Chapter 4

Propagation in Bayesian networks

This chapter presents the algorithm used in HUGIN for probability updating in Bayesian networks. The algorithm does not work directly on the Bayesian network, but on a so-called junction tree which is a tree of clusters of variables. The clusters are also called cliques because they are cliques in a triangulated graph, which is a special graph constructed over the network. Each clique holds a table over the configurations of its variables, and HUGIN propagation consists of a series of operations on these tables. The subjects in this chapter are rather mathematical, and the reader interested in the results rather than in the reasoning behind them can jump directly to the summary in Section 4.7, which should give sufficient background for the reading of Chapters 5 and 6.

In Section 4.1 we define the multiplication and division of tables to be used in the algorithm. Section 4.2 gives methods for entering evidence and updating probabilities provided the full joint probability table is available, and in Section 4.3 we give the architecture of the algorithm when the cluster tree is available. Section 4.4 defines the concept junction tree, and we prove the correctness of the algorithm when applied on a junction tree. Section 4.5 is devoted to the construction of a junction tree from the Bayesian network.

The HUGIN algorithm yields the exact updated probabilities, but if you are unlucky, the algorithm will require so much space or time that the task is intractable. In Section 4.6 we present a technique, stochastic simulation, which can be used to get approximate probabilities when this happens.

4.1 An algebra of belief tables

Before we treat probability updating, we will introduce more formally the multiplication of belief tables, which we have used implicitly already.
We shall use the following proposition later.

**Proposition 4.1**

Let \( t(c^*) \) be a table over \( W \) and \( t' \) be a table over \( V \). Then

\[
\sum_{v} (t_w \cdot t_v) = t_w \cdot \sum_{v} t_v.
\]

**4.1.1 Multiplication and division**

Let \( t \) and \( t' \) be two tables over the same variables. Then the product \( t \cdot t'(c^*) = t(c^*) \cdot t'(c^*) \) for all configurations \( c^* \).

Table 4.1 gives an example.

If the two tables are over different sets of variables we can also perform a multiplication.

Let \( t_{AB} \) be a table over \( \{A, B\} \), and let \( t_{AC} \) be a table over \( \{A, C\} \). Then \( t_{AB} \) and \( t_{AC} \) are multiplied by constructing a table \( t_{ABC} \) over \( \{A, B, C\} \), and letting

\[
t_{AB} \cdot t_{AC}(a, b, c) = t_{AB}(a, b) \cdot t_{AC}(a, c)
\]

for all configurations \( (a, b, c) \).

See Table 4.2 for an example.

**4.1.2 Marginalization**

Let \( t_{V} \) be a table over \( V \), and let \( W \) be a subset of \( V \). A table \( t_{W} \) over \( W \) can be constructed by marginalization. For each configuration \( v^* \), let \( t_{W}(v^*) \) be the sum of all \( t_{V}(v^*) \), where \( v^* \) is a configuration of \( V \) coinciding with \( v^* \). The notation is

\[
t_{W} = \sum_{v \in V \setminus W} t_{V}.
\]

We shall use the following proposition later.

**Proposition 4.1**

Let \( W \) and \( V \) be disjoint sets of variables, and let \( t_{W} \) and \( t_{V} \) be tables over \( W \) and \( V \). Then

\[
\sum_{V} (t_w \cdot t_v) = t_w \cdot \sum_{V} t_v.
\]

That is, tables containing only variables over which you do not marginalize can be taken out of marginalization. See Table 4.3 for an example.

**4.2 Probability updating in joint probability tables**

Let \( A \) be a variable with \( P(A) = (x_1, \ldots, x_n) \). Assume we get the information \( e \) that \( A \) can only be in states \( i \) and \( j \). This statement says that all states except \( i \) and \( j \) are impossible, and we have the belief \( P(A, e) = (0, \ldots, 0, x_i, 0, \ldots, x_j, 0, \ldots, 0) \). Note that \( P(e) \), the prior probability of \( e \), is \( x_i + x_j \), the sum of the probabilities of the possible states. To calculate \( P(A | e) \) we use the fundamental rule:

\[
P(A | e) = \frac{P(A, e)}{P(e)} = \frac{P(A, e)}{\sum_{A} P(A, e)}.
\]

The way that \( e \) is entered can be interpreted as a multiplication of \( P(A) \) with the table \( e = (0, \ldots, 0, 1, 0, \ldots, 0, 0, \ldots) \) resulting in \( P(A, e) \).

**Definition.** Let \( A \) be a variable with \( n \) states. A **finding** on \( A \) is an \( n \)-dimensional table of zeros and ones.

Semantically, a finding is a statement that certain states of \( A \) are impossible.

Now, let \( U \) be a universe of variables, and assume that we have easy access to \( P(U) \), the joint probability table. Then, \( P(B) \) for any variable \( B \) in \( U \) is easy to calculate:

\[
P(B) = \sum_{U \setminus \{B\}} P(U).
\]

Suppose we wish to enter the above finding. Then \( P(U, e) \) is the table resulting from \( P(U) \) by giving all entries with \( x^* \) in state \( i \) or \( j \) the value zero and leaving the other entries unchanged. Again, \( P(e) \) is the sum of all entries in \( P(U, e) \) and

\[
P(U | e) = \frac{P(U, e)}{P(e)} = \frac{P(U, e)}{\sum_{U} P(U, e)}.
\]
Note that \( P(U, e) \) is the product of \( P(U) \) and the finding \( e \). If \( e \) consists of several findings \( \{f_1, \ldots, f_m\} \) each finding can be entered separately, and \( P(U, e) \) is the product of \( P(U) \) and the findings \( f_i \). We can express the considerations above in the following theorem.

**Theorem 4.1** Let \( U \) be a universe of variables and let \( e = \{f_1, \ldots, f_m\} \). Then

\[
P(U, e) = P(U) \cdot \prod_{f_i}^m f_i \quad \text{and} \quad P(U, e) = \frac{P(U \mid e)}{P(e)}
\]

where

\[
P(e) = \sum_U P(U, e).
\]

Theorem 4.1 says that if we have access to \( P(U) \), then we can enter evidence and perform probability updating. However, even for small sets of variables, the table \( P(U) \) is intractably large, and we have to find a smaller representation.

### 4.3 Cluster trees

As shown in Section 2.3.7 (the chain rule), a Bayesian network over \( U \) is a representation of \( P(U) \). This means that we can, in principle, calculate \( P(U) \) as the product of all conditional probabilities from the network. The question then is, whether we can enter evidence and perform probability updating in Bayesian networks without being forced to calculate \( P(U) \). It has turned out to be rather difficult.

Instead we can work with another representation called cluster trees.

**Definition.** A cluster tree over \( U \) is a tree of clusters of variables from \( U \). The nodes are subsets of \( U \), and the union of all nodes is \( U \). (A tree is an undirected graph without cycles.)

The links are labelled with *separators* which consist of the intersection of the adjacent nodes.

Each node and separator holds a real numbered table over the configurations of its variable set.

In Figure 4.1 we give a cluster tree for the network \( M_{\text{min}} \).

![Figure 4.1](Image)

**Figure 4.1** The Bayesian network \( M_{\text{min}} \) and a corresponding cluster tree. Separators are in square boxes.

Now, let \( BN \) be a Bayesian network over \( U \). A cluster tree corresponding to \( BN \) is constructed in the following way:

- form a family of nodes such that for each variable \( A \) with parent set \( \text{pa}(A) \) there is at least one node \( V \) such that \( \text{pa}(A) \cup \{A\} \subseteq V \);
- organize the nodes as a tree with separators (so far there is no restriction on how you organize the tree);
- give all nodes and separators a table of ones.
- for each variable \( A \) choose exactly one node \( V \) containing \( \text{pa}(A) \cup \{A\} \) and multiply \( P(A \mid \text{pa}(A)) \) on \( V \) table.

Then the product of all node tables in the cluster tree is the product of all conditional probability tables in \( BN \), and therefore we have the following theorem.

**Theorem 4.2** Let \( BN \) be a Bayesian network over \( U \). Then any cluster tree corresponding to \( BN \) is a representation of \( P(U) \), and \( P(U) \) is the product of all cluster tables divided by the product of all separator tables.

**Remark.** In Theorem 4.2 we divide the product of all cluster tables by the product of all separator tables. This does not do any harm, because the separator tables consist of ones, but the reader may wonder why. The reason is that, when we now start to move the information around in the cluster tree, then the product of all cluster tables divided by all separator tables is invariant, and thereby the tree remains a representation of \( P(U) \).

It is easy to insert findings into a cluster tree. Let \( e \) be a finding on \( A \). Multiply \( e \) on the table of any node containing \( A \). Then, by the chain rule and Theorem 4.1, the product of all node tables is \( P(U \cdot e) = P(U, e) \).

To calculate \( P(B, e) \) for an arbitrary variable \( B \) is not as easy, and the coming sections are devoted to this problem.

#### 4.3.1 Absorption in cluster trees

We introduce an operation in cluster trees. It has the effect of re-arranging the information stored in the tables.

**Definition.** Let \( V \) and \( W \) be neighbours in a cluster tree, let \( S \) be their separator, and let \( t_V \), \( t_W \) and \( t_S \) be their tables. The operation absorption is the result of the following procedure:

- calculate \( t_S^* = \sum_{V \in S} t_V \);
- give \( S \) the table \( t_S^* \);
- give \( W \) the table \( t_W^* = t_W t_S^* \).

We then say that \( W \) has absorbed from \( V \) or that \( W \) calibrates to \( V \).
Theorem

Then any zero-entry in tables divided by the product of all separator tables is invariant under absorption.

Proof

We have $t_S = \sum_{V \setminus S} t_V$, where $t_V = t_W \cdot t_S$, then absorption does not change anything. We then say that the link is consistent. If all links in the cluster tree are consistent we say that the tree is consistent. If a tree is consistent, then absorption does not have any effect at all.

Assume that the link is consistent, but now some evidence changes are supportive since the separator tables have no zero-entries. The idea behind absorption is that the information which is unchanged. We have $t_W = t_V$, and this is what we need to prove.

$T$ is consistent, then absorption does not have any effect at all.

Assume that the link is consistent, but now some evidence changes. Then after $W$ has absorbed from $V$, the three tables all hold the same information on $S$.

$T$ is consistent, then absorption does not have any effect at all.

Lemma 4.1 Supportiveness is preserved under absorption.

Proof. Let $W$ absorb from $V$ through the separator $S$. Then $t^*_S = \frac{t^*_V}{t^*_S}$, where $t^*_S = \sum_{V \setminus S} t_V$. Then any zero-entry in $t^*_S$ is also a zero-entry in $t^*_W$. This clearly also holds for $t^*_S$.

Theorem 4.3 Let $T$ be a supportive cluster tree. Then the product of all cluster tables divided by the product of all separator tables is invariant under absorption.

Proof. When $W$ absorbs from $V$ through the separator $S$, only the tables of $W$ and $S$ are unchanged. Therefore it is enough to prove that the fraction of combined tables is invariant under absorption.

$t^*_W = \frac{t^*_W \cdot t^*_S}{t^*_S \cdot t^*_V} = \frac{t^*_W}{t^*_V}$

Theorem 4.3 ensures that if we start with a Bayesian network over $U$, construct a corresponding cluster tree $T$, and then perform a series of absorptions, then $T$ remains a representation of $P(U)$, and $P(U)$ can be calculated as the product of all cluster tables divided by the product of all separator tables.

4.3.2 Message passing in cluster trees

The next question is how many absorptions can we perform, and can they help us in transforming the tables in a cluster tree into a form where it is easy to calculate $P(A)$ for single variables?

We can think of absorptions as messages passed between the nodes in the tree. That is, a node $V$ sends a message to its neighbour $W$ when $W$ absorbs from $V$.

Message passing scheme. A node $V$ can send exactly one message to a neighbour $W$, and it may only be sent when $V$ has received a message from each of its other neighbours.

Consider, for example, the cluster tree in Figure 4.3. The leaves of the tree (the nodes $A, B, C, D$) can send to their single neighbour (1). Then $E$ can send to $G$, and $H$ can send to $F$ (2). Next, $G$ can send to $F$, and $F$ can send to $G$ (3). $F$ can send to $H, B$ and $C$, and $G$ can send to $E$ (4). Finally $E$ can send to $A$ and $H$ to $D$ (5). Now each node has sent to all of its neighbours.

As can be seen, the message passing algorithm is not sequential, and a good way of thinking of it is that each variable is busy waiting, eager to send messages. Each time it receives a message it updates its own table and sends a message to the eligible neighbours (if any).

Theorem 4.4 Let $T$ be a supportive cluster tree, and suppose that messages are passed according to the message passing scheme. Then:
Let the first message to be passed over 

\( \text{ts} = \sum_{\text{V}} \sum_{\text{W}} \text{tw} \). Next, when the message from \( V \) and \( W \) has to be passed, the tables for \( S \) and \( W \) have not been changed (\( W \) has not received further messages). Let the table for \( V \) be \( t^*_V \). After message passing we have

\[
\text{ts}^*_V = \sum_{\text{V}} t^*_V \quad \text{and} \quad \text{tw} = \text{tw} \cdot \text{ts}.
\]

Now

\[
\sum_{\text{V}} \sum_{\text{W}} \text{tv} = \sum_{\text{V}} \sum_{\text{W}} \sum_{\text{S}} \text{tv} = \sum_{\text{V}} \sum_{\text{W}} \sum_{\text{S}} \text{tw} = \sum_{\text{V}} \sum_{\text{W}} \sum_{\text{S}} \text{ts}.
\]

Therefore the link is consistent.

## 4.4 Junction trees

Let \( T \) be a cluster tree over \( U \), let \( A \) be a variable in \( U \), and suppose that \( A \) is an element of the nodes \( V \) and \( W \). If \( T \) is consistent we would expect \( \sum_{V \cup A} \text{tv} = \sum_{W \cup A} \text{tw} \). Certainly this is so if \( V \) and \( W \) are neighbours, but otherwise there is no guarantee. See Figure 4.4 for an example.

We say that a consistent cluster tree is **globally consistent** if for any nodes \( V \) and \( W \) with intersection \( I \) we have

\[
\sum_{\text{V}} \text{tv} = \sum_{\text{W}} \text{tw}.
\]

As Figure 4.4 indicates, the reason why consistence does not imply global consistence is that a variable \( A \) can be placed in two locations in the tree such that information on \( A \) cannot be passed between the two locations. To ensure global consistence we must add a requirement to cluster trees.

**Definition.** A cluster tree is a junction tree if, for each pair of nodes \( V, W \), all nodes on the path between \( V \) and \( W \) contain the intersection \( V \cap W \).

**Theorem 4.5** A consistent junction tree is globally consistent.

**Proof:** Exercise 4.7.

The following theorems will show that if we construct a junction tree corresponding to a Bayesian network, then we have good algorithms for insertion of evidence as well as probability updating. When we construct a cluster tree corresponding to a Bayesian network we have several degrees of freedom, and we shall use them for constructing a junction tree. However, it is not easy. For example, with the clusters in Figure 4.4 it is impossible to construct a tree with the junction tree property. We will leave this problem here, and return to it in Section 4.5.

**Theorem 4.6** Let \( T \) be a consistent junction tree over \( U \), and let \( \text{tv} \) be the product of all node tables divided by the product of all separator tables. Let \( V \) be a node with table \( t_V \). Then

\[
\text{tv} = \sum_{U \cup V} \text{tv}.
\]

**Proof.** Induction on the number of nodes.

Clearly the theorem holds when \( T \) consists of a single node.

Now, assume the theorem to hold for any junction tree with \( n \) nodes, and let \( T \) be a consistent junction tree with \( n + 1 \) nodes. Let \( V \) be a leaf of \( T \) linked to \( W \) and with separator \( S \) (see Fig. 4.5). Let \( T' \) be the junction tree resulting from removing \( V \) (and \( S \)), and let \( T' \) have the universe \( U' \). Then

\[
\text{ts}_V = \text{ts}_V \cdot \frac{\text{tv}}{\text{ts}_S},
\]

where \( \text{tv} \) is the product of all node tables in \( T' \) divided by the separator tables in \( T' \). Let \( D \) be the set of variables \( V \setminus S \), and let \( H \) be \( W \setminus S \). From the junction tree property we have that \( D \cap U' = \emptyset \).
Since $T$ is consistent we have

$$\sum_{D} t_V = t_S = \sum_{W} t_W.$$  

Now

$$\sum_{D} t_U = \sum_{W} t_U = \frac{t_V}{t_S} = \frac{t_U}{t_S} = \frac{t_U}{t_S} \sum_{W} t_W.$$  

Therefore, by the induction hypothesis we have

$$\sum_{U \cup S} t_U = t_U$$

for all $V_i$ in $T$.

Furthermore,

$$\sum_{U \cup S} t_U = \sum_{U \cup S} t_U \cdot \frac{t_V}{t_S} = \sum_{W} t_W \cdot \frac{t_V}{t_S} = \frac{t_V}{t_S} \sum_{W} t_W \cdot \frac{t_V}{t_S} = \frac{t_V}{t_S}.$$  

The considerations above are summarized in the following theorem.

**Theorem 4.7** Let $BN$ be a Bayesian network representing $P(U)$, and let $T$ be a junction tree corresponding to $BN$. After a full round of message passing in $T$, we have for each node $V$ and each separator $S$ that

$$t_V = \sum_{U \cup V} P(U) = P(V)$$

and

$$t_S = P(S).$$

**Proof.** By Theorem 4.2, $P(U)$ is the product of the initial node tables divided by the separator tables. Theorems 4.3 and 4.4 give that after a full round of message passing $T$ is consistent, and $P(U)$ is the product of all node tables divided by all separator tables. Theorems 4.5 and 4.6 yield the conclusion.

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**JUNCTION TREES**

**Theorem 4.8** Let $BN$ be a Bayesian network representing $P(U)$, and let $T$ be a junction tree corresponding to $BN$. Let $e = \{f_1, \ldots, f_m\}$ be findings on the variables $\{A_1, \ldots, A_m\}$. For each $i$ find a node containing $A_i$ and multiply its table with $f_i$. Then, after a full round of message passing we have for each node $V$ and separator $S$ that

$$t_V = P(V, e)$$

and

$$t_S = P(S, e) = \sum_V t_V.$$

**Proof.** Use Theorem 4.1 and proceed as in the proof of Theorem 4.7.

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**4.4.1 HUGIN propagation**

Assume that we have a consistent junction tree, and now a single node $V$ receives evidence. Then half of the messages can be avoided: $V$ sends messages to all of its neighbours who recursively send messages to all neighbours except the one from which the message came (see Fig. 4.6). We call this algorithm $\text{DistributeEvidence}$.

Now, suppose that we are only interested in the certainty of one node, $V$. Then half of the certainty updating messages can be avoided: $V$ asks all its neighbours to send it a message, and if they are not allowed to do so, they recursively pass the request to all neighbours except the one from which the request came (see Fig. 4.7). We call this algorithm $\text{CollectEvidence}$.

The two algorithms $\text{DistributeEvidence}$ and $\text{CollectEvidence}$ can be used for a more organized message passing scheme. No matter the amount of evidence entered, take any variable $V$. Call $\text{CollectEvidence}$ from $V$ and after that call $\text{DistributeEvidence}$ from $V$. The result is that all messages have been passed, and they were passed when permitted (see Fig. 4.8 and Exercise 4.5).
Figure 4.8 Updating through CollectEvidence(V) followed by DistributeEvidence(V).

Figure 4.9 Evidence ev has been entered at the righthand side of S. ew has been entered at the lefthand side of S. C is used as a root for the propagation.

HUGIN propagation uses corresponding junction trees, and the operations CollectEvidence and DistributeEvidence. A node Rt in the junction tree is chosen as a root, and whenever a propagation takes place, CollectEvidence(Rt) is called followed by a call of DistributeEvidence(Rt). When the calls are finished, the tables are normalized so that they sum to one.

HUGIN propagation has a nice side effect, namely that it gives access to various probabilities of sets of entered findings.

Let us use Theorem 4.8 to have a closer look at what is actually communicated in the propagation algorithm. The general situation is described in Figure 4.9.

A call of CollectEvidence(C) will cause a call of CollectEvidence(V), and by Theorem 4.8 this will result in \( t' = P(V, ev) \). This gives that \( P(ev) \) can be calculated without further propagations. Unfortunately, the situation is not symmetric. In the DistributeEvidence phase the message passed from W to S is \( P(S, e) \).

4.5 Construction of junction trees

In this section we shall give a method for constructing junction trees for DAGs.

4.5.1 Singly connected DAGs

A DAG is singly connected if the graph you get by dropping the directions of the links is a tree (see Fig. 4.10).

For singly connected DAGs it is easy to construct junction trees. For each variable A with \( pa(A) \neq \emptyset \) you form the cluster \( pa(A) \cup \{A\} \). Between any two clusters with a non-empty intersection you add a link with the intersection as a separator. The resulting graph is called a junction graph. All separators consist of a single variable, and if the junction graph has cycles, then all separators on the cycle contain the same variable. Therefore any of the links can be removed to break the cycle, and by removing links until you have a tree, you get a junction tree (see Fig. 4.11).

We know that when we construct a cluster tree corresponding to a DAG, then for all variables A there must be a cluster V containing \( pa(A) \cup \{A\} \). We can illustrate this on a graph by having a link between any pair of variables which must appear in the same cluster. This means that we take the DAG, add a link between any pair of variables with a common child, and drop the directions of the original links. The resulting graph is called the moral graph. From the moral graph you can read the clusters to consider, namely the cliques in the graph (maximal sets of variables that are all pairwise linked). In Figure 4.12 we give an example of the construction.
4.5.2 Coping with cycles

Consider the junction graph in Figure 4.13. The intersection of the two clusters of variables is \((AB)\), and a junction tree is easily found.

Consider the DAG in Figure 4.14(a) with its moral graph in Figure 4.14(b). Sticking to the approach that the clusters are the cliques in the moral graph, we see that if we join \(A\), \(B\), and \(C\), then we get a junction tree.

The DAG in Figure 4.15 is more problematic. The cycle in the junction graph cannot be broken.

The propagation problem is that coupled information (on \((DE)\)) is decoupled but meets again under propagation. This can also be seen from the cycle \(D - E - C - A - B - D\) in the moral graph. A way to solve the problem is to add so-called fill-ins to the moral graph: add a link between \(C\) and \(D\) and one between \(B\) and \(C\). The result is shown in Figure 4.16 together with the resulting junction tree.

The general rule for filling-in the moral graph is that any cycle with more than three variables shall have a chord. In this case the graph is called triangulated.

In Figures 4.17 and 4.18 there is another example of the process from DAG to junction tree. Note that without the fill-in \((B - D)\) the cycle \(A - B - F - D - A\) does not have a chord.
4.5.3 From DAG to junction tree

In this section we present, without proofs, algorithms for triangulation of graphs and for construction of junction trees from triangulated graphs. Proofs of Theorems 4.9 and 4.10 are given in Appendix A.

Definition. An undirected graph is triangulated if any cycle of length $> 3$ has a chord.

Definition. A node $A$ is eliminated by adding links such that all of its neighbours are pairwise linked and then removing $A$ together with its links.

Note that if a node $A$ can be eliminated without adding links, then $A$ cannot be part of a chordless cycle of length $> 3$.

Theorem 4.9 A graph is triangulated if and only if all of its nodes can be eliminated one by one without adding any link.

Theorem 4.9 yields a method for triangulation as well as a test for whether a graph is triangulated. The method consists of eliminating the nodes in some order (adding links, if necessary) and when this is done the resulting graph is triangulated. An example is given in Figure 4.20.

Note that there are several triangulations of the graph. Intuitively, triangulations with as few fill-ins as possible are preferred. However, optimality is connected to the resulting junction tree and the computational complexity of the propagation algorithm. We shall return to the question of optimality later.

Definition. A junction graph for an undirected graph $G$ is an undirected, labelled graph. The nodes are the cliques in $G$. Every pair of nodes with a non-empty intersection has a link labelled by the intersection.

There is an easy way of identifying the cliques in a triangulated graph $G$. Let $A_1, \ldots, A_n$ be an elimination sequence for $G$, and let $C_i$ be the set of variables containing $A_i$ and all its neighbours at the time of elimination (neighbours with higher numbers). Then every clique of $G$ is a $C_i$ for some $i$.

The reader may check that the cliques of the graphs in Figure 4.20(a) are $C_1, C_2, C_3, C_4$, and that the cliques of the graph in Figure 4.20(b) are $C_1, C_2, C_3, C_4$.

The junction tree we are aiming at will be a subgraph of the junction graph. Since message passing will be restricted to links in the junction tree we are not allowed to remove a link from the junction graph if thereby some kind of information cannot be passed. If, for example, the clusters $U$ and $V$ have the variable $A$ in common, they have a link with label $A$. If this link is removed, there shall be another path in the remaining graph through which information on $A$ can be passed from $U$ to $V$. So, let us recall the following definition.

Definition. A spanning tree of a junction graph is a junction tree if it has the property that for each pair of nodes, $U, V$, all nodes on the path between $U$ and $V$ contain $U \cap V$. (A subtree of a graph is a spanning tree if all nodes of the graph are nodes in it.)
Theorem 4.10 An undirected graph is triangulated if and only if its junction graph has a junction tree.

Definition. The weight of a link in a junction graph is the number of variables in the label. The weight of a junction tree is the sum of the weights of the labels.

Theorem 4.11 (Without proof.) A subtree of the junction graph of a triangulated graph is a junction tree if and only if it is a spanning tree of maximal weight.

Theorem 4.11 provides an easy way of constructing junction trees, namely Kruskal's algorithm: choose successively a link of maximal weight unless it creates a cycle.

There are other ways of constructing junction trees. In particular, if an elimination sequence for the triangulated graph is known, very efficient algorithms exist (see Exercise 4.8). So, if the graph is triangulated then the construction of a junction tree is rather fast.

The only problematic step in the process from DAG to junction tree is the triangulation. Since any elimination sequence will produce a triangulation it may not seem a problem, but for the propagation algorithm it is. In HUGIN, propagation the cliques in the junction graph shall have joint probability tables attached to them. The size of the table is the product of the number of states of the variables. So, the size increases exponentially with the size of the clique. A good triangulation, therefore, is a triangulation yielding small cliques, or to be more precise, yielding small probability tables. The problem of determining an optimal triangulation is NP-complete. However, there is a heuristic algorithm which has proven to give fairly good results. It is a version of the greedy approach: eliminate repeatedly a node not requiring fill-ins and if this is not possible, eliminate a node yielding the smallest table. In Figure 4.23 an example is given.

4.6 Stochastic simulation

The propagation method requires tables for the cliques in the triangulated graph. These cliques may be very large, and it happens that the space requirements cannot be met by the hardware available. In this case an approximate method would be satisfactory.

In this section we shall give a flavour of an approximate method called stochastic simulation. The idea behind the simulation is that the causal model is used to simulate the flow of impact. When impact from a set of variables to a variable A is simulated, a random generator is used to decide the state of A.

To illustrate the technique, consider the Bayesian network in Figure 4.24 with the conditional probabilities specified in Table 4.4.
Figure 4.22 Junction trees for the junction graphs in Figure 4.20.

Figure 4.23 A heuristic elimination sequence is $E, D$ (and $A, B, C$).

Figure 4.24 An example network. All variables have the states $y$ and $n$.

Table 4.4 The conditional probabilities for the example network. $P(A) = (0.4, 0.6)$.

<table>
<thead>
<tr>
<th></th>
<th>$A$</th>
<th>$A$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$B$</td>
<td>$y$</td>
<td>$n$</td>
</tr>
<tr>
<td>$C$</td>
<td>$y$</td>
<td>$n$</td>
</tr>
<tr>
<td>$D$</td>
<td>$y$</td>
<td>$n$</td>
</tr>
<tr>
<td>$E$</td>
<td>$y$</td>
<td>$n$</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>$P(B \mid A)$</th>
<th>$P(C \mid A)$</th>
<th>$P(D \mid B)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$y$</td>
<td>$(0.3, 0.7)$</td>
<td>$(0.999, 0.001)$</td>
<td>$(0.999, 0.001)$</td>
</tr>
<tr>
<td>$n$</td>
<td>$(0.7, 0.3)$</td>
<td>$(0.001, 0.999)$</td>
<td>$(0.001, 0.999)$</td>
</tr>
</tbody>
</table>

Table 4.5 A set of 100 configurations of $(A, B, C, D, E)$ sampled from the network in Figure 4.24 and Table 4.4.

<table>
<thead>
<tr>
<th>$AB$</th>
<th>$CDE$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$yy$</td>
<td>$yyn$</td>
</tr>
<tr>
<td>$yy$</td>
<td>4</td>
</tr>
<tr>
<td>$yn$</td>
<td>2</td>
</tr>
<tr>
<td>$ny$</td>
<td>9</td>
</tr>
<tr>
<td>$nn$</td>
<td>0</td>
</tr>
</tbody>
</table>

The idea now is to draw a random configuration of the variables $(A, B, C, D, E)$, and to do this a sufficient number of times.

A random configuration is selected by successively sampling the states of the variables. First the state of $A$ is sampled. A random generator (with even distribution) is asked to give a real number between zero and one. If the number is less than 0.4 the state is $y$, if not the state is $n$. Assume that the result is $y$. From the conditional probability table $P(B \mid A)$ we have that $P(B \mid y) = (0.3, 0.7)$. The random generator is asked again, and if the number is less than 0.3, the state of $B$ is $y$. This procedure is repeated to get the state of $C, D,$ and $E$, and a configuration is determined.

The next configuration is sampled through the same procedure, and the procedure is repeated until $m$ configurations are sampled. In Table 4.5 an example set of configurations is given.

The probability distributions for the variables are calculated by counting in the sample set (see Exercise 4.12). For 39 of the samples in Table 4.5 the first state is $y$, and this gives an estimated probability $P(A) = (0.39, 0.61)$.

The method above, called forward sampling, does not require a triangulation of the network, and it is not necessary to store the sampled configurations (like Table 4.5); it is enough to store the counts for each variable. Whenever a sampled configuration has been determined, the counts of all variables are updated, and the sample can
be discarded. This method saves a great deal of space, and each configuration is determined in a time linear to the number of variables. The cost is accuracy and time.

So far only the initial probabilities are calculated. When evidence arrives, it can be handled by simply discarding the configurations which do not conform to it. That is, a new series of stochastic simulations are started, and whenever a state of an observed variable is drawn, you stop simulating if the state drawn is not the observed one.

Unfortunately, this method has a serious drawback. Assume in the example above that the observations for the network are \( B = n \) and \( E = n \). The probability for \((B = n, E = n)\) is 0.00282. This means that in order to get 100 configurations you should for this tiny example, expect to perform more than 35,000 stochastic simulations.

Methods have been constructed for dealing with this problem. A promising method is called Gibbs sampling.

In Gibbs sampling you start with some configuration consistent with the evidence (for example determined by forward sampling), and then you randomly change the state of the variables in causal order. In one sweep through the variables you determine a new configuration, and then you use this configuration for a new sweep, etc.

In the example let \( B = n \) and \( E = n \) be the evidence, and let the starting configuration be yyyny. Now, calculate the probability of \( A \) given the other states of that configuration. That is, \( P(A \mid B = n, C = y, D = y, E = n) \). From the network we see that it is sufficient to calculate \( P(A \mid B = n, C = y) \). It is easily done by Bayes’ rule: it is (0.8, 0.2). We draw a number from the random generator, and let us assume that the number is 0.456 resulting in \( A = y \). The next free variable is \( C \). We calculate

\[
P(C \mid A = y, B = n, D = y, E = n) = P(C \mid A = y, D = y, E = n)
\]

\[
= (0.996, 0.04).
\]

We draw from the random generator, and assume we keep \( C = y \).

In general the calculation goes as follows. Let \( A \) be a variable in a Bayesian network \( BN \), let \( B_1, \ldots, B_n \) be the remaining variables, and let \( b^* = (b_1, \ldots, b_n) \) be a configuration of \((B_1, \ldots, B_n)\). Then \( P(A \mid b^*) \) is the product of all conditional tables of \( BN \) with \( B_i \) instantiated to \( b_i \). Therefore \( P(A, b^*) \) is proportional to the product of the tables involving \( A \), and \( P(A \mid b^*) \) is the result of normalizing this product. Note that the calculation of \( P(A \mid b^*) \) is a local task.

Back to the example. The next variable is \( D \). We follow the same procedure and assume that the result is \( D = y \). Then the configuration from the first sweep is unaltered, i.e. yyyny.

The next sweep follows the same procedure. Assume the result for \( A \) is that the state is changed to \( n \). Then we shall calculate \( P(C \mid A = n, D = y, E = n) \), and so forth.

In this way a large sample of configurations consistent with the observations are produced. The question is whether the sample is representative for the probability distribution. It is not always so. It may be that the initial configuration is rather improbable, and therefore the first samples, likewise, are out of the mainstream. Therefore you usually discard the first 5–10% of the samples. This is called burn-in.

Another problem is that you may be stuck in certain “areas” of the configurations. Perhaps there is a set of very likely configurations, but in order to reach them from the one you are in, a variable should change to a state which is highly improbable given the remaining configuration (see Exercise 4.13).

A third serious problem is that it may be very hard to find a starting configuration. In fact, it is \( NP \)-hard (see Exercise 4.14).

We shall not deal with these problems, but refer the interested reader to the literature.

4.7 Summary of Sections 4.2–4.5

Junction trees

The nodes of a junction tree are sets of variables, they are called cliques. Each clique and separator holds a real numbered table over the configurations of its variable set.

The junction tree property. For each pair \( V, W \) of cliques, all cliques on the path between \( V \) and \( W \) contain the intersection \( V \cap W \). A junction tree is said to represent the Bayesian network \( BN \) over the variables \( U \) if:

(i) for each variable \( A \), there is a clique containing \( pa(A) \cup \{A\} \);

(ii) \( P(U) \) is the product of all clique tables divided by all separator tables.

Construction of junction trees

Let \( BN \) be a Bayesian network over the variables \( U \).

(i) Construct the moral graph: the undirected graph with a link between all variables in \( pa(A) \cup \{A\} \) for all \( A \).

(ii) Triangulate the moral graph: add links until all cycles consisting of more than three links have a chord.

(iii) The nodes of the junction tree are the cliques of the triangulated graph.

(iv) Connect the cliques of the triangulated graph with links such that a junction tree is constructed.

(v) First give all cliques and separators a table consisting of only ones. Then, for each variable \( A \) find a clique containing \( pa(A) \cup \{A\} \), and multiply \( P(A \mid pa(A)) \) on its table.

The resulting junction tree represents \( BN \).
Junction tree corresponding to $S \{A_1, \ldots, A_m\}$.

Correctness of sorbs from them. when all the called neighbours have finished, recursively. An arbitrary clique $HUGIN$- be represented as a table of zeros and ones with a zero at the places for impossible states. A finding on a variable $A$ is entered into a clique $V$ containing $A$ by multiplying $Vs$ table by the table for the finding.

Absorption in junction trees

Definition. Let $V$ and $W$ be neighbours in a junction tree, let $S$ be their separator, and let $t_V$, $t_W$ and $t_S$ be their tables. The operation absorption is the result of the following procedure:

- calculate $t'_S = \sum_{S \setminus S} t_V$;
- give $S$ the table $t'_S$;
- give $W$ the table $t'_W = t_W t'_S$.

We then say that $W$ has absorbed from $V$. (See Fig. 4.25.)

**HUGIN** propagation

An arbitrary clique $Rt$ in the junction tree is chosen as a root. The operation CollectEvidence is called in $Rt$ followed by a call of DistributeEvidence in $Rt$.

CollectEvidence($Rt$) asks all neighbours to CollectEvidence and they proceed down the tree recursively. When all the called neighbours have finished, $Rt$ absorbs from them. DistributeEvidence($Rt$) makes all its neighbours absorb from $Rt$, and afterwards recursively DistributeEvidence to its neighbours (except $Rt$). See Figure 4.26.

Correctness of **HUGIN** propagation

**Theorem 4.8** Let $BN$ be a Bayesian network representing $P(U)$, and let $T$ be a junction tree corresponding to $BN$. Let $e = \{f_1, \ldots, f_m\}$ be findings on the variables $\{A_1, \ldots, A_m\}$. For each $f_i$ find a node containing $A_i$ and multiply its table with $f_i$.

Then, after a full round of message passing we have for each node $V$ and separator $S$ that

$$t_V = P(V, e)$$
$$t_S = P(S, e)$$
$$P(e) = \sum_V t_V.$$

**Side effect of Hugin Propagation**

Let $Rt$ be the root for HUGIN propagation, and let $W$ and $V$ be neighbours with separator $S$. Assume that $W$ is closer to $Rt$ than $V$. Then $S$ divides the entered evidence in $ev$ and $ew$ (see Fig. 4.27).

A call of CollectEvidence($Rt$) results in the table $P(S, ev)$ being communicated from $V$ to $S$. By marginalization you can calculate $P(ev)$.

**4.8 Bibliographical notes**

A version of probability updating in singly connected DAGs through message passing was presented by Kim & Pearl (1983). HUGIN propagation was proposed by Jensen et al. (1990). It is a modification of an algorithm proposed by Lauritzen & Spiegelhalter (1988). Similar methods were used for pedigree analysis by Cannings et al. (1978). Shafer & Shenoy (1990) propose a different message-passing method for junction trees. Other propagation methods for multiply connected DAGs exist, e.g. arch reversal proposed by Shachter (1986) or conditioning proposed by Pearl (1986a).

The concepts of triangulated graphs and junction trees have been discovered and rediscovered with various names. In Bertele & Broisch (1972) they are used for dynamic programming, and Beeri et al. (1983) use them for data base management. A good reference on triangulated graphs is Golumbic (1980). Tarjan & Yannakakis (1984) gives various triangulation methods and very efficient methods for testing whether a graph is triangulated. Jensen & Jensen (1994) contains a proof of **Theorem 4.8** together with a method for constructing optimal junction trees from triangulated graphs.

Forward sampling was proposed by Henrion (1988). Gibbs sampling was originally

Exercises

Exercise 4.1 For Table 4.6, calculate $t_v$ and $t_w$.

<table>
<thead>
<tr>
<th></th>
<th>$a_1$</th>
<th>$a_2$</th>
<th>$a_3$</th>
<th>$c_1$</th>
<th>$c_2$</th>
<th>$c_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$b_1$</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>6</td>
<td>12</td>
<td>24</td>
</tr>
<tr>
<td>$b_2$</td>
<td>3</td>
<td>2</td>
<td>1</td>
<td>18</td>
<td>6</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>$tv$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>$tw$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Exercise 4.2 For the universe $U$ over the ternary variables $(A, B, C)$ with the joint probability Table 4.7 we get the findings $f_1$: “$A$ is in state $a_1$”, and $f_2$: “$C$ is in state $c_1$ or $c_3$”.

<table>
<thead>
<tr>
<th></th>
<th>$a_1$</th>
<th>$a_2$</th>
<th>$a_3$</th>
<th>$b_1$</th>
<th>$b_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$b_1$</td>
<td>(2,4,3)</td>
<td>(1,4,8)</td>
<td>(5,0,7)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$b_2$</td>
<td>(5,10,4)</td>
<td>(2,3,3)</td>
<td>(1,5,4)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$b_3$</td>
<td>(1,5,6)</td>
<td>(3,3,3)</td>
<td>(0,6,2)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 4.7 Table for Exercise 4.2.

$P(A, B, C)$ multiplied by ten.

Calculate $P(B | f_1, f_2), P(C | f_1, f_2), P(f_1), P(f_2)$ and $P(f_1, f_2)$.

Exercise 4.3 Prove that the anarchistic message passing algorithm formulated in Section 4.3.2 never runs into a deadlock: as long as there are unused message channels at least one variable can send a message. (Hint: Induction on the number of nodes and the fact that any sending sequence must start with a leaf sending.)

Exercise 4.4 Let $B$ be independent of $C$ given $A$, and let $P(A, B)$ and $P(A, C)$ be consistent. What is $P(A, B, C)$?

Exercise 4.5 Prove that a call of CollectEvidence in any node followed by a call of DistributeEvidence in the same node will result in a full propagation (all messages passed and passed when permitted).

Exercise 4.6 Construct the moral graph and a junction tree for the singly connected DAG below.

Exercise 4.7 Show that a consistent junction tree is globally consistent.
Table 4.8 Table for Exercise 4.10(i).

<table>
<thead>
<tr>
<th>A = y</th>
<th>A = n</th>
<th>A = y</th>
<th>A = n</th>
</tr>
</thead>
<tbody>
<tr>
<td>B = y</td>
<td>0.2</td>
<td>0.5</td>
<td>C = y</td>
</tr>
<tr>
<td>B = n</td>
<td>0.8</td>
<td>0.5</td>
<td>C = n</td>
</tr>
</tbody>
</table>

Exercise 4.11 (Conditioning.) Propagation methods for singly connected DAGs have existed for a long time. A propagation method for multiply connected DAGs consists of reducing a DAG to a set of singly connected DAGs.

(i) Consider the DAG (a) below with $P(A)$, $P(B \mid A)$, $P(C \mid A)$ and $P(D \mid B, C)$ given. Assume that $A = a$. Show that the DAG is reduced to the DAG (b) with $P(B \mid a)$, $P(C \mid a)$, and $P(D \mid B, C)$ given.

(ii) Show that $P(D, a) = P(D \mid b, c)P(B \mid a)P(C \mid a)$.

(iii) Assume that for all states $a$ of $A$ we have a reduced DAG as in (i). Let evidence $e$ be entered and propagated in all the reduced DAGs, yielding $P(B, e \mid a)$, $P(C, e \mid a)$, $P(D, e \mid a)$ for all $a$. Calculate $P(B, e)$ and $P(A, e)$.

The procedure above is called conditioning on $A$.

(iv) Reduce the DAG by conditioning on $B$. Show that the tables are $P(A \mid b)$, $P(C \mid A)$ and $P(D \mid C, b)$.

(v) Show that conditioning on $D$ does not result in a singly connected DAG.

Conditioning over several variables can be performed stepwise.

(vi) Determine a minimal set of conditioning variables for the DAG shown below to reduce it to singly connected DAGs.

(vii) The numbers attached to the variables indicate the number of states. Determine a conditioning resulting in a minimal number of singly connected DAGs.

Table 4.9 Table for Exercise 4.10(ii).

<table>
<thead>
<tr>
<th>B = y</th>
<th>B = n</th>
</tr>
</thead>
<tbody>
<tr>
<td>C = y</td>
<td>(0.1, 0.4)</td>
</tr>
<tr>
<td>C = n</td>
<td>(0.3, 0.5)</td>
</tr>
</tbody>
</table>

Exercise 4.12 Calculate the marginals from the sample in Table 4.5, and compare the result with the exact marginals.

Exercise 4.13 The binary variables $A$ and $B$ are parents of the binary variable $C$. $P(A) = P(B) = (0.5, 0.5)$, and the conditional probability table is an exclusive or table: $C = y$ if and only if exactly one of $A$ and $B$ is in the state $y$.

Show that Gibbs sampling on this structure will give either $P(C = y) = 1$ or $P(C = n) = 1$.

Exercise 4.14 Given a Bayesian network over $U$ with evidence $e$ entered, show that it is NP-hard to find a configuration $U^*$ such that $P(U^*, e) > 0$.

(Hint. Look at Exercise 3.16.)
Appendix A

Construction of junction trees (proofs)

This appendix contains proofs of the crucial theorems in Section 4.5.

Definitions. Let $G$ be an undirected graph with node set $N$.

If in $G$ there is a link between $A$ and $B$ they are said to be neighbours. Sometimes the word adjacent is used. A path in $G$ is a sequence $A_1, \ldots, A_n$ of distinct nodes where $A_i$ and $A_{i+1}$ are neighbours. A cycle is a path where $A_1 = A_n$ and all other nodes are distinct. A chord in a cycle $A_1, \ldots, A_n$ is a link between two nodes $A_i$ and $A_j$, where $i$ and $j$ are not consecutive numbers.

$G$ is triangulated if any cycle of length $> 3$ has a chord.

A subset $S$ of $N$ is complete if each pair of nodes in $S$ are neighbours. A node is simplicial if its neighbour set is complete.

A node $A$ is eliminated from $G$ by adding links to $G$ such that $A$ becomes simplicial and then removing it together with its links. The result is denoted $G_A$. $G$ is said to be eliminatable if all nodes can be successively eliminated without adding extra links.

Theorem A.1 Any eliminatable graph is triangulated.

Proof. Induction on the number of nodes.

Clearly, a graph consisting of one node is both triangulated and eliminatable.

Assume the theorem to hold for all graphs consisting of $< n$ nodes, and let $G$ be an eliminatable graph with $n$ nodes. Since $G$ is eliminatable it must have at least one simplicial node $A$. By the induction hypothesis, $G_A$ is triangulated. Since $A$ is simplicial, any cycle of length $> 3$ containing $A$ must have a chord (see Fig. A.1). Hence $G$ is triangulated.

To prove that any triangulated graph is eliminatable we need a lemma.

Lemma A.1 (Decomposition lemma.) Let $G$ be a noncomplete triangulated graph with at least three nodes, and with node set $N$. Then there is a complete subset $S$ of $N$ such that $G \setminus S$ is disconnected.
Appendix A

Construction of junction trees (proofs)

This appendix contains proofs of the crucial theorems in Section 4.5.

Definitions. Let $G$ be an undirected graph with node set $N$.

If in $G$ there is a link between $A$ and $B$ they are said to be neighbours. Sometimes the word adjacent is used. A path in $G$ is a sequence $A_1, \ldots, A_n$ of distinct nodes where $A_i$ and $A_{i+1}$ are neighbours. A cycle is a path where $A_1 = A_n$ and all other nodes are distinct. A chord in a cycle $A_1, \ldots, A_n$ is a link between two nodes $A_i$ and $A_j$, where $i$ and $j$ are not consecutive numbers.

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Assume the theorem to hold for all graphs consisting of $<n$ nodes, and let $G$ be an eliminatable graph with $n$ nodes. Since $G$ is eliminatable it must have at least one simplicial node $A$. By the induction hypothesis, $G_A$ is triangulated. Since $A$ is simplicial, any cycle of length $>3$ containing $A$ must have a chord (see Fig. A.1). Hence $G$ is triangulated.

To prove that any triangulated graph is eliminatable we need a lemma.

Lemma A.1 (Decomposition lemma.) Let $G$ be a noncomplete triangulated graph with at least three nodes, and with node set $N$. Then there is a complete subset $S$ of $N$ such that $G \setminus S$ is disconnected.
Now, assume the claim to be true for any graph with \( < n \) nodes and let \( G \) have \( n \) nodes. If \( G \) is complete then any two nodes are simplicial. So, suppose that \( G \) is not complete. Then the decomposition lemma yields a complete subset \( S \) separating \( G \) into at least two components.

Let \( G_A \) and \( G_B \) be defined as in the proof of the decomposition lemma. They are both triangulated (any chordless cycle in \( G_A \) is also a chordless cycle in \( G \)). If \( G_A \) is not complete we can choose two nonadjacent simplicial nodes \( A_1 \) and \( A_2 \). Since they cannot both be members of \( S \), we choose \( A_1 \) outside \( S \) to ensure that \( A_1 \) is simplicial in \( G \). If \( G_A \) is complete, choose any node outside \( S \). The same can be done for \( G_B \). Since both nodes are chosen outside \( S \) they are nonadjacent.

Let us recall the following.

Definitions. A junction graph for an undirected graph \( G \) is an undirected, labelled graph. The nodes are the cliques of \( G \), and every pair of nodes with a nonempty intersection has a link labelled by that intersection.

A spanning tree of \( G \)'s junction graph is a junction tree for \( G \) if it has the property that for each pair \( U, V \) of nodes, all the nodes in the path between \( U \) and \( V \) contain the intersection \( U \cap V \). The labels on the links in a junction tree are called separators.

Theorem A.3 A connected undirected graph \( G \) is triangulated if it has a junction tree.

Proof. Induction on the number of nodes in the graph.

The theorem is trivially true for any graph with at most two nodes.

Assume the theorem to hold for all graphs with \( < n \) nodes. Let \( G \) be a connected graph with \( n \) nodes, and let \( T \) be a junction tree, for \( G \). Since \( T \) is a tree there is a clique \( C \) with only one neighbour \( C' \) in \( T \). Let \( A \in C \setminus C' \). Since \( T \) is a junction tree, \( A \) can only be a member of one clique, namely \( C \). Then all neighbours of \( A \) are members of \( C \) and hence pairwise linked. This means that \( A \) is simplicial in \( G \).

Now, remove \( A \) from \( C \) (if thereby the new clique becomes a subset of \( C' \) then remove \( C \) from \( T \)). The resulting tree \( T^* \) is a junction tree for \( G_A \).

By the induction hypothesis \( G_A \) is triangulated, and hence \( G \) is also triangulated.

Theorem A.4 Any connected triangulated graph has a junction tree.

Proof. Induction on the number of nodes of the graph.

Trivially, the theorem is true for any graph with at most two nodes.

Assume the theorem to hold for any graph with \( < n \) nodes, and let \( G \) be a connected triangulated graph with \( n \) nodes. By the proof of Theorem A.2 \( G \) has at least one simplicial node \( A \). Then \( A \) together with its neighbours form a clique \( C \) in \( G \). Also, \( G \) and \( G_A \) have the same cliques except for \( C \). \( G_A \) may instead have a clique consisting of \( C \setminus \{A\} \). Also, \( G_A \) must be triangulated because \( G \) is, and by the induction hypothesis - \( G_A \) has a junction tree \( T^* \).

Now, construct \( T \) from \( T^* \) in the following way.
Appendix B

Value of information (proofs)

This appendix contains proofs omitted in Section 5.5.

Proposition B.1 If \( V(P(H)) = \sum_{h \in H} a_h P(h) \) then \( EV(T) = V(P(H)) \).

Proof.

\[
EV(T) = \sum_{t \in T} P(t) V(P(H | t)) = \sum_{t \in T} P(t) \sum_{h \in H} a_h P(h | t) = \sum_{t \in T} \sum_{h \in H} a_h P(h, t)
\]

\[
= \sum_{h \in H} \sum_{t \in T} a_h P(h, t) = \sum_{h \in H} a_h P(h) = V(P(H)).
\]

Proposition B.2 (Jensen's inequality)

(Jensen is the most frequent Danish name. This Jensen lived from 1859 to 1925.)

Let \( V : \mathbb{R}^n \to \mathbb{R} \) be a convex function. Let \( x_1, \ldots, x_n \in \mathbb{R}^n \) and let \( a_1, \ldots, a_n \in [0, 1] \), such that \( \sum_{i=1}^n a_i = 1 \). Then

\[
V \left( \sum_{i=1}^n a_i x_i \right) \leq \sum_{i=1}^n a_i V(x_i).
\]

Proof. Induction on \( n \). The basis \( n = 2 \) is precisely the convexity property.

Assume that the inequality holds for \( n - 1 \).

Put \( t = \sum_{i=1}^{n-1} a_i \), then \( a_n = 1 - t \), and assume \( t \neq 0 \).

\[
V \left( \sum_{i=1}^n a_i x_i \right) = V \left( t \left( \sum_{i=1}^{n-1} \frac{a_i}{t} x_i \right) + (1-t)x_n \right)
\]

\[
\leq tV \left( \sum_{i=1}^{n-1} \frac{a_i}{t} x_i \right) + (1-t) V(x_n)
\]

\[
\leq t \sum_{i=1}^{n-1} a_i V(x_i) + a_n V(x_n)
\]

\[
= \sum_{i=1}^n a_i V(x_i).
\]