTING AND ACTION MEMORANDUM

ROUTING

TO: (1) Network Sciences Division (lyer, Purush)

Report is available for review

(2) Proposal Files Proposal No.: 54231NSMUR.175

DESCRIPTION OF MATERIAL

CONTRACT OR GRANT NUMBER: W911NF-08-1-0242

INSTITUTION: University of Washington

PRINCIPAL INVESTIGATOR: Pedro Domingos

TYPE REPORT: Manuscript

DATE RECEIVED: 9/23/14 4:36PM

PERIOD COVERED: through

TITLE: Gradient Boosting for Conditional Random Fields

ACTION TAKEN BY DIVISION

(x) Report has been reviewed for technical sufficiency and IS [x] IS NOT [] satisfactory.

(x) Material has been given an OPSEC review and it has been determined to be non sensitive and, except for manuscripts and progress reports, suitable for public release.

Approved by SSL\PURUSH.S.IYER on 9/26/14 10:55AM

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16. SECURI	TY CLASSIFICA	ATION OF:	17. LIMITATION C)F	15. NUMB	ER	19a. NAME OF RESPONSIBLE PERSON
a. REPORT	b. ABSTRACT	c. THIS PAGE	ABSTRACT		OF PAGES		Pedro Domingos
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Report Title

Gradient Boosting for Conditional Random Fields

ABSTRACT

In this paper, we present a gradient boosting algorithm for tree-shaped conditional random fields (CRF). Conditional random fields are an important class of models for accurate structured prediction, but effective design of the feature functions is a major challenge when applying CRF models to real world data. Gradient boosting, which can induce and select functions, is a natural candidate solution for the problem. However, it is non-trivial to derive gradient boosting algorithms for CRFs, due to the dense Hessian matrices introduced by variable dependencies. We address this challenge by deriving a Markov Chain mixing rate bound to quantify the dependencies, and introduce a gradient boosting algorithm that iteratively optimizes an adaptive upper bound of the objective function. The resulting algorithm induces and selects features for CRFs via functional space optimization, with provable convergence guarantees. Experimental results on three real world datasets demonstrate that the mixing rate based upper bound is effective for training CRFs with non-linear potentials.

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Gradient Boosting for Conditional Random Fields

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Abstract

In this paper, we present a gradient boosting algorithm for tree-shaped conditional random fields (CRF). Conditional random fields are an important class of models for accurate structured prediction, but effective design of the feature functions is a major challenge when applying CRF models to real world data. Gradient boosting, which can induce and select functions, is a natural candidate solution for the problem. However, it is non-trivial to derive gradient boosting algorithms for CRFs, due to the dense Hessian matrices introduced by variable dependencies. We address this challenge by deriving a Markov Chain mixing rate bound to quantify the dependencies, and introduce a gradient boosting algorithm that iteratively optimizes an adaptive upper bound of the objective function. The resulting algorithm induces and selects features for CRFs via functional space optimization, with provable convergence guarantees. Experimental results on three real world datasets demonstrate that the mixing rate based upper bound is effective for training CRFs with non-linear potentials.

1 Introduction

Many problems in machine learning involve structured prediction, i.e. predicting a group of outputs 033 that depends on each other. Conditional random fields [9] are among the most successful solutions 034 to these problem. Variants of tree-shaped conditional random fields have been proposed and widely applied to structured prediction problems in domains such as natural language processing [9, 16], computer vision [7, 15] and bio-informatics [21]. As opposed to classification models that assume 037 independent output variables, CRF models capture the dependency pattern between output and input via potential functions. Potential functions are usually defined using a linear combination of carefully engineered features of the input and the output variables. These feature functions are crucial for learning accurate models. Thus, it is important to ask whether we can induce arbitrary poten-040 tial functions automatically (via functional space optimization), instead of manually crafting them 041 and/or restricting them to linear combinations. 042

043 Gradient boosting [4], which performs additive training in functional spaces, is a natural candidate 044 for this problem. Effective gradient boosting algorithms, such as LogitBoost and its variants [5, 11, 17], have been proposed for inducing feature functions for (independent) multi-class classification problems. The key ingredient in these methods is the effective use of second order information 046 via diagonal approximation of Hessian matrices. Unfortunately, it is non-trivial to develop such 047 boosting methods for CRFs, since variable interdependencies introduce dense Hessian matrices that 048 make gradient boosting infeasible due to the computational complexity. Instead, existing boosting approaches either optimize approximate objectives [20, 12] or only take first order information into account when optimizing exact likelihood [1]. Unfortunately, the convergence of this method is 051 guaranteed only with small step sizes. 052

053 In this paper, we present a novel gradient boosting algorithm for inducing non-linear feature functions for tree-shaped CRFs. The CRF training is performed via iteratively optimizing adaptive upper

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bounds of the loss function, to address the challenge of dense Hessians. The adaptive bounds, which
are derived using Markov Chain mixing rates, measure the dependency between variables, and accordingly control the conservativeness of the updates. The resulting gradient boosting algorithm,
which can be viewed as generalization of LogitBoost to structure prediction problems, optimizes
the CRF objective with provable convergence guarantees. Experimental results on three real world
datasets demonstrate the effectiveness and efficiency of our bound and the proposed algorithm.

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2 Overview of Method

Model Formalization Given input x, a CRF model describes a distribution over the outputs as

$$P(y|x) = \frac{\exp(\Phi(y,x))}{\sum_{y' \in \mathcal{V}} \exp(\Phi(y',x))}$$
(1)

where \mathcal{Y} is the set of possible output combinations, and $\Phi(y, x)$ captures the dependency between the input and output variables. The model Φ usually factorizes as a sum of unary and pairwise (edge) *potential functions* ϕ_i between individual output variables, which can be expressed as follows:

$$\Phi(y,x) = \sum_{i=1}^{m} \phi_i(x)\mu_i(y), \ \phi_i \in \mathcal{F}, \mu_i \in N \cup \mathcal{E} \quad \text{subject to } \phi_i = \phi_j \text{ for } (i,j) \in \mathcal{C},$$
(2)

073 where $N = \{\mathbf{1}(y_t = k)\}, \mathcal{E} = \{\mathbf{1}(y_s = k_1, y_t = k_2)\}$ are the sets of indicator functions of each 074 node and edge state. Each μ_i corresponds to a event $y_t = k$ or $y_s = k_1, y_t = k_2$ (depending on 075 whether ϕ_i is node or edge potential). We use μ as short hand for $\mu(y)$ and view it as a vector of 076 random variables. The family of all possible node and edge potential functions is $\mathcal{F} = \mathcal{F}_n \cup \mathcal{F}_e$, 077 whose size could be infinite. C represents equivalence classes in different parts of the model that we 078 use to capture parameter sharing, which is common in most applications of CRFs. In standard linear-079 chain CRFs with linear potentials, \mathcal{F} contains linear functions of x, and C can be used to constrain the node and edge potential functions in different position to be the same. On the other hand, LogitBoost considers arbitrary \mathcal{F} , however constrains the model to contain only node potentials 081 (there are no edge potentials). In this paper we are interested in arbitrary function families \mathcal{F} , and focus on tree-shaped \mathcal{E} that allow exact inference of marginals. 083

Training Objective Using functions ϕ in Eq.(2) as the model parameters allows us to induce potential functions automatically through functional space optimization. In particular, we generalize the standard CRF objective over training data $\mathcal{D} = \{(y, x)\}$ to the following:

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$$L(\phi) = \sum_{y,x\in\mathcal{D}} l(y,x,\phi) + \sum_{c} \Omega(\phi_c) = -\sum_{y,x\in D} \ln P(y|x) + \sum_{c} \Omega(\phi_c).$$
(3)

Here l is the negative log-likelihood function over each data point. $\sum_{c} \Omega(\phi_c)$ is a regularization term that measures the complexity of the learned function, defined as a sum over the equivalence class defined by C. In standard CRFs, for example, Ω is often the square of L_2 norm of the parameter vector. This generalized objective function encourages us to select *predictive* (i.e. optimizes l) and *simple* (i.e. optimizes Ω) functions as potentials of a CRF.

Challenges for Function Learning Since the model parameters in this formulation are functions, Eq.(3) cannot be directly optimized using traditional optimization techniques. Instead, we train the model additively: at each iteration t, our proposed algorithm first searches over the functional space \mathcal{F} to find functions $\delta = [\delta_1, \delta_2, \dots, \delta_m]$ that optimize the objective function $L(\phi^{(t)} + \delta)$, and then adds them to the current model $\phi^{(t+1)} \leftarrow \phi^{(t)} + \delta$. However, due to the complex nature of the objective function, directly performing such a brute-force search requires a large amount of computation and is thus infeasible. In the same spirit as LogitBoost [5, 11] for multi-class prediction, we consider the second order Taylor expansion of the negative log-likelihood $l(y, x, \phi)$:

$$l(y, x, \phi + \delta) \simeq l(y, x, \phi) + \delta^T \mathbf{G}(y, x) + \frac{1}{2} \delta^T \mathbf{H}(y, x) \delta.$$
(4)

The gradient **G** and Hessian **H** in Eq.(4) are given by the following equation, where p_i and p_{ij} are short hand notations for $p_i \triangleq P(\mu_i = 1|x), p_{ij} \triangleq P(\mu_i \mu_j = 1|x)$:

$$\mathbf{G}_i = \mu_i(y) - p_i, \quad \mathbf{H}_{ij} = p_{ij} - p_i p_j. \tag{5}$$

108 Note that Eq.(5) holds for all i, j pairs, including two special cases: (1) $\mathbf{H}_{ii} = p_i(1-p_i)$ when 109 i = j, and (2) $\mathbf{H}_{ij} = -p_i p_j$ when μ_i and μ_j are mutual events. Intuitively, \mathbf{H}_{ij} measures the 110 correlation between two events, and is nonzero due to the dependencies in the CRF model. These dense elements of the Hessian make direct optimization of Eq. (4) still very costly. An existing 111 112 approach to functional optimization for CRFs, presented in [1], resorts to first order approximation to the loss, and can only guarantee convergence when the step size is small. An alternative is to 113 iteratively update one δ_i for $i \in \{1, \dots, m\}$ at a time. This approach would require m inference 114 steps per iteration, and is simply not applicable when the constraint C exists. 115

Our Approach In this paper, we consider an upper bound of Eq. (4) instead. The intuition behind this approach, which will be formalized in following sections, is as follows: each variable in the CRF depends weakly on variables that are "far" from it. This motivates the use of a diagonal upper bound of the Hessian to construct loss functions, given by the following Lemma.

Lemma 2.1. Let \mathcal{U} be a index set of potential functions we want to update, and let γ be a function that satisfies the following inequality

$$\gamma_i(y, x) \mathbf{H}_{ii}(y, x) \ge \sum_{j \in \mathcal{U}} |\mathbf{H}_{ij}(y, x)| \tag{6}$$

Then for $\delta \in \{ [\delta_1, \delta_2, \dots, \delta_m] \mid \delta_i = 0 \text{ for } i \notin \mathcal{U} \}$, the following inequality holds,

$$l(y, x, \phi + \delta) \le l(y, x, \phi) + \sum_{i \in \mathcal{U}} [\delta_i \mathbf{G}_i(y, x) + \frac{1}{2} \gamma_i(y, x) \mathbf{H}_{ii} \delta_i^2(y, x)] + o(\delta^2).$$
(7)

The detailed proof of the lemma is given in supplementary material. For a given γ that satisfies the condition, we can iteratively optimize $\tilde{L}(\phi, \delta)$, which is an upper bound of $L(\phi)$, defined by

$$\tilde{L}(\phi,\delta) = L(\phi) + \sum_{i \in \mathcal{U}} \left[\left(\sum_{x,y \in \mathcal{D}} \mathbf{G}_i(y,x) \right) \delta_i(y,x) + \frac{1}{2} \left(\sum_{x,y \in \mathcal{D}} \gamma_i(y,x) \mathbf{H}_{ii} \right) \delta_i^2(y,x) + \Omega(\phi_i + \delta_i) - \Omega(\phi_i) \right].$$
(8)

135 $\hat{L}(\phi, \delta)$ is composed of $|\mathcal{U}|$ independent loss functions with a regularization term, and can be used 136 to guide the common function search (such as regression tree learning). Iteratively optimizing \tilde{L} will 137 result in a gradient boosting algorithm that ensures the convergence of $L(\phi)$ (Proof in Section 4). 138 Furthermore, the form of \tilde{L} allows the search of δ_i for $i \in \mathcal{U}$ to be done *in parallel* for each equivalent 139 class defined by C, which gives us further computational benefits. In the next two sections, we will 140 discuss how we can efficiently estimate γ when \mathcal{U} is the index set of all node potentials, and when it 141 is the index set of all edge potentials, using a mixing rate of Markov chain.

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3 Upper Bound Derivation using a Markov Chain Mixing Rate

145 In this section, we will discuss how we can estimate γ when \mathcal{U} is the index set of all node potentials, and the index set of all edge potentials. Conceptually, the choice of γ should be related to the inter-146 dependency of variables in the current model. When the variables in the model are independent from 147 each other, γ should be small, and when the variables in the model have strong dependencies, γ 148 should become larger. We want to quantitatively measure the dependencies in the CRF. Specifically, 149 we will connect the dependency level to the mixing rate of a Markov chain defined by the conditional 150 distribution of outputs on input P(y|x). To begin with, we re-express the right side of Eq. (6) using 151 total variation distance, defined by $||P - Q||_{tv} = \frac{1}{2} \sum_{x} |P(x) - Q(x)|.$ 152

Lemma 3.1. Let \mathcal{U} correspond to the set of all node potentials $\mathcal{U} = \{j | \phi_j \in N\}$, assuming index *i* corresponds to the event $y_t = k$ (i.e. $\mu_i = \mathbf{1}(y_t = k)$), then

$$\sum_{j \in \mathcal{U}} |\mathbf{H}_{ij}| = 2p_i \sum_{s} ||P(y_s|x, \mu_i = 1) - P(y_s|x)||_{tv}.$$
(9)

Lemma 3.2. Let \mathcal{U} correspond to the set of all edge potentials $\mathcal{U} = \{j | \phi_j \in \mathcal{E}\}$, then

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$$\sum_{j \in \mathcal{U}} |\mathbf{H}_{ij}| = 2p_i \sum_{(s,v) \in \mathcal{E}} ||P(y_s, y_v|x, \mu_i = 1) - P(y_s, y_v|x)||_{tv}.$$
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Note that we abuse the notation slightly here, by using \mathcal{E} to indicate the index set of edges in CRF.

The proof is a re-arrangement of terms, and is provided in the supplementary material. Intuitively, the total variation terms in Lemma 3.1 and 3.2 measure how dependent y_s is on the event $y_t = k$. When y_s is only weakly dependent on y_t , the distance will be small. The complexity of calculating Eq. (9) for all i is quadratic in the number of nodes, which is too expensive to be calculated directly for most applications. We need an algorithm that scales linearly with the number of nodes.

Total variation distance allows us to approach the problem in terms of dependencies between vari-ables. Intuitively, we expect the dependencies between y_s and y_t to become smaller as we change s to get away from t. We formally state this in the following theorem:

Theorem 3.1. Mixing rate bound for Markov chain in CRF

Assume y_t , y_s and y_v form a Markov chain $y_t \to y_s \to y_v$, conditioned on x, i.e. $P(y_v|y_s, y_t, x) =$ $P(y_v|y_s, x)$ holds. Define $d(s, t, k) \triangleq ||P(y_s|x, y_t = k) - P(y_s|x)||_{tv}$. Then, the total variation d(v, t, k) can be bounded by

$$d(v,t,k) \le [1 - \sum_{i} \min_{i} P(y_v = j | y_s = i, x)] d(s,t,k) \triangleq \alpha_{s,v} d(s,t,k).$$
(10)

Proof. Define notation: $M_{ij} \triangleq P(y_v = j | y_s = i, x), Q_j \triangleq \min_i M_{ij}$, then

$$2d(v,t,k) = \sum_{j} |P(y_v = j|y_t = k, x) - P(y_v = j|x)|$$

$$= \sum_{j} |\sum_{i} M_{ij}P(y_s = i|y_t = k, x) - \sum_{i} M_{ij}P(y_s = i|x)|$$

$$= \sum_{j} |\sum_{i} (M_{ij} - Q_j)[P(y_s = i|y_t = k, x) - P(y_s = i|x)]|$$

$$\leq \sum_{j} \sum_{i} (M_{ij} - Q_j)|P(y_s = i|y_t = k, x) - P(y_s = i|x)|$$

$$= \sum_{i} (1 - \sum_{j} Q_j)|P(y_s = i|y_t = k, x) - P(y_s = i|x)| = 2\alpha_{s,v}d(s, t, k) \square$$

The derivation of Theorem 3.1 is inspired, in spirit, by the mixing rate bounds of time homogeneous Markov Chains $[10]^1$. Intuitively, Theorem 3.1 shows the dependency decays exponentially as s moves away from t. The following corollary holds as the direct consequence of the theorem.

Corollary 3.1. Let $q = [q(1), q(2), \dots, q(n)]$ be the path sequence in \mathcal{E} from t to s (i.e. q(1) =t, q(n) = s) then we can bound d(t, s, k) using d(t, t, k) times the decay ratio α along the path,

$$d(s,t,k) \le \prod_{i=1}^{n-1} \alpha_{q(i),q(i+1)} d(t,t,k).$$
(11)

In the case when \mathcal{E} is a chain, Corollary 3.1 simplifies to $d(s,t,k) \leq \prod_{h=t}^{s-1} \alpha_{h,h+1} d(t,t,k)$ when s > t, and $d(s, t, k) \leq \prod_{h=s+1}^{t} \alpha_{h,h-1} d(t, t, k)$ when s < t. An important property of Theorem 3.1 is that the position specific mixing rate $\alpha_{s,v}$ can be computed efficiently (complexity analysis in Sec 4). We still need to calculate d(t, t, k), which is given by the following Lemma

Lemma 3.3. Let M correspond to the index set of μ_i such that μ_i, μ_j are mutual to each other (i.e. $\mu_i \mu_j = 0$ for $i \neq j, i, j \in M$), and $\sum_{j \in M} P(\mu_i = j | x) = 1$. Then the following identity holds

$$\frac{1}{2}\sum_{j\in M} |P(\mu_j = 1|\mu_i = 1, x) - P(\mu_j = 1|x)| = 1 - P(\mu_i = 1|x)$$
(12)

The proof is given in supplementary material. From Lemma 3.3, it follows that $d(t, t, k) = 1 - p_i$. We will make use of Lemma 3.3 and Corollary 3.1 to efficiently estimate γ in next section.

Gradient Boosting for CRF

In this section, we will present our gradient boosting algorithm. We will give estimation of γ for \mathcal{U} to be the index set of all node potentials and edge potentials, given by the following two theorems.

¹Note that our proof is actually for a time inhomogeneous Markov Chain.

Algorithm 1 Gradient Boosting for CRF
repeat
for $\mathcal{U} \in \{N, \mathcal{E}\}$ do
for $y, x \in \mathcal{D}$ in parallel do
{inference of p_i, γ_i are done using dynamic programming}
Infer $\mathbf{G}_i(y, x) \leftarrow \mu_i(y) - p_i, \mathbf{H}_{ii}(y, x) \leftarrow p_i(1 - p_i)$ for each $i \in \mathcal{U}$
Infer $\gamma_i(y, x)$ using Theorem 4.1 and 4.2 for each $i \in \mathcal{U}$
end for
for $[c] \subset \mathcal{U}$ in parallel do
$\{We use [c] \text{ to enumerate over set of equivalent index defined by } C \text{ in } U\}$
$\delta_c \leftarrow \operatorname{argmin}_{\delta_c \neq \tau} \Omega(\phi_i + \delta) + \sum \sum_{i=\tau} \left[\mathbf{G}_i(y, x) \delta(y, x) + \gamma_i(y, x) \mathbf{H}_{ii}(y, x) \delta^2(y, x) \right]$
$ \sum_{i \in [c]} \sum_{y,x \in \mathcal{D}} \left[\sum_{i \in [c]} \sum_{y,x \in \mathcal{D}} \left[\sum_{i \in [c]} \sum_{y,x \in \mathcal{D}} \sum_{y,x \in D$
$\phi_c \leftarrow \phi_c + \epsilon \delta_c$
end for
end for
until convergence

Theorem 4.1. Let \mathcal{U} be the index set of all node potentials, assume $\mu_i = \mathbf{1}(y_t = k)$, and define \mathcal{Q}_t to be the set of all paths that start from t, then

$$\gamma_i(y,x) = 2(1 + \sum_{q \in \mathcal{Q}_t} \sum_{s=2}^{len(q)} \prod_{i=1}^{s-1} \alpha_{q(i),q(i+1)})$$
(13)

satisfies Eq.(6). We use len(q) as the length of the path, α is defined in Theorem 3.1. **Theorem 4.2.** Let \mathcal{U} be the index the set of all edge potentials, assume $\mu_i = \mathbf{1}(y_t = k_1, y_{t+1} = k_2)$, and define $\mathcal{Q}_{t,t+1}$ to be the set of all path that start from t and t + 1 and do not cross (t, t+1), then

$$\gamma_i(y,x) = 2\left(1 + \sum_{q \in \mathcal{Q}_{t,t+1}} \left(1 + \sum_{s=2}^{len(q)} \prod_{i=1}^{s-1} \alpha_{q(i),q(i+1)}\right)\right)$$
(14)

satisfies Eq.(6), with the same definition of len(q) and α as in Theorem 4.1.

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Both theorems can be proved by using Corollary 3.1 and Lemma 3.3 to bound the total variation distance. We give the detailed proof in supplementary material. Based on Theorem 4.1 and 4.2, we can get an efficient gradient boosting algorithm for CRF (GBCRF), which is presented in Algorithm 1. Here ϵ is a shrinkage term used to control overfitting. Our algorithm adaptively estimates γ via the mixing rate calculation at each iteration. At the beginning of training, where each node variable is independent from each other, we will have a γ that is close to 2 (and thus the updates are aggressive). γ increases as the variables become dependent on each other (inducing more conservative updates).

The calculation of γ can be performed using dynamic programming. To explain the idea more clearly, let us consider the case when \mathcal{E} is a chain. In this case, Eq. (13) specializes into a calculation of $\beta_t^+ \triangleq \sum_{s=t}^n \prod_{i=t}^{s-1} \alpha_{i,i+1}$ and $\beta_t^- \triangleq \sum_{s=1}^t \prod_{i=s+1}^t \alpha_{i,i-1}$, and both can be calculated efficiently using the following recursive formula

$$\beta_t^+ = \alpha_{t,t+1}(1+\beta_{t+1}^+), \ \beta_t^- = \alpha_{t,t-1}(1+\beta_{t-1}^-).$$
(15)

Similarly, we can use dynamic programming for any tree-shaped \mathcal{E} (using up-down recursion). A direct consequence of Theorem 4.1 is that we can bound the loss using estimation by number of nodes in CRF. Though this bound is usually worse than the bound using mixing rate.

Corollary 4.1. When \mathcal{U} is the index set of node potentials, $\gamma_i = 2n$ satisfies Eq. (6), where n is number of nodes in CRF.

Relation to LogitBoost Our algorithm can be viewed as a generalization of multi-class classification
 using LogitBoost [5]. When the variables in each position are independent (no edge potentials), the
 estimation of γ equals 2, and our algorithm becomes identical to LogitBoost. When the variables
 are dependent on each other, which is common in structured prediction, our model estimates the
 dependency level via the Markov Chain mixing rate to guide the boosting objective in each iteration.

270 **Time Complexity** The time complexity for the gradient boosting statistics collection in Algorithm 1 271 is $O(|\mathcal{D}|nK^2)$, where K is the number of states in each node and n is the average number of 272 nodes (e.g. length of sequence) in each instance. This is due to the fact that estimation of γ can 273 be done in $O(|\mathcal{D}|nK^2)$ complexity, using a dynamic programming algorithm. This complexity is 274 same as the complexity for traditional training methods for linear CRF. The time complexity of entire algorithm is $O(|\mathcal{D}|nK^2 + g(|\mathcal{D}|, n))$, where $g(|\mathcal{D}|, n)$ is cost of function learning given the 275 statistics. For learning trees, the complexity of function learning is usually $O(|\mathcal{D}|n\log(|\mathcal{D}|n))$. 276 Thus our approach extends CRFs to non-linearity with only an additional log factor. 277

Convergence Analysis In this section, we analyze the convergence of our algorithm. The advantage of our method is that it makes use of second order information, and guarantees convergence.

Theorem 4.3. $L(\phi)$ converges with the procedure described by Algorithm 1 for $\epsilon \leq 1$.

282 Proof. During each iteration, assume δ^* is the function that optimizes $\tilde{L}(\phi, \delta)$ defined in Eq. (8),

$$L(\phi + \epsilon\delta^*) \le \tilde{L}(\phi, \epsilon\delta^*) \le \tilde{L}(\phi, \mathbf{0}) = L(\phi)$$
(16)

Thus the loss function L decreases after each boosting step, and the algorithm converges to a minima (possibly local minima when \mathcal{F} is nonlinear) of L.

5 Related Work

Conditional random fields [9] are among the most successful solutions to structured prediction prob-290 lems. Variants of conditional random fields have been proposed and widely applied for structured 291 prediction in domains such as natural language processing [9, 16], computer vision [7, 15] and bio-292 informatics [21]. Most popular instantiations assume linear potential functions and improve the 293 performance by carefully engineering features. Our work focuses on learning probabilistic models for tree-shaped CRFs with nonlinear potential functions. When there are loops in the CRF and infer-295 ence is intractable, relaxation of the objective can be done to use approximate inference and learn-296 ing [6, 13, 3]. A similar dependency based term is also used in the approximate inference [13, 3], 297 but is usually set be a constant value across all instances and training iterations. As a future work, 298 it would be interesting to explore whether our adaptive Markov Chain mixing rate bound can be 299 applied to this more general setting.

300 Gradient boosting [4], which performs additive optimization in the functional space, has been suc-301 cessfully applied to classification problems that assume independent outputs conditioned on the 302 input [5]. Most existing attempts to "boost" CRF models optimize approximate objectives [20, 12]. 303 TreeCRF algorithm [1] is similar to our approach in that it directly optimizes the log-likelihood func-304 tion defined using non-linear potential functions, however they only take first order information into 305 account during optimization, requiring a decreasing step size. On the other hand, our method makes 306 use of second order information, and guarantees convergence with fixed step size. Our method can 307 also be viewed as a generalization of LogitBoost [5] for CRF. It is worth noting that the recent improvements of LogitBoost, which make use of adaptive base function [11, 17], can also potentially 308 be combined with our method to make further improvements. 309

6 Experiments

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313 In this section, we evaluate our method on named entity recognition, hand written character recogni-314 tion, and protein secondary structure prediction. We compare the following methods: (1) GBCRF is the proposed method in this paper. We set \mathcal{F}_n to be a set of regression trees, and \mathcal{F}_e to be linear func-315 tions of basic transition features between states; (2) LogitBoost is a gradient boosting method for 316 multi-class classification [5] that does not support the dependencies between outputs; (3) TreeCRF 317 is a gradient boosting method that only takes *first-order* information [1]. We use the same family of 318 edge and node potentials as GBCRF; (4) Linear CRF is the standard CRF model with linear edge 319 and node potentials [9]. For all the methods, the training parameters are selected using a validation 320 set or cross validation, depending on the specific setup of each dataset.

Named Entity Recognition We first test our methods on the natural language task of named entity
 recognition (NER) using the CoNLL-2003 shared task benchmark dataset [19]. The dataset contains around 20K sentences, and defines a standard split into 14K as training set, 3.3K as validation

324 325	Table 1: F1 Measure of Name Entity Recognition on CoNLL-2003 Dataset. We denote validation set, and subscript <i>test</i> to denote test set.
326	Word Embedding only Word Features + Embed

	Word Embedding only		Word Features + Embedding		
Method	$F1_{val}$	$F1_{test}$	$F1_{val}$	$F1_{test}$	
Linear CRF	0.8452	0.7943	0.8952	0.8475	
LogitBoost	0.8532	0.7887	0.8717	0.8197	
TreeCRF	0.8630	0.8060	0.8846	0.8399	
GBCRF	0.8801	0.8269	0.9015	0.8635	

Table 2: Cross Validation Error on

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Handwritten Character Recognition Dataset.

Table 3: Predictive Q8 Accuracy on Protein Secondary Structure Dataset.

use subscript val to

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Method	Error	Method	Accuracy
Linear CRF	0.1292 ± 0.0080	Linear CRF	0.614
LogitBoost	0.0967 ± 0.0049	LogitBoost	0.710
TreeCRF	0.0699 ± 0.0040	TreeCRF	0.718
GBCRF	0.0464 ± 0.0027	GBCRF	0.722
NeuroCRF (Do et al.[2])	0.0444	SC-GSN (Zhou et al.[22])	0.711
		SC-GSN with "kick-start" ([22])	0.721

set (also called development set), and 3.5K sentences as test set. Traditional approaches for NER
 involve a lot of time-consuming feature engineering that requires domain expertise, and build a
 Linear CRF over these features. Instead, in our experiment, we explore whether it is possible to
 perform minimal feature engineering, and use a representation learned from data for prediction.

346 Specifically, we take the word embedding vectors from Mikolov et.al [14], which is learned from 347 Google news corpus, and train the models on this representation. In this setting, each word is 348 represented by a 300 dimensional vector that captures the "semantics" of the word. For each position 349 in the sentence, we take the embedding vector of the previous, current, and next word as input to node 350 potential function. We call this setting "word embedding only". We further perform minimal feature engineering to *only* generate the unigram features (word, postag and case pattern of current word). 351 We use these basic features to train a weak linear model, then use additive training to boost the base 352 model using the word embedding representation. We call this setting "word feature+embedding". 353

The results of token-wise F1 evaluation for these models are shown in Table 1. From the result, we see that GBCRF works better than Linear CRF in both settings. The gap between LogitBoost and GBCRF indicates the importance of introducing edge potentials to this problem. We also find that taking second order information into account helps us obtain a more accurate model.

Handwriting Character Recognition We also evaluate our method on a handwriting recognition dataset². The dataset consists of 6877 words and corresponds to about 52 thousand handwritten characters [8, 18], each represented by a binary pixel vector of 128 dimensions, and belonging to one of 26 alphabets. The dataset is randomly split into 10 folds for cross validation. We train the models on 9 folds, test on 1 fold, and use the cross validation error to compare the methods.

The experiment results are shown in Table 2. Both our method and TreeCRF outperforms CRF with linear potential functions, which indicates the effectiveness of introducing a non-linear potential function into the CRF on this dataset. The gap between LogitBoost and models that consider dependencies indicate the importance of incorporating structure information of the outputs into the model. Our results are also comparable to NeuroCRF [2], which uses a deep neural network as a potential function whose weights are initialized by Restricted Boltzmann Machines.

369 Protein Secondary Structure Prediction We also conduct an experiment on protein secondary 370 structure prediction. The task is to predict 8-state secondary structure labels for a given amino-acid 371 sequence of a protein. We use the protein secondary structure data-set recently introduced by Zhou 372 et al.[22], which is the largest publicly available protein secondary structure prediction dataset. The 373 dataset contains 6128 proteins, with average sequence length around 208. We use exactly the same 374 features and data split step as [22]. The resulting data set contains 5600 sequences as training set, 256 sequences as validation set and 272 sequences as test set. Each position of the protein sequence 375 contains 46 dimension features (22 for PSSM, 22 for sequence and 2 for terminals) for prediction. 376

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²http://www.seas.upenn.edu/ taskar/ocr/

Mixing Rate based Estimation Brute Force Estimation

Average Length based Estimatio 381 382 loalikel 383 Vegativ 384 385 386 387 200 Iteration 200 250 300 100 150 250 300 350 388 (a) Evolution of Negative-loglikelihood (b) Evolution of average γ estimation 389 390 Figure 1: Convergence of GBCRF on hand written character dataset. (a) Convergence comparison 391 between GBCRF and TreeCRF, with shrinkage rate of both algorithm set to 1. GBCRF converges 392 faster than TreeCRF; (b) Evolution of different γ estimations on the CRF model in each round based on 500 sequences. Mixing rate based estimation has the same trend as brute force estimation, and 393 provides a tighter estimation than the length based estimation. 394

- GBCRF TreeCR

395 To train the models, we take the concatenation of feature vectors within 3 positions of the target 396 position as input to the node potential, resulting in 322 input features in each position. 397

The performance of the model is measured by the accuracy of predictions on the test set (denoted as 398 Q8). We train the models with parameters discovered using the validation set, and report the results 399 in Table 3. From the table, we find that using trees as potential functions leads to better performance 400 than restricting the model to using linear functions. Our results are comparable to the state-of-401 art result in this dataset, produced by Zhou et al. [22] (SC-GSN-3layer). The result is generated 402 by a deep convolutional generative stochastic network model to perform secondary structure label 403 prediction, optimized with a "kick-start" initialization scheme.

404 **Convergence of GBCRF** We further analyze the convergence of GBCRF on the handwritten char-405 acter dataset. We plot the convergence of negative log-likelihood function of GBCRF and TreeCRF 406 in Fig. 1(a). We find that GBCRF converges faster than TreeCRF, demonstrating that taking second 407 order information into account not only gives theoretical guarantee of convergence, but also helps 408 the method to converge faster in practice. 409

We also investigate the tightness of γ estimation. Figure 1(b) gives the average of different γ esti-410 mations on models trained by GBCRF in each round. Mixing rate based estimation is the method 411 proposed in this paper. We perform Brute Force estimation to compute γ exactly using Eq. (9); 412 the complexity of this estimation is quadratic in the number of nodes and outputs in the CRF, and 413 hence cannot be used for most real-world sequences. Average Length based estimation is a naive 414 estimation using 2 times number of nodes in CRF, and provides an valid estimation of γ since it 415 upper bounds Eq. (13), as we show in Corollary 4.1. We restrict this evaluation to the shortest 500 416 sequences, due to the computation cost of brute force estimation. From the figure, we find that mix-417 ing rate based estimation exhibits the same trend as the brute force estimation, and is at most 2.3 times higher than the brute force estimation. Further, the mixing rate based bound is consistently 418 lower than the fixed bound computed by the length based estimation. These results indicate that our 419 mixing rate based estimation captures the changes in the dependencies in the model during training 420 correctly. Hence our proposed mixing rate based approach is indeed useful to estimate γ efficiently. 421

7 Conclusion

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425 In this paper, we present novel gradient boosting algorithm for CRF. It is non-trival to design an effective gradient boosting for CRF, mainly due to the dense Hessian matrices introduced by variable 426 interdependency. To solve the problem, we make use of a Markov Chain mixing rate to derive an 427 efficiently computable adaptive upper bound of the loss function, and construct a gradient boosting 428 algorithm that iteratively optimizes the bound. The resulting algorithm can be viewed as a gener-429 alization of LogitBoost to CRF, thus introducing non-linearity in CRFs at only a log factor cost. 430 Experimental results demonstrate that our method is both efficient and effective. As future work, it 431 is interesting to explore whether we can generalize the result to loopy models.

432 References 433

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Supplementary Material

A Proof for Lemma 2.1

Proof. The following inequality holds for γ that satisfies the condition

$$\sum_{i \in \mathcal{U}} \gamma_i \mathbf{H}_{ii} \delta_i^2 \ge \sum_{i \in \mathcal{U}} \sum_{j \in \mathcal{U}} |\mathbf{H}_{ij}| \delta_i^2 = \frac{1}{2} \sum_{i \in \mathcal{U}} \sum_{j \in \mathcal{U}} |\mathbf{H}_{ij}| (\delta_i^2 + \delta_j^2) \ge \sum_{i \in \mathcal{U}} \sum_{j \in \mathcal{U}} |\mathbf{H}_{ij}| \delta_i \delta_j$$

Applying it to Talyor expansion in Eq (4), we have

$$l(y, x, \phi + \delta) = l(y, x, \phi) + \sum_{i \in \mathcal{U}} \delta_i \mathbf{G}_i(y, x) + \frac{1}{2} \sum_{i \in \mathcal{U}} \sum_{j \in \mathcal{U}} |\mathbf{H}_{ij}| \delta_i \delta_j + o(\delta^2)$$

$$\leq l(y, x, \phi) + \sum_{i \in \mathcal{U}} \delta_i \mathbf{G}_i(y, x) + \frac{1}{2} \sum_{i \in \mathcal{U}} \gamma_i \mathbf{H}_{ii} \delta_i^2 + o(\delta^2).$$

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B Proof for Lemma 3.3

Proof. Taking the fact that μ_i and μ_j are mutual for $j \neq i$, we have

$$\begin{split} &\sum_{j \in M} |P(\mu_j = 1 | \mu_i = 1, x) - P(\mu_j = 1 | x)| \\ &= |P(\mu_i = 1 | \mu_i = 1, x) - P(\mu_i = 1 | x)| + \sum_{j \neq i} |P(\mu_j = 1 | \mu_i = 1, x) - P(\mu_j = 1 | x)| \\ &= |1 - P(\mu_i = 1 | x)| + \sum_{j \neq i} |0 - P(\mu_j = 1 | x)| \\ &= (1 - P(\mu_i = 1 | x)) + \sum_{j \neq i} P(\mu_j = 1 | x) \\ &= 2(1 - P(\mu_i = 1 | x)) \end{split}$$

C Proof for Lemma 3.1 and 3.2

Proof. The proof is exactly the same for both node and potential case, we present the proof for \mathcal{U} to be all node potentials here. Recall the definition of \mathbf{H} : $\mathbf{H}_{ij} = p_{ij}$. Note that p_i and p_{ij} are short hand notations for $p_i \triangleq P(\mu_i = 1|x)$, $p_{ij} \triangleq P(\mu_i \mu_j = 1|x)$, we have

$$\frac{1}{2p_i} \sum_{j \in \mathcal{U}} |\mathbf{H}_{ij}| = \sum_j |p_{ij}/p_i - p_j|$$

= $\sum_{j \in \mathcal{U}} |P(\mu_j = 1 | \mu_i = 1, x) - P(\mu_j = 1 | x)|$
= $\sum_{s,k'} |P(y_s = k' | y_t = k, x) - P(y_s = k' | x)|$
= $\sum_{s,k'} ||P(\mu_s | x, \mu_s = k) - P(\mu_s | x)||$

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$$= \sum_{s} \|P(y_s|x, y_t = k) - P(y_s|x)\|_{tv}$$

⁵⁴⁰ D Proof for Theorem 4.1 and 4.2

Proof. **Proof for Theorem 4.1** Basically, we want to bound the total variation distance given by Eq. (9) in Lemma 3.1,

$$2p_i \sum_{s} \|P(y_s|x, y_s = k) - P(y_s|x)\|_{tv} = 2p_i \sum_{s} d(s, t, k)$$

$$= 2p_i [d(t, t, k) + \sum_{q \in \mathcal{Q}_t} \sum_{s=2}^{len(q)} d(q(s), t, k)]$$

$$\leq 2p_i [d(t, t, k) + \sum_{q \in \mathcal{Q}_t} \sum_{s=2}^{len(q)} d(t, t, k) \prod_{i=1}^{s-1} \alpha_{q(i), q(i+1)}]$$

$$= 2p_i (1 - p_i) [1 + \sum_{q \in \mathcal{Q}_t} \sum_{s=2}^{len(q)} \prod_{i=1}^{s-1} \alpha_{q(i), q(i+1)}]$$

Here the inequality is given by Corollary 3.1 $(d(q(s), t, k) \leq d(t, t, k) \prod_{i=1}^{s-1} \alpha_{q(i),q(i+1)})$, and last equality is given by Lemma 3.3 $(d(t, t, k) = 1 - p_i)$. Recall that $\mathbf{H}_{ii} = p_i(1 - p_i)$, we have proved Theorem 4.1.

Proof. **Proof for Theorem 4.2** In this proof, we will reduce the total variation distance between joint distribution of edge states into total variation distance of marginal distribution over nodes, as in Theorem 4.1. Assume in edge pairs are $(y_t, y_{t+1}), (y_s, y_{s+1})$, and y_s is closer to y_{t+1} (without loss of generality), then

$$P(y_s, y_{s+1}|y_t, y_{t+1}, x) = P(y_{s+1}|y_s, x)P(y_s|y_{t+1}, x)$$

We can convert total variation by

$$\|P(y_{s}, y_{s+1}|y_{t}, y_{t+1}, x) - P(y_{s}, y_{s+1}|x)\|_{tv} = \sum_{y_{s}, y_{s+1}} |P(y_{s}, y_{s+1}|y_{t}, y_{t+1}, x) - P(y_{s}, y_{s+1}|x)|$$

$$= \sum_{y_{s}, y_{s+1}} P(y_{s+1}|y_{s}, x)|P(y_{s}|y_{t+1}, x) - P(y_{s}|x)|$$

$$= \sum_{y_{s}} |P(y_{s}|y_{t+1}, x) - P(y_{s}|x)|$$

$$= \|P(y_{s}|y_{t+1}, x) - P(y_{s}|x)\|_{tv}$$
(17)

576 Now the case become same as node potential, we can make use of Corollary 3.1 bound the total 577 variation. Specifically, let $q \in Q_{t,t+1}$ (i.e. $q(1) \in \{t, t+1\}, q(i) \notin \{t, t+1\}$ for i > 1) 578 len(q)

$$\sum_{i=1}^{len(q)} \|P(y_{q(i)}, y_{q(i+1)}|y_{t} = k_{t}, y_{t+1} = k_{t+1}, x) - P(y_{q(i)}, y_{q(i+1)}|x)\|_{tv}$$

$$= \sum_{i=1}^{len(q)} \|P(y_{q(i)}|y_{q(1)} = k_{q(1)}, x) - P(y_{q(i)}|x)\|_{tv}$$

$$\leq \|P(y_{q(1)}|y_{q(1)} = k_{q(1)}, x) - P(y_{q(1)}|x)\|_{tv} + \sum_{i=2}^{len(q)} \|P(y_{q(1)}|y_{q(1)} = k_{q(1)}, x) - P(y_{q(1)}|x)\|_{tv} \prod_{j=1}^{i-1} \alpha_{q(j),q(j+1)}$$

$$= [1 - P(y_{q(1)} = k_{q(1)}|x)](1 + \sum_{i=2}^{len(q)} \prod_{j=1}^{i-1} \alpha_{q(j),q(j+1)})$$

$$\leq [1 - P(y_{t} = k_{t}, y_{t+1} = k_{t+1}|x)](1 + \sum_{i=2}^{len(q)} \prod_{j=1}^{i-1} \alpha_{q(j),q(j+1)}).$$

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$$\leq [1 - P(y_t = k_t, y_{t+1} = k_{t+1}|x)](1 + \sum_{i=2}^{t \in n(q)} \prod_{j=1}^{i-1} \alpha_{q(j),q(j+1)})$$

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Here the first inequality is due to Corollary 3.1. Summing the results over all $q \in Q_{t,t+1}$ will give us Eq. (14).