Data Mining Graphical Models for Discrete Data Undirected Graphs (2)

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Overview of Coming Two Lectures

- Introduction
- Independence and Conditional Independence
- Graphical Representation of Conditional Independence
- Log-linear Models
 - Hierarchical
 - Graphical
 - Decomposable
- Maximum Likelihood Estimation
- Model Testing
- Model Selection

2×2 Table

The probability function P_{12} of bivariate Bernoulli random vector (X_1, X_2) is determined by

$$P(x_1,x_2) = p(x_1,x_2)$$

where $p(x_1, x_2)$ is the table of probabilities:

$p(x_1,x_2)$	$x_2 = 0$	$x_2 = 1$	Total
$x_1 = 0$	p(0,0)	p(0,1)	$p_1(0)$
$x_1 = 1$	p(1,0)	p(1, 1)	$p_1(1)$
Total	$p_2(0)$	$p_{2}(1)$	1

Probability function for 2×2 Table

Again we can write this as a single formula:

$$P(x_1, x_2) = p(0,0)^{(1-x_1)(1-x_2)}p(0,1)^{(1-x_1)x_2}p(1,0)^{x_1(1-x_2)}p(1,1)^{x_1x_2}$$

Taking logarithms and collecting terms in x_1 , x_2 , and x_1x_2 gives:

$$\log P(x_1, x_2) = \log p(0, 0) + \log \frac{p(1, 0)}{p(0, 0)} x_1 + \log \frac{p(0, 1)}{p(0, 0)} x_2 + \log \frac{p(1, 1)p(0, 0)}{p(0, 1)p(1, 0)} x_1 x_2$$

Verify this using elementary properties of logarithms:



Log-linear expansion

Reparameterizing the right hand side leads to the so-called *log-linear* expansion

$$\log P(x_1, x_2) = u_{\emptyset} + u_1 x_1 + u_2 x_2 + u_{12} x_1 x_2$$

The coefficients, u_{\emptyset} , u_1 , u_2 , u_{12} are known as the *u*-terms.

For example, the coefficient of the product x_1x_2 ,

$$u_{12} = \log \frac{p(1,1)p(0,0)}{p(0,1)p(1,0)} = \log \operatorname{cpr}(X_1, X_2)$$

is the logarithm of the cross product ratio of X_1 and X_2 .

Independence and *u*-terms

Claim:

$$X_1 \perp \!\!\! \perp X_2 \Leftrightarrow u_{12} = 0$$

Proof: the factorisation criterion states that $X_1 \perp \!\!\! \perp X_2$ iff there exist two functions g and h such that

$$\log P(x_1, x_2) = g(x_1) + h(x_2)$$
 for all (x_1, x_2)

If $u_{12} = 0$, we get

$$\log P(x_1, x_2) = u_{\emptyset} + u_1 x_1 + u_2 x_2,$$

SO

$$g(x_1) = u_{\emptyset} + u_1 x_1$$
 $h(x_2) = u_2 x_2$

suffices. If $u_{12} \neq 0$, no such decomposition is possible.



Three Dimensional Bernoulli

The joint distribution of three binary variables can be written:

$$P(x_1, x_2, x_3) = p(0, 0, 0)^{(1-x_1)(1-x_2)(1-x_3)} \cdots p(1, 1, 1)^{x_1x_2x_3}$$

Log-linear expansion

$$\log P(x_1, x_2, x_3) = u_{\emptyset} + u_1 x_1 + u_2 x_2 + u_3 x_3 + u_{12} x_1 x_2 + u_{13} x_1 x_3 + u_{23} x_2 x_3 + u_{123} x_1 x_2 x_3$$

With

$$u_{123} = \log \frac{p(1,1,1)p(1,0,0)}{p(1,1,0)p(1,0,1)} - \log \frac{p(0,1,1)p(0,0,0)}{p(0,1,0)p(0,0,1)}$$
$$= \log \frac{\operatorname{cpr}(X_2, X_3 | X_1 = 1)}{\operatorname{cpr}(X_2, X_3 | X_1 = 0)}$$

Independence and the *u*-terms

Observation:

$$X_2 \perp \!\!\! \perp X_3 | X_1 \Leftrightarrow u_{23} = 0$$
 and $u_{123} = 0$

Proof: use factorisation criterion.

 $X_2 \perp \!\!\! \perp X_3 | X_1 \Leftrightarrow$ there are functions $g(x_1,x_2)$ and $h(x_1,x_3)$ such that

$$\log P(x_1, x_2, x_3) = g(x_1, x_2) + h(x_1, x_3)$$

This is only possible when $u_{23} = 0$ (so the term x_2x_3 drops out), and $u_{123} = 0$ (so the term $x_1x_2x_3$ drops out).

Why the log-linear representation?

Why do we use the log-linear representation of the probability table?

- We are interested in expressing conditional independence constraints.
- There is a straightforward correspondence between such constraints being satisfied, and the elimination of certain collections of u-terms from the log-linear expansion.
- **3** This correspondence is established by applying the factorisation criterion: $X \perp \!\!\! \perp Y \mid Z$ if and only if there exist functions g and h such that

$$\log P(x,y,z) = g(x,z) + h(y,z)$$

Log-linear expansion: non-binary variables

For a 2×2 table the log-linear expansion is given by:

$$\log P(x_1, x_2) = u_{\emptyset} + u_1 x_1 + u_2 x_2 + u_{12} x_1 x_2$$

for $x \in \{0, 1\}^2$.

What if the x_i have more than two levels? In that case the u terms become functions of x rather than constants:

$$\log P(x_1, x_2) = u_{\emptyset} + u_1(x_1) + u_2(x_2) + u_{12}(x_1, x_2)$$

Log-linear expansion: non-binary variables

Suppose $x \in \{0, 1, 2\}$. We can write

$$P(x) = p(1)^{\delta_{x=1}} p(2)^{\delta_{x=2}} p(0)^{(1-\delta_{x=1}-\delta_{x=2})},$$

where δ_A is the indicator function, that is,

$$\delta_A = \left\{ \begin{array}{ll} 1 & \text{if } A \text{ is true} \\ 0 & \text{otherwise} \end{array} \right.$$

Taking logarithms left and right, we get

$$\begin{split} \log P(x) &= \delta_{x=1} \log p(1) + \delta_{x=2} \log p(2) + (1 - \delta_{x=1} - \delta_{x=2}) \log p(0) \\ &= \delta_{x=1} \log p(1) + \delta_{x=2} \log p(2) + \log p(0) - \delta_{x=1} \log p(0) - \delta_{x=2} \log p(0) \\ &= \log p(0) + \log \frac{p(1)}{p(0)} \delta_{x=1} + \log \frac{p(2)}{p(0)} \delta_{x=2} \\ &= u_{\emptyset} + u(x) \end{split}$$

Log-linear expansion: non-binary variables

Where $u_{\emptyset} = \log p(0)$ and

$$u(x) = \begin{cases} \log \frac{p(1)}{p(0)} & \text{if } x = 1\\ \log \frac{p(2)}{p(0)} & \text{if } x = 2\\ 0 & \text{if } x = 0 \end{cases}$$

Similar rules apply to the case of multiple non-binary variables.

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Log-linear expansion: general

The log-linear expansion of the probability distribution P_K is

$$\log P_K(x) = \sum_{a \subseteq K} u_a(x_a)$$

where the sum is taken over all possible subsets a of $K = \{1, 2, ..., k\}$.

- We get a (set of) u-term(s) for each subset of the variables.
- Code the values of X_i as $\{0, 1, \dots, d_i 1\}$, where d_i is size of the domain of X_i .
- Set $u_a(x_a) = 0$ whenever $x_i = 0$ for any X_i with $i \in a$.
- This is analogous to the case where X is binary.
- There are as many u-terms in the full log-linear expansion as there are cells in the contingency table.

Independence and the u-terms

Let

- $X = (X_1, X_2, \dots, X_k)$
- $a, b, c \subseteq \{1, 2, \dots, k\}$
- a, b, c disjoint and $a \cup b \cup c = \{1, 2, \dots, k\}$

Independence and the u-terms

We have

$$X_b \perp \!\!\! \perp X_c \mid X_a$$

if and only if all u-terms in the log-linear expansion with coordinates in both b and c, are equal to zero.

Independence and the u-terms: Example

Let $X = (X_1, ..., X_5)$, and $a = \{1, 3\}, b = \{4\}, c = \{2, 5\}$. According to the factorization criterion, the conditional independence

$$X_4 \perp \!\!\! \perp (X_2, X_5) \mid (X_1, X_3)$$

holds if, and only if, there are functions g and h such that

$$\log P(x_1,\ldots,x_5) = g(x_1,x_3,x_4) + h(x_1,x_2,x_3,x_5)$$

For this to be possible, the u-terms that contain elements from both the sets $\{4\}$ and $\{2,5\}$ have to be zero. So all u-terms $u_{24}, u_{45}, u_{124}, u_{145}, \ldots, u_{12345}$ have to be zero.

Independence and the u-terms

Proof: Let s be an arbitrary subset of $a \cup b \cup c = \{1, 2, \dots, k\}$.

If all u-terms, u_s , are zero whenever

- \bullet $s \not\subseteq a \cup b \ (s \cap c \neq \emptyset)$, and

(i.e. whenever s contains coordinates from both b and c), then

$$\log P_{\mathcal{K}}(x) = \sum_{s \subseteq a \cup b} u_s(x_s) + \sum_{s \subseteq a \cup c} u_s(x_s) - \sum_{s \subseteq a} u_s(x_s)$$

But this function is of the form $g(x_a, x_b) + h(x_a, x_c)$ and hence $X_b \perp \!\!\! \perp X_c \mid X_a$ by the factorisation criterion.

Hierarchical Log-Linear Models

- In most applications, it does not make sense to include the three-way association u_{123} unless the two-way associations u_{12} , u_{13} and u_{23} are all present as well.
- A log-linear model is said to be *hierarchical* if the presence of a term implies that all lower-order terms are also present. That is, if $u_A(x_A)$ is present, then for all $a \subseteq A$, $u_a(x_a)$ must be present as well.
- Hence, a hierarchical model is uniquely identified by listing its highest order interaction terms.

Hierarchical Models for three dimensions

Model	Omitted	Interpretation
123	none	saturated
12,13,23	u_{123}	homogeneous association
12,13	u_{123}, u_{23}	$X_2 \perp \!\!\! \perp X_3 \mid X_1$
12,23	u_{123}, u_{13}	$X_1 \perp \!\!\! \perp X_3 \mid X_2$
13,23	u_{123}, u_{12}	$X_1 \perp \!\!\! \perp X_2 \mid X_3$
12,3	u_{123}, u_{13}, u_{23}	$(X_1,X_2) \perp \!\!\! \perp X_3$
13,2	u_{123}, u_{12}, u_{23}	$(X_1,X_3) \perp \!\!\! \perp X_2$
23,1	u_{123}, u_{12}, u_{13}	$(X_2,X_3) \perp \!\!\! \perp X_1$
1,2,3	$u_{123}, u_{12}, u_{13}, u_{23}$	mutual independence

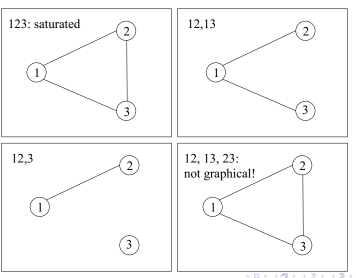
Graphical Log-Linear Model

Given its independence graph G = (K, E), the log-linear model for the random vector X is a graphical model for X if the distribution of X is arbitrary apart from constraints of the form that for all pairs of coordinates not in the edge set E, the u-terms containing the selected coordinates are equal to zero.

All constraints of a graphical model can be read from the independence graph.

A graphical model is a hierarchical model in which the highest order interaction terms correspond to the cliques in the graph.

Hierarchical models and their independence graphs



Estimating Models from Data: Maximum Likelihood

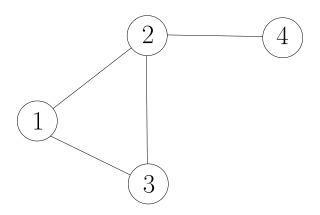
- ML estimator of graphical log-linear model M returns estimates of the cell probabilities that maximize the probability of the observed data, subject to the constraint that the conditional independencies of M are satisfied by the estimates.
- ullet ML estimator of graphical log-linear model M satisfies the likelihood equations

$$\hat{n}_a^M = N\hat{P}_a^M = n_a$$

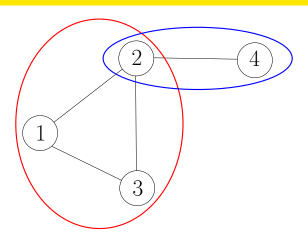
whenever the subset of vertices a in the graph form a clique.

- Slogan: Observed = Fitted for every marginal table corresponding to a complete subgraph.
- The same likelihood equations hold for all hierarchical models, where the margins a correspond to the highest order interaction terms in the model.

ML: Determine the cliques



ML: Observed=Fitted for margins corresponding to cliques



$$\hat{n}(x_1, x_2, x_3) = n(x_1, x_2, x_3)$$

 $\hat{n}(x_2, x_4) = n(x_2, x_4)$

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ML: Observed=Fitted for margins corresponding to cliques

We can see as follows why this has to be the case:

- If there are no constraints to fit an observed table of counts, then the parameter estimates that yield fitted counts equal to the observed counts maximize the likelihood function. For example, the saturated model will yield fitted counts identical to the observed counts.
- ② By definition, a graphical model is arbitrary (has no constraints) except for the constraints that can be read from the independence graph.
- **3** Suppose a forms a clique in the independence graph. Now consider the partitioning $X = (X_a, X_b)$ where b contains all variables not in a. We can write (product rule):

$$P(X) = P(X_a)P(X_b \mid X_a)$$

Since $P(X_a)$ is not constrained by the model (complete graph), all model constraints apply only to $P(X_b \mid X_a)$. Therefore, the maximum likelihood estimates will yield $\hat{n}_a = n_a$.

Maximum Likelihood Estimation: Example

$$\begin{split} \hat{P}(x_1, x_2, x_3, x_4) &= \hat{P}(x_1, x_3, x_4 | x_2) \hat{P}(x_2) & \text{(product rule)} \\ &= \hat{P}(x_1, x_3 | x_2) \hat{P}(x_4 | x_2) \hat{P}(x_2) & \text{($X_4 \perp \!\!\! \perp (X_1, X_3) \mid X_2$)} \\ &= \hat{P}(x_1, x_3 | x_2) \hat{P}(x_2, x_4) & \text{(product rule)} \\ &= \frac{\hat{P}(x_1, x_2, x_3) \hat{P}(x_2, x_4)}{\hat{P}(x_3)} & \text{(product rule)} \end{split}$$

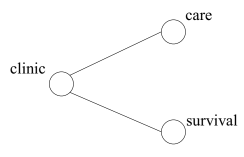
In terms of counts we have:

$$\begin{split} \hat{n}(x_1, x_2, x_3, x_4) &= \frac{\hat{n}(x_1, x_2, x_3) \hat{n}(x_2, x_4)}{\hat{n}(x_2)} \\ &= \frac{n(x_1, x_2, x_3) n(x_2, x_4)}{n(x_2)} \end{split} \tag{fitted} = \text{observed for complete subgraph})$$

In this case we have a closed form solution for the maximum likelihood fitted counts.

ML Estimation: Numeric Example

$\overline{n(c, ca, s)}$	survival		
clinic	care	no	yes
clinic 1	less	3	176
	more	4	293
clinic 2	less	17	197
	more	2	23



Margin Constraints and Sufficient Statistics

The margin constraints are:

- ② $\hat{n}(\text{clinic}, \text{survival}) = n(\text{clinic}, \text{survival})$

$\overline{n(c,ca)}$	care		
clinic	less more		
clinic 1	179	297	
clinic 2	214	25	

n(c,s)	survival		
clinic	no	yes	
clinic 1	7	469	
clinic 2	19	220	

Fitted values

The maximum likelihood fitted counts are given by:

$$\hat{n}(\text{clinic}, \text{care}, \text{survival}) = \frac{n(\text{clinic}, \text{care})n(\text{clinic}, \text{survival})}{n(\text{clinic})}$$

For example:

$$\hat{n}(\mathsf{clinic}\ 1,\mathsf{less},\mathsf{yes}) = \frac{n(\mathsf{clinic}\ 1,\mathsf{less})n(\mathsf{clinic}\ 1,\mathsf{yes})}{n(\mathsf{clinic}\ 1)} = \frac{179\times469}{476} = 176.37$$

$\hat{n}(c, ca, s)$	survival		
clinic	care	no	yes
clinic 1	less	2.63	176.37
	more	4.37	292.63
clinic 2	less	17.01	196.99
	more	1.99	23.01

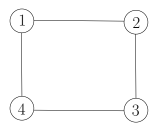
Model seems to fit very well!

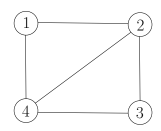
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Decomposable Graphical Models

- Decomposable graphical models have explicit formulas for the maximum likelihood estimates.
- Decomposable models have *triangulated* independence graphs: they have no *chordless cycles* of length greater than three.
- A cycle is chordless if only the *successive* pairs of vertices in the cycle are connected by an edge. (There is no "shortcut" in the cycle).

Example





- The left graph is *not* decomposable because it contains the chordless cycle 1-2-3-4-1.
- The graph on the right is decomposable. The cycle 1-2-3-4-1 is no longer chordless because 2 and 4 are adjacent in the graph but not successive in the cycle.

Maximum Likelihood Estimation for Decomposable Models

An ordering C_1, \ldots, C_m of the cliques of the graph has the running intersection property (RIP) iff

$$C_j \cap (C_1 \cup \ldots \cup C_{j-1}) \subseteq C_i$$

for some i < j, and for j = 2, ..., m.

We define the corresponding separator sets

$$S_j = C_j \cap (C_1 \cup \ldots \cup C_{j-1}),$$

with $S_1 = \emptyset$.

ML Estimation for Decomposable Models

Let C_1, \ldots, C_m be a RIP ordering of the cliques, with separator sets S_1, \ldots, S_m .

The maximum likelihood fitted counts are given by

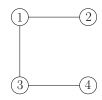
$$\hat{n}(x) = \frac{\prod_{j=1}^{m} n(x_{C_j})}{\prod_{j=2}^{m} n(x_{S_j})}$$

where $n(x_{\emptyset}) = N$.

Likewise, the maximum likelihood fitted probabilities are given by

$$\hat{P}(x) = \frac{\prod_{j=1}^{m} n(x_{C_j})}{\prod_{j=1}^{m} n(x_{S_j})}$$

Example



The clique ordering $C_1 = \{1,2\}$, $C_2 = \{3,4\}$, $C_3 = \{1,3\}$ does not have the running intersection property and does therefore not produce correct estimates.

j	C_j	S_j	RIP?
1	$\{1, 2\}$	Ø	_
2	$\{3,4\}$	Ø	√
3	$\{1, 3\}$	$\{1, 3\}$	 X

Example



The clique ordering $C_1 = \{1, 2\}$, $C_2 = \{1, 3\}$, $C_3 = \{3, 4\}$ does have the running intersection property:

j	C_j	S_j	RIP?
1	{1,2}	Ø	_
2	{1,3}	{1}	 √
3	{3,4}	{3}	√

The corresponding ML fitted counts are:

$$\hat{n}(x_1, x_2, x_3, x_4) = \frac{n(x_1, x_2)n(x_1, x_3)n(x_3, x_4)}{n(x_1)n(x_3)}$$

Iterative Proportional Fitting (IPF)

- If the model is not decomposable, there is no closed form solution (formula) for the maximum likelihood estimates.
- We need iterative algorithms to compute the fitted counts/probabilities.
- Iterative Proportional Fitting (IPF) is such an algorithm.

IPF: Example

Fit independence model to

$n(x_1,x_2)$	$x_2 = 0$	$x_2 = 1$	$n_1(x_1)$
$x_1 = 0$	30	10	40
$x_1 = 1$	30	30	60
$n_2(x_2)$	60	40	100

The margin constraints for the independence model are

$$\hat{n}_1(x_1) = n_1(x_1)$$
, and

$$\hat{n}_2(x_2) = n_2(x_2)$$

IPF makes the fitted counts agree with the observed counts on each margin in turn.

We begin with a table $\hat{n}^{(0)}$ of uniform counts (left)

First step: fit to row margin

$$\hat{n}(x_1, x_2)^{(1)} = n_1(x_1) \times \frac{\hat{n}(x_1, x_2)^{(0)}}{\hat{n}_1(x_1)^{(0)}}$$

We compute (row 1):

$$\hat{n}(0,0)^{(1)}=40\times\frac{1}{2}=20$$

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We begin with a table $\hat{n}^{(0)}$ of uniform counts (left)

First step: fit to row margin

$$\hat{n}(x_1, x_2)^{(1)} = n_1(x_1) \times \frac{\hat{n}(x_1, x_2)^{(0)}}{\hat{n}_1(x_1)^{(0)}}$$

We compute (row 1):

$$\hat{n}(0,0)^{(1)}=40\times\frac{1}{2}=20$$

$$\hat{n}(0,1)^{(1)} = 40 \times \frac{1}{2} = 20$$

$$\begin{array}{ccccc}
\hat{n}^{(0)} & 0 & 1 \\
0 & 1 & 1 & 2 \\
1 & 1 & 1 & 2 \\
& 2 & 2 &
\end{array}$$

$$\begin{array}{c|ccc}
0 & 1 \\
0 & 20 & 20 & 40 \\
1 & & & 60
\end{array}$$

First step continued (row 2):

$$\hat{n}(1,0)^{(1)}=60\times\frac{1}{2}=30$$

$$\hat{n}(1,1)^{(1)}=60\times\frac{1}{2}=30$$

which yields $\hat{n}^{(1)}$:

$$\begin{array}{cccc}
\hat{n}^{(1)} & 0 & 1 \\
0 & 20 & 20 & 40 \\
1 & 30 & 30 & 60 \\
\hline
& 50 & 50 & 50
\end{array}$$

$$\begin{array}{cccc}
\hat{n}^{(1)} & 0 & 1 \\
0 & 20 & 20 & 40 \\
1 & 30 & 30 & 60 \\
\hline
& 50 & 50 & & & \\
\end{array}$$

$$\begin{array}{c|c} 0 & 1 \\ 0 & \\ 1 & \\ \hline 60 & 40 \end{array}$$

Second step: fit to column margin

$$\hat{n}(x_1,x_2)^{(2)} = n_2(x_2) \times \frac{\hat{n}(x_1,x_2)^{(1)}}{\hat{n}_2(x_2)^{(1)}}$$

Which gives (first column):

$$\hat{n}(0,0)^{(2)} = 60 \times \frac{20}{50} = 24$$

$$\hat{n}(1,0)^{(2)} = 60 \times \frac{30}{50} = 36$$

$$\begin{array}{c|cccc}
0 & 1 \\
0 & 24 \\
1 & 36 \\
\hline
60 & 40
\end{array}$$



Likewise for the second column:

$$\hat{n}(0,1)^{(2)} = 40 \times \frac{20}{50} = 16$$
 $\hat{n}(1,1)^{(2)} = 40 \times \frac{30}{50} = 24$

This yields $\hat{n}^{(2)}$:

$$\begin{array}{ccccc}
\hat{n}^{(2)} & 0 & 1 \\
0 & 24 & 16 & 40 \\
1 & 36 & 24 & 60 \\
\hline
& 60 & 40 & 0
\end{array}$$

Notice that the row totals are still 40 and 60, so we have simultaneously satisfied the conditions

$$\hat{n}_1(x_1) = n_1(x_1)$$
 and $\hat{n}_2(x_2) = n_2(x_2)$

so we have converged.

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IPF: General Algorithm Sketch

Say we have m margins $\{a_1, a_2, \ldots, a_m\}$ to be fitted $(\cup_i a_i = K)$.

We have to find a table $\hat{n}(x)$ that agrees with the observed table n(x) on the m margins corresponding to the subsets a_i .

The algorithm cycles through the list of subsets

$$a=a_i, \qquad \qquad i=1,2,\ldots,m$$

fitting $\hat{n}(x)$ to each margin in turn.

This is repeated until convergence is reached, i.e. all margin constraints are (approximately) satisfied simultaneously.

IPF updating rule

To fit to the margin a, the observed count $n_a(x_a)$ on x_a is distributed over the cells $x = (x_a, x_b)$ of the full table according to

$$\hat{n}_{ab}(x_a, x_b)^{(t+1)} = n_a(x_a)\hat{P}(x_b|x_a)^{(t)}$$

where b is the complement of a, and

$$\hat{P}(x_b|x_a)^{(t)} = \frac{\hat{n}_{ab}(x_a, x_b)^{(t)}}{\hat{n}_a(x_a)^{(t)}},$$

is the current estimate of the conditional probability $P(X_b = x_b | X_a = x_a)$.

IPF Pseudocode

Algorithm 1 IPF(n(x), A)

```
1: t \leftarrow 0
 2. for all values x of X do
         \hat{n}(x)^{(t)} \leftarrow 1
 3. end for
 4: repeat
 5:
         for all margins a \in \mathcal{A} do
            for all values x_a of X_a do
 6:
                for all values x_b of X_b do
 7.
                    \hat{n}_{ab}(x_a, x_b)^{(t+1)} \leftarrow n_a(x_a) \left( \frac{\hat{n}_{ab}(x_a, x_b)^{(t)}}{\hat{n}_a(x_a)^{(t)}} \right)
 8.
                end for
            end for
 9:
10:
         t \leftarrow t + 1
         end for
11.
12: until convergence
```

Some properties of IPF

- If the cliques of a decomposable model are presented in RIP order to IPF, then the algorithm will converge in one iteration (one cycle through all cliques).
- Otherwise IPF will converge in two iterations on decomposable models.

The Log-Likelihood score of a model

The likelihood score of a model M is

$$L^M = \prod_x \hat{P}^M(x)^{n(x)},$$

where $\hat{P}^{M}(x)$ is the fitted probability of cell x according to model M.

Hence, the likelihood score of model M is the probability of the observed data using the fitted cell probabilities according to model M.

Likewise, the log-likelihood score of a model M is

$$\mathcal{L}^{M} = \sum_{x} n(x) \log \hat{P}^{M}(x)$$

Example

Suppose we have data

n(x)	$x_2 = 0$	$x_2 = 1$	$n(x_1)$
$x_1 = 0$	30	10	40
$x_1 = 1$	30	30	60
$n(x_2)$	60	40	100

The independence model gives probability estimates:

$$\hat{P}(0,0) = 0.24$$
, $\hat{P}(0,1) = 0.16$, $\hat{P}(1,0) = 0.36$, $\hat{P}(1,1) = 0.24$.

The probability of the observed data according to this model is

$$0.24^{30}\times0.16^{10}\times0.36^{30}\times0.24^{30}$$

This is the likelihood score of the model given the data.

The corresponding log-likelihood score is

$$\mathcal{L} = 30\log 0.24 + 10\log 0.16 + 30\log 0.36 + 30\log 0.24 \approx -134.6$$

Example (continued)

n(x)	$x_2 = 0$	$x_2 = 1$	$n(x_1)$
$x_1 = 0$	30	10	40
$x_1 = 1$	30	30	60
$n(x_2)$	60	40	100

The saturated model gives probability estimates:

$$\hat{P}(0,0) = 0.3$$
, $\hat{P}(0,1) = 0.1$, $\hat{P}(1,0) = 0.3$, $\hat{P}(1,1) = 0.3$.

The probability of the observed data according to this model is

$$0.3^{30}\times0.1^{10}\times0.3^{30}\times0.3^{30}$$

The corresponding log-likelihood score is

$$\mathcal{L} = 30\log 0.3 + 10\log 0.1 + 30\log 0.3 + 30\log 0.3 \approx -131.4$$

Of course this is better than the independence model.

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Model Deviance

Since for the saturated (unconstrained) model

$$\hat{P}(x) = \frac{n(x)}{N},$$

the log-likelihood score of the saturated model is

$$\mathcal{L}^{\mathsf{sat}} = \sum_{x} n(x) \log \frac{n(x)}{N}$$

The deviance of a fitted model compares the log-likelihood score of the fitted model to that of the saturated model.

The larger the model deviance, the poorer the fit.

Model Deviance

Deviance of M is 2 (log-likelihood of the saturated model - log-likelihood of M):

$$\begin{aligned} \operatorname{dev}(M) &= 2(\mathcal{L}^{\mathsf{Sat}} - \mathcal{L}^{M}) \\ &= 2\left(\sum_{x} n(x) \log \frac{n(x)}{N} - \sum_{x} n(x) \log \hat{P}^{M}(x)\right) \\ &= 2\left(\sum_{x} n(x) \left(\log \frac{n(x)}{N} - \log \hat{P}^{M}(x)\right)\right) \\ &= 2\sum_{x} n(x) \log \frac{n(x)}{N\hat{P}^{M}(x)} \end{aligned}$$

which can be summarised by the slogan

$$2\sum_{\mathsf{cells}}\mathsf{observed}\times\mathsf{log}\,\frac{\mathsf{observed}}{\mathsf{fitted}}$$

Deviance difference

Let $M_0 \subseteq M_1$, that is M_0 is the simpler model (the *u*-terms present in M_0 are a subset of the *u*-terms present in M_1).

The deviance difference between M_0 and M_1 is

$$\operatorname{dev}(M_0) - \operatorname{dev}(M_1) = -2\mathcal{L}^{M_0} + 2\mathcal{L}^{M_1} = 2(\mathcal{L}^{M_1} - \mathcal{L}^{M_0})$$

For large N we have that:

$$2(\mathcal{L}^{M_1}-\mathcal{L}^{M_0})\approx_{M_0}\chi^2_{\nu}$$

 χ^2_{ν} : chi-square distribution with ν degrees of freedom. ν : number of *additional* restrictions (zero *u*-terms) of M_0 compared to M_1 .

Likelihood Ratio Test

We reject the null hypothesis that M_0 is the true model when

$$\boxed{2(\mathcal{L}^{M_1}-\mathcal{L}^{M_0})>\chi^2_{\nu;\alpha},}$$

where α is the significance level of the test, and $P(X^2 > \chi^2_{\nu;\alpha}) = \alpha$, that is, $\chi^2_{\nu;\alpha}$ is the *critical value*.

The test is called a likelihood ratio test because

$$\log \frac{L^{M_1}}{L^{M_0}} = \log L^{M_1} - \log L^{M_0} = \mathcal{L}^{M_1} - \mathcal{L}^{M_0}$$

Model Testing: example

Does

give a good fit of the observed table? Test against the saturated model.

Compute the deviance

$$2\sum_{\text{cells}} \text{observed} \times \log \frac{\text{observed}}{\text{fitted}} \approx 0.082$$

$$\chi^2_{2;0.05}\approx 6$$

So we "accept" the model.

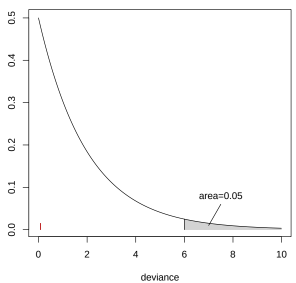
Note: since M_1 is the saturated model, the deviance difference between M_0 and M_1 is just the deviance of M_0 .

Fitted Counts and Observed Counts

$\hat{n}(\text{clinic, care, survival})$		survival		
clinic	care	no	yes	
clinic 1	less	2.63	176.37	
	more	4.37	292.63	
clinic 2	less	17.01	196.99	
	more	1.99	23.01	

n(clinic, care, survival)		survival	
clinic	care	no	yes
clinic 1	less	3	176
	more	4	293
clinic 2	less	17	197
	more	2	23

Test of survival $\perp \perp$ care clinic; χ_2^2 distribution.



Model Testing: example

Does the mutual independence model give a good fit of the observed table? Test against the saturated model.

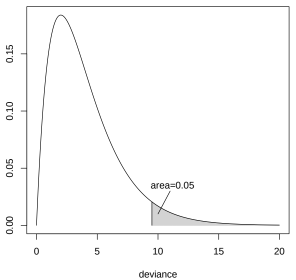
Compute the deviance

$$2\sum_{\text{cells}} \text{observed} \times \log \frac{\text{observed}}{\text{fitted}} \approx 211$$

$$\chi^2_{4;0.05}\approx 9.5$$

So we reject the mutual independence model.

Test of Independence Model; χ_4^2 distribution.



Model Selection

The Problem: find a good model for a high-dimensional table when little prior knowledge is available.

Solution: Search the space of possible models.

Two approaches:

- Use significance testing
- Use a quality function

Quality Functions: AIC and BIC

Two components:

- the lack-of-fit of the model
- complexity of the model

Akaike's Information Criterion assigns quality to model M as follows

$$\mathsf{AIC}(M) = \mathsf{dev}(M) + 2 \; \mathsf{dim}(M)$$

where dim(M) is the number of parameters (the number of u-terms) of the model.

Bayesian Information Criterion assigns quality to model M as follows

$$\mathsf{BIC}(M) = \mathsf{dev}(M) + \mathsf{log}(N) \; \mathsf{dim}(M)$$

Search

Exhaustive search is usually not feasible.

A straightforward approach is local search with hill climbing:

- pick some initial model
- consider the quality of all neighbors of the current model
- if they all have lower quality, stop and return the current model.
- otherwise move to the neighbor with highest quality and return to 2.

Example: Decomposable Graphical Models

Hill climbing local search with decomposable graphical models and AIC scoring.

- 1 pick an initial model, e.g. the empty graph
- neighbors
 - add an edge
 - delete an edge

(without creating a chordless cycle of length > 3)

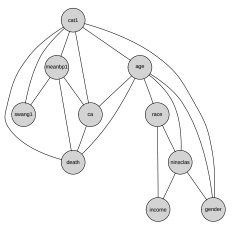
- o if all neighbors have higher AIC, stop and return the current model.
- otherwise move to the neighbor with lowest AIC and return to 2.

Local search in gRim

```
# fit initial model (empty graph)
> rhc.init <- dmod(~.^1,data=rhc.dat)</pre>
# display some info about this model
> summary(rhc.init)
is graphical=TRUE; is decomposable=TRUE
generators (glist):
  : "cat1"
  :"death"
  :"swang1"
  : "gender"
  :"race"
  :"ninsclas"
  :"income"
  :"ca"
  : "age"
  : "meanbp1"
```

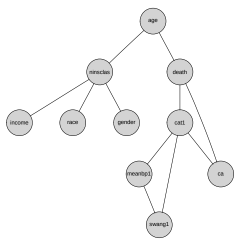
Result of Search (AIC, decomposable)

- $\hbox{\tt\# perform stepwise search of decomposable models using AIC (only add edges)}$
- > rhc.step1 <- stepwise(rhc.init,direction="forward")</pre>
- > plot(rhc.step1)



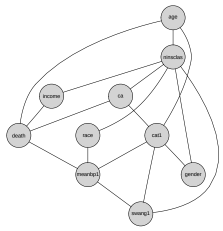
Result of Search (BIC, decomposable)

- > rhc.step2 <- stepwise(rhc.init,direction="forward",k=log(nrow(rhc.dat)))</pre>
- > plot(rhc.step2)



Result of Search (BIC, unrestricted)

- > plot(rhc.step3)



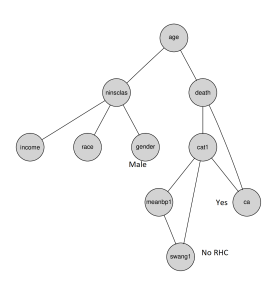
Trace of Search (AIC, unrestricted)

```
> rhc.step4 <- stepwise(rhc.init,direction="both",</pre>
                type="unrestricted",details=1)
STEPWISE:
 criterion: aic (k = 2)
direction: both
type : unrestricted
 search : all
 steps : 1000
. FORWARD: type=unrestricted search=all, criterion=aic(2.00), alpha=0.00
. Initial model: is graphical=TRUE is decomposable=TRUE
  change.AIC -3061.4959 Edge added: ninsclas age
  change.AIC -1685.0212 Edge added: cat1 ca
  change.AIC -1347.0960 Edge added: income ninsclas
  change.AIC -420.0886 Edge added: swang1 cat1
  change.AIC -306.3228 Edge added: race ninsclas
  change.AIC -285.8656 Edge added: age cat1
  change.AIC -253.7602 Edge added: death ca
  change.AIC -497.6596 Edge added: age death
  . . .
```

Model Use: Inference and Prediction with gRain

```
# prepare selected model for use in gRain
> rhc.mod2 <- grain(as(rhc.step2, "graphNEL"), rhc.dat)</pre>
# perform inference on "death" from evidence on "gender", "ca", and "swang1"
> predict(rhc.mod2, c("death"), c("gender", "ca", "swang1"),
  data.frame(gender="Male",ca="Yes",swang1="No RHC"), type = "dist")
$pred
$pred$death
            Nο
                     Yes
[1,] 0.2235305 0.7764695
# change cancer to "Metastatic"
> predict(rhc.mod2, c("death"), c("gender", "ca", "swang1"),
  data.frame(gender="Male",ca="Metastatic",swang1="No RHC"), type = "dist")
$pred
$pred$death
             Nο
                      Yes
[1.] 0.09421555 0.9057845
```

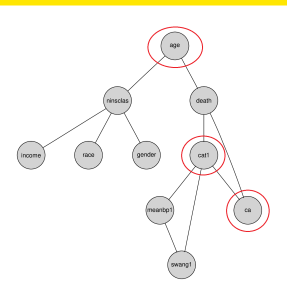
Evidence entered



Model Use: Inference and Prediction with gRain

```
# predict death (in-sample) from its Markov blanket
> death.pred <- predict(rhc.mod2, c("death"), c("ca","cat1","age"),</pre>
                        rhc.dat, type = "class")
> table(rhc.dat$death,death.pred$pred$death)
        No Yes
  Nο
      730 1283
  Yes 524 3198
> (730+3198)/nrow(rhc.dat)
[1] 0.6849172
# model is a little better than just predicting the majority class
> summary(rhc.dat$death)
  No Yes
2013 3722
> 3722/nrow(rhc.dat)
[1] 0.6489974
```

Markov Blanket of Death



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