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UNIVERSITY OF CALIFORNIA,  
IRVINE

Approximate Inference in Graphical Models

DISSERTATION

submitted in partial satisfaction of the requirements  
for the degree of

DOCTOR OF PHILOSOPHY

in Computer Science

by

Sholeh Forouzan

Dissertation Committee:  
Professor Alexander Ihler, Chair  
Professor Rina Dechter  
Professor Charless Fowlkes

2015



# DEDICATION

To my loving and ever-supportive husband, Reza  
and to my mother, father and brother

AND

To Dina who showed me the true meaning of courage

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# CURRICULUM VITAE

Sholeh Forouzan

## EDUCATION

**Ph.D in Computer Science**

University of California, Irvine

**2015**

*Irvine, CA*

**M.S in Computer Science**

University of California, Irvine

**2011**

*Irvine, CA*

**M.S in AI and Robotics**

University of Tehran

**2008**

*Tehran, Iran*

**B.S in Computer Engineering**

Shahid Beheshti University

**2006**

*Tehran, Iran*

# ABSTRACT OF THE DISSERTATION

Approximate Inference in Graphical Models

By

Sholeh Forouzan

Doctor of Philosophy in Computer Science

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Professor Alexander Ihler, Chair

Graphical models have become a central paradigm for knowledge representation and reasoning over models with large numbers of variables. Any useful application of these models involves inference, or reasoning about the state of the underlying variables and quantifying the models' uncertainty about any assignment to them. Unfortunately, exact inference in graphical models is fundamentally intractable, which has led to significant interest in approximate inference algorithms.

In this thesis we address several aspects of approximate inference that affect its quality. First, combining the ideas from variational inference and message passing on graphical models, we study how the regions over which the approximation is formed can be selected more effectively using a content-based scoring function that computes a local measure of the improvement to the upper bound to log partition function. We then extend this framework to use the available memory more efficiently, and show that this leads to better approximations. We propose different memory allocation strategies and empirically show how they can improve the quality of the approximation to the upper bound. Finally, we address the optimization algorithms used in approximate inference tasks. Focusing on maximum *a posteriori* (MAP) inference and linear programming (LP), we show how the Alternating Direction Method of Multipliers (ADMM) technique can provide an elegant algorithm for finding the saddle point

of the augmented Lagrangian of the approximation, and present an ADMM-based algorithm to solve the primal form of the MAP-LP whose closed form updates are based on a linear approximation technique.

## Introduction

Graphical models are a powerful paradigm for knowledge representation and reasoning. Well known examples of graphical models include Bayesian networks, Markov random fields, constraint networks and influence diagrams. An early application of graphical models in computer science is medical diagnostics, in which medical specialists are interested in diagnosing the disease a patient might have by reasoning about the possible causes of a set of observed symptoms, or evaluate which future tests might best resolve the patient's underlying diseases. Another popular example application of graphical models is in computer vision, such as image segmentation and classification, where each image might consist of thousands of pixels and the goal is to figure out what type of object each pixel corresponds to.

To model such problems using graphical models, we represent them by a set of random variables, each of which represent some facet of the problem. Our goal is then to capture the uncertainty about the possible states of the world in terms of the joint probability distribution over all assignments to the set of random variables.

One of the main characteristics of such models is that there is going to be some significant uncertainty about the correct answer. Probability theory is used to deal with such uncertainty in a principled way by providing us with probability distributions as a declarative representation with clear semantics, accompanied by a toolbox of powerful reasoning pat-

terns like conditioning, as well as a range of powerful learning methodologies to learn the models from data.

While probability theory deals with modeling the uncertainty in such problems, in most cases we are still faced with another complexity: the very large number of variables to reason about. Even for the simplest case where each random variable is binary, for a system with  $n$  variables the joint distribution will have  $2^n$  states, which requires us to deal with representations that are intrinsically exponentially large. For these computational reasons, we exploit ideas from computer science, specifically graphs, to encode structure within this distribution and exploit the structure to represent and manipulate the distribution efficiently. The resulting graphical representation gives us an intuitive and compact data structure to encode high dimensional probability distributions, as well as a suite of methods that exploit the graphical structure for efficient reasoning. At the same time, the graph structure allows the parameters of the probability distribution to be encoded compactly, representing high dimensional probability distributions using a small number of parameters and allowing efficient learning of the parameters from data.

From this point of view, graphical models combine ideas from probability theory and ideas from computer science to provides powerful tools for describing the structure of a probability distribution and to organize the computations involved in reasoning about it. As a result, this framework has been used in a broad range of applications in areas including coding and information theory, signal and image processing, data mining, computational biology and computer vision.

## ■ 1.1 Examples of graphical models

**Protein side chain prediction.** Predicting side-chain conformation given the backbone structure is a central problem in protein-folding and molecular design. Proteins are chains of simpler molecules called amino acids. All amino acids have a common structure - a central carbon atom (C $\alpha$ ) to which a hydrogen atom, an amino group (NH $_2$ ) and a carboxyl group (COOH) are bonded. In addition, each amino acid has a chemical group called the side-chain, bound to C $\alpha$ . This group distinguishes one amino acid from another and gives its distinctive properties. Amino acids are joined end to end during protein synthesis by the formation of peptide bonds. An amino acid unit in a protein is called a residue. The formation of a succession of peptide bonds generate the backbone (consisting of C $\alpha$  and its adjacent atoms, N and CO, of each residue), upon which the side-chains are hung [Yanover and Weiss, 2003].

The goal of molecular design is then to predict the configuration of all the side-chains relative to the backbone. The standard approach to this problem is to define an energy function and use the configuration that achieves the global minimum of the energy as the prediction. To model this problem, a random variable is defined for each residue and the state of it represents the configuration of the side-chain of that residue. The factors in graphical model then capture the constraints and the energy of the interactions with the goal of finding a configuration that achieves the global minimum of the energy defined over the factors [Yanover and Weiss, 2003].

**Genetic linkage analysis.** In human genetic linkage analysis, the haplotype is the sequence of alleles at different loci inherited by an individual from one parent, and the two haplotypes (maternal and paternal) of an individual constitute that individual's genotype. However, this inheritance process is not easily observed. Measurement of the genotype of an individual typically results in a list of unordered pairs of alleles, one pair for each locus. This

information must be combined with pedigree data (a family tree of parent/offspring relationships) in order to estimate the underlying inheritance process [Fishelson and Geiger, 2002]. A graphical model representation of given pedigree of individuals with marker information (alleles at different loci) takes the form of a Bayesian network with variables representing the genotypes, phenotypes, and selection of maternal or paternal allele for each individual and locus. Finding the haplotype of individuals translates to an optimization task of finding the most probable explanation (mpe) of the Bayesian network. Another central task is linkage analysis, which seeks to find the loci on the chromosome that are associated with a given disease. This question can be answered by finding the probability of evidence over a very similar Bayesian network [Fishelson and Geiger, 2002].

## ■ 1.2 Inference

Inference in graphical models refers to reasoning about the state of the underlying variables and quantifying the model's uncertainty about any assignment to the random variables. For example, given the graphical model, we might be interested in finding the most likely configuration of variables and its value. This is an inference task that comes up in protein side-chain prediction, where the goal is then to find a configuration that achieves the maximum value of the objective function and recover optimal amino acid side-chain orientations in a fixed protein backbone structure.

Another equally important inference task is computing summations (marginalizing) over variables. Such inference task comes up when computing the marginal probability of a variable being in any of its states and computing the partition function. Both of these tasks are essential parts of training conditional random fields for image classification where we need to compute the partition function and the marginal distributions in order to evaluate the likelihood and its derivative. More importantly, because both of these quantities should

be computed for each training instance every time the likelihood is computed, we need to have efficient methods for it.

Unfortunately, like many interesting problems, inference in graphical models is NP-hard [Koller and Friedman, 2009a] as there are exponentially large number of possible assignments to the variables in the models. Despite such complexity, inference can be performed exactly in polynomial time for some graphical models that have simple structure like trees. The most popular exact algorithm, the junction tree algorithm, groups variables into clusters until the graph becomes a tree. Once an equivalent tree has been constructed, its marginals can be computed using exact inference algorithms that are specific to trees.

However, many interesting problems that arise ubiquitously in scientific computation are not amenable to such simple algorithm. For many real-world problems the junction tree algorithm is forced to make clusters which are very large and the inference procedure still requires exponential time in the worst case. Such complexity have inspired significant interest in approximate inference algorithms and had led to significant advances in approximation methods in the past decade.

### ■ 1.3 Approximate Inference

The complexity of exact inference have led to significant interest in approximate inference algorithms. Monte Carlo algorithms and variational methods are the two main classes that received the most attention. Monte Carlo algorithms are stochastic algorithms that attempt to produce samples from the distribution of interest. Variational algorithms on the other hand convert the inference problem into an optimization problem, trying to find a simple approximation that most closely matches the intractable marginals of interest. Generally, Monte Carlo algorithms are unbiased and given enough time, are guaranteed to sample from

the distribution of interest. However, in practice, it is generally impossible to know when that point of time has been reached. Variational algorithms, on the other hand, tend to be much faster, but they are inherently biased. In other words, no matter how much computation time they are given, they can not lessen the error that is inherent to the approximation.

## ■ 1.4 Summary of Contributions

Our ultimate goal is to advance the computational capabilities of reasoning algorithms for intelligent systems using the graphical models framework. We consider graphical models that involve discrete random variables and develop methods that allow us to improve on the existing approximate inference algorithms in several dimensions. We focus our attention to several areas that affect the quality of approximate inference:

- finding better approximations to the exponentially hard exact inference problem
- finding more efficient ways to use the available memory to improve the approximation
- finding better optimization algorithms to solve the approximate inference task faster and more accurately

As stated earlier an inherent source of error in variational methods is the approximation itself and thus finding better approximations is the key to reducing the error in the inference task. We show how such an approximation can be built incrementally in the context of weighted mini-bucket elimination for computing the partition function. We also study how approximate inference algorithms based on mini-bucket elimination use the available memory and develop algorithms that allow efficient use of the amount of memory available to construct better approximations and hence better approximate inference. Finally we focus our attention on the algorithms used to solve the optimization task for approximate inference

and develop an algorithm that uses Alternative direction method of multipliers (ADMM), a state of the art optimization algorithm, to find better solutions to the approximate inference problem for computing the maximizing assignment to variables.

To do so, this thesis first presents some background on problems and methods for graphical models in Chapter 2, then describes our contributions in three parts:

### **Chapter 3:**

- We describe how to use the message passing framework of Weighted Mini-bucket Elimination(WMBE) to select better regions to define the approximation and improve the bound
- We introduce a new merging heuristic for (weighted) mini-bucket elimination that uses message passing optimization of the bound, and variational interpretations, in order to construct a better heuristic for selecting moderate to large regions in an intelligent, energy-based way
- We propose an efficient structure update procedure that incrementally updates the join graph of mini-bucket elimination after new regions are added in order to avoid starting from scratch after each merge.

### **Chapter 4:**

- We describe how controlling the complexity of inference using *ibound* can result in inefficient use of resources and propose memory-aware alternatives
- We propose several memory allocation techniques to bypass the choice of a single control parameter for content-based MBE approximation that allows a more flexible control of memory during inference

- We expand the incremental region selection algorithm for weighted mini-bucket elimination to use a more fine tuned memory budget rather than a fixed *ibound*
- We perform an empirical evaluation of different allocation schemes, characterizing their behavior in practice
- We propose practical guidelines for choosing the memory allocation scheme for different classes of problems

## **Chapter 5:**

- We present an algorithm based on the Alternating Direction Method of Multipliers (ADMM) for approximate MAP inference using its linear programming relaxation
- We characterize different formulations of such problem using a graphical approach and discuss the challenges
- We propose a linear approximation to ADMM that allows solving the optimization efficiently

# Background

Graphical models capture the dependencies among large numbers of random variables by explicitly representing the independence structure of the joint probability distribution. It is useful to represent probability distributions using the notion of factor graphs. Such a representation allows the inference algorithms to be applied equally well to Bayesian networks and Markov random fields, as both of those can be easily converted to factor graphs.

For example, the factor graph in Figure 2.1 represents the joint distribution over random variables  $X_1, \dots, X_5$  as a collection of factors  $f_{ij}(X_i, X_j)$  over pairs of variables. Each random variable  $X_i$  can take one of several possible values  $x_i \in \mathcal{X}_i$ , where  $\mathcal{X}_i$  is called the domain and  $|\mathcal{X}_i|$  is called the cardinality of the variable. A factor  $f(X_1, \dots, X_N)$  is then a function (or table) that takes a set of random variables  $\{X_1, \dots, X_N\}$  as input, and returns a value for every assignment,  $(x_1, \dots, x_N)$ , to those random variables in the cross product space

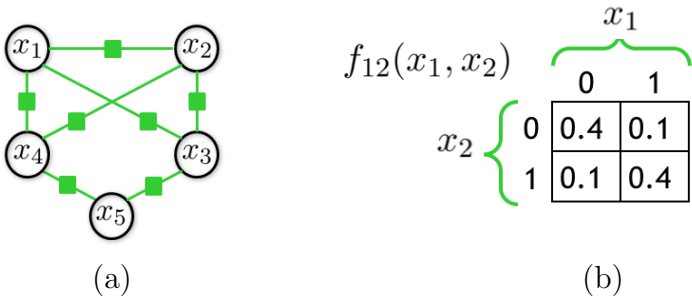


Figure 2.1: (a) Factor graph (b) Factor

$\mathcal{X} = \mathcal{X}_1 \times \dots \times \mathcal{X}_N$ . In the sequel, we use  $\text{var}(f)$  to refer to the set of input random variables,  $\{X_1, \dots, X_N\}$ , for factor  $f(X_1, \dots, X_N)$ . By this definition, a factor can be normalized, i.e.  $\sum_{\mathbf{x}_\alpha} f(\mathbf{x}_\alpha) = 1$  and  $f(\mathbf{x}_\alpha) \geq 0$  for all assignments  $\mathbf{x}_\alpha$ , or may be un-normalized so that it does not necessarily correspond to a probability distribution. These factors are the fundamental building blocks in defining distributions in a high dimensional space. A probability distribution over a large set of variables is then formed by multiplying factors:

$$p(x_1, \dots, x_N) = p(\mathbf{x}) = \frac{1}{Z} \prod_{\alpha \in \mathcal{I}} f_\alpha(\mathbf{x}_\alpha) \quad \text{where} \quad Z = \sum_{\mathbf{x}} \prod_{\alpha} f_\alpha(\mathbf{x}_\alpha)$$

Here  $\mathbf{x}_\alpha$  indicates the subset of variables that appear as arguments to factor  $f_\alpha$ , and  $Z$  is a constant which serves to normalize the distribution (called the partition function). We assume  $f(\mathbf{x}_\alpha) \geq 0$ . As shown in Figure 2.1, we can then associate  $p(\mathbf{x})$  with a graph  $G = (V, E)$ , where each variable  $X_i$  is associated with a node of the graph  $G$ . The node corresponding to  $X_i$  is connected to  $X_j$  if both variables are arguments of some factor  $f_\alpha(\mathbf{x}_\alpha)$ . The set  $\mathcal{I}$  is then a set of fully connected cliques in  $G$ .

Given such a representation, common inference tasks include finding the most likely or maximum a posteriori (MAP) configuration of  $p(\mathbf{x})$ , a combinatorial optimization problem, or computing the partition function  $Z$  or the marginal distributions of variables, a combinatorial summation problem:

$$\mathbf{x}^* = \arg \max_{\mathbf{x}} \prod_{\alpha \in \mathcal{I}} f_\alpha(\mathbf{x}_\alpha) \quad Z = \sum_{\mathbf{x}} \prod_{\alpha} f_\alpha(\mathbf{x}_\alpha) \quad b(x_i) = \sum_{\mathbf{x} \setminus x_i} p(\mathbf{x})$$

## ■ 2.1 Elimination Based Inference

Unfortunately, inference tasks such as computing the partition function or finding the most likely configuration of the variables are often computationally intractable for many real-world

problems. Elimination based methods such for exact inference, such as variable or ‘bucket’ elimination [Dechter, 1999] directly eliminate (by summing or maximizing) the variables, one at a time, along a predefined elimination ordering. The complexity of such an elimination procedure is exponential in the tree-width of the model, leading to a spectrum of approximations and bounds subject to computational limits. We first introduce the bucket elimination method [Dechter, 1999] in Section 2.1.1 and then the mini-bucket elimination method [Dechter and Rish, 2003] for approximate inference. We mainly focus our discussion on marginal inference, but the same methods can be applied to maximization tasks by replacing the sum operator with max.

### ■ 2.1.1 Bucket Elimination

Bucket elimination (BE) [Dechter, 1999] is an exact algorithm that directly eliminates variables in sequence. Given an elimination order, BE collects all factors that include variable  $X_i$  as their earliest-eliminated argument in a *bucket*  $B_i$ , then takes their product and eliminates  $X_i$  to produce a new factor over later variables, which is placed in the bucket of its “parent”  $\pi_i$ , associated with the earliest uneliminated variable:

$$\lambda_{i \rightarrow \pi_i}(\mathbf{x}_{i \rightarrow \pi_i}) = \sum_{x_i} \prod_{f_\alpha \in B_i} f_\alpha(\mathbf{x}_\alpha) \prod_{\lambda_{j \rightarrow i} \in B_i} \lambda_{j \rightarrow i}(\mathbf{x}_{j \rightarrow i})$$

where  $\mathbf{x}_{i \rightarrow \pi_i}$  is the set of variables that remain in the scope of the intermediate functions  $\lambda_{i \rightarrow \pi_i}$  after  $X_i$  is eliminated. This calculation can be conveniently organized as a “Bucket Elimination” procedure shown in Algorithm 2.1.

As an example, for the toy model of Figure 2.1, the BE algorithm takes the following steps: Given the elimination order  $o = [1, 2, 3, 4, 5]$ , the bucket elimination algorithm starts by eliminating variable  $X_1$  by grouping all factors containing  $X_1$  in their scope, in bucket  $B_1$

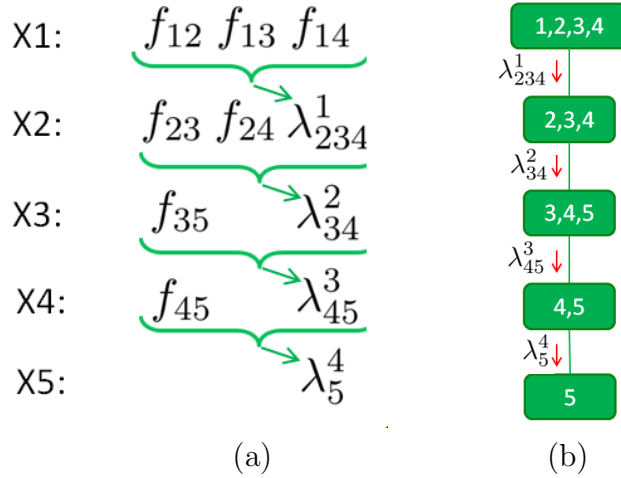


Figure 2.2: (a) Bucket Elimination (b) The Cluster Tree for Bucket Elimination

and eliminating  $X_1$  by

$$\lambda_{1 \rightarrow \pi_1}(x_2, x_3, x_4) = \sum_{x_1} f_{12} f_{13} f_{14},$$

where we use  $f_{ij}$  as shorthand for  $f_{ij}(x_i, x_j)$ . The elimination results in the intermediate function,  $\lambda_{1 \rightarrow \pi_1}$ . This intermediate function is then passed to bucket  $B_2$  (associated with variable  $X_2$ ), which is the first variable that will be eliminated from this function based on the elimination order  $o$ . Figure 2.2(a) shows how the original factors are grouped together in the buckets and how the messages generated by the algorithm are assigned to each bucket.

Execution of Bucket elimination algorithm induces a tree structure,  $CT = (V, E)$ , known as a cluster tree. The cluster tree then associates a node  $i \in V$  to each bucket  $B_i$ . An edge  $(i, j) \in E$  connects the two nodes  $i$  and  $j$  if the intermediate function  $\lambda_{i \rightarrow \pi_i}$  produced by processing bucket  $B_i$  is passed to bucket  $B_j$ .

From the cluster tree perspective, the functions  $\lambda_{i \rightarrow \pi_i}$  constructed during the elimination process can be interpreted as *messages* that are passed downward in a cluster tree representation of the model [Ihler et al., 2012]; see Figure 2.2(b). We will refer to bucket  $B_2$  as being

---

**Algorithm 2.1** Bucket/Variable Elimination for computing the partition function  $Z$ 

---

**Input:** Set of factors of a graphical model  $\mathbf{F} = \{f_\alpha(\mathbf{x}_\alpha)\}$ , an elimination order  $o = [x_1, \dots, x_N]$

**Output:** The partition function  $Z$

**for**  $i \leftarrow 1$  to  $N$  **do**

Find the set (bucket)  $\mathbf{B}_i$  of factors  $f_\alpha$  and intermediate messages  $\lambda_{j \rightarrow i}$  over variable  $x_i$ .

Eliminate variable  $x_i$ :

$$\lambda_{i \rightarrow \pi_i}(\mathbf{x}_{i \rightarrow \pi_i}) = \sum_{x_i} \prod_{f_\alpha \in \mathbf{B}_i} f_\alpha(\mathbf{x}_\alpha) \prod_{\lambda_{j \rightarrow i} \in \mathbf{B}_i} \lambda_{j \rightarrow i}(\mathbf{x}_{j \rightarrow i})$$

Update the factor list  $\mathbf{F} \leftarrow \{\mathbf{F} - \mathbf{B}_i\} \cup \{\lambda_{i \rightarrow \pi_i}(\mathbf{x}_{i \rightarrow \pi_i})\}$ :

**end for**

**Return:** partition function  $Z = \prod_{\lambda_{i \rightarrow \emptyset} \in \mathbf{F}} \lambda_{i \rightarrow \emptyset}$

---

the parent of bucket  $B_1$  and bucket  $B_1$  as being a child of bucket  $B_2$ . Note that a bucket may have multiple children, but can only have one parent.

Note that while computing the partition function of the probability distribution over the variables  $\{X_1 \dots X_5\}$  using a brute-force procedure requires a sum over a 5 dimensional tensor with computational complexity  $O(k^5)$ , where  $k$  is the number of possible states for  $X_i$ , the bucket elimination complexity is only  $O(k^4)$ .

In general, the space and time complexity of BE are exponential in the *induced width* of the graph along the elimination order, which is the maximum number of variables that appear together in a bucket and need to be reasoned about jointly. While good elimination orders can be identified using various heuristics [see e.g., Kask et al., 2011], this exponential dependence often makes direct application of BE intractable for many problems of interest.

## Computing Marginal Probabilities

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**Algorithm 2.2** Bucket/Variable Elimination to compute marginals  $p(x_i)$  [Dechter, 1999]

---

**Input:** Set of factors of a graphical model  $\mathbf{F} = \{f_\alpha(\mathbf{x}_\alpha)\}$ , an elimination order  $o = [x_1, \dots, x_N]$

**Output:** The partition function  $Z$  and the marginal distributions  $\{p(x_{B_i})\}$

**Forward Pass:** Use Algorithm 2.1 to calculate  $Z$  and all the messages  $\lambda_{i \rightarrow \pi_i}$  for all buckets  $B_i$

**Backward Pass:**

Initialize  $p(\mathbf{x}_{B_i}) = \prod_{f_\alpha \in B_i} f_\alpha(\mathbf{x}_\alpha) \prod_{\lambda_{j \rightarrow i} \in B_i} \lambda_{j \rightarrow i}(\mathbf{x}_{j \rightarrow i})$

**for**  $j \leftarrow N - 1$  to  $1$  **do**

    Compute a message from  $B_j$  to  $B_i$ ; where  $B_j$  is the bucket that receives the message  $\lambda_{i \rightarrow \pi_i}$  in the forward elimination

$$\lambda_{j \rightarrow i}(\mathbf{x}_{j \rightarrow i}) = \sum_{x_{B_j} \setminus x_{B_i}} \frac{p(\mathbf{x}_{B_j})}{\lambda_{i \rightarrow \pi_i}(\mathbf{x}_{i \rightarrow \pi_i})}$$

    Compute the marginal over variables in  $B_i$ :

$$p(\mathbf{x}_{B_i}) = \lambda_{j \rightarrow i}(\mathbf{x}_{j \rightarrow i}) p(\mathbf{x}_{B_i})$$

**end for**

**Return:** partition function  $Z$  and the marginals  $\{p(\mathbf{x}_{B_i})\}$

*Remark: Marginals over single variables can be computed by further marginalizing over the variables in the bucket. e.g.  $p(x_i) = \sum_{\pi_i} p(\mathbf{x}_{B_i})$*

---

In many cases we are interested in calculating the marginal probabilities,  $p(x_i)$ . To do so we can simply fix the value  $x_i$  and run the BE Algorithm 2.1 for all possible values of  $x_i$ . While simple, such approach is very inefficient as we often need to compute  $p(x_i)$  for all variables and all their possible values which repeat many of the same calculations. By sharing the repeated calculations, one can derive a more efficient algorithm. Dechter [1999] shows how an additional backward elimination can be used to compute the marginal probabilities recursively; this procedure is shown as Algorithm 2.2.

### ■ 2.1.2 Mini-bucket Elimination.

To avoid the complexity of bucket elimination, Dechter and Rish [2003] proposed an approximation in which the factors in bucket  $B_i$  are grouped into partitions  $Q_i = \{q_i^1, \dots, q_i^p\}$ , where each partition  $q_i^j \in Q_i$ , also called a *mini-bucket*, includes no more than  $ibound + 1$  variables. The user-selected bounding parameter  $ibound$  then serves as a way to control the complexity of elimination, as the elimination operator is applied to each mini-bucket separately. The  $ibound$  parameter then provides a flexible method to trade off between complexity and accuracy.

For the running example of Figure 2.1, the elimination process for  $X_1$ ,  $\sum_{x_1} f_{12}f_{13}f_{14}$ , can be approximated by the following upper and lower bounds to the exact elimination [Dechter and Rish, 2003]:

$$\sum_{x_1} f_{12}f_{13} \min_{x_1} f_{14} \leq \sum_{x_1} f_{12}f_{13}f_{14} \leq \sum_{x_1} f_{12}f_{13} \max_{x_1} f_{14},$$

which holds for all values of  $(x_2, x_3, x_4)$ . Mini-bucket approximation eliminates  $X_1$  from  $f_{12}f_{13}$  and  $f_{14}$  separately, instead of eliminating  $X_1$  from their product  $f_{12}f_{13}f_{14}$ ; this reduces the computational complexity from  $O(k^4)$  to  $O(k^3)$ , as the separate eliminations operate over smaller functions. Figure 2.3 (a) shows mini-bucket elimination with  $ibound = 2$  and Figure 2.3 (b) shows the cluster tree generated along the elimination procedure.

More generally, this approximation can be applied within each elimination step of Bucket Elimination (Algorithm 2.1), which results in a general mini-bucket elimination (MBE) algorithm given in Algorithm 2.3 [Dechter and Rish, 2003]. At each elimination step, MBE first splits the functions in the bucket  $B_i$  into smaller mini-buckets (partitions) and then eliminates  $X_i$  from each mini-bucket separately. The results are then passed to buckets later in the elimination order.

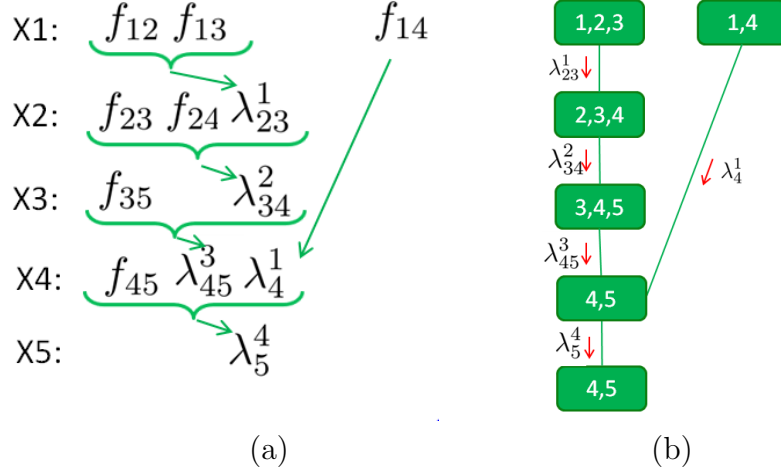


Figure 2.3: (a) Mini-Bucket Elimination (b) The Cluster Tree for Mini-Bucket Elimination

In general, using the inequality

$$\sum_{x_i} \prod_{f_\alpha \in B_i} f_\alpha \leq \left[ \sum_{x_i} \prod_{f_\alpha \in q_i^1} f_\alpha \right] \cdot \left[ \max_{x_i} \prod_{f_\alpha \in q_i^2} f_\alpha \right], \quad (2.1)$$

MBE gives an upper bound on the true partition function, where its time and space complexity are exponential in the user-controlled *ibound*. A similar inequality, with the max operator replaced by min can be used to find a lower bound on the exact elimination.

$$\sum_{x_i} \prod_{f_\alpha \in B_i} f_\alpha \geq \left[ \sum_{x_i} \prod_{f_\alpha \in q_i^1} f_\alpha \right] \cdot \left[ \min_{x_i} \prod_{f_\alpha \in q_i^2} f_\alpha \right], \quad (2.2)$$

Clearly, the complexity of MBE is reduced to  $O(k^{ibound})$  instead of  $O(k^{tw})$  of exact bucket elimination, where *tw* is the induced width of the graph along the elimination order. Smaller *ibound* values result in lower computational cost, but are typically less accurate; higher *ibound* values give more accurate results, but are computationally more expensive to compute.

Unfortunately the bounds defined by MBE are relatively loose and require use of a high *ibound* to achieve good results in practice. To overcome this drawback, Liu and Ihler [2011]

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**Algorithm 2.3** Mini-Bucket Elimination for computing the partition function  $Z$  [Dechter and Rish, 2003]

---

**Input:** Set of factors of a graphical model  $\mathbf{F} = \{f_\alpha(\mathbf{x}_\alpha)\}$ , an elimination order  $o = [x_1, \dots, x_N]$ , and an *ibound*

**Output:** An upper (lower) bound on partition function  $Z$

**for**  $i \leftarrow 1$  to  $N$  **do**

Find the set (bucket)  $\mathbf{B}_i$  of factors  $f_\alpha$  and intermediate messages  $\lambda_{j \rightarrow i}$  over variable  $x_i$ .

Partition  $\mathbf{B}_i$  into  $p$  subgroups  $q_i^1, \dots, q_i^p$  such that  $\cup_{j=1}^p q_i^j = \mathbf{B}_i$  and

$$|\cup_{f \in q_i^j} \text{var}(f)| \leq \text{ibound} + 1 \text{ for all } j = 1, \dots, p$$

Eliminate variable  $x_i$ :

**for**  $m = 1 \dots p$  **do**

$$\lambda_{i \rightarrow \pi_i}(\mathbf{x}_{i \rightarrow \pi_i}) = \begin{cases} \sum_{x_i} (\prod_{f_\alpha \in q_i^m} f_\alpha(\mathbf{x}_\alpha) \prod_{\lambda_{j \rightarrow i} \in q_i^m} \lambda_{j \rightarrow i}(\mathbf{x}_{j \rightarrow i})), & \text{if } m = 1 \\ \max_{x_i} (\prod_{f_\alpha \in q_i^m} f_\alpha(\mathbf{x}_\alpha) \prod_{\lambda_{j \rightarrow i} \in q_i^m} \lambda_{j \rightarrow i}(\mathbf{x}_{j \rightarrow i})), & \text{if } m \neq 1 \end{cases}$$

**end for**

Update the factor list  $\mathbf{F} \leftarrow \{\mathbf{F} - \mathbf{B}_i\} \cup \{\lambda_{i \rightarrow \pi_i}(\mathbf{x}_{i \rightarrow \pi_i})\}$ :

**end for**

**Return:** the bound on partition function  $Z = \prod_{\lambda_{i \rightarrow \emptyset} \in \mathbf{F}} \lambda_{i \rightarrow \emptyset}$

*Remark:* A lower bound can be obtained by replacing the *max* in elimination step with *min*

---

introduced *weighted mini-bucket elimination* which uses a more general bound based on Höllder's inequality that can give tighter bounds for small *ibound* values. The details of weighted MBE are explained in section 2.1.3.

It is important to highlight the relationship between mini-bucket elimination and message passing, as mini-bucket elimination can be interpreted in the context of a cluster tree and message passing on it. The corresponding cluster tree for a particular partitioning then

contains a cluster for each mini-bucket, with its scope being the union of the scope of all the factors in that mini-bucket, and MBE bound corresponds to a problem relaxation in which a copy of shared variables between the mini-buckets is made for each, and the result of eliminations  $\lambda_{i \rightarrow \pi_i}$ s correspond to the messages in a cluster tree defined on the augmented model over the variable copies; see Figure 2.3(b). This cluster tree is guaranteed to have induced-width *ibound* or less. The problems are equivalent if all copies of  $X_i$  are constrained to be equal; otherwise, the additional degrees of freedom lead to a relaxed problem and thus can generate an upper bound. This connection allows us to apply any of the message passing paradigms to our inference problem and inspires combining message passing techniques with MBE based approximate inference techniques and balance the positive properties of both.

Another factor that affects the mini-bucket bound significantly is the choice of partitions  $Q_i$ . In Chapter 3, we explain the existing heuristics that have been used to guide the choice of partitions, and introduce a new general partitioning heuristic within weighted mini-bucket that results in better approximation quality.

### ■ 2.1.3 Weighted Mini-bucket

A recent improvement to mini-bucket generalizes the MBE bound with a “weighted” elimination [Liu and Ihler, 2011]. Compared to standard MBE, which approximates the intractable summation operators with upper/lower bounds based on max/min operators, weighted mini-bucket builds an approximation based on the more general powered sum with a “weighted” elimination step. An upper or lower bound can then be computed depending on the signs of the weights.

Weighted mini-bucket applies Hölder’s and reverse Hölder inequalities, which provide general tools for constructing bounds or approximations to the sum of products, to form the foundation for of a mini-bucket approximation [Liu and Ihler, 2011]. For a set of positive

functions  $f_\alpha(x_\alpha)$ ,  $\alpha = 1, \dots, n$  defined over discrete variables  $x$ , and a set of non-zero weights  $w = [w_1, \dots, w_n]$ , Hölder's inequality results in an upper bound

$$\sum_x \prod_\alpha f_\alpha \leq \prod_r \left[ \sum_{x_i} f_{\alpha^{w_r}}^{\frac{1}{w_r}} \right]^{w_r} \quad \text{for } w \in \mathcal{W}^+ \quad (2.3)$$

and reverse Hölder's inequality results in a lower bound

$$\sum_x \prod_\alpha f_\alpha \geq \prod_r \left[ \sum_{x_i} f_{\alpha^{w_r}}^{\frac{1}{w_r}} \right]^{w_r} \quad \text{for } w \in \mathcal{W}^- \quad (2.4)$$

where

$$\mathcal{W}^+ = \left\{ w : \sum_r w_r = 1 \text{ and } w_r > 0, \forall r = 1, \dots, n \right\}$$

$$\mathcal{W}^- = \cup_{k=1}^n \mathcal{W}_k^- \quad \text{where} \quad \mathcal{W}_k^- = \left\{ w : \sum_r w_r = 1 \text{ and } w_k > 0, w_r < 0, \forall r \neq k \right\}$$

so that  $\mathcal{W}^+$  corresponds to a probability simplex on the weights  $w_r$ , and  $\mathcal{W}^-$  corresponds to a normalized weights with exactly one positive element. Given equations (2.3) and (2.4), the sum of products of functions can be approximated using products of powered sums over individual functions and can be used to provide upper and lower bounds for the partition function in a manner similar to mini-bucket elimination.

For the running example of Figure 2.1, weighted mini-bucket approximation results in the following elimination, where  $w_1$  and  $w_2$  are weights satisfying  $w_1 + w_2 = 1$  and the direction of inequality depends on the signs of the weights  $[w_1, w_2]$ .

$$\sum_{x_1} f_{12} f_{13} f_{14} \begin{array}{c} \geq \\ \leq \end{array} \left[ \sum_{x_1} (f_{12} f_{13})^{\frac{1}{w_1}} \right]^{w_1} \left[ \sum_{x_1} f_{14}^{\frac{1}{w_2}} \right]^{w_2}$$

It is interesting to note that the powered sum  $\sum_x^w f(x) = \left( \sum_x f(x)^{\frac{1}{w}} \right)^w$  approaches  $\max_x f(x)$

when  $w \rightarrow 0+$  and  $\min_x f(x)$  as  $w \rightarrow 0$ . Consider the upper bound of equation (2.3) for a partition  $q_i^1, q_i^2$ , which takes the form

$$\sum_{x_i} \prod_{f_\alpha \in B_i} f_\alpha \leq \left[ \sum_{x_i} \prod_{f_\alpha \in q_i^1} f_\alpha^{\frac{1}{w_1}} \right]^{w_1} \cdot \left[ \sum_{x_i} \prod_{f_\alpha \in q_i^2} f_\alpha^{\frac{1}{w_2}} \right]^{w_2},$$

where  $w_i > 0$  and  $w_1 + w_2 = 1$ . It is easy to see that this bound generalizes the mini-bucket upper bound where  $w_1 = 1$  and  $w_2 \rightarrow 0^+$ . The same argument holds for the lower bound (2.4), as it generalizes the mini-bucket lower bound where  $w_1 = 1$  and  $w_2 \rightarrow 0^-$ .

Based on these Hölder inequality bounds, Liu and Ihler [2011] then generalized mini-bucket elimination to *weighted mini-bucket elimination* (WMB), presented in Algorithm 2.4, by replacing the mini-bucket bound with Hölder’s inequality. The general procedure is the same, except that the sum/max operators are replaced with weighted sums where the weights are normalized to sum to one for each variable.

Liu and Ihler [2011] also show that the resulting bound is equivalent to a class of bounds based on tree reweighted (TRW) belief propagation [Wainwright et al., 2005], or more generally conditional entropy decompositions (CED) [Globerson and Jaakkola, 2007a], on a join-graph defined by the mini-bucket procedure. This connection is used to derive fixed point reparameterization updates, which change the relative values of the factors  $f_\alpha$  while keeping their product constant in order to tighten the bound. Section 2.2 gives some background on variational methods for inference and further explains this connection.

## Variable Splitting Perspective

It is useful to consider the effect of partitioning of functions into different mini-buckets and eliminating in each mini-bucket separately. This procedure effectively splits a variable into one or more replicates, one for each mini-bucket. From this perspective, mini-bucket

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**Algorithm 2.4** Weighted Mini-Bucket Elimination for computing the partition function  $Z$  [Liu and Ihler, 2011]

---

**Input:** Set of factors of a graphical model  $\mathbf{F} = \{f_\alpha(\mathbf{x}_\alpha)\}$ , an elimination order  $o = [x_1, \dots, x_N]$ , and an *ibound*

**Output:** An upper (lower) bound on partition function  $Z$

**for**  $i \leftarrow 1$  to  $N$  **do**

Find the set (bucket)  $\mathbf{B}_i$  of factors  $f_\alpha$  and intermediate messages  $\lambda_{j \rightarrow i}$  over variable  $x_i$ .

Partition  $\mathbf{B}_i$  into  $p$  subgroups  $q_i^1, \dots, q_i^p$  such that  $\cup_{j=1}^p q_i^j = \mathbf{B}_i$  and

$$|\cup_{f \in q_i^j} \text{var}(f)| \leq \text{ibound} + 1 \text{ for all } j = 1, \dots, p$$

Assign a weight  $w_j$  to each partition  $q_i^j$  such that  $\sum_j w_j = 1$ .

Eliminate variable  $x_i$  from each partition:

**for**  $m = 1 \dots p$  **do**

$$\lambda_{i \rightarrow \pi_i}(\mathbf{x}_{i \rightarrow \pi_i}) = \left[ \sum_{x_i} \prod_{f_\alpha \in q_i^m} f_\alpha^{\frac{1}{w_{m_i}}} \right]^{w_{m_i}}$$

**end for**

Update the factor list  $\mathbf{F} \leftarrow \{\mathbf{F} - \mathbf{B}_i\} \cup \{\lambda_{i \rightarrow \pi_i}(\mathbf{x}_{i \rightarrow \pi_i})\}$ :

**end for**

**Return:** the bound on partition function  $\bar{Z} = \prod_{\lambda_{i \rightarrow \emptyset} \in \mathbf{F}} \lambda_{i \rightarrow \emptyset}$

*Remark: An upper or lower bound can be computed based on the sign of the weights  $w_{m_i}$*

---

elimination (Algorithm 2.3) is identical to bucket elimination (Algorithm 2.1), but executes on the replicas of variables in different mini-buckets.

WMB uses this perspective to define the bounds on an augmented model defined by splitting variables which characterizes the weighted mini-bucket bound as an explicit function of the augmented model parameters and the weights. This formulation then allow the development of efficient algorithms to optimize the parameters and the weights to obtain the tightest bound.

To make this formulation clear, note that the process of partitioning the functions assigned to the mini-buckets can be interpreted as replicating the variables that appear in different mini-buckets. Let  $\bar{\mathbf{x}}_i = \{x_i^r\}_{r=1}^{R_i}$  be the set of  $R_i$  copies of variable  $x_i$ . Also let  $\bar{\mathbf{w}}_i = \{w_i^r\}_{r=1}^{R_i}$  be the corresponding collection of weights for each replicate such that  $\sum_{r=1}^{R_i} w_i^r = 1$ . The sets  $\bar{\mathbf{x}} = \{\bar{\mathbf{x}}_1, \dots, \bar{\mathbf{x}}_n\}$  and  $\bar{\mathbf{w}} = \{\bar{\mathbf{w}}_1, \dots, \bar{\mathbf{w}}_n\}$  thus represent the collection of variable copies and the weights. The elimination order  $o = [1, \dots, n]$  on the original variables  $\mathbf{x}$  can then be simply extended to the set  $\bar{\mathbf{x}}$  as  $\bar{o} = [1^1, \dots, 1^{R_1}, \dots, n^1, \dots, n^{R_n}]$ . Let  $\bar{\boldsymbol{\theta}} = \{\bar{\theta}_\alpha : \alpha \in \mathcal{I}\}$  be the set of factors defined over the set of variable copies  $\bar{\mathbf{x}}$  where  $\bar{\theta}_\alpha = \log \bar{f}_\alpha$ . Such formulation allows computing the log-partition function as a sequential powered sum,

$$\Phi(\bar{\boldsymbol{\theta}}, \bar{\mathbf{w}}) = \log \sum_{\bar{x}_m}^{\bar{w}_m} \cdots \sum_{\bar{x}_1}^{\bar{w}_1} \prod_{\alpha \in \mathcal{I}} \exp(\bar{\theta}_\alpha(\bar{\mathbf{x}}_\alpha)) \quad (2.5)$$

It is therefore possible to jointly optimize  $\bar{\boldsymbol{\theta}}$  and  $\bar{\mathbf{w}}$  to get the tightest bound. Computing the tightest upper bound then requires solving the following optimization problem

$$\min_{\bar{\boldsymbol{\theta}}, \bar{\mathbf{w}}} \Phi(\bar{\boldsymbol{\theta}}, \bar{\mathbf{w}}) \text{ s.t. } \bar{\boldsymbol{\theta}} \in \boldsymbol{\Theta}, \bar{\mathbf{w}} \in \mathcal{W}^+ \quad (2.6)$$

Here  $\boldsymbol{\Theta}$  is the set of augmented natural parameters  $\bar{\boldsymbol{\theta}} = \{\bar{\theta}_{\bar{\alpha}} : \bar{\alpha} \in \bar{\mathcal{I}}\}$  that are consistent with the original model in that  $\sum_{\bar{\alpha} \in \bar{\mathcal{I}}} \bar{\theta}_{\bar{\alpha}}(\bar{\mathbf{x}}_\alpha) = \sum_{\alpha \in \mathcal{I}} \theta_\alpha(\mathbf{x}_\alpha)$  for all the values  $\bar{x}_{ir} = x_i$ . And  $\mathcal{W}^+$  is the set of weights that makes the corresponding Hölder inequality to hold, namely,

$$\mathcal{W}^+ = \{\bar{\mathbf{w}} : \sum_r \bar{w}_{ir} = 1, \bar{w}_{ir} \geq 1, \forall i, r\}$$

This optimization is convex and its global optimum can be calculated efficiently. Liu and Ihler [2011] show how the forward-backward Algorithm 2.5 can be used to calculate  $\bar{\Phi}(\bar{\boldsymbol{\theta}}, \bar{\mathbf{w}})$  in a forward pass which is identical to weighted mini-bucket Algorithm 2.4. A backward pass then calculates the approximate marginals. These marginals can then be used to compute

the derivatives of  $\bar{\Phi}(\bar{\boldsymbol{\theta}}, \bar{\mathbf{w}})$  with respect to  $\bar{\boldsymbol{\theta}}$  and  $\bar{\mathbf{w}}$  as:

$$\frac{\partial}{\partial \bar{\theta}_\alpha} \bar{\Phi}(\bar{\boldsymbol{\theta}}, \bar{\mathbf{w}}) = \bar{p}_w(\bar{\mathbf{x}}_\alpha), \quad (2.7)$$

$$\frac{\partial}{\partial \bar{w}_k} \bar{\Phi}(\bar{\boldsymbol{\theta}}, \bar{\mathbf{w}}) = H_w(\bar{x}_k | \bar{\mathbf{x}}_{k+1:n}; \bar{p}_w) \quad (2.8)$$

In principle, having the values  $\bar{\Phi}(\bar{\boldsymbol{\theta}}, \bar{\mathbf{w}})$  and derivatives as output from Algorithm 2.5, we could directly apply black-box optimization routines to optimize the bound. However since  $\bar{\Phi}(\bar{\boldsymbol{\theta}}, \bar{\mathbf{w}})$  is calculated using a relatively expensive forward-backward message-passing algorithm, it seems most efficient to update  $\bar{\boldsymbol{\theta}}$  and  $\bar{\mathbf{w}}$  while computing messages.

Moment matching conditions are used to update  $\bar{\boldsymbol{\theta}}$  given the positive weights  $\bar{\mathbf{w}} > 0$ . To do so, first an "average marginal"  $p_m(x_i)$  is computed using the geometric mean of the marginals of the replicates  $\{\bar{p}_w(\bar{x}_{ir})\}$  and then  $\bar{\boldsymbol{\theta}}$  is adjusted to correct for the difference between the marginal of each replicate and the average marginal as follows:

$$p_m(x_i) = \left[ \prod_r \bar{p}_w(\bar{x}_{ir} = x_i)^{\bar{w}_{ir}} \right]^{\frac{1}{\sum_r \bar{w}_{ir}}}$$

$$\bar{\theta}_{c_{ir}} = \bar{\theta}_{c_{ir}} + \bar{w}_{ir} \log \frac{p_m(x_i)}{\bar{p}_w(\bar{x}_{ir})}$$

Entropy matching can then be used to update  $\{\bar{w}_{ir}\}$  in  $\bar{\mathcal{W}}^+$ , fixing the augmented natural parameters  $\bar{\boldsymbol{\theta}}$ , giving the following updates:

$$\bar{w}_{ir} = \bar{w}_{ir} \exp \left[ -\epsilon \bar{w}_{ir} (H_{ir|\prec} - \sum_r \bar{w}_{ir} H_{ir|\prec}) \right], \quad \forall r = 1, \dots, R_i$$

$$\bar{w}_{ir} = \frac{\bar{w}_{ir}}{\sum_r \bar{w}_{ir}}, \quad \forall r = 1, \dots, R_i$$

Very important aspect of such optimization which directly impacts the tightness of the bound is the structure of the cluster tree over which the augmented model  $\bar{p}(\bar{\mathbf{x}})$  is formed. Such structure can be formed by running the mini-bucket elimination Algorithm 2.3 which requires partitioning the functions that belong to a bucket into smaller mini-buckets. Different partitioning strategies result in different structures than can affect the tightness of the bound and have been previously studied for selecting better mini-buckets for MBE algorithm [Rollon and Dechter, 2010]. In Chapter 3 we study one such partitioning strategy which allows us to add new regions to the cluster tree  $\bar{\mathcal{G}}$  incrementally to achieve tighter bounds.

---

**Algorithm 2.5** Calculating the WMB bound  $\bar{\Phi}(\bar{\theta}, \bar{w})$  and its derivatives [Liu and Ihler, 2011]

---

**Input:** Set of augmented factors of a graphical model  $\{\bar{f}_{c_k}(\bar{\mathbf{x}}_{c_k})\}$  and their cluster tree  $\bar{\mathcal{G}}$ . A weight vector  $\bar{\mathbf{w}} = [\bar{w}_1, \dots, \bar{w}_n]$ . An elimination order  $\bar{o} = [1^1, \dots, 1^{R_1}, \dots, n^1, \dots, n^{R_n}]$

**Output:**  $\bar{\Phi}(\bar{\theta}, \bar{w})$  and its derivatives

**Forward pass:**

**for**  $i \leftarrow 1$  to  $\bar{n}$  **do**

$$\lambda_{i \rightarrow l}(\bar{\mathbf{x}}_{i \rightarrow l}) = \left[ \sum_{\bar{\mathbf{x}}_i} (f_{c_i}(\bar{\mathbf{x}}_i)) \prod_{j \in \text{child}(i)} \lambda_{j \rightarrow i}(\bar{\mathbf{x}}_{j \rightarrow i})^{\frac{1}{w_i}} \right]^{w_i}$$

where  $l = \pi_i$  is the parent of  $c_i$  and  $\text{child}(i)$  is the set of nodes that have  $i$  as their parents.

**end for**

**Backward pass:**

**for**  $i \leftarrow \bar{n}$  to 1 **do**

$$\lambda_{k \rightarrow i}(\bar{\mathbf{x}}_{k \rightarrow i}) = \left[ \sum_{\bar{\mathbf{x}}_k \setminus \bar{\mathbf{x}}_i} (f_{c_k}(\bar{\mathbf{x}}_k)) \prod_{j \in \delta(i)} \lambda_{j \rightarrow i}(\bar{\mathbf{x}}_{j \rightarrow i})^{\frac{1}{w_i}} \lambda_{i \rightarrow k}(\bar{\mathbf{x}}_{i \rightarrow k})^{-\frac{1}{w_i}} \right]^{w_{m_i}}$$

where  $\delta(i)$  is all the nodes connected to  $i$  in the cluster tree  $\bar{\mathcal{G}}$  including its parent  $k$

**end for**

**Compute the bound:**

$$\bar{\Phi}(\bar{\theta}, \bar{w}) = \log \prod_k \left( \sum_{\bar{\mathbf{x}}_k}^{\bar{w}_k} [\bar{f}_{c_k}(\bar{\mathbf{x}}_k) \prod_{i \in \delta(k)} \lambda_{i \rightarrow k}(\bar{\mathbf{x}}_{i \rightarrow k})] \right)$$

where  $k$  lists all the clusters with no parents in  $\bar{\mathcal{G}}$ .

**Compute the approximate marginals:**

$$\bar{p}_w(\bar{\mathbf{x}}_{c_k}) \propto [\bar{f}_{c_k}(\bar{\mathbf{x}}_k) \prod_{i \in \delta(k)} \lambda_{i \rightarrow k}(\bar{\mathbf{x}}_{i \rightarrow k})]^{\frac{1}{w_k}}$$


---

## ■ 2.2 Variational Methods

All inference methods discussed so far use a sequence of variable elimination steps to compute the partition function. In contrast, variational methods convert the inference problem into an optimization task over the space of distributions. The goal is then to find a distribution minimizing a divergence measure to the target distribution over which we can solve the inference problem more efficiently. To review the variational inference framework next we introduce the exponential family form for graphical models and the variational form for the log-partition function.

### ■ 2.2.1 Exponential Family and Marginal Polytope

The factorized distribution  $p(\mathbf{x}) = \frac{1}{Z} \prod_{\alpha} f_{\alpha}(\mathbf{x}_{\alpha})$  can be written into an overcomplete exponential family form [Wainwright and Jordan, 2008b],

$$p(\mathbf{x}; \boldsymbol{\theta}) = \exp(\boldsymbol{\theta}(\mathbf{x}) - \Phi(\boldsymbol{\theta})) \quad \boldsymbol{\theta}(\mathbf{x}) = \sum_{\alpha \in \mathcal{I}} \theta_{\alpha}(\mathbf{x}_{\alpha}) \quad (2.9)$$

where  $\theta_{\alpha}(\mathbf{x}_{\alpha}) = \log(f_{\alpha}(\mathbf{x}_{\alpha}))$  are the log of the factors  $f_{\alpha}$  in our graphical model. The values  $\theta_{\alpha}$  are called the *natural parameters* of the exponential family and the vector  $\boldsymbol{\theta} = \{\theta_{\alpha}(\mathbf{x}_{\alpha}) : \alpha \in \mathcal{I}, \mathbf{x}_{\alpha} \in \mathcal{X}_{\alpha}\}$  is a vector formed by concatenating all the natural parameters.  $\Phi(\boldsymbol{\theta})$  is then the log-partition function, that normalizes the distribution

$$\Phi(\boldsymbol{\theta}) = \log \sum_{\mathbf{x}} \exp(\boldsymbol{\theta}(\mathbf{x})) \quad (2.10)$$

The framework of variational inference converts the problem of computing the log-partition function (2.10) to an optimization problem:

$$\Phi(\boldsymbol{\theta}) = \max_{q \in \mathcal{P}_x} \{\mathbb{E}_q(\boldsymbol{\theta}(\mathbf{x})) + H(\mathbf{x}; q)\} \quad (2.11)$$

where  $\mathcal{P}_x$  is the set of all possible distributions defined on  $\mathbf{x}$  which can be represented as

$$\mathcal{P}_x = \{q(\mathbf{x}) : \sum_{\mathbf{x}} q(\mathbf{x}) = 1 \text{ and } q(\mathbf{x}) \geq 0, \forall \mathbf{x}\} \quad (2.12)$$

and  $H(\mathbf{x}; q) = -\mathbb{E}_q(\log q(\mathbf{x}))$  is the entropy of distribution  $q$ .

This representation has several interesting properties[Wainwright and Jordan, 2008b]:

1.  $\Phi(\boldsymbol{\theta})$  is a convex function of  $\boldsymbol{\theta}$ .
2. The maximum is obtained at  $q^*(\mathbf{x}) = p(\mathbf{x}) = \exp(\theta(\mathbf{x}) - \Phi(\boldsymbol{\theta}))$ .
3. The derivatives of  $\Phi(\boldsymbol{\theta})$  with respect to the natural parameters  $\theta_\alpha(\mathbf{x}_\alpha)$  equal the marginal distribution of  $p(\mathbf{x})$ , as

$$\frac{\partial \Phi(\boldsymbol{\theta})}{\partial \theta_\alpha(\mathbf{x}_\alpha)} = p(\mathbf{x}_\alpha), \quad \forall \alpha \in \mathcal{I}, \mathbf{x}_\alpha \in \mathcal{X}_\alpha \quad (2.13)$$

An important simplification is possible by observing that  $\mathbb{E}_q(\theta(\mathbf{x}))$  can be decomposed into smaller terms defined only over the factor scopes  $\alpha$  as:

$$\mathbb{E}_q(\theta(\mathbf{x})) = \sum_{\alpha} \mathbb{E}_q(\theta_\alpha(\mathbf{x}_\alpha)) = \sum_{\alpha} \sum_{\mathbf{x}_\alpha} q(\mathbf{x}_\alpha) \theta_\alpha(\mathbf{x}_\alpha)$$

This simplification allows us to represent each  $q(\mathbf{x}) \in \mathcal{P}_x$  which is intrinsically high dimensional with  $(\prod_i |\mathcal{X}_i| - 1)$  values by  $\sum_{\alpha} (\prod_{X_i \in \alpha} |\mathcal{X}_i| - 1)$ .

Additionally it allows defining the *marginal polytope*  $\mathcal{M}(\mathcal{I})$  as the set of all possible marginal distributions  $\boldsymbol{\mu} := \{\mu_\alpha : \alpha \in \mathcal{I}, \mathbf{x}_\alpha \in \mathcal{X}_\alpha\}$  that are consistent with the joint distribution on  $\mathbf{x}$ . More specifically:

$$\mathcal{M}(\mathcal{I}) = \{\boldsymbol{\mu} : \exists q(\mathbf{x}) \in \mathcal{P}_x \text{ such that } \mu_\alpha(\mathbf{x}_\alpha) = \sum_{\mathbf{x} \setminus \mathbf{x}_\alpha} q(\mathbf{x}) \quad \forall \alpha \in \mathcal{I}, \mathbf{x}_\alpha \in \mathcal{X}_\alpha\}$$

As a result the optimization (2.11) can now be defined as an optimization over the marginal polytope  $\mathcal{M}(\mathcal{I})$  as:

$$\Phi(\boldsymbol{\theta}) = \max_{\boldsymbol{\mu} \in \mathcal{M}(\mathcal{I})} \{\langle \boldsymbol{\mu}, \boldsymbol{\theta} \rangle + H(\mathbf{x}; \boldsymbol{\mu})\} \quad (2.14)$$

where  $\langle \boldsymbol{\mu}, \boldsymbol{\theta} \rangle$  is the inner product,

$$\langle \boldsymbol{\mu}, \boldsymbol{\theta} \rangle = \sum_{\alpha \in \mathcal{I}} \sum_{\mathbf{x}_\alpha} \boldsymbol{\mu}(\mathbf{x}_\alpha) \boldsymbol{\theta}_\alpha(\mathbf{x}_\alpha)$$

As a result both the partition function and marginals can be computed from a single continuous optimization (2.14) instead of the sum inference task.

Unfortunately, simply casting the log-partition function as an optimization problem does not cause the computation to be tractable and the optimization in (2.14) remains intractable for two reasons: (1) the entropy  $H(\mathbf{x}; \boldsymbol{\mu})$  is intractable to compute in general, and (2) the marginal polytope,  $\mathcal{M}$ , is difficult to characterize exactly, requiring an exponential number of linear constraints in general.

Approximating the variational form (2.14) efficiently then requires three components: (1) approximating the marginal polytope  $\mathcal{M} \approx \bar{\mathcal{M}}$ ; (2) approximating the entropy  $H(\mathbf{x}; \boldsymbol{\mu}) \approx \bar{H}(\mathbf{x}; \boldsymbol{\mu})$ ; and (3) solving the continuous optimization with those approximations.

As a result, some of the important problems in variational approximations include the selection of an appropriate entropy approximation  $\bar{H}$ , and the choice of regions, or subsets of variables whose beliefs are represented directly which often affects both the accuracy of  $\bar{H}$  and the form of the polytope approximation  $\bar{\mathcal{M}}$ .

## □ 2.2.2 WMBE - The Variational View

As described in section 2.1.3, weighted mini-bucket elimination (WMBE) can be used to compute an upper or lower bound of the log-partition function by passing weighted messages forward along an elimination order. The weighted summation and Hölder's inequality were then used to compute its bounds (2.3) and (2.4). Liu and Ihler [2011] show that this bound can be interpreted from a variational perspective by first approximating the marginal polytope by a marginal polytope on the cluster tree formed by the partitions and then bounding the exact entropy using a weighted conditional entropy.

To make this connection clear, note that the process of partitioning the functions assigned to the mini-buckets can be interpreted as replicating the variables that appear in different mini-buckets. Let  $\bar{\mathbf{x}}_i = \{x_i^r\}_{r=1}^{R_i}$  be the set of  $R_i$  copies of variable  $x_i$ . Also let  $\bar{\mathbf{w}}_i = \{w_i^r\}_{r=1}^{R_i}$  be the corresponding collection of weights for each replicate such that  $\sum_{r=1}^{R_i} w_i^r = 1$ . The sets  $\bar{\mathbf{x}} = \{\bar{\mathbf{x}}_1, \dots, \bar{\mathbf{x}}_n\}$  and  $\bar{\mathbf{w}} = \{\bar{\mathbf{w}}_1, \dots, \bar{\mathbf{w}}_n\}$  thus represent the collection of variable copies and the weights. The elimination order  $o = [1, \dots, n]$  on the original variables  $\mathbf{x}$  can then be simply extended to the set  $\bar{\mathbf{x}}$  as  $\bar{o} = [1^1, \dots, 1^{R_1}, \dots, n^1, \dots, n^{R_n}]$ . Let  $\bar{\boldsymbol{\theta}} = \{\bar{\theta}_\alpha : \alpha \in \mathcal{I}\}$  be the set of factors defined over the set of variable copies  $\bar{\mathbf{x}}$ . Such formulation allows, the primal WMB bound to be

$$\Phi(\boldsymbol{\theta}) \leq \Phi(\bar{\boldsymbol{\theta}}, \bar{\mathbf{w}}) = \log \sum_{\bar{\mathbf{x}}_m}^{\bar{w}_m} \cdots \sum_{\bar{\mathbf{x}}_1}^{\bar{w}_1} \prod_{\alpha \in \mathcal{I}} \exp(\bar{\theta}_\alpha(\bar{\mathbf{x}}_\alpha)) \quad (2.15)$$

From the variational perspective, the above replaces the marginal polytope  $\mathcal{M}$  on the original variables, with the marginal polytope  $\bar{\mathcal{M}}$  on the variable replicates which means the search for the mean vector  $\boldsymbol{\mu} \in \mathcal{M}$  is replaced with searching for some extended mean vector  $\bar{\boldsymbol{\mu}} \in \bar{\mathcal{M}}$ . The entropy can be bounded using a weighted sum of the conditional entropies,

computed using the marginals  $\bar{\boldsymbol{\mu}}$ , on the mini-bucket graph

$$H(\boldsymbol{\mu}) \leq \bar{H}_{\bar{\mathbf{w}}}(\bar{\boldsymbol{\mu}}) = \sum_{i \in \bar{o}} \bar{w}_i H(\bar{x}_i | \bar{\mathbf{x}}_{i+1:\bar{n}}; \bar{\boldsymbol{\mu}}) \quad (2.16)$$

Plugging the approximations to the marginal polytope  $\bar{\mathcal{M}}$  and conditional entropy  $\bar{H}_{\bar{\mathbf{w}}}(\bar{\boldsymbol{\mu}})$  in to (2.14) results in the dual WMB bound to the log partition function

$$\Phi(\boldsymbol{\theta}) \leq \max_{\bar{\boldsymbol{\mu}} \in \bar{\mathcal{M}}} \{\langle \bar{\boldsymbol{\mu}}, \bar{\boldsymbol{\theta}} \rangle + \bar{H}_{\bar{\mathbf{w}}}(\bar{\boldsymbol{\mu}})\} \quad (2.17)$$

In principle, we could use a variety of methods to directly optimize (2.17). However, as Liu and Ihler [2011] show, the primal WMB bound in (2.15) can be optimized efficiently via simple message passing updates described in Algorithm 2.5 which is the route we follow in our later experiments involving WMB.

### ■ 2.2.3 Variational Methods for Maximization

The variational form (2.14) casts the problem of computing the log-partition function as a continuous optimization problem. A similar representation exists for the maximization tasks in graphical models as:

$$\Phi(\boldsymbol{\theta}) = \max_{\boldsymbol{\mu} \in \mathcal{M}(\mathcal{I})} \{\langle \boldsymbol{\mu}, \boldsymbol{\theta} \rangle\} \quad (2.18)$$

which provides a powerful toolkit for the max-inference; a combinatorial optimization problem. It is interesting to note that the form (2.18) and (2.14) differ only by the addition of the entropy term in the latter. Intuitively for the maximization we expect the marginals of the optimal distribution to correspond to a single assignment while for the summation the entropy term causes the optimal distribution to spread across many high-probability

assignment to increase the entropy.

Approximating the marginal polytope  $\mathcal{M}$  in (2.18) with a more manageable set such as the local consistency polytope  $\mathcal{L}$  results in an approximation known as a relaxation of *linear programming relaxation* for MAP inference. Many efficient algorithms have been developed to solve the linear program (2.18), including max-product linear programming (MPLP) [Globerson and Jaakkola, 2007b] and dual decomposition [Komodakis et al., 2011, Sontag et al., 2010]. In Chapter 5 we discuss how to use augmented Lagrangian methods combined with the Alternating Direction Method of Multipliers to solve this optimization more efficiently.

# Incremental Region Selection for Mini-bucket Elimination Bounds

### ■ 3.1 Region Choice for MBE

The popularity of mini-bucket elimination [Dechter and Rish, 2003] (discussed in detail in Chapter 2) has led to its use in many reasoning tasks; MBE is often used to develop heuristic functions for search and combinatorial optimization problems [Dechter and Rish, 2003, Kask and Dechter, 2001, Marinescu and Dechter, 2007, Marinescu et al., 2014], as well as to provide bounds on weighted counting problems such as computing the probability of evidence in Bayesian networks [Rollon and Dechter, 2010, Liu and Ihler, 2011].

A critical component of the mini-bucket approximation is the set of partitions formed during the elimination process. By bounding the size of each partition and eliminating within each separately, MBE ensures that the resulting approximation has bounded computational complexity, while providing an upper or lower bound. A single control variable, the *ibound*, allows the user to easily trade off between accuracy and computational complexity (including both memory and time). The partitioning of a bucket into mini-buckets of bounded size can be accomplished in many ways, each resulting in a different accuracy. From the variational

perspective, this corresponds to the critical choice of *regions* in the approximations, defining which sets of variables will be reasoned about jointly.

Traditionally, MBE is guided only by the graph structure, using a *scope-based* heuristic [Dechter and Rish, 2003] to minimize the number of buckets. However, this ignores the importance of the function *values* on the bound. More recent extensions such as Rollon and Dechter [2010] have suggested ways of incorporating the function values into the partitioning process, with mixed success. A more bottom-up construction technique is the *relax-compensate-recover* (RCR) method of Choi and Darwiche [2010], which constructs a sequence of mini-bucket-like bounds of increasing complexity.

Variational approaches typically use a greedy, bottom-up approach termed *cluster pursuit*. Starting with the smallest possible regions, the bounds are optimized using message passing, and then new regions are added greedily from an enumerated list of clusters such as triplets [e.g., Sontag et al., 2008, Komodakis and Paragios, 2008]. This technique is often very effective if only a few regions can be added, but the sheer number of regions considered often creates a computational bottleneck and prevents adopting large regions [see, e.g., Batra et al., 2011].

We propose a hybrid approach that is guided by the graph structure in a manner similar to the mini-bucket construction, but takes advantage of the iterative optimization and scoring techniques of cluster pursuit. In practice, we find that our methods work significantly better than either the partitioning heuristics of Rollon and Dechter [2010], or a pure region pursuit approach. We also discuss the connections of our work to RCR [Choi and Darwiche, 2010]. We validate our approach with experiments on a wide variety of problems drawn from recent UAI approximate inference competitions [Elidan et al., 2012].

### ■ 3.1.1 Partitioning Methods

As discussed above, mini-bucket elimination and its weighted variant compute a partitioning over each bucket  $B_i$  to bound the complexity of inference and compute an upper bound on the partition function  $Z$ . However, different partitioning strategies will result in different upper bounds. Rollon and Dechter [2010] proposed a framework to study different partitioning heuristics, and compared them with the original scope based heuristic proposed by Dechter and Rish [1997]. Here we summarize several approaches.

**Scope-based Partitions.** Proposed by Dechter and Rish [1997], scope-based partitioning is a simple but effective top-down approach that tries to minimize the number of mini-buckets in  $B_i$  by including as many functions as possible in each mini-bucket  $q_i^k$ . To this end, it first orders the factors in  $B_i$  by decreasing number of arguments. Starting from the largest, each factor  $f_\alpha$  is then merged with the first available mini-bucket that satisfies the computational limits, i.e., where  $|\text{var}(f) \cup \text{var}(q_i^j)| \leq \text{ibound} + 1$  where  $\text{var}(f)$  represents the scope of function  $f$ . If there are no mini-buckets available that can include the factor, a new mini-bucket is created and the scheme continues until all factors are assigned to a mini-bucket.

Scope-based partitioning provides an efficient way to quickly decide which regions to include in the cluster graph. However, because it relies solely on the function arguments, it ignores a significant source of information: the functions values themselves.

Consider an example when eliminating variable  $X_1$  from a bucket containing four factors,  $\{f_{14}, f_{17}, f_{123}, f_{156}\}$  with  $\text{ibound} = 3$ . Suppose that the two functions  $f_{123}$  and  $f_{156}$  return 1 for every assignment to the variables in their scope, i.e.  $f_{123} = \mathbb{1}$  and  $f_{156} = \mathbb{1}$ . Eliminating  $X_1$  from the bucket without partitioning is then proportional to eliminating  $X_1$  from the

$$\begin{array}{ccc}
\left. \begin{array}{c} \underbrace{f_{14} \ f_{17} \ f_{123} \ f_{156}} \\ C \times \sum_{\mathbf{x}} f_{14} \ f_{17} \\ \text{(a)} \end{array} \right| & \left. \begin{array}{c} \underbrace{f_{14} \ f_{17}} \ \underbrace{f_{123}} \ \underbrace{f_{156}} \\ C \times \sum_{\mathbf{x}} f_{14} \ f_{17} \\ \text{(b)} \end{array} \right| & \left. \begin{array}{c} \underbrace{f_{123} \ f_{14}} \ \underbrace{f_{156} \ f_{17}} \\ C \times \sum_{\mathbf{x}} f_{14} \ \max_{\mathbf{x}} f_{17} \\ \text{(c)} \end{array} \right.
\end{array}$$

Figure 3.1: Comparing approximations. Suppose  $f_{123} = f_{156} = \mathbb{1}$ ; then (a) exact elimination provides the same answer as (b) an approximation using three mini-buckets, while (c) the scope-based approximation of two mini-buckets may give a loose upper bound.

product  $f_{14}f_{17}$  as

$$\sum_{x_1} f_{14}f_{17}f_{123}f_{156} = C \times \sum_{x_1} f_{14}f_{17} \tag{3.1}$$

Two ways of partitioning these four functions into different mini-buckets are shown in Figure 3.1(b) and (c). Scope-based partitioning aims to minimize the number of mini-buckets, and so groups the functions into two mini-buckets,  $\{f_{123}, f_{14}\}$  and  $\{f_{156}, f_{17}\}$ . However, another choice would be to use three partitions  $\{f_{123}\}$ ,  $\{f_{156}\}$ , and  $\{f_{14}, f_{17}\}$ . Comparing the results of elimination from these two partitionings,

$$\sum_{x_1} f_{17}f_{123} \max_{x_1} f_{17}f_{156} = C \times \left( \sum_{x_1} f_{14} \max_{x_1} f_{17} \right) \tag{3.2}$$

$$\sum_{x_1} f_{17}f_{14} \max_{x_1} f_{123} \max_{x_1} f_{156} = C \times \sum_{x_1} f_{14}f_{17} \tag{3.3}$$

with the exact elimination (3.1), we can see that the scope-based partitioning results in a potentially loose upper bound, while eliminating from the three partitions (3.3) provides the same value as exact elimination. In other words, the particular values of  $f_{123}$  and  $f_{156}$  cause a partitioning that looks less accurate (three mini-buckets) to actually provide a much more accurate approximation.

**Content-based Partitions.** Since looking at function values can help choose better mini-

buckets, Rollon and Dechter [2010] explored informed selection of variable groupings into mini-buckets based on content-based distance functions, and proposed various *content-based partitioning heuristics* that seek to find a partitioning that is closest to the true bucket function,  $g_i = \sum_{x_i} \prod_{\alpha \in B_i} f_\alpha$ .

Rollon and Dechter [2010] frame this selection process as an optimization problem,

$$Q^* = \arg \min_Q \text{dist}(g_i^Q, g_i), \quad (3.4)$$

where  $Q = \{q_i^1, \dots, q_i^p\}$  is a partitioning of  $B_i$  with bounding parameter *ibound* and

$$g_i^Q = \prod_{j=1}^p \sum_{x_i} \prod_{\alpha \in q_i^j} f_\alpha$$

is the function represented by the partitioning  $Q$ . Rollon and Dechter [2010] studied the effectiveness of several different distance functions across multiple problem instances; however, no single distance was found to consistently outperform scope-based partitioning.

It is important to note that we can not efficiently compute all possible partitionings and score them to decide which one to choose, so Rollon and Dechter [2010] proposed a greedy approach that organizes the space of partitionings into a lattice using the refinement relation between partitions. Also, in general, computing the distance (3.4) is exponential in the number of arguments of  $g_i$ , since we need to measure the distance of the resulting approximation function  $g_i^Q$  relative to the intractably large true function  $g_i$ . However, for some types of distance functions like relative error, they show how to derive a local measure that only requires computing functions in the two candidate mini-buckets to be merged and can be interpreted as the penalty or error associated with keeping them separated. In this chapter, we use the ideas presented in [Rollon and Dechter, 2010] and extend them to form a new hybrid partitioning heuristic.

**Relax-Compensate-Recover.** Choi and Darwiche [2010] indirectly address the problem of partition selection within their *Relax, Compensate and Recover* framework, in which certain equality constraints in the graph are first relaxed in order to reduce the computational complexity of inference. New auxiliary factors are then introduced to compensate for the relaxation and enforce a weaker notion of equivalence. The recovery process then aims to identify those equivalence constraints whose relaxation were most damaging and recover them. Choi and Darwiche [2010] proposed a number of recovery heuristics, including mutual information and residual recovery.

### ■ 3.1.2 Variational bounds.

The variational viewpoint of inference corresponds to optimizing an objective function over a collection of *beliefs* that are constrained to lie within the marginal polytope, or set of marginal probabilities that can be achieved by some joint distribution. Efficient approximations are developed by relaxing the optimization to enforce only a subset of the constraints – for example that the beliefs be consistent between overlapping cliques. In the case of the log partition function, we also approximate the entropy term in the objective; for example, the weighted mini-bucket bound is:

$$\log Z \leq \max_{b_\alpha \in \mathbb{L}} \sum_{\alpha} \mathbb{E}_{b_\alpha}[\log f_\alpha] + \sum_{i,\alpha} w_{i\alpha} H(x_i | x_{\alpha \setminus i}; b_\alpha)$$

where  $\sum_{\alpha} w_{i\alpha} = 1$  for all  $i$ . Like mini-bucket bounds, the quality of variational bounds depends significantly on the choice of regions, which determine both the constraints that will be enforced and the form of the entropy approximation. Traditionally, research on variational approximations for the log partition function has focused more on the optimization of the bound through message passing than the region selection aspect. Often, regions are chosen to match the original model factors, and then may be improved using methods like cluster pursuit, described next.

**Cluster Pursuit.** Sontag et al. [2008] studied the problem of region selection for MAP inference in the context of *cluster-based* dual decomposition relaxations. They developed a bottom-up approach in which regions (typically cycles or triplets) are added incrementally: First, the dual decomposition bound is optimized through message passing. Then, a pre-defined set of clusters, such as triplets or the faces of a grid, are scored by computing a lower bound on their potential improvement to the dual objective; the scoring function used measures the difference between independently maximizing each pairwise factor, versus jointly maximizing over the triplet. After adding the best-scoring cluster, or a small number of them, the procedure repeats. Similar cycle repair processes were also proposed by Komodakis and Paragios [2008] and Werner [2008], and related cluster pursuit methods have also been applied to summation problems [Welling, 2004, Hazan et al., 2012]. However, scoring all possible clusters often becomes a computational bottleneck; for example, to make the process more efficient, Batra et al. [2011] proposed pre-selection heuristics to reduce the number of clusters considered.

## ■ 3.2 A Hybrid Approach

To summarize, existing elimination based approaches avoid the space and time complexity of exact inference by using a top-down partitioning approach that mimics the construction of a junction tree and allows large regions to be added to the approximation quickly. In contrast, message passing algorithms often use cluster pursuit methods to select regions, a bottom-up approach in which a predefined set of clusters (such as triplets) are scored based on their potential improvement to the bound, and incrementally added.

It is important to highlight the relationship between mini-bucket elimination and message passing, since mini-bucket elimination can be interpreted in the context of cluster graph and message passing on it. This connection inspires combining message passing techniques with

mini-bucket elimination based approximate inference techniques and balance the positive properties of both. To balance the effectiveness of both approaches, being able to add larger regions while taking into account their potential improvement to the upper bound for the log partition function, our hybrid scheme, like mini-bucket, uses the graph structure to guide region selection, while also taking advantage of the iterative optimization and scoring techniques of cluster pursuit.

Cluster pursuit algorithms use the function values, and more concretely the improvement to the bound produced by them, in order to select regions that tighten the upper bound effectively. However, there are often prohibitively many clusters to consider: for example, in a fully connected pairwise model, there are  $O(n^3)$  triplets,  $O(n^4)$  possible 4-cliques, etc., to score at each step. For this reason, cluster pursuit methods typically restrict their search to a predefined set of clusters, such as triplets [Sontag et al., 2008]. Our proposed approach uses the graph structure to guide the search for regions, restricting our search to merging pairs of existing clusters within a single bucket at a time. This allows us to constrain the complexity of the search and add larger regions more effectively.

In contrast, the content-based heuristics for region selection of Rollon and Dechter [2010] use the graph structure as a guide, but their scoring scheme only takes into account the messages from the earlier buckets in the elimination order. Our proposed hybrid approach uses iterative optimization on the junction tree in order to make more effective partitioning decisions. Algorithm 3.1 describes the overall scheme of our hybrid approach, which is explained in detail next.

### ■ 3.2.1 Initializing a join tree

Given a factor graph  $G$  and a bounding parameter  $ibound$ , we start by initializing a join graph, using a min-fill elimination ordering [Dechter, 2003] (which we denote  $\mathbf{o} = \{x_1, \dots, x_n\}$ )

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**Algorithm 3.1** Incremental region selection for WMBE

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**Input:** factor graph ( $\mathcal{G}$ ), bounding parameter  $ibound$  and maximum number of iterations  $T$

**Initialize**  $wmb$  to a join graph using e.g. a min-fill ordering  $o$ , uniform weights and uniform messages

**for** each bucket  $B_i$  following the elimination order **do**

**repeat**

$(q_i^m, q_i^n) \leftarrow \text{SelectMerge}(Q_i)$

$\mathcal{R} \leftarrow \text{AddRegions}(wmb, o, q_i^m, q_i^n)$

$wmb \leftarrow \text{MergeRegions}(wmb, \mathcal{R})$

**for**  $iter = 1$  **to**  $T$  **do**

      // pass forward messages and reparameterize:

$wmb \leftarrow \text{msgForward}(wmb)$

      // pass backward messages:

$wmb \leftarrow \text{msgBackward}(wmb)$

**end for**

**until** no more merges possible

**end for**

---

without loss of generality) and  $ibound = 1$ . For any given bucket  $B_i$ , this results in each mini-bucket (or region)  $q_i^k \in B_i$  containing a single factor  $f_\alpha$ . In the sequel, to simplify the notation, we refer to the mini-bucket  $q_i^k$  simply as region  $k$  when the context is clear. We denote the result of the elimination as  $\lambda_{k \rightarrow l}$  and the variables in its scope as  $\text{var}(\lambda_{k \rightarrow l})$ .  $\lambda_{k \rightarrow l}$  is then a message that is sent to the bucket  $B_j$  of its first-eliminated argument in  $\mathbf{o}$ . Here,  $l = \pi_k$  denotes the parent region of  $k$  which can be one of the initial mini-buckets in  $B_j$  if  $\text{var}(\lambda_{k \rightarrow l}) \subseteq \text{var}(l)$ , or be a new mini-bucket containing a single factor,  $f_l = \mathbb{1}$ , that assigns the value one to every assignment to the variables in  $\text{var}(\lambda_{k \rightarrow l})$ . In our implementation we choose  $l \in B_j$  to be the mini-bucket with the largest number of arguments,  $|\text{var}(l)|$ , such that  $\text{var}(\lambda_{k \rightarrow l}) \subseteq \text{var}(l)$ .

Using the weighted mini-bucket elimination scheme of Algorithm 2.4, we initialize the mini-bucket weights  $w^r$  uniformly within each bucket  $B_i$ , so that for  $q_i^r \in Q_i$ ,  $w^r = \frac{1}{|Q_i|}$ , which ensures  $\sum_{q_i^r \in Q_i} w^r = 1$ .

### □ 3.2.2 Message Passing

We use iterative message passing on the join graph to guide the region selection decision. Having built an initial join graph, we use the weighted mini-bucket messages [Liu and Ihler, 2011] to compute forward and backward messages, and perform reparameterization of the functions  $f_\alpha$  to tighten the bound.

Let  $k$  be a region of the mini-bucket join graph, and  $l$  its parent,  $l = \pi_k$ , with weights  $w^k$  and  $w^l$ , and  $f_k(\mathbf{x}_k)$  the product of factors assigned to region  $k$ .  $\mathbf{x}_k$  is then the union of all the variables in the scope of the factors assigned to region  $k$ . Then we compute the forward messages as,

**Forward Messages:**

$$\lambda_{k \rightarrow l}(\mathbf{x}_l) = \left[ \sum_{\mathbf{x}_k \setminus \mathbf{x}_l} [f_k(\mathbf{x}_k) \prod_{t: l = \pi_t} \lambda_{t \rightarrow l}(\mathbf{x}_l)]^{\frac{1}{w^k}} \right]^{w^k} \quad (3.5)$$

and compute the upper bound using the product of forward messages computed at roots of the join graph,

**Upper bound on Z:**

$$Z \leq \prod_{k: \pi_k = \emptyset} \lambda_{k \rightarrow \emptyset} \quad (3.6)$$

In order to tighten the bound, we compute backward messages in the join graph,

**Backward Messages:**

$$\lambda_{l \rightarrow k}(\mathbf{x}_k) = \left[ \sum_{\mathbf{x}_l \setminus \mathbf{x}_k} [f_l(\mathbf{x}_l) \prod_{t \in \delta(l)} \lambda_{t \rightarrow l}(\mathbf{x}_l)]^{\frac{1}{w^l}} [\lambda_{k \rightarrow l}(\mathbf{x}_l)]^{-\frac{1}{w^k}} \right]^{w^k}$$

where  $\delta(l)$  is the set of all neighbors (parent and children) of region  $l$  in the cluster tree. We then use these incoming messages to compute a weighted belief at region  $k$ , and repara-

parameterize the factors  $f_k$  for each region  $k$  in a given bucket  $B_i$  (e.g.,  $k \in Q_i$ ) to enforce a weighted moment matching condition:

**Reparameterization:**

$$\begin{aligned}
 b_k(x_i) &= \sum_{\mathbf{x}_k \setminus x_i} [f_k(\mathbf{x}_k) \prod_{t \in \delta(l)} \lambda_{t \rightarrow k}(\mathbf{x}_k)]^{\frac{1}{w^k}} \\
 \bar{b}(x_i) &= \left[ \prod_{k \in Q_i} b_k(x_i) \right]^{1/\sum_k w^k} \\
 f_k(\mathbf{x}_k) &\leftarrow f_k(\mathbf{x}_k) [\bar{b}(x_i)/b_r(x_i)]^{w^k}
 \end{aligned}$$

In practice, we usually match on the variables present in all mini-buckets  $k \in Q_i$ , e.g.,  $\cap_{k \in Q_i} \mathbf{x}_k$ , rather than just  $x_i$ ; this gives a tighter bound for the same amount of computation.

### ■ 3.2.3 Adding new regions

New regions are added to the initial join tree after one or more rounds of iterative optimization. To bound the complexity of the search over clusters, we restrict our attention to pairs of mini-buckets to merge within each bucket and use the elimination order  $o$  to guide our search, processing each bucket  $B_i$  one at a time.

Given bucket  $B_i$  and a current partitioning  $Q_i = \{q_i^1, \dots, q_i^k\}$ , we score the merge for each allowed pair of mini-buckets  $(q_i^m, q_i^n)$ , e.g., those with  $|\text{var}(q_i^m) \cup \text{var}(q_i^n)| \leq \text{ibound} + 1$ , using an estimate of the benefit to the bound that may arise from merging the pair:

$$S(q_i^m, q_i^n) = \max_{\mathbf{x}} \log [\lambda_{m \rightarrow \pi_m}(\mathbf{x}_{\pi_m}) \times \lambda_{n \rightarrow \pi_n}(\mathbf{x}_{\pi_n}) \div \lambda_{r \rightarrow \pi_r}(\mathbf{x}_r)] \quad (3.7)$$

This score can be justified as a lower bound on the decrease in the approximation to  $\log Z$ , since it corresponds to adding region  $\pi_r$  with weight  $w^{\pi_r} = 0$ , while reparameterizing the parents  $\pi_m, \pi_n$  to preserve their previous beliefs. This procedure leaves the bound unchanged

except for the contribution of  $\pi_r$ ; eliminating with  $w^{\pi_r} = 0$  is equivalent to the max operation. For convenience, we define  $S(q_i^m, q_i^n) < 0$  for pairs which violate the *ibound* constraint. Then, having computed the score between all pairs, we choose the pair with maximum score to be merged into a new clique. In Algorithm 3.1, the function  $\text{SelectMerge}(\cdot)$  denotes this scoring and selection process.

### ■ 3.2.4 Updating graph structure

Having found which mini-buckets to merge, we update the join graph to include the new clique  $r = q_i^m \cup q_i^n$ . Our goal is to add the new region in such a way that it affects the scope of the existing regions in the join tree as little as possible. Adding the new clique is done in two steps.

First, we initialize a new mini-bucket in  $B_i$  with its scope matching,  $\text{var}(r)$ , the scope of the new merged region  $r$ . Eliminating variable  $x_i$  from this new mini-bucket results in the message  $\lambda_{r \rightarrow \pi_r}$ . The earliest argument of  $\lambda_{r \rightarrow \pi_r}$  in the elimination order determines the bucket  $B_j$  containing mini-buckets that can potentially be the parent,  $\pi_r$ , of the new region. To find  $\pi_r$  in  $B_j$  we seek a mini-bucket  $q_j^k$  that can contain  $r$ , i.e.,  $\text{var}(\lambda_{r \rightarrow \pi_r}) \subseteq \text{var}(q_j^k)$ . If such a mini-bucket exists, we set  $\pi_r$  to  $q_j^k$ ; otherwise, we create a new mini-bucket  $q_j^{|Q_j|+1}$  and add it to  $Q_j$ , with a scope that matches  $\text{var}(\lambda_{r \rightarrow \pi_r})$ . The same procedure is repeated after eliminating  $x_j$  from  $q_j^{|Q_j|+1}$  until we either find a mini-bucket already in the join tree that can serve as the parent, or  $\text{var}(\lambda_{r \rightarrow \pi_r}) = \emptyset$  in which case the newly added mini-bucket is a root. Algorithm 3.2 describes these initial structural modifications.

Having added the new regions, we then try to remove any unnecessary mini-buckets, and update both the join tree and the function values of the newly added regions to ensure that the bound is improved. To this end, we update every new mini-bucket  $r$  that was added to the join tree in the previous step as follows. For mini-bucket  $r \in Q_i$ , we first find any

---

**Algorithm 3.2** AddRegions: find regions to add for merge

---

**Input:** The join graph  $wmb$ , elimination order  $o$ , and mini-buckets  $q_i^m$  and  $q_i^n$  to be merged

**Output:** a list of newly added mini-buckets  $\mathcal{R}$

**Initialize** new region  $q^r$  with  $\text{var}(q^r) = \text{var}(q_i^m \cup q_i^n)$  and add it to  $Q_i$

**repeat**

**Update**  $\mathcal{R} = \mathcal{R} \cup q^r$

**Set** new clique  $C = \text{var}(q^r) \setminus x_i$

**if**  $C = \emptyset$  **then**

$done \leftarrow True$

**else**

**Find**  $B_j$  corresponding to the first un-eliminated variable in  $C$  based on elimination order  $o$

**for** each mini-bucket region  $q_j^k \in Q_j$  **do**

**if**  $C \subseteq \text{var}(q_j^k)$  **then**

                // forward message fits in existing mini-bucket:

$done \leftarrow True$

**end if**

**end for**

**end if**

**if** not  $done$  **then**

        // Create a new region to contain forward message:

**Initialize** new region  $q^r$  with  $\text{var}(q^r) = C$  and add it to  $Q_j$

**end if**

**until** done

---

mini-buckets  $s \in Q_i$  that can be subsumed by  $r$ , i.e.,  $\text{var}(s) \subseteq \text{var}(r)$ . For each of these mini-buckets  $s$ , we connect all of  $s$ 's children (mini-buckets  $t$  such that  $\pi_t = s$ ) to  $r$ , e.g., set  $\pi_t = r$ . We also merge the factors associated with  $r$  and  $s$ , so that  $f_r \leftarrow f_r \times f_s$ .

Next, we reparameterize several other functions in the join graph in order to preserve or improve the current bound value. Specifically, removing  $s$  changes the incoming, forward messages to its parent,  $\pi_s = \text{pa}(s)$ , which changes the bound. By reparameterizing the factor at  $\pi_s$ ,

$$f_{\pi_s} \leftarrow f_{\pi_s} \times \lambda_{s \rightarrow \pi_s} \qquad f_{\pi_r} \leftarrow f_{\pi_r} \div \lambda_{s \rightarrow \pi_s}$$

we keep the overall distribution unchanged, but ensure that the bound is strictly decreased.

Finally we remove  $s$  from  $Q_i$ , completing the merge of mini-buckets  $s$  and  $r$ . This process is given in Algorithm 3.3 and depicted in Figure 3.2 for a small portion of join-graph.

Every merge decision is followed by one or more iterations of message passing, followed by rescoreing the mini-buckets in  $B_i$ . The process of message passing and merging continues until no more mini-buckets of  $B_i$  can be merged while still satisfying the bounding parameter *ibound*.

Continuing along the elimination order, the same procedure is repeated for the mini-buckets in each bucket  $B_i$ , and the final upper bound to the partition function is computed using Eq. (3.6).

**Computational Complexity of Incremental region selection for WMBE.** Rollon and Dechter [2010] provide an analysis of the computational complexity of their content-based partitioning heuristic. Following the same framework, next we analyze the computational complexity of adding new regions in our hybrid approach.

**Proposition 3.1.** *The computational complexity of merging all pair of regions  $(q^m, q^n)$  in Algorithm 3.1, is at most  $O( R \cdot L^2 \cdot \exp(z) + R \cdot L \cdot z \cdot \exp(z) + T \cdot R^2 \cdot \exp(z) )$ , where  $L$  is the maximum number of initial regions in a bucket (which can be bounded by the largest degree of any variable),  $z$  is the *ibound* and  $R = \sum_{\alpha} |\alpha|$  bounds the number of possible regions in the cluster graph.*

*Proof.* Let  $R$  be as defined, and denote by  $O( S )$  the computational complexity of a single merge. There cannot be more than  $R$  merges, which means that the complexity of all possible merges is  $O(R \cdot S)$ . The complexity of a single merge can then be bounded as  $O( S ) = O( L^2 \cdot \exp(z) + L \cdot z \cdot \exp(z) + T \cdot R \cdot \exp(z) )$ . The first term in  $O( S )$  captures the complexity of computing the score, which is  $O(\exp(z))$ , for any pair of regions. The algorithm computes the score for every pair of regions in a bucket, which yields  $O( L^2 \cdot \exp(z) )$  for score computation.

---

**Algorithm 3.3** MergeRegions: merge and parameterize newly added regions to improve bound

---

**Input:** The join graph  $wmb$  and a list of newly added mini-buckets  $\mathcal{R}$

**for all**  $r \in \mathcal{R}$  **do**

**Initialize** new region  $r$  in  $B_i$  with  $f_r(x_r) = 1$

**Find** regions  $\{s \mid s \in Q_i \ \& \ \text{var}(s) \subseteq \text{var}(r)\}$

*// Remove / merge contained regions  $s$ :*

**for all** found regions  $s$  **do**

        Connect all children of  $s$  to  $r$

$f_r = f_r \times f_s$    *// merge factors and*

*// preserve belief at parent  $\pi_s$ :*

$f_{\pi_s} = f_{\pi_s} \times \lambda_{s \rightarrow \pi_s}$

$f_{\pi_r} = f_{\pi_r} \div \lambda_{s \rightarrow \pi_s}$

        Remove  $s$  from  $Q_i$

**end for**

**end for**

---

The second term in  $O(S)$  is the complexity bound of the merge process. For each merge we have to add the new merged region and all its parents, which is bounded by  $z$ . After adding these new regions, we need to check for subsumptions of old regions of the cluster graphs with the new ones which involves  $z \cdot L$  tests. If one region can be subsumed by the other, we need to update the messages sent to them by their children which requires  $O(\exp(z))$  computations for each update. Putting it all together, the computational complexity of the merge process is bounded by  $O(L \cdot z \cdot \exp(z))$ . The third component accounts for the complexity of  $T$  iterations of messages passing after each merge, which is bounded by  $O(T \cdot R \cdot \exp(z))$  when we have at most  $R$  regions in the cluster graph. Having the complexity of a single merge as  $O(S) = O(L^2 \cdot \exp(z) + L \cdot z \cdot \exp(z) + T \cdot R \cdot \exp(z))$ , we can then bound the complexity of doing all merges as  $O(R \cdot S) = O(R \cdot L^2 \cdot \exp(z) + R \cdot L \cdot z \cdot \exp(z) + T \cdot R^2 \cdot \exp(z))$ .  $\square$

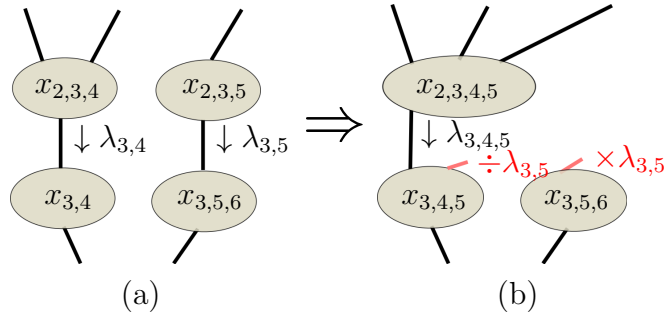


Figure 3.2: Merge and post-merge reparameterization operations. (a) A portion of a join-graph corresponding to the elimination of  $x_2$  and  $x_3$ , each with two mini-buckets. (b) Merging cliques  $(2, 3, 4)$  and  $(2, 3, 5)$  produces a new clique  $(3, 4, 5)$ , which subsumes and removes clique  $(3, 4)$ . Having removed parent  $(2, 3, 5)$ , we reparameterize the new clique functions by the original message  $\lambda_{3,5}$  (red) to preserve the original belief at  $(3, 5, 6)$  and ensure that the bound is tightened. See text for more detail.

### ■ 3.3 Discussion

Our method is similar to content-based mini-buckets, with the main difference being that message passing performed on the simpler graph is used to reparameterize the functions before the merge scores are computed.

Our method can also be viewed as a cluster pursuit approach, in which we restrict the clusters considered, to unions of the current minibuckets at the earliest bucket  $B_i$ , and merge up to our computational limit before moving on to later buckets. These restrictions serve to reduce the number of clusters considered, but in addition, appear to lead to better regions than a purely greedy region choice – in the experiments (Section 3.4), we compare our approach to a more “cluster pursuit-like” method, in which pairs of regions in *any* bucket  $B_i$  are considered and scored. Perhaps surprisingly, we find that this greedy approach actually gives significantly worse regions overall, suggesting that processing the buckets in order can help by avoiding creating unnecessary regions.

Finally, our method is also closely related to RCR [Choi and Darwiche, 2010]. From this

perspective, we “relax” to a low-*bound* minibucket, “compensate” by variational message passing, and “recover” by selecting regions that will tighten the variational bound defined by the join graph. Compared to RCR, we find a number of differences in our approach: (1) RCR selects constraints to recover anywhere in the graph, similar to a greedy cluster pursuit; as noted, this appears to work significantly less well than an ordered recovery process. (2) RCR makes its recovery updates to the relaxed graph, then (re)builds a (new) join tree over the relaxed graph; in contrast, we incrementally alter the join graph directly, which avoids starting from scratch after each merge. (3) Our method is solidly grounded in the theory of variational bounds and message passing, ensuring that both the message passing and region merging steps are explicitly tightening the same bound. From this perspective, for example, it becomes clear that RCR’s “residual recovery” heuristic is unlikely to be effective, since after message passing, the reparameterization updates should ensure that all mini-buckets containing a variable  $x_i$  will match on their marginal beliefs. In other words, residual recovery is making its structure (region) choices using a criterion that actually measures mismatches that can be resolved by message passing.

### ■ 3.4 Empirical Evaluation

To show our method’s effectiveness compared to previous region selection strategies for MBE, we tested our incremental approach on a number of real world problems drawn from past UAI approximate inference challenges, including linkage analysis, protein side chain prediction, and segmentation problems. We compare our hybrid region selection method against the scope-based heuristic of Dechter and Rish [1997] and the content-based heuristic of Rollon and Dechter [2010].

Table 3.1: **UAI Segmentation Instances.** Different columns show the bound achieved using each partitioning heuristic, where “Scp”, “Cont” and “Hyb” represent the naïve scope based partitioning for MBE [Dechter and Rish, 1997], the context (or energy) based heuristic of Rollon and Dechter [2010] and our hybrid approach interleaving iterative optimization with partitioning, respectively. In all but one case, our proposed construction provides tighter bounds.

Instance	<i>ibound</i> = 5			<i>ibound</i> = 10		
	Scp	Cont	Hyb	Scp	Ctxt	Hyb
2-17-s	-31.3197	-33.4840	<b>-49.5670</b>	-38.9801	-42.1524	<b>-52.507</b>
2-17-s- <i>opt</i>	-46.9314	-45.4286	<b>-49.6432</b>	-48.661	-48.4306	<b>-52.5633</b>
8-18-s	-54.9884	-60.9899	<b>-85.6518</b>	-72.3045	-72.284	<b>-87.2385</b>
8-18-s- <i>opt</i>	-80.6527	-81.1391	<b>-85.6694</b>	-83.0398	-79.0921	<b>-87.2556</b>
9-24-s	-51.2897	-49.3903	<b>-55.6046</b>	-54.9325	-55.0151	<b>-55.615</b>
9-24-s- <i>opt</i>	<b>-56.0513</b>	-53.7852	-55.6241	-54.9325	-55.0151	<b>-55.615</b>
17-4-s	-58.7953	-59.0758	<b>-81.5415</b>	-80.267	-79.3323	<b>-85.3712</b>
17-4-s- <i>opt</i>	-71.7959	-76.8213	<b>-81.6079</b>	-83.2573	-82.9646	<b>-85.3865</b>
7-11-s	-59.8250	-57.2773	<b>-72.7178</b>	-71.1296	-70.1542	<b>-75.2869</b>
7-11-s- <i>opt</i>	-70.7037	-68.8255	<b>-72.9556</b>	-74.6424	-73.9855	<b>-75.2905</b>

## Experimental Setup

For each set of experiments, we initialize a join tree using WMB elimination with  $ibound = 1$ . We use an elimination ordering found using the min-fill heuristic [Dechter, 2003] and set the weights uniformly in each bucket. As a result, each mini-bucket  $q_i^k$  contains a single factor  $f_\alpha$  as described in section 3.2.1.

From this initial setup, we then use Algorithm 3.1 to merge mini-buckets incrementally and compute the upper bound as in Eq. (3.6).

**Segmentation.** To evaluate the different methods on pairwise binary problems we used a set of segmentation models from the UAI08 approximate inference challenge. These models have  $\approx 230$  binary variables and  $\approx 850$  factors. We used varying  $ibounds$  for comparison

Table 3.2: **UAI Pedigree Instances.** Different columns show the bound achieved using each partitioning heuristic; again, “Scp”, “Cont” and “Hyb” are scope based partitioning [Dechter and Rish, 1997], the content-based heuristic [Rollon and Dechter, 2010] and our proposed, hybrid approach. In all cases, our proposed construction provides stronger bounds, both before and after full optimization using message passing.

Instance	<i>ibound</i> = 5			<i>ibound</i> = 10		
	Scp	Cont	Hyb	Scp	Ctxt	Hyb
ped23	-67.8848	-69.9015	<b>-71.9677</b>	-75.6057	-78.4033	<b>-79.4649</b>
ped23- <i>opt</i>	-71.6951	-71.6988	<b>-72.0670</b>	-76.0531	-78.7646	<b>-79.4669</b>
ped20	-35.6986	-40.1787	<b>-44.7230</b>	-51.2648	-54.4136	<b>-57.6506</b>
ped20- <i>opt</i>	-42.3024	-42.8980	<b>-44.7501</b>	-52.6043	-56.2193	<b>-57.7841</b>
ped42	-41.6656	-43.5206	<b>-51.0000</b>	-55.0681	-57.5755	<b>-61.3504</b>
ped42- <i>opt</i>	-49.0089	-50.0585	<b>-51.1018</b>	-57.3718	-59.2170	<b>-61.3560</b>
ped38	-79.4742	-89.6906	<b>-92.7643</b>	-98.6339	-101.1178	<b>-113.6004</b>
ped38- <i>opt</i>	-83.0351	-91.8510	<b>-93.0615</b>	-101.0715	-104.1031	<b>-113.8926</b>
ped19	-58.9234	-63.2737	<b>-80.6488</b>	-90.7840	-93.9027	<b>-100.3230</b>
ped19- <i>opt</i>	-72.3311	-77.7023	<b>-80.7167</b>	-92.8916	-96.2846	<b>-100.3388</b>

and report the results on two values,  $ibound \in [5, 10]$ . Table 3.1 compares the upper bound on the log partition function for a representative subset of instances in this category, for two different computational limits,  $ibound = 5$  and  $ibound = 10$ . Different columns show the bound achieved using different partitioning heuristics:

- (1) **Scp** represents naïve scope-based partitioning;
- (2) **Cont** represents the energy based heuristic of Rollon and Dechter [2010]; and
- (3) **Hyb** represents our hybrid approach, interleaving iterative optimization with partitioning.

The results show clear improvement in the upper bound using our hybrid approach, indicating the effectiveness of iterative message passing and optimization in guiding region selection. To further study the effectiveness of the merged regions in the context of message passing and optimization, we then fully optimized the join-graphs generated by the three region

selection schemes using iterative message passing until convergence. The upper bounds after this optimization process are denoted by *inst-opt* for each problem instance, *inst*. As might be expected, this additional optimization step improves the bounds of the scope-based and content-based heuristics more dramatically than our hybrid method; however, even after full optimization of the bounds, we find that the hybrid method’s bounds remain better in all of the 6 instances except one, indicating that our method has identified fundamentally better regions than the previous approaches.

Of course any improvement in the bound from the content-based partitioning and our hybrid method comes with an additional computational overhead. While our implementations for these algorithms are in MATLAB and are not optimized for timing experiments, it is important to consider the effect of this computation on time. In each set of these experiments, computing the bound using content-based partitioning took on average 10 times more than scope-based partitioning. Our hybrid approach then requires one round of message passing after each merge and took on average 10 times more than the content-based to compute the upperbound.

**Linkage Analysis.** To compare the various methods on models with non-pairwise factors and higher cardinalities of variables, we studied pedigree models. The pedigree linkage analysis models from the UAI08 approximate inference challenge have  $\approx 300 - 1000$  variables, whose cardinalities vary in the range of  $[2, \dots, 5]$ ; the induced width of the models are typically  $\approx 20 - 30$ . We used varying *ibounds* for comparison and report the results on two values  $ibound \in [5, 10]$ .

Table 3.2 shows the upper bounds on a subset of pedigree problems, again showing the effectiveness of the hybrid method: we find that again, the hybrid method consistently outperforms the other two region selection approaches, and results in better fully optimized bounds in all of the 22 instances when  $ibound = 5$  and all but two cases when  $ibound = 10$ .

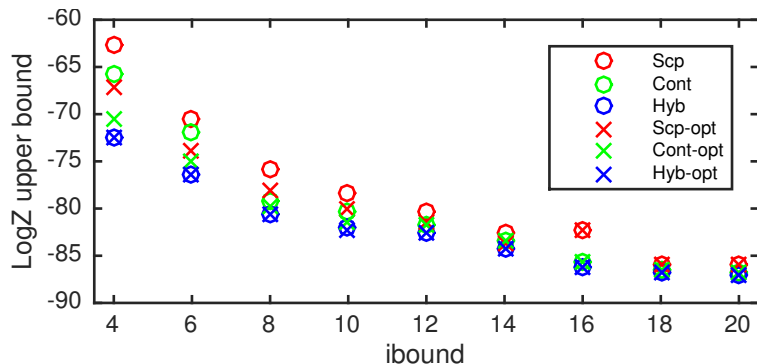


Figure 3.3: The upper bound achieved by the three partitioning heuristics for pedigree23 instance over *ibound* range between 4 to 20.

As for the segmentation instances, computing the upperbound using our hybrid approach is on average 10 times slower than using the content-based partitioning and using content-based partitioning is on average 10 times slower than using the scope-based heuristic.

**Effect of *ibound*.** We also studied the results of the three partitioning methods across a range of *ibounds*, to understand how our method’s effectiveness changes as a function of the allowed region sizes. Figure 3.3 shows the results for an instance of pedigree dataset. Empirically, our method is more effective on smaller *ibounds*, where there are a large number of possible merges and finding the best one results in a greater improvement to the upper bound. For larger *ibounds*, where the resulting cluster trees are relatively close to that of exact variable elimination, the upper bounds produced by all three heuristics are also fairly close.

**Protein Side-Chain Prediction.** Finally, to examine models over high-cardinality variables, we look at a subset of the protein side chain prediction models, originally from Yanover and Weiss [2003] and Yanover et al. [2006]. These models contain  $\approx 300 - 1000$  variables with cardinalities between 2 and 81, with pairwise potential functions. For these problems, we only ran our experiments using *ibound* = 2, due to the high number of states for each variable. Table 3.3 shows the results of the three partitioning methods, which again agrees with the

previous experiments: our hybrid method outperforms the other two in all 44 instances in the problem set, often dramatically, both before and after the bound is fully optimized.

For these set of instances where the variables have large cardinalities, computing the upper bound using our hybrid approach is on average 10 times slower than using the content-based partitioning and using content-based partitioning is on average 20 times slower than using the scope-based heuristic.

**Greedy vs. Elimination Order Based Merging.** As discussed before, we restrict the clusters considered for merges to unions of the current minibuckets at the earliest bucket  $B_i$ , and merge up to our computational limit before moving on to later buckets, which serves to reduce the number of clusters considered. We compare our choice of clusters with a purely greedy region choice in which pairs of regions in *any* bucket  $B_i$  are considered and scored.

Interestingly, the upper bounds achieved using the greedy approach were not better than the top-down merging based on elimination order. A possible reason for this behavior is that the top-down approach allows large regions generated by mini-buckets early in the elimination ordering to be processed by buckets later in the order; the greedy approach disrupts this flow and results in extra regions that cannot be merged with any other region while respecting the *ibound*.

Table 3.3: **Protein side-chain prediction.** Here we show results for only  $ibound = 2$ , due to the high number of states in each variable. Our method often produces dramatically better partitionings than scope- or content-based mini-bucket partitions.

<b>Instance</b>	<b>Scp</b>	<b>Cont</b>	<b>Hyb</b>
1crz	-242.2036	-284.865	<b>-451.598</b>
1crz <i>-opt</i>	-528.5348	-495.514	<b>-545.929</b>
2cav	71.2802	-26.0052	<b>-148.637</b>
2cav <i>-opt</i>	-156.5387	-240.289	<b>-272.606</b>
1kk1	89.5527	46.8216	<b>-121.723</b>
1kk1 <i>-opt</i>	-115.4737	-105.894	<b>-143.447</b>
1e4f	40.6686	-6.1785	<b>-190.943</b>
1e4f <i>-opt</i>	-212.4607	-202.547	<b>-240.27</b>
1ehg	71.3308	14.768	<b>-149.158</b>
1ehg <i>-opt</i>	-169.8435	-147.333	<b>-211.678</b>

Table 3.4: **Top-down vs. Greedy Merging.** We examine the effect of using a “fully greedy” merging procedure closer to standard cluster pursuit, in which we merge the best-scoring cluster in any bucket at each step. We find that following the top-down ordering actually results in significantly better bounds. Results shown are for  $ibound = 5$ .

<b>Instance</b>	<b>Top-Down</b>	<b>Greedy</b>
ped23	<b>-71.9677</b>	-67.9094
ped23 <i>-opt</i>	<b>-72.0670</b>	-67.9094
ped20	<b>-44.7230</b>	-38.0717
ped20 <i>-opt</i>	<b>-44.7501</b>	-38.0718
ped42	<b>-49.9955</b>	-37.8576
ped42 <i>-opt</i>	<b>-50.0469</b>	-37.8582
ped38	<b>-92.7643</b>	-79.9144
ped38 <i>-opt</i>	<b>-93.0615</b>	-79.9144
ped19	<b>-80.6488</b>	-48.6900
ped19 <i>-opt</i>	<b>-80.7167</b>	-48.6904

## ■ 3.5 Conclusion

We presented a new merging heuristic for (weighted) mini-bucket elimination that uses message passing optimization of the bound, and variational interpretations, in order to construct a better heuristic for selecting moderate to large regions in an intelligent, energy-based way. Additionally we showed how to incrementally update the join graph of mini-bucket elimination after new regions are added in order to avoid starting from scratch after each merge.

Our approach inherits the advantages of both cluster pursuit in variational inference, and (weighted) mini-bucket elimination perspectives to produce a tight bound. We validated our approach with experiments on a wide variety of problems drawn from a recent UAI approximate inference competition. In practice, we find that our methods work significantly better than either existing partitioning heuristics for mini-bucket [Rollon and Dechter, 2010], or a pure region pursuit approach. We expect this construction to improve our ability to search and solve large problems. However, our method does involve additional computational overhead compared to, say, scope-based constructions, in order to evaluate and make merge decisions. We did not focus here on any-time performance; a more nuanced balance of time, memory, and bound quality is one direction of potential future study.

# Improving Resource Usage in Mini-bucket Elimination

## ■ 4.1 Introduction

In Chapter 3, we presented an incremental form of mini-bucket elimination (MBE) that proposed new heuristics for selecting the regions, or mini-buckets, used in the approximation. As with standard mini-bucket, we used a single control parameter, the *ibound*, to manage the computational (time) and representational (memory) complexity of the approximation by limiting the number of variables in each region. One of the main reasons MBE is so successful is the easy tradeoff between memory and quality: Using more memory (larger regions) almost always provides a significant improvement in quality, especially on summation problems (e.g., computing the log partition function); see Liu and Ihler [2011].

However, when applying MBE to real-world problems, we need to run it on a wide array of problem types with very different properties, in terms of graph structures and cardinalities of variables. Consider an example of protein side-chain prediction, where the model is dense and pairwise, on perhaps 350 variables with cardinalities that vary between 2 and 81; choosing an *ibound* = 4 for MBE with scope-based partitioning requires 17 GB of memory. In contrast,

for a linkage analysis (pedigree) instance, the graph is relatively sparse, and the 448 variables have cardinalities that vary between 2 and 5, and we can compute an approximation with  $i_{bound} = 16$  and using only 1 GB of memory. As a result, to be able to apply MBE to these different problems, we need robust methods that can find good approximations in the face of this kind of diversity.

A central challenge is then to automatically and quickly build approximations that satisfy the resource constraints, and in particular the amount of memory available. As we discuss in Section 4.2, finding the largest  $i_{bound}$  that respects the memory limits is easy and efficient for scope-based heuristics. However, for content-based heuristics, the regions are selected based on the values of the functions and the incoming messages, which means that the exact memory requirement of a particular execution is not easily evaluated without fully computing the bound. If the resulting approximation uses too much memory, this work is wasted; if we have memory left over, we may need to repeat the effort. For this reason, it would be useful to be able to estimate and control the amount of memory used by an approximation on the fly, as it is built.

In this chapter we first review the state of the art and discuss the challenges of estimating the memory needs of content-based MBE. Then we propose two improvements that allow us to better control how limited memory resources are used, and as a result often provide a better bound on the partition function. Our goal is to more fully utilize the available memory resources, while also allowing us to also use content-based selection methods that typically give tighter bounds. We evaluate our framework on various real world problems and compare different memory management strategies within content-based mini-bucket approximation.

## ■ 4.2 State of the Art

In this section, we study the role that limited memory resources play during the design of a mini-bucket based inference approximation and how current systems address the memory allocation problem. Memory limits are a common and important form of constraint placed on approximations; typically, the more memory is available, the better the approximation quality.

One advantage of scope-based mini-bucket elimination is that it is relatively easy to design an approximation that can fit in the available memory. For scope-based partitions, the elimination process can be simulated very quickly for various *ibound* values. Using the scopes of the intermediate functions, but without actually computing any of the functions themselves, one can quickly compute the used memory, and select the largest *ibound* that complies with the memory constraints; see Otten et al. [2012] for an example. Also, since the amount of memory grows exponentially as *ibound* is increased, it is easy to search over different values of *ibound* as the range of values in the search space is relatively small. However, using the scope-based simulation with a particular *ibound* value to estimate the memory required by an approximation has two main disadvantages, discussed next.

### ■ 4.2.1 Content-based region choices

For content-based mini-bucket, ensuring bounded memory use by searching over *ibound* is not as easy as scope-based mini-bucket elimination. In particular, the partitioning process depends on the values of the functions and incoming messages, which means that it cannot be simulated without fully computing the functions and messages.

From this perspective, an incremental approach (e.g., the method described in Chapter 3) is a useful paradigm. However several issues should be addressed. For example, in Chapter

3 we showed that it was best to do all merges within a single bucket before moving on to a later bucket. However, allowing all the memory to be used by the earlier buckets in the elimination order is unlikely to be an effective use of the available memory. This suggests that some “forecasting” of memory use in future buckets is probably useful.

A possibly naïve way to estimate the amount of memory required by the content-based partitioning process is to compute the amount of memory required by the scope-based mini-bucket and use it as an estimate for the memory used by the content-based process. However for models with even moderate variation in the domain size of the variables, different partitioning choices can result in very different memory use. Then, when processing for a given *ibound*, if the available memory is exceeded the procedure will not provide a usable approximation; if it uses less memory than is available, it may result in a poor quality one.

Figure 4.1 (a) shows an example of this difference in memory use for protein side-chain prediction instances, where WMBE is used to compute an upper bound to the log partition function with *ibound* = 2, using scope- and content-based heuristics. The memory required by the content-based approximation is compared to the memory required by the scope-based approximation by computing the difference,  $(M(Ctnt) - M(Scp)) / M(Scp)$ . Here  $M(Ctnt)$  is the memory required by the content-based partitioning and  $M(Scp)$  is the memory required by scope-based partitioning. For some instances, the content-based partitioning required more memory than scope-based, while for the others, the relationship is reversed. This property makes it difficult to use the scope-based *ibound* value, and its associated memory use, as an accurate estimate for a content-based partitioning. Hence we would like an “online”, memory-aware partitioning procedure that can estimate and manage the amount of memory it requires while building the approximation.

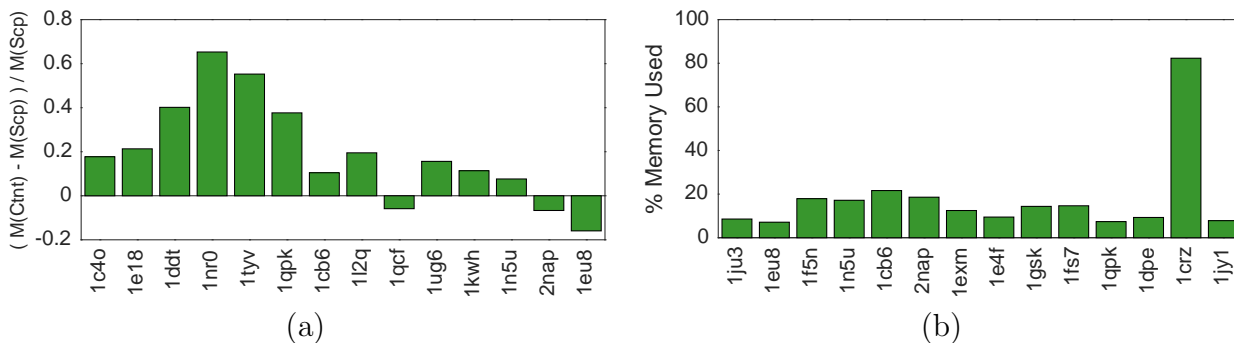


Figure 4.1: (a) Memory used by content-based partitioning, compared to the memory used by scope-based partitioning in WMBE. Different partitioning decisions can lead to significantly varying memory requirements. We show the relative difference in memory use,  $\frac{M(Cnt) - M(Scp)}{M(Scp)}$ , across several problem instances, where  $M(Cnt)$  is the memory required by weighted mini-bucket elimination using content-based partitioning and  $M(Scp)$  is the memory required by scope-based partitioning. Positive values indicate the content-based partitioning required more memory. (b) Percentage of memory used by scope-based mini-bucket elimination when the *ibound* parameter is set to the largest value for which the mini-bucket cluster tree fits in memory, across several protein examples. For many instances, a large fraction of the available memory is left unused.

## ■ 4.2.2 Inefficient memory allocation

Another aspect of estimating the required memory for any mini-bucket approximation is the relationship between *ibound* and the memory required by the resulting mini-bucket approximation. Standard practice finds an *ibound* such that MBE using *ibound*+1 does not fit in the memory. However, the *ibound* is a very coarse mechanism for controlling how much memory is used, as any change in the *ibound* changes the amount of required memory exponentially. This exponential growth makes searching over *ibound* values efficient, but also means that the approximation built with the largest *ibound* may use only a fraction of the available memory. This effect is more pronounced in models with larger domain sizes of the variables. Figure 4.1 (b) shows this effect for instances of protein side-chain prediction problems. Each bar shows the percent of memory used by scope-based mini-bucket elimination when *ibound* is set to the largest value for which the cluster tree fits in memory. For many instances, a large fraction of the available memory is unused. This effect argues for a more fine-grained approach, or possibly a coarse-to-fine memory allocation procedure. Again, forecasting the

memory requirements for an approximation as new regions are added to the approximation would be useful, as some buckets could use higher *ibound* values than others if we can afford it.

To this end, we propose a set of *budget-based* memory allocation strategies that allow finer control over how the available memory is used. We frame these approaches in terms of the basic incremental cluster tree construction proposed in Chapter 3, then propose and study empirically several mechanisms for setting the initial budget values, and updating the budgets as the cluster tree is built.

### ■ 4.3 Memory awareness

A key component of our framework is the incremental perspective to adding new regions; to this end we will rephrase existing elimination based methods within our incremental viewpoint. First, we define some of the the notation that we will require throughout the chapter.

Let  $wmb$  be a valid mini-bucket cluster tree for a graphical model consisting of variables  $X = \{X_1, \dots, X_n\}$  and factors  $F = \{f_\alpha(X_\alpha)\}$ , indexed by their scopes  $\alpha \subseteq \{1, \dots, n\}$ , with respect to ordering  $\mathbf{o} = [X_1, \dots, X_n]$ . Then,  $wmb$  consists of a set of regions (or mini-buckets)  $\mathcal{R}$  organized into buckets  $B_1, \dots, B_n$ , with the properties:

1. Each factor  $f_\alpha$  can be assigned to a region that contains its scope:

$$\forall \alpha, \exists r \in \mathcal{R} \text{ s.t. } X_\alpha \subseteq \text{var}(r)$$

where  $\text{var}(r)$  is the set of variables in region  $r$ .

2. Each region  $r \in \mathcal{R}$  is associated with the bucket corresponding to its earliest eliminated

variable,

$$r \in B_i \text{ iff } X_i \in \text{var}(r) \text{ and } \forall X_j \in \text{var}(r), j \geq i$$

where  $j \geq i$  means that  $X_j$  is eliminated later than  $X_i$  in ordering  $\mathbf{o}$ .

3. Each region  $r$  with more than one variable is identified with a parent region  $\pi_r$  to which its downward message (eliminating  $X_i$ ) can be assigned:

$$\text{if } |\text{var}(r)| > 1 \text{ and } r \in B_i, \exists \pi_r \in \mathcal{R} \text{ with } \text{var}(r) \setminus X_i \subseteq \text{var}(\pi_r)$$

Now, we define several useful functions for measuring the computational resources associated with the cluster tree  $wmb$ , bucket  $B_i$ , or region  $r$ :

$$\text{size}(r) = |\text{var}(r)| \quad : \text{ the number of variables in region } r$$

$$\text{size}(B_i) = \max_{r \in B_i} \text{size}(r) \quad : \text{ the size of the largest region in } B_i$$

$$\text{size}(wmb) = \max_i \text{size}(B_i) \quad : \text{ the size of the largest region in } \mathcal{R}$$

$$M(r) = \prod_{X_i \in \text{var}(r)} |\mathcal{X}_i| \quad : \text{ the number of entries in a table over the variables in } r$$

$$M(B_i) = \sum_{r \in B_i} M(r) \quad : \text{ the total number of entries in the tables for regions in bucket } i$$

$$M(wmb) = \sum_i M(B_i) \quad : \text{ the total number of entries in all tables for the regions in } \mathcal{R}$$

Then, “ $\text{size}(\cdot)$ ” corresponds to measuring the inherent complexity or memory requirements of a region, bucket, or cluster tree according to its scope (as is typical for standard mini-bucket formulations), while “ $M(\cdot)$ ” corresponds to measuring its complexity in terms of the amount of memory required to represent an arbitrary function over those variables. We also define the total memory budget (the memory available for inference) as  $\mathcal{M}$ .

Two key steps of the incremental perspective are: (1) forming the initial cluster tree, and

(2) merging new regions and updating the graph structure.

**Initializing the cluster tree.** Given the factor graph  $\mathcal{G}$ , our incremental approach starts from an “initial” cluster tree, where each factor is a region, with descendants corresponding to partial eliminations of that factor. We construct this initial cluster tree,  $wmb$ , using a min-fill elimination ordering  $\mathbf{o}$  and  $ibound = 1$ . For any given bucket  $B_i$ , this results in each mini-bucket (region)  $r \in B_i$  containing a single factor. We represent the result of the elimination as a message,  $\lambda_{r \rightarrow \pi_r}$ , with scope  $\text{var}(\lambda_{r \rightarrow \pi_r})$ , from region  $r$  to its parent  $\pi_r$  in a later bucket  $B_j$ , corresponding to its first-eliminated argument in  $\mathbf{o}$ . The parent region  $\pi_r$  may be one of the initial mini-buckets in  $B_j$ , if a region in  $B_j$  with sufficient scope already exists, or be a new mini-bucket. In our implementation we choose  $\pi_r \in B_j$  to be the mini-bucket over the most variables, such that  $\text{var}(\lambda_{r \rightarrow \pi_r}) \subseteq \text{var}(\pi_r)$ . To enable memory awareness we also evaluate  $M(r)$  and  $M(B_i)$  for every region  $r$  and bucket  $B_i$  in the initial cluster tree to keep track of the memory used.

**Merge Operation.** Algorithm 4.1 is used to score pairs of regions in each bucket and to select the best region to be added to the current cluster tree based on a predefined scoring function. Having selected which two regions to merge, we can then use the subroutines `AddRegion(.)` (Algorithm 3.2) and `MergeRegions(.)` (Algorithm 3.3) defined in Chapter 3 to add the new region to the cluster tree. However, to be memory-aware, we need to check if the updated cluster tree after the merge still respects the memory limit  $\mathcal{M}$ . To do so, `SelectMerge(.)` simulates the process of adding the newly merged region and computes the additional memory required by the merge. This additional memory accounts for adding the new regions to the cluster tree  $wmb$  and updating its structure. To do so, Algorithm 4.1 first simulates the merge process using Algorithm 4.6 and scores pairs of mini-buckets only if the resulting cluster tree can still fit in the available memory.

To simulate the process of merging regions  $r_j$  and  $r_k$  in bucket  $B_i$  and adding the new region

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**Algorithm 4.1** SelectMerge: Scoring all possible merges in a bucket while taking into account the total memory  $\mathcal{M}$

---

**Input:** cluster tree  $wmb$ , bucket  $B_i$ , elimination order  $o$ , total memory available  $\mathcal{M}$   
**for** every pair  $(r_j, r_k)$  in  $B_i$  such that  $|\text{var}(r_j) \cup \text{var}(r_k)| < ibound + 1$  **do**  
    *// check if merge respects the memory budget :*  
    Let  $C = \text{var}(r_j) \cup \text{var}(r_k)$  be a clique containing all variables from  $r_j$  and  $r_k$   
     $M_{new} \leftarrow \text{CheckMem}(wmb, o, C, \mathcal{M})$   
    **if**  $M_{new} < \mathcal{M}$  **then**  
         $S(j, k) \leftarrow \text{score}(r_j, r_k)$   
    **else**  
         $S(j, k) \leftarrow -\text{inf}$   
    **end if**  
**end for**  
    *// select the best merge:*  
     $(j^*, k^*) = \arg \max_{j, k} S(j, k)$   
**Return:**  $(r_{j^*}, r_{k^*})$

---

$r_n$ , Algorithm 4.6 uses the clique  $C = \text{var}(r_j) \cup \text{var}(r_k)$ , which contains the variables in the two regions  $r_j$  and  $r_k$ . The newly merged region  $r_n$  then has  $C$  as its scope. The memory required by bucket  $B_i$  after the merge,  $M_{new}(B_i)$ , is updated to account for the new region. To do so, the memory required by the new region,  $M(r_n)$ , is added to the current amount of memory used by  $B_i$ ,  $M(B_i)$ . The process also checks for any other regions in bucket  $B_i$  that can be subsumed by the new region  $r_n$  during the merge, and adds the memory used by those to  $M_{old}$ . To account for the memory that is going to be available when these regions are subsumed by  $r_n$ ,  $M_{old}$  is then subtracted from  $M_{new}$ . The total memory required by the cluster tree after adding this new region to bucket  $B_i$  is then updated in  $M_{used}$  and is compared with the total available memory  $\mathcal{M}$ . If the new region can be added, we then need to account for the memory required for updating the cluster tree structure. Eliminating variable  $X_i$  from the new region  $r_n$  results in a message  $\lambda_{r \rightarrow \pi_r}$  which has  $C = C \setminus X_i$  as its scope. This message is sent to the bucket  $B_m$  corresponding to the first variable of  $C$  in the elimination order. If  $B_m$  has any region,  $r_k$ , that can subsume the message ( e.g.  $C \subseteq \text{var}(r_k)$  ), then no new regions need to be added to the cluster tree. If it can not be subsumed, then a new region  $r_n$  should be added to the cluster tree and the memory required for it should

---

**Algorithm 4.2 WMBE-IB:** Memory-aware Incremental region selection for WMBE using  $ibound$

---

**Input:** factor graph ( $\mathcal{G}$ ),  $ibound$ , and the available memory  $\mathcal{M}$   
**Initialize**  $wmb$  to a cluster tree using e.g. a min-fill ordering  $o$ , uniform weights  
*// pass forward messages along the elimination order*  
 $wmb \leftarrow \text{msgForward}(wmb)$   
**for** each bucket  $B_i$  following the elimination order **do**  
  **repeat**  
     $(r_m, r_n) \leftarrow \text{SelectMerge}(B_i, ibound, \mathcal{M})$   
     $\mathcal{R} \leftarrow \text{AddRegions}(wmb, o, r_m, r_n)$   
     $wmb \leftarrow \text{MergeRegions}(wmb, \mathcal{R})$   
    *// pass forward messages along the merge path  $\mathcal{R}$  until we reach a root*  
    **set**  $r$  and  $\pi_r$  to the first and second regions in  $\mathcal{R}$  ( $\pi_r$  is parent of  $r$  in cluster tree)  
    **while**  $\pi_r \neq \emptyset$  **do**  
      *// pass forward message*  
       $\lambda_{r \rightarrow \pi_r}(\mathbf{x}_{\pi_r}) = \left[ \sum_{\mathbf{x}_r \setminus \mathbf{x}_{\pi_r}} [f_r(\mathbf{x}_r) \prod_{t: \pi_r = \pi_t} \lambda_{t \rightarrow \pi_r}(\mathbf{x}_{\pi_r})] \right]^{\frac{1}{w^{\pi_r}}}$   
      **end while**  
    **until** no more merges possible  
  **end for**  
**Return:** Upper bound on  $Z$ :  $\hat{Z} = \prod_{r: \pi_r = \emptyset} \lambda_{r \rightarrow \emptyset}$

---

be added to  $M_{used}$ . This process continues until either the message can be subsumed by a region that is already in the cluster tree or  $C \setminus X_m$  is empty which means that the previously added region has no parents. Algorithm 4.6 shows details of this process and returns the total memory required to add the new region  $r_n$  to the current cluster tree  $wmb$ .

### ■ 4.3.1 Baseline Methods

Having a scoring and merge process that takes into account the memory constraints, we can now give the basic weighted mini-bucket algorithms in our incremental, memory-aware framework, detailed in Algorithm 4.2. The scoring function to choose the pair of regions to add then depends on the partitioning heuristic. For scope based partitioning, we can define,

$$S(r_m, r_n) = |\text{var}(r_m) \cup \text{var}(r_n)|,$$

that computes the number of variables in the resulting merged region and for content based partitioning we can use the scoring function (3.7), defined in Chapter 3.

**Computational Complexity of WMBE-IB.** As in Rollon and Dechter [2010] next we analyze the computational complexity of adding new regions using our incremental memory-aware approach.

**Proposition 4.1.** *The computational complexity of merging all mergable pairs of regions  $(r_m, r_n)$  in WMBE-IB (Algorithm 4.2), is at most  $O( R \cdot L^2 \cdot \exp(z) + R \cdot L^3 \cdot z + R \cdot L \cdot z \cdot \exp(z) + R \cdot n \cdot \exp(z) )$ , where  $L$  is the maximum number of initial regions in a bucket (which can be bounded by the maximum degree of any variable),  $z$  is the ibound,  $n$  is the number of variables in the graphical model and  $R = \sum_{\alpha} |\alpha|$  upper bounds the maximum possible regions in the cluster graph.*

*Proof.* Let  $R$  be as defined, and denote by  $O( S )$  the computational complexity of a single merge. We cannot make more than  $R$  merges, which means that the complexity of all possible merges is  $O(R \cdot S)$ . The complexity of a single merge can then be bounded as  $O( S ) = O( L^2 \cdot \exp(z) + L^3 \cdot z + L \cdot z \cdot \exp(z) + n \cdot \exp(z) )$ . The first term in  $O( S )$  captures the complexity of computing the score, which is  $O(\exp(z))$ , for any pair of regions. The algorithm computes the score for every pair of regions in a bucket, which yields  $O( L^2 \cdot \exp(z) )$  for score computation. The second term in  $O( S )$  is the complexity of `CheckMem(.)` that simulates the merge process to decide if a merge fits in memory. For each merge we have to add the new merged region and all its parents, which is bounded by  $z$ . After adding these new regions, we need to check for subsumptions of old regions of the cluster graphs with the new ones which involves  $z \cdot L$  tests. Checking for the required memory using `CheckMem(.)` should be done for all merges in the bucket, and results in total computational complexity of  $O( L^3 \cdot z )$ . The third term then computes the bound for updating the graph structure for the pair that is selected to be merged. As before the complexity of

adding the new merged region and its parents is bounded by  $z$ , with  $z \cdot L$  possible merges between the old regions and the new ones. For every such merge, if one region can be subsumed by the other, we need to update the messages sent to them by their children which requires  $O(\exp(z))$  computations for each update. Putting it all together, the computational complexity of the merge process is bounded by  $O(L \cdot z \cdot \exp(z))$ . Having updated the cluster graph structure, the complexity of computing the messages from the newly added region to the root is bounded by  $n \cdot \exp(z)$ . Having the complexity of a single merge as  $O(S) = O(L^2 \cdot \exp(z) + L^3 \cdot z + L \cdot z \cdot \exp(z) + n \cdot \exp(z))$ , we can then bound the complexity of doing all merges as  $O(R \cdot L^2 \cdot \exp(z) + R \cdot L^3 \cdot z + R \cdot L \cdot z \cdot \exp(z) + R \cdot n \cdot \exp(z))$ .  $\square$

This memory-aware incremental built have several advantages: (1) It allows assessing the memory required for each new region, and only adds those regions that result in an updated cluster tree that still fits in the available memory. As a result, the algorithm never runs out of memory, merging as many regions as possible along the elimination order, and always produces an upper bound. (2) When the *ibound* found by simulating the scope-based partitioning for mini-bucket elimination leaves a large portion of memory unused, we can use the memory-aware incremental region selection of Algorithm 4.2, WMBE-IB, with  $ibound+1$  and allow it to use as much memory until it cannot add any new regions to the approximation. We will study this memory allocation strategy empirically in section 4.4.1.

However, using a larger *ibound* to compensate for the large amount of unused memory is not the only possible solution. In the next section we define the concept of “memory budget” which allows us to have finer control over how the available memory is used by different buckets, and describe how the memory-aware algorithm 4.2 can be updated to use the memory budget instead of a fixed *ibound*.

### ■ 4.3.2 Memory Budget for Weighted MBE

Given a graphical model and a fixed memory limit  $\mathcal{M}$ , our goal is to define and evaluate several different online memory allocation schemes that allow us to have finer control over how the available memory is used. We would like to be able to use the available memory as efficiently as possible and achieve tight approximations.

To do so we define the *memory budget* for bucket  $B_i$ ,  $MB(B_i)$ , as the amount of memory allocated to bucket  $B_i$  which can be used by all the mini-buckets included in that bucket. This means that  $\sum_{r \in B_i} M(r) \leq MB(B_i)$  should hold for every bucket  $B_i$ . We represent the memory budget for the cluster tree  $wmb$  as  $MB(wmb) = \{MB(B_i) \text{ for } B_i \in wmb\}$ . The total memory available is then distributed between different buckets such that  $\mathcal{M} = \sum_{B_i \in wmb} MB(B_i)$ .

The memory budget for bucket  $B_i$  can then replace the fix *ibound* to control the amount of memory used by  $B_i$ . A *memory allocation scheme* then defines how the available memory,  $\mathcal{M}$ , should be distributed among different buckets along the elimination order. There are two questions that need to be addressed by each memory allocation scheme; 1) How to initialize the memory budget for each bucket and 2) how to update the initial estimate as we merge mini-buckets using different partitioning schemes.

However, in order to use any memory allocation scheme, we need to be able to keep track of the available memory and the memory used by different mini-buckets as we add more regions to mini-bucket approximation. To do so, we modify Algorithm 4.2, to use a memory budget instead of *ibound*. Algorithm 4.3 shows the details of memory-aware incremental region selection using memory budget. The general procedure to select and add regions is as before, when using *ibound* and total available memory  $\mathcal{M}$ . The only difference is that the new procedure keeps track of the memory budget and how it is used at each bucket, which is explained in detail next.

---

**Algorithm 4.3 WMBE-MB:** Memory-aware Incremental region selection for WMBE using memory budget

---

**Input:** factor graph ( $\mathcal{G}$ ), available memory  $\mathcal{M}$   
**Initialize**  $wmb$  to a cluster tree using e.g. a min-fill ordering  $o$ , uniform weights  
*// pass forward messages along the elimination order*  
 $wmb \leftarrow \text{msgForward}(wmb)$   
**Initialize Memory Budget**  $MB(wmb) \leftarrow \text{InitMB}(wmb)$   
**for** each bucket  $B_i$  following the elimination order **do**  
  **repeat**  
     $(r_m, r_n, M(r_m, r_n)) \leftarrow \text{SelectMergeMB}(B_i, MB(wmb))$   
     $\mathcal{R} \leftarrow \text{AddRegions}(wmb, o, r_m, r_n)$   
     $wmb \leftarrow \text{MergeRegions}(wmb, \mathcal{R})$   
    *// pass forward messages along the merge path  $\mathcal{R}$  until we reach a root*  
    **set**  $r$  and  $\pi_r$  to the first and second regions in  $\mathcal{R}$  ( $\pi_r$  is parent of  $r$  in cluster tree)  
    **while**  $\pi_r \neq \emptyset$  **do**  
      *// pass forward message*  
       $\lambda_{r \rightarrow \pi_r}(\mathbf{x}_{\pi_r}) = \left[ \sum_{\mathbf{x}_r \setminus \mathbf{x}_{\pi_r}} [f_r(\mathbf{x}_r) \prod_{t: \pi_t = \pi_r} \lambda_{t \rightarrow \pi_r}(\mathbf{x}_{\pi_r})]^{\frac{1}{w^{\pi_r}}} \right]^{w^{\pi_r}}$   
    **end while**  
    *// update available memory :*  
     $MB(wmb) \leftarrow \text{UpdateBudgetMerge}(MB(wmb), M(r_m, r_n))$   
  **until** no more merges possible  
  *// update available memory :*  
   $MB(wmb) \leftarrow \text{UpdateBudget}(MB(wmb))$   
**end for**  
**Return:** Upper bound on  $Z$ :  $\hat{Z} = \prod_{r: \pi_r = \emptyset} \lambda_{r \rightarrow \emptyset}$

---

**Initialization.** Given the factor graph  $\mathcal{G}$  and the elimination order  $o$ , as before, first we initialize a cluster tree  $wmb$  with  $ibound = 1$ . Next, we initialize a memory budget  $MB(wmb)$  to manage memory allocation and keep track of the memory used as new regions are added to the approximation. To do so, taking into account the amount of memory used by the initial cluster tree,  $M(wmb)$ , the available memory is updated as  $\mathcal{M} = \mathcal{M} - \sum_{B_i \in wmb} M(B_i)$  and the memory budget  $MB(wmb)$  is initialized by distributing available memory among different buckets based on a memory allocation scheme (described in section 4.3.3). This step is done by  $\text{InitMB}(wmb)$  in Algorithm 4.3.

---

**Algorithm 4.4** SelectMergeMB: Scoring all possible merges in a bucket while taking the memory budget into account

---

**Input:** cluster tree  $wmb$ , bucket  $B_i$ , elimination order  $o$ , memory budget  $MB(wmb)$

**for** every pair  $(r_j, r_k)$  in  $B_i$  **do**

Set  $C = \text{var}(r_j) \cup \text{var}(r_k)$  to a clique that contains all variables from  $r_j$  and  $r_k$

*// check if merge respects the memory budget :*

$(fits, M(r_m, r_n)) \leftarrow \text{CheckMemMB}(wmb, C, MB(wmb))$

**if**  $fits$  **then**

$S(j, k) \leftarrow \text{score}(r_j, r_k)$

*// keep track of the memory it requires*

$M(j, k) \leftarrow M(r_j, r_n)$

**else**

$S(j, k) \leftarrow -\text{inf}$

**end if**

**end for**

*// select the best merge:*

$(j^*, k^*) = \arg \max_{j,k} S(j, k)$

**Return:**  $(r_{j^*}, r_{k^*})$  and the corresponding memory profile  $M(r_{j^*}, r_{k^*})$

---

**Algorithm 4.5** UpdateBudgetMerge: Updating memory budget after merging  $(r_m, r_n)$

---

**Input:** memory budget  $MB(wmb)$ , memory profile of the merge  $M(r_m, r_n)$

*// compute the extra memory we have along the merge path  $B_m$  :*

$ME = \sum_{B_m} MB(r_m, r_n)$

*// update the budget along the merge path  $B_m$  :*

**for**  $B_i \in B_m(\text{end} : -1 : 1)$  **do**

**if**  $MB(B_i) < M(B_i)$  **then**

$MB(B_i) = M(B_i)$

**end if**

$ME = ME - MB(B_i)$

**end for**

$MB(B_1) = ME$

---

**Selecting Mini-buckets to Merge.** To select the pair of mini-buckets to merge, our budget based algorithm needs to compare the required memory for the merge at each bucket with the memory budget allocated to them. We update the two subroutines, SelectMerge(.) and CheckMem(.), to use the memory budget. The updated subroutine, SelectMergeMB(.), Algorithm 4.4, then uses CheckMemMB(.), Algorithm 4.7, to compute the memory required for the merge and score the pairs that can be successfully added to the current cluster tree  $wmb$ . Having a memory budget  $MB(wmb)$ , which defines a limit on how much memory can

be used in each bucket,  $\text{CheckMemMB}(\cdot)$  needs to check if the pair of regions  $(r_m, r_n)$  can be successfully added to the current cluster tree while respecting the memory budget  $MB(B_i)$  at every bucket that is affected by the merge. The pair is then scored only if it respects the budget. After scoring all pairs that respect the memory budget, the best is selected to be added as the next region to the cluster tree.

**Checking Memory Requirement of a Merge.** Checking for the memory required by a merge is done by simulating the process of adding the new merged region to the current cluster tree  $wmb$ . If a merge is possible, the subroutine  $\text{CheckMemMB}(\cdot)$  returns a memory profile,  $M(C)$ , for the merge which contains the memory required at each bucket,  $M(B_i)$ , that is affected by the merge and contains the amount of memory used by that bucket after the merge. Algorithm 4.7 details all the steps involved.

The procedure to compute the memory required at each bucket,  $M_{new}(B_i)$ , is similar to WMBE-IB. However instead of comparing the total memory used after adding the new regions ( $M_{used}$ ) with the total memory available ( $\mathcal{M}$ ) the algorithm compares the memory used at each bucket ( $M(B_i)$ ) with the memory budget allocated to it ( $MB(B_i)$ ). To do so, when merging the two regions  $r_j$  and  $r_k$  into the new region  $r_n$ , the algorithm first checks if the required memory at  $B_i$  after the merge,  $M_{new}(B_i)$ , respects the memory budget  $MB(B_i)$  and continues only if the required memory is less than the available budget for the bucket.

Adding the new region  $r_n$  to the cluster tree might require adding additional regions to the buckets later in the elimination order as well.  $\text{CheckMemMB}(\cdot)$  keeps track of these buckets in  $B_m = \{B_i; \text{if } \exists \text{ new } r_n \in B_i \text{ after the merge}\}$ . To decide if there is enough memory for the merge, the sum of the required memory for all the buckets in  $B_m$  is compared with the total memory available to those by the memory budget, as  $\sum_{B_i \in B_m} M_{new}(B_i) \leq \sum_{B_i \in B_m} MB(B_i)$ . This comparison assumes that any extra memory that is allocated to the buckets earlier in the elimination order and is not used, can be shifted to the buckets later in the elimination

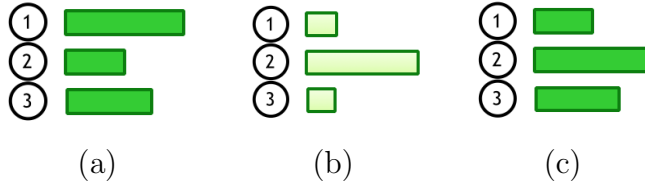


Figure 4.2: Updating the memory budget after a merge: (a) Memory budget before merge (b) Memory required by the merge (c) Memory budget after merge

order.

The simulation continues until the new region can be merged with an existing region in the cluster tree or the new region has no parents. The algorithm then returns if the merge can be done and the memory required at each bucket  $B_i \in B_m$  after the merge, in a memory profile  $M(C)$ .

**Updating graph structure and memory budget.** Having selected the two regions to add, the two subroutines, `AddRegions(.)` and `MergeRegions(.)`, perform the merge exactly as before (see Algorithm 3.2 and Algorithm 3.3). After adding the new regions in  $\mathcal{R}$  to  $wmb$ , WMBE-MB algorithm updates the memory budget for buckets affected by the merge ( $B_m$  in `CheckMemMB(.)`) before continuing with another round of adding more regions. This memory budget update is required because `CheckMemMB(.)` compares the sum of the required memory for all the buckets in  $B_m$  against the total memory available for those by the memory budget, e.g.  $\sum_{B_i \in B_m} M_{new}(B_i) \leq \sum_{B_i \in B_m} MB(B_i)$ , to decide if a merge is possible. This comparison is based on the assumption that any amount of available memory can be shifted to later buckets along the elimination order. As a result a merge can be approved if  $\sum_{B_i \in B_m} M_{new}(B_i) \leq \sum_{B_i \in B_m} MB(B_i)$  while the required memory at some bucket  $B_j \in B_m$  is larger than the memory budget at that bucket, e.g.  $M_{new}(B_j) > MB(B_j)$ . Figure 4.2 shows such an example where (a) is the memory budget assigned to the three buckets before the merge, (b) is the required memory at each bucket for the merged region to be added to the cluster tree and (c) is the updated memory budget for the three buckets after the

merge. The required memory at bucket  $B_2$  is more than what is available in the memory budget,  $\text{MB}(B_2)$ , but since bucket  $B_1$  has some extra memory that is not used after adding the new region, a portion of it can be shifted to bucket  $B_2$  to make the merge possible.  $\text{UpdateBudgetMerge}(\cdot)$ , Algorithm 4.5, makes sure that the memory budget after the merge reflects the actual available memory in each bucket after the merge is done by shifting the available memory along the elimination order, from buckets with extra memory to those that require more.

**Memory Reallocation.** After all possible merges are done for bucket  $B_i$ ,  $\text{UpdateBudget}(\cdot)$  shifts all the unused memory that was assigned to  $B_i$  to the buckets later in the elimination order. This final memory budget update is required only for some memory allocation schemes, but results in better memory use and a better approximation in all. Section 4.3.3 describes different strategies for reallocating the unused memory.

**Computational Complexity of WMBE-MB.** Next we analyze the computational complexity of adding new regions using our budget based incremental memory-aware approach.

**Proposition 4.2.** *The computational complexity of merging all mergable pairs of regions  $(r_m, r_n)$  in WMBE-MB (Algorithm 4.3), is at most  $O( R \cdot L^2 \cdot M + R \cdot L^3 \cdot \log M + R \cdot L \cdot M \cdot \log M + R \cdot M )$ , where  $L$  is the maximum number of initial regions in a bucket (which is bounded by the maximum degree of any variable),  $M$  is the total memory and  $R = \sum_{\alpha} |\alpha|$  upper bounds the number of regions in the cluster graph.*

*Proof.* Let  $R$  be as defined and denote by  $O( S )$  the computational complexity of a single merge. Again, we cannot have more than  $R$  merges, which means that the complexity of all possible merges is  $O(R \cdot S)$ . The complexity of a single merge can then be bounded as  $O( S ) = O( L^2 \cdot M + L^3 \cdot \log M + L \cdot M \cdot \log M + M )$ . The first term in  $O( S )$  captures the complexity of computing the score, which is  $O( M )$ , for any pair of regions. The algorithm

computes the score for every pair of regions in a bucket, which yields  $O(L^2 \cdot M)$  for score computation. The second term in  $O(S)$  is the complexity of  $\text{CheckMem}(\cdot)$  that simulates the merge process to decide if a merge fits in memory. For each merge we have to add the new merged region and all its parents, which is bounded by  $\log M$ . After adding these new regions, we need to check for subsumptions of old regions of the cluster graphs with the new ones which involves  $\log M \cdot L$  tests. Checking for the required memory using  $\text{CheckMem}(\cdot)$  should be done for all merges in the bucket, and results in total complexity of  $O(L^3 \cdot \log M)$ . The third term then computes the bound for updating the graph structure for the pair that is selected to be merged. Since each elimination reduces the table size geometrically, the complexity of adding the new merged region and its parents is bounded by  $\log M$ , with  $\log M \cdot L$  possible merges between the old regions and the new ones. For every such merge, if one region can be subsumed by the other, we need to update the messages sent to them by their children; this requires  $O(M)$  computations for each update. Putting it all together, the computational complexity of the merge process is bounded by  $O(L \cdot M \cdot \log M)$ . Having updated the cluster graph structure, the complexity of computing the messages from the newly added region to the root is bounded by  $M$ . Having the complexity of a single merge as  $O(S) = O(L^2 \cdot M + L^3 \cdot \log M + L \cdot M \cdot \log M + M)$ , we can then bound the complexity of doing all merges as  $O(R \cdot L^2 \cdot M + R \cdot L^3 \cdot \log M + R \cdot L \cdot M \cdot \log M + R \cdot M)$ .  $\square$

### ■ 4.3.3 Memory Allocation Schemes

Using the above memory aware algorithm, different memory allocation schemes proceed by using different methods to initialize the memory budget and update it at each step. Different schemes studied in this thesis are as follows:

**Waterfall Memory Distribution.** Perhaps the simplest way to allocate available memory is to allow each bucket to use as much memory as it needs, adding the largest regions possible to

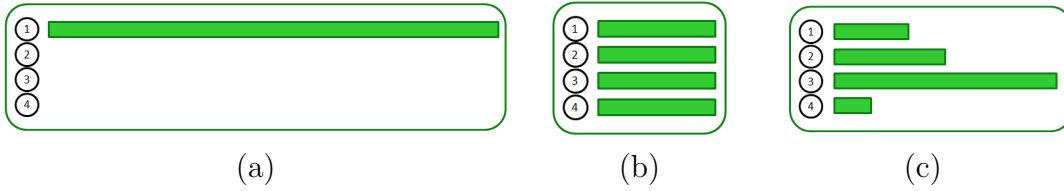


Figure 4.3: Different memory allocation schemes: (a) Waterfall Distribution where the total available memory is assigned to the first bucket and any extra memory after all merges at each bucket is redistributed between the buckets later in the elimination order. (b) Uniform Distribution where the total available memory is equally split between different buckets; (c) Proportional Distribution where the total available memory is distributed between different buckets proportional to the memory used by scope-based partitioning;

the approximation until no more memory is available. In this scheme, all available memory is initially allocated to the first bucket; see Figure 4.3 (a). Any memory that is unused by the first bucket can then be shifted along the elimination order to become available to the next bucket. For this simple allocation scheme, we can define  $\text{InitMB}(wmb)$  as:

$$\begin{aligned}
 MB(B_1) &= \mathcal{M} \\
 MB(B_i) &= 0 \quad \text{for } i = 2 \dots N
 \end{aligned}$$

To update the memory budget after each bucket is processed, we pass any remaining memory into the next bucket's budget, defining  $\text{UpdateBudget}(wmb)$  as

$$\begin{aligned}
 MB(B_{i+1}) &\leftarrow MB(B_{i+1}) + (MB(B_i) - M(B_i)) \\
 MB(B_i) &\leftarrow M(B_i)
 \end{aligned}$$

where  $B_{i+1}$  is the bucket after  $B_i$  in the elimination order. This scheme is efficient in the sense that almost all available memory will be allocated to some mini-bucket. However, it will typically result in large regions early in the elimination order, and many small regions in buckets later in the order, which can lead to poor approximation quality.

**Uniform Memory Distribution.** A simple way to encourage a more equitable distribution of memory to different buckets is to divide the total memory uniformly among buckets; see Figure 4.3 (b). More formally, for a graphical model with  $N$  variables such allocation uses an initial memory distribution,  $\text{InitMB}(wmb)$ :

$$MB(B_i) = \frac{\mathcal{M}}{N} \text{ for } i = 1 \dots N$$

While this allocation strategy is simple to use, it tends to be very conservative. Some parts of the model are likely to be less complex than others; empirically, we observe that buckets early in the elimination order often need less memory compared to buckets later in the order, which often involve larger regions (a consequence of the graph structure and the elimination ordering chosen).

As a result, uniform memory distribution may not use the available memory as efficiently as the waterfall distribution. To some extent, we can remedy this limitation during the  $\text{UpdateBudget}(\cdot)$  step by shifting any unused memory from earlier buckets to the later ones, similarly to the waterfall scheme; strategies for performing such memory shifts are discussed in the sequel.

**Proportional Memory Distribution.** A final option is to use information from a “pilot” estimate of the complexity of different portions of the inference process. Since estimating the structure, and hence memory requirements of, scope based partitioning is easy and efficient, we can use this scope-only structure to inform our budgets. After searching for the largest *ibound* such that a scope-based partitioning will fit in our total memory, we evaluate the memory usage of each bucket  $B_i$ . Then, we initialize our memory budget proportionally to these values:

$$MB(B_i) = \frac{M_s(B_i)}{\sum_i M_s(B_i)} \times \mathcal{M},$$

where  $M_s(B_i)$  is the memory used by bucket  $B_i$  in the scope-based partitioning and  $\mathcal{M}$  is our total memory ceiling. The proportional scheme provides a better estimate of the required memory at each bucket, while assuming that the memory needs of the content-based partitioning will be similar to that of scope-based partitioning at each level. Figure 4.3 (c) shows an illustration of this allocation scheme.

**Memory Reallocation.** In all memory allocation schemes discussed above, the resulting regions in one bucket may use less memory than allocated to that bucket’s budget. This excess memory can then be redistributed among the later buckets in the elimination order, in the `UpdateBudget(.)` step in Algorithm 4.3. The waterfall redistribution, in which any extra memory is placed in the next bucket, is one such method. Alternatively, however, the extra memory in each bucket can also be distributed among later buckets in other ways – for example, uniformly among the later buckets:

$$MB(B_j) \leftarrow MB(B_j) + (MB(B_i) - M(B_i))/(N - i) \text{ for } j > i$$

or, proportionally to the expected needs of the later buckets:

$$MB(B_j) \leftarrow MB(B_j) + (MB(B_i) - M(B_i)) \frac{MB(B_j)}{\sum_{j>i} MB(B_j)} \text{ for } j > i$$

and reduce the “budget” of bucket  $i$  to reflect its actual use:  $MB(B_i) = M(B_i)$ .

Given these various initialization and update methods, we can combine them to generate a number of different memory allocation strategies:

- (1) **UF**: uniform initial distribution with no reallocation;

- (2) **UU**: uniform initial distribution with uniform reallocation;
- (3) **UP**: uniform initial distribution with proportional reallocation;
- (4) **UW**: uniform initial distribution with waterfall reallocation;
- (5) **PF**: proportional initial distribution with no reallocation;
- (6) **PU**: proportional initial distribution with uniform reallocation;
- (7) **PP**: proportional initial distribution with proportional reallocation;
- (8) **PW**: proportional initial distribution with waterfall reallocation;
- (9) **WW**: waterfall initial distribution with waterfall reallocation.

---

**Algorithm 4.6 CheckMem:** Checks if a new clique  $C$  can be added to the current cluster tree  $wmb$  and returns the memory required for the updated cluster tree after merge

---

**Input:** The cluster tree  $wmb$ , elimination order  $o$ , total memory available  $\mathcal{M}$  and clique  $C$

**Output:**  $M_{used} = M(wmb)$  after the merge is done

**Initialize:**  $B_m$  to the bucket corresponding to the first un-eliminated variable in  $C$  based on elimination order  $o$

**Initialize:**  $X_m$  to the variable that is being eliminated at  $B_m$

**Initialize:**  $M_{used} = \sum_i M(B_i)$  to keep track of the memory used by current cluster tree  $wmb$

**Initialize:**  $done \leftarrow False$ ;

**while** not  $done$  **do**

**Consider** new region  $r_n$  with  $\text{var}(r_n) = C$  and set  $M_{old} \leftarrow 0$

**Find** all regions  $r_m \in B_m$  that can be subsumed by  $r_n$  ( $C \subseteq \text{var}(r_m)$ ) and add their memory,  $M(r_m)$ , to  $M_{old}$

**Set**  $M_{new} = M(B_m) + M(r_n) - M_{old}$

**Update**  $M_{used} = M_{used} - M(B_m) + M_{new}$

**if**  $M_{used} > \mathcal{M}$  **then**

        // memory required to add the new region exceeds memory limit:

$done \leftarrow True$

**end if**

**if** not  $done$  **then**

        // Remove  $X_m$  from  $C$  to get the scope of the outgoing message from the new region:

**Set** new clique  $C = \text{var}(r_n) \setminus X_m$

**if**  $C = \emptyset$  **then**

            // new region  $r_n$  is a root and has no outgoing messages:

$done \leftarrow True$

**else**

**Find**  $B_m$  corresponding to the first un-eliminated variable in  $C$  based on elimination order  $o$

**for** each mini-bucket region  $r_m \in B_m$  **do**

**if**  $C \subseteq \text{var}(r_m)$  **then**

                    // forward message fits in existing mini-bucket:

$done \leftarrow True$

**end if**

**end for**

**end if**

**end while**

**return**  $M_{used}$

---

---

**Algorithm 4.7 CheckMemMB:** Checks if a new clique  $C$  can be added to the current cluster tree  $wmb$  under the memory budget  $MB$

---

**Input:** The cluster tree  $wmb$ , elimination order  $o$ , memory budget  $MB$  and clique  $C$   
**Output:**  $fits \leftarrow True$  if  $C$  can be added to  $wmb$ , and the memory required for the merge  $MC$   
**Initialize:**  $B_m = \{B_j\}$  to the bucket corresponding to the first un-eliminated variable in  $C$ , based on elimination order  $o$  and  $X_j$  to the variable that is being eliminated at  $B_j$   
**Initialize:**  $fits \leftarrow True$ ;  
**Consider** new region  $r_n$  with  $\text{var}(r_n) = C$  and set  $M_{old} \leftarrow 0$   
**Find** all mini-bucket region  $r_j \in B_j$  that can be subsumed by  $r_n$  and add their memory to  $M_{old}$   
**Set**  $M_{new}(B_j) = M(B_j) + M(r_n) - M_{old}$   
**if**  $M_{new}(B_j) > MB(B_j)$  **then**  
     $fits \leftarrow False$  // New clique doesn't fit in the budget for bucket  $B_j$   
**end if**  
**if**  $fits$  **then**  
    **repeat**  
        **Set** new clique  $C = \text{var}(r_n) \setminus X_j$   
        **if**  $C = \emptyset$  **then**  
             $done \leftarrow True$   
        **else**  
            **Find**  $B_j$  corresponding to the first un-eliminated variable in  $C$  based on elimination order  $o$  and set  $X_j$  to the variable that is being eliminated at  $B_j$   
            **for** each mini-bucket region  $r_j \in B_j$  **do**  
                **if**  $C \subseteq \text{var}(r_j)$  **then**  
                     $done \leftarrow True$  // forward message fits in existing mini-bucket  
                **end if**  
            **end for**  
        **end if**  
        **if** not  $done$  **then**  
             $B_m \leftarrow B_m \cup \{B_j\}$  // New region needs to be added to contain forward message  
            **Consider** new region  $r_n$  with  $\text{var}(r_n) = C$  and set  $M_{old} \leftarrow 0$   
            **Find** all mini-bucket region  $r_j \in B_j$  that can be subsumed by  $r_n$  and add their memory to  $M_{old}$   
            **Set**  $M_{new}(B_j) = M(B_j) + M(r_n) - M_{old}$   
            **if**  $\sum_{B_i \in B_m} M_{new}(B_i) > \sum_{B_i \in B_m} MB(B_i)$  **then**  
                // Total budget along these buckets is not enough to add the new clique  $B_j$ :  
                 $fits \leftarrow False$  and  $done \leftarrow True$   
            **end if**  
        **end if**  
    **until**  $done$   
**end if**  
**return** ( $fits, M(C) = \{M_{new}(B_i) \text{ for } B_i \in B_m\}$ )

---

## ■ 4.4 Empirical Evaluation

In this section, we compare the proposed memory allocation schemes on two domains of problems, linkage analysis and protein side chain prediction, from past UAI approximate inference challenges and study their ability to effectively use the available memory, and the quality of the bounds they produce on the log-partition function using content-based partitioning.

The two problem domains from which our instances are drawn have very different statistics and operating points. The protein side-chain instances have variables with cardinalities ranging from 2 to 81. On these instances, we typically find very low feasible *ibound* values for typical memory bounds (e.g., 1GB), and memory use changes in large jumps, so that many instances use 20% or less of the available memory. On the other hand, the pedigree (linkage analysis) instances have variables with cardinality only between 2 and 5; on these instances, the *ibound* selected is typically much higher, and a standard scope-based mini-bucket uses, on average, more than half the available memory.

In the sequel, we will refer to the largest *ibound* value for which scope-based partitioning fits into our memory budget as “*iscope*”.

### ■ 4.4.1 Content Based Partitioning

First, we explore the effect of different memory allocation schemes on the quality of the bound. To this end, we compare the content-based approximation of WMBE-IB with the approximations from WMBE-MB under various memory budget strategies. For all methods, when adding new regions, we use the scoring function introduced in Chapter 3, Eq. (3.7), and merge the pair with the maximum score.

**Protein Side-chain prediction.** For the protein side-chain prediction examples, we find

Table 4.1: **UAI Protein Side-chain Prediction.** Each column shows the improvement in the upper bound for that memory allocation scheme, over using an *ibound* control on complexity (see section 4.3.3). The improvement is computed as  $\log \hat{Z}_{iB} - \log \hat{Z}_M$ , where  $\log \hat{Z}_M$  is the upper bound to the log partition function computed using WMBE-MB and  $\log \hat{Z}_{iB}$  is the upper bound computed by WMBE-IB. Larger values indicate improved approximation using the budget based WMBE-MB, compared to the *ibound* based algorithm, WMBE-IB.

<b>Inst</b>	<b>UF</b>	<b>UU</b>	<b>UP</b>	<b>UW</b>	<b>PF</b>	<b>PU</b>	<b>PP</b>	<b>PW</b>	<b>WW</b>
2cav	67.98	119.15	119.15	153.30	67.91	165.43	149.42	168.21	66.68
1ehg	30.32	63.69	63.69	78.16	15.12	66.97	48.16	77.02	-95.58
1exm	43.51	69.57	69.57	66.81	42.58	71.14	54.77	68.32	-61.66
1i24	122.73	111.82	111.82	144.42	81.87	115.32	94.90	114.35	-76.40
1ewf	100.62	107.53	107.53	116.20	72.31	119.73	106.30	121.71	-116.66

$iscope \leq 5$  in general. This is much smaller than the induced width of the models (typically  $\geq 20$ ). Using a weighted min-fill elimination order, we initialize a cluster tree using the individual factors in each model. We then use WMBE-MB (Algorithm 4.3) with different memory allocation schemes to upper bound the log partition function, and compare these bounds with that of WMBE-IB (Algorithm 4.2) when  $ibound = iscope$ .

In all 44 protein instances, only one content-based WMBE-IB used all available memory before reaching the last bucket. However, in 65% of instances, content-based WMBE-IB used more memory than scope-based WMBE, but did not reach the memory limit.

Table 4.1 shows the improvement in the bounds of WMBE-MB over WMBE-IB (i.e., the difference of their bounds) on a randomly selected subset of these models. Each column corresponds to a different budget management strategy. The values in the table correspond to the difference  $\log \hat{Z}_{iB} - \log \hat{Z}_M$ , where  $\log \hat{Z}_M$  is the upper bound computed by WMBE-MB and  $\log \hat{Z}_{iB}$  is the upper bound computed by WMBE-IB.

Figure 4.4 (a) summarizes the results for each memory allocation scheme across all instances. For each allocation strategy, a box plot summarizes the improvement in the upper bound to the log partition function, e.g.,  $\log \hat{Z}_{iB} - \log \hat{Z}_M$ , for that allocation scheme; positive values

indicate that WMBE-MB found a tighter bound. Additionally, Figure 4.4 (b) shows a summary of the proportion of the available memory used by each allocation scheme. Strategies employing “fixed” reallocation (e.g., no reallocation of memory) used the smallest fraction of memory; strategies with some reallocation all averaged  $\approx 80\%$  or higher usage.

On these set of instances, all allocation schemes except waterfall allocation (WW) give an improved upper bound compared to *iboundcontrol*. WW fares poorly, since in practice it allows all possible merges in each bucket until the available memory is exhausted (see Figure 4.4 (b)). This results in fully merged buckets and exact elimination for variables early in the elimination order, followed by almost no merges (very small mini-buckets) after the memory is used; this gives poor quality approximations. Other schemes bound the maximum amount of memory available to each bucket, preventing early buckets from consuming all the memory and improving the approximation.

As expected, memory shifting schemes (UU, UP, UW and PU, PP, PW) allow better memory allocation and improve the approximation more than the corresponding fixed memory allocation schemes (UF and PF). It is interesting to note that in general, shifting the extra memory using a waterfall distribution (UW and PW) improves the approximation more than uniform and proportional distribution of extra memory (UU and PU, PU and PP). This may be because uniform memory shifting tends to overestimate the memory needed by buckets later in the elimination order, limiting the available memory to earlier buckets and leaving more memory unused at the end. On the other hand, waterfall redistribution of unused memory allows the extra memory to be used as needed at each bucket when determining merges. Unlike the full waterfall strategy (WW), here the initial uniform or proportional distribution limits the available memory to early buckets and avoids too uneven a distribution of the available memory.

**Alternative Initial Structure.** One alternative to consider is to use a “coarse to fine”

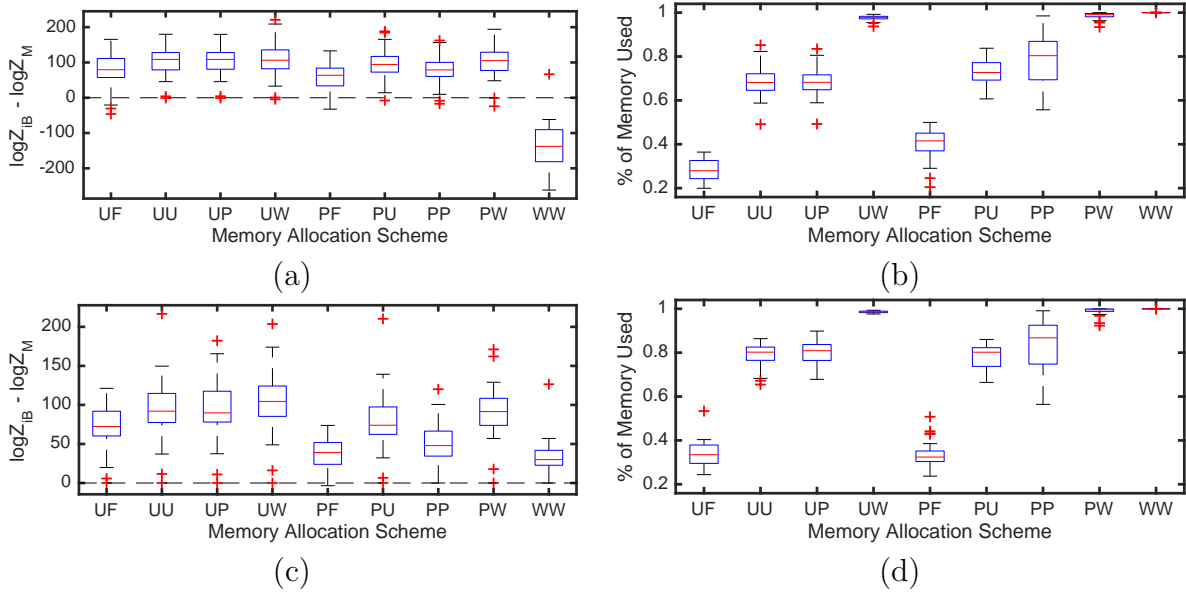


Figure 4.4: **Protein side-chain prediction.** Summarizing (a),(c) the improvement in the upper bound, and (b),(d) the proportion of memory used, for different memory allocation schemes across all instances. The horizontal axis represents different allocation schemes (see Section 4.3.3). The vertical axis shows the improvement  $\log \hat{Z}_{iB} - \log \hat{Z}_M$ . (a) Each box plot summarizes the improvement in the upper bound for the corresponding scheme when WMBE-MB starts from an initial factor graph. Positive values indicate better approximation by content-based WMBE-MB compared to content-based WMBE-IB. (b) Proportion of the memory used by different allocation schemes when WMBE-MB starts from an initial factor graph. (c) Each box plot summarizes the improvement to the upper bound for the corresponding scheme when WMBE-MB is initialized with the cluster tree of content-based WMBE-IB when  $ibound = iscope$ . (d) Proportion of the memory used when WMBE-MB is initialized with the cluster tree of content-based WMBE-IB when  $ibound = iscope$ .

memory allocation. Instead of initializing our cluster tree to individual factors in WMBE-MB, we could first use WMBE-IB with  $ibound = iscope$  to compute an approximation and then execute WMBE-MB to refine the resulting cluster tree. Then, WMBE-MB can distribute any unused memory among different buckets. Figure 4.4 (c) and (d) show a summary of the improvement in the approximation and proportion of memory used for different memory allocation schemes in this case. With the exception of WW, the qualitative results are similar to starting WMBE-MB from individual factors. For WW, however, starting from the cluster tree with induced width of  $iscope$  avoids using all the memory early, giving a bound that is at least as good as using no memory allocation.

Table 4.2: **UAI Protein Side-change Prediction.** The upper bound computed by WMBE-MB is compared for different initializations.  $\log \hat{Z}_{FG}$  is the upper bound computed when WMBE-MB is initialized with the factor graph and  $\log \hat{Z}_{IB}$  is the upper bound computed when WMBE-MB is initialized with the resulting cluster tree from WMBE-IB using  $ibound = iscope$ . Each entry shows the proportion of instances for which one approximation is better than the other.

UB\MemScheme	UF	UU	UP	UW	PF	PU	PP	PW	WW
$\log \hat{Z}_{FG} < \log \hat{Z}_{IB}$	63%	63%	65%	55%	83%	80%	78%	68%	0%
$\log \hat{Z}_{FG} > \log \hat{Z}_{IB}$	37%	37%	35%	45%	17%	20%	22%	32%	100%

Figure 4.5 compares the results of the best two memory allocation schemes,  $UW$  and  $PW$ , for the two possible initializations (individual factors, or  $ibound$ -based), and Table 4.2 shows the proportion of instances for which each initialization results in a tighter upper bound. The results show that initializing WMBE-MB with the factor graph results in better approximation in most cases.

Initializing WMBE-MB using the factor graph allows more flexibility in how the available memory is used, and seems to result in generally tighter bounds. Starting from the cluster tree of WMBE-IB with  $ibound = iscope$  often results in mini-buckets that cannot be merged unless the memory available to the bucket is increased significantly. As an example, consider a factor graph with three factors on  $X_1$ :  $\{X_1, X_2, X_3\}$ ,  $\{X_1, X_4, X_5\}$ , and  $\{X_1, X_7\}$ . Using  $ibound = 4$ , we can merge the first two mini-buckets, resulting in two mini-buckets,  $\{X_1, X_2, X_3, X_4, X_5\}$  and  $\{X_1, X_7\}$ , one of which is over five variables. However if  $ibound = 3$  was used to initialize the cluster tree first, resulting in  $\{X_1, X_2, X_3, X_7\}$  and  $\{X_1, X_4, X_5\}$ , we are not able to perform any further merging if  $ibound$  is increased to 4, yielding a largest mini-bucket over four variables.

**Choice of  $ibound$ .** A simple technique to use a larger fraction of available memory is to use  $ibound = iscope + 1$  in content-based WMBE-IB, especially when  $iscope$  leaves large portions of the memory unused. Memory-aware WMBE-IB then allows merges up to  $iscope + 1$ , until

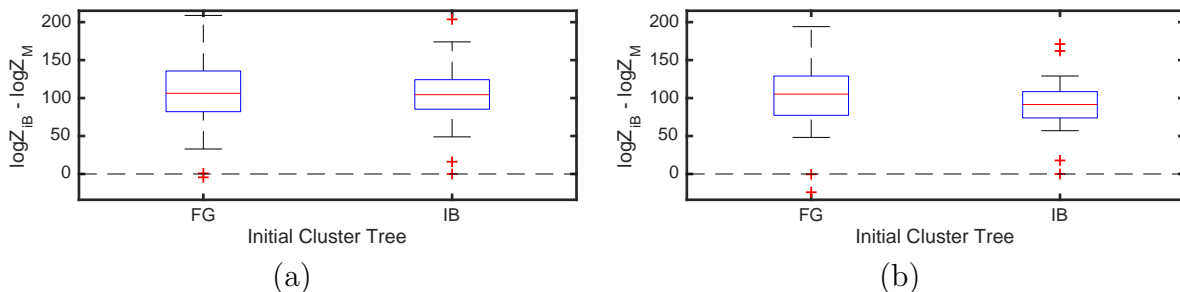


Figure 4.5: Comparing the improvement in the bound when WMBE-MB is initialized with (1) factor graph (FG), and (2) the resulting cluster tree from WMBE-IB (IB) when  $ibound = iscope$ . (a) shows the improvement in the approximation for  $UW$  and (b) shows the improvement in the approximation for  $PW$  memory allocation scheme. The improvement is computed as  $\log \hat{Z}_{iB} - \log \hat{Z}_M$ , where  $\log \hat{Z}_{iB}$  is the upper bound computed using WMBE-IB and  $\log \hat{Z}_M$  is the upper bound computed using WMBE-MB with different initializations. Although the improvement given the two initialization is very close, starting from the initial factor graph gives more flexibility in using the available memory and results in tighter bounds in general.

the total memory limit is reached. Unfortunately, this simple strategy does not consistently result in better approximations. In approximately half of protein instances, content-based WMBE-IB with  $iscope + 1$  uses all the available memory before reaching the buckets later in the elimination order, giving an upper bound that is less tight than WMBE-BI with  $iscope$ . In general, memory-based allocation performs better – for all instances, using WMBE-MB with  $PW$  memory allocation gave a better approximation than WMBE-IB with  $ibound = iscope + 1$ . Figure 4.6 compares the upper bounds computed using WMBE-MB using  $UW$  and  $PW$  allocation schemes with the upper bound computed with WMBE-IB.

**Linkage Analysis.** We also analyze performance on the linkage analysis (pedigree) models, where scope-based WMBE typically uses a higher proportion of the available memory. Pedigree instances have variables whose cardinalities vary in the range of  $[2, \dots, 5]$ , and the induced width of the model is typically  $\approx 20 - 30$ . The  $iscope$  found for these models is, on average, about half the induced width.

Content-based partitioning with WMBE-MB behaves quite differently on these instances, compared to the protein examples where the approximation using  $iscope$  left a large portion

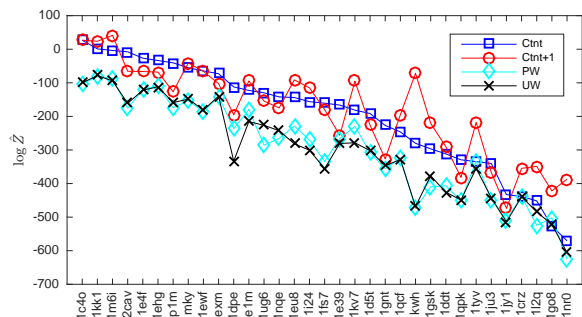


Figure 4.6: **Protein side-chain prediction** The upper bound computed by WMBE-MB using  $UW$  and  $PW$  memory allocation schemes is compared with the upper bound computed using WMBE-IB when  $ibound$  is set to  $iscope$  (Cnt) and to  $iscope + 1$  (Cnt+1) respectively. The approximations using  $ibound = iscope + 1$  use as much memory as the approximations using  $UW$  and  $PW$  but in all cases the memory allocation schemes allow us to compute better approximation with tighter bounds. Here the x-axis show different instances of protein side-chain prediction and the y-axis shows the upper bound.

of memory unused. Figure 4.7 (a) summarizes the improvement of the upper bound for different memory allocation schemes over all instances, a random subset of which is shown in Table 4.3. Figure 4.7 (b) shows the proportion of memory used by each memory allocation scheme. For 25% of instances, content-based WMBE-IB with  $ibound = iscope$  used more memory than scope-based WMBE, but did not reach the memory limit before processing the final bucket.

For linkage analysis models, the memory used by different buckets varies considerably, which is why initializing the memory proportionally to scope-based WMBE usage proves to be more useful than initializing uniformly. The results generally agree with those from the protein problems, although the improvement is not as pronounced for these instances. The main difference is that while we could only fit protein models with very low  $ibound$  values (2 or 3) in 1GB of memory, here we can use much higher  $ibound$  values within the memory limit. Any improvement in the approximation from using more memory is less pronounced, compared to models with a much smaller  $ibound$ . Again, although the waterfall distribution uses almost all memory available, it does not provide a tight bound because it distributes the memory unevenly among buckets and leaves many, small mini-buckets later in the elimination order.

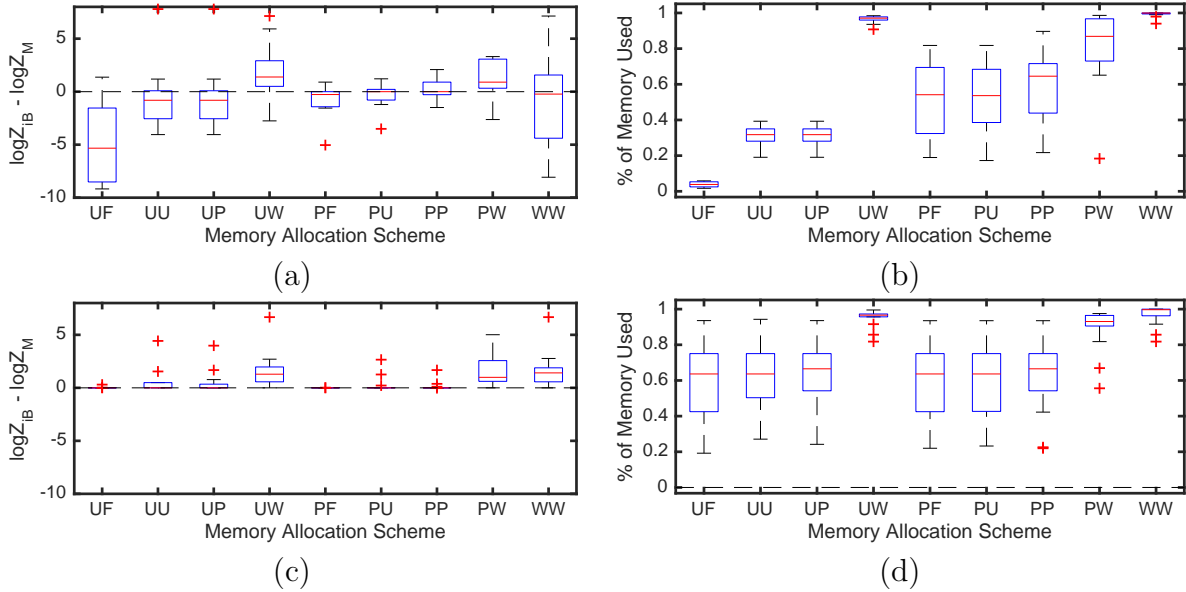


Figure 4.7: **UAI Linkage Analysis.** Summarizing (a),(c) the improvement in the upper bound, and (b),(d) the proportion of memory used, for different memory allocation schemes over all linkage instances. For each allocation scheme, the vertical axis shows the improvement  $\log \hat{Z}_{iB} - \log \hat{Z}_M$ . (a) Box plot summarizing the improvement in the upper bound for each scheme when WMBE-MB starts from an initial factor graph. Positive values indicate better bounds from WMBE-MB than WMBE-IB. (b) Proportion of the memory used by different allocation schemes when WMBE-MB starts from an initial factor graph. (c) Box plots summarizing the improvement for the corresponding scheme when WMBE-MB is initialized with the cluster tree of content-based WMBE-IB with  $ibound = iscope$ . (d) Proportion of memory used when WMBE-MB is initialized with the cluster tree of WMBE-IB with  $ibound = iscope$ .

$UW$  and  $PW$  appear to distribute the available memory more effectively compared to the other schemes and result in an improved upper bound in all but 2 instances.

Figure 4.7 (c) and (d) show the improvement in the upper bound and the memory used by each allocation scheme when WMBE-MB is initialized using the cluster tree built by WMBE-IB with  $ibound = iscope$ . As expected, all memory allocation schemes result in an upper bound that is at least as good as the initial approximation by WMBE-IB. Here, since the amount of extra memory that can be distributed among different buckets is not as large, under many schemes ( $UF$  and  $PF$ ) no new regions can be added to the approximation and the bound does not change. However shifting the extra memory along the elimination order

Table 4.3: **UAI Linkage Analysis.** Different columns show the improvement on the upper bound achieved using each memory allocation scheme (see section 4.3.3). The improvement in the bound is computed as  $\log \hat{Z}_{iB} - \log \hat{Z}_M$ , where  $\log \hat{Z}_M$  is the upper bound to log partition function computed by WMBE-MB and  $\log \hat{Z}_{iB}$  is the upper bound computed by WMBE-IB. Larger values then show better approximation using budget based algorithm compared to a fixed *ibound*.

<b>Ins</b>	<b>UF</b>	<b>UU</b>	<b>UP</b>	<b>UW</b>	<b>PF</b>	<b>PU</b>	<b>PP</b>	<b>PW</b>	<b>WW</b>
ped13	-0.62	-2.55	-2.55	1.57	-1.55	1.21	1.23	1.08	-4.04
ped37	0.55	7.79	7.79	7.14	-0.28	-0.07	-0.28	2.71	7.14
ped38	-6.47	-0.45	-0.45	4.80	-0.26	-0.04	-0.23	3.31	5.09
ped42	1.37	-0.67	-0.67	0.51	-1.16	0.22	0.22	2.14	1.09
ped7	-1.54	1.19	1.19	2.39	-1.53	-1.16	-0.97	3.10	-4.39

Table 4.4: **UAI Linkage Analysis.** The upper bound computed by WMBE-MB is compared for different initializations.  $\log \hat{Z}_{FG}$  is the upper bound computed when WMBE-MB is initialized to the factor graph and  $\log \hat{Z}_{IB}$  when initialized with the cluster tree from WMBE-IB using *ibound = iscope*. Each entry shows the proportion of instances for which that approximation was tighter.

<b>UB\MemScheme</b>	<b>UF</b>	<b>UU</b>	<b>UP</b>	<b>UW</b>	<b>PF</b>	<b>PU</b>	<b>PP</b>	<b>PW</b>	<b>WW</b>
$\log \hat{Z}_{FG} < \log \hat{Z}_{IB}$	15%	22%	22%	65%	36%	50%	65%	58%	36%
$\log \hat{Z}_{FG} > \log \hat{Z}_{IB}$	85%	78%	78%	35%	64%	50%	35%	42%	64%

(as in *UW* and *PW*), seems to make better use of the resources to improve the bound.

Figure 4.8 compares the improvement in the upper bound over the best allocation schemes, *UW* and *PW*, when WMBE-MB is initialized with the factor graph and the cluster tree of WMBE-IB with *ibound = iscope* and Table 4.4 shows the proportion of instances for which each initialization results in a tighter upper bound. As before when the most effective allocation schemes, *UW* and *PW*, are used for WMBE-MB, starting from the initial factor graph results in tighter approximation in most cases.

**iBound vs. Memory** Comparing the effectiveness of WMBE-MB with the *ibound*-based WMBE-IB over the two problem sets highlights an important difference between the two.

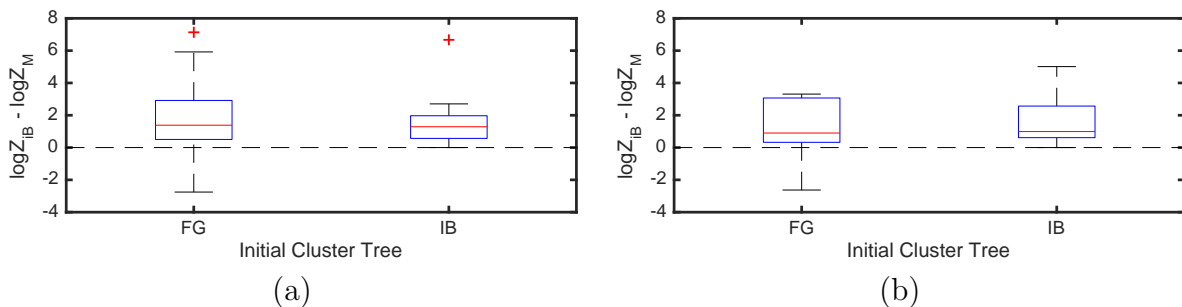


Figure 4.8: **UAI Linkage Analysis.** Comparing the improvement in the bound when WMBE-MB is initialized to the factor graph (FG), or the cluster tree from WMBE-IB when *ibound = iscope* (IB). (a) The improvement in the approximation for *UW*, and (b) improvement in the approximation for *PW* memory allocation scheme. Although the improvement for each of the two initialization is very similar, starting from the initial factor graph gives more flexibility in using the available memory and seems to result in generally tighter bounds.

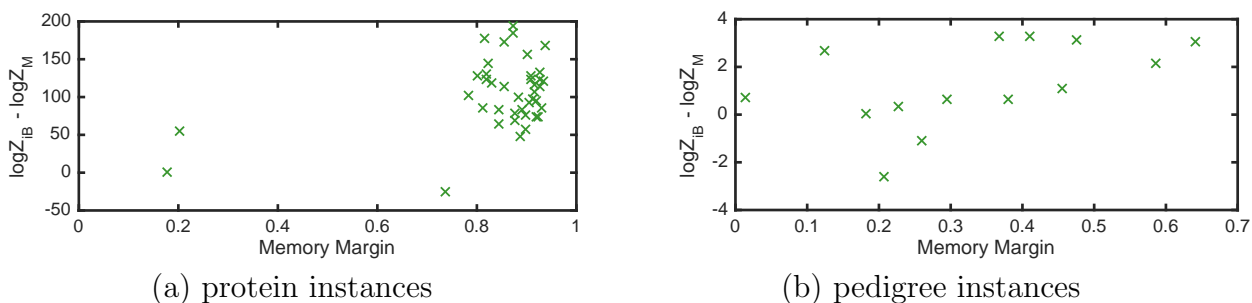


Figure 4.9: Relationship between the proportion of the memory that is not used by WMBE-IB (memory margin) and the improvement of the upper bound when WMBE-MB with *PW* memory allocation scheme is used for (a) protein side-chain prediction and (b) pedigree (linkage analysis) problems (note different scales). Clearly the larger memory margins allow larger potential improvement of the bound.

For protein problems, WMBE-IB typically uses around 30% of the available memory, while for linkage problems, the proportion of memory used is closer to 70%. As a result budget-based algorithms are more effective on the protein instances, and improve the upper bound significantly. Figure 4.9 shows the connection between the fraction of memory that is not used by WMBE-IB (memory margin) and the improvement of the upper bound when WMBE-MB is used with *PW* memory allocation scheme for (a) protein side-chain prediction and (b) linkage analysis problems.

Clearly, larger memory margins allow a greater potential improvement of the bound, since

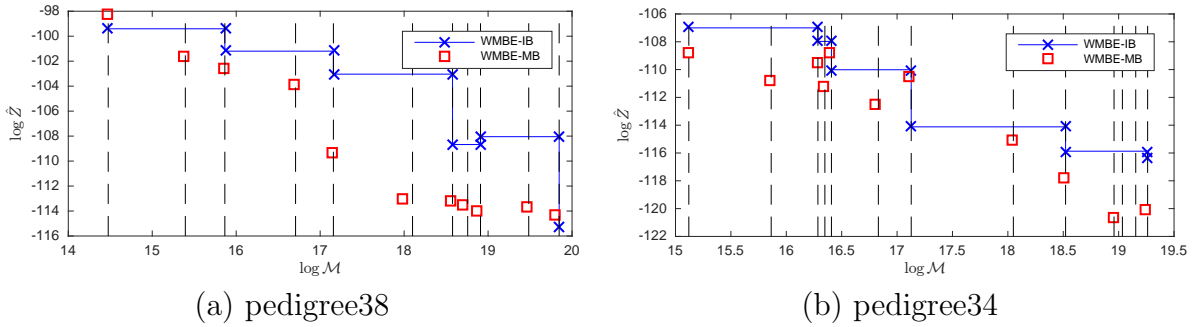


Figure 4.10: Comparing the upper bound computed using WMBE-MB using  $UW$  memory allocation with the upper bound of WMBE-IB as the memory limit is varied. As memory is increased (x-axis, log scale), WMBE-IB improves only when the *ibound* can be increased, resulting in a piecewise constant behavior. In contrast, the fine-grain allocation of WMBE-MB allows better approximation at memory budgets in between, and often even at points where WMBE-IB uses the entire available memory.

the budget-based methods' finer grain control over memory allow them to use larger regions if they fit. But how much benefit does this provide? Additionally, what if the scope-based methods do use all the memory – can budget based methods still more effective?

In order to analyze these questions, we evaluated the bounds produced by WMBE-IB at a series of *ibound* values, and computed the method's memory usage at these points. The performance of WMBE-IB, as a function of memory, is essentially piecewise-constant: if *ibound* = 10 fits into memory, then as the memory budget is increase, the bound will not improve until *ibound* = 11 manages to fit as well. Figure 4.10 traces this step-wise behavior (blue) on two different pedigree instances. Then, we ran WMBE-MB using  $UW$  budget allocation (red), with maximum allowed memory determined by each of these transition points, as well as at memory bounds placed halfway in between. Interestingly, the performance of WMBE-MB is often (though not always) significantly better than WMBE-IB, even when WMBE-IB uses the entire available memory budget. The points halfway in between changes in the *ibound* serve to emphasize the increased flexibility of WMBE-MB to improve its performance as resources are increased by even small amounts.

#### ■ 4.4.2 Content Based Partitioning with Message Passing

Finally, we study the effect of our memory allocation schemes when message passing steps are interleaved with the content-based partitioning. We initialize WMBE-MB using the factor graph, and use each allocation scheme to initialize and update the budget as regions are added to the cluster tree. After each merge we do one round of forward and backward message passing on the WMB cluster tree, similar to Algorithm 3.1, before updating the scores and adding new regions.

Figure 4.11 (a) shows the improvement in the approximation, compared to WMBE-IB (also with message-passing), for the protein instances. For 1 of the 44 instances, WMBE-IB used all the available memory before reaching the final bucket. In 60% of instances it used more memory compared to scope-based WMB with fixed *iscope*, but still used only  $\approx 30\%$  of available memory. On these instances, since *iscope* is very small compared to the induced width of each instance, message passing improves the upper bound significantly. The improvement achieved through memory allocation is therefore typically less than was observed during single pass content-based WMB, but the general ordering of the different schemes and their use of available memory is the same. *UW* and *PW* use the memory more efficiently, merging mini-buckets early in the elimination order while ensuring that there is enough memory left to allow merging later as well. Figure 4.11 (b) shows the same results for linkage analysis instances. As before, the general trends remain the same as without message passing; we see that *UW* and *PW* appear most effective, and achieve tighter bounds compared to WMBE-IB in all but one instance.

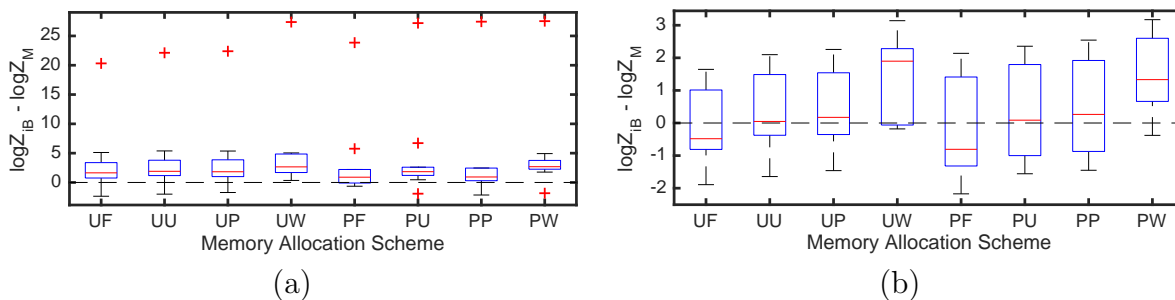


Figure 4.11: Summarizing the results for different memory allocation schemes over all instances of (a) protein side-chain prediction and (b) uai linkage analysis. The horizontal axis represents different allocation schemes (see section 4.3.3) and each box plot summarizes the improvement in the approximation for the corresponding scheme. Positive values indicate better approximation by WMBE-MB compared to WMBE-IB.

## ■ 4.5 Discussion and Future Work

We proposed a framework for using any of several memory allocation schemes to bypass the choice of a single control parameter in content-based WMBE approximation. We studied several memory distribution schemes and evaluated their effectiveness on variety of real world problems. There are two main choices to be made when selecting any of the proposed schemes: first, how to initialize the memory available to each bucket; and second, how to redistribute the extra memory in case some buckets do not use all the memory allocated to them. We showed how different initialization techniques and redistribution of extra memory affect the upper bound on the log partition function, and compared empirical results on two classes of problem instances, which representing two different regimes of models.

Our results show how effective allocation of memory can improve the approximation compared to using a fixed *ibound* to control complexity, especially on models with larger or more variation in the domain size of the variables, and large induced width. It is important to note that using as much memory as possible is not always the best policy for allocating the memory to different buckets; for example, the waterfall initialization fails badly because it fails to save any memory for use with variables late in the elimination ordering. Memory

allocation techniques like  $UW$  and  $PW$  that ensure large regions can be added both to buckets early in the elimination order and those at the end prove to be more effective. The uniform or proportional initial distributions reserve some of the available memory to be used in later buckets, while using waterfall redistribution of extra memory allows excess memory to accumulate and be used to add larger regions along the way. Uniform initial distribution with other redistribution techniques does not use the available memory as effectively, and results in upper bounds that are less tight.

We also discussed how the improvement achieved by different allocation schemes depends on the amount of extra memory that is available to be distributed among different mini-buckets. For instances where there is a large memory margin between the memory used by a scope-based partitioning and the available limit, such as protein problems, all memory allocation schemes result in significant improvements, with  $UW$  and  $PW$  providing in the tightest approximations. However when the memory margin is smaller, as in linkage analysis instances, only  $UW$  and  $PW$  consistently result in tighter upper bounds compared to WMBE-IB.

For problems where there is small memory margin, we showed that running WMBE-MB initialized to the cluster tree of WMBE-IB with  $ibound = iscope$  results in an upper bound that is at least as good as basic WMBE-IB, for all allocation schemes. However initializing WMBE-MB with a factor graph can allow us to find tighter bounds when the most effective allocation schemes,  $UW$  and  $PW$ , are used.

Finally, we studied the interplay between message passing and memory allocation. Not surprisingly, message passing and optimization steps interleaved with region selection can result in tighter approximations compared to the single-pass content based partitioning (this effect was studied in detail in Chapter 3). Efficient allocation of memory resources can still give an additional improvement to the upper bound, with  $UW$  and  $PW$  allocation schemes

being empirically most effective across our various problem types and instances.

The interplay between the message passing steps and efficient use of memory by adding larger regions highlights an important aspect of approximate inference which was not addressed directly in this chapter. So far, we have focused on two aspects to improve the approximation: first, by designing a better content-based partitioning framework for region selection (Chapter 3), and second by using the available memory more effectively via memory-aware partitioning schemes. While these methods give better bounds on the log partition function, any of these methods involve more work compared to a simple scope-based weighted mini-bucket elimination. Given a fixed amount of time, it could be more effective to quickly compute a scope-based cluster tree and use the additional time to run more message passing; or it could be more effective to spend the time scoring and select better regions. Which is the better choice may depend significantly on the problem instance, and may require a more detailed study of the interplay between these different aspects affecting approximation quality.

# Linear Approximation to ADMM for MAP inference

## ■ 5.1 Approximate Inference for MAP

Another fundamental inference task in graphical models is to find the most likely configuration of variables, called maximum a *posteriori* (MAP) inference, equivalent to the “most probable explanation” (MPE) task in Bayesian network literature. Unfortunately, MAP/MPE inference is in general an NP-hard problem, and cannot be solved exactly for many problems of interest.

One successful class of approximate inference algorithms are based on linear programming (LP) relaxations. These algorithms come in several different forms. One approach is to solve the dual of the LP using coordinate descent, employed by MPLP [Globerson and Jaakkola, 2007b] and MSD [Werner, 2007]. Although such methods show good empirical behavior, they are not guaranteed to reach the global optimum of the LP relaxation. Approaches based on variants of subgradient descent [Komodakis et al., 2011, Jojic et al., 2010] are guaranteed to converge globally but are typically slower than coordinate descent approaches in practice [Jojic et al., 2010].

Introduced by Gabay and Mercier [1976], the Alternating Direction Method of Multipliers (ADMM) has recently become popular as an easy-to-apply method for distributed convex optimization with good empirical performance on variety of problems [Lin et al., 2011, Boyd et al., 2011]. However, a direct application of ADMM to the MAP-LP relaxation involves solving a non-trivial quadratic program at each iteration of the algorithm. To circumvent this difficulty, two different globally convergent algorithms based on ADMM have been proposed for MAP-LP relaxations: APLP/ADLP [Meshi and Globerson, 2011] and DD-ADMM [Martins et al., 2011].

Both of these methods avoid solving the non-trivial quadratic program by introducing additional auxiliary variables. In particular, Martins et al. [2011] gives a closed-form update for binary pairwise factors and special “logical constraint” factors; the resulting DD-ADMM algorithm works by converting a general model into this form before solving it. In contrast, the APLP and ADLP algorithms (which correspond to optimizing the primal and dual MAP-LP form, respectively) work on general graphs, but introduce multiple copies of the variables associated with each factor to provide a closed-form update. As a result, both the DD-ADMM and APLP/ADLP approaches increase the number of variables being updated at each iteration and their constraints, which can significantly slow convergence and increases the required memory.

In this work, we choose a different approach to overcome the difficulty of variable updates. Like DD-ADMM [Martins et al., 2011] we use the augmented Lagrangian of the primal MAP-LP problem. However, instead of binarizing the graph to obtain closed form updates, we replace the difficult quadratic local terms by their first order Taylor approximation, allowing closed form updates at each iteration. Such approximations have been previously applied to ADMM in problems such as Low-Rank Representation [Lin et al., 2011]. We show that in practice our algorithm produces better bounds during optimization and improves convergence time for models with general (non-pairwise or non-binary) factors.

## ■ 5.2 MAP and LP Relaxations

Section 2.2.3 introduced the variational view of MAP inference as well as overcomplete exponential family form. Here we discuss an approximate method based on *linear programming relaxation*.

Let  $\mathbf{x} \triangleq \{x_1, \dots, x_N\}$  be a vector of discrete random variables, where each  $x_i \in \mathcal{X}_i$ , with  $\mathcal{X}_i$  a finite set, and let

$$P(\mathbf{x}; \theta) \propto \exp \left( \sum_{f \in F} \theta_f(\mathbf{x}_f) + \sum_{i \in V} \theta_i(x_i) \right)$$

be a probability distribution over  $\mathbf{x}$ , expressed in terms of “factors”  $\theta_f$  each defined over a subset of the variables,  $\mathbf{x}_f$ . We abuse notation to use  $i \in f$  to indicate those variables  $x_i$  that are part of  $\mathbf{x}_f$ , and  $d_i$  to be the number of factors  $f$  in which variable  $i$  participates.

It is helpful to also use the overcomplete exponential family form of undirected graphical models [Wainwright and Jordan, 2008a, Koller and Friedman, 2009b], so that each factor  $\theta_i(x_i)$  can be represented as a vector  $\boldsymbol{\theta}_i^T \boldsymbol{\delta}_{x_i}$  where  $\boldsymbol{\delta}_{x_i}$  is a binary indicator vector with one element,  $\delta_{x_i;s}$ , for each state  $s \in \mathcal{X}_i$ , and  $\delta_{x_i;s}$  takes value one when  $x_i = s$  and zero otherwise. We similarly define  $\boldsymbol{\theta}_f$  and  $\boldsymbol{\delta}_{\mathbf{x}_f}$  over the configurations of  $\mathbf{x}_f$ . Finding the most probable assignment (or MAP configuration),

$$\mathbf{x}^* \triangleq \arg \max_{\mathbf{x} \in \mathcal{X}} P(\mathbf{x}; \theta)$$

can then be framed as an integer linear program over the  $\boldsymbol{\delta}_{x_i}$ ,  $\boldsymbol{\delta}_{\mathbf{x}_f}$ .

$$\begin{aligned}
& \max_{(\boldsymbol{\delta}_{x_f}, \boldsymbol{\delta}_{x_i})} \sum_{f \in F} \boldsymbol{\theta}_f^T \boldsymbol{\delta}_{x_f} + \sum_{i \in V} \boldsymbol{\theta}_i^T \boldsymbol{\delta}_{x_i} \\
& \text{s.t. } \boldsymbol{\delta}_i \in \{0, 1\} \quad \forall \alpha \in \{F, V\} \\
& \quad \sum_s \delta_{x_\alpha; s} = 1 \quad \forall \alpha \in \{F, V\} \\
& \quad \sum_{\mathbf{x}_f \setminus x_i} \boldsymbol{\delta}_{\mathbf{x}_f} = \boldsymbol{\delta}_{x_i} \quad \forall f \in F, x_i \in f
\end{aligned}$$

One of the methods most often used to tackle this NP-hard, combinatorial discrete optimization is linear programming. To perform this relaxation, we introduce the marginal variables  $\boldsymbol{\mu}_i$  and  $\boldsymbol{\mu}_f$ , which are constrained to the so-called *local polytope*  $\mathcal{L}(\mathcal{G})$ :

$$\mathcal{L}(\mathcal{G}) = \left\{ (\boldsymbol{\mu}_i, \boldsymbol{\mu}_f) \left| \begin{array}{l} \sum_{\mathbf{x}_f \setminus x_i} \mu_f(\mathbf{x}_f) = \mu_i(\mathbf{x}_i) \\ \sum_{x_i} \mu_i(\mathbf{x}_i) = 1 \\ \mu_i(\mathbf{x}_i) \geq 0 \wedge \mu_f(\mathbf{x}_f) \geq 0 \end{array} \right. \right\}$$

Since every assignment to  $\boldsymbol{\delta}_{x_i}$  and  $\boldsymbol{\delta}_{x_f}$  in the integer problem also satisfies the constraints of the linear program, the resulting linear programming relaxation

$$\max_{(\boldsymbol{\mu}_i, \boldsymbol{\mu}_f) \in \mathcal{L}(\mathcal{G})} \sum_{f \in F} \boldsymbol{\theta}_f^T \boldsymbol{\mu}_f + \sum_{i \in V} \boldsymbol{\theta}_i^T \boldsymbol{\mu}_i \tag{5.1}$$

is an upper bound to the original integer problem. For a more thorough treatment, see Wainwright and Jordan [2008a] or Koller and Friedman [2009b].

There are a number of different approaches to solve the LP relaxation (5.1). One approach is to solve the dual of (5.1) using coordinate descent algorithms, as in MPLP [Globerson and Jaakkola, 2007b, Werner, 2007]. Another approach is based on variants of subgradient

descent [Komodakis et al., 2011, Jojic et al., 2010]; unlike the coordinate descent algorithms, these are guaranteed to converge to a global optimum. Subgradient descent algorithms such as dual decomposition [Komodakis et al., 2011] solve the dual by separating its objective into simple local problems, and using Lagrange multipliers to force the local solutions to agree on a consensus. In practice it usually takes many iterations to reach a consensus for variables that appear in many local problems and convergence can be slow.

Using the ADMM framework to solve the LP relaxation combines the advantages of both approaches, leading to an algorithm that can converge to the global optimum in a time comparable to coordinate descent algorithms. However, there are a number of different strategies to solve the LP relaxation using ADMM framework. In the next two sections, we give an overview of the ADMM framework for convex optimization, and describe its application to the MAP LP relaxation, including two existing algorithms.

### ■ 5.3 Alternating Direction Method of Multipliers

The Alternating Direction Method of Multipliers, or ADMM, has gained recent popularity as an easy-to-use and effective technique for convex optimization [Boyd et al., 2011]. Consider an optimization over convex functions  $f$  and  $g$ :

$$\min_{x,z} f(x) + g(z) \quad s.t. \quad Ax + Bz = c \quad (5.2)$$

The ADMM algorithm uses an augmented Lagrangian,

$$L_\rho(x, z, y) = f(x) + g(z) + y^T(Ax + Bz - c) + \frac{\rho}{2} \|Ax + Bz - c\|^2 \quad (5.3)$$

to enforce the linear constraints, where  $y$  is a vector of Lagrange multipliers and  $\rho > 0$  is a quadratic penalty coefficient. The benefit of including the quadratic penalty is that the dual

function can be shown to be differentiable under rather mild conditions [Boyd et al., 2011].

The solution to (5.2) is then obtained as

$$\max_y \min_{x,z} L_\rho(x, z, y),$$

and the ADMM method provides an elegant algorithm for finding this saddle point.

ADMM performs iterative updates of the variables, using coordinate descent over the  $x$  and  $z$  and subgradient descent over the Lagrange multipliers  $y$  at each iteration  $t$ :

$$x^{(t+1)} = \arg \min L_\rho(x, z^{(t)}, y^{(t)}) \quad (5.4a)$$

$$z^{(t+1)} = \arg \min L_\rho(x^{(t+1)}, z, y^{(t)}) \quad (5.4b)$$

$$y^{(t+1)} = y^t + \rho(Ax^{(t+1)} + Bz^{(t+1)} - c) \quad (5.4c)$$

Choosing the step size equal to the quadratic penalty  $\rho$  ensures the algorithm is monotonic.

## ■ 5.4 ADMM for MAP-LP

The ADMM algorithm can be applied to the MAP-LP relaxation (5.1) in variety of ways, depending on how we define the functions  $f(x)$ ,  $g(z)$  and how the constraints are enforced.

A simple and direct application of (5.3) to the LP (5.1) is:

$$\begin{aligned} & \max_{(\boldsymbol{\mu}_i, \boldsymbol{\mu}_f) \in \mathcal{P}(\boldsymbol{\mu})} \sum_{f \in F} \boldsymbol{\theta}_f^T \boldsymbol{\mu}_f + \sum_{i \in V} \boldsymbol{\theta}_i^T \boldsymbol{\mu}_i \\ & + \sum_{\substack{f \in F \\ i \in f}} \lambda_{if}^T (\mathbf{A}_{if} \boldsymbol{\mu}_f - \boldsymbol{\mu}_i) - \sum_{\substack{f \in F \\ i \in f}} \frac{\rho}{2} \|\mathbf{A}_{if} \boldsymbol{\mu}_f - \boldsymbol{\mu}_i\|^2 \end{aligned} \quad (5.5)$$

where  $\boldsymbol{\mu}_f$ ,  $\boldsymbol{\mu}_i$  take the roles of the  $x$ ,  $z$  in ADMM, and we enforce that the  $\boldsymbol{\mu}$  ( $\boldsymbol{\mu}_i$  and  $\boldsymbol{\mu}_f$ )

live in the probability simplex  $\mathcal{P}(\boldsymbol{\mu}) = \{\boldsymbol{\mu} > 0 | \mathbf{1}^T \boldsymbol{\mu} = 1\}$ . Lagrange multipliers enforce local consistency ( $\mathcal{L}$ ) among the  $\boldsymbol{\mu}$ , and  $\mathbf{A}_{if} \boldsymbol{\mu}_f$  marginalizes  $\boldsymbol{\mu}_f$  with respect to variable  $i$ .

Unfortunately, (5.5) is difficult to optimize. Updating  $\boldsymbol{\mu}_i$  and  $\boldsymbol{\mu}_f$  at each iteration involves solving the following kind of subproblems, where  $h(x)$  is a linear function of  $x$ :

$$\min_{x \in \mathcal{P}(x)} h(x) + \frac{\rho}{2} \|Ax - w\|^2 \quad (5.6)$$

When fixing  $\boldsymbol{\mu}_f$  variables, to update  $\boldsymbol{\mu}_i$  variables, optimization (5.6) involves a quadratic term  $\sum_f \|\mathbf{w}_{if} - \boldsymbol{\mu}_i\|^2$ ; its solution can be easily computed in closed form using a partitioning technique [Duchi et al., 2008], described for completeness Section 5.4.3. However when fixing  $\boldsymbol{\mu}_i$ , optimizing for  $\boldsymbol{\mu}_f$  involves the quadratic term  $\sum_i \|\mathbf{A}_{if} \boldsymbol{\mu}_f - \mathbf{w}_i\|^2$  under the constraint  $\boldsymbol{\mu}_f \in \mathcal{P}(x)$ , for which a general closed-form solution is not easy to compute.

### ■ 5.4.1 APLP/ADLP

To overcome these difficulties, a common strategy is to introduce auxiliary variables and reformulate the optimization such that it involves solving only QPs with identity mappings at each step. The APLP algorithm [Meshi and Globerson, 2011] uses this strategy and formulates a primal MAP-LP relaxation with auxiliary variables. APLP keeps a copy  $\bar{\boldsymbol{\mu}}_{if}$  of the factor marginals  $\boldsymbol{\mu}_f$  for each variable  $i \in f$  and enforces marginalization constraints over these copies:

$$\begin{aligned} & \max_{\substack{(\boldsymbol{\mu}_i, \boldsymbol{\mu}_f) \in \mathcal{P}(\boldsymbol{\mu}) \\ \bar{\boldsymbol{\mu}}_{if}}} \sum_{f \in F} \boldsymbol{\theta}_f^T \boldsymbol{\mu}_f + \sum_{i \in V} \boldsymbol{\theta}_i^T \boldsymbol{\mu}_i \\ & + \sum_{\substack{f \in F \\ i \in f}} \boldsymbol{y}_f^T (\boldsymbol{\mu}_f - \bar{\boldsymbol{\mu}}_{if}) - \sum_{\substack{f \in F \\ i \in f}} \frac{\rho}{2} \|\boldsymbol{\mu}_f - \bar{\boldsymbol{\mu}}_{if}\|^2 \\ & + \sum_{\substack{f \in F \\ i \in f}} \boldsymbol{\lambda}_{if}^T (\boldsymbol{\mu}_i - \mathbf{A}_{if} \bar{\boldsymbol{\mu}}_{if}) - \sum_{\substack{f \in F \\ i \in f}} \frac{\rho}{2} \|\boldsymbol{\mu}_i - \mathbf{A}_{if} \bar{\boldsymbol{\mu}}_{if}\|^2 \end{aligned} \quad (5.7)$$

Then, updating  $\boldsymbol{\mu}_i$  or  $\boldsymbol{\mu}_f$  involves solving an identity-mapping QP constrained to the probability simplex; this can be done efficiently via partitioning (see section 5.4.3). Moreover,  $\bar{\boldsymbol{\mu}}_{if}$  can also be computed efficiently, since it requires inverting  $Q = I + \mathbf{A}_{if}^T \mathbf{A}_{if}$ , a block-diagonal binary matrix (its entries are zero or one) whose inverse can be computed efficiently in closed form. For more details see Meshi and Globerson [2011].

Although introducing such auxiliary variables makes each step of the algorithm more efficient, the increased number of variables in the optimization, and increased number of constraints to enforce, means that ADMM often needs more iterations to converge. This effect can be seen in our the experiments section.

Meshi and Globerson (2011) also introduced an ADLP algorithm that formulates the dual of the LP (5.1) as an ADMM optimization problem. Formulating the dual problem requires introducing fewer auxiliary variables; this tends to make ADLP significantly faster to converge than the primal APLP [Meshi and Globerson, 2011]. Because of this, in our experiments we compare to ADLP rather than APLP.

## ■ 5.4.2 DD-ADMM

Another approach to making each update of ADMM for the MAP-LP tractable is given by Martins et al. [2011] in the DD-ADMM algorithm. They formulate the primal LP as:

$$\begin{aligned} \max_{(\bar{\boldsymbol{\mu}}_i, \boldsymbol{\mu}_{if}, \boldsymbol{\mu}_f) \in \mathcal{L}(\mathcal{G})} & \sum_{f \in F} \left( \boldsymbol{\theta}_f^T \boldsymbol{\mu}_f + \sum_{i \in V} \frac{1}{d_i} \boldsymbol{\theta}_i^T \boldsymbol{\mu}_{if} \right) \\ & + \sum_{\substack{f \in F \\ i \in f}} \lambda_{if}^T (\boldsymbol{\mu}_{if} - \bar{\boldsymbol{\mu}}_i) - \sum_{\substack{f \in F \\ i \in f}} \frac{\rho}{2} \|\boldsymbol{\mu}_{if} - \bar{\boldsymbol{\mu}}_i\|^2 \end{aligned} \quad (5.8)$$

They then note that the resulting quadratic forms can be solved in closed form for two specific cases corresponding to binary-valued  $x_i$ : when the factors are pairwise (involves only two variables), or when they take on specific logical constraints, such as enforcing an

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**Algorithm 5.1** Efficient projection on to the  $l_1$  ball

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**Input:** A vector  $\mathbf{v} \in \mathbb{R}^n$  and a scalar  $z > 0$

Sort  $\mathbf{v}$  into  $\nu : \nu_1 \geq \nu_2 \geq \dots \geq \nu_p$

Find  $J$ , the largest  $j$  such that  $\nu_j - \frac{1}{j} \left( \sum_{r=1}^j \nu_r - z \right) > 0$

Define  $S = \frac{1}{J} \left( \sum_{i=1}^J \nu_i - z \right)$

**Output:**  $\mathbf{w}$  s.t.  $w_i = \max \{v_i - S, 0\}$

---

“exclusive-or”. They propose to apply this to general graphical models by “binarizing” the model, creating a binary variable for each variable and state  $x_i = s$ , and for each clique state  $x_f = (s_1 \dots s_{|f|})$ . They argue that the overhead in terms of number of factors and time per update is minimal.

However, what is not obvious is that these many inter-related variables also create many more dependencies to be enforced. Consequently, although the time per iteration is similar to coordinate descent or subgradient methods, the number of iterations required to converge may increase. In our experiments, we find this effect can be significant.

### ■ 5.4.3 Quadratic Programs and Identity Matrix

Several components of ADMM based frameworks require optimizing a quadratic form with identity matrix over a probability simplex, i.e., a Euclidean projection onto the simplex. Duchi et al. [2008] describe an efficient algorithm for this projection, which can be more formally described as:

$$\min_{\mathbf{w}} \frac{1}{2} \|\mathbf{w} - \mathbf{v}\|_2^2 \quad \text{s.t.} \quad w_i \geq 0, \quad \sum_{i=1}^n w_i = z \quad (5.9)$$

with  $z = 1$  for the probability simplex. The solution to (5.9) can be found using Algorithm 5.1.

## ■ 5.5 Linearized ADMM Algorithm

Ideally, since auxiliary variables increase the number of iterations required for convergence, we would prefer to solve the original, direct application of ADMM in (5.5). As discussed, the major obstacle in doing so is a difficult quadratic program when updating the  $\boldsymbol{\mu}_f$ . In this section, we will sidestep this difficulty using a proximal linearization technique, and derive a Linearized Augmented Primal LP (LAPLP) algorithm with fewer auxiliary variables and faster convergence.

Consider again the primal MAP LP relaxation (5.5). This leads to the following updates at each iteration:

$$\boldsymbol{\mu}_f^{(t+1)} = \arg \max_{\boldsymbol{\mu}_f \in \mathcal{P}(\boldsymbol{\mu})} \mathbf{w}_f^{(t)T} \boldsymbol{\mu}_f - \frac{\rho}{2} \boldsymbol{\mu}_f^T \mathbf{Q}_f \boldsymbol{\mu}_f \quad (5.10a)$$

$$\mathbf{w}_f^{(t)} = \boldsymbol{\theta}_f + \sum_{f:i \in f} \mathbf{A}_{if}^T (\boldsymbol{\lambda}_{if}^{(t)} + \rho \boldsymbol{\mu}_i^{(t)})$$

$$\mathbf{Q}_f = \sum_{i \in f} \mathbf{A}_{if}^T \mathbf{A}_{if}$$

$$\boldsymbol{\mu}_i^{(t+1)} = \arg \max_{\boldsymbol{\mu}_i \in \mathcal{P}(\boldsymbol{\mu})} \mathbf{w}_i^{(t+1)T} \boldsymbol{\mu}_i - \frac{\rho d_i}{2} \boldsymbol{\mu}_i^T \boldsymbol{\mu}_i \quad (5.10b)$$

$$\mathbf{w}_i^{(t+1)} = \boldsymbol{\theta}_i + \sum_{f:i \in f} \left( -\boldsymbol{\lambda}_{if}^{(t)} + \rho \mathbf{A}_{if} \boldsymbol{\mu}_f^{(t+1)} \right)$$

$$\boldsymbol{\lambda}_{if}^{(t+1)} = \boldsymbol{\lambda}_{if}^{(t)} - \rho \left( \mathbf{A}_{if} \boldsymbol{\mu}_f^{(t+1)} - \boldsymbol{\mu}_i^{(t+1)} \right) \quad (5.10c)$$

Optimization (5.10b) is a quadratic program with an identity mapping, constrained to the probability simplex, which can be solved efficiently via partitioning (see section 5.4.3). However, as discussed, computing a closed form solution to optimization (5.10a) with the non-

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**Algorithm 5.2** Linearized APLP
 

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**Input:** factor graph  $(\mathcal{G})$ , penalty parameter  $\rho$  and maximum iterations  $T$   
**Initialize**  $\lambda_{if} = 0$  for all factors  $f \in F$  and all  $i \in f$   
**Initialize**  $\mu_f = MAP(\theta_f)$  for all factors  $f \in F$   
**Initialize**  $\mu_i = MAP(\theta_i)$  for all variables  $i \in V$   
**for**  $t = 1$  **to**  $T$  **do**  
   **for each**  $f \in F$  **do**  
     Update  $\mu_f^{(t+1)} = Quad(\mu_f)$  by solving (5.12)  
     Update  $\mu_i^{(t+1)} = Quad(\mu_i)$  by solving (5.10b)  
     Update  $\lambda_{if}^{(t+1)} = \lambda_{if}^{(t)} - \rho (\mathbf{A}_{if} \mu_f^{(t+1)} - \mu_i^{(t+1)})$   
   **end for**  
**end for**

---

identity mapping  $\mathbf{Q}_f$  and linear constraints on  $\mu_f$  is not trivial. In order to find a closed form solution to optimization (5.10a) without introducing auxiliary variables, we rewrite (5.10a) as:

$$\mu_f^{(t+1)} = \arg \min_{\mu_f \in \mathcal{P}(\mu)} -\theta_f^T \mu_f + \frac{\rho}{2} \sum_{i:i \in f} \|\mathbf{A}_{if} \mu_f - \mu_i - \frac{1}{\rho} \lambda_{if}\|^2$$

The quadratic term can be approximated by a first order Taylor expansion around the current estimate, plus a proximal term (e.g., Martinet 1970, Rockafellar 1976), giving:

$$\begin{aligned} \mu_f^{(t+1)} &= \arg \min_{\mu_f \in \mathcal{P}(\mu)} -\theta_f^T \mu_f & (5.11) \\ &+ \sum_{i:i \in f} \langle \mu_f - \mu_f^{(t)}, \mathbf{A}_{if}^T (\rho (\mathbf{A}_{if} \mu_f^{(t)} - \mu_i^{(t)}) - \lambda_{if}^{(t)}) \rangle \\ &+ \sum_{i:i \in f} \frac{\rho \eta_{A_{if}}}{2} \|\mu_f - \mu_f^{(t)}\|^2 \end{aligned}$$

where  $\langle \cdot \rangle$  is the vector product and  $\eta_{A_{if}} > 0$  is a proximal coefficient that will influence the

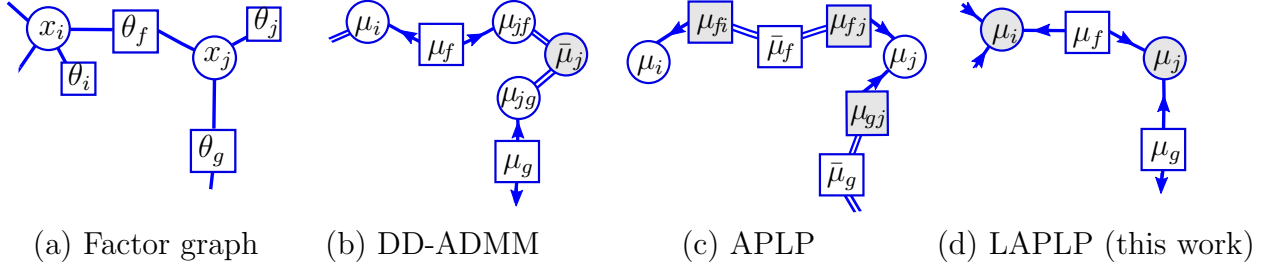


Figure 5.1: Auxiliary variables and updates in different frameworks. Double lines indicate enforced equality (an identity quadratic term); arrows indicate enforced marginal equality (a non-identity term for  $\mu_f$ 's update). ADMM alternates between updating all shaded nodes, then all unshaded nodes. (a) A portion of the original factor graph. (b) DD-ADMM binarizes each variable (not shown) and creates a copy  $\bar{\mu}_i$  of variable marginals  $\mu_i$  on which it enforces probability simplex constraints. (c) APLP creates a copy  $\mu_{fi}$  of joint marginals  $\mu_f$  for each variable  $i \in f$ , since a single outgoing non-identity marginalization constraint can be enforced in closed form. (d) Our linearized algorithm creates no copies, and linearizes the resulting non-trivial quadratic term on  $\mu_f$  due to more than one outgoing marginalization constraint (arrow).

convergence of the algorithm. Eq. (5.11) can be further simplified to

$$\begin{aligned} \boldsymbol{\mu}_f^{(t+1)} &= \arg \min_{\boldsymbol{\mu}_f \in \mathcal{P}(\boldsymbol{\mu})} \mathbf{w}_f^T \boldsymbol{\mu}_f + \frac{\rho \eta_A}{2} \|\boldsymbol{\mu}_f - \boldsymbol{\mu}_f^{(t)}\|^2 & (5.12) \\ \mathbf{w}_f &= -\boldsymbol{\theta}_f + \sum_{i:i \in f} (\rho(\mathbf{A}_{if} \boldsymbol{\mu}_f^{(t)} - \boldsymbol{\mu}_i^{(t)}) - \lambda_{if}^{(t)})^T \mathbf{A}_{if} \\ \eta_A &= \sum_{i:i \in f} \eta_{A_{if}} \end{aligned}$$

which is a QP with an identity mapping, that (as before) we can solve efficiently via partitioning. The procedure is summarized in Algorithm 5.2.

A similar linearization technique was used by Lin et al. [2011] to solve ADMM updates for a low-rank representation problem, a type of subspace clustering task. Linearization has significant advantages: it makes the auxiliary variables unnecessary, saving memory and avoiding updates to those variables. Moreover, without the extra constraints introduced by the auxiliary variables, the convergence (in terms of number of iterations) is also faster.

We illustrate the number of auxiliary variables introduced, along with the ADMM update pattern, for a small part of a factor graph in Figure 5.1. Figure 5.1(a) shows a factor graph, with variables as circles and factors as squares. Figure 5.1(b)–(d) illustrate the dependence and updates of the DD-ADMM, APLP, and LAPLP algorithms. The alternating ADMM updates are shown using shaded and unshaded nodes; squares indicate marginals over clique configurations ( $\boldsymbol{\mu}_f$ ) and circles indicate marginals over variable configurations ( $\mu_i$ ). Equality constraints are indicated using double lines, and marginalization constraints using arrows, pointing in the direction of the marginalization. The LAPLP update has significantly less variable duplication (some of the duplication of DD-ADMM is not visualized); its difficult quadratic update (5.10a) is visible as squares (e.g.,  $\boldsymbol{\mu}_f$ ) with more than one outgoing arrow.

## ■ 5.6 Performance Analysis

### ■ 5.6.1 Parameter Selection

Our linearized ADMM is guaranteed to converge to the global optimum if  $\eta_{A_{if}} \geq \|\mathbf{A}_{if}\|^2$ ; see Lin et al. [2011]. For this reason, we usually set  $\eta_A$  in (5.12) as

$$\eta_A = \|\mathbf{A}\|^2 = \sum_{i \in f} \|\mathbf{A}_{if}\|^2;$$

However, our experiments show that in practice and for the range of quadratic penalty terms  $\rho$  that are of interest, linearized ADMM converges to the global optimum even when  $\eta_A$  is set to smaller values. We compare the results for setting  $\eta_A = \|\mathbf{A}\| = (\sum_{i \in f} \|\mathbf{A}_{if}\|^2)^{\frac{1}{2}}$  and  $\eta_A = 2\|\mathbf{A}\|$  as well. Our experiments show that choosing smaller values for  $\eta_A$  results in faster convergence to global optimum. However, special care needs to be made when choosing the penalty  $\rho$  to make sure the algorithm converges to the global optimum.

Selecting the penalty parameter  $\rho$  is an important issue when using any of the ADMM based algorithms. Setting  $\rho$  very small or very large makes ADMM based algorithms slow. In our experiments we studied the effect of choosing the penalty by cross validation. To do so, we run the ADMM based algorithms on a small number of instances in a problem class using a range of penalty terms  $\rho$ , and select the best penalty on those for the remaining instances in the same class. Figure 5.3 (bottom) and Figure 5.4 (bottom) compare the relative convergence times when using this selected penalty for all problems, compared to the case where the best value of penalty is chosen for each problem independently. As the results show, the relative convergence time of ADMM based algorithms does not change in the two different settings. However, this choice of  $\rho$  slows down ADMM based algorithms compared to MPLP.

## ■ 5.6.2 Experimental Results

To evaluate our linearized augmented primal LP (LAPLP) algorithm, we compare it with the two other ADMM based algorithms for finding approximate MAP solutions, DD-ADMM by Martins et al. [2011] and ADLP by Meshi and Globerson [2011] (since it is faster than the more comparable APLP updates from the same work) as well as the coordinate descent algorithm MPLP [Globerson and Jaakkola, 2007b]. For ADLP, we use the implementation provided in the Darwin C++ framework [Gould, 2012]. For DD-ADMM, we use the code provided online by the authors<sup>1</sup>. Note that the DD-ADMM code includes some basic “message scheduling” heuristics, updating only those variables whose neighbors have changed significantly at each iteration. Since the ADLP implementation does not perform scheduling, we disabled this aspect of DD-ADMM and did not include scheduling in our own implementation of LAPLP.

We evaluate the algorithms on different sets of problems including Potts models, pedigree

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<sup>1</sup><http://www.ark.cs.cmu.edu/AD3>

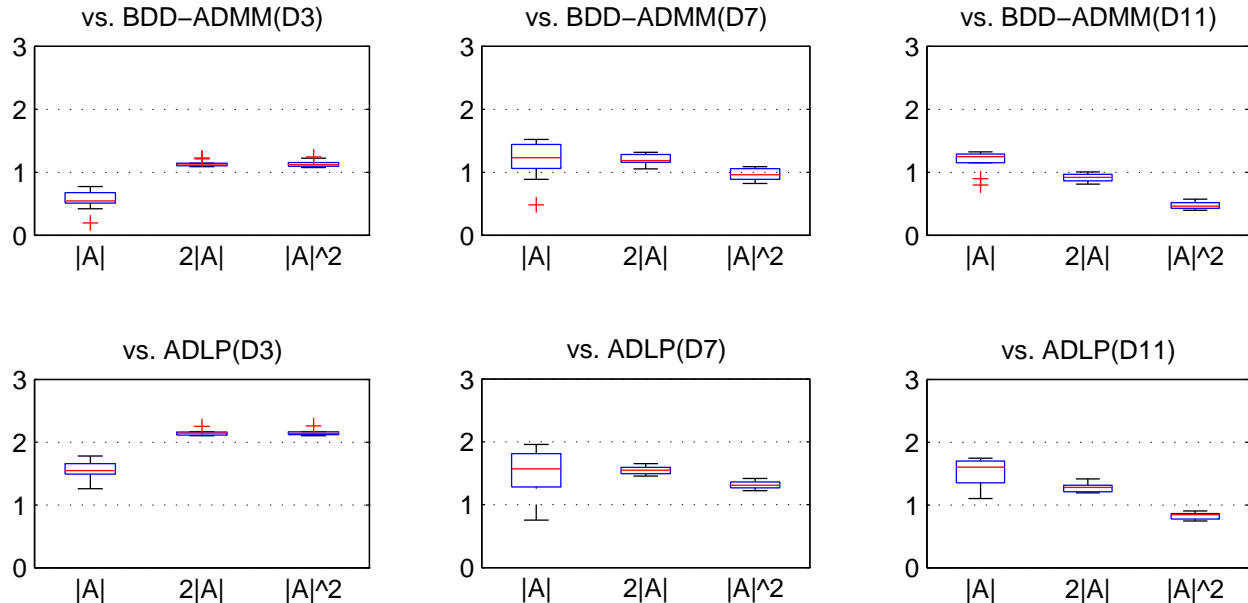


Figure 5.2: Comparison of convergence time of LAPLPL with DD-ADMM (binarized) and ADLP for different Potts models. Log relative convergence time  $-\log(t_c(\text{LAPLPL})/t_c(\text{XLP}))$  is used for comparison, where  $t_c(\text{LAPLPL})$  is the convergence time of the LAPLPL algorithm (tolerance= $1e-4$ ) and  $t_c(\text{XLP})$  is the convergence time of XLP. Here XLP represents any of the algorithms DD-ADMM, ADLP, or MPLP. MPLP converges to local optimum in these experiments.

trees and protein side-chain prediction. To compare different methods we use relative convergence time  $-\log(t_c(\text{LAPLPL})/t_c(\text{XLP}))$ , where  $t_c(\text{LAPLPL})$  is the convergence time of LAPLPL algorithm (tolerance= $1e-4$ ) and  $t_c(\text{XLP})$  is the convergence time of XLP algorithm, where XLP is one of DD-ADMM, ADLP or MPLP.

**Potts Models** To compare different algorithms on Potts models, we generated 20x20 Potts models with single node log-potentials chosen as  $\theta_i(x_i) \sim \mathcal{U}[-1, 1]$  and edge log potentials as  $\theta_{i,j}(x_i, x_j) \sim \mathcal{U}[-5, 5]$  if  $x_i \neq x_j$  and 0 otherwise. We generated models with different variable cardinalities (3, 7 and 11) to study the effect of model size on different algorithms.

Figure 5.2 compares the convergence time of LAPLPL to DD-ADMM (binarized) and ADLP, averaged over 10 models of the same size (models with multi-valued variables with 3, 7

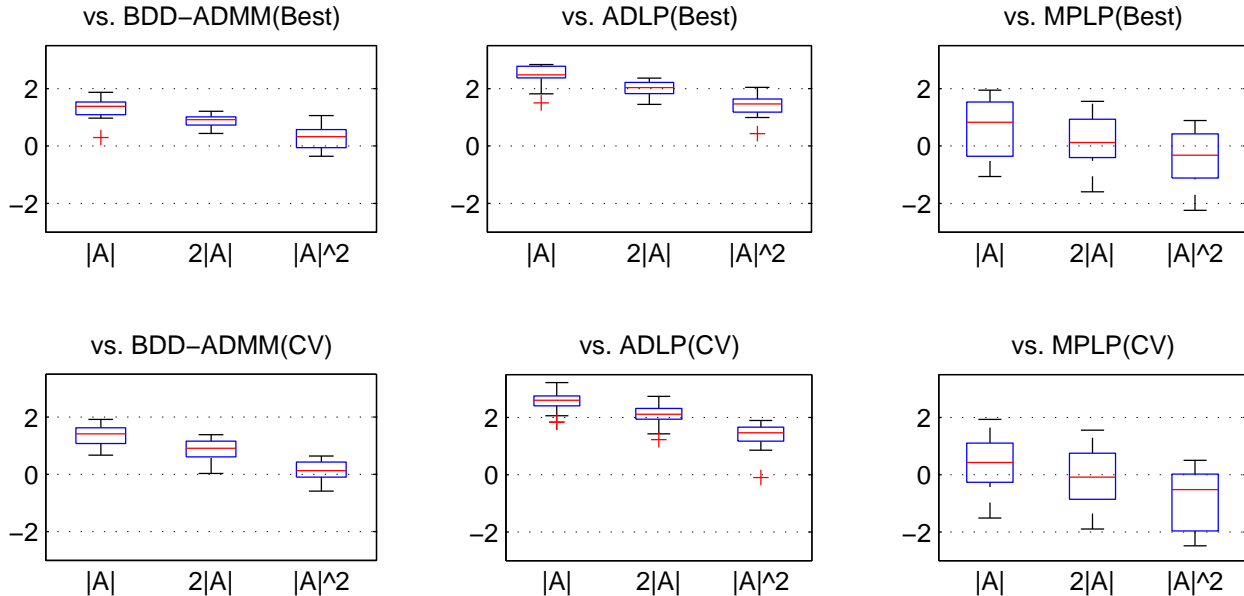


Figure 5.3: Comparison of convergence time of LAPLP with DD-ADMM (binarized) and ADLP on pedigree trees when using the best penalty  $\rho$  for each model (top) and choosing the penalty  $\rho$  using cross validation. Log relative convergence time  $-\log(t_c(\text{LAPLP})/t_c(\text{XLP}))$  is used for comparison, where  $t_c(\text{LAPLP})$  is the convergence time of the LAPLP algorithm (tolerance= $1e-4$ ) and  $t_c(\text{XLP})$  is the convergence time of XLP. Here XLP represents any of the algorithms DD-ADMM, ADLP, or MPLP.

and 11 different values respectively). As can be seen LAPLP is faster than both ADLP and DD-ADMM. Its important to note that MPLP often converges to local optima in these experiments, while ADMM based algorithms are able to find the global optimum.

**Pedigree Models** We also compared the algorithms on pedigree models from UAI 2008 biological linkage analysis problems. These models involve non-pairwise factors with variables that have cardinalities between 2 and 7. Of the total 19 pedigree models, MPLP converged to local optima in 3 experiments. Figure 5.3 (top) compares the convergence time of LAPLP to DD-ADMM, ADLP and MPLP. As shown here, MPLP can converge faster than ADMM based algorithms in some models but it has the potential to converge to local optima. As in our other experiments, the linearized ADMM converges faster than existing globally convergent approaches. This shows that linearizing the quadratic term helps improve convergence

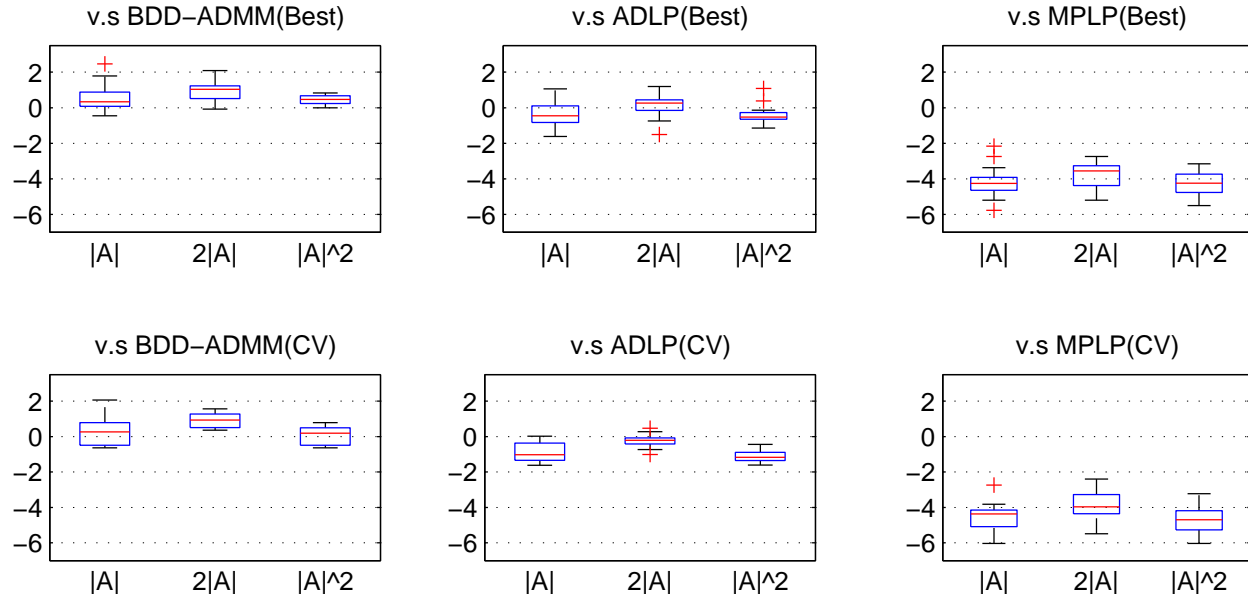


Figure 5.4: Comparison of convergence time of LAPLPL with DD-ADMM(binarized) and ADLP on protein side-chain prediction when using the best penalty  $\rho$  for each model (top) and choosing the penalty  $\rho$  using cross validation. Log relative convergence time  $-\log(t_c(\text{LAPLPL})/t_c(\text{XLP}))$  is used for comparison, where  $t_c(\text{LAPLPL})$  is the convergence time of the LAPLPL algorithm (tolerance=1e-4) and  $t_c(\text{XLP})$  is the convergence time of XLP. Here XLP represents any of the algorithms DD-ADMM, ADLP, or MPLP.

time by avoiding the introduction of auxiliary variables and their corresponding constraints.

**Protein Side-chain Prediction** Finally we evaluate the algorithms on protein side-chain prediction problems from Yanover and Weiss [2003] and Yanover et al. [2006]<sup>2</sup>. We use the set of “large” model instances, containing 20 problems of between 300 – 1000 amino acids (variables), each with 2 – 81 possible states (average cardinality  $\approx 20$ ) and pairwise potential functions. These results show that ADLP’s convergence time is less affected by model size compared to DD-ADMM (binarized) and LAPLPL, and convergence time of the three algorithms are comparable in half of the experiments. MPLP convergence time is much faster when it finds the global optimum. The results of these experiments are summarized in Figure 5.4 (top).

<sup>2</sup><http://cyanover.fhcrc.org/proteinMRFs.html>

Figure 5.5 compares the behavior of LAPLP, ADLP and DD-ADMM across the three sets of problems. Since different models have different energy values and times to convergence, to plot average performance we compute the normalized energy for each algorithm, consisting of the percentage increase in energy over the optimal value of the LP at convergence, and plot it against the percentage of time used compared to the convergence time for our LAPLP method on that model.

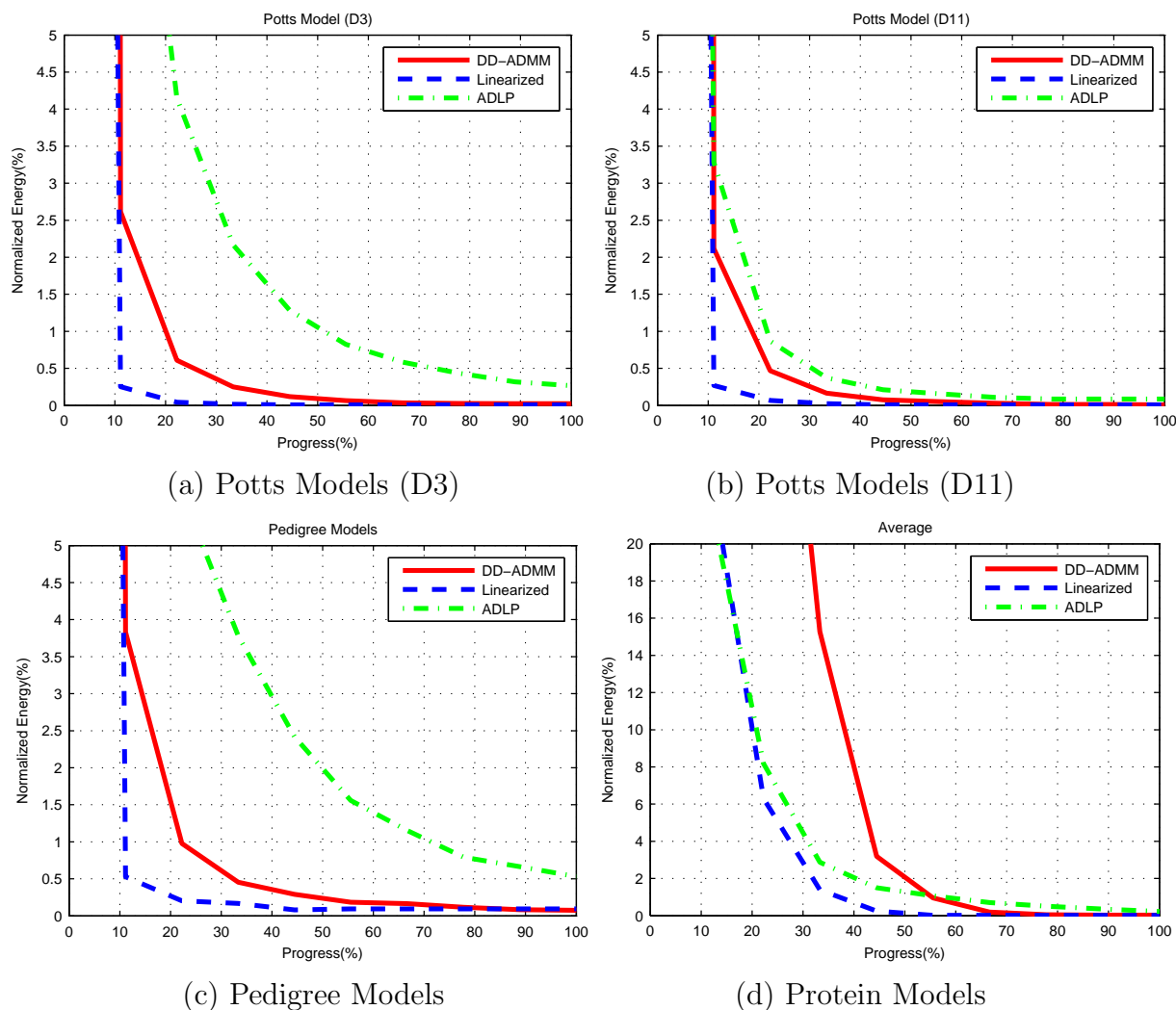


Figure 5.5: Comparing average run time of different ADMM algorithms. We show the percent increase in energy over the optimal LP value, relative to the percentage of time to our LAPLP algorithms convergence, averaged across problem instances. Here LAPLP is fastest, closely followed by ADLP, with the binarized DD-ADMM slower for much of the runtime but catching up near the end.

## ■ 5.7 Discussion

In this chapter, we presented an algorithm based on the Alternating Direction Method of Multipliers (ADMM) for approximate MAP inference using its linear programming relaxation. Our algorithm is based on augmenting the primal MAP-LP with a quadratic term that enforces strict convexity of the Lagrangian, and solving this quadratic form by linearization with an additional proximal term. Importantly, we find that performing such approximate solutions does not significantly affect the convergence time of the ADMM algorithm. We compared our algorithm with two existing ADMM-based algorithms on Potts models, pedigree trees and protein side-chain prediction problems for approximate MAP inference, showing that our linearized primal MAP-LP algorithm can solve MAP inference faster than methods based on auxiliary variables in models with non-binary variables. We also showed that a cross validation procedure can be used to choose the penalty term  $\rho$  for a problem class.

Several practical improvements can be considered over our basic algorithm. One is to use an adaptive penalty parameter  $\rho$ , which may improve convergence in practice. However, the theoretical convergence guarantees of ADMM may no longer hold. Another potential improvement is to use a scheduling method [Elidan et al., 2006, Tarlow et al., 2011] to select which sub-problems to solve during each iteration of ADMM. As a simple example, we need only solve each local sub-problem  $\mu_f$  if some neighboring consensus variable  $\mu_i$  has been changed at the previous iteration, since otherwise the previous results can be simply re-used.

# Conclusions and Future Directions

Despite the fact that exact inference in graphical models is fundamentally difficult, the ubiquity and importance of graphical models for knowledge representation and reasoning in a wide variety of domains has meant an increasing reliance on approximate inference algorithms. In this thesis, we have focused on several aspects of approximate inference, and in particular, variational bounding techniques, that affect the quality of the approximation.

In Chapter 3, we focused on finding better regions, a key component of many approximate inference bounds. Mini-bucket elimination avoids the space and time complexity of exact inference by using a top-down partitioning approach that mimics the construction of a junction tree and aims to minimize the number of regions subject to a bound on their size; however, standard mini-bucket approaches rarely take into account the functions' values. In contrast, message passing algorithms often use "cluster pursuit" methods to select regions, a bottom-up approach in which a pre-defined set of clusters (such as triplets) is scored and incrementally added. We developed a hybrid approach that balances the advantages of both perspectives, providing larger regions chosen in an intelligent, energy-based way, by defining a scoring function that computes a local estimate of the bound's improvement to select better regions. We combined our proposed scoring function with the message passing framework of weighted mini-bucket elimination in order to better estimate the true impact of the regions on the approximation. Finally, we proposed an efficient structure update procedure that

incrementally updates the join graph of mini-bucket elimination after new regions are added in order to avoid starting from scratch after each merge.

In Chapter 4, we studied how efficient use of available memory can improve the quality of inference. We described how controlling the complexity of inference using *ibound* can result in an inefficient use of resources and proposed memory-aware alternatives. By using our incremental construction of the join graph, we proposed to track the memory requirement of the approximation as it is built; we then extended this framework to use a more flexible set of controls on the join graph complexity, expressed in terms of a single or set of memory budgets, and proposed a number of ways of setting the initial memory budgets and of re-allocating during construction. Together, these give a more fine-grained control over the approximation complexity than a single *ibound* parameter. We showed experimentally that using an allocation technique that first distributes the memory between different buckets proportionally to the scope-based MBE construction, and then shifts any extra memory along the elimination order, can use the available memory more efficiently and give tighter bounds by allowing larger regions to be added to the approximation.

In Chapter 5, we focused on maximum a *posteriori* (MAP) inference and its linear programming (LP) relaxation, a commonly used and successful class of approximate inference algorithms. We discussed how the augmented Lagrangian method can be used to overcome a lack of strict convexity in LP relaxations, and how the Alternating Direction Method of Multipliers (ADMM) provides an elegant algorithm for finding the saddle point of the augmented Lagrangian. We characterized different formulations of the ADMM-based algorithm using a graphical approach, discussed the challenges and presented an ADMM-based algorithm to solve the primal form of the MAP-LP, using closed form updates based on a linear approximation technique. We showed how our technique’s efficient, closed form updates converge to the global optimum of the LP relaxation and compared our algorithm to two existing ADMM-based MAP-LP methods, showing that our technique is faster on general,

non-binary or non-pairwise models.

To conclude, in this thesis we have focused on several problems related to the quality of approximate inference algorithms in graphical models, including choosing better-quality regions for the approximation, using the available memory more efficiently, and finally using more efficient optimization algorithms. In light of our results, several interesting directions are opened for future research, discussed next.

**Models with determinism.** Budget based memory allocation for content-based WMBE can be particularly useful when reasoning about models with a significant amount of determinism. In these models, particular assignments to variables have zero probability; in this setting, a sparse representation of factors (keeping track of only non-zero entries, rather than the full table) may offer advantages. For these models, scope-based partitioning methods, and complexity control using an *ibound* parameter, do not provide a realistic estimate of the memory required by the model. With content-based WMBE however, since we are using the function values to decide which regions to add to the approximation, we can also evaluate the memory required by a merge, taking into account the amount of determinism present in the factors. Budget-based memory allocation schemes are then also applicable, and by providing a more accurate estimate of the required memory, could allow more or larger regions to be added to the approximation. However, the amount of determinism in the models directly affects the potential improvement from a sparse representation of factors and requires further study.

**Memory-aware region selection.** Another potential improvement to the content-based memory-aware WMBE comes by defining scoring functions that also take into account the amount of memory used by each merge. Our current framework uses an estimate of the memory required by a merge only to decide if a merge should be scored or not; but several “small” merges could be more useful than a single “large” merge. A very simple way to

include the memory used by a merge into the scoring function would be to use it as a tiebreaker when two different merge choices have the same potential improvement to the bound. More sophisticated methods of including such information to score and select regions is a potential future direction for research.

**Anytime bounds.** Finally, it is important to look at the anytime behavior of different approaches. The original mini-bucket elimination is a one-time, non iterative approach, and does not provide an upper bound until it is done with all possible merges, and used an amount of memory corresponding to its final size. While an anytime version can be constructed by simply building mini-buckets of increasing *ibound*, this is inefficient – the previous bound is discarded at each step, and much of the work may be repeated when constructing the next bound. On the other hand, our incremental framework sidesteps this limitation; it naturally allow us to compute an approximation early (corresponding to a low *ibound*), and then continues to provide steadily improving bound values as the region merging process continues. An important question is then how to balance between message passing (improving the bound for a fixed set of regions), and merging (tightening the bound, but resulting in more computational complexity). In particular, how we can rapidly move from one set of regions to the next in a way that results in good anytime behavior? Again, the significant diversity of models across different domains requires extensive study to develop a robust procedure.

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