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# 15 Lifted First-Order Probabilistic Inference

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Most probabilistic inference algorithms are specified and processed on a propositional level, even though many domains are better represented by first-order specifications that compactly stand for a class of propositional instantiations. In the last fifteen years, many algorithms accepting first-order specifications have been proposed. However, these algorithms still perform inference on a mostly propositional model, generated by the instantiation of first-order constructs. When this is done, the rich and useful first-order structure is not explicit anymore. This first-order representation and structure allow us to perform *lifted* inference, that is, inference on the first-order representation directly, manipulating not only individuals but also groups of individuals. This has the potential of greatly speeding up inference. We precisely define the problem and present an algorithm that generalizes variable elimination and manipulates first-order representations in order to perform lifted inference.

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## 15.1 Introduction

Probabilistic inference algorithms are widely employed in artificial intelligence. Among those, graphical models such as Bayesian and Markov networks (BNs and MNs respectively) ([8]) are among the most popular. These models are specified by a set of conditional probabilities (for BNs) or factors, also called potential functions (for MNs). Both conditional probabilities and factors are defined over particular subsets of the available random variables, and map assignments of those random variables to positive real numbers (called *potentials* in MNs). For our purposes, it will be helpful to think of graphical models in general and simply consider conditional probabilities as a type of factor.

For example, in an application for document subject classification, one can specify a dependence between the random variables *subject\_apple*, *word\_mac* (which

indicate that the subject of the document is “apple” and that the word “mac” is present in it) by defining a factor on their assignments. The higher the potential for a given assignment to these random variables, the more likely it will be in the joint distribution defined by the model.

A limitation of graphical models arises when the same dependence holds between different subsets of random variables. For example, we might declare the dependence above to hold also between *subject\_microsoft*, *word\_windows*. In traditional graphical models, we must use separate potential functions to do so, even though the dependence is the same. This brings redundancy to the model and possibly wasted computation. It is also an ad hoc mechanism since it does not cover other sets of random variables exhibiting the same dependence (in this case, some other company and product).

The root of this limitation is that graphical models are *propositional* (random variables can be seen as analogous to propositions in logic), that is, they do not allow quantifiers and parameterization of random variables by objects. A *first-order* or *relational* language, on the other hand, does allow for these elements. With such a language, we can specify a potential function that applies, for example, to *all* tuples of random variables obtained by instantiating  $X$  and  $Y$  in the tuple

$$subject(X), company(X), product(X, Y), word(Y). \quad (15.1)$$

This way we not only cover both cases presented before, but also unforeseen ones, with a single compact specification.

In the last fifteen years, many proposals for probabilistic inference algorithms accepting first-order specifications have been presented ([7, 6, 1, 4, 10, 11], among many others), most of which based on the theoretic framework of Halpern [5]. However, these solutions still perform inference at a mostly propositional level; they typically instantiate potential functions according to the objects relevant to the present query, thus obtaining a regular graphical model on propositional random variables, and then using a regular inference algorithm on this model. In domains with a large number of objects this may be both costly and essentially unnecessary. Suppose we have a medical application about the health of a large population, with a random variable per person indicating whether they are sick with a certain disease, and with a potential function representing the dependence between a person being sick and that person getting hospitalized. To answer the query “what is the probability that *someone* will be hospitalized?”, an algorithm that depends on propositionalization will instantiate *a random variable per person*. However this is not necessary since one can calculate the same probability by reasoning about individuals on a general level, simply using the population size, in order to answer that query in a much shorter time. In fact, the latter calculation would not depend on the population size at all.

Naturally, it is possible to reformulate the problem so that it is solved in a more efficient manner. However, this would require manual devising of a process *specific* to the model or query in question. It is desirable to have an algorithm that

can receive a *general* first-order model and *automatically* answer queries like these without computational waste.

A first step in this direction was given by Poole [9], which proposes a generalized version of the variable elimination algorithm [12] that is *lifted*, that is, deals with groups of random variables at a first-order level. The algorithm receives a specification in which parameterized random variables stand for all of their instantiations and then eliminates them in a way that is equivalent to, but much cheaper than, eliminating all their instantiations at once. For the parameterized potential function (15.1), for example, one can eliminate  $product(X, Y)$  in a single step that would be equivalent to eliminating all of its instantiations.

The algorithm in Poole [9], however, applies only to certain types of models because it uses a single elimination operation that can only eliminate parameterized random variables containing all parameters present in the potential function (the method can eliminate  $product(X, Y)$  from (15.1) but not  $company(X)$  because the latter does not contain the parameter  $Y$ ). As we will see later, Poole's algorithm uses the operation we call *inversion elimination*. In addition to inversion elimination, we have developed further operations (the main ones called *counting elimination* and *partial inversion*) that broaden the applicability of lifted inference to a greater extent ([2, 3]). These operations are combined to form the first-order variable elimination (FOVE) algorithm presented in this chapter. The cases to which lifted inference applies can be roughly summarized as those containing dependencies where the set of parameters of each parameterized random variable are disjoint or, when this is not the case, where there is a set of parameters whose instantiations create independent solvable cases. We specify these conditions in more detail when explaining the operations, and further discuss applicability in section 15.6. When no lifted inference operation applies to a specific part of a model, FOVE can still apply standard propositional methods to that part, assuring completeness and limiting propositional inference to only some parts of the model.

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## 15.2 Language, Semantics and Inference problem

Like Markov networks, first-order probabilistic models (FOPMs) are essentially defined by a set of factors. However, unlike them, these factors are defined over *parameterized* random variables, and for this reason we call them **parfactors** (following [9]).

Given a universe of objects over which the parameters range, we can generate regular propositional factors from a parfactor by replacing its parameters by particular objects. A parfactor is therefore a compact representation of a set of regular factors, and a FOPM is a compact representation of a Markov network composed by all instantiations of all of its parfactors.

Based on the correspondence to logic concepts, we call a parameterized random variable an **atom**, and a parameter a **logical variable** (as opposed to *random* variables). We also call the functors of atoms **predicates**. Even though we infor-

mally refer to atoms as “parameterized random variables”, they are not, technically speaking, random variables, but stand for classes of them. A ground atom, however, denotes a random variable. Sometimes we call random variables **ground** to emphasize their correspondence to ground atoms.

Logical variables are typed, with each type being a finite set of objects. We denote the **domain**, or **type**, of a logical variable  $X$  by  $D_X$  and its cardinality by  $|X|$ . In our examples, unless noted, all logical variables have the same type. Each predicate  $p$  also has its domain,  $D_p$ , which is the set of values that each of the random variables with that predicate can take.

Formally, a **parfactor**  $g$  is a tuple  $(\phi_g, A_g, C_g)$ , where  $\phi_g$  is a potential function defined over atoms  $A_g$  to be instantiated by all substitutions of its logical variables satisfying a constraint  $C_g$ . A **constraint** is a pair  $(F, V)$  where  $F$  is an equational formula on logical variables and  $V$  is the set of logical variables to be instantiated (some of them may not be in the formula). We sometimes denote a constraint by its formula  $F$  alone, when the set of logical variables  $V$  is clear from context. Tautological formulas are represented by  $\top$ . For example, the parfactor  $(\phi, (p(X), q(X, Y)), (X \neq a, \{X, Y\}))$  applies  $\phi$  to all instantiations of  $(p(X), q(X, Y))$  by substitutions of  $X$  and  $Y$  satisfying  $X \neq a$ . We denote the set of substitutions satisfying  $C$  by  $[C]$ .

While we are neutral as to how the potential functions are actually specified, logical formulas seem to be a convenient choice. For example, a weighted formula  $0.7 : \text{epidemic}(D) \Rightarrow \text{sick}(P, D)$  might represent a potential function  $\phi(\text{epidemic}(D), \text{sick}(P, D))$  with potential 0.7 for assignments in which the formula is true. This allows us to specify FOPMs by sets of weighted logical formulas that are intuitive and simple to read, and is the approach taken by Markov logic networks ([11]).

The **projection**  $C|_L$  of a constraint  $C = (F, V)$  **onto a set of logical variables**  $L$  is a constraint equivalent to  $(\exists L' F, L)$  for  $L' = V \setminus L$ . Intuitively,  $C|_L$  describes the conditions posed by  $C$  on  $L$  alone, that is, the possible substitutions on  $L$  that are part of substitutions in  $[C]$ . For example,  $(X \neq a \wedge X \neq Y \wedge Y \neq b, \{X, Y\})|_{\{X\}} = (X \neq a, \{X\})$ . FOVE uses a constraint solver which is able to solve several constraint problems, such as determining the number of solutions of a constraint and its projection onto sets of logical variables.

In certain contexts we wish to describe the class of random variables instantiated from an atom with constraints on its logical variables (for example, the set of random variables instantiated from  $p(X, Y)$ , with  $X \neq a$ ). We call such pairs of atoms and constraints **constrained atoms**, or **c-atoms**. The **c-atoms of a parfactor** is the set of c-atoms formed by its atoms and its constraint.

Let  $\alpha$  be a parfactor, c-atom, constraint or a set of those. We define  $RV(\alpha)$  to be the set of (ground) random variables specified by  $\alpha$ , and  $\alpha\theta$  denotes the result of applying a substitution  $\theta$  to  $\alpha$ .  $[C_g]$  is also denoted by  $\Theta_g$ .

A FOPM is specified by a set of parfactors  $G$  and the types of its logical variables. Its semantics is a joint distribution defined on  $RV(G)$  by the Markov network formed by all the instantiations of parfactors. Thus it is proportional to the product

of all instantiated parfactors:

$$P(RV(G)) \propto \prod_{g \in G} \prod_{\theta \in \Theta_g} g\theta.$$

For convenience, we denote  $\prod_{\theta \in \Theta_g} g\theta$  by  $\Phi(g)$ , and  $\prod_{g \in G} \Phi(g)$  by  $\Phi(G)$ . Therefore we can write the above as  $P(RV(G)) \propto \Phi(G)$ .

The most important inference task in graphical models is marginalization. For FOPMs, it takes the following form: given a set of ground random variables  $Q$ , calculate

$$P(Q) \propto \sum_{RV(G) \setminus Q} \Phi(G), \quad (15.2)$$

where the summation ranges over all assignments to  $RV(G) \setminus Q$ . Posterior probabilities can be calculated by representing evidence as additional parfactors on the evidence atoms.

The FOVE algorithm makes the simplifying assumption that the FOPM is **shattered** w.r.t the query  $Q$ . A set of c-atoms is **shattered** if the instantiations of any pair of its elements are either identical or disjoint. A parfactor, or set of parfactors, is shattered if the set of their c-atoms is shattered. A FOPM is shattered w.r.t. a query  $Q$  if the union of its c-atoms and those of the query is shattered. For example, we can have c-atoms  $(p(X), X \neq a), (p(Y), Y \neq a)$  and  $p(a)$  in a model, but not  $p(Y)$  and  $p(a)$ , because  $RV(p(a)) \subset RV(p(Y))$  but  $RV(p(a)) \neq RV(p(Y))$ . When a FOPM and query are not shattered, we can replace them by equivalent shattered FOPM and query through the process of **shattering**, detailed in section 15.5.2.

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### 15.3 The First-Order Variable Elimination (FOVE) algorithm

Computing (15.2) directly is intractable since it would take exponential time in the number of random variables in  $RV(G) \setminus Q$ . This is the case even for the propositional case, which is the reason why algorithms have been developed that take advantage of independences represented in the model in order to compute marginals more efficiently. One of these algorithms is variable elimination (VE) [12]. First-order variable elimination (FOVE) is a first-order generalization of VE. While VE eliminates a random variable at a time, FOVE eliminates a c-atom, or set of c-atoms, at each step. By eliminating a c-atom, we implicitly eliminate all of its instantiations at the same time. Let  $E$  be a set of c-atoms to be eliminated from a FOPM with a set  $G$  of parfactors. Let  $G_E, G_{\neg E} \subseteq G$  be the sets of parfactors depending and not depending on  $E$ , respectively. Then

$$\sum_{RV(G) \setminus Q} \Phi(G) = \sum_{(RV(G) \setminus RV(E)) \setminus Q} \Phi(G_{\neg E}) \sum_{RV(E)} \Phi(G_E).$$

We later show operations computing a parfactor  $g'$  such that  $\sum_{RV(E)} \Phi(G_E) = \Phi(g')$ . Once we have  $g'$ , the right-hand side of the above is equal to

$$\sum_{(RV(G) \setminus RV(E)) \setminus Q} \Phi(G_{-E}) \Phi(g') = \sum_{(RV(G) \setminus RV(E)) \setminus Q} \Phi(G_{-E} \cup \{g'\}) = \sum_{RV(G') \setminus Q} \Phi(G')$$

where  $G' = G_{-E} \cup \{g'\}$ . In other words, we have reduced the original marginalization to a smaller instance that does not include  $RV(E)$ . This is repeated until only  $Q$  is left.

A crucial difference between VE and FOVE is elimination ordering. VE eliminates random variables according to an ordering given a priori. In FOVE, eliminating certain c-atoms may require eliminating some other c-atoms first, so it may be the case that some c-atoms are not eliminable at all times (these conditions will be clarified later). Because parfactors and c-atoms are sometimes changed and reorganized during the algorithm, it is not a simple matter to choose an ordering in advance. Instead, the elimination ordering is dynamically determined.

Before we move on to explaining the operations for calculating  $\sum_{RV(E)} \Phi(G_E) = \Phi(g')$ , we mention that, in fact, they only calculate  $\sum_{RV(E)} \Phi(g)$  for a single parfactor  $g$ . This is not a problem because the operation of **fusion**, covered in section 15.5.1, calculates  $g$  such that  $\Phi(g) = \Phi(G)$  for any set of parfactors  $G$ .

### 15.3.1 Counting Elimination

We first show counting elimination on a specific example and later generalize it. Consider the summation

$$\sum_{RV(p(X))} \prod_{X,Y} \phi(p(X), p(Y)),$$

where  $p$  is a boolean predicate. (Note that the  $X$  used under the summation is not the same  $X$  used by the product.  $RV(p(X))$  is shorthand for all assignments over the set  $\{p(X) : X \in D_X\}$ , so  $X$  is locally used. In fact, we could have written  $RV(p(Y))$ , or even  $RV(p(Z))$ , to the same effect. We choose to use  $X$  or  $Y$  to make the link with the atom in the parfactor more obvious.)

Counting elimination is based on the following insight: because a parfactor will typically only evaluate to a few different potentials, large groups of its instantiations will evaluate to the same potential. So the summation is rewritten

$$\sum_{RV(p(X))} \phi(0,0)^{|(0,0)|} \phi(0,1)^{|(0,1)|} \phi(1,0)^{|(1,0)|} \phi(1,1)^{|(1,1)|},$$

where  $|v_1, v_2|$  indicates the number of possible choices for  $X$  and  $Y$  so that  $p(X) = v_1$  and  $p(Y) = v_2$  given the current assignment to  $RV(p(X))$ . These partition sizes can be calculated by a combinatorial, or counting, argument. Assume we know  $\vec{N}_p$ , a vector of integers that indicates how many random variables in  $RV(p(X))$  are currently assigned a particular value, that is,  $\vec{N}_{p,i} = |\{r \in RV(p(X)) : r = i\}|$

for each  $i \in D_p$ . Naturally,  $\sum_i \vec{N}_{p,i} = |RV(p(X))|$ . Then there are  $\vec{N}_{p,v_1}$  possible values for  $X$  (so that  $p(X) = v_1$ ) and  $\vec{N}_{p,v_2}$  distinct possible values for  $Y$  (so that  $p(Y) = v_2$ ), so  $|(v_1, v_2)| = \vec{N}_{p,v_1} \vec{N}_{p,v_2}$ .

We take advantage of the fact that the values  $|(v_1, v_2)|$  do not depend on the particular assignments to  $RV(p(X))$ , but only on  $\vec{N}_p$ . This allows us to iterate over the *groups* of assignments with the same  $\vec{N}_p$  and do the calculation for the entire group. We also take into account the group size, which is provided by the binomial coefficient of  $\vec{N}_p$ ,  $\binom{|RV(p(X))|}{\vec{N}_{p,0}}$  (or, equivalently,  $\binom{|RV(p(X))|}{\vec{N}_{p,1}}$ ). We then have

$$\sum_{\vec{N}_p} \binom{|RV(p(X))|}{\vec{N}_{p,0}} \prod_{(v_1, v_2)} \phi(v_1, v_2)^{\vec{N}_{p,v_1} \vec{N}_{p,v_2}}$$

which has a number of terms linear in  $|RV(p(X))|$ , as opposed to the previous exponential number.

Counting elimination is not a universal method. The counting argument presented above requires that there be little interaction between the logical variables of atoms. If a parfactor is on  $p(X, Y), q(X, Z)$ , for example, the counting argument does not work because the choices for  $(X, Z)$  depend on the particular  $X$  chosen for  $p(X)$ ; we can no longer compute number of choices using counters alone but need to know the particular assignment to  $RV(p(X))$ . Generally, under counting elimination, choices for one atom cannot constrain the choices for another atom (there are exceptions to this rule, as for example *just-different* atoms, presented in [3]).

We now give the formal account of counting elimination, starting with some preliminary definitions.

First, we define the notion of **independent atoms given a constraint**. Intuitively, this happens when choosing a substitution for the logical variables of one atom does not change the possible choices of substitutions for the other atom. Let  $\bar{X}_1$  and  $\bar{X}_2$  be two sets of logical variables such that  $\bar{X}_1 \cup \bar{X}_2 \subseteq V$ .  $\bar{X}_1$  is *independent from  $\bar{X}_2$  given  $C$*  if, for any substitution  $\theta_2 \in [C]_{\bar{X}_2}$ ,  $C|_{\bar{X}_1} \Leftrightarrow (C\theta_2)|_{\bar{X}_1}$ .  $\bar{X}_1$  and  $\bar{X}_2$  are *independent given  $C$*  if  $\bar{X}_1$  is independent from  $\bar{X}_2$  given  $C$  and vice-versa. Two *atoms*  $p_1(\bar{X}_1)$  and  $p_2(\bar{X}_2)$  are *independent given  $C$*  if  $\bar{X}_1$  and  $\bar{X}_2$  are independent given  $C$ .

Finally, we define **multinomial counters**. Let  $a$  be a c-atom with domain  $D_a$ . Then the *multinomial counter* of  $a$ ,  $\vec{N}_a$ , is a vector where  $\vec{N}_{a,j}$  indicates how many instantiations of  $a$  are assigned the  $j$ -th value in  $D_a$ . The *multinomial coefficient*  $\vec{N}_a! = \frac{(\vec{N}_{a,1} + \dots + \vec{N}_{a,|D_a|})!}{\vec{N}_{a,1}! \dots \vec{N}_{a,|D_a|}!}$  is a generalization of binomial coefficients and indicates how many assignments to  $RV(a)$  exhibit the particular value distribution counted by  $\vec{N}_a$ .

Counters can be applied to *sets* of c-atoms with the same general meaning. The set of multinomial counters for a set of c-atoms  $A$  is denoted  $\vec{N}_A$ , and the product  $\prod_{a \in A} \vec{N}_a!$  of their multinomial coefficients is denoted  $\vec{N}_A!$ .

**Theorem 15.1 Counting Elimination**

Let  $g$  be a shattered parfactor and  $E = \{E_1, \dots, E_k\}$  be a subset of  $A_g$  such that  $RV(E)$  is disjoint from  $RV(A_g \setminus E)$ ,  $A' = A_g \setminus E$  are all ground, and where each pair of atoms is independent given  $C_g$ . Then

$$\sum_{RV(E)} \prod_{\theta \in \Theta_g} \phi(A_g \theta) = \sum_{\vec{N}_E} \vec{N}_E! \prod_{v \in D_E} \phi(v, A')^{\prod_{i=1}^k \vec{N}_{E_i, v_i}}.$$

The theorem's proof reflects the argument given above. Counting elimination brings a significant computational advantage because iterating over assignments is exponential in  $|RV(E)|$  while doing so over groups of assignments is only polynomial in it.

It is important to notice that  $E$  must contain all non ground c-atoms in  $g$ . Also, if all c-atoms in  $g$  are ground,  $E$  can be any subset of them and we will have a simple propositional summation, the same used in VE (counters over 1-random variable c-atoms reduce to ordinary assignments).

**15.3.2 Inversion**

Counting elimination requires a parfactor's atoms to be independent given its constraints. In particular, logical variables shared between atoms may render them dependent on each other. In some of these cases, the operation of *inversion* can be applied. In fact, even in cases in which counting elimination can be applied, it is advantageous to apply inversion first, if possible, for efficiency reasons.

Let us consider a couple of examples before we formalize inversion. Consider the following:

$$\begin{aligned} & \sum_{RV(q(X,Y))} \prod_{XY} \phi(p(X), q(X, Y)) \\ &= \sum_{q(o_1, o_1)} \sum_{q(o_1, o_2)} \cdots \sum_{q(o_n, o_n)} \phi(p(o_1), q(o_1, o_1)) \phi(p(o_1), q(o_1, o_2)) \cdots \phi(p(o_n), q(o_n, o_n)) \end{aligned}$$

(by observing that  $\phi(p(o_i), q(o_i, o_j))$  depends on  $q(o_i, o_j)$  only)

$$= \sum_{q(o_1, o_1)} \phi(p(o_1), q(o_1, o_1)) \cdots \sum_{q(o_n, o_n)} \phi(p(o_n), q(o_n, o_n))$$

(by observing that only  $\phi(p(o_i), q(o_i, o_j))$  depends on  $q(o_i, o_j)$ )

$$\begin{aligned} &= \left( \sum_{q(o_1, o_1)} \phi(p(o_1), q(o_1, o_1)) \right) \cdots \left( \sum_{q(o_n, o_n)} \phi(p(o_n), q(o_n, o_n)) \right) \\ &= \prod_{XY} \sum_{q(X,Y)} \phi(p(X), q(X, Y)) \end{aligned}$$



(by observing that only the summation is the same for all  $q(X, Y)$ )

$$= \prod_{XY} \phi'(p(X)).$$

Inversion works by establishing a one-to-one correspondence between parfactor instantiations and summations. If the summation were on the instantiations of  $p(X)$ , such correspondence would not be possible because there would be less summations ( $|X|$  of them) than parfactor instantiations ( $|X| * |Y|$  of them).

Another condition for inversion is that the c-atom being inverted not have different instances in the same instance of the parfactor. For example, we cannot use inversion on  $p(X, Y)$  for a parfactor on  $p(X, Y), p(Y, X)$  because for any pair of objects  $o_i, o_j$ , neither of the parfactor instantiations  $p(o_i, o_j)$  and  $p(o_j, o_i)$  can be factored out of the innermost of  $\sum_{p(o_i, o_j)}$  and  $\sum_{p(o_j, o_i)}$ . This breaks the one-to-one correspondence between summations and instantiated parfactors.

In the case above, the resulting inner summation was propositional. Inversion resulting in propositional summations were called *inversion elimination* in our earlier work [2, 3]. In the next example, the inner summation is one computed by counting elimination.

Suppose we want to calculate

$$\begin{aligned} & \sum_{RV(p(X, Y))} \prod_{X, Y, Z} \phi(p(X, Y), p(X, Z)) \\ &= \sum_{RV(p(X, Y))} \prod_X \prod_{Y, Z} \phi(p(X, Y), p(X, Z)) \\ &= \sum_{RV(p(o_1, Y))} \cdots \sum_{RV(p(o_n, Y))} \prod_{Y, Z} \phi(p(o_1, Y), p(o_1, Z)) \cdots \prod_{Y, Z} \phi(p(o_n, Y), p(o_n, Z)) \\ &= \left( \sum_{RV(p(o_1, Y))} \prod_{Y, Z} \phi(p(o_1, Y), p(o_1, Z)) \right) \cdots \left( \sum_{RV(p(o_n, Y))} \prod_{Y, Z} \phi(p(o_n, Y), p(o_n, Z)) \right) \\ &= \prod_X \sum_{RV(p(X, Y))} \prod_{Y, Z} \phi(p(X, Y), p(X, Z)) \end{aligned}$$

Because  $X$  is now bound before the summation, it works as a constant (whose exact identity is irrelevant), and so it is not included in the counting argument. The counting argument now involves only  $Y$  and  $Z$  and is actually very similar to our original counting argument example. For that reason, the above is equal to  $\prod_X \phi'()$ , for  $\phi'$  the result of counting elimination.

Inversions resulting in counting elimination problems only invert on a subset of the parfactor's logical variables. For this reason, they have been called *Partial inversions* in our previous work. Note however that, since propositional sums are a trivial case of counting elimination, both inversion operations can be unified into one. This is what we do in the formalization below.

### 15.3.2.1 Uniform Solution Counting Partition (USCP)

Before we present the theorem formalizing inversion, we touch a last issue. Consider the inversion of  $X$  resulting in the expression

$$\prod_X \sum_{RV(p(X,Y))} \prod_{Y \neq X, Z \neq X, Y \neq a, Z \neq a} \phi(p(X,Y), p(X,Z)).$$

The summation can be done by counting elimination since  $X$  is bound. However, it will depend on  $|RV(p(X,Y))|$ , but that depends on whether  $X = a$  or not. One needs to split the expression according to cases  $X = a$  and  $X \neq a$ :

$$\begin{aligned} & \left( \sum_{RV(p(a,Y))} \prod_{Y \neq a, Z \neq a} \phi(p(a,Y), p(a,Z)) \right) \\ & \times \left( \prod_{X \neq a} \sum_{RV(p(X,Y))} \prod_{Y \neq X, Z \neq X, Y \neq a, Z \neq a} \phi(p(X,Y), p(X,Z)) \right) \end{aligned}$$

and then proceed as usual.

In general, one needs to consider the **uniform solution counting partition (USCP)** of the inverted logical variables with respect to an original constraint system. The USCP  $U_L(C)$  of a set of logical variables  $L$  with respect to a constraint  $C$  is a set of constraints  $\{C_1, \dots, C_k\}$  such that  $\{[C_i]\}_i$  forms a partition of  $[C]_L$  and

$$\forall i \cdot \forall \theta', \theta'' \in [C_i] \cdot |[C\theta']| = |[C\theta'']|,$$

that is, the number of solutions for the constraint conditioned on  $L$  is the same for each of the components  $C_i$ .

### 15.3.2.2 Inversion Formalization

#### Theorem 15.2 Inversion

Let  $g$  be a shattered parfactor,  $L$  a set of logical variables and  $E$  a set of c-atoms such that  $RV(E)$  and  $RV(A_g \setminus E)$  are disjoint. If

1.  $\forall e_k, e_l \in E \cdot \forall \theta_i, \theta_j \in [C_{g|L}] \cdot \theta_i \neq \theta_j \Rightarrow RV(e_k \theta_i) \cap RV(e_l \theta_j) = \emptyset$ .
2.  $\forall \theta_i, \theta_j \in [C_{g|L}] \cdot \theta_i \neq \theta_j \Rightarrow RV(E \theta_i) \cap RV(E \theta_j) = \emptyset$ .

then

$$\sum_{RV(E)} \Phi(g) = \prod_{C \in U_L(C_g)} \Phi(g_C),$$

where  $g_C$  is the parfactor  $(\phi_{g'}, A_{g'}, C \wedge C_{g'})$  and using  $g'$  defined by the recursive computation  $g'\theta = \sum_{RV(E\theta)} \Phi(g\theta)$ , for  $\theta$  an arbitrary element of  $[C]$  (by the definition of USCP, it does not matter which).

**Proof** Let  $E = \{e_1, \dots, e_n\}$ ,  $[C_g|L] = \{\theta_1, \dots, \theta_m\}$ . Below, we decompose  $C_g$  into the part w.r.t.  $L$  and the remaining logical variables:

$$\begin{aligned}
\sum_{RV(E)} \Phi(g) &= \sum_{RV(E)} \prod_{\theta \in \Theta_g} \phi(A_g \theta) \\
&= \sum_{RV(E)} \prod_{\theta \in [C_g|L]} \prod_{\theta' \in [C_g \theta]} \phi(A_g \theta \theta') \\
&= \sum_{RV(e_1)} \cdots \sum_{RV(e_n)} \prod_{\theta \in [C_g|L]} \prod_{\theta' \in [C_g \theta]} \phi(A_g \theta \theta') \\
&= \sum_{RV(e_1 \theta_1)} \cdots \sum_{RV(e_n \theta_1)} \cdots \sum_{RV(e_1 \theta_m)} \cdots \sum_{RV(e_n \theta_m)} \left( \prod_{\theta' \in [C_g \theta_1]} \phi(A_g \theta_1 \theta') \right) \cdots \left( \prod_{\theta' \in [C_g \theta_m]} \phi(A_g \theta_m \theta') \right) \\
&= \left( \sum_{RV(e_1 \theta_1)} \cdots \sum_{RV(e_n \theta_1)} \prod_{\theta' \in [C_g \theta_1]} \phi(A_g \theta_1 \theta') \right) \cdots \left( \sum_{RV(e_1 \theta_m)} \cdots \sum_{RV(e_n \theta_m)} \prod_{\theta' \in [C_g \theta_m]} \phi(A_g \theta_m \theta') \right) \\
&= \prod_{\theta \in [C_g|L]} \sum_{RV(e_1 \theta)} \cdots \sum_{RV(e_n \theta)} \left( \prod_{\theta' \in [C_g \theta]} \phi(A_g \theta \theta') \right) \\
&= \prod_{\theta \in [C_g|L]} \sum_{RV(E \theta)} \left( \prod_{\theta' \in [C_g \theta]} \phi(A_g \theta \theta') \right) \\
&= \prod_{C \in U_L(C_g)} \prod_{\theta \in [C]} \sum_{RV(E \theta)} \Phi(g \theta) = \prod_{C \in U_L(C_g)} \prod_{\theta \in [C]} \Phi(g_C \theta) = \prod_{C \in U_L(C_g)} \Phi(g_C).
\end{aligned}$$

Note that condition 1 is used to ensure the summations on  $\sum_{RV(e_1 \theta_1)} \cdots \sum_{RV(e_n \theta_m)}$  are indeed distinct. Condition 2 ensures that the innermost products are on distinct sets of random variables and can therefore be factored out as shown. ■

### 15.3.3 The Algorithm

Figure 15.1 shows the main pseudocode for FOVE. The algorithm consists of successively choosing eliminations  $(E, \{L_1, \dots, L_k\})$ , consisting of a collection of atoms  $E$  to eliminate after performing a series of inversions based on sets  $L_1, \dots, L_k$  of logical variables. A possible way of choosing eliminations is presented in figure 15.2. It is presented separately from the main algorithm for clarity, but because these two phases have many operations in common, actual implementations will typically integrate them more tightly.

There are potentially many ways to choose eliminations. The one we present starts by choosing an atom and checking if its inversion will produce a propositional summation, since this is the most efficient case. If not, we successively add atoms to  $E$  until  $G_E$  forms a parfactor where all atoms with logical variables are part of  $E$  (because counting elimination requires it). Then, for efficiency and to avoid shared logical variables between atoms, we try to determine as many inversions as possible, coded in the sequence  $L_1, \dots, L_k$ , to be done before counting elimination (or explicit summation when counting cannot be done).

FUNCTION *FOVE*( $G, Q$ )

$G$  a set of parfactors,  $Q \subseteq RV(G)$ ,  $G$  shattered against  $Q$  (section 15.5.2).

1. If  $RV(G) = Q$ , return  $G$ .
2.  $(E, \{L_1, \dots, L_k\}) \leftarrow \text{CHOOSE-ELIMINATION}(G, Q)$ .
3.  $g_E \leftarrow fs(G_E)$  (fusion, section 15.5.1).
4.  $G' \leftarrow \text{ELIMINATE}(g_E, E)$ .
5. Return  $\text{FOVE}(G' \cup G_{\neg E}, Q)$ .

FUNCTION *ELIMINATE*( $g, E, \{L_1, \dots, L_k\}$ )

1. If  $k = 0$  (no inversion)  
return *SUMMATION-WITHOUT-INVERSION*( $g, E$ ).
2.  $E_1 \leftarrow \{e \in E : LV(e) \cap L_1 \neq \emptyset\}$  (get inverted atoms).
3. Return  $\bigcup_{C_1 \in U_L(C_g)} \text{ELIMINATE-GIVEN-UNIFORMITY}(g, E_1, C_1, \{L_2, \dots, L_k\})$ .

FUNCTION *ELIMINATE-GIVEN-UNIFORMITY*( $g, E_1, C_1, \{L_2, \dots, L_k\}$ )

1. Choose  $\theta_1 \in [C_1]$  (bind inverted logical variables arbitrarily).
2.  $G' \leftarrow \text{ELIMINATE}(g\theta_1, E_1\theta_1, \{L_2, \dots, L_k\})$ .
3.  $G'' \leftarrow \bigcup_{g'\theta_1 \in G'} (\phi_{g'}, A_{g'}, C_1 \wedge C_{g'})$ .
4. Return  $\bigcup_{g'' \in G''} \text{SIMPLIFICATION}(g'')$  (simplification, section 15.5.3).

FUNCTION *SUMMATION-WITHOUT-INVERSION*( $g, E$ )

1. If  $E = \{E_1, \dots, E_k\}$  atoms are independent given  $C_g$  and  $A_g \setminus E$  is ground  
return  $(\sum_{\vec{N}_E} \vec{N}_E! \prod_v \phi_g(v, A_g \setminus E) \prod_{i=1}^k \vec{N}_{E_i, v_i}, A_g \setminus E, \top)$  (counting, section 15.3.1).
2. Return  $(\sum_{RV(E)} \prod_{\theta \in \Theta_g} \phi_g(A_g\theta_g), A_g \setminus E, C_g)$  (propositional elimination).

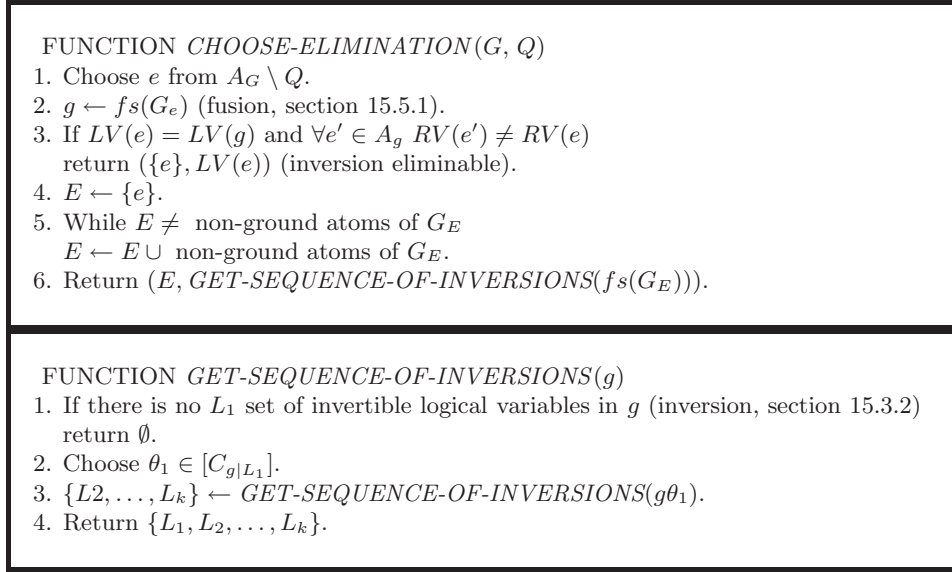
Notation:

- $LV(\alpha)$ : logical variables in object  $\alpha$ .
- $g\theta$ : parfactor  $(\phi_g, A_g\theta, C_g\theta)$ .
- $U_L(C_g)$ : USCP of  $L$  with respect to  $C_g$  (section 15.3.2.1).
- $C_{|L}$ : constraints projected to a set of logical variables  $L$ .
- $G_E$ : subset of parfactors  $G$  which depend on  $RV(E)$ .
- $G_{\neg E}$ : subset of parfactors  $G$  which do not depend on  $RV(E)$ .
- $\top$ : tautology constraint.

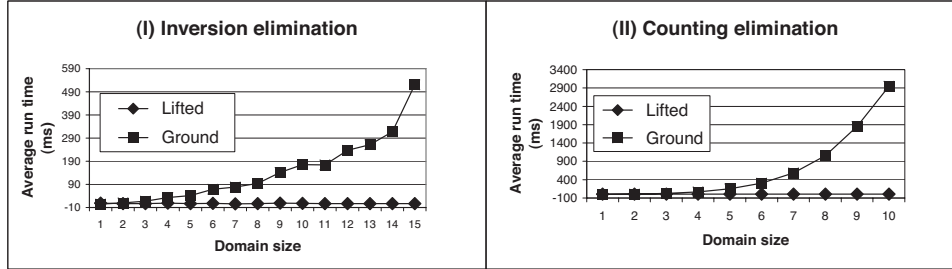
Figure 15.1 The FOVE algorithm.

## 15.4 An experiment

We use the implementation available at <http://l2r.cs.uiuc.edu/~cogcomp> to compare average run times between lifted and propositional inference (which produce the exact same results) for two different models while increasing the number



**Figure 15.2** One possible way of choosing an elimination.



**Figure 15.3** (I) Average run time for answering query  $P(p)$  from a parfactor on  $(p, q(X))$ , using inversion elimination, with domain size  $|X|$  being gradually increased. (II) Average run time for answering query  $P(r)$  from a parfactor on  $(p(X), p(Y), r)$ , using counting elimination, with domain size  $|X| = |Y|$  being gradually increased.

of objects in the domain. The first one, (I) in figure 15.3, answers the query  $P(p)$  from a parfactor on  $(p, q(X))$  and uses inversion elimination only. The inference in (II) answers query  $P(r)$  from a parfactor on  $(p(X), p(Y), r)$  and uses counting elimination only. In both cases propositional inference starts taking very long before any noticeable variation in lifted inference run times.

## 15.5 Auxiliary operations

### 15.5.1 Fusion

We have assumed in section 15.3 that we have operations to calculate  $\sum_{RV(E)} \Phi(G_E)$ , but elimination operations calculate  $\sum_{RV(E)} \Phi(g)$ , for  $g$  a *single* parfactor. *Fusion* bridges this gap by computing, for any set of parfactors  $G$ , a single parfactor  $fs(G)$  such that  $\Phi(G) = \Phi(fs(G))$ .

Fusion works by replacing the constraints of all parfactor in the set by a single, common constraint which is the conjunction of them all. This guarantees that all parfactories get instantiated by the same set of substitutions on a single set of logical variables, which allows their products (in the expression for  $\Phi(G)$ ) to be unified under a single product. Note that not all parfactories contain all the logical variables, and will be instantiated to the same ground factor by distinct substitutions (those agreeing on the logical variables present in the parfactor, but disagreeing on some of the others). In other words, some of the parfactories will have their number of instantiations increased by this unification. For this reason, we also exponentiate the potential function to the inverse of how many times the number of instantiations was increased, keeping the final result the same as before.

This is illustrated in the example below:

$$\begin{aligned} \left( \prod_{D,P} \phi_1(e(D), s(D, P)) \right) \left( \prod_{D'} \phi_2(e(D')) \right) &= \prod_{D,P,D'} \phi_1(e(D), s(D, P)) \phi_2^{\frac{1}{|D,P|}}(e(D')) \\ &= \prod_{D,P,D'} \phi_3(e(D), s(D, P), e(D')). \end{aligned}$$

(note that logical variables in different parfactories must be standardized apart.)  
Formally, we have the fusion theorem below.

**Theorem 15.3 Fusion**

Let  $G$  be a set of parfactories. Define  $C_G = \bigwedge_{g \in G} C_g$ ,  $\Theta_G = [C_G]$  and  $A_G = \bigcup_{g \in G} A_g$ . Let  $fs(G)$  be the parfactor  $(\prod_{g \in G} \phi_g^{|\Theta_g|/|\Theta_G|}, A_G, C_G)$ . Then  $\Phi(G) = \Phi(fs(G))$ .

**Proof**

$$\begin{aligned} \Phi(G) &= \prod_{g \in G} \prod_{\theta \in \Theta_g} \phi_g(A_g \theta) = \prod_{g \in G} \prod_{\theta \in \Theta_G} \phi_g(A_g \theta)^{|\Theta_g|/|\Theta_G|} \\ &= \prod_{\theta \in \Theta_G} \prod_{g \in G} \phi_g(A_g \theta)^{|\Theta_g|/|\Theta_G|} = \prod_{\theta \in \Theta_G} \phi_{fs(G)}(A_G \theta) = \Phi(fs(G)) \end{aligned}$$

■

While the above is correct, it is rather unnatural to have  $e(D)$  and  $e(D')$  be distinct atoms. If a set of logical variables has the same possible substitutions, like  $D$  and  $D'$  here, we can do something better:

$$\begin{aligned}
\left(\prod_{D,P} \phi_1(e(D), s(D, P))\right) \left(\prod_{D'} \phi_2(e(D'))\right) &= \left(\prod_D \prod_P \phi_1(e(D), s(D, P))\right) \left(\prod_{D'} \phi_2(e(D'))\right) \\
&= \prod_{D''} \left(\left(\prod_P \phi_1(e(D''), s(D'', P))\right) (\phi_2(e(D''))) \right) \\
&= \prod_{D''} \left(\prod_P \phi_1(e(D''), s(D'', P)) \phi_2(e(D''))^{\frac{1}{|P|}}\right) \\
&= \prod_{D'', P} \phi_1(e(D''), s(D'', P)) \phi_2^{\frac{1}{|P|}}(e(D'')) \\
&= \prod_{D'', P} \phi_3(e(D''), s(D'', P)).
\end{aligned}$$

Formally, this process is similar to inversion with respect to  $D''$ . However, it does require the additional previous step of unifying distinct logical variables (but with identical sets of possible substitutions) into a single one first (in the example,  $D$  and  $D'$  are replaced by  $D''$ ). For lack of space we omit the details of this improvement.

### 15.5.2 Shattering

In section 15.3 we mentioned the need for *shattering*, which we now discuss in more detail. This need arises from c-atoms representing overlapping, but not identical, classes of random variables. Consider the following marginalization over parfactors  $g_1$  and  $g_2$  with potential functions  $\phi_1$  and  $\phi_2$  respectively:

$$\sum_{RV(p(X,Y))} \left(\prod_{X,Y} \phi_1(p(X,Y), q)\right) \prod_Y \phi_2(p(a,Y))$$

If we pick  $E = p(a, Y)$ ,  $G_E = \{g_1, g_2\}$ . However, only some instantiations of  $g_1$  depend on  $p(a, Y)$  (the ones with  $X = a$ ). Moreover, the operations we later talk about require any pair of c-atoms in  $G_E$  to represent either identical classes of random variables, or those classes to be disjoint. This is violated by  $RV(p(a, Y))$  being a subset of  $RV(p(X, Y))$ . Picking  $E = p(X, Y)$  also violates this requirement.

The solution is to **split** parfactor  $g_1$  into two different parfactors. The union of instantiations of  $(\phi_1, (p(X, Y), q), X \neq a)$  and  $(\phi_1, (p(a, Y), q), \top)$  is identical to the set of instantiations of  $g_1$ , so the summation can be simply rewritten as

$$\begin{aligned}
&\sum_{RV(p(X,Y))} \left(\prod_{X,Y:X \neq a} \phi_1(p(X,Y), q)\right) \left(\prod_Y \phi_1(p(a,Y), q)\right) \prod_Y \phi_2(p(a,Y)) \\
&= \sum_{RV(p(X,Y):X \neq a)} \left(\prod_{X,Y:X \neq a} \phi_1(p(X,Y), q)\right) \sum_{p(a,Y)} \left(\prod_Y \phi_1(p(a,Y), q)\right) \prod_Y \phi_2(p(a,Y))
\end{aligned}$$

Now  $E = p(a, Y)$  satisfies the operations' requirements. Picking  $E = p(X, Y)$ ,  $X \neq a$  would work equally well.

Splitting parfactors is done by pairwise comparisons of atoms of the same predicate. We split parfactors  $g_1 = (\phi_1, A_1, C_1)$  and  $g_2 = (\phi_2, A_2, C_2)$  around atoms  $a_1 \in A_1$  and  $a_2 \in A_2$  by replacing them by parfactors  $(\phi_1, A_1, C_1 \wedge a_1 = a_2)$ ,  $(\phi_1, A_1, C_1 \wedge a_1 \neq a_2)$ ,  $(\phi_2, A_2, C_2 \wedge a_1 = a_2)$  and  $(\phi_2, A_2, C_2 \wedge a_1 \neq a_2)$ , after standardizing apart their logical variables. In fact, we only need to keep those whose constraint is satisfiable. (This is why  $g_2$  does not need to be broken in the example above – that would only produce itself and another parfactor with zero instantiations.) In particular, if  $RV(a_1) = RV(a_2)$ , we end up obtaining the original parfactors.

The uniformity requirement is met by **shattering** the FOPM in advance, that is, by successively splitting the parfactors of each pair of c-atoms, including the query atoms, until no overlapping non-identically grounded pair remains. The query atoms need to be involved in shattering because if a c-atom includes query and non-query random variables, it needs to be split so that the non-query ones can be eliminated.

As pointed out by Poole [9], splitting parfactors resembles the role of unification in first-order resolution, which determines the conditions for two atoms to match. In probabilistic inference, however, we are interested not only in the overlapping of atoms but also in the *residual* parfactors that originate from the matching. The reason for this difference is that the *number* of instantiations of a parfactor matters for the final joint distribution. In regular resolution, the original clauses are kept because their redundancy with the clauses resulting from resolution makes no difference, while here we need to discount them and replace the originals with the non-matching cases.

### 15.5.3 Irrelevant Logical Variable Simplification

Inversion often produces parfactors with constraints with logical variables not present in its atoms. The first inversion example produces the expression below. We can simplify it by observing that the actual value of  $Y$  is irrelevant inside the product. Only the number  $|Y|$  of possible values for  $Y$  will make a difference. Therefore we can write

$$\prod_{XY} \phi'(p(X)) = \prod_X \phi'(p(X))^{|Y|} = \prod_X \phi''(p(X)).$$

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## 15.6 Applicability of lifted inference

As explained in the previous sections, the lifted operations of FOVE are not always applicable, each of them requiring certain conditions to be satisfied in advance. Therefore a natural question is to what kinds of FOPMs we can apply FOVE in an exclusively lifted manner.

It is not clear at this point whether it is possible to tell in advance if a FOPM can be solved with lifted operations alone. The main reason for this is that lifted



operations will be applied to parfactors resulting from previous operations, so we do not know them in advance. It may be that two parfactors satisfying the lifted operations conditions fuse to form one which does not. (This is similar to the fact that an elimination ordering is not computed in advance but only as the algorithm proceeds.)

As a summarization, the conditions for applying lifted operations to eliminate a set  $RV(E)$  from a parfactor  $g$  are the following: for counting elimination, the atoms in  $g$  must be independent given its  $C_g$ ; for inversion on  $L \subseteq LV(g)$ ,

1.  $\forall e_k, e_l \in E \cdot \forall \theta_i, \theta_j \in [C_{g|L}] \cdot \theta_i \neq \theta_j \Rightarrow RV(e_k \theta_i) \cap RV(e_l \theta_j) = \emptyset$ .
2.  $\forall \theta_i, \theta_j \in [C_{g|L}] \cdot \theta_i \neq \theta_j \Rightarrow RV(E \theta_i) \cap RV(E \theta_j) = \emptyset$ .

When lifted operations do not apply, FOVE uses non-lifted operations to calculate  $\sum_{RV(E)} G_E$ . These non-lifted methods could be propositionalization, sampling etc, but with the advantage of being restricted to a subset of the model only.

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## 15.7 Future Directions

There are several possible directions for further development of FOVE. One of the main ones is the incorporation of function symbols, both random (the color of an object, for example) and interpreted (summation over integers), which will greatly increase its expressivity and applicability.

In applications involving evidence over many objects (for example, the facts about all the words in an English document), shattering may take a long time because all parfactors have to be checked against it. The large number of objects involved may create the need for numerous parfactor splittings. This is unfortunate because often only some objects are truly relevant to the query. For example, analyzing only some words and phrases in a document will often be enough to determine its subject. Therefore a variant of FOVE that does only the necessary shattering, guided by the inference process, is of great interest.

Finally, lifted FOVE operations do not cover all possible cases and explicit summation may be required at times, so increasing their coverage is an important direction.

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## 15.8 Conclusion

Intuitive descriptions of models very often include first-order elements. When these models are probabilistic, the dominant approach has been that of grounding the model to a propositional one and solving it with a regular propositional algorithm. This strategy loses the explicit representation of the model's first-order structure, which can be used to great computational advantage, and which is computationally hard to retrieve from the grounded model.

We presented FOVE, a first-order generalization of the popular VE propositional inference algorithm. Like VE, FOVE successively eliminates random variables from the model by summing them out while taking advantage of independences for efficiency. Unlike VE, FOVE directly manipulates first-order representations, eliminating c-atoms that stand for potentially large sets of random variables at once. This can in some cases exponentially (in the domain size) speed inference up.

There are important directions in which FOVE needs to be extended, such as incorporating function symbols, avoiding unnecessary shattering, and extending operations for as of yet uncovered cases. However FOVE is already applicable, and especially useful, in domains with large objects about which we have identical knowledge. More than that, it is a general framework to be expanded and help close the gap between logic and probabilistic reasoning.

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