MODEL SELECTION AND ACCOUNTING FOR MODEL UNCERTAINTY IN GRAPHICAL MODELS USING OCCAM'S WINDOW

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TECHNICAL REPORT No. 213
July 1991

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July 22, 1991

Abstract

We consider the problem of model selection and accounting for model uncertainty in high-dimensional contingency tables, motivated by expert system applications. The approach most used currently is a stepwise strategy guided by tests based on approximate asymptotic P-values leading to the selection of a single model; inference is then conditional on the selected model. The sampling properties of such a strategy are complex, and the failure to take account of model uncertainty leads to underestimation of uncertainty about quantities of interest. In principle, a panacea is provided by the standard Bayesian formalism which averages the posterior distributions of the quantity of interest under each of the models, weighted by their posterior model probabilities. However, this has not been used in practice because computing the posterior model probabilities is hard and the number of models is very large (often greater than $10^{11}$).

We argue that the standard Bayesian formalism is unsatisfactory and we propose an alternative Bayesian approach that, we contend, takes full account of the true model uncertainty by averaging over a much smaller set of models. An efficient search algorithm is developed for finding these models. We consider two classes of models that arise in expert systems: the recursive causal models and the decomposable log-linear models. For each of these, we develop efficient ways of computing exact Bayes factors and hence posterior model probabilities. For the decomposable log-linear models, this is based on properties of chordal graphs and hyper Markov prior distributions and the

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resultant calculations can be carried out locally. The end product is an overall strategy for model selection and accounting for model uncertainty that searches efficiently through the very large classes of models involved.

Three examples are given. The first two concern data sets which have been analysed by several authors in the context of model selection. The third addresses a urological diagnostic problem.

KEYWORDS: Chordal graph; Contingency table; Decomposable log-linear model; Expert system; Hyper Markov distribution; Recursive causal model.

Contents

1 Introduction 2

2 Model Selection Strategy 5
  2.1 General Principles and Occam’s Razor 5
  2.2 Model Selection Strategy 7
  2.3 Occam’s Window 8

3 The Directed Case—Recursive Causal Model Selection 10
  3.1 Implementation 10
  3.2 Examples 12
    3.2.1 Coronary Heart Disease Risk Factors 12
    3.2.2 Women and Mathematics 15

4 The Undirected Case—Decomposable Model Selection 17
  4.1 Implementation 17
  4.2 Examples 19
    4.2.1 Coronary Heart Disease Risk Factors 19
    4.2.2 Women and Mathematics 20
    4.2.3 Scrotal Swellings 21

5 Expert Systems 24
  5.1 Elicitation for Decomposable Model Selection 24
  5.2 Model Priors 27

6 Discussion 28

1 Introduction

Fruitful approaches to inference in high-dimensional contingency tables all involve choosing a broad class of models to be considered and then comparing them on the basis of how well
they predict the data. Typically, the model classes are huge and inference in the presence of the many competing models is not easy.

Here we consider two classes of models: the recursive causal models of Knijfer et al. (1984) and the decomposable log-linear models introduced by Goodman (1970) and Haberman (1974). This work is motivated by applications in expert systems which use a belief network to represent knowledge and perform inference. Lauritzen and Spiegelhalter (1988) have described a method for constructing such belief networks. A recursive causal model is elicited from an “expert” in the form of an acyclic directed graph with nodes representing random variables and directed links representing conditional dependence assumptions: we will refer to this as the “qualitative layer” of the model. Next, the joint distribution of the random variables being modeled is elicited; this is the “quantitative layer”. Finally, through a series of graphical and numerical operations, the recursive causal model is converted to a decomposable, and hence graphical, log-linear model. We assume throughout that all links in the graph corresponding to the recursive causal model are directed. It is important that the required probabilities be elicited in the directed recursive framework, because elicitation of joint distributions in the undirected decomposable framework is usually not feasible.

Potentially the most important advantage of constructing expert systems in this fashion is the system’s ability to modify itself as data becomes available. In a series of recent papers, Spiegelhalter and Lauritzen (1990a, 1990b), Dawid and Lauritzen (1989) and Spiegelhalter and Cowell (1991) have addressed the issue of updating the quantitative layer of the model. Building on this work, we address the issue of updating the qualitative layer—how can the graphical structure itself be updated as data becomes available? If we succeed in our objective we will have truly constructed an expert system which can learn.

Currently, the most used approach to model selection in contingency tables is a stepwise one, adapted from stepwise regression by Goodman (1971); see also Bishop, Fienberg and Holland (1975, Section 4.5 and Chapter 9). This consists of sequentially adding and deleting terms on the basis of approximate asymptotic likelihood ratio tests, leading to the selection of a single model. Inference about the quantities of interest is then made conditionally on the selected model.

There are several difficulties with this approach. The sampling properties of the overall strategy are complex because it involves multiple tests and, at least implicitly, the comparison of non-nested models (Fenech and Westfall, 1988). The use of $P$-values themselves is controversial, even when there are only two models to be compared, because of the so-called “conflict between $P$-values and evidence” discussed by Berger and Sellke (1987) and Berger
and Delampady (1987). One aspect of this is that tests based on $P$-values tend to reject even apparently satisfactory models when the sample size is large; a dramatic example of this was discussed by Raftery (1986b). On the other hand, when the sample size is small and the table sparse, the asymptotic approximations on which the $P$-values are based tend to break down.

Perhaps most fundamentally, conditioning on a single selected model ignores model uncertainty and so leads to underestimation of the uncertainty about the quantities of interest. This underestimation can be large, as was shown by Regal and Hook (1991) in the contingency table context and by Miller (1984) in the regression context. One bad consequence is that it can lead to decisions that are too risky (Hodges, 1987).

In principle, the standard Bayesian formalism provides a panacea for all these difficulties. If $\Delta$ is the quantity of interest, such as a structural characteristic of the system being studied, a future observation, or the utility of a course of action, then its posterior distribution given data $D$ is

$$
\text{pr}(\Delta \mid D) = \sum_{k=1}^{K} \text{pr}(\Delta \mid M_k, D)\text{pr}(M_k \mid D) .
$$

(1)

This is an average of the posterior distributions under each of the models, weighted by their posterior model probabilities. In equation (1), $M_1, \ldots, M_K$ are the models considered and

$$
\text{pr}(M_k \mid D) = \frac{\text{pr}(D \mid M_k)\text{pr}(M_k)}{\sum_{i=1}^{K} \text{pr}(D \mid M_i)\text{pr}(M_i)} ,
$$

(2)

where

$$
\text{pr}(D \mid M_k) = \int \text{pr}(D \mid \theta_k, M_k)\text{pr}(\theta_k \mid M_k)d\theta_k
$$

(3)

is the marginal likelihood of model $M_k$, $\theta_k$ is the (vector) parameter of $M_k$, $\text{pr}(\theta_k \mid M_k)$ is the prior distribution of $\theta_k$, $\text{pr}(D \mid \theta_k, M_k)$ is the likelihood, and $\text{pr}(M_k)$ is the prior probability of $M_k$.

However, this approach has not been adopted in practice. This appears to be because (a) the posterior model probabilities $\text{pr}(M_k \mid D)$ are hard to compute since they involve the very high-dimensional integrals in equation (3), and (b) the number of models in the sum in equation (1) is huge. For example, with just 10 variables (small by expert system standards) there are approximately $4.2 \times 10^{18}$ recursive causal models and $1.9 \times 10^{11}$ decomposable models.

One might hope that most of the posterior probability would be accounted for by a small number of models so that the sum in equation (1) would be well approximated by a small number of terms. Unfortunately, this does not typically appear to be the case because,
although a small number of models do have much higher posterior probabilities than all the others, the very many models with small posterior probabilities contribute substantially to the sum. For example, Moulton (1991) reported a regression example with $2^{12} = 4096$ models where about 800 models were needed to account for 90% of the posterior probability.

We argue that the standard Bayesian formalism of equation (1) is flawed. Adopting standard methods of scientific investigation, we show that accounting for the true model uncertainty involves averaging over a much smaller set of models. We develop simple and efficient ways of computing the posterior model probabilities for the two model classes considered. Our approach is to take advantage of the graphical structure to calculate the required probabilities very quickly, while representing prior opinion in an easily elicitable form. We also describe an efficient algorithm for searching the very large model space.

Putting all this together gives us a simple and computationally efficient way of selecting the best models and accounting for model uncertainty in recursive causal models and decomposable log-linear models. To demonstrate the generality of our approach, our discussion will be in the context of conventional statistical model selection rather than expert systems, although we will return to some expert system specific issues in Section 5. In Section 2 we describe the principles underlying our approach to model selection. In Section 3 we apply those principles to the recursive causal models, while in Section 4 we consider the decomposable models.

2 Model Selection Strategy

2.1 General Principles and Occam's Razor

We argue that equation (1) does not accurately represent model uncertainty. Science is an iterative process in which competing models of reality are compared on the basis of how well they predict what is observed; models that predict much less well than their competitors are discarded. Most of the models in equation (1) have been discredited in the sense that they predict the data far less well than the best models and so they should be discarded; there is no uncertainty about this in any real sense. Hence they should not be included in equation (1).

In our approach, if a model predicts the data far less well than the best model in the class it will be discarded, so that initially we exclude from equation (1) those models not
belonging to the set

\[ A' = \left\{ M_k : \frac{\max_i \{ \text{pr}(M_i \mid D) \}}{\text{pr}(M_k \mid D)} \leq C \right\} \tag{4} \]

for some constant \( C \). The value of \( C \) used will depend on the context. In our examples we used \( C = 20 \), by analogy with the popular .05 cutoff for \( P \)-values; Jeffreys (1961, Appendix B) would suggest some number between 10 and 100, while Evett (1991) suggests a value of 1000 for forensic evidence in criminal cases.

Next we appeal to one of the most widely accepted norms of scientific investigation, namely Occam’s razor. Let \( E \) represent the evidence and \( \text{pr}(H \mid E) \) the probability of a specified hypothesis \( H \) given the evidence \( E \). Occam’s razor states that if:

\[ \text{pr}(H_1 \mid E) = \text{pr}(H_2 \mid E) = \ldots = \text{pr}(H_k \mid E) \]

for hypotheses \( H_1, \ldots, H_k \), then the simplest among \( H_1, \ldots, H_k \) is to be preferred (Kotz and Johnson, 1985).

Thus we also exclude from equation (1) models belonging to the set

\[ B = \left\{ M_k : \exists M_i \in A, M_i \subset M_k, \frac{\text{pr}(M_i \mid D)}{\text{pr}(M_k \mid D)} > 1 \right\} \tag{5} \]

and equation (1) is replaced by

\[ \text{pr}(\Delta \mid D) = \frac{\sum_{M_k \in A} \text{pr}(\Delta \mid M_k, D)\text{pr}(D \mid M_k)\text{pr}(M_k)}{\sum_{M_k \in A} \text{pr}(D \mid M_k)\text{pr}(M_k)} \tag{6} \]

where

\[ A = A' \setminus B. \tag{7} \]

This greatly reduces the number of models in the sum in equation (1) and hence simplifies the model uncertainty problem a great deal. Note that our argument is not an approximation adopted for computational convenience, but rather an exact solution based on the way science works. Note also that our approach in equation (6) will not necessarily give an answer close to that given by equation (1) because, due to the very large number of models in the class, the models discarded may have a large total posterior probability \( \sum_{M_k \in A} \text{pr}(M_k \mid D) \), even though each individual model discarded has a very small posterior probability.

The problem thus reduces to finding the set \( A \), and we now outline a computational strategy for doing this.
2.2 Model Selection Strategy

Our approach to model selection is a variant of the greedy-search algorithm. The essentials of the approach are the same for the recursive causal models and the decomposable models and could readily be applied to more general graphical models. Posterior model probabilities are used as a metric to guide the search. The strategy proceeds out into model space away from the opening set of models, comparing models via ratios of posterior model probabilities in a series of nested comparisons. The extent of the search is easily controlled and will depend on the resources available for specific applications. In what follows, $M_0$ will denote the smaller of the two models being compared and $M_1$ will denote the larger. In fact, $M_0$ and $M_1$ will differ by just one link throughout. We now describe the elements of our approach.

Edwards and Havránek (1985) proposed a model search procedure which is based on the following rules:

- If a model is rejected, then all its submodels are rejected.

- If a model is accepted, then all models that include it are considered accepted. (We use the term accepted in place of the more correct non-rejected.)

These rules were first suggested by Gabriel (1969). He coined the term coherence for testing procedures satisfying these rules. However, Gabriel's arguments are predicated on the use of a monotone metric for model testing. A test statistic $Z(M)$ is said to be monotone if $M_i \subset M_j \Rightarrow Z(M_i) \geq Z(M_j)$. Likelihood ratio test statistics are monotone but posterior model probabilities clearly are not. We now argue that, in this context, the second rule is inappropriate.
Consider the (undirected) example in Figure 1. Suppose that we start with the saturated model $[ABC]$ of Figure 1(a), and that when we compare it with the model of conditional independence $[AC][BC]$ of Figure 1(b), we reject the smaller model decisively. Then we are precisely rejecting the conditional independence of $A$ and $B$ given $C$. This conditional independence also holds in all the submodels of $[AC][BC]$ and so we reject all of those as well, including the model $[A][BC]$ of Figure 1(c). Thus, if we reject a model, we reject all its submodels, which is the first of the two rules of Edwards and Havránek (1985) above.

Now, working in the opposite direction, suppose that we start with the model $[A][BC]$ of Figure 1(c). Comparing it with $[AC][BC]$ let us suppose that we decisively reject the smaller model. We are precisely rejecting the marginal independence of $A$ in favour of the conditional independence of $A$ and $B$ given $C$. However, we have learned nothing about the model $[ABC]$. Indeed, its probability could be even lower than that of $[A][BC]$! It follows that the second rule of Edwards and Havránek is inappropriate in the present context.

### 2.3 Occam's Window

A crucial aspect of the strategy concerns the interpretation of the ratio of posterior model probabilities when comparing two models. Again we appeal to Occam's razor which, translated into the language of model fitting, we implement as follows:

- If the log posterior odds is positive, i.e., the data provides evidence for the smaller model, then we reject $M_1$ and consider $M_0$. We could generalize this by requiring the log posterior odds to be greater than some positive constant $O_0$ before rejecting $M_1$.

- If the log posterior odds is small and negative, providing evidence against the smaller model which is not "very strong" (Jeffreys, 1961), then we consider both models.

- If the log posterior odds is large and negative, i.e., smaller than $O_L = -\log(C)$ where $C$ is defined by equation (7), we reject $M_0$ and consider $M_1$.

Thus there are three possible actions following each comparison—see Figure 2.

Now that the various elements of the strategy are in place, we outline the search technique. The search can proceed in two directions: "Up" from each starting model by adding links, or "Down" from each starting model by dropping links. When starting from a non-saturated, non-empty model, we first execute the "Down" algorithm. Then we execute the "Up" algorithm, using the models from the "Down" algorithm as a starting point. Experience to date suggests that the ordering of these operations has little impact on the final set of
Figure 2: Occam’s Window: Interpreting the log posterior odds

models. Let $\mathcal{A}$ and $\mathcal{C}$ be subsets of model space $\mathcal{M}$, where $\mathcal{A}$ denotes the set of “acceptable” models and $\mathcal{C}$ denotes the models under consideration. For both algorithms, we begin with $\mathcal{A} = \emptyset$ and $\mathcal{C} =$ set of starting models.

**BGMS-Down Algorithm**

1. Select a model $M$ from $\mathcal{C}$
2. $\mathcal{C} \leftarrow \mathcal{C} - M$ and $\mathcal{A} \leftarrow \mathcal{A} + M$
3. Select a submodel $M_0$ of $M$ by removing a link from $M$
4. Compute $B = \log \frac{\text{pr}(M_0|D)}{\text{pr}(M|D)}$
5. If $B > O_R$ then $A \leftarrow A - M$ and if $M_0 \notin \mathcal{C}, \mathcal{C} \leftarrow \mathcal{C} + M_0$
6. If $O_L \leq B \leq O_R$ then if $M_0 \notin \mathcal{C}, \mathcal{C} \leftarrow \mathcal{C} + M_0$
7. If there are more submodels of $M$, go to 3
8. If $\mathcal{C} \neq \emptyset$, go to 1

**BGMS-Up Algorithm**

1. Select a model $M$ from $\mathcal{C}$
2. $\mathcal{C} \leftarrow \mathcal{C} - M$ and $\mathcal{A} \leftarrow \mathcal{A} + M$
3. Select a supermodel $M_1$ of $M$ by adding a link to $M$
4. Compute \( B = \log \frac{\text{pr}(\mathcal{M}_1 | D)}{\text{pr}(\mathcal{M}_0 | D)} \)

5. If \( B < O_L \) then \( \mathcal{A} \leftarrow \mathcal{A} - \mathcal{M} \) and if \( M_1 \notin \mathcal{C} \), \( \mathcal{C} \leftarrow \mathcal{C} + M_1 \)

6. If \( O_L \leq B \leq O_R \) then if \( M_1 \notin \mathcal{C} \), \( \mathcal{C} \leftarrow \mathcal{C} + M_1 \)

7. If there are more supermodels of \( \mathcal{M} \), go to 3

8. If \( \mathcal{C} \neq \emptyset \), go to 1

Upon termination, \( \mathcal{A} \) contains the set of potentially acceptable models. Finally, we remove all the models which satisfy equation (5), where \( I \) is replaced by \( \exp(O_R) \), and those models \( M_k \) for which

\[
\max_i \{ \frac{\text{pr}(M_i | D)}{\text{pr}(M_k | D)} \} > C. \tag{8}
\]

\( \mathcal{A} \) now contains the acceptable models.

3 The Directed Case—Recursive Causal Model Selection

3.1 Implementation

Implementation for the recursive causal models proceeds in a straightforward fashion. Consider a recursive causal model for a set of random variables \( X_v, v \in V \). The model is represented by a directed graph where each variable in \( V \) is represented by a node in the graph. For each variable \( v \in V \) we define \( \text{pa}(v) \) to be the set of parent nodes of \( v \), i.e. nodes \( w \) for which there exists a directed link from \( w \) to \( v \). The assumptions of the model imply that the joint distribution of \( X_v, v \in V \), which we denote \( \text{pr}(V) \), is given by

\[
\text{pr}(V) = \prod_{v \in V} \text{pr}(v | \text{pa}(v)).
\]

In early implementations, \( \text{pr}(v | \text{pa}(v)) \) was assumed to be fully specified for all \( v \) by the expert/data analyst. Spiegelhalter and Lauritzen (1990a) introduced a parametrisation for \( \text{pr}(v | \text{pa}(v)) \) whereby the relationship between a node \( v \) and its parents \( \text{pa}(v) \) is fully specified by \( \theta_v \subset \Theta_v \). This leads to a conditional distribution for \( V \):

\[
\text{pr}(V | \theta) = \prod_{v \in V} \text{pr}(v | \text{pa}(v), \theta_v).
\]

where \( \theta \) is a general parameter with components \( \theta_v \).
Spiegelhalter and Lauritzen (1990a) make two key assumptions which greatly simplify subsequent analysis. The first assumption is that of global independence whereby the parameters $\theta_v$ are assumed mutually independent a priori. This assumption alone allows us to calculate the likelihood for a single case:

$$p_r(v) = \int p_r(v, \theta)d\theta = \int \prod_i p_r(v|pa(v), \theta_v)p_r(\theta_v)d\theta_v = \prod_i p_r(v|pa(v))$$

where

$$p_r(v|pa(v)) = \int p_r(v|pa(v), \theta_v)p_r(\theta_v)d\theta_v.$$

The second assumption is that of local independence whereby the parameter $\theta_v$ breaks into components corresponding to the elements of the state space of $pa(v)$. These components are assumed to be mutually independent a priori.

Now consider a conditional probability distribution $p_r(v|pa(v)^+, \theta_v^v) = \theta_v^v$ for a specific state $pa(v)^+$ of $pa(v)$. We assume that $\theta_v^v$ has a Dirichlet distribution $D[\lambda_1^v, ..., \lambda_k^v]$, where $k$ is the number of states of $v$. Then we can show that

$$p_r(v|pa(v)^+) = \lambda^v/\sum_i \lambda_i^v.$$

If we observe $v$ to be in state $x_v$ and the parent state to be $pa(v)^+$, we have

$$\theta_v^v|v \sim D[\lambda_1^v, ..., \lambda_j^v + 1, ..., \lambda_k^v].$$

This provides a straightforward method for sequentially calculating the ratios of posterior model probabilities required for the model selection strategy. The elicitation of the required Dirichlet priors is feasible provided the cardinality of $pa(v)$ is not too large. Computer-based methods for eliciting Dirichlet prior distributions have been described by Chaloner and Duncan (1987). If $pa(v)$ is not observed the updating becomes more complex—see Spiegelhalter and Lauritzen (1990a,1990b) for details.

A considerable computational saving is obtained by noting that the sequential updating of the distribution of $\theta_v$ depends on the states of $v$ and $pa(v)$ only. Therefore the likelihood for all qualitative layers (graphs) having the same set $pa(v)$ of parent nodes of $v$ will have identical contributions from $v$. For example, consider the two recursive causal models of Figure 3. When calculating the likelihood for the model of Figure 3(a), we store the likelihood of each node/parent combination separately. Now when subsequently calculating the likelihood for the model of Figure 3(b), only the likelihood for node $B$ requires recalculation as the sets of parent nodes of $A$ and $C$ have not changed.
To implement the model selection strategy described in Section 2 for the recursive causal models an ordering of the nodes must be pre-specified by the expert/data analyst. If \( v_i \) precedes \( v_j \) in the ordering, then a directed link from \( v_j \) to \( v_i \) is prohibited. In certain applications it may be possible to search over all possible orderings but this will typically not be the case. Pearl's IC-algorithm (Pearl and Verma, 1991) induces directed "causal" structures from data. An ordering of the nodes is not required, but for each pair of nodes \( v_i \) and \( v_j \), the algorithm does involve searching amongst all subsets of \( V - \{v_i, v_j\} \) for cutsets between \( v_i \) and \( v_j \) (sets which when conditioned on, render \( v_i \) and \( v_j \) independent.) Cooper and Herskovits (1991) provide a review of other approaches.

### 3.2 Examples

#### 3.2.1 Coronary Heart Disease Risk Factors

Firstly we consider a data set which has been previously analysed by Edwards and Havrânek (1985). The data concerns 1,841 men cross-classified according to six coronary heart disease risk factors. The data is reproduced in Table 1. The risk factors are as follows: 
- \( A \), smoking;
- \( B \), strenuous mental work;
- \( C \), strenuous physical work;
- \( D \), systolic blood pressure;
- \( E \), ratio of \( \beta \) and \( \alpha \) proteins;
- \( F \), family anamnesis of coronary heart disease.

Their likelihood ratio-based model selection strategy selected two graphical log-linear models: 
\[
[AC][ADE][BC][BE][F]
\]
which is not decomposable and therefore is not equivalent to any recursive causal model, and 
\[
[ACE][ADE][BC][F]
\]
which is decomposable. A striking feature of both models is the independence of \( F \), family anamnesis. The models are shown in Figure 4.
Table 1: Risk factors for Coronary Heart Disease

<table>
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<tr>
<th>F</th>
<th>E</th>
<th>D</th>
<th>A</th>
<th>B No</th>
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</table>

Figure 4: Models Selected by Edwards and Havránek
To implement the Bayesian graphical model selection procedure, we started from the saturated model and used the “Down” algorithm only (starting from the empty model and using the “Up” algorithm produced the same set of models). All qualitative structures were assumed equally likely a priori. We adopted a standard Jeffreys prior density throughout. A natural partial ordering of the variables suggests itself: $F, (B, C), A, (E, D)$. $B, F$ or $C$ could not be “influenced” by the other factors and must be exogenous, although the ordering of $B$ and $C$ is unclear. Similarly, $D$ or $E$ could hardly influence $A$, although the ordering of $E$ and $D$ is unclear. The four corresponding complete orderings produced strong evidence for the precedence of $E$ over $D$, and indifference as to the ordering of $B$ and $C$. Several further orderings were tried, but this “natural” ordering resulted in the models with highest posterior probabilities. The selected models are shown in Figure 5 and their posterior probabilities in Table 2.

Two models, those shown in Figures 5(a) and 5(b), account for most of the posterior probability. They are rather similar in that both contain the $C - B, C - A, A - E, E - D$ and $A - D$ links. The main difference between them lies in the way they describe the effect of strenuous mental work ($B$) and strenuous physical work ($C$) on the ratio of $\beta$ and $\alpha$ proteins.
Table 2: Coronary Heart Disease: Posterior Model Probabilities for Recursive Causal Models

<table>
<thead>
<tr>
<th>Figure</th>
<th>Log marginal likelihood</th>
<th>Posterior probability %</th>
</tr>
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<tr>
<td>5(a)</td>
<td>-6723.0</td>
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<td>5(b)</td>
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<td>5(c)</td>
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<td>5(d)</td>
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</table>

(E). Model 5(a) says that C affects E both directly and indirectly via A, whereas model 5(b) says that the effect of C on E is solely indirect, being mediated by B and A. There is also some uncertainty about the presence of a link from smoking (A) to systolic blood pressure (D). There is decisive evidence in favour of the marginal independence of family anamnesis of coronary heart disease (F).

The four models selected correspond very closely to the models of Figure 4 above. We note that the A – D link (smoking and systolic blood pressure) is present in both of the models of Figure 4 and also in models (a) and (b) of Figure 5, but it is absent from models (c) and (d) of Figure 5. In fact, the exact test for zero partial association of A and D reported by Edwards and Havránek (1985) had a significance level of 0.04 which was the largest of any of the links accepted at the 5% level.

3.2.2 Women and Mathematics

Our second example concerns a survey which was reported in Fowlkes et al. (1988) concerning the attitudes of New Jersey high-school students towards mathematics. The data has been further analysed by Upton (1991). A total of 1190 students in eight schools took part in the survey. Data on six dichotomous variables was collected:

A. Lecture Attendance; attended or did not attend;

B. Sex; female or male;

C. School Type; suburban or urban;

D. “I’ll need mathematics in my future work”; agree or disagree;

E. Subject Preference; maths/science or liberal arts;
Figure 6: Women and Mathematics: Recursive Causal Model Selected

F. Future Plans; college or job;

Upton (1991) reports that a model selection procedure based on the AIC criterion (Akaike, 1973) selects \([ABCE][CDF][BCD][DEF]\) while a procedure based on the BIC criterion (Raftery, 1986a) selects the much simpler \([A][BE][CE][CF][BD][DE][DF]\). Clearly an important difference between these two models is the treatment of \(A\).

The Bayesian graphical model selection procedure started from the empty model and used the “Up” algorithm. It is clear that \(B\) (Sex) cannot be influenced by other variables and must be exogenous. Initially it was also assumed that \(C\) (School Type) was exogenous. An exhaustive search over all consequent orderings produced the single model shown in Figure 6.

The selected model is similar to the model selected by Upton’s BIC procedure. The model selected by AIC clearly over-fits the data (Upton, 1991). It is of interest to note the direction of the link from \(D\) to \(F\). Both Upton (1991) and Fowlkes et al. (1988) treat \(D\) as a response variable and Upton’s path diagram shows a directed link from \(F\) to \(D\). However, the data provides strong evidence that the direction of the influence is from \(D\) to \(F\), i.e. that students’ attitudes towards mathematics influence their future plans, rather than the other way around. The ability of the selected model to predict is unaffected by the direction of the \(E - D\) link.

Further analysis removed the restriction that \(C\) be exogenous. The data now provides some support for the presence of a link from \(E\) to \(C\) although its interpretation is somewhat unclear.
4 The Undirected Case—Decomposable Model Selection

4.1 Implementation

To implement the strategy for the decomposable models, we rely heavily on a recent fundamental paper by Dawid and Lauritzen (1989), hereafter DL. We consider three issues which are specific to model selection for the decomposable models. Firstly, how should we add and remove links whilst efficiently ensuring that all the models created are decomposable? Secondly, given any two decomposable models $M$ and $M^*$, is it possible to generate $M^*$ from $M$, adding or removing only one edge at a time but staying within the class of decomposable models? Finally, how do we calculate the required posterior model probabilities?

The first two issues are addressed by the following two lemmas:

**Lemma 1** Let $G = (V, E)$ be a chordal graph with vertices $V$ and edges $E$ and let $G' = (V, E')$ be a chordal subgraph of $G$ with exactly one edge, $e$, less. Then $e$ is contained in exactly one clique of $G$.

**Proof.** This follows from Lemma 3 of Frydenberg and Lauritzen (1989).

Therefore the model selection strategy must remove only links which are members of a single clique. When adding links, the strategy must not create any chordless four-cycles.

**Lemma 2** Let $G = (V, E)$ and $G' = (V, E')$ both be chordal graphs such that $E' \subseteq E$ and $G'$ has $k$ less edges than $G$. Then there is an increasing sequence $G' = G_0 \subseteq \ldots \subseteq G_k = G$ of chordal graphs that differ by exactly one edge.

**Proof.** This follows from Lemma 5 of Frydenberg and Lauritzen (1989).

Now we address the calculation of the posterior model probabilities. Following DL, we consider a decomposable model $M$ for a set of random variables $X_v, v \in V$. Let $I$ denote the set of possible configurations of $X$. Let the quantitative layer of $M$ be specified by $\theta$. Then the distribution of $\theta$ is determined by the clique marginal probability tables $\theta_C = (\theta_C)_{c \in C}$.
where \( C \) denotes the set of cliques of \( M \):

\[
\theta(i) = \frac{\prod_{c \in C} \theta_C(i \cap c)}{\prod_{s \in S} \theta_S(i \cap s)}, \quad i \in I.
\]

\( S \) denotes the system of separators in an arbitrary perfect ordering of \( C \).

For each clique \( C \in C \), let

\[
\lambda_C = (\lambda_C(i_C))_{i_C \in I_C}
\]

be a given table of arbitrary positive numbers and let \( D(\lambda_C) \) denote the Dirichlet distribution for \( \theta_C \) with density

\[
\pi(\theta_C|\lambda_C) \propto \prod_{i_C \in I_C} \theta_C(i_C)^{\lambda_C(i_C)-1},
\]

where \( \sum_{i_C} \theta_C(i_C) = 1 \) and \( \theta(i_C) > 0 \).

Now let us suppose that the collection of specifications \( D(\lambda_C), C \in C \) are constructed in such a way that for any two cliques \( C \) and \( D \) in \( C \) we have:

\[
\lambda_C(i_C \cap D) = \lambda_D(i_C \cap D).
\]

Then DL show that there exists a unique strong hyper Markov distribution for \( \theta \) over \( M \) that has density \( D(\lambda_C) \) for all \( C \in C \). DL call this the hyper Dirichlet distribution for \( \theta \).

A distribution for \( \theta \) is strong hyper Markov if and only if \( \theta_{A|B}, \theta_{B|A} \) and \( \theta_{A \cap B} \) are mutually independent whenever \( A \cap B \) is complete and separates \( A \) from \( B \). It follows that by letting \( \lambda_0 = \sum_{i \in I} \lambda_i \), the likelihood for a single case is given by:

\[
pr(v) = \frac{\prod_{c \in C} \lambda_C}{\lambda_0^{|S| \prod_{s \in S} \lambda_S}}.
\]

From Lemma 3 we have that updating can be carried out one clique at a time:

**Lemma 3** If the prior distribution \( \mathcal{L}(\theta) \) is strong hyper Markov, the posterior distribution of \( \theta \) is the unique hyper Markov distribution \( \mathcal{L}^* \) specified by the clique-marginal distributions \( \{\mathcal{L}_C : C \in C\} \), where \( \mathcal{L}_C^* \) is the posterior distribution of \( \theta_C \) based on its prior distribution \( \mathcal{L}_C \) and the clique-specific data \( X_C = x_C \).

**Proof.** This is Corollary 9 of DL.

The posterior distribution for \( \theta_C \) given data \( n_C \) from the marginal table corresponding to clique \( C \) is \( D(\lambda_C + n_C) \).
Consider the Bayes factor
\[ B_{01} = \frac{\text{pr}(D|M_0)}{\text{pr}(D|M_1)} \]
where \( M_0 \) and \( M_1 \) are decomposable and \( M_0 \) is obtained from \( M_1 \) by deleting one edge \( e \) linking \( u \) with \( v \). From Lemma 1 we have that \( e \) is contained in a single clique, \( C \) say, of \( M_1 \). Let \( C_u = C - \{v\}, C_v = C - \{u\}, C_0 = C - \{u, v\} \). Then DL show that the Bayes factor is given by:
\[ B_{01} = \frac{pc_u(D_{C_u})pc_v(D_{C_v})}{pc_0(D_{C_0})pc(D_C)}. \]
Thus, the required decomposable model comparisons can be carried out very rapidly with calculations local to single cliques.

### 4.2 Examples

#### 4.2.1 Coronary Heart Disease Risk Factors

Firstly we consider again the coronary heart disease risk factor data of Edwards and Havránek (1985) which is shown in Table 1. We note that the model of Figure 4(a) which was selected by the Edwards and Havránek procedure is not decomposable and hence will not be selected by our procedure.

The selection procedure started from the saturated model and used the "Down" algorithm. All qualitative structures were assumed equally likely \textit{a priori}. A standard Jeffreys prior was adopted for \( \theta_C, C \in C \). Just two models were selected and they are shown in Figure 7. Starting from the empty model and using the "Up" algorithm resulted in the same two models. The corresponding posterior probabilities are shown in Table 3.
### Table 3: Coronary Heart Disease: Posterior Model Probabilities for Decomposable Models

<table>
<thead>
<tr>
<th>Figure</th>
<th>Model</th>
<th>Log marginal likelihood</th>
<th>Posterior probability %</th>
</tr>
</thead>
<tbody>
<tr>
<td>7(a)</td>
<td>$[BC][ACE][ADE][F]$</td>
<td>-6719.5</td>
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<tr>
<td>7(b)</td>
<td>$[ABC][ABE][ADE][F]$</td>
<td>-6721.9</td>
<td>8</td>
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</table>

### Table 4: Women and Mathematics: Posterior Model Probabilities for Decomposable Models

<table>
<thead>
<tr>
<th>Figure</th>
<th>Model</th>
<th>Log marginal likelihood</th>
<th>Posterior probability %</th>
</tr>
</thead>
<tbody>
<tr>
<td>8(a)</td>
<td>$[A][BDE][CDF]$</td>
<td>-4492.4</td>
<td>75</td>
</tr>
<tr>
<td>8(b)</td>
<td>$[A][BDE][DF][CF]$</td>
<td>-4493.5</td>
<td>25</td>
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</table>

The model of Figure 7(a) was also selected by the directed model selection procedure and by Edwards and Havránek (1985). The model of Figure 7(b) is essentially a decomposable version of the directed model of Figure 5(b) and Edwards and Havránek's model of Figure 4(a). It is interesting to note that a model identical to Figure 7(a) except for the $A \rightarrow D$ link falls just outside Occam's window in the undirected selection.

Over all, the model selection exercise indicates that there is very strong evidence for the $B \rightarrow C$, $A \rightarrow C$, $A \rightarrow E$ and $D \rightarrow E$ links, with evidence for the $A \rightarrow D$ link that is strong but somewhat less so. There is also some evidence for the $C \rightarrow E$ and $B \rightarrow E$ links, but it seems that one of these alone is enough to describe the data, and it is not fully clear which one is better. Again, as in the directed case, there is decisive evidence for the marginal independence of $F$.

#### 4.2.2 Women and Mathematics

We consider again the survey data previously analysed by Fowlkes et al. (1988) and Upton (1991). We note the models selected in Upton (1991) are not graphical and hence will not be selected by our procedure. The procedure adopted was identical to that adopted for the example of Section 4.2.1. The two models selected are shown in Figure 8 and the corresponding posterior probabilities are shown in Table 4.

As in the directed case, the selected models are close to the models selected by the BIC model selection procedure carried out by Upton (1991). However there is uncertainty about
the $C-D$ link (School Type and "I’ll need mathematics in my future work") which is not apparent in Upton’s analysis. The data strongly supports the marginal independence of $A$.

4.2.3 Scrotal Swellings

Our final example concerns the diagnosis of scrotal swellings. Data on 299 patients was gathered at the Meath Hospital, Dublin, Ireland under the supervision of Mr. M.R. Butler. We consider a cross-classification of the patients according to one disease class, Hernia ($H$), and 7 binary indicators as follows: $A$, possible to get above the swelling; $B$, swelling transilluminates; $C$, swelling separate from testes; $D$, positive valsalva/stand test; $E$, tender; $F$, pain; $G$, evidence of other urinary tract infections. The data is reproduced in Table 5. There are 28 possible links to be considered by the selection procedure in this example. In the absence of prior expert opinion, computation times can be prohibitive. Clearly, if the starting point for the selection procedure were close to the models for which the data provides evidence, this problem could be overcome. With this objective we adopted the following heuristic procedure: firstly, Bayes factors for each of the 28 links are calculated by comparing the saturated model with the 28 sub-models generated by removing single links. Links for which the data provides evidence in this manner are then used as a starting point for the selection procedure (if the model thus constructed is not decomposable, some of the links may be removed or additional ones may be added.) The starting model is shown in Figure 9.

Now the “Up” algorithm is executed, followed by the “Down” algorithm (or vice versa). Note that if the starting links are badly chosen, the complete procedure has the opportunity to remove them, although in this example, the final model contains all the links from the
Table 5: Scrotal Swelling data

<table>
<thead>
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<th>Hernia</th>
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</tbody>
</table>

Figure 9: Starting Model for Scrotal Swelling Example
starting model. Two models were selected by this procedure and they are shown in Figure 10. The corresponding posterior probabilities are shown in Table 6.

The result of primary interest here is the importance of $A$ (possible to get above swelling) and $D$ (valsalva/stand test) with respect to Hernia diagnosis. Both indicants can be established through simple procedures at physical examination. The only real model uncertainty which is exhibited concerns the relationship between $C$ (swelling separate from testes) and $E$ (tender). Analysis of further cross-classifications extracted from this database also yield similarly sparse models.

Table 6: Scrotal Swellings: Posterior Model Probabilities for Decomposable Models

<table>
<thead>
<tr>
<th>Figure</th>
<th>Model</th>
<th>Log marginal likelihood</th>
<th>Posterior probability %</th>
</tr>
</thead>
<tbody>
<tr>
<td>8(a)</td>
<td>[AH][DH][BDE][CDE][EF][EG]</td>
<td>-806.2</td>
<td>75</td>
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<tr>
<td>8(b)</td>
<td>[AH][DH][BDE][CD][EF][EG]</td>
<td>-807.3</td>
<td>25</td>
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</table>
5 Expert Systems

As discussed in the introduction, model updating for expert system applications can be carried out in either the recursive causal model framework or the undirected decomposable framework. However, while the decomposable model selection strategy of Section 4 can be efficiently implemented and avoids the computational problems associated with orderings in the directed models, direct elicitation of the required Dirichlet distributions, $D(\lambda_C)$, will typically be intractable. In this section we outline a simple procedure for eliciting the required priors. This had previously been a barrier to the application of the approach of DL. We also address briefly the use of prior model probabilities to control the search in very large model spaces.

5.1 Elicitation for Decomposable Model Selection

Our objective is to use the initial recursive causal model as a framework in which to elicit the quantitative layer and ensure that the resultant prior distribution is hyper Dirichlet. We note that for both the recursive causal and the decomposable model selection strategies, the hyper Dirichlet framework provides a straightforward mechanism for evolving prior distributions as links are added and deleted.

Consider a simple recursive causal model with two binary variables as shown in Figure 11(a). This model is decomposable and the corresponding undirected representation is given in Figure 11(b).

Using the approach outlined in Section 3, the model of Figure 11(a) would require the elicitation of prior Dirichlet distributions for $pr(B|A)$, $pr(B|\overline{A})$ and $pr(A)$ with a total of 6 parameters. The model of Figure 11(b) however has only one clique $\{A, B\}$ and its prior hyper Dirichlet distribution has just 4 parameters. In general, the Dirichlet prior representation considered in Section 3 is over-parametrised relative to the hyper Dirichlet distribution.
Figure 12: Lauritzen and Spiegelhalter’s Dyspnoea Example

Our approach is to regard the parameters $\lambda_{(v|\text{pa}(v))}$ as "equivalent prior samples" which are elicited subject to constraints which ensure consistency. The implied equivalent prior sample for $\lambda$ satisfies the independence relationships of the recursive causal model and specifies the complete hyper Dirichlet distribution. Marginalising onto each clique $C \in \mathcal{C}$ provides the clique prior distributions.

Thus for the example of Figure 11, the equivalent prior sample is an unconstrained two-by-two table. Having elicited a Dirichlet prior for $\text{pr}(A)$, there are only two degrees of freedom remaining for the prior distributions of $\text{pr}(B|A)$ and $\text{pr}(B|\bar{A})$. Effectively the size of the equivalent prior sample and its breakdown between $A$ and $\bar{A}$ is fixed by the elicitation of the prior for $\text{pr}(A)$, and only one further parameter is now required for each of $\text{pr}(B|A)$ and $\text{pr}(B|\bar{A})$. We envisage software which would display all three prior distributions simultaneously, and maintain the required constraints interactively.

Since for all $v \in V$, $\{v \cup \text{pa}(v)\} \subseteq C$ for some $C \in \mathcal{C}$, implementation of the procedure in more general models is straightforward. We consider a fictitious example previously considered by Lauritzen and Spiegelhalter (1988) and provide a detailed description of the elicitation procedure. The (elicited) recursive causal model is shown in Figure 12.

Elicitation of the hyper Dirichlet prior and the marginal prior distributions for the cliques of the corresponding decomposable model proceeds as follows:

1. Elicit distributions for $\text{pr}(\alpha), \text{pr}(\tau|\alpha)$.

   Degrees of freedom : 4
2. Elicit distribution for $\text{pr}(\tau, \lambda)$. The marginal distribution of $\text{pr}(\tau)$ has already been specified in 1. Furthermore the model implies $\tau \perp \lambda$.

Degrees of freedom : 5

3. Elicit distribution for $\text{pr}(\xi|\epsilon)$. The marginal distribution of $\text{pr}(\epsilon)$ has already been specified in 2.

Degrees of Freedom : 2

4. Elicit distribution for $\text{pr}(\lambda|\sigma)$. The marginal distribution of $\text{pr}(\lambda)$ has already been specified in 2.

Degrees of Freedom : 2

5. Elicit distribution for $\text{pr}(\beta|\sigma)$. The marginal distribution of $\text{pr}(\sigma)$ has already been specified in 4.

Degrees of Freedom : 2

5a. Since the model implies $\lambda \perp \beta|\sigma$, we have that:

$$\text{pr}(\lambda, \beta, \sigma) = \frac{\text{pr}(\lambda, \sigma)\text{pr}(\beta, \sigma)}{\text{pr}(\sigma)}$$

and hence we derive the distribution of $\text{pr}(\lambda, \beta, \sigma)$ and the marginal distribution of $\text{pr}(\lambda, \beta)$.

5b. Since the model implies $\epsilon \perp \beta|\lambda$, we have that:

$$\text{pr}(\epsilon, \beta, \lambda) = \frac{\text{pr}(\epsilon, \lambda)\text{pr}(\beta, \lambda)}{\text{pr}(\lambda)}$$

and hence we derive the distribution of $\text{pr}(\epsilon, \beta, \lambda)$ and the marginal distribution of $\text{pr}(\epsilon, \beta)$.

6. Elicit distribution for $\text{pr}(\delta|\epsilon, \beta)$.

Degrees of Freedom : 4

26
5.2 Model Priors

In the examples considered above, the prior model probabilities \( \text{pr}(M) \) were assumed equal (Cooper and Herskovits, 1991, also assume that models are equally likely \textit{a priori}). In general this can be unrealistic and may also be expensive and we will want to penalise the search strategy as it moves further away from the model(s) provided by the expert(s). Ideally one would elicit prior probabilities for all possible qualitative structures from the expert but this will be feasible only in trivial cases.

For models with fewer than 15 to 20 nodes, prior model probabilities may be approximated by eliciting prior probabilities for the presence of every possible link and assuming that the links are mutually independent, as follows: Let \( \mathcal{E} = \mathcal{E}_P \cup \mathcal{E}_A \) denote the set of all possible links for the nodes of model \( M \), where \( \mathcal{E}_P \) denotes the set of links which are present in model \( M \) and \( \mathcal{E}_A \) denotes the absent links. For every link \( e \in \mathcal{E} \) we elicit \( \text{pr}(e) \), the prior probability that link \( e \) is included in \( M \). The prior model probability is then approximated by

\[
\text{pr}(M) \propto \prod_{e \in \mathcal{E}_P} \text{pr}(e) \prod_{e \in \mathcal{E}_A} (1 - \text{pr}(e)).
\]

Prior link probabilities from multiple experts are treated as independent sources of information and are simply multiplied together to give pooled prior model probabilities. Clearly, the contribution from each expert could be weighted.

For applications involving a larger number of nodes or where the elicitation of link probabilities is not possible, we could assume that the "evidence" in favour of each link included by the expert(s) in the elicited qualitative structure(s) is "substantial" or "strong" but not "very strong" or "decisive" (Jeffreys, 1961). For example, we could assume that the evidence in favour of an included link lies at the center of Occam's window corresponding to a prior link probability for all \( e \in \mathcal{E}_P \) of

\[
\text{pr}(e) = \frac{1}{1 + \exp(\frac{\Theta_P + \Theta_A}{2})}.
\]

Similarly, the prior link probabilities for \( e \in \mathcal{E}_A \) are given by

\[
\text{pr}(e) = \frac{\exp(\frac{\Theta_P + \Theta_A}{2})}{1 + \exp(\frac{\Theta_P + \Theta_A}{2})}.
\]

In the directed case it may be possible to construct a prior distribution on the space of orderings—see Critchlow (1985) for further discussion.
6 Discussion

We have outlined an overall strategy for model selection and accounting for model uncertainty in two important classes of models for high-dimensional contingency tables. This involves a redefinition of the Bayesian model uncertainty formalism, an efficient way of computing exact Bayes factors that exploits the graphical structure, and an algorithm for quickly searching through the very large model classes involved. The resulting procedure is quite efficient: for the example of Section 4.2.1, approximately 3,000 model comparisons per minute can be carried out on a Sun IPC.

There is a considerable literature on model selection for multidimensional contingency tables; this is generally concerned with the selection of a single "best" model. Most of it is based on the asymptotic properties of goodness-of-fit statistics (Wermuth (1976), Havránek (1984), Whittaker (1984), Edwards and Havránek (1985) or Fowlkes et al. (1988)). There are also approaches based on information criteria and discrepancy measures (Gokhale and Kullback, 1978; Sakamoto, 1984; Linhart and Zucchini, 1986). A recent review is provided by Upton (1991) who advocates the use of the BIC statistic. The calculation of Bayes factors for contingency table models has been considered by Spiegelhalter and Smith (1982), Raftery (1986a, 1988), Spiegelhalter and Lauritzen (1990a) and Spiegelhalter and Cowell (1991).

Pearl and Verma (1991) and Glymour et al. (1987) have proposed strategies for recovering causal structure from data. While these authors' objectives differ from ours, their procedures for selecting directed graphical structures have much in common with our recursive causal model selection strategy.

Cooper and Herskovits (1991) and Anderson et al. (1991) have examined model selection in the context of probabilistic expert systems. In both cases, model selection is based solely on data analysis and the incorporation of prior expert opinion is not considered. Cooper and Herskovits (1991) outline a Bayesian strategy which seeks out the "best" recursive causal model for the qualitative layer, where "best" is taken to mean the single model with maximum probability. The algorithm starts with a model with no links and at each stage adds the directed link which most increases the model probability. The user must pre-specify an ordering of the nodes. Anderson et al. (1991) carry out their search in the undirected graphical model framework using a method introduced by Kreiner (1987). The difficulties with large sparse tables mentioned above are avoided by using exact tests when comparing models.

While we believe that the methods we provide a workable approach to qualita-
tive updating in expert systems, some issues remain. Spiegelhalter and Lauritzen (1990a) and other authors have expressed concerns about automatically updating the qualitative structure without reference to the domain expert. Such concerns need to be addressed in the context of real expert systems. Extension of the methods to include the more general graphical models of Wermuth and Lauritzen (1990) and Edwards (1990) will also be important for expert system applications. Missing data will frequently be a problem and we are currently exploring a number of techniques for the incorporation of missing data in the model selection strategy.

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Marcel Dekker.


**MODEL SELECTION AND ACCOUNTING FOR MODEL UNCERTAINTY IN GRAPHICAL MODELS USING OCCAM'S WINDOW**

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**CONTRACT OR GRANT NUMBER(s)**
N00014-88-K-0265

**PROGRAM ELEMENT, PROJECT, TASK, WORK UNIT NUMBERS**
NR-661-003

**REPORT DATE**
July 1991

**NUMBER OF PAGES**
33

**DISTRIBUTION STATEMENT (of this Report)**
APPROVED FOR PUBLIC RELEASE: DISTRIBUTION UNLIMITED.

**KEY WORDS**
(Continue on reverse side if necessary and identify by block number)

**ABSTRACT**
(Continue on reverse side if necessary and identify by block number)
See Reverse side
We consider the problem of model selection and accounting for model uncertainty in high-dimensional contingency tables, motivated by expert system applications. The approach most used currently is a stepwise strategy guided by tests based on approximate asymptotic $P$-values leading to the selection of a single model; inference is then conditional on the selected model. The sampling properties of such a strategy are complex, and the failure to take account of model uncertainty leads to underestimation of uncertainty about quantities of interest. In principle, a panacea is provided by the standard Bayesian formalism which averages the posterior distributions of the quantity of interest under each of the models, weighted by their posterior model probabilities. However, this has not been used in practice because computing the posterior model probabilities is hard and the number of models is very large (often greater than $10^{11}$).

We argue that the standard Bayesian formalism is unsatisfactory and we propose an alternative Bayesian approach that, we contend, takes full account of the true model uncertainty by averaging over a much smaller set of models. An efficient search algorithm is developed for finding these models. We consider two classes of models that arise in expert systems: the recursive causal models and the decomposable log-linear models. For each of these, we develop efficient ways of computing exact Bayes factors and hence posterior model probabilities. For the decomposable log-linear models, this is based on properties of chordal graphs and hyper Markov prior distributions and the resultant calculations can be carried out locally. The end product is an overall strategy for model selection and accounting for model uncertainty that searches efficiently through the very large classes of models involved.

Three examples are given. The first two concern data sets which have been analysed by several authors in the context of model selection. The third addresses a urological diagnostic problem.

KEYWORDS: Chordal graph; Contingency table; Decomposable log-linear model; Expert system; Hyper Markov distribution; Recursive causal model.