Abstract

We propose the use of hyperedge replacement graph grammars for factor graphs, or factor graph grammars (FGGs) for short. FGGs generate sets of factor graphs and can describe a more general class of models than plate notation, dynamic graphical models, case–factor diagrams, and sum–product networks can. Moreover, inference can be done on FGGs without enumerating all the generated factor graphs. For finite variable domains (but possibly infinite sets of graphs), a generalization of variable elimination to FGGs allows exact and tractable inference in many situations. For finite sets of graphs (but possibly infinite variable domains), a FGG can be converted to a single factor graph amenable to standard inference techniques.

1 Introduction

Graphs have been used with great success as representations of probability models, both Bayesian and Markov networks (Koller and Friedman, 2009) as well as latent-variable neural networks (Schulman et al., 2015). But in many applications, especially in speech and language processing, a fixed graph is not sufficient. The graph may have substructures that repeat a variable number of times: for example, a hidden Markov model (HMM) depends on the number of words in the string. Or, part of the graph may have several alternatives with different structures: for example, a probabilistic context-free grammar (PCFG) contains many trees for a given string.

Several formalisms have been proposed to fill this need. Plate notation (Buntine, 1994), plated factor graphs (Obermeyer et al., 2019), and dynamic graphical models (Bilmes, 2010) address the repeated-substructure problem, but only for sequence models like HMMs. Case–factor diagrams (McAllester et al., 2008) and sum–product networks (Poon and Domingos, 2011) address the alternative-substructure problem, so they can describe PCFGs, but only for fixed-length inputs.

More general formalisms like probabilistic relational models (Getoor et al., 2007) and probabilistic programming languages (van de Meent et al., 2018) address both problems successfully, but because of their generality, tractable exact inference in them is often not possible.

Here, we explore the use of hyperedge replacement graph grammars (HRGs), a formalism for defining sets of graphs (Bauderon and Courcelle, 1987; Habel and Kreowski, 1987; Drewes et al., 1997). We show that HRGs for factor graphs, or factor graph grammars (FGGs) for short, are expressive enough to solve both the repeated-substructure and alternative-substructure problems, and constrained enough to allow exact and tractable inference in many situations. We make three main contributions:

- We define FGGs and show how they generalize the constrained formalisms mentioned above (§3).
- We define a conjunction operation that enables one to modularize a FGG into two parts, one which defines the model and one which defines a query (§4).
- We show how to perform inference on FGGs without enumerating the (possibly infinite) set of graphs they generate. For finite variable domains, we generalize variable elimination to FGGs.

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We draw a factor graph. Although the elements of a variable domain must be finite sets of node labels and edge labels, and assume there is a function \( \text{type} : L^E \to (L^V)^* \), which says for each edge label what the number and labels of the endpoint nodes must be.

**Definition 1.** A hypergraph (or simply a graph) is a tuple \((V, E, \text{att}^V, \text{lab}^V, \text{lab}^E)\), where

- \( V \) is a finite set of nodes.
- \( E \) is a finite set of hyperedges (or simply edges).
- \( \text{att} : E \to V^* \) maps each edge to zero or more endpoint nodes, not necessarily distinct.
- \( \text{lab}^V : V \to L^V \) assigns labels to nodes.
- \( \text{lab}^E : E \to L^E \) assigns labels to edges.
- For all \( e \), \(|\text{att}(e)| = |\text{type}(\text{lab}^E(e))|\), and if \( \text{att}(e) = v_1 \cdots v_k \) and \( \text{type}(\text{lab}^E(e)) = \ell_1 \cdots \ell_k \), then \( \text{lab}^V(v_i) = \ell_i \) for \( i = 1, \ldots, k \).

Although the elements of \( V \) and \( E \) can be anything, we assume in our examples that they are natural numbers. If a node \( v \) has label \( \ell \), we draw it as a circle with \( \ell \) inside it. We draw a hyperedge as a square with lines to its endpoints. In principle, we would need to indicate the ordering of the endpoints somehow, but we omit this to reduce clutter.

**2.1 Hypergraphs**

**Definition 2.** A factor graph (Kschischang et al., 2001) is a hypergraph \((V, E, \text{att}^V, \text{lab}^V, \text{lab}^E)\) together with mappings \( \Omega \) and \( F \), where

- \( \Omega \) maps node labels to sets of possible values. For brevity, we write \( \Omega(v) \) for \( \Omega(\text{lab}^V(v)) \).
- \( F \) maps edges labels to functions. For brevity, we write \( F(e) \) for \( F(\text{lab}^E(e)) \). For every edge \( e \) with \( \text{att}(e) = v_1 \cdots v_k \), \( F(e) \) is of type \( \Omega(v_1) \times \cdots \times \Omega(v_k) \to \mathbb{R}_{\geq 0} \).

A node \( v \) together with its domain \( \Omega(v) \) is called a variable. An edge \( e \) together with its function \( F(e) \) is called a factor.

We draw a factor \( e \) as a small square, but instead of writing its label, we write \( F(e) \) next to it, as an expression in terms of its endpoints. As shorthand, we often write Boolean expressions, which are implicitly converted to real numbers (true = 1 and false = 0).

**Example 3.** Although HMMs are defined for sentences of arbitrary length, factor graphs force us to choose a fixed length; below is a HMM for sentences of length 3. (Here, \( T \) and \( W \) are node labels, \( \Omega(T) \) is the set of possible tags, and \( \Omega(W) \) is the set of possible words.)

\[
\begin{array}{cccccccc}
T_0 = \text{BOS} & p(T_1 | T_0) & p(T_3 | T_1) & p(T_5 | T_3) & p(T_7 | T_5) & T_7 = \text{EOS} \\
& p(W_2 | T_1) & p(W_4 | T_3) & p(W_6 | T_5) & \\
W_2 & W_4 & W_6
\end{array}
\]
As with a CFG, we can abstract away from the ordering of replacement steps using a derivation tree. Although a left-hand side $X$ has external nodes associated with it, if a nonterminal $X$ is replaced with the right-hand side of a rule in a derivation, we simply write $X$ for the set of all replacements of $H$. In a factor graph with no factors, every assignment has weight 1. A factor graph with no variables has exactly one assignment. Factor graphs are general enough to represent Bayesian networks and Markov networks. They can also represent stochastic computation graphs (SCGs), introduced by Schulman et al. (2015) for latent-variable neural networks.

### 2.3 Hyperedge Replacement Graph Grammars

Hyperedge replacement graph grammars (HRGs) were introduced by Bauderon and Courcelle (1987) and Habel and Kreowski (1987), and surveyed by Drewes et al. (1997). They generate graphs by using a context-free rewriting mechanism that replaces nonterminal-labeled edges with graphs. In this section, we provide a brief definition of HRGs, with a minor extension for node labels.

**Definition 4.** If $H$ is a factor graph, define an assignment $\xi$ of $H$ to be a mapping from nodes to values: $\xi(v) \in \Omega(v)$. We write $\Xi_H$ for the set of all assignments of $H$. The weight of an assignment $\xi$ is given by

$$w_H(\xi) = \prod_{e \text{ edges with } \text{att}(e) = v_1 \cdots v_k} F(e)(\xi(v_1), \ldots, \xi(v_k)).$$

In a factor graph with no factors, every assignment has weight 1. A factor graph with no variables has exactly one assignment.

**Definition 5.** A hypergraph fragment is a tuple $(V, E, \text{att}, \text{lab}^V, \text{lab}^E, \text{ext})$, where

- $(V, E, \text{att}, \text{lab}^V, \text{lab}^E)$ is a hypergraph,
- $\text{ext} \in V^*$ is a sequence of zero or more external nodes.

In our figures, we draw external nodes as black nodes. In principle, we would need to indicate their ordering somehow, but we omit this to reduce clutter.

**Definition 6.** A hyperedge replacement graph grammar (HRG) is a tuple $(N, T, P, S)$, where

- $N \subseteq L^E$ is a finite set of nonterminal symbols.
- $T \subseteq L^E$ is a finite set of terminal symbols, such that $N \cap T = \emptyset$.
- $P$ is a finite set of rules of the form $(X \rightarrow R)$, where
  - $X \in N$.
  - $R$ is a hypergraph fragment with edge labels in $N \cup T$.
  - If $R$ has external nodes $x_1 \cdots x_k$, then $\text{type}(X) = \text{lab}^V(x_1) \cdots \text{lab}^V(x_k)$.
  - $S \in N$ is a distinguished start nonterminal symbol with $\text{type}(S) = \epsilon$.

Although a left-hand side $X$ is formally just a nonterminal symbol, we draw it as a hyperedge labeled $X$ inside, with replicas of the external nodes as its endpoints. On right-hand sides, we draw an edge $e$ with nonterminal label $X$ as a square with $X_e$ inside. If $R$ is the empty graph, we write $\emptyset$.

Intuitively, a HRG generates graphs by starting with a hyperedge labeled $S$ and repeatedly selecting an edge $e$ labeled $X$ and a rule $X \rightarrow R$ and replacing $e$ with $R$. (See Figure 1a for an example, where $H = R$.) Replacement stops when there are no more nonterminal-labeled edges.

As with a CFG, we can abstract away from the ordering of replacement steps using a derivation tree, in which the nodes are labeled with HRG rules, and an edge from parent $\pi_1$ to child $\pi_2$ has a label indicating which edge in the right-hand side of $\pi_1$ is replaced with the right-hand side of $\pi_2$.

**Definition 7.** Let $G$ be a HRG. For all nonterminals $X$, define the set $D(G, X)$ of $X$-type derivation trees (or simply $X$-type derivations) of $G$ to be the smallest set containing all finite, unordered, edge-labeled trees of the form shown in Figure 1b, where $\pi = (X \rightarrow R)$ is a rule in $G$, $R$ has nonterminal-labeled edges $e_1, \ldots, e_k$ with labels $X_1, \ldots, X_k$, and for $i = 1, \ldots, k$, $D_i$ is an $X_i$-type derivation. We simply write derivation for $S$-type derivation, and we let $D(G) = D(G, S)$.

The derived graph of a derivation $D$ is the graph formed as follows. If $D$ is as shown in Figure 1b, then for $i = 1, \ldots, k$, let $H_i$ be the derived graph of $D_i$. In (a copy of) $R$, replace $e_i$ with $H_i$, making the $j$th endpoint of $e_i$ and the $j$th external node of $H_i$ into the same node ($j = 1, \ldots, |\text{att}(e_i)|$). The resulting node is external iff the $j$th endpoint was. All other nodes are kept distinct. (Again, see Figure 1a for an example with $X = X_i$ and $H = H_i$.)
Figure 1: (a) Example of replacing a hyperedge labeled $X$ with a hypergraph fragment $H$. Here $|\text{type}(X)| = 3$, but in general, there could be any number of endpoint/external nodes, including zero. (b) A derivation tree.

![Diagram](image)

(a) (b)

Figure 2: (a) A FGG generating the infinite set of unrollings of a HMM, one for each sequence length. Each rule is labeled $\pi_i$ for use in the derivation tree. (b) Derivation tree of the factor graph of Example 3. An edge from parent $\pi$ with label $X$ to child $\pi'$ means that the right-hand side of $\pi'$ replaces the edge labeled $X$ in the right-hand side of $\pi$.

From now on, when we mention a derivation $D$ in a context where a graph would be expected, the derived graph of $D$ is to be understood.

## 3 Factor Graph Grammars

**Definition 8.** A HRG for factor graphs, or a factor graph grammar (FGG) for short, is a HRG together with mappings $\Omega$ and $F$, as in the definition of factor graphs (Definition 2), except that $F$ is defined on terminal edge labels only.

**Example 9.** Figure 2 shows a FGG which is equivalent to a HMM. It generates an infinite number of graphs, one for each string length. Also shown is the derivation tree of the factor graph of Example 3.

Example 19 in Appendix A shows how to simulate a PCFG in Chomsky normal form as a FGG.

The graphs generated by a FGG can be viewed, together with $\Omega$ and $F$, as factor graphs, each of which defines a (not necessarily normalized) distribution over assignments. Moreover, the whole language of the FGG defines a (not necessarily normalized) distribution over derivations and assignments to the variables in them. If $D \in D(G)$, then

$$w_G(D, \xi) = w_D(\xi).$$

FGGs can simulate several other formalisms for dynamically-structured models. As mentioned above (§1), they can solve two problems that previous formalisms have addressed separately.

FGGs can generate repeated substructures like plate notation (Buntine, 1994; Obermeyer et al., 2019) and dynamic graphical models (Bilmes, 2010) can. There are some structures that plate notation can describe that a FGG cannot – like the set of all restricted Boltzmann machines, which have two
fully-connected layers of nodes. But these are the same structures that Obermeyer et al. (2019) try to avoid, because inference on them is (believed) intractable. FGG rules these structures out naturally.

FGGs can generate alternative substructures like case–factor diagrams (McAllester et al., 2008) and valid sum–product networks (Poon and Domingos, 2011) can; in particular, they can simulate PCFGs in such a way that inference is equivalent to the cubic-time inside and Viterbi algorithms.

**Theorem 10.** All of the following can be converted into an equivalent FGG:

1. Plated factor graphs for which the sum–product algorithm of Obermeyer et al. (2019) succeeds.
2. Dynamic graphical models.
3. Case–factor diagrams.
4. Valid sum–product networks.

**Proof.** See Appendix B.

## 4 Conjunction

The preceding examples show how to use FGGs to model the probability of all tagged strings or all trees generated by a grammar. But it’s common for queries to constrain some variables to fixed values, sum over some variables, and get the distribution of the remaining variables. How do such queries generalize to FGGs? For example, in a HMM, how do we compute the probability of all taggings of a given string? Or, how do we compute the marginal distribution of the second-to-last tag?

To answer such questions, we need to be able to specify a set of nodes across the graphs of a graph language, like the second-to-last tag. Our only means of doing this is to specify a particular node in a particular right-hand side, which could correspond to zero, one, or many nodes in the derived graphs. And we can modify a FGG so that a particular node in a particular right-hand side is always (say) the second-to-last tag. But we propose to factor such modifications into a separate FGG, keeping the FGG describing the model unchanged. Then the modifications can be applied using a conjunction operation, which we describe in this section.

Conjunction is closely related to synchronous HRGs (Jones et al., 2012), and, because HRG derivation trees are generated by regular tree grammars, to intersection/composition of finite tree automata/transducers (Comon et al., 2007). It is also similar to the PRODUCT operation on weighted logic programs (Cohen et al., 2011).

**Definition 11.** Two FGG rules are *conjoinable* if they can be written in the form

\[
X_1 \to R_1 = (V, E_N \cup E_1, att_N \cup att_1, lab^V, lab_1^E, ext)
\]

\[
X_2 \to R_2 = (V, E_N \cup E_2, att_N \cup att_2, lab^V, lab_2^E, ext),
\]

Figure 3: Illustration of the conjunction operation (Example 12). In rules with \(i\) in their nonterminals, \(i\) ranges from 1 to \(n\) where \(n = |w|\).
where

- $E_N$ contains only nonterminal edges, and $att_N$ is defined on $E_N$.
- $E_1, E_2$ contain only terminal edges, and $att_1, att_2$ are defined on $E_1, E_2$, respectively.
- $type(X_1) = type(X_2)$, and for $e \in E_N$, $type(lab_1^E(e)) = type(lab_2^E(e))$.

Then their conjunction is

$$\langle X_1, X_2 \rangle \rightarrow R$$

$$R = (V, E_N \cup E_1 \cup E_2, att, lab^V, lab^E, ext)$$

where $\cup$ means that all edges in $E_1$ and $E_2$ are kept distinct while taking their union, and

$$lab^E(e) = \begin{cases} (lab_1^E(e), lab_2^E(e)) & \text{if } e \in E_N \\ lab_1^E(e) & \text{if } e \in E_1 \\ lab_2^E(e) & \text{if } e \in E_2 \end{cases}$$

att(e) = \begin{cases} att_N(e) & \text{if } e \in E_N \\ att_1(e) & \text{if } e \in E_1 \\ att_2(e) & \text{if } e \in E_2. \end{cases}

The conjunction of two FGGs $G_1$ and $G_2$, written as $G_1 \cap G_2$, is the FGG containing the conjunction of all conjoinable pairs of rules from $G_1$ and $G_2$.

**Example 12.** Our FGG for HMMs (Example 9) is repeated in Figure 3 as $G$. We can constrain the $W$ variables to an observed string $w$ using another FGG, $G_w$, which has the same variables as $G$ but different factors; its nonterminal edges are the same as $G$ but with different labels. This FGG generates just one graph, whose $W$ nodes spell out the string $w$. The conjunction of these two FGGs is shown in the last column ($G \cap G_w$). It combines the factors and nonterminal labels of $G$ and $G_w$ and generates just one graph, the HMM for string $w$.

**Example 13.** To compute the distribution of the second-to-last tag, we need a way of identifying the variable for the second-to-last tag across all graphs. We can do this by conjoining with the FGG:

$S \xrightarrow{\pi_1} T_1, X_1 \quad T_1 \xrightarrow{\pi_2} T_2, X_1 \quad T_1 \xrightarrow{\pi_3} T_2, Y_1 \quad T_2 \xrightarrow{\pi_4} T_1 \quad T_1 \xrightarrow{\pi_5} T_2 \quad T_2 \xrightarrow{\pi_6} T_1 \quad T_1 \xrightarrow{\pi_7} T_2 \quad T_2 \xrightarrow{\pi_8} W_1 \quad W_2$

Then the second-to-last tag is always node $T_1$ in the right-hand side of rule $\pi_3$. The methods of the following section can then be used to compute the distribution of this node.

Example 20 in Appendix A shows how to use conjunction to constrain a PCFG to a single input string.

## 5 Inference

Given a FGG $G$, we want to be able to efficiently compute its sum–product,

$$Z_G = \sum_{D \in D(G)} \sum_{\xi \in \Xi_D} w_G(D, \xi).$$

We can answer a wide variety of queries by using the conjunction operation to constrain variables based on observations, and then computing the sum–product in various semirings: ordinary addition and multiplication would sum over assignments to the remaining variables, and the expectation semiring (Eisner, 2002) would compute expectations with respect to them. The Viterbi (max–product) semiring would find the highest-weight derivation and assignment, not necessarily the highest-weight graph and assignment, which is NP-hard (Lyngsø and Pedersen, 2002).

We consider three cases below: finite variable domains, but possibly infinite graph languages ($\S$5.1); finite graph languages, but possibly infinite variable domains ($\S$5.2); and infinite variable domains and graph languages ($\S$5.3). To help characterize these cases and their subcases, we introduce the following definitions.

**Definition 14.** A FGG is recursive if it has an $X$-type derivation that contains an $X$-type derivation as a proper subtree; otherwise, it is nonrecursive. A nonrecursive FGG generates a finite set of graphs; this is a common case, because the conjunction of any FGG with a nonrecursive FGG (e.g., one describing a finite-sized observation) is nonrecursive.
A recursive FGG is *nonlinearly recursive* if it has an $X$-type derivation that contains two disjoint $X$-type derivations as proper subtrees; otherwise, it is *linearly recursive*.

A FGG is *nonreentrant* if no derivation contains two different $X$-type derivations as subtrees. Every nonreentrant FGG is nonrecursive, and any nonrecursive FGG can be made nonreentrant by duplicating rules and renaming nonterminals (though this may cause an exponential blowup in the size of the grammar).

### 5.1 Finite variable domains

When a HRG generates a graph, the derivation tree is isomorphic to a tree decomposition of the graph: each derivation tree node $\pi = (X \rightarrow R)$ corresponds to a bag of the tree decomposition containing the nodes in $R$. It follows that a HRG whose right-hand sides have at most $(k + 1)$ nodes generates graphs with treewidth at most $k$ (Bodlaender, 1998, Theorem 37). So if a FGG $G$ generates a graph $H$, computing the sum–product of $H$ by variable elimination (VE) takes time linear in the size of $H$ and exponential in $k$.

In this section, we generalize VE to compute the sum–product of all graphs generated by $G$ without enumerating them. If $G$ is nonrecursive, this is (like VE) linear in the size of $G$ and exponential in $k$; in the envisioned typical use-case, we have a fixed FGG $G$ representing the model and different FGGs $G'$ representing different observations; since conjunction cannot increase $k$, we may regard $k$ as fixed, so computing the sum–product of $G \cap G'$ takes time linear in the size of $G \cap G'$.

**Theorem 15.** Let $G = (N, T, P, S)$ be a FGG such that for all $v$ in $G$, $|\Psi(v)| \leq m$. Let $|G|$ be the number of rules in $G$, and let $k$ be such that every right-hand side in $G$ has at most $(k + 1)$ nodes. Then $Z_G$ is the least solution of a monotone system of polynomial equations, and in particular:

1. If $G$ is nonrecursive, $Z_G$ can be computed in $O(|G|m^{k+1})$ time.
2. If $G$ is linearly recursive, $Z_G$ can be computed in $O(|G|^3m^{3(k+1)})$ time in the worst case.

**Proof.** The computation of the sum–product is closely analogous to the sum–product of a PCFG (Stolcke, 1995; Nederhof and Satta, 2008). We introduce some shorthand for assignments. If $\xi$ is an assignment and $v_1 \cdots v_l$ is a sequence of nodes, we write $\xi(v_1) \cdots \xi(v_l)$ for $\xi(v_1) \cdot \cdots \cdot \xi(v_l)$. If $X$ is a nonterminal and $\text{type}(X) = \ell_1 \cdots \ell_k$, we define $\Xi_X = \Omega(\ell_1) \times \cdots \times \Omega(\ell_k)$, the set of assignments to the endpoints of an edge labeled $X$.

Next, we define a system of equations whose solution gives the desired sum–product. The unknowns are $\psi_X(\xi)$ for all $X \in N$ and $\xi \in \Xi_X$, and $\tau_R(\xi)$ for all rules $(X \rightarrow R)$ and $\xi \in \Xi_X$. For all $X \in N$, let $P^X$ be the rules in $P$ with left-hand side $X$. For each $\xi \in \Xi_X$, add the equation

$$\psi_X(\xi) = \sum_{(X \rightarrow R) \in P^X} \tau_R(\xi).$$

For each right-hand side $R = (V, E_N \cup E_T, \text{att}, \text{lab}^V, \text{lab}^E, \text{ext})$, where $E_N$ contains only nonterminal edges and $E_T$ contains only terminal edges, and for each $\xi \in \Xi_X$, add the equation

$$\tau_R(\xi) = \sum_{\xi' \in \Xi_R} \prod_{e \in E_T} F(e)(\xi'(\text{att}(e))) \prod_{e \in E_N} \psi_{\text{lab}^e(e)}(\xi'(\text{att}(e))).$$

Then $\sum_{\xi \in \Xi_X} \psi_X(\xi)$ represents the sum–product of all $X$-type derivations. In particular, the sum–product of the FGG is $\psi_S()$.

To solve these equations, construct a directed graph over nonterminals with an edge from $X$ to $Y$ iff there is a rule $X \rightarrow R$ where $R$ contains an edge labeled $Y$. For each connected component $C$ of this graph in reverse topological order:

1. If $C = \{X\}$, compute $\psi_X$ and substitute it into the other equations.
2. Else if the equations for $\psi_X$ and $\tau_R$ where $X \in C$ and $(X \rightarrow R) \in P$ are linear, solve them and substitute into the other equations (Stolcke, 1995; Goodman, 1999).
3. Else, the equations can be approximated iteratively (Goodman, 1999; Nederhof and Satta, 2008).
If \( G \) is nonrecursive, the graph of nonterminals is acyclic, so case (1) always applies. The total running time is \( O(|G|n^k+1) \).

If \( G \) is linearly recursive, then case (2) may also apply. In the worst case, the nonterminal graph is one connected component, corresponding to \( O(|G|n^k+1) \) unknowns. Solving the equations could involve inverting a matrix of this size, which takes \( O(|G|^2n^{2(k+1)}) \) time.

If \( G \) is nonlinearly recursive, any of the three cases may apply. For case (3), each iteration takes \( O(|G|n^k+1) \) time (fixed-point iteration method) or \( O(|G|^3n^{3(k+1)}) \) time (Newton’s method), but the number of iterations depends on \( G \).

Finally, we note that we can reduce the sizes of the right-hand sides of a FGG by a process analogous to binarization of CFGs (Gildea, 2011; Chiang et al., 2013):

**Proposition 16.** For any hypergraph fragment \( R \), let \( \bar{R} \) be the hypergraph formed by adding a hyperedge connecting \( R \)'s external nodes. Let \( G \) be a HRG, \( n_G \) be the total number of nodes in its right-hand sides, and \( k \) be such that for every right-hand side \( R \), the treewidth of \( \bar{R} \) is at most \( k \). Then there is an equivalent HRG with at most \( n_G \) rules whose right-hand sides have at most \( (k+1) \) nodes.

**Proof.** See Appendix C.

### 5.2 Finite graph languages

Next, we show that a nonrecursive FGG can also be converted into an equivalent factor graph, such that the sum–product of the factor graph is equal to the sum–product of the FGG. This makes it possible to use standard graphical model inference techniques for reasoning about the FGG, even with infinite variable domains. However, the conversion increases treewidth in general, so when the method of Section 5.1 is applicable, it should be preferred.

The construction is similar to constructions by Smith and Eisner (2008) and Pynadath and Wellman (1998) for dependency parsers and PCFGs, respectively. Their constructions and ours encode a set of possible derivations as a graphical model, using hard constraints to ensure that every assignment to the variables corresponds to a valid derivation.

**Theorem 17.** Let \( G = (N, T, P, S) \) be a nonreentrant FGG. Let \( n_G \) and \( m_G \) be the total number of nodes and edges in the right-hand sides of \( G \) respectively. Then \( G \) can be converted into a factor graph with \( O(n_G) \) variables and \( O(n_G + m_G) \) factors which gives the same sum–product.

**Proof.** We construct a factor graph that encodes all derivations of \( G \). (Example 31 in Appendix D shows an example of this construction for a toy FGG.) First, we add binary variables (with label \( B \) where \( \Omega(B) = \{ \text{true}, \text{false} \} \)) that switch on or off parts of the factor graph (somewhat like the gates of Minka and Winn (2008)). For each nonterminal \( X \in N \), we add \( B_X \), indicating whether \( X \) is used in the derivation, and for each rule \( \pi \in P \), we add \( B_{\pi} \), indicating whether \( \pi \) is used.

Next, we create factors that constrain the \( B \) variables so that only one derivation is active at a time. We write \( P^X \) for the set of rules with left-hand side \( X \), and \( P_{\rightarrow X} \) for the set of rules which have a right-hand side edge labeled \( X \). Define the following function:

\[
\text{CondOne}_l(B, B_1, \ldots, B_l) = \begin{cases} 
\exists! i \in \{1, \ldots, l\}, \ B_i & \text{if } B = \text{true} \\
\neg(B_1 \lor \cdots \lor B_l) & \text{if } B = \text{false} 
\end{cases}
\]

Then we add these factors, which ensure that if one of the rules in \( P_{\rightarrow X} \) is used (or \( X = S \)), then exactly one rule in \( P^X \) is used; if no rule in \( P_{\rightarrow X} \) is used (and \( X \neq S \)), then no rule in \( P^X \) is used.

- For the start symbol \( S \), add a factor \( e \) with \( \text{att}(e) = B_S \) and \( F(e)(B_S) = (B_S = \text{true}) \).
- For \( X \in N \setminus \{S\} \), let \( P_{\rightarrow X} = \{\pi_1, \ldots, \pi_l\} \) and add a factor \( e \) with \( \text{att}(e) = B_X B_{\pi_1} \cdots B_{\pi_l} \) and \( F(e) = \text{CondOne}_l \).
- For \( X \in N \), let \( P^X = \{\pi_1, \ldots, \pi_l\} \) and add a factor \( e \) with \( \text{att}(e) = B_X B_{\pi_1} \cdots B_{\pi_l} \) and \( F(e) = \text{CondOne}_l \).

Next, define the function:

\[
\text{Cond}(B, x) = \begin{cases} 
x & \text{if } B = \text{true} \\
1 & \text{otherwise.}
\end{cases}
\]
For each rule \( \pi \in P \), where \( \pi = (X \rightarrow R) \) and \( R = (V, E_N \cup E_T, \text{att}, \text{lab}^V, \text{lab}^E) \), we construct a “cluster” \( C_\pi \) of variables and factors:

- For each \( v \in V \), add a variable \( v' \) with the same label to \( C_\pi \). Also, add a factor with endpoints \( B_\pi \) and \( v' \) and function \( \text{CondNormalize}_{v'}(B_\pi, v') \), defined to equal \( \text{Cond}(\neg B_\pi, p(v')) \), where \( p \) is any probability distribution over \( \Omega(v') \). This ensures that if \( \pi \) is not used, then \( v' \) will sum out of the sum–product neatly.

- For each \( e \in E_T \) where \( \text{att}(e) = v_1 \ldots v_k \), add a new edge \( e' \) with \( \text{att}(e') = B_\pi v_1' \ldots v_k' \) and function \( \text{CondFactor}_{e'}(B_\pi, v_1', \ldots, v_k') \), defined to equal \( \text{Cond}(B_\pi, F(e)(v_1', \ldots, v_k')) \).

Next, for each \( X \in N \), let \( l = |\text{type}(X)| \). We create a cluster \( C_X \) containing variables \( v_{X,i} \) for \( i = 1, \ldots, l \), which represent the endpoints of \( X \), such that \( \text{lab}^V(v_{X,i}) = \text{type}(X)_i \). We give each an accompanying factor with endpoints \( B_\pi \) and \( v_{X,i} \) and function \( \text{CondNormalize}_{v_{X,i}} \).

These clusters are used by the factors below, which ensure that if two variables are identified during rewriting, they have the same value. Define \( \text{CondEquals}(B, v, v') = \text{Cond}(B, v = v') \).

- For each \( \pi \in P^{-X} \), let \( v_1, \ldots, v_l \) be the endpoints of the edge in \( \pi \) labeled \( X \). (By non-reentrancy, there can be only one such edge.) For \( i = 1, \ldots, l \), create a factor \( e \) where \( \text{att}(e) = B_\pi v_{X,i} v_i \) and \( F(e) = \text{CondEquals} \).

- For each \( \pi \in P^X \), let \( \text{ext} \) be the external nodes of \( \pi \). For \( i = 1, \ldots, l \), create a factor \( e \) where \( \text{att}(e) = B_\pi v_{X,i} \text{ext}_i \) and \( F(e) = \text{CondEquals} \).

The resulting graph has \( |N| + |P| \) binary variables, \( n_G \) variables in the clusters \( C_\pi \), and \( \sum_{X \in N} |\text{type}(X)| \leq n_G \) variables in the clusters \( C_X \), so the total number of variables is in \( O(n_G) \). It has \( m_G \) \( \text{CondFactor}_{e} \) factors, \( n_G + \sum_{X \in N} |\text{type}(X)| \leq 2n_G \) \( \text{CondNormalize}_{v} \) factors, \( 2|N| \) \( \text{CondOne} \) factors, and \( 2 \sum_{(X \rightarrow R) \in P} |\text{ext}_R| \leq 2n_G \) \( \text{CondEquals} \) factors, so the total number of factors is in \( O(n_G + m_G) \).

Appendix D contains more information on this construction, including an example, a detailed proof that the sum–product is preserved, and a discussion of inference on the resulting graph. \( \square \)

### 5.3 Infinite variable domains, infinite graph languages

Finally, if we allow both (countably) infinite domains and infinite graph languages, then computing the sum–product is undecidable. This has already been observed even for single factor graphs with infinite variable domains (Dreyer and Eisner, 2009), but we show further that this can be done using a minimal inventory of factors.

**Theorem 18.** Let \( G \) be a FGG whose variable domains are \( \mathbb{N} \) and whose factors only use the successor relation and equality with zero. It is undecidable whether the sum–product of \( G \) is zero.

**Proof.** By reduction from the halting problem for Turing machines. See Appendix E. \( \square \)

### 6 Conclusion

Factor graph grammars are a powerful way of defining probabilistic models that permits practical inference. We plan to implement the algorithms described in this paper as differentiable operations and release them as open-source software. We will also explore techniques for optimizing inference in FGGs, for example, by automatically modifying rules to reduce their treewidth (Bilmes, 2010) or reducing the cost of matrix inversions in Theorem 15 (Nederhof and Satta, 2008). Another important direction for future work is the development of approximate inference algorithms for FGGs.

**Broader Impact**

This research is of potential benefit to anyone working with structured probability models, including latent-variable neural networks. As this research is purely theoretical, we are not aware of any direct negative impacts.
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A Simulating PCFGs

Example 19. Below is a FGG for derivations of a PCFG in Chomsky normal form. The start symbol of the FGG is $S'$ and the start symbol of the PCFG is $S$. Random variables $N$ range over nonterminal symbols of the PCFG, and random variables $W$ range over terminal symbols.

Example 20. We can conjoin the FGG of Example 19 with the following FGG to constrain it to an input string $w$, with $n = |w|$, $0 \leq i < j < k \leq n$, and $1 \leq l \leq n$:

The resulting rules have a total of $O(n^3)$ variables in their right-hand sides. The largest right-hand side has 3 variables, so $k = 2$. The variables range over nonterminals, so $m = |N|$ where $N$ is the CFG’s nonterminal alphabet. Therefore, running the algorithm of Theorem 15 on this FGG takes $O(nGm^{k+1}) = O(|N|^3n^3)$ time, which is the same as the CKY algorithm. This construction generalizes easily to CFGs not in Chomsky normal form; applying Lemma 16 would keep the inference complexity down to $O(n^3)$ (or $O(n^2)$ for a linear CFG).

B Relationship to other formalisms

B.1 Plate diagrams

Plate diagrams are extensions of graphs that describe repeated structure in Bayesian networks (Buntine, 1994) or factor graphs (Obermeyer et al., 2019). A plate is a subset of variables/factors, together with a finite set $B$ of plates and a function $P : V \cup E \rightarrow 2^B$ that assigns each variable and factor to a set of plates. If $b \in P(v)$ and $e$ is incident to $v$, then $b \in P(e)$.

The unrolling of $H$ by $M : B \rightarrow \mathbb{N}$ is the factor graph that results from making $M(b)$ copies of every node $v$ such that $b \in P(v)$ and every edge $e$ such that $e \in P(e)$.

Obermeyer et al. (2019) give an algorithm for computing the sum–product of a PFG. It only succeeds on some PFGs. An example for which it fails is the set of all restricted Boltzmann machines (fully-connected bipartite graphs); one of their main results is to characterize the PFGs for which their algorithm succeeds. Below, we show how to convert these PFGs to FGGs.

Proposition 22. Let $H$ be a PFG. If the sum–product algorithm of Obermeyer et al. (2019) succeeds on $H$, then there is a FGG $G$ such that for any $M : B \rightarrow \mathbb{N}$, there is a FGG $G_M$ such that $G \cap G_M$ generates one graph, namely the unrolling of $H$ by $M$.

Proof. We just describe how to construct $G \cap G_M$ directly; hopefully, it should be clear how to construct $G$ and $G_M$ separately ($G$ has factors but not counts; $G_M$ has counts but not factors). Algorithm 1 converts $H$ and $M$ to $G \cap G_M$. It has the same structure as the sum–product algorithm of Obermeyer et al. (2019) and therefore works on the same class of PFGs. \qed
Algorithm 1 Procedure for converting a PFG $H$ and count assignment $M$ to a FGG.

while $E \neq \emptyset$ do
  let $e = \arg \max_{e \in E} |P(e)|$ (breaking ties arbitrarily) and $L = P(e)$
  let $H_L$ be the subgraph of nodes and edges of $H$ whose plate set is $L$
  for each connected component $H_c$ of $H_L$, do
    let $V_f$ be the variables not in $H_c$ but incident to factors in $H_c$
    let $L' = \cup_{v \in V_f} P(v)$
    if $L = L'$ then
      error
    let $X$ be a fresh nonterminal
    let $n = \prod_{b \in L \setminus L'} M(b)$
    replace $H_c$ with an edge with label $X^n$ and endpoints $V_f$
    for $i \leftarrow n, \ldots, 1$ do
      create rule $X^i \rightarrow R$ where $R$ has:
      • internal nodes and edges from $H_c$
      • external nodes $V_f$
      • an edge with label $X^{i-1}$ and endpoints $V_f$
    create rule $X^0 \rightarrow R$ where $R$ has external nodes $V_f$ and no other nodes/edges
  create rule $S \rightarrow H$

If the algorithm of Obermeyer et al. (2019) fails on a PFG, there might not be an equivalent FGG. In particular, FGGs cannot generate the set of RBMs, because a $m \times n$ RBM has treewidth $\min(m, n)$, so the set of all RBMs has unbounded treewidth and can’t be generated by a HRG.

Although, in this respect, FGGs are less powerful than PFGs, we view this as a strength, not a weakness. Because FGGs inherently generate graphs of bounded treewidth, our sum–product algorithm (Theorem 15) works on all FGGs, and no additional constraints are needed to guarantee efficient inference.

Example 23. The following PFG is from Obermeyer et al. (2019):

Converting to a FGG produces the following rules (in order of their construction by the above algorithm):

1. $X \rightarrow A \rightarrow Y_2 
   \rightarrow X_1 \rightarrow Y_3 \rightarrow X_2 \rightarrow A^{j-1} 
   1 \leq j \leq J$

2. $X \rightarrow A \rightarrow Y_2 
   \rightarrow X_1 \rightarrow Y_3 
   1 \leq i \leq I$

3. $X \rightarrow B \rightarrow X_1 
   \rightarrow X_2 \rightarrow A^{j-1} \rightarrow Y_2 
   \rightarrow G 
   1 \leq j \leq J$

4. $S \rightarrow F \rightarrow X_1 \rightarrow B^{j-1} \rightarrow X_2$
B.2 Dynamic graphical models

For simplicity, we only consider binary factors, which we draw as directed edges, and we ignore edge labels.

**Definition 24.** A dynamic graphical model or DGM (Bilmes, 2010) is a tuple $(H_1, H_2, H_3, E_{12}, E_{22}, E_{23})$, where the $H_i = (V_i, E_i)$ are factor graphs and the $E_{ij} \subseteq V_i \times V_j$ are sets of edges from $H_i$ to $H_j$.

A DGM specifies how to construct, for any length $n \geq 2$, a factor graph

$$H^n = (V_1 \cup V_2 \times \{1, \ldots, n\} \cup V_3, E),$$

where $E$ is defined by:

- If $(u, v) \in E_{12}$, add an edge from $u$ to $(v, 1)$.
- If $(u, v) \in E_{22}$, add an edge from $(u, i - 1)$ to $(v, i)$ for all $1 < i \leq n$.
- If $(u, v) \in E_{23}$, add an edge from $(u, n)$ to $v$.

**Proposition 25.** Given a DGM $D = (H_1, H_2, H_3, E_{12}, E_{22}, E_{23})$, there is a FGG $G$ such that for any count $n \geq 2$, there is another FGG $G_n$ such that $G \cap G_n$ generates exactly one graph, the unrolling of $D$ by $n$.

**Proof.** Again, we give an algorithm for constructing $G \cap G_n$, and hopefully, it should be clear how to construct $G$ and $G_n$ separately. Create the following rules:

- $S \rightarrow R$, where $R$ contains
  - Nodes and edges from $H_1, H_2$, and $E_{12}$
  - An edge labeled $A^{n-1}$ and endpoints $\{u \mid (u, v) \in E_{22}\}$.
- $A^i \rightarrow R$, where $R$ contains
  - Nodes and edges from $H_2$
  - If $(u, v) \in E_{22}$, $R$ has an external node $u'$
  - For each $(u, v) \in E_{22}$, an edge from $u'$ to $v$
  - An edge labeled $A^{i-1}$ and endpoints $\{u \mid (u, v) \in E_{22}\}$.
- $A^1 \rightarrow R$, where $R$ contains
  - Nodes and edges from $H_2, H_3$, and $E_{23}$
  - If $(u, v) \in E_{22}$, $R$ has an external node $u'$
  - For each $(u, v) \in E_{22}$, an edge from $u'$ to $v$.

**Example 26.** Bilmes (2010) give the following example of a DGM. All factors have two endpoints, and we draw them as directed edges instead of the usual squares. We draw the edges in $E_{22}$ with dotted lines.

![Diagram of a DGM example]

The resulting FGG:
where, in the middle rule, $1 < i < n$.

Running the algorithm of Theorem 15 would not be guaranteed to achieve the same time complexity as that of (Bilmes and Bartels, 2003), which searches through alternative ways of dividing the unrolled factor graph into time slices.

### B.3 Case–factor diagrams and sum–product networks

Case–factor diagrams (McAllester et al., 2008) and sum–product networks (Poon and Domingos, 2011) are compact representations of probability distributions over assignments to Boolean variables. They generalize both Markov networks and PCFGs.

Both formalisms represent models as rooted directed acyclic graphs (DAGs), with edges directed away from the root, in which some nodes mention variables. If $D$ is a DAG, for any node $v \in D$, let $\text{scope}(v)$ be the set of variables mentioned in $v$ or any descendant of $v$.

**Definition 27.** A case–factor diagram (CFD) model is a pair $(D, \Psi)$, where $D$ is a rooted DAG with root $r$, each of whose nodes is one of the following:

- **case**$(x)$ with two children $v_1$ and $v_2$, where $x$ is a variable not in $\text{scope}(v_1) \cup \text{scope}(v_2)$.
- **factor** with two children $v_1$ and $v_2$, where $\text{scope}(v_1) \cap \text{scope}(v_2) = \emptyset$.
- **unit** with no children.
- **empty** with no children.

And $\Psi : \text{scope}(r) \rightarrow \mathbb{R}_{\geq 0}$ assigns a cost to each variable in $\text{scope}(r)$.

A CFD model defines a probability distribution over assignments to its variables. We compute quantities $q(v, \xi)$ and $Z(v)$ for each node $v$ as follows. Let $v_1, v_2$ be the children of $v$, if any.

\[
\begin{align*}
    v &= \text{case}(x) & q(v, \xi) &= \begin{cases} e^{-\Psi(x)} q(v_1, \xi) & \text{if } \xi(x) = 1 \\ q(v_2, \xi) & \text{if } \xi(x) = 0 \end{cases} & Z(v) &= e^{-\Psi(x)} Z(v_1) + Z(v_2) \\
    v &= \text{factor} & q(v, \xi) &= q(v_1, \xi) q(v_2, \xi) & Z(v) &= Z(v_1) Z(v_2) \\
    v &= \text{unit} & q(v, \xi) &= 1 & Z(v) &= 1 \\
    v &= \text{empty} & q(v, \xi) &= 0 & Z(v) &= 0
\end{align*}
\]

Define $q(\xi) = q(r, \xi)$ and $Z = Z(r)$. Then $P(\xi) = q(\xi)/Z$.

**Proposition 28.** If $(D, \Psi)$ is a CFD model, there is a FGG $G$ such that $(D, \Psi)$ and $G$ have the same sum–product, and for any assignment $\xi$ of $(D, \Psi)$, there is a FGG $G_\xi$ such that the sum–product of $G \wedge G_\xi$ equals $q(\xi)$.
Proof. Given a CFD, we can construct a FGG where each node \( v \) of the CFD becomes a different nonterminal symbol \( D_v \):

| node           | G              | \( G_\xi \)          |
|----------------|-----------------|-----------------------|
| \( v = \text{case}(x) \) | \( D_v \rightarrow \xi \) | \( D_v \rightarrow \xi \) |
|                | \( x = 1 \) | \( x = \xi(x) \) |
| \( v = \text{factor} \) | \( D \rightarrow D_{v_1} D_{v_2} \) | \( D \rightarrow D_{v_1} D_{v_2} \) |
| \( v = \text{unit} \) | \( \text{unit} \rightarrow \emptyset \) | \( \text{unit} \rightarrow \emptyset \) |

We do not create any rule with left-hand side empty, so that any derivations that generate empty fail. \( \square \)

The number of rules in \( G \) is the number of nodes in \( D \). Computing its sum–product is linear in the number of rules, just as computing the sum–product of \( D \) is linear in the number of nodes.

**Definition 29.** A valid sum–product network (SPN) is a rooted DAG whose nodes are each either:
- \( \text{sum}(\lambda_1, \lambda_2) \) with two children \( v_1 \) and \( v_2 \), where \( \text{scope}(v_1) = \text{scope}(v_2) \).
- \( \text{product} \) with two children \( v_1 \) and \( v_2 \) such that no variable appears in one and negated in the other.
- \( x \) or \( \bar{x} \) with no children.

A valid SPN defines a distribution over assignments to its variables. For each node \( v \), let \( v_1, v_2 \) be the children of \( v \), if any.

\[
\begin{align*}
\text{sum}(\lambda_1, \lambda_2) & \quad q(v, \xi) = \lambda_1 q(v_1, \xi) + \lambda_2 q(v_2, \xi) \\
\text{product} & \quad q(v, \xi) = q(v_1, \xi) q(v_2, \xi) \\
x & \quad q(v, \xi) = \xi(x) \\
\bar{x} & \quad q(v, \xi) = 1 - \xi(x)
\end{align*}
\]

Converting a valid SPN to a FGG is straightforward, but the resulting FGG has a separate node for each occurrence of a variable \( x \). The syntactic constraints in the definition of valid SPN ensure that in any graph with nonzero weight, all occurrences of \( x \) have the same value.

**Proposition 30.** Any valid SPN \( S \) can be converted into a FGG \( G \) such that \( S \) and \( G \) have the same sum–product, and for any assignment \( \xi \) of \( S \), there is a FGG \( G_\xi \) such that the sum–product of \( G \land G_\xi \) equals \( q(\xi) \).
**Proof.** We construct a FGG where each node $v$ becomes a different nonterminal symbol $D_v$:

| node | $G$ | $G_\xi$ |
|------|-----|---------|
| $v = x$ | $D_v \rightarrow x \quad x = 1$ | $D_v \rightarrow x \quad x = \xi(x)$ |
| $v = \bar{x}$ | $D_v \rightarrow \bar{x} \quad x = 0$ | $D_v \rightarrow \bar{x} \quad x = \xi(x)$ |
| $v = \text{sum}(\lambda_1, \lambda_2)$ | $D_v \rightarrow D_{v_1} \quad \lambda_1 \quad D_v \rightarrow D_{v_1}$ | $D_v \rightarrow D_{v_2} \quad \lambda_2 \quad D_v \rightarrow D_{v_2}$ |
| $v = \text{product}$ | $D_v \rightarrow D_{v_1} \quad D_{v_2}$ | $D_v \rightarrow D_{v_1} \quad D_{v_2}$ |

The number of rules in $G$ is the number of nodes in $S$. Computing its sum–product is linear in the number of rules, just as computing the sum–product of $S$ is linear in the number of nodes.

Further variations of SPNs have been proposed, in particular to generate repeated substructures (Stuhlmüller and Goodman, 2012; Melibari et al., 2016). Factored SPNs (Stuhlmüller and Goodman, 2012) are especially closely related to FGGs, in that they allow one part of a SPN to “reference” another, which is analogous to a nonterminal-labeled edge in a FGG.

CFDs and SPNs present a rather different, lower-level view of a model than the other formalisms surveyed here do. Whereas factor graphs and the other formalisms represent the model’s variables and the dependencies among them, CFDs and SPNs (including factored SPNs) represent the computation of the sum–product. For instance, converting a factor graph $H$ to a CFD or SPN requires forming a tree decomposition of $H$ (McAllester et al., 2008), and the resulting CFD/SPN’s structure is that of the tree decomposition, not of $H$.

FGGs, in a sense, combine both points of view. Their derived graphs represent a model’s variables and dependencies, while their derivation trees represent the computation of the sum–product. Thus, a factor graph $H$ can be trivially converted into a FGG $S \rightarrow H$, and, as can be seen in the translations given above, a CFD or SPN can also be converted to a FGG while preserving its structure.

### C Proof of Proposition 16

Let $H = (V, E)$ be a hypergraph. Recall that a tree decomposition of $H$ is a tree whose nodes are called bags, to each of which is associated a set of nodes, $V_B \subseteq V$, and (nonstandardly) a set of edges, $E_B \subseteq E$. The bags must satisfy the properties:

- **Node cover:** $\bigcup B V_B = V$.
- **Edge cover:** for every edge $e \in E$, there is exactly one bag $B$ such that $e \in E_B$ and $\text{att}(e) \subseteq V_B$.
- **Running intersection:** if $v \in V_{B_1}$ and $v \in V_{B_2}$, then for every bag $B$ between $B_1$ and $B_2$, $v \in V_B$.

The width of a tree decomposition is $\max_B |V_B| - 1$, and the treewidth of $H$ is the minimum width of any tree decomposition of $H$. A tree decomposition can always be made to have at most $n$ nodes without changing its width (Bodlaender, 1993).

Chiang et al. (2013) give a parsing algorithm for HRGs that matches right-hand sides incrementally using their tree decompositions. They observe that this is related to the concept of binarization of context-free grammars. Here, we make this connection explicit by showing how to factorize a HRG.
For every rule \((X \rightarrow R)\), where \(\bar{R}\) has \(n_R\) nodes and treewidth at most \(k\), form a tree decomposition of \(\bar{R}\) with \(n_R - k \leq n_R\) bags. Let the root of the tree decomposition be the bag containing all the external nodes of \(\bar{R}\). For each bag \(B\), construct a rule \(X_B \rightarrow R_B\) as follows.

- If \(B\) is the root bag, \(X_B = X\); otherwise, \(X_B\) is a fresh nonterminal symbol.
- Add all nodes in \(V_B\) and edges in \(E_B\) to \(R_B\).
- If \(B\) is the root bag, \(R_B\)'s external nodes are the same as \(R\)'s; if \(B\) has parent \(P\), let \(R_B\)'s external nodes be \(V_P \cap V_B\).
- For each child bag \(B_i\), add a hyperedge with label \(X_{B_i}\) and endpoints \(V_B \cap V_{B_i}\).

This new FGG generates the same language as \(G\). The number of rules is at most \(\sum_{(X \rightarrow R) \in G} n_R = n_G\). Every right-hand side has at most \((k + 1)\) nodes.

### D Supplement to Theorem 17

#### D.1 An example

**Example 31.** We show how to construct the factor graph corresponding to the following simple, nonreentrant FGG:

\[
\begin{align*}
S &\xrightarrow{\pi_1} A_1 \quad B_2 \\
S &\xrightarrow{\pi_2} A_1 \quad Y_3 \quad B_2 \\
\end{align*}
\]

\[
\begin{align*}
A_1 &\xrightarrow{\pi_3} f(A_1, A_4) \\
A_4 &\xrightarrow{\pi_4} g(A_1, B_2) \\
\end{align*}
\]

This grammar generates just two graphs:

\[
\begin{align*}
A_1 &\xrightarrow{\pi_3} f(A_1, A_4) \\
A_4 &\xrightarrow{\pi_4} g(A_4, B_2) \\
\end{align*}
\]

\[
\begin{align*}
A_1 &\xrightarrow{\pi_4} g(A_1, B_2) \\
B_2 &\xrightarrow{\pi_3} f(A_1, A_4) \\
\end{align*}
\]

Applying the construction from Theorem 17 gives the factor graph shown in Figure 4.
Figure 4: The FGG of Example 31, converted to a single factor graph using the construction of Theorem 17. Some detail has been omitted to reduce clutter. The edges from clusters to \( B \) variables are “meta-tentacles” that stand for a tentacle from every factor inside the cluster to the \( B \) variable. We draw CondNormalize and CondEquals factors as smaller squares and omit their names. Lastly, rather than writing out “CondFactor”, we use the name of the original factor function (\( f \) or \( g \)).
D.2 Complexity of inference

As noted in Section 5.2, the purpose of this conversion to a single factor graph is to make inference possible with infinite variable domains; after converting to a factor graph, existing, possibly approximate, inference methods can be applied. But with finite variable domains, an algorithm like variable elimination would not be appropriate because this conversion has the potential to increase treewidth dramatically.

In the proof of Theorem 15, we constructed the nonterminal graph, which has a node for every nonterminal and an edge from X to Y iff there is a rule X → R where R has an edge labeled Y. For a nonreentrant FGG, the nonterminal graph is always a DAG. If, for each X ∈ N \ S, X appears in the right-hand side of exactly one rule, then the nonterminal graph is a tree.

When the nonterminal graph is a tree, we can construct a tree decomposition by making one bag with assignments to D. Then: all but 1·|B| variables are consistent with some derivation, they all have value 1. (Note that the product over \(\xi\) includes many possible settings of the B variables. But the CondOne factors tell us that, if the assignment to the B variables does not give us a valid derivation, then the weight of that assignment will be 0. Therefore, we only need to sum over assignments to the B variables which represent a valid derivation, and so we can express the sum–product using a sum over derivations rather than a sum over assignments to B variables. Let \(\xi_B\) represent the assignment to the B variables. Then:

\[
Z_H = \sum_{D \in D(G)} \sum_{\xi_B \in \Xi_H \text{ consistent with } D} \prod_{e \in H} F(e)(\xi(e)).
\]

(Note that the product over \(e \in H\) can ignore all CondOne factors, since when the assignment to the B variables is consistent with some derivation, they all have value 1.)

We can associate a derivation D with the subset of clusters in H corresponding to the nonterminals and rules which were used in the derivation; call this \(C_D\). For any D, all the variables in H are divided into three parts: those that belong to clusters in \(C_D\) (call this \(V_D\)), those that belong to clusters not in \(C_D\) (call this \(V_{\overline{D}}\)), and the B variables (which don’t belong to any cluster). Let \(\xi_{B,D}\) be the unique assignment to the B variables that is consistent with D. Let \(\Xi_D\) be the set of all assignments extending \(\xi_{B,D}\) with assignments to \(V_D\), and let \(\Xi_{\overline{D}}\) be the set of all assignments extending \(\xi_{B,D}\) with assignments to \(V_{\overline{D}}\).
Let $E_D$ be the set of factors involving a variable in $V_D$, and let $E_{\Pi}$ be the set of factors involving a variable in $V_{\Pi}$. Because any factors between $V_D$ and $V_{\Pi}$ are CondEquals factors with value 1 (since their $B$ variable is false), we can ignore them. Similarly, the only factors which don’t involve either $V_D$ or $V_{\Pi}$ are the CondOne factors, which we are already ignoring. This allows us to rewrite the sum–product as

$$Z_H = \sum_{D \in D} \left( \sum_{\xi \in \Xi_D} \prod_{e \in E_D} F(e)(\xi(e)) \right) \left( \sum_{\xi \in \Xi_{\Pi}} \prod_{e \in E_{\Pi}} F(e)(\xi(e)) \right).$$

Consider $Z_{\Pi}$ first. All CondFactor and CondEquals factors in $E_{\Pi}$ have value 1 and can be ignored, leaving only CondNormalize factors. Because these place a probability distribution $p_v$ on each variable $v$ in an unused cluster, those variables all sum out:

$$Z_{\Pi} = \sum_{\xi \in \Xi_{\Pi}} \prod_{C \in C_{\Pi}, \xi \in C_{\Pi}} \prod_{v \in C_{\Pi}} \text{CondNormalize}_v(B_{\Pi}, \xi)$$

$$= \sum_{\xi \in \Xi_{\Pi}} \prod_{C \in C_{\Pi}, \xi \in C_{\Pi}} \prod_{v \in C_{\Pi}} p_v(\xi(v))$$

$$= \prod_{C \in C_{\Pi}, \xi \in C_{\Pi}} \left( \sum_{x \in \Omega(v)} p_v(x) \right) \prod_{v \in C_{\Pi}} \left( \sum_{x \in \Omega(v)} p_v(x) \right)$$

$$= 1.$$

Now consider $Z_D$. All CondNormalize factors in $E_D$ have value 1 and can be ignored, leaving only CondEquals and CondFactor factors. Let $H_D$ be the derived graph of $D$. We can think of the derivation as merging pairs of nodes in $V_D$, so that a single node $v \in H_D$ may correspond to several “copies” in $V_D$. However, the CondEquals constraints ensure that all copies of $v$ have the same value. Therefore, instead of summing over the assignments to $V_D$, we can simply sum over the assignments to $H_D$ (and omit CondEquals factors):

$$Z_D = \sum_{\xi \in \Xi_{H_D}} \prod_{C_{\epsilon} \in C_{H_D}, \xi \in \epsilon} \text{CondFactor}_{\epsilon}(B_{\Pi}, \xi(\text{att}(\epsilon)))$$

$$= \sum_{\xi \in \Xi_{H_D}} \prod_{C_{\epsilon} \in C_{H_D}, \xi \in \epsilon} F(e)(\xi(\text{att}(\epsilon)))$$

$$= \sum_{\xi \in \Xi_{H_D}} \prod_{e \in H_D} F(e)(\xi(\text{att}(\epsilon))).$$

So, finally, the sum–product of $H$ can be rewritten as:

$$Z_H = \sum_{D \in D(G)} \sum_{\xi \in \Xi_{H_D}} \prod_{e \in H_D} F(e)(\xi(\text{att}(\epsilon)))$$

$$= \sum_{D \in D(G)} \sum_{\xi \in \Xi_{H_D}} w_G(D, \xi)$$

$$= Z_G.$$

E  Proof of Theorem 18

Let $\Gamma$ be a finite alphabet containing a blank symbol ($\omega$), and let $k = |\Gamma|$. Number the symbols in $\Gamma$ as $\gamma_0 = \omega, \gamma_1, \gamma_2, \ldots, \gamma_{k-1}$. Define an encoding for strings over $\Gamma$:

$$\langle e \rangle = 0$$

$$\langle \gamma_i w \rangle = i + k \cdot \langle w \rangle.$$

Note that strings that differ only in the number of trailing blanks have the same encoding.
We write $x / k$ for $\lfloor x/k \rfloor$ and $x \% k = x - x / k \cdot k$.

Let $M$ be a Turing machine with doubly-infinite tape, input alphabet $\Sigma$, tape alphabet $\Gamma$, start state $q_0$, transition function $\delta$, accept state $q_{\text{accept}}$, and reject state $q_{\text{reject}}$. For any input string $w \in \Sigma^*$, construct the following rules, where the $q$ nodes track the Turing machine’s state, the $u$ nodes track the reverse of the tape to the left of the head, and the $v$ nodes track the tape from the head rightward:

For each transition $\delta(q, a) = (r, b, L)$:

For each transition $\delta(q, a) = (r, b, R)$:

The sum-product of this FGG is 1 if $M$ halts on $w$, 0 otherwise. Therefore, computing the sum-product of an FGG is undecidable.

The operations $+, \cdot, /, \%$ and $=$ can be further reduced to just the successor relation and equality with zero, as shown below.
Integer division and remainder can both be computed using the rule:

where \( x_7 \) is the dividend, \( x_2 \) is the divisor, \( x_1 \) is the quotient, and \( x_3 \) is the remainder.