Diverse M-Best Solutions in Markov Random Fields

Dhruv Batra¹, Payman Yadollahpour¹, Abner Guzman-Rivera², and Gregory Shakhnarovich¹

 $^{1}\ \, {\rm TTI\text{-}Chicago} \\ ^{2}\ \, {\rm UIUC}$

Abstract. Much effort has been directed at algorithms for obtaining the highest probability (MAP) configuration in probabilistic (random field) models. In many situations, one could benefit from additional high-probability solutions. Current methods for computing the M most probable configurations produce solutions that tend to be very similar to the MAP solution and each other. This is often an undesirable property. In this paper we propose an algorithm for the Diverse M-Best problem, which involves finding a diverse set of highly probable solutions under a discrete probabilistic model. Given a dissimilarity function measuring closeness of two solutions, our formulation involves maximizing a linear combination of the probability and dissimilarity to previous solutions. Our formulation generalizes the M-Best MAP problem and we show that for certain families of dissimilarity functions we can guarantee that these solutions can be found as easily as the MAP solution.

1 Introduction

The introduction of sophisticated discrete optimization tools for inference in Markov Random Fields (MRFs) over the last two decades has allowed optimal or provably approximate solutions to certain vision problems previously deemed intractable. For instance, using max-flow/min-cut methods [7,13], one can find the globally optimal solution for a (submodular) foreground-background segmentation problem on a 2 Megapixel image within seconds, effectively searching $2^{2,000,000}$ possible segmentations.

However, optimization error is only one component of generalization error of a learning algorithm [4]. Even when exact inference in MRFs is efficient, the maximum a posteriori (MAP) solution could be far from the ground truth. The source of this discrepancy may be approximation error, due to the limitations of the model class (e.g. pairwise binary submodular MRFs), or estimation error i.e. error made because parameters are learnt from a finite training set.

Indeed, recent empirical studies [20, 32] have repeatedly found that MAP solutions of existing models are of much poorer quality than the ground-truth on vision problems like segmentation, stereo, optical flow, denoising, etc. Equivalently, the ground-truth has lower probability than the MAP solution under existing models. This discrepancy has been the driving force behind the search for more accurate (higher-order) models; [16, 23, 27] are just a few examples.

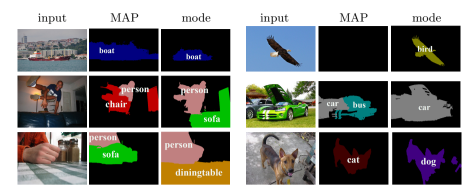


Fig. 1. Examples of category-level segmentation results on test images from VOC 2010. For each image, "mode" above is the best of 10 solutions obtained with *Div*MBEST.



Fig. 2. (Left) Interactive segmentation setup; (Right) Pose tracking. "Mode" above refers to the solution obtained with *DivMBEST*.

In addition to this quest for better models, we believe that another way to mitigate this problem is to look beyond obtaining a single MAP solution – to look for a diverse set of highly probable solutions instead from our existing models. Even if the MAP solution alone is of poor quality, a diverse set of highly probably hypotheses might still enable accurate predictions. Note that in contrast to the M-Best MAP problem [9, 22, 38] that involves finding the top M most probable solutions under a probabilistic model, our approach emphasizes diversity. We seek to produce highly probable solutions that are qualitatively different from the MAP and from each other. This is an important distinction because the literal definition of M-best MAP is not expected to work well in practice. In a large state-space problem $(e.g.\ 2^{2,000,000})$ any reasonable setting of M (10-50) would return solutions nearly identical to the MAP. Ideally, we would like to find the modes of the distribution learnt by our probabilistic model. Figs. 1, 2 show examples of these diverse solutions extracted for different tasks.

Overview. In this paper, we introduce the *Diverse M-Best* problem, which involves finding a diverse set of highly probable solutions under a discrete probabilistic model. Our formulation assumes access to a dissimilarity function $\Delta(\cdot, \cdot)$ measuring the difference between two solutions. We present an integer programming formulation for the Diverse M-Best problem, study its Lagrangian

relaxation. We show that this relaxation has an interesting interpretation as the Δ -augmented energy minimization problem, which minimizes a linear combination of the energy and similarity to previous solutions, thereby producing solutions with low energy and high diversity. For simplicity we sometimes will refer to these solutions as modes, although they are not guaranteed to be actual modes of the Gibbs distribution.

Contributions. Our principal contribution is the *first* principled optimization formulation for extracting diverse high-probability solutions in MRFs. It includes the familiar M-Best MAP problem as a special case. Perhaps most crucial for a practitioner, we show that for certain families of Δ -functions (that we discuss in Section 5), the Δ -augmented energy minimization problem is as easily solvable as the original MAP problem. Thus if exact or provably approximate algorithms exist for finding the MAP solution in a model, (e.g. graph-cuts [7, 13]), those same algorithms are applicable for finding Diverse M-Best solutions.

Applications. Our algorithm could be applicable whenever multiple hypotheses can be effectively used to infer a solution to the problem. In this paper, we show applications on three vision tasks: (i) an interactive application (object-cutout) where multiple diverse cutouts are presented to a user to minimize interaction time (ii) category-level image segmentation, where multiple highly-probable solutions could be ranked using a secondary mechanism [19] not amenable to inference, and finally (iii) human pose tracking in video, where multiple solutions per frame can be used for pose-tracking via a Viterbi-like decoding scheme that exploits temporal context [25]. In all three of these tasks we demonstrate that Diverse M-Best solutions can be used to significantly improve performance.

2 Related Work

M-Best MAP. Most directly relevant to our work is literature on the M-Best MAP problem. Lawler [18] proposed a general algorithm to compute the top M solutions for a large family of discrete optimization problems, and the ideas used in Lawler's algorithm form the basis of most algorithms for the M-Best MAP problem. The first family of algorithms for M-Best MAP [22, 30] were junctiontree based exact algorithms, thus feasible only for low-treewidth graphs. Dechter and colleagues [8,21] have recently provided dynamic-programming algorithms for M-Best MAP, but these are exponential in treewidth as well. Yanover and Weiss [38] proposed an algorithm that requires access only to max-marginals. Thus, for certain classes of MRFs that allow efficient exact computation of max-marginals, e.g. binary pairwise supermodular MRFs [12], M-Best solutions can be found for arbitrary treewidth graphs. Moreover, approximate M-Best solutions may be found by approximating the max-marginal computation, e.g. via max-product BP. More recently, Fromer and Globerson [9] provided a Linear Programming (LP) view of the M-Best MAP problem by studying the assignment-excluding marginal polytope. We show in Section 4 that M-best MAP is a special case of our formulation, with a particular kind of $\Delta(\cdot, \cdot)$.

Sampling Based Approaches. A common alternative is to use sampling [1,26, 35] to produce multiple solutions, which may then be refined or checked for diversity. Such approaches typically exhibit long wait times to transition from one mode to another. Moreover, in contrast to our work, there is no direct mechanism to require the multiple solutions (samples) obtained in this way to be diverse. An interesting approach by Papandreou and Yuille [24] shows that approximate samples may be drawn from the Gibbs distribution by injecting a certain noise into the parameters and solving for MAP on these perturbed parameters. Our approach can be seen as a deterministic counterpart to their stochastic perturbation – we always perturb the parameters in a fixed way, and the perturbation does not produce iid samples, rather Diverse M-Best solutions.

Diverse Solutions. The need for diverse solutions arises in a number of problems in computer vision and more broadly in machine learning. Yu and Joachims [39] proposed to learn a predictor that selects a topic-diverse subset of documents. Diversity is also a key goal in the context of non-maximal suppression for object detection [2, 3]. Recently, Park and Ramanan [25] applied the maxmarginal algorithm of [38] to decode multiple solutions from a deformable parts model, with an added constraint forcing at least one non-overlapping part. Their approach is also contained in our formulation as a special case (see Section 4).

3 Preliminaries: MAP Inference in MRFs

Notation. For any positive integer n, let [n] be shorthand for the set $\{1, 2, \ldots, n\}$. We consider a set of discrete random variables $\mathbf{x} = \{x_i \mid i \in [n]\}$, each taking value in a finite label set, $x_i \in X_i$. For a set $A \subseteq [n]$, we use x_A to denote the tuple $\{x_i \mid i \in A\}$, and X_A to be the cartesian product of the individual label spaces $\times_{i \in A} X_i$. For ease of notation, we use x_{ij} as a shorthand for $x_{\{i,j\}}$.

MAP. Let $G = (\mathcal{V}, \mathcal{E})$ be a graph defined over these variables, *i.e.* $\mathcal{V} = [n], \mathcal{E} \subseteq \binom{\mathcal{V}}{2}$, and let $\theta_A : X_A \to \mathbb{R}$, $(\forall A \in \mathcal{V} \cup \mathcal{E})$ be functions defining the energy at each node and edge for the labeling of variables in scope. The goal of MAP inference is to find the labeling \mathbf{x} of the variables that minimizes this real-valued energy function:

$$\min_{\mathbf{x} \in X_{\mathcal{V}}} \sum_{A \in \mathcal{V} \cup \mathcal{E}} \theta_A(x_A) = \min_{\mathbf{x} \in X_{\mathcal{V}}} \sum_{i \in \mathcal{V}} \theta_i(x_i) + \sum_{(i,j) \in \mathcal{E}} \theta_{ij}(x_i, x_j). \tag{1}$$

The techniques proposed in this paper are naturally applicable to higher-order MRFs. However, to simplify the exposition we only consider pairwise energies.

MAP Integer Program. MAP inference is typically set up as an integer programming problem over boolean variables. For each node and edge $A \in \mathcal{V} \cup \mathcal{E}$, let $\mu_A = \{\mu_A(s) \mid s \in X_A, \ \mu_A(s) \in \{0,1\}\}$, be a vector of indicator variables encoding all possible configurations of x_A . If $\mu_A(s)$ is set to 1, this implies that x_A takes label s. Moreover, let $\theta_A = \{\theta_A(s) \mid s \in X_A\}$ be a vector holding energies for all possible configurations of x_A , and $\mu = \{\mu_A \mid A \in \mathcal{V} \cup \mathcal{E}\}$ be a vector holding the entire configuration. Using this notation, the MAP inference integer program can be written as:

$$\min_{\boldsymbol{\mu}_{i}, \boldsymbol{\mu}_{ij}} \quad \sum_{i \in \mathcal{V}} \boldsymbol{\theta}_{i} \cdot \boldsymbol{\mu}_{i} + \sum_{(i,j) \in \mathcal{E}} \boldsymbol{\theta}_{ij} \cdot \boldsymbol{\mu}_{ij}$$
 (2a)

s.t.
$$\sum_{s \in X_i} \mu_i(s) = 1 \qquad \forall i \in \mathcal{V} \qquad (2b)$$

$$\sum_{s \in X_i} \mu_{ij}(s,t) = \mu_j(t), \quad \sum_{t \in X_j} \mu_{ij}(s,t) = \mu_i(s) \qquad \forall \{i,j\} \in \mathcal{E}$$
 (2c)

$$\mu_i(s), \mu_{ij}(s,t) \in \{0,1\}.$$
 (2d)

Here (2b) and (2c) enforce that exactly one label is assigned to a variable, and that assignments are consistent across edges. To be concise, we will use $\mathcal{L}(G)$ to denote the set of μ that satisfy these two constraints. Thus, the above problem

(2) can be written concisely as:
$$\min_{\boldsymbol{\mu} \in \mathcal{L}(G), \; \mu_A(s) \in \{0,1\}} \; \sum_{A \in \mathcal{V} \cup \mathcal{E}} \boldsymbol{\theta}_A \cdot \boldsymbol{\mu}_A.$$

Problem (2) is known to be NP-hard in general. A number of techniques solve a Linear Programming (LP) relaxation of this problem [36], which is given by relaxing the boolean constraints (2d) to the unit interval, *i.e.* $\mu_i(s)$, $\mu_{ij}(s,t) \geq 0$.

4 Diverse M-Best: Formulation

We now describe our proposed Diverse M-Best formulation. Recall that the goal is to produce a diverse set of low-energy solutions. We approach this problem with a greedy algorithm, where the next solution is defined as the lowest energy state with at least some minimum dissimilarity from the previously chosen solutions. To do so, we assume access to a dissimilarity function $\Delta(\mu^1, \mu^2)$ between solutions. Let μ^m denote the m^{th} -best mode. Thus μ^1 is the MAP, μ^2 is the second-best mode and so on. Let us first search for the second mode. We propose the following straightforward yet fairly general formulation:

$$\mu^{2} = \underset{\mu \in \mathcal{L}(G), \ \mu_{A}(s) \in \{0,1\}}{\operatorname{argmin}} \sum_{A \in \mathcal{V} \cup \mathcal{E}} \theta_{A} \cdot \mu_{A}$$
 (3a)

s.t.
$$\Delta(\boldsymbol{\mu}, \boldsymbol{\mu}^1) \ge k$$
. (3b)

We refer to the above formulation as $Div2Best(\Delta, k)$. Note that it is parametrized by the two design choices Δ and k (both of which we will discuss in detail). Intuitively, we can see that the above formulation searches for the lowest energy solution that is at least k-units away from the MAP solution under $\Delta(\cdot, \cdot)$. The extension from $Div2Best(\Delta, k)$ to $DivMBest(\Delta, k)$ is fairly simple: we search for the lowest energy solution at least k_m -units away from each of the previously found (M-1) solutions, i.e. $\Delta(\mu, \mu^m) \geq k_m$, $\forall m \in \{1, \ldots, M-1\}$, where $k = \{k_m \mid m \in [M-1]\}$ is the vector of different distance thresholds.

This formulation is general enough to contain existing ones as special cases:

Special Case: M-Best MAP is obtained when Δ is a 0-1 dissimilarity (*i.e.* $\Delta(\mu, \mu^1) = [\![\mu \neq \mu^1]\!]$, where $[\![\cdot]\!]$ is an indicator function), and k=1. Thus (3b) simply forces the next best solution to not be identical to MAP. Other choices of $\Delta(\cdot, \cdot)$ are also possible to express the M-Best MAP problem.

¹ Whenever we refer to MAP we mean exact or approximate MAP, as produced by the inference engine for the model at hand.

Special Case: N-Best Maximal Decoding of Park and Ramanan [25] corresponds to $\Delta(\mu, \mu^1) = \max_{i \in \mathcal{V}} \Delta_i(\mu_i, \mu_i^1)$ and k = 1. Intuitively, their approach defines local dissimilarity functions at each node, and forces at least one node label to be non-identical (under Δ_i) to the MAP label at that node.

5 Diverse M-Best: Lagrangian Relaxation

In general, Div MBEST (Δ, \mathbf{k}) is at least as hard to solve the MAP inference problem, which is NP-hard. Moreover, the dissimilarity constraints obfuscate some of the structure in the problem typically exploited by MAP inference algorithms. Thus we study the Lagrangian relaxation of Div MBEST (Δ, \mathbf{k}), formed by dualizing the dissimilarity constraints $\Delta(\mu, \mu^m) \geq \mathbf{k}$:

$$f(\lambda) = \min_{\boldsymbol{\mu} \in \mathcal{L}(G), \ \boldsymbol{\mu}_A(s) \in \{0,1\}} \sum_{A \in \mathcal{V} \cup \mathcal{E}} \boldsymbol{\theta}_A \cdot \boldsymbol{\mu}_A - \sum_{m=1}^{M-1} \lambda_m \left(\Delta(\boldsymbol{\mu}, \boldsymbol{\mu}^m) - k_m \right).$$
(4)

Here $\lambda = \{\lambda_m \mid m \in [M-1]\}$ is the set of Lagrange multipliers, that determine the weight of the penalty imposed for violating the constraints by a solution. Intuitively, we can see that the Lagrangian relaxation minimizes a linear combination of the energy of the MRF and similarity (negative dissimilarity) to the MAP solution, with the weighting given by the Lagrange multipliers. Formally, we can state the following theorem, proven in [5]:

Proposition 1. For all values of $\lambda \geq 0$, $f(\lambda)$ is a lower-bound on the value of the primal problem DivMBEST (Δ, \mathbf{k}) . Moreover, $f(\lambda)$ is a piece-wise linear function and concave in λ .

5.1 Supergradient Ascent on the Lagrangian Dual

The tightest lower-bound is obtained by setting up the Lagrangian dual problem: $\max_{\lambda \geq 0} f(\lambda)$. Since f is a non-smooth concave function, this can be achieved by the supergradient ascent algorithm, analogous to the subgradient descent for minimizing non-smooth convex functions [31]. Since λ is a constrained variable, we follow the projected supergradient ascent algorithm: iteratively updating the Lagrange multipliers according to the following update rule: $\lambda^{(t+1)} \leftarrow [\lambda^{(t)} + \alpha_t \nabla f(\lambda^{(t)})]_+$, where $\nabla f(\lambda^{(t)})$ is the supergradient of fat $\lambda^{(t)}$, α_t is the step-size and $[\cdot]_+$ is the projection operator that projects a vector onto the positive orthant. If the sequence of multipliers $\{\alpha_t\}$ satisfies $\alpha_t \geq 0$, $\lim_{t\to\infty} \alpha_t = 0$, $\sum_{t=0}^{\infty} \alpha_t = \infty$, then projected supergradient ascent converges to the optimum of the lagrangian dual [31].

To find the supergradient of $f(\lambda)$, note that f is a point-wise minimum of linear functions: i.e. $f(\lambda) = \min_{\mu} a_{\mu} \cdot \lambda + b_{\mu}$. It can be easily shown that the supergradient of f is given by $\nabla f(\lambda) = a_{\hat{\mu}(\lambda)}$, where $\hat{\mu}(\lambda) = \operatorname{argmin}_{\mu} a_{\mu} \cdot \lambda + b_{\mu}$. We omit the proof due to space constraints.

Mapping this definition to (4), we can see that the supergradient of f our formulation is given by (5): $\nabla f(\lambda) = -\begin{bmatrix} \Delta(\hat{\mu}(\lambda), \mu^1) - k_1 \\ \cdots \\ \Delta(\hat{\mu}(\lambda), \mu^{M-1}) - k_{M-1} \end{bmatrix}$ (5)

where $\hat{\boldsymbol{\mu}}(\boldsymbol{\lambda})$ is the optimal primal solution of (4) for the current setting of $\boldsymbol{\lambda}$. This supergradient (and the update procedure) has an intuitive interpretation. Recall that the Lagrangian relaxation minimizes a linear combination of the energy and similarity to the MAP solution, with the weighting given by $\boldsymbol{\lambda}$. If $\hat{\boldsymbol{\mu}}(\boldsymbol{\lambda}^{(t)})$ violates one of the diversity constraints, *i.e.* is less than k_m units away from a previous solution $\boldsymbol{\mu}^m$, then the supergradient w.r.t. $\lambda_m^{(t)}$ will be positive and the cost for violating the constraint will increase after the update, thus encouraging the next solution $\hat{\boldsymbol{\mu}}(\boldsymbol{\lambda}^{(t+1)})$ to satisfy the constraints. Conversely, if the diversity constraints are satisfied, the supergradient is negative indicating that $\lambda_m^{(t)}$ may be over-penalizing for violations and may be reduced to allow lower energy solutions.

5.2 Computing the Supergradient

Note that computing the supergradient requires solving the Δ -augmented energy minimization problem (4). At a high-level, this setup is a similar to cutting-plane methods for training Structured-SVMs [34], where the generation of each additional cutting plane requires solving the loss-augmented energy minimization problem. As we discuss next, for some classes of Δ -functions, we can solve the Δ -augmented energy minimization problem simply by reusing the same algorithms used for finding the MAP, by modifying the energy of the MRF. This allows all the developments in the MAP inference literature to be directly translated to the Diverse M-Best problem, without any changes.

Example: Dot-Product Dissimilarity: $\Delta(\mu, \mu^1) = -\sum_{i \in \mathcal{V}} \mu_i^T W \mu_i^1$, i.e. the sum of bilinear forms of labellings at nodes. For discrete solutions $\mu(s), \mu^1(s) \in \{0, 1\}$, setting W to the identity matrix (I) makes this dissimilarity function equivalent to the Hamming distance between the two solutions. In the presence of non-identity W, this is a weighted-Hamming distance, where W incorporates cross-label similarity. Moreover, note that $\sum_{A \in \mathcal{V} \cup \mathcal{E}} \theta_A \cdot \mu_A - \sum_{m=1}^{M-1} \lambda_m (\Delta(\mu, \mu^m) - k_m) = \sum_{i \in \mathcal{V}} (\theta_i + \sum_{m=1}^{M-1} \lambda_m W \mu_i^m) \cdot \mu_i + \sum_{(i,j) \in \mathcal{E}} \theta_{ij} \cdot \mu_{ij}$. Thus, $f(\lambda)$ becomes the same as the MAP problem with modified unary energies. When W = I, the modification simply increases the local cost of each of the previous states μ_i^m by λ_m . Non-identity W "smears" the affect of μ_i^m . For this Δ -function, we can use any existing MAP inference algorithm to solve this problem. Perhaps the most attractive feature is that the edge-energies are left unaffected. For instance, if they were submodular in the original model, they continue to be submodular. This allows us to use efficient graph-cut algorithms [7, 13].

Example: Higher-Order Dissimilarity. Another example of a useful dissimilarity function is one that decomposes into functions of subsets of variables, i.e. $\Delta(\mu, \mu^1) = \sum_{A \in \mathcal{V}} \Delta_A(\mu_A, \mu_A^1)$. If each of the these terms $\Delta_A(\cdot, \cdot)$ has some structure, e.g. cardinality potentials [33] or lower linear-envelope potentials [10] or sparse (pattern-based) higher-order potentials [14, 28], that allows for messages to be efficiently computed, this Δ -augmented energy minimization can be

performed via dual-decomposition based message-passing algorithms. The details can be found in the supplementary material (from the authors' webpages).

Finally, we note that successive supergradient computations (at $\lambda^{(t)}$ and $\lambda^{(t+1)}$) require solving fairly similar inference problems, which may be warm-started from the solutions of the previous iterations – either by re-using the search trees in graph-cuts [11] or by reusing messages in dual-decomposition.

5.3 Tightness of the Lagrangian Dual

Recall that the Lagrangian dual involves finding the tightest lower-bound for the primal, i.e. $\max_{\lambda \geq 0} f(\lambda) \leq Div \text{MBEST}(\Delta, \mathbf{k})$. One important question to ask is – when is the relaxation tight? To answer this question, we state the following theorem (see supplementary material from the authors' webpages for the proof):

Theorem 1. First, the Lagrangian dual $\max_{\lambda \geq 0} f(\lambda)$ is equivalent to solving (i.e. has zero duality gap with) the following primal relaxation of DivMBEST (Δ, \mathbf{k}) :

$$\min_{\boldsymbol{\mu}} \quad \sum_{A \in \mathcal{V} \cup \mathcal{E}} \boldsymbol{\theta}_A \cdot \boldsymbol{\mu}_A \tag{6a}$$

s.t.
$$\boldsymbol{\mu} \in Co\Big\{\mu_A(s) \in \{0,1\} \mid \boldsymbol{\mu} \in \mathcal{L}(G)\Big\}$$
 (6b)

$$\Delta(\boldsymbol{\mu}, \boldsymbol{\mu}^m) \ge k_m \qquad \forall m \in \{1, \dots, M-1\}$$
 (6c)

where $Co\{\cdot\}$ denotes the convex hull of the discrete solutions.

Second, for certain specific Δ -functions (e.g. the 0-1 function of M-Best MAP), the convex hull can be replaced by the discrete solutions themselves, i.e. (6b) may be replaced by $\mu_A(s) \in \{0,1\}$, $\mu \in \mathcal{L}(G)$. Thus for such specific families, the Lagrangian relaxation is tight.

However, if no assumptions are made on Δ , the Lagrangian relaxation is not guaranteed to be tight and may leave a duality gap.

Although the dot-product dissimilarity does not lead to a tight Lagrangian relaxation, we show in our experiments that even the relaxed solutions achieved by the Lagrangian dual could be very useful for the applications.

5.4 How Much Diversity?

Our DivMBEST formulation provides a principled way to trade off diversity vs. optimizing the energy. One issue of practical concern is how the tradeoff parameter k is chosen. The answer is related to the topology of the energy landscape. If k is set too low the next solution may not be able to escape the energy valley (and thus will not be a real "mode"). If k is set too high, then several modes will be ignored. An appropriate value of k is problem-dependent and must be tuned as a free parameter. Moreover, $\hat{\lambda}(k) = \operatorname{argmin}_{\lambda \geq 0} f(\lambda)$ the optimum setting of lambda is different for different values of k. Thus, instead of performing grid search on k and running supergradient ascent for each value of k, we can directly perform grid search on λ . Intuitively, this corresponds to learning from

data a linear weighting between the energy of the model and a diversity-inducing term. This is computationally much more efficient, both at the validation and testing stage. For our experiments, we learn the appropriate degree of diversity by tuning λ on a validation data set. This is similar in principle to the treatment of, say, a regularizer in learning. In the primal form, the regularization parameter limits the norm of the solution, but in practice the Lagrangian multiplier is treated as a norm penalty coefficient, and directly tuned via cross-validation.

6 Experiments

We apply our Diverse M-Best formulation, with (uniformly weighted) Hamming (dot product) dissimilarity Δ , to three scenarios:

- 1. An interactive segmentation (object cutout) setup in Section 6.1.
- 2. Category-level segmentation on PASCAL VOC 2010 data in Section 6.2.
- 3. Human articulated pose tracking in video in Section 6.3.

These scenarios use very different models (a binary pairwise submodular flat CRF, a multi-label hierarchical CRF with global factors, and a multi-label tree-CRF), with different MAP inference algorithms (max-flow/min-cut, α -expansion, dynamic programming). Despite the differences, our approach can naturally use the inference algorithm in each model to find the Diverse M-Best solutions.

We can compare Div MBEST against two alternatives for producing multiple solutions: **M-Best-MAP**, that produces low energy solutions without a focus on diversity, and perturbation-based techniques that simply produce diverse solutions without optimization within the original model. Specifically, given a MAP solution, we can produce additional solutions by changing assignments of a subset of nodes. The subset can be chosen either randomly (denoted **Random**), or based on the estimated confidence in the MAP labels; a natural measure of confidence is the entropy of the estimated min-marginals (**Confidence**). To make for a fair comparison, for each solution produced by Div MBEST that differs from MAP in S nodes, we generate a perturbation-based solution that changes exactly S nodes as well, thus achieving equal amount of diversity with Div MBEST.

We show that both these alternatives fall short in all scenarios. The M-Best-MAP produces redundant solutions with little extra information over the MAP, despite significant computational effort. The sampling-based methods produce solutions that are diverse, but improbable and inaccurate. In contrast, DivMBEST provides a principled way to trade-off both goals.

6.1 Interactive Segmentation

Interactive segmentation is a task where a user is interested in cutting out a foreground object of interest from an image via annotations like scribbles [6] or a coarse bounding box [29]. This is typically treated as a figure/ground segmentation task, with the MAP solution presented to the user. The user

MAP	DivMBest-dot prod.	DivMBest-HOP	M-Best	Random	Confidence
Acc.(%)91.542	95.16	93.82	91.59	91.68	93.17

Table 1. Interactive segmentation: pixel accuracies averaged over 50 test images

then provides additional supervision, leading to updated MAP solutions, till the MAP solution is acceptable. In order to minimize user interactions, the interface could show not a single cutout, but a set of possible cutouts for the user to simply select from. To make the most of this setup, this list of solutions must be small, diverse, and the algorithm that generates it must be efficient.

Dataset, Features, Energies, Inference. We simulate this scenario on 100 images from the PASCAL VOC 2010 dataset, and manually provided scribbles on objects contained in them; Fig. 2 shows examples. For each image, we set up a 2-label pairwise CRF, with a node term for each superpixel in the image and an edge term for each adjacent pair of superpixels. At each superpixel, we extract colour and texture features, and given the foreground/background scribbles on a single input image, train a Transductive SVM on these features. The node terms for the image are derived from the output of these TSVMs. The edge terms are contrast-sensitive Potts. A detailed description of the pipeline, the features and the parameter setting is in the supplementary materials. Fifty of the images were used for tuning the parameters, and the other 50 for reporting testing accuracies.

Baselines. The 2-label contrast-sensitive Potts model results in a submodular energy function so we can efficiently compute the exact MAP solution using graph-cuts implementation [13]. Moreover, with Hamming dissimilarity, we can efficiently and optimally solve the Δ -augmented energy minimization problem to compute the Diverse M-Best solutions using graph-cuts as well. As a first baseline, we implemented the M-best MAP algorithm of Yanover and Weiss [38], which requires repeated computation of min-marginals. We computed exact min-marginals using the dynamic graph-cuts algorithm of Kohli and Torr [12]; these were used to produce the **Confidence** baseline as well. Finally, we tested DivMBEST with a higher-order potential (HOP) dissimilarity; we include the results in Table 1 and give additional details in supplementary material.

Results. For each of the 50 test images in our dataset we generated MAP and 5 additional solutions using the methods described above. Table 1 shows the max accuracy of these 6 solutions for each method, averaged over 50 images. We can see that DivMBEST solutions result in the highest improvement over MAP. Figure 2 shows some example segmentations. Notice that the 2^{nd} -best MAP solution is nearly identical to the MAP solution whereas the solution from DivMBEST is qualitatively different, and could be significantly closer to the ground-truth labeling when MAP makes a mistake. In one case, the 2^{nd} mode found another instance of the object that MAP had missed, and in another, it completed a thin long structure (the arm of the person). These results suggest that the segmentation model is not completely accurate but still contains useful

segmentations as other modes of the CRF distribution. *DivMBest* provides a principled way of extracting these other segmentations.

6.2 Category Level Segmentation

Problem. In the second experiment we consider the problem of category-level segmentation, *i.e.* labeling each pixel in an image with one of 20 object categories or the background. This task is part of the PASCAL VOC comp5 challenge.

Model. The model we consider for this problem is the Associative Hierarchical CRF of Ladicky et~al.~[15], which achieved competitive results in recent years' VOC challenges. We used the publicly available implementation by the authors – the Automatic Labeling Environment (ALE) [17]. ALE includes many kinds of potentials: unary potentials based on textonboost features, P^n Potts terms (between superpixel nodes and pixel nodes) and a global co-occurence potential [16]. In a sequence of papers [15, 16], the authors developed a MAP inference algorithm for this model, and we are directly able to utilize this algorithm to compute DivMBEST solutions, by rerunning it with modified node energies.

Baselines. The same three baselines as in Section 6.1 are applicable here, however we found the M-best MAP algorithm [38] to be infeasibly slow for this model. Unlike the previous application, energy in this model is not submodular and does not allow for efficient computation of min-marginals. According to our estimates, it would take 10 years of CPU-time to compute each additional M-Best MAP solution for each image. On the other hand, computation of each additional solution for DivMBEST takes the same time as MAP. We also report the **Random** (averaged over ten runs) and **Confidence** baselines.

Results. We evaluated all methods on the VOC 2010 dataset, consisting of train, validation and test partitions with 964 images each. We use the standard PASCAL pixelwise "intersection / union" performance measure for each category. Since labels for test are not directly available, we first compare different methods for obtaining multiple solutions on validation (having trained the model on train), and then report accuracy on test (having trained on trainval).

Given a set of candidate segmentations, one could *select* a single solution from the set, or *combine* them, obtaining a solution not equal to any of the candidates. We discuss means of achieving this automatically below; here we consider an "oracle" evaluation protocol designed to measure the upper bound on performance of any eventual selection mechanism.

On validation data, for which we have ground truth, we obtained the oracle accuracy automatically by selecting, for each image, the solution with the highest pixel label accuracy averaged over categories. Table 2 shows the result of this evaluation for Div MBEST and the two sampling baselines, with M=10. On test we manually selected the solution that was visually perceived to be the best among the M solutions. Since the baselines are clearly inferior, we only evaluated the oracle on test with Div MBEST, M=10, with the results in Table 3. With larger values of M on validation (Fig. 3) the oracle bound goes

	Backgr.	Plane	Bicycle	Bird	Boat	Bottle	Bus	Car	Cat	Chair	Cow	D. Table	Dog	Horse	M.bike	Person	Plant	Sheep	Sofa	Train	TV.Mo.	Average
MAP	78.5	35.1	5.2	20.3	20.8	11.8	39.4	38.2	25.8	8.9	14.1	30.2	10.0	12.3	37.6	33.5	10.3	24.2	16.2	28.7	20.5	24.8
Conf.	78.5	35.1	5.3	20.1	20.7	12.6	39.4	37.9	26.8	8.9	14.1	30.2	10.3	12.2	39.5	33.4	10.6	24.2	17.3	28.4	20.5	25.1
Random	74.9	32.4	6.4	16.1	14.7	12.3	34.3	32.6	22.6	8.0	13.2	21.1	8.7	10.4	32.9	28.9	7.8	20.6	10.8	23.5	17.3	21.4
Div 10Best	85.6	53.9	14.6	36.9	33.6	33.2	64.2	56.3	47.7	16.1	30.3	46.8	29.1	28.7	59.0	50.0	32.5	46.7	31.2	52.9	39.0	42.3

Table 2. VOC 2010 Validation set accuracies

Table 3. VOC 2010 Test set accuracies

	Backgr.	Plane	Bicycle	Bird	Boat	Bottle	Bus	Car	Cat	Chair	Cow	D.Table	Dog	Horse	M. bike	Person	Plant	Sheep	Sofa	Train	TV.Mo.	Average
MAP Div10Best					$\frac{21.0}{34.5}$																	

further up and with M=30 the average accuracy on validation reaches about 48%, a near 7%-point improvement² over state of the art!

Ranking. The oracle results above indicate that the small set of diverse solutions from DivMBEST has a lot of potential; to realize this potential one needs a ranking and/or combination mechanism. A method recently proposed in [19] is designed to rank and combine a very large pool of segments (single label masks) most of which are of poor quality. This is different from the situation here (very small number of full image segmentations, some of which are of high quality) but a similar method may be applicable, and we are investigating this direction.

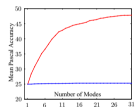


Fig. 3. Oracle accuracy vs. M on VOC2010 validation. Red: DivMBEST, blue: Confidence

6.3 Pose Estimation

Problem. We applied our formulation to the problem of tracking and estimating the pose of people in video sequences – a challenging problem due to appearance variation and articulation. We follow the setup of Park and Ramanan [25], where M candidate human poses are generated for every frame, and a single smooth track is selected using temporal context. We compare different methods that produce M-best pose estimates with respect to the quality of the tracks.

² The winning entry of VOC2010 competition achieved 40.1% on test. Accuracies on validation and test aren't directly comparable, but performance of most techniques is higher on test than val, and the purpose of the comparison here is to give a qualitative idea of relative performance.

Model. The model we consider for this problem is the articulated part-based model of Yang and Ramanan [37], which has demonstrated competitive performance on various benchmarks. The variables in the model are part (head, body, etc) locations and type. The graph-structure is a tree and (exact) inference is performed by dynamic programming. We used the authors' implementation, and modified node potentials to produce Diverse M-Best solutions.

The tracking model [25] is a chain-CRF, where each frame is a node whose label is the choice of the mode. Node potentials prefer low energy modes and edge-potentials prefer smooth transitions between successive frames. Exact MAP inference on the chain-CRF was also performed via dynamic-programming.

Dataset, Baselines. We used the dataset of [25] which consists of four video sequences (walking, pitching, lola1, lola2) of varying lengths for which a few frames have been manually annotated with ground-truth limb locations. We compared DivMBEST against the NBest method of [25], which is a special case of our formulation (as we describe in Section 4). Note that in [25] NBest was shown to outperform a number of sampling-based algorithms. We also compare to a Confidence baseline, where parts are repositioned to the next best position in the min-marginal table that hasn't been used for previously generated candidates. We omit Random since it is clearly inferior.

Results. Each algorithm was used to generate M candidate poses for all frames in every sequence. A track was computed using the chain-CRF to select a single pose for each frame. Algorithms were evaluated on the recovered tracks by computing Percentage of Correct Parts (PCP) scores (avg-CRF-PCP), which is shown in Fig. 4. DivMBest produced the best tracks for the vast majority of sequence-M combinations. For sequences lola1 and lola2 DivMBest significantly outperforms NBest by 10% and 4% points respectively.

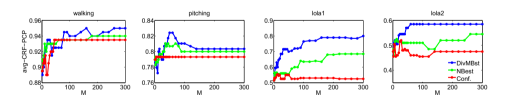


Fig. 4. Average PCP scores for the pose tracks vs. M (avg-CRF-PCP)

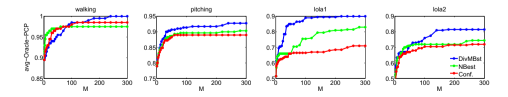


Fig. 5. Oracle PCP scores averaged across frames vs. M(avg-Oracle-PCP)

We also evaluate "oracle" accuracies by selecting for each frame the most accurate (w.r.t. the ground-truth) pose among M (avg-Oracle-PCP), which is shown in Fig. 5. In general, we observed that Div MBEST was able to produce better sets of hypotheses than NBest and Confidence across all values of M.

Finally, additional analysis, including average Hamming distance of the modes to the MAP solution, is included in the supplementary materials.

7 Discussions and Conclusion

We have presented the first algorithm for the Diverse M-Best MAP problem, which involves finding a diverse set of highly probable solutions under a discrete probabilistic model. Our Lagragian relaxation formulation involves solving the Δ -augmented energy minimization problem, minimizing a linear combination of the energy and similarity to previous solutions. We showed that this formulation is a generalization of the M-best MAP problem and that for certain classes of the Δ -function, DivMBEST can be computed using the same algorithms as those developed for computing the MAP solution. With some of the models and inference algorithms commonly used in vision, this can be tremendously useful.

Currently researchers have to design sophisticated high-order models for images and clever optimization methods to allow for reasonably efficient inference under such models. Our work suggests a different paradigm: use simpler models in which exact or approximate MAP inference, and thus DivMBEST inference with sufficiently nice Δ , is tractable, and obtain a set of diverse solutions. Then merely evaluate the more complex high-order model on these solutions to rank or otherwise combine them to provide the final output.

In this vein, our results bring into focus the problem of ranking a small set of diverse highly probably solutions. This is seen most dramatically in the results on VOC segmentation data; if one could only pick the most accurate segmentation in a pool of 30 segmentations for an image, one would improve the state of the art on VOC 2010 by 10%-points.

As future work we would like to investigate the performance and implications of other Δ -functions, apply them to higher-order energy functions and also apply this method to speed up cutting-plane methods for training Structural SVMs.

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