ISPOR Europe 2019

Copenhagen, 2-6 November

Short course:

Probabilistic graphical models with OpenMarkov, an open-source tool

Francisco Javier Díez

Dept. Artificial Intelligence. UNED Madrid, Spain

www.ia.uned.es/~fjdiez www.cisiad.uned.es

Probabilistic graphical models with OpenMarkov, an opensource tool

Short course at <u>ISPOR Europe 2019</u> (Copenhagen, Denmark)

Presenter:

Francisco Javier Díez, PhD Dept. Artificial Intelligence, UNED, Madrid, Spain

Description

This course will introduce probabilistic graphical models (PGMs), such as Bayesian networks, influence diagrams and decision analysis networks, and discuss their advantages over traditional techniques; for example, influence diagrams and decision analysis networks are equivalent to decision trees containing thousands of branches, Markov influence diagrams can model state-transition problems without multiplying the number of states and decision analysis networks can evaluate large models with unordered decisions. OpenMarkov, an open-source tool, allows building PGMs for complex problems using a graphical user interface, without writing any code, such as spreadsheet formulas, macros or functions. For beginners, the course will be an introduction to medical diagnosis and decision making. For participants who have already conducted cost-effectiveness analyses it will be interesting to see that building and evaluating a PGM is easier, faster and less error-prone than building and debugging an equivalent model using a spreadsheet, a (Markov) decision tree or a programming language, such as R, MATLAB or C++. Participants are invited to bring their own laptops with OpenMarkov installed.

Content

- Outline and schedule (PDF)
- Slides (PDF, 13 MB) [they will be updated a few days before the course]
- Hands-on exercises (PDF)

Attendants must install version 0.3.2 of OpenMarkov.

We recommend printing the outline and the exercices (on paper) and practicing with OpenMarkov's tutorial before attending the course.

Additional information

OVERVIEW

- 1. Introduction
- 2. Probabilistic diagnosis
- 3. Bayesian networks
- 4. Unicriterion decision analysis
- 5. Multicriteria decision analysis
- 6. Sensitivity analysis
- 7. Temporal models
- 8. Conclusion

1. Introduction: history of PGMs

History of PGMs

- Markov chains: Andrey Markov, 1906
- Bayesian networks for genetics: Sewall Wright, 1921
- Markov decision processes (MPDs): Richard Bellman, 1957
- Naïve Bayes method: three independent papers, 1963
- Partially observable MDPs (POMDPs): Karl Åström, 1965
- Influence diagrams: Ronald Howard, James Matheson, 1980, 1984
- Bayesian networks: Judea Pearl, 1982, 1986, 1988
- Dynamic Bayesian networks: Thomas Dean, Keiji Kanazawa, 1989
- Factored MDPs: Craig Boutilier et al., 1995, 2000
- Factored POMDPs: Craig Boutilier, David Poole, 1996
- Decision analysis networks: Javier Díez et al., 2012
- Markov influence diagrams: Javier Díez et al., 2015, 2017

2. Probabilistic diagnosis

2.1. Bayes theorem

Bayes theorem

We knew that

$$P(x|y) = \frac{P(x,y)}{P(y)}$$
 by the definition of $P(x|y)$

$$P(x, y) = P(x) \cdot P(y \mid x)$$
 by the definition of $P(y|x)$

$$P(y) = \sum_{x} P(y|x) \cdot P(x)$$
 by the theorem of total prob.

Combining these results:

$$P(x|y) = \frac{P(x,y)}{P(y)} = \frac{P(x) \cdot P(y|x)}{P(y)} = \frac{P(x) \cdot P(y|x)}{\sum_{x'} P(x') \cdot P(y|x')}$$

• It means that knowing P(x) and P(y|x) we compute P(x|y).

Predictive value of a finding

◆ Positive predictive value: P(+e/+h)

$$P(+e|+h) = \frac{P(+e) \cdot P(+h|+e)}{P(+e) \cdot P(+h|+e) + P(\neg e) \cdot P(+h|\neg e)}$$

$$PPV = \frac{prev \cdot sens}{prev \cdot sens + (1 - prev) \cdot (1 - spec)}$$

◆ Negative predictive value: P(¬e/+h)

$$P(\neg e | \neg h) = \frac{P(\neg e) \cdot P(\neg h | \neg e)}{P(+e) \cdot P(\neg h | + e) + P(\neg e) \cdot P(\neg h | \neg e)}$$

$$NPV = \frac{(1-prev) \cdot spec}{prev \cdot (1-sens) + (1-prev) \cdot spec}$$

Hands-on exercise 1

◆ Example:

- Prevalence of a disease: 14%
- Sensitivity of a test: 70%
- Specificity of the test: 91%

Questions:

- What is the positive predictive value (PPV)?
 - If the test is positive, what is the probability that the patient has the disease?
- What is the <u>negative predictive value</u> (NPV)?
 - If the test is negative, what is the probability that the patient does not have the disease?

Probabilistic diagnosis with two findings

◆ Example:

- Prevalence of the disease: 14%
- Sensitivity of test C: 70%
- Specificity of test C: 91%
- Sensitivity of test E: 90%
- Specificity of test E: 93%

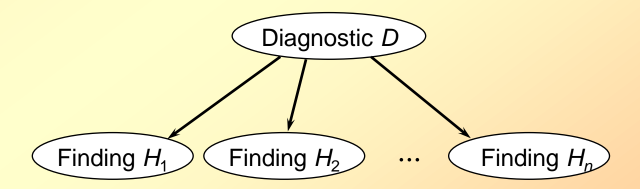
Question:

What is the posterior probability for each combination of findings?

2.2. The naïve Bayes method

The naïve Bayes method

- Two hypotheses:
 - Diagnostics are mutually exclusive
 every patient has at most one disease
 - Findings are conditionally independent given the diagnostics
- Graphical representation:



Succesfull applications of the naïve-Bayes

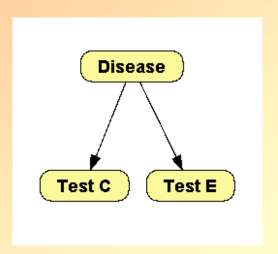
- Lodwick GS, Haun CL, Smith WE, et al., "Computer diagnosis of primary bone tumors: A preliminary report" Radiology 80 (1963) 273-275.
- Overall JE, Williams CM, "Conditional probability program for diagnosis or thyroid function" JAMA 183 (1963) 307-313.
- Toronto AF, Veasy LG, Warner HR, "Evaluation of a computer program for diagnosis
 of congenital heart disease" Progress in Cardiovascular Diseases 5 (1963) 362-377.
 - Warner HR, Toronto AF, Veasy LG, "Experience with Bayes' theorem for computer diagnosis of congenital heart disease" *Annals New York Acad. Sciences* **115** (1964) 558-567.
- de Dombal FT, Leaper JR, Staniland JR, et al., "Computer-aided diagnosis of <u>acute</u> <u>abdominal pain</u>" BMJ 2 (1972) 9-13.
- Gorry GA, Kassirer JP, Essig A, Schwartz WB, "Decision analysis as the basis for computer-aided management of <u>acute renal failure</u>" *Amer. J Med* 55 (1973) 473-484.
- Gorry GA, Silverman H, Pauker SG, "Capturing clinical expertise: A computer program that considers clinical responses to <u>digitalis</u>" Amer. J. Med 64 (1978) 452-460.

More accurate than medical doctors (in restricted domains).

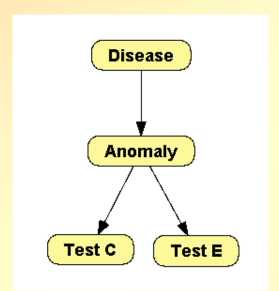
Limitations of the naïve Bayes

- ◆ In general the <u>diagnostics</u> are <u>not mutually exclusive</u>.
- ♦ In general, findings are not conditionally independent.

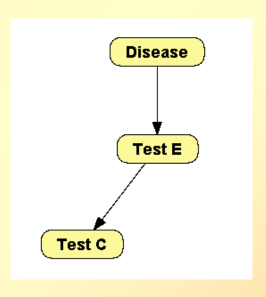
Three solutions for the two-test example



Test results are conditionally independent given the disease



Correlation, even when the disease is known to be present or absent



Test C is conditionally independent of the disease given test E

In the three cases the sensitivity and specificity of the tests (wrt the disease) are the same, but the posterior probabilities are different

Impact of correlation on the posterior prob.

One extreme case:

test results are conditionally independent given the disease

- P(+d/+c, +e) = 0.9421
- maximum increase in the posterior probability
- Opposite extreme case:
 test C is conditionally independent of the disease given test E:
 - P(+d/+c, +e) = P(+d/+e) = 0.6767
 - \triangleright no increase in the posterior probability \equiv no new information
- Intermediate cases: correlation among findings
 - > 0.6767 < P(+d/+c, +e) < 0.9421
 - > the bigger the correlation, the smaller the information contributed.

Prob. diagnosis with two findings (revisited)

◆ Example:

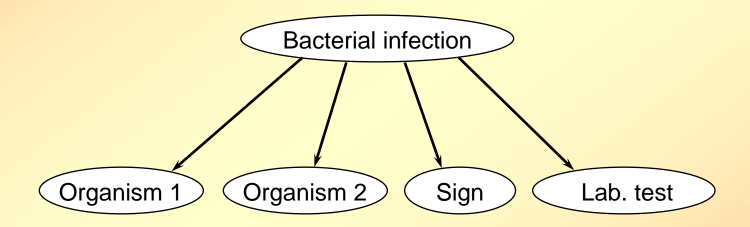
- Prevalence of the disease: 14%
- Sensitivity of test C: 70%
- Specificity of test C: 91%
- Sensitivity of test E: 90%
- Specificity of test E: 93%

Question:

- What is the posterior probability for each combination of findings?
- Answer: The problem is ill-specified
 - The solution depends on the correlation between findings

Limitations of the naïve Bayes

- In general the <u>diagnostics</u> are <u>not mutually exclusive</u>.
- In general, <u>findings</u> are <u>not conditionally independent</u>.



- These limitations are rarely discussed in the books of medical decision analysis and evidence-based medicine.
- This is the only method presented in those books.

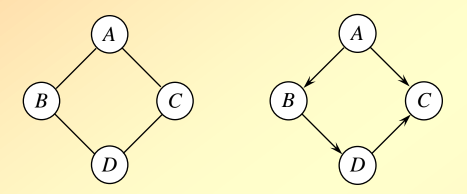
3. Bayesian networks

Probabilistic graphical models

- Elements of a PGM
 - Qualitative component (structure): a graph
 - Links usually represent causal relations
 - Quantitative components (parameters): potentials
 - A conditional probability for each chance node
 - A value function for each value node
- Relation between the graph and the prob. distribution
 - Every node in the graph represents a variable of the prob.
 - > The graph represents the dependencies of the prob. distr.
- Examples of PGMs: BNs, IDs, POMDPs, etc.

3.1. Definition of BN

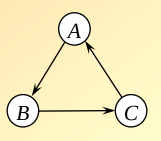
Notions about graphs

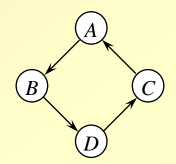


- Basic concepts
 - Definition: a set of nodes and links (vertices and edges)
 - Two types of links: directed / undirected
 - Open path (A-B, A-B-C-D), closed path (A-B-C-D-A),
- In directed graphs:
 - parent, child, ancestor, descendant.

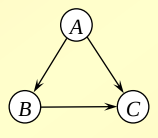
Directed graphs: cycles and loops

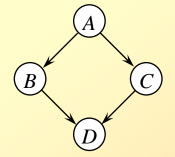
◆ Cycles

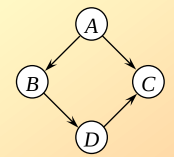




◆ Loops





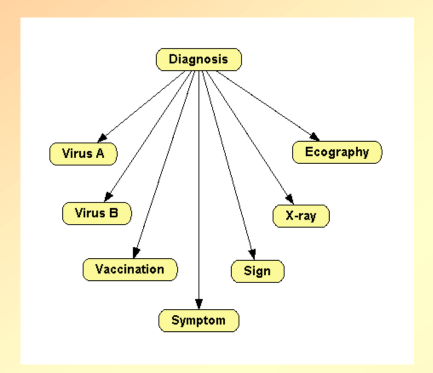


Definition of Bayesian network

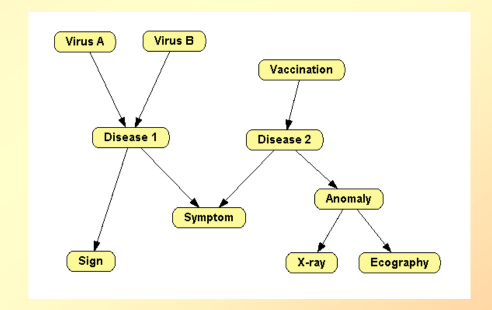
- ◆ Elements:
 - \triangleright a set of variables $\{X_i\}$
 - > an acyclic directed graph
 - every node in the graph represents a variable X_i
 - \triangleright a conditional probability distribution (usually a table) for each variable: $P(x_i | pa(x_i))$
 - for a node without parents: $P(x_i | pa(x_i)) = P(x_i)$
- ◆ Result: join probability for the network

$$P(x_1, \dots, x_n) = \prod_{i=1}^n P(x_i | pa(x_i))$$

Naïve Bayes



Bayesian network



3.2. Examples of BNs

Examples of BNs

- Medical Bayesian networks we have built
 - DIAVAL: echocardiography (valvulopathies)
 F. J. Díez' thesis, 1994
 - Prostanet: urology (prostate cancer)
 Carmen Lacave's thesis, 2003
 - Nasonet: nasopharyngeal cancer spread Severino Galán's thesis, 2003
 - HEPAR II: liver diseases
 Agnieszka Onisko's thesis, 2003
 - Catarnet: Cataract surgery Nuria Alonso's thesis, 2009

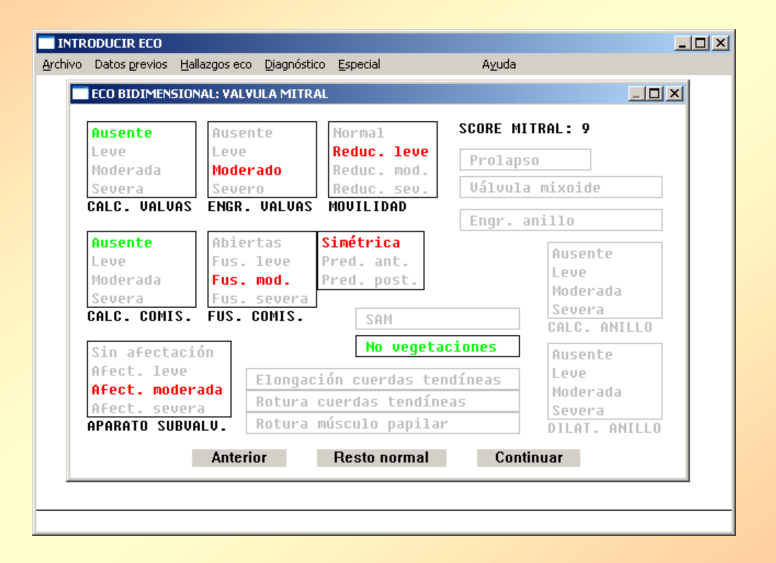
DIAVAL

INTRODUCIR ECO	_UX								
Archivo Datos previos Hallazgos eco Diagnóstico Especial Ayuda									
■ DATOS ADMINISTRATIVOS	×								
Eco número: 104382 Fecha: 29 10 03 Transtorácico: S Cinta: 512 Hora grabación: 1.23.56 Transesofágico: N	- I I								
Nombre: MARIA Apellidos: PEREZ GARCIA									
Sexo: MUJER DNI: 123456 Edad: 51 años Peso: 58 Kg Estatura: 158 cm Sup. corporal: 1.58 m²									
* Solicitante: CARDIOLOGIA Situación: INGRESADO Sector: 3 Cama: 512A									
Continuar									
Introducir los datos del paciente.									

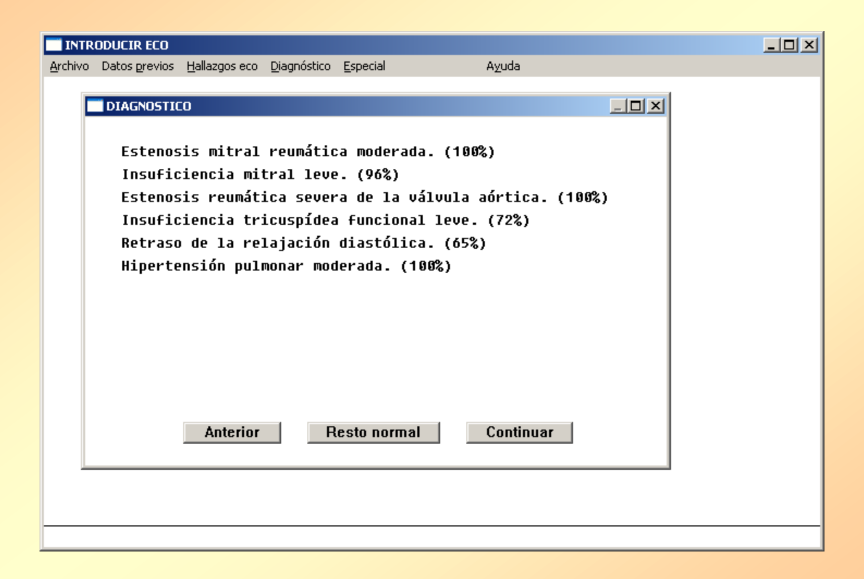
DIAVAL: numeric findings

INTR	ODUCIR ECO								×	
<u>A</u> rchivo	Datos <u>p</u> revios <u>H</u> a	llazgos eco <u>D</u> i	agnóstico	<u>E</u> special		A <u>y</u> uda				
PARAMETROS DEL ECO DOPPLER (M y T)										
	? E			cm/s	"+105%"	"mod.	aumentada"			
	? A	- 544		cm/s						
	? Cociente ? T.R.IV.	9 E/H		ms						
		celeración		MS						
	_			1.1.3						
	_	áx. mitral		mmHg			moderada"			
	? Grad. me	ed. mitral	7.0	mmHg		"lev.	aumentado"			
	? T.H.P. r			MS	-		aumentado"			
	? Area mit	tral (THP)	0.9	CM2	''-76%''	"este	n. crítica"			
	? Vel. máx	k. tric.		cm/s						
	_	íx. tric.		mmHg						
	? Grad. me	ed. tric.		mmHg						
	_				o 1					
			Anterio	or	Continuar					
Pulsar "?" para obtener más información sobre un parámetro.										

DIAVAL: qualitative findings

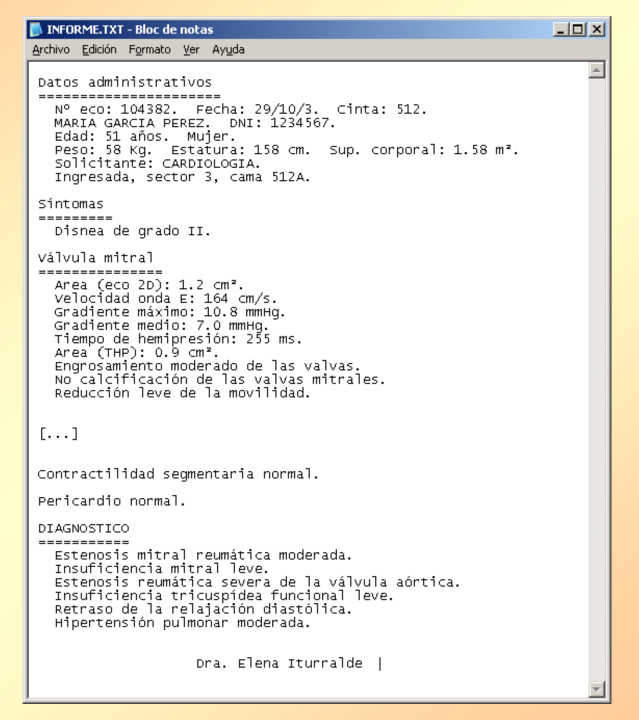


DIAVAL: diagnostics

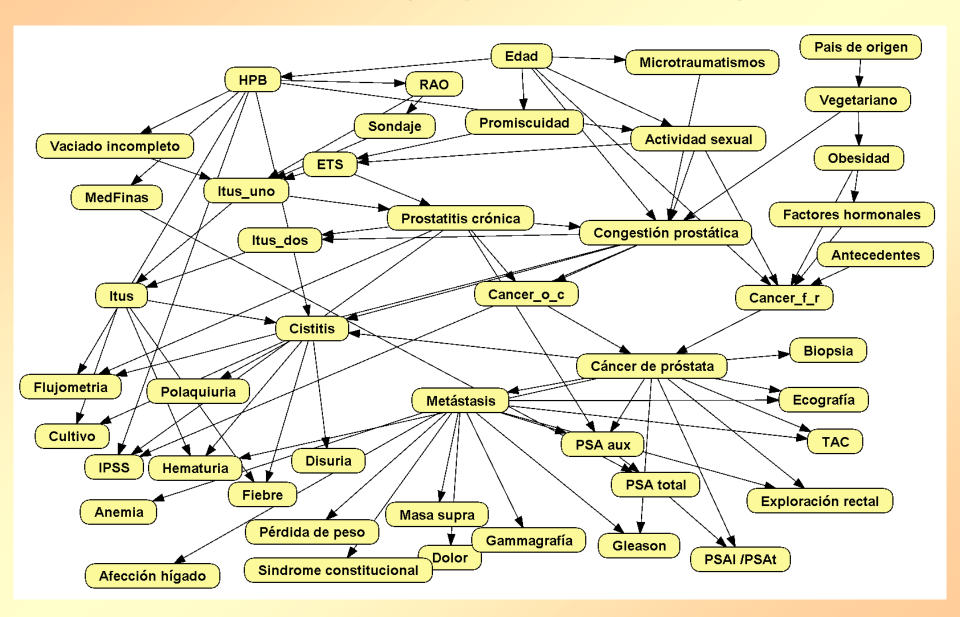


DIAVAL: final report

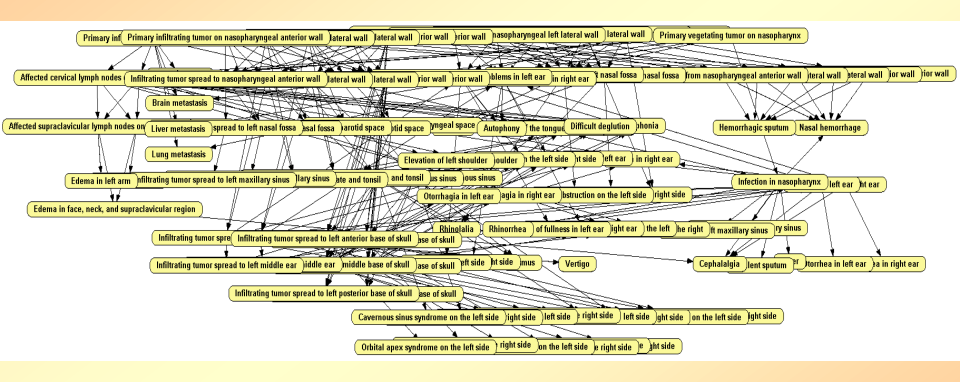
in a text editor



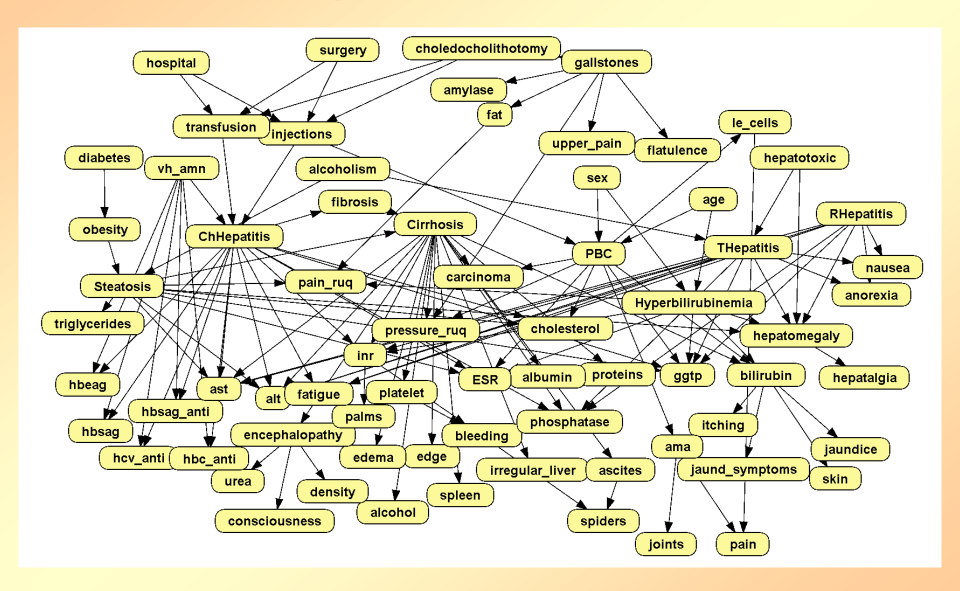
Prostanet (for prostate diseases)



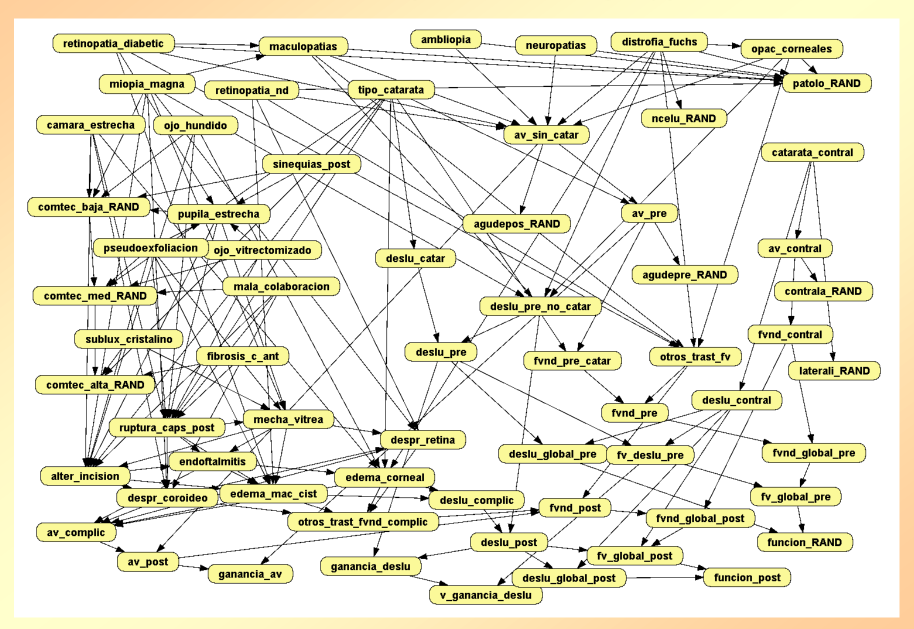
Nasonet (nasopharyngeal cancer spread)



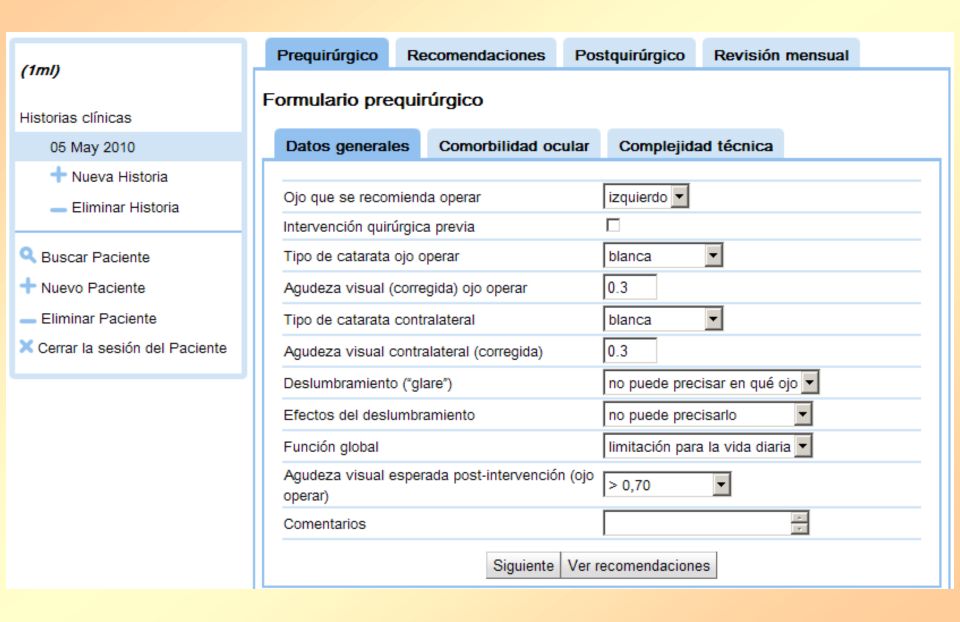
Hepar II (liver diseases)



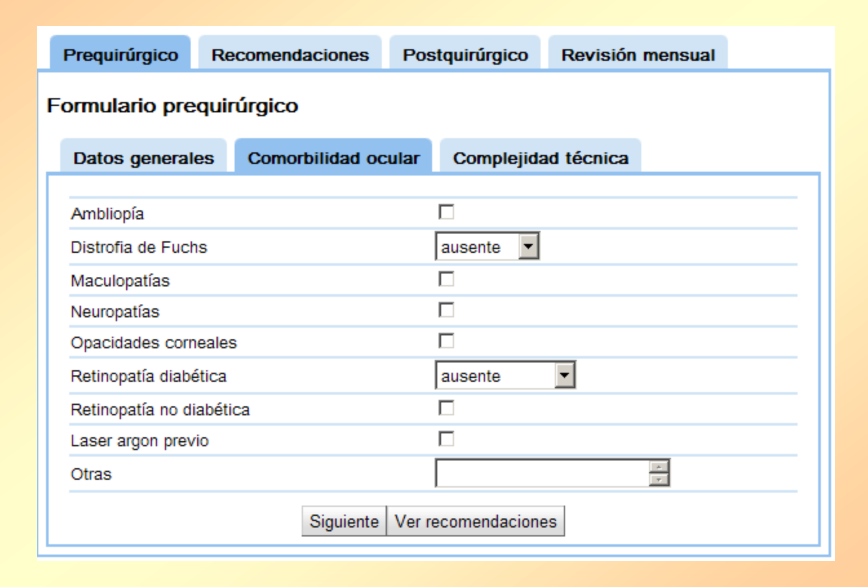
Catarnet (cataract surgery)



Input: 1. General data



Input: 2. Ocular comorbidity



Input: 3. Surgical complexity

Prequirúrgico	Recomendaciones	Postquirúrgico	Revisión mensual			
Formulario prequirúrgico						
Datos general	es Comorbilidad od	cular Complejida	ad técnica			
Cámara estrecha	1					
Fibrosis de la cá	psula anterior					
Mala colaboració	on del paciente (prevista)					
Miopía magna		~				
Ojo hundido	Ojo hundido					
Ojo vitrectomiza	do					
Pseudoexfoliació	ón					
Pupila estrecha Sinequias posteriores						
Subluxación de d	cristalino					
Otras			× v			
	Ver	recomendaciones				

Output: 1. Expert panel's recommendations

Prequirúrgico

Recomendaciones

Postquirúrgico

Revisión mensual

Recomendaciones de SAD-Catar

Panel de expertos

Recomendación: Facoemulsificación apropiada

Mediana de las puntuaciones (1 a 9): 8,5

Grado de acuerdo: Acuerdo

▼ Escenario

Explicación

Variable	Valor
A.V. contralateral	$\geq 0.2 \text{ y} \leq 0.4$
A.V. previa en el ojo a operar	≥ 0,2 y ≤ 0,4
Patología asociada a la catarata	Catarata simple
Lateralidad de la catarata	Bilateral
Complejidad técnica	Moderada por presencia de: " miopía magna (leve) " catarata blanca (moderada)
Función visual	Dificultades en las actividades de la vida diaria

Output: 2. BN recommendation

Red bayesiana CatarNet

Recomendación: 9 (Totalmente recomendada)

Mejoría en A.V. (máx. 6): 5,2

Mejoría en deslumbramiento (máx. 5): 1,7

▼ Probabilidades

Función visual post-intervención	Probabilidad
Sin problemas	0,057
Dificultades para el ocio	0,830
Dificultades para la vida diaria	0,113
AV post-intervención	Probabilidad
≤ 0,15	0,029
> 0,15 y ≤ 0,4	0,088
$> 0.4 \text{ y} \le 0.7$	0,047
> 0,7	0,836
Deslumbramiento post-intervención	Probabilidad
Deslumbramiento	0,544
Complicaciones	Probabilidad
Desprendimiento de coroides	0,001
Desprendimiento de retina	0,080
Edema corneal	0,042
Edema macular cistoide	0,020
Endoftalmitis	0.002

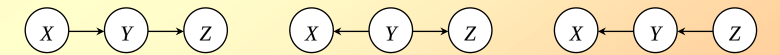
3.3. BNs and causality

Two interpretations of BNs

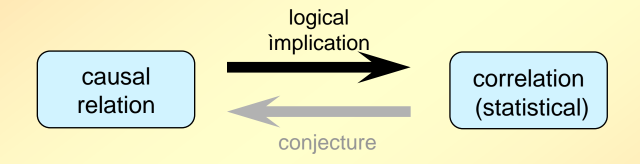
- Semantics of a Bayesian network:
 - As a mathematical model: probabilistic independencies
 - As a model of the real world: they usually represent causality
- Two models are mathematically equivalent when they represent the same set of independencies.
- But two different BNs can never have the same causal meaning.
- Example 1



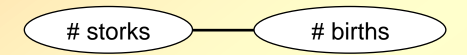
Example 2

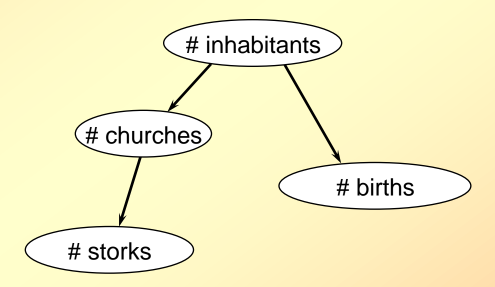


Correlation does not imply causality

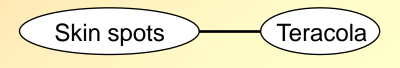


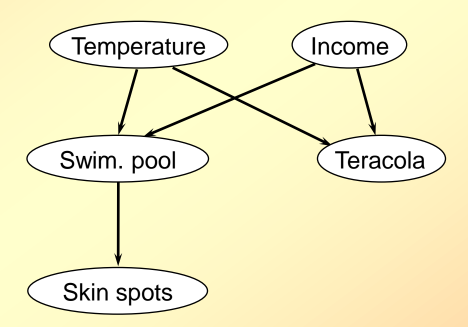
Correlation does not imply causality (example 1)



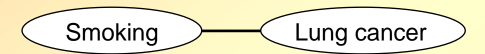


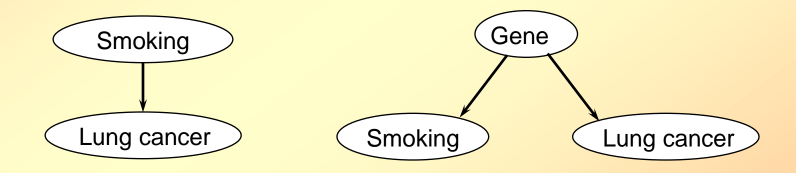
Correlation does not imply causality (example 2)





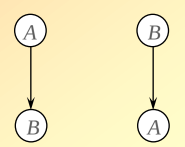
Correlation does not imply causality (example 3)





Several types of correlation

Direct cause

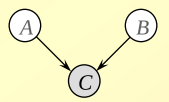


◆ Common cause

C

B

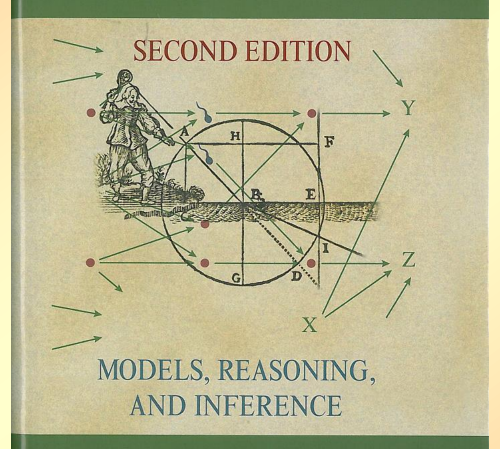
♦ Selection bias



(example: Berkson bias)

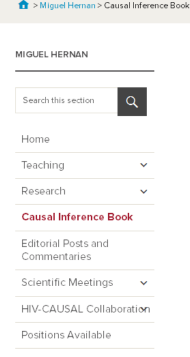
Correlation
without
direct causality

CAUSALITY



JUDEA PEARL

Miguel Hernan



Causal Inference Book

My colleague Jamie Robins and I are working on a book that provides a cohesive presentation of concepts of, and methods for, causal inference. Much of this material is currently scattered across journals in several disciplines or confined to technical articles. We expect that the book will be of interest to anyone interested in causal inference, e.g., epidemiologists, statisticians, psychologists, economists, sociologists, political scientists, computer scientists... The book is divided in 3 parts of increasing difficulty: causal inference without models, causal inference with models, and causal inference from complex longitudinal data.

We are making drafts of selected book sections available on this website. The idea is that interested readers can submit suggestions or criticisms before the book is published. To share any comments, please email me or visit @causalinference on Facebook. To cite the book, please use "Hernán MA, Robins JM (2018). Causal Inference. Boca Raton: Chapman & Hall/CRC, forthcoming."

Follow the links below to access different parts of the book:

- Part I, Chapters 1-10 (updated 4 October 2017)
- Dart II Chantere 11_17 (undated 5 March 2017)

6	Graphical representation of causal effects	69
	6.1 Causal diagrams	69
	6.2 Causal diagrams and marginal independence $\ \ldots \ \ldots \ \ldots$	72
	6.3 Causal diagrams and conditional independence	73
	6.4 Graphs, counterfactuals, and interventions	75
	6.5 A structural classification of bias	77
	6.6 The structure of effect modification	78
7	Confounding	83
	7.1 The structure of confounding	83
	7.2 Confounding and identifiability of causal effects	85
	7.3 Confounders	86
	7.4 Confounding and exchangeability	89
	7.5 How to adjust for confounding	92
8	Selection bias	95
	8.1 The structure of selection bias	95
	8.2 Examples of selection bias	97
	8.3 Selection bias and confounding	99
	8.4 Selection bias and identifiability of causal effects $\dots \dots \dots$	101
	8.5 How to adjust for selection bias	102
	8.6 Selection without bias	106
9	Measurement bias	109
	9.1 Measurement error	109
	9.2 The structure of measurement error	110
	9.3 Mismeasured confounders	111

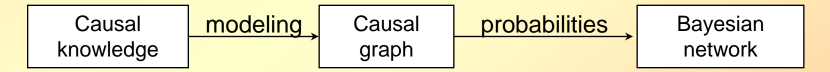
3.4. Building BNs

How to build a Bayesian network

From a database



- There are many algorithms, several new algorithms every year
- Similar to statistical methods (logistic regression, neural nets...)
- With a human expert's help

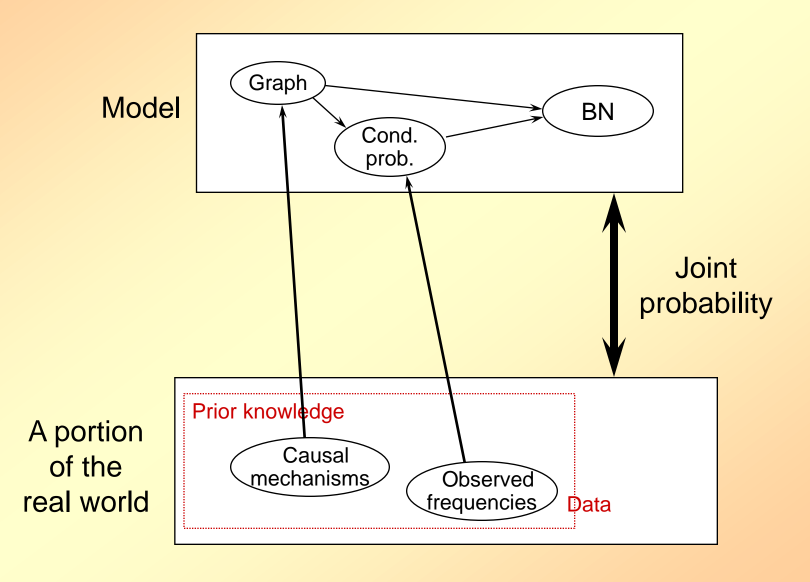


- Hybrid methods:

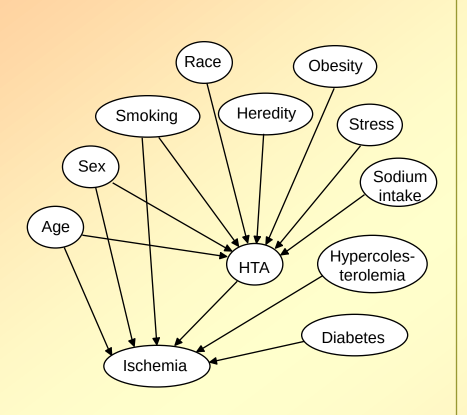
 - Experts → initial model; new cases → refine the probabilities

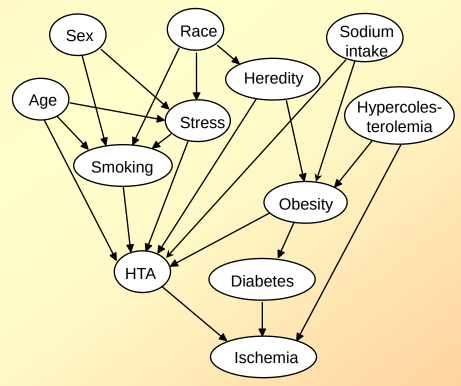
3.4.1. Building BNs with causal knowledge

Building BNs with causal knowledge



PROBLEMS DUE TO LACK OF CAUSAL KNOWLEDGE (1)





Where do the probabilities come from?

Epidemiological studies

- advantage: we obtain directly the parameters we need
- difficulties: time and cost; biases (e.g. selection biases)

Medical literature

- advantages: reliable, inexpensive
- difficulties: few qualitative data, few direct probabilities, different criteria, population-dependent, publication biases

Databases

- advantages: fast, inexpensive
- difficulties: small databases, selection biases

Subjective estimates

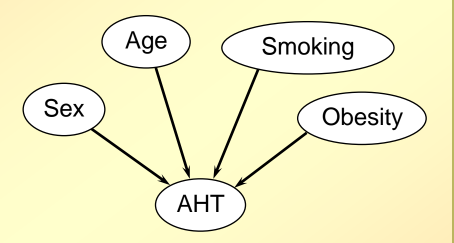
- advantage: relatively inexpensive
- difficulties: unavailability of experts, psychological biases

3.4.1.1. Canonical models

Canonical models

General model

- Probability table:
 - $P(y | x_1, \ldots, x_n)$
- ◆ Factors that influence the prob. of X

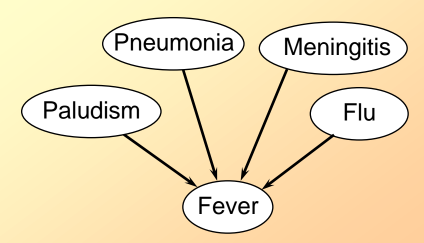


Noisy OR

Efficiency of each link:

 C_i

Causes that can produce X



The noisy OR (hypotheses)

- Each cause, by itself, is able to produce the effect, with a certain probability
 - which is less than 100% when there are inhibitors.
- 2. The effect is absent when no cause has produced it
 - i.e., when every cause is either absent or inhibited
- If a cause has produced the effect, then the effect is present (regardless of the other causes)
- 4. Independence of causal influences
 - ⇒ there is no interaction between the causes (or its inhibitors) when producing the effect
 - ⇒ the probability of the effect is the probability that the first cause has produced it, plus the probability that the second cause produces it when the first has not, plus...

Application of the noisy OR when building BNs

- Advantages of the noisy OR
 - Easier to build, because it requires fewer parameters
 - from a database: more cases to estimate each parameter
 - from a human expert: fewer parameters and more intuitive
 - The computation of probability is more efficient (faster)
 - Possibility of explaining the reasoning: differencial diagnosis (explaining away)
- Two ways to establish the noisy OR
 - From a statistical study
 - Knowing the causal mechanisms

Canonical Probabilistic Models for Knowledge Engineering

Francisco J. Díez

FJDIEZ@DIA.UNED.ES

Dept. Inteligencia Artificial, UNED Juan del Rosal, 16, 28040 Madrid, Spain

Marek J. Druzdzel

MAREK@SIS.PITT.EDU

Decision Systems Laboratory, School of Information Sciences and Intelligent Systems Program University of Pittsburgh, Pittsburgh, PA 15260, USA

Abstract

The hardest task in knowledge engineering for probabilistic graphical models, such as Bayesian networks and influence diagrams, is obtaining their numerical parameters. Models based on acyclic directed graphs and composed of discrete variables, currently most common in practice, require for every variable a number of parameters that is exponential in the number of its parents in the graph, which makes elicitation from experts or learning from databases a daunting task. In this paper, we review the so called canonical models, whose main advantage is that they require much fewer parameters. We propose a general framework for them, based on three categories: deterministic models, ICI models, and simple canonical models. ICI models rely on the concept of independence of causal influence and can be subdivided into noisy and leaky. We then analyze the most common families of canonical models (the OR/MAX, the AND/MIN, and the noisy XOR), generalizing them and offering criteria for applying them in practice. We also briefly review temporal canonical models.

Contents

1	Inti	roduction	3
	1.1	Overview of the paper	4
2	\mathbf{Pre}	liminaries	5
	2.1	Notation	5
	2.2	Systems, models, variables, and probability distributions	6
	2.3	Bayesian networks and influence diagrams	7
	2.4	Causality and network structure	8
3	Ger	neral framework 1	0
	3.1	Deterministic models	0
	3.2	ICI models	2
		3.2.1 Noisy ICI models	2
		3.2.2 Leaky ICI models	4
		3.2.3 Probabilistic ICI models	7
	33	Simple canonical models	Q

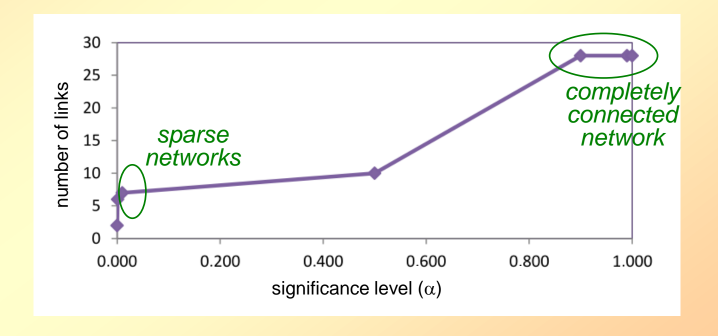
3.4.2. Learning BNs from data

Learning BNs from data

- Two possibilities of learning
 - automatic, interactive
- ◆ Two main algorithms:
 - Search-and-score
 - search
 - depart from a network with no links
 - add/remove/invert a link in each iteration
 - score
 - use a metric (there are several metrics available) to quantify how well the model matches the data
 - > PC
 - depart from a fully-connected undirected graph
 - when two variables are independent, remove the link
 - more precisely, when the correlation is not statistically significant $(p < \alpha)$
 - when two variables are conditionally indep., remove the link
 - orient the remaining links to obtain a directed graph

The role of significance in the PC algorithm

- lacktriangle We set the value of the significance, α
- ♦ For each link, when $p > \alpha$ we assume that the <u>correlation</u> in the database is <u>spurious</u> (i.e., due randomness) and <u>remove</u> the link
- ♦ Low value of $\alpha \Rightarrow$ removing many links \Rightarrow sparse network
- lacktriangle High value of $\alpha \Rightarrow$ keeping many links \Rightarrow dense network



Advantages of interactive learning

- ◆ The system proposes, the user decides
 - Very useful for tuition
 - Useful for combining data with expert knowledge
 - Useful for debugging new algorithms (workbench)
- See <u>www.openmarkov.org/docs/tutorial</u>.

A comparison of both methods for building BNs

- Automatic learning from databases
 - Advantage: faster (graph + probabilities)
 - Limitation: medical databases are usually incomplete
 - Missing values → problem of imputation (rarely missing at random)
 - Missing variables → spurious correlations
 - Blackbox algorithm that returns non-causal models
 - ⇒ Human experts are reluctant to accept their advice
- With expert knowledge ("manual" method)
 - Only method possible when there is not a good-enough database
 - Difficulty in practice: getting the collaboration of experts
 - Building the structure of the causal is sometimes difficult
 - Obtaining the probabilities is even more difficult.

Summary: BNs vs. the naïve Bayes

- BNs can diagnose <u>several diseases</u> simultaneously.
- BNs do not assume conditional independence of findings.
- ♦ BNs are usually <u>causal</u> models
 - closer to doctors' reasoning: explanation of reasoning
 - probabilities are in general easier to obtain
- ◆ Three types of <u>reasoning</u>: abductive, deductive, inter-causal.
- They can combine <u>data</u> (from databases), <u>epidemiological studies</u> (scientific literature) and <u>expert knowledge</u> (doctors).

In spite of these advantages, BNs are almost unknown <u>in medicine</u>. No book for medical doctors mentions them!

4. Unicriterion decision analysis

4.1. Introductory examples

Medical example (1)

◆ Three variables

➤ Chance variable: $X \rightarrow$ bacterial infection; P(+x) = 0.14

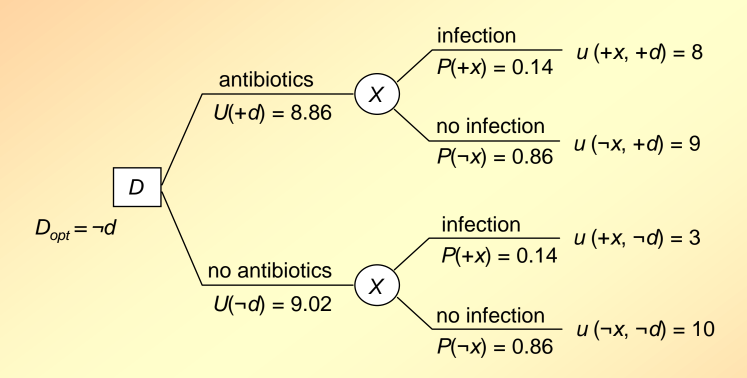
ightharpoonup Decision: D o give antibiotics

ightharpoonup Utility (value): $U \rightarrow$ effectiveness

<i>u</i> (<i>x</i> , <i>d</i>)	+x	$\neg x$
+d	8	9
$\neg d$	3	10

- When making the decision we do not know whether the patient is infected with the bacteria.
- Question: Should we give antibiotics?

Decision tree (1)



Optimal decision: $D_{opt} = \neg d \Rightarrow \text{do not give antibiotics}$

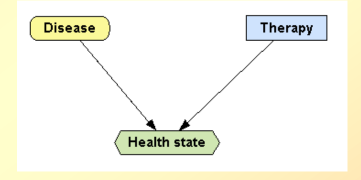
Prognosis: $U = \max (U(+d), U(\neg d)) = \max (8.86, 9.02) = 9.02$

Influence diagram

Disease Therapy

Health state

DAN (decision analysis network)



- Both models are identical.
- ◆ They generate the same decision tree.

Utility as a function of prevalence

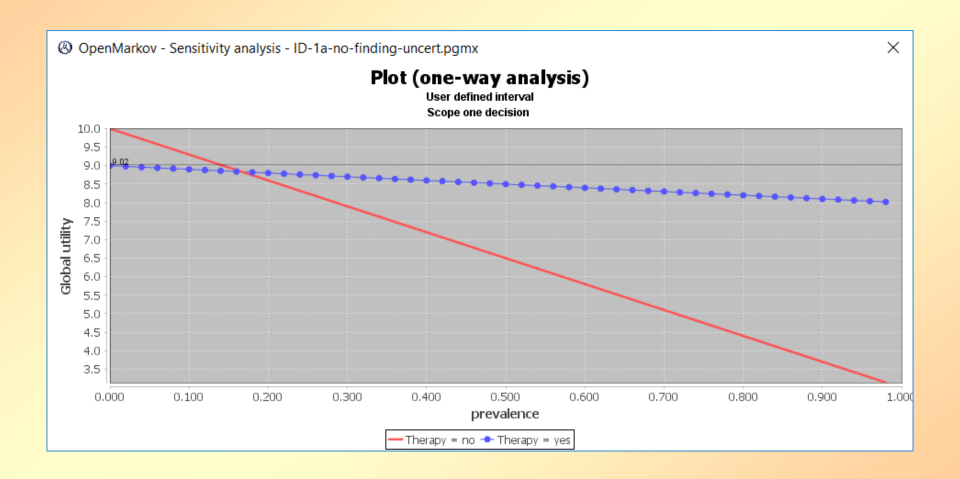
$$U(d) = \sum_{x} u(x,d) \cdot P(x)$$

$$D_{opt} = \arg \max(U(+d), U(\neg d))$$

$$U = \max(U(+d), U(\neg d))$$

P(+x)	U(+d)	$U(\neg d)$	D_{opt}	U	
0'00	9'00	10'00	$\neg d$	10'00	
0'05	8'95	9'65	$\neg d$	9'79	
0'14	8'86	9'02	$\neg d$	9'02	decision
0'17	8'83	8'81	+d	8'83	threshold
0'40	8'60	7'20	+d	8'60	
0'75	8'25	4'75	+d	8'25	
1'00	8'00	3'00	+d	8'00	

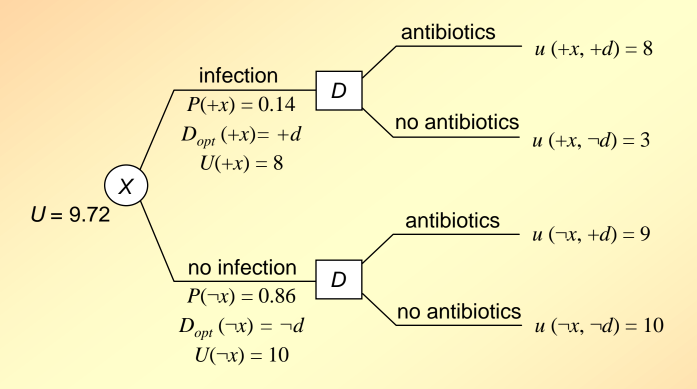
Utility as a function of prevalence



Medical example (2)

- In the previous scenario, what should we do if we knew with certainty whether the patient has the disease?
 - Question 1: What to do when infection is present?
 - Question 2: What to do when infection is absent?
- What is the average utility in this sub-population?

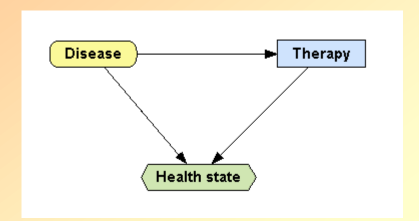
Decision tree (2)



Optimal decision: infection (+x) \Rightarrow give antibiotics (+d) no infection $(\neg x)$ \Rightarrow do not give antibiotics $(\neg d)$

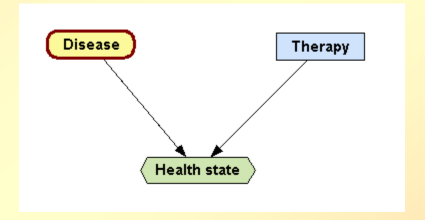
Expected utility: $U = 8 \times 0.14 + 10 \times 0.86 = 9.72$

Influence diagram



We have added an information link.

DAN



We have marked *Disease* as always-observed.

- ◆ Two different ways of saying that the value of *Disease* is known when making the decision *Therapy*.
- Both models are equivalent: they generate the same decision tree.

Medical example (3): The value of information

◆ Test Y for detecting X

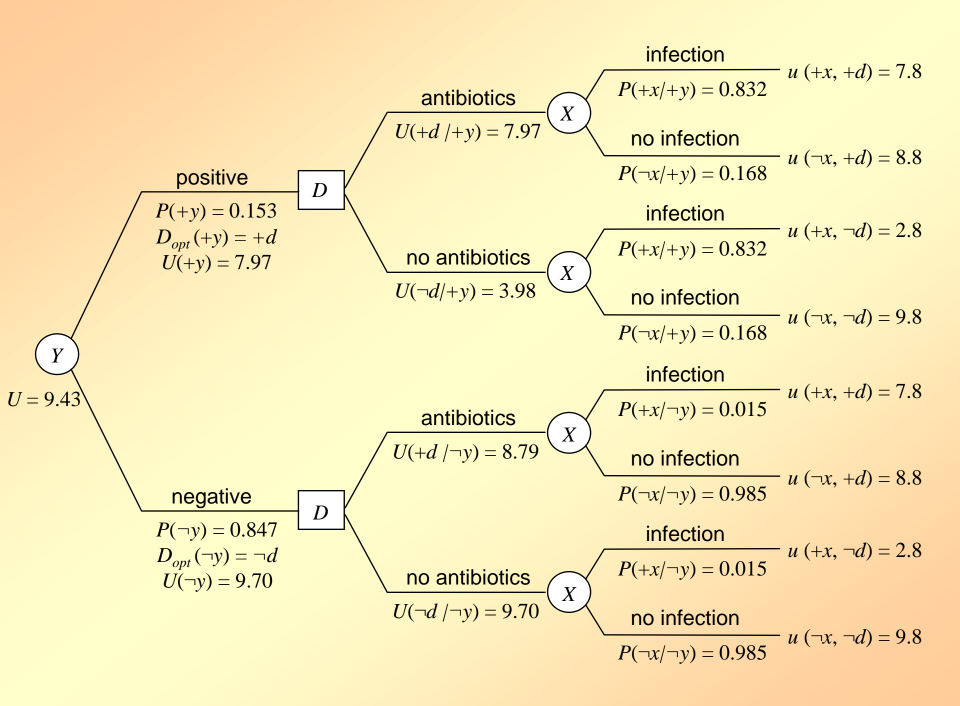
> sensitivity: P(+y/+x) = 0.91

> specificity: $P(\neg y/\neg x) = 0.97$

 ν cost: $u_{\text{test}}(x, d) = u_{\text{no test}}(x, d) - 0.2$

<i>u</i> (<i>x</i> , <i>d</i>)	+x	$\neg \chi$
+d	7'8	8'8
$\neg d$	2'8	9'8

- When making the decision we do know the result of the test.
- Question: Should we give antibiotics?



Policy and prognosis

Policy:

- When Y is positive: give antibiotics
- When Y is negative: do not give antibiotics

◆ Prognosis

When Y is positive: U(+y) = 7.97

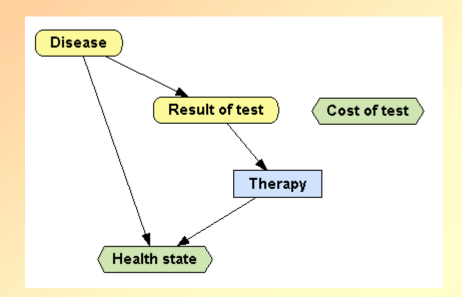
When Y is negative: $U(\neg y) = 9.70$

Global prognosis (average utility)

$$U_{\text{with test}} = U(+y) \times P(+y) + U(\neg y) \times P(\neg y)$$

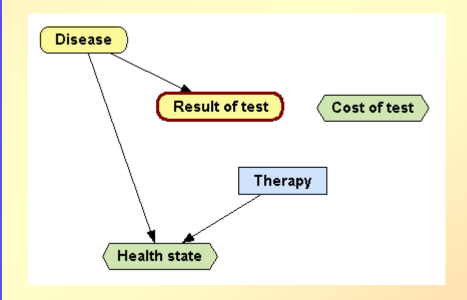
= 7.97 × 0.153 + 9.69 × 0.847
= 9.43

Influence diagram



An **information link** from *Result of test* to *Therapy*

DAN

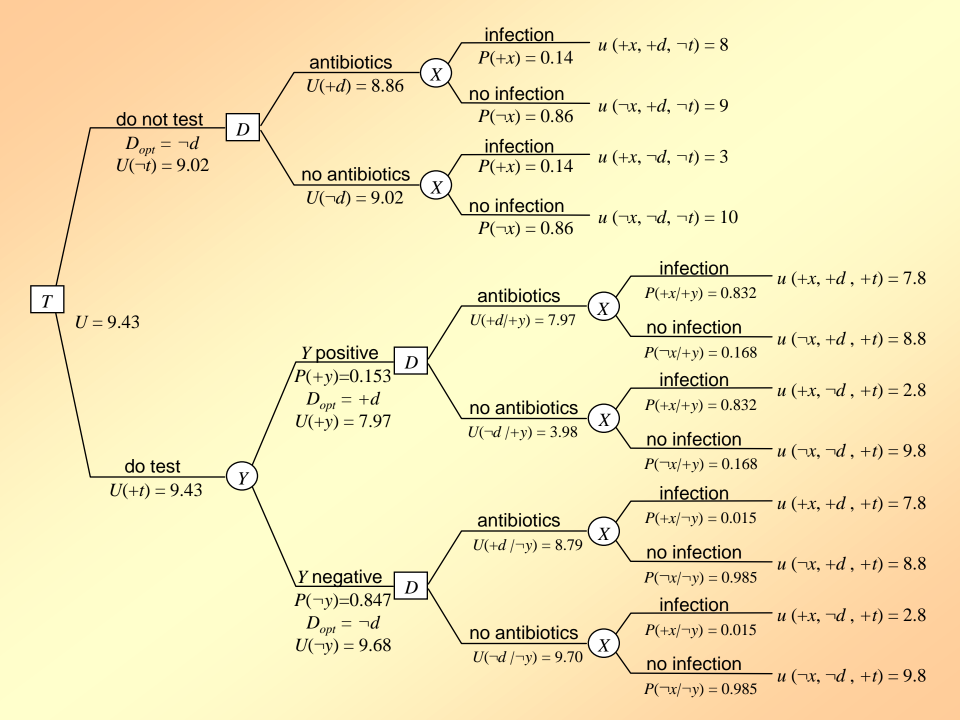


Result of test is marked as always-observed.

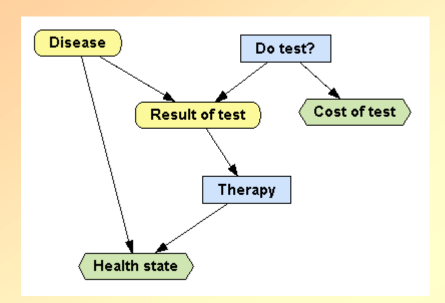
- Different ways of indicating the flow of information.
- Both models generate the same decision tree.

Medical example (4): deciding about a test

- Test Y
 - Advantage: gives information
 - Disadvantage: has a cost
- Is it worth doing the test?
- ◆ Three possible policies:
 - 1. Give the therapy to all patients, preventively
 - 2. Never apply the therapy
 - 3. Do test *Y*; apply the therapy only when it is positive



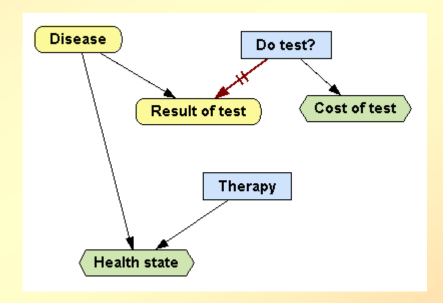
Influence diagram



An **information link**.

Total ordering of the decisions

DAN



Restrictions. Revelation link.
The decisions are not ordered.

- Different ways of indicating the flow of information.
- ◆ The decision trees are different but <u>equivalent</u>: the same probabilities, utilities, and policies.

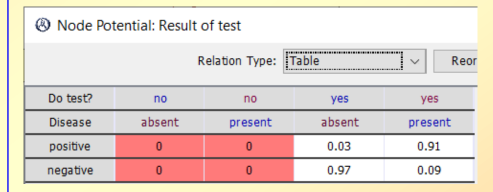
Conditional prob. for Result of test

in the ID

Node Potential: Result of test				
Relation Type: Table V				
Do test?	no	no	yes	yes
Disease	absent	present	absent	present
positive	0	0	0.03	0.91
negative	0	0	0.97	0.09
not done	1	1	0	0

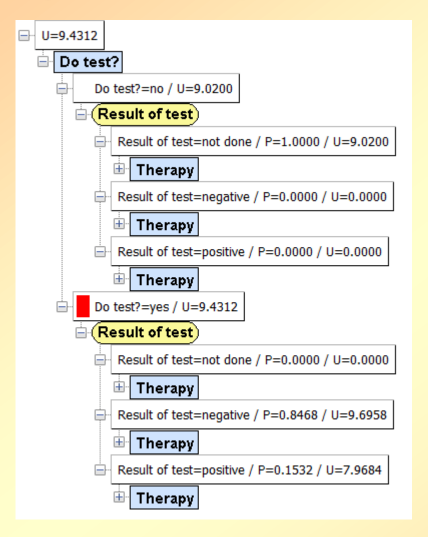
 dummy value: test not done

in the DAN

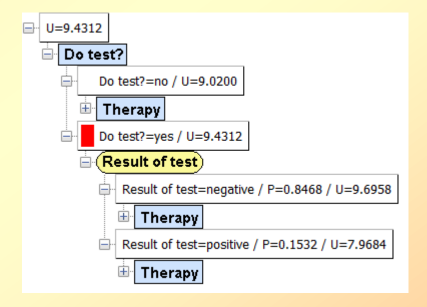


- restrictions
- no dummy value

Decision tree generated by the ID



Decision tree generated by the DAN



symmetric

asymmetric

Hands-on exercise 3

Exercise: Optimal stratety for two tests

Test	sensitivity	specificity	discomfort
Α	0.60	0.92	0.0003 QALY
В	0.80	0.91	0.0001 QALY

Disease →	absent	present
therapy	38 QALY	30 QALY
no therapy	40 QALY	20 QALY

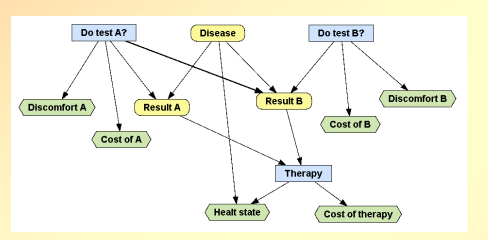
Question: What is the most effective strategy?

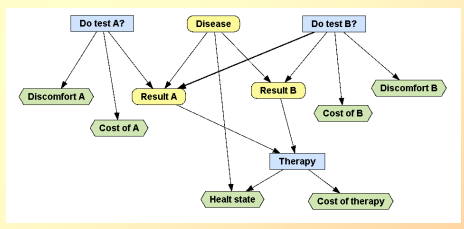
Medical example (5): optimal order of two tests

- Two tests
 - Different sensitivity, specificity and cost
- Many different strategies:
 - 1. No test, no therapy
 - 2. No test, always therapy
 - 3. Test 1; if positive, therapy; if negative, no therapy
 - 4. Test 2; if positive, therapy; if negative, no therapy
 - 5. Test 1; if positive, (test 2; if positive, therapy; if negative...); if...
 - 6. Test 2; if positive, (test 1; if positive, therapy; if negative...); if...
 - 7. etc.

Solution with influence diagrams

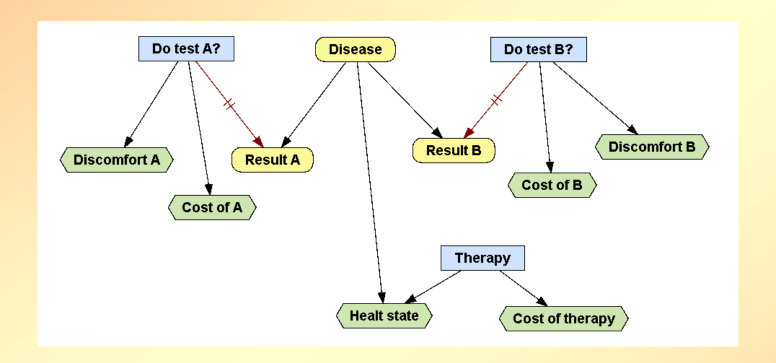
- ◆ IDs require a total ordering of the decisions
- It is not possible to represent this problem with one ID
- ◆ Trick: use two influence diagrams





- ◆ We choose the order (the ID) with the higher expected utility.
- This trick does not work for more than two tests.

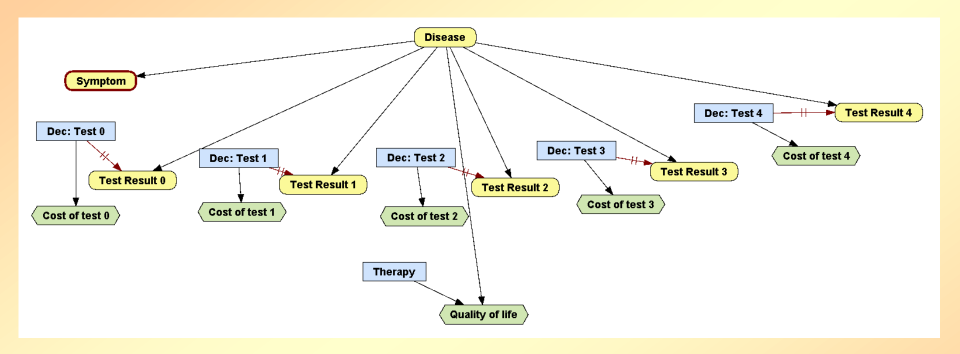
Solution with a DAN



Advantages wrt IDs

- > one network is sufficient
- no dummy states, such as "test not done"
- can accommodate any number of tests

The *n*-test problem



- Computationally hard: n! possible orderings of the tests.
- We have developed an any-space algorithm for this problem
- and a fast algorithm (9 minutes for the 7-test problem).
- We are developing more efficient algorithms.



Contents lists available at ScienceDirect

International Journal of Approximate Reasoning



www.elsevier.com/locate/ijar

Decision analysis networks







ARTICLE INFO

Article history:
Received 7 July 2017
Received in revised form 15 December 2017
Accepted 21 February 2018
Available online 27 February 2018

Keywords:
Decision analysis
Decision trees
Influence diagrams
Probabilistic graphical models
Asymmetric decision problems

ABSTRACT

This paper presents decision analysis networks (DANs) as a new type of probabilistic graphical model. Like influence diagrams (IDs), DANs are much more compact and easier to build than decision trees and can represent conditional independencies. In fact, for every ID there is an equivalent symmetric DAN, but DANs can also represent asymmetric problems involving partial orderings of the decisions (order asymmetry), restrictions between the values of the variables (domain asymmetry), and conditional observability (information asymmetry). Symmetric DANs can be evaluated with the same algorithms as IDs. Every asymmetric DAN can be evaluated by converting it into an equivalent decision tree or, much more efficiently, by decomposing it into a tree of symmetric DANs. Given that DANs can solve symmetric problems as easily and as efficiently as IDs, and are more appropriate for asymmetric problems—which include virtually all real-world problems—DANs might replace IDs as the standard type of probabilistic graphical model for decision support and decision analysis. We also argue that DANs compare favorably with other formalisms proposed for asymmetric decision problems. In practice, DANs can be built and evaluated with OpenMarkov, a Java open-source package for probabilistic graphical models.

© 2018 Elsevier Inc. All rights reserved.

1. Introduction

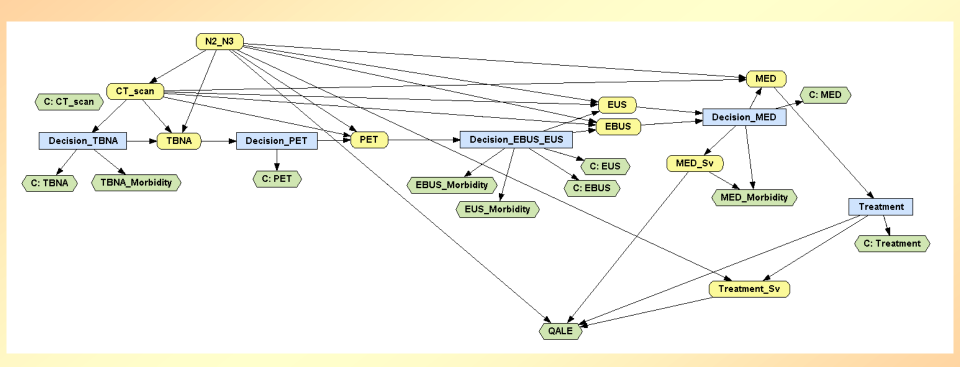
The two formalisms most widely used for the representation and analysis of decision problems are decision trees (DTs) [31] and influence diagrams (IDs) [15]. DTs have the advantage of almost absolute flexibility, but also have three drawbacks: their size grows exponentially with the number of variables, they cannot represent conditional independencies, and they

DANs vs. IDs

- ◆ DANs can replace IDs as the standard decision analysis tool (in AI, MDM, operations research...) because:
 - For every ID there is an equivalent symmetric DAN
 - but for many DANs there is no equivalent ID
 - Virtually all real-world problems are asymmetric.
 - There many problems that cannot be modeled with IDs.
 - Even if a problem can be modeled with an ID, a DAN is usually better because it does not need dummy states.

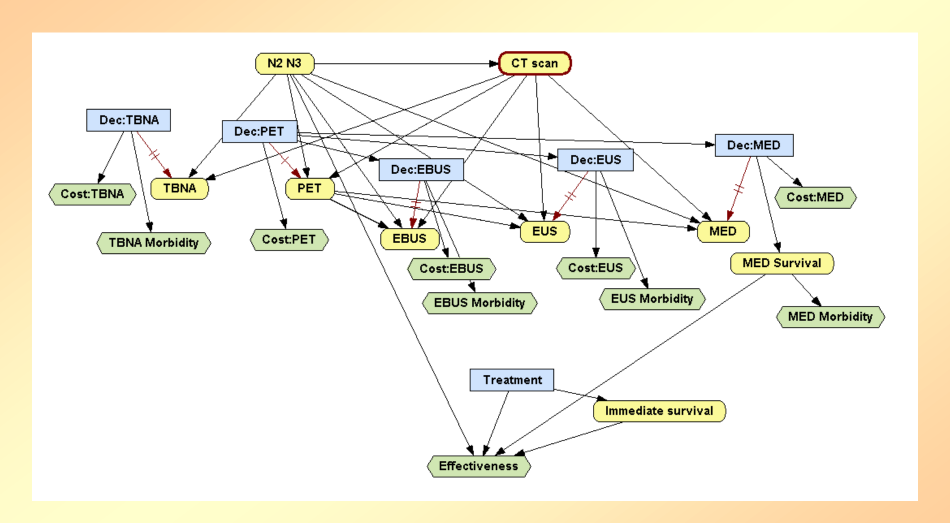
4.2. Examples of decision models for real-world problems

Mediastinet, an ID for lung cancer



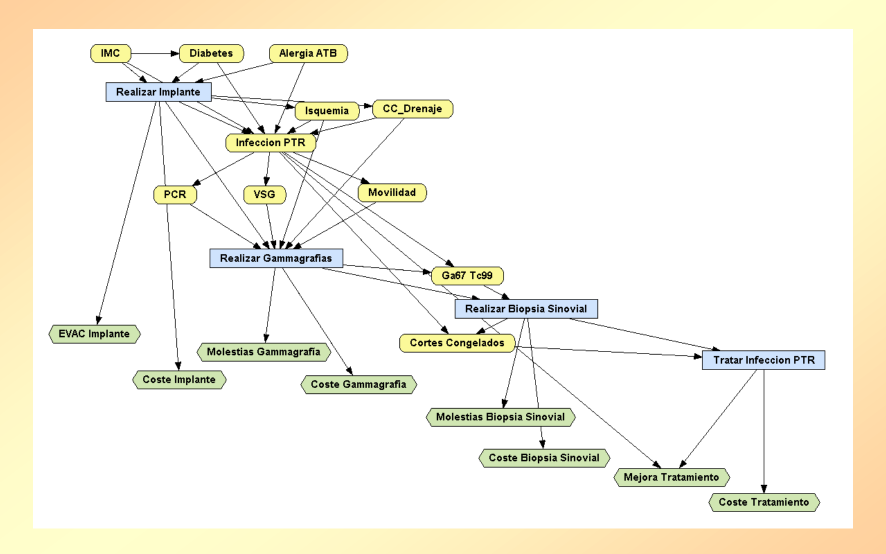
Equivalent to a decision tree containing ~10⁴ branches.

Mediastinet (DAN version)



Decisions are partially ordered.

Arthronet, an ID for total knee arthroplasty



Equivalent to a decision tree containing ~10⁴ branches.

4.3. Advantages and limitations of influence diagrams

Advantages of influence diagrams (1/3)

- ◆ IDs are more <u>compact</u> than decision trees
 - > An ID having *n* binary nodes ~ a DT having 2ⁿ branches
- IDs transform automatically into decision trees
 - but the reverse is not true (no general algorithm)
 - If you build a decision tree, you only have a decision tree.
 - If you build an ID, you have both.
- ◆ IDs are <u>much easier to build</u> than decision trees
 - ➤ IDs use direct probabilities (prevalence, sensitivity, specificity...) and costs (mortality, morbidity, economic cost...)
 - ➤ ID can use canonical models (noisy OR, noisy AND, etc.)
 - Each parameter appears only once in the ID
 - in many cases it is not necessary to have parametric variables
 - > IDs can have several value nodes: more clarity, separate criteria

Advantages of influence diagrams (2/3)

- ◆ No external pre-calculation of probabilities is required
- Having all the information, no debugging is usually needed
 - ➤ On the contrary, "all trees have bugs" (Primer on MDA, at MDM journal)
- ◆ IDs are much easier to modify than decision trees
 - Refine the model with new decisions and chance variables
 - Structural sensitivity analysis
 - Can adapt to different regional settings
 - Can adapt to patient's medical characteristics and preferences
- Explicit representation of <u>causality</u>
 - > a link indicates causal influence
 - ▶ the absence of a link means "no causal influence" (hypothesis)

Advantages of influence diagrams (3/3)

- ◆ Two possibilities of evaluation:
 - 1. expansion of an equivalent decision tree
 - exponential complexity (time and space)
 - equivalent to the brute-force method for Bayesian networks
 - many problems can <u>not</u> be solved with this method
 - 2. operations on the ID (recursive reduction of the ID)
 - direct manipulation of the graph and/or potentials of the ID
 - similar to the best algorithms for Bayesian networks
 - canonical models and the separation of utility nodes can lead to more efficient evaluations
- More possibilities of explanation of reasoning
 - computation of posterior probabilities on the ID (as if it were a BN)
 - > value of information (EVPI and other measures) can be computed easily
 - other methods from Bayesian networks and qualitative prob. networks.

These methods can be used to debug/refine IDs.

Decision Analysis

Vol. 2, No. 4, December 2005, pp. 229–231 ISSN 1545-8490 | EISSN 1545-8504 | 05 | 0204 | 0229



DOI 10.1287/deca.1050.0054 © 2005 INFORMS

The Influence of Influence Diagrams on Artificial Intelligence

Craig Boutilier

Department of Computer Science, University of Toronto, Toronto, Ontario, M5S 3G5 Canada, cebly@cs.toronto.edu

Howard and Matheson's article "Influence Diagrams" has had a substantial impact on research in artificial intelligence (AI). In this perspective, I briefly discuss the importance of influence diagrams as a model for decision making under uncertainty in the AI research community; but I also identify some of the less direct, but no less important, influences this work has had on the field.

Key words: influence diagrams; decision theory; artificial intelligence; value of information; graphical models; perspective, the focus on graphical modeling research

History: Received on November 14, 2005. Accepted by Eric Horvitz on November 23, 2005, without revision.

Howard and Matheson's (1984/2005) "Influence Diagrams" has had a profound impact on developments in artificial intelligence. Some of these influences have been quite direct; others are more indirect, but in many ways, more substantial. The paper itself in representative of developments that had been

vision (Binford and Levitt 2003), dialog management, user interface design, multiagent systems, and game theory (Koller and Milch 2003), to name but a few.

Another reasonably direct impact of "Influence Diagrams" derives from its role in the development

Decision Analysis

Vol. 2, No. 4, December 2005, pp. 238–244 ISSN 1545-8490 | EISSN 1545-8504 | 05 | 0204 | 0238



DOI 10.1287/deca.1060.0060 © 2005 INFORMS

The Influence of Influence Diagrams in Medicine

Stephen G. Pauker, John B. Wong

Division of Clinical Decision Making, Informatics and Telemedicine, Department of Medicine, Tufts–New England Medical Center, Tufts University School of Medicine, 750 Washington St., NEMC 302, Boston, Massachusetts 02111 {spauker@tufts-nemc.org, jwong@tufts-nemc.org}

A lthough influence diagrams have used medical examples almost from their inception, that graphical representation of decision problems has disseminated surprisingly slowly in the medical literature and among clinicians performing decision analyses. Clinicians appear to prefer decision trees as their primary modeling metaphor. This perspective examines the use of influence diagrams in medicine and offers explanations and suggestions for accelerating their dissemination.

Key words: decision analysis; influence diagrams; clinical decision making; medicine *History*: Received December 12, 2005. Accepted by Eric Horvitz on January 5, 2006, after 1 revision.

Introduction

Two decades after Howard's landmark paper (Howard and Matheson 1984/2005) that introduced the concept of the influence diagram and three decades since Miller's initial report (Miller et al. 1976), *Decision Analysis* reproduced that paper in 2005 and solicited a set of commentaries. This paper

modeling paradigm slowly spread from Stanford, both with courses offered at meetings of the Society for Medical Decision Making (Society for Medical Decision Making 2005) and with the development of software that could conveniently capture and evaluate such models.

Clinical practice guidelines (CPGs)

- Construction of CPGs
 - Usually: expert opinion or consensus of experts
 - Another possibility: probabilistic graphical models
 - Sanders, Nease, Owens: several papers on building CPGs from IDs.
- Advantages of a PGM wrt a traditional CPG
 - explicit decision model
 - combines expert opinions and evidence (statistical data)
 - helps in difficult cases, in which the policy is not evident for experts
 - flexibility: can be extended and adapted, as mentioned above
 - can include patients' preferences
 - the physician plays an active role, he/she is not a passive user of CPGs developed by others.

A proverb

- Don't give a man a <u>fish</u>;
 give him a <u>rod</u>
 and teach him how to fish.
- Don't give a doctor a written <u>CPG</u>; give him/her a <u>DAN</u> and teach him/her how to use OpenMarkov.

IDs in the literature on MDM (1/3)

- Books that mention decision trees but do <u>not</u> mention IDs
 - Weinstein, Fineberg. Clinical Decision Making. 1980.
 - Sloan (ed.). Valuing Health Care. 1995.
 - Gold et al. Cost-Effectiveness in Health and Medicine. 1996.
 - Sackett et al. Evidence-Based Medicine. 1997 (and three other books on EBM).
 - Petitti. Meta-Analysis, Decision Analysis and CEA. 2nd ed., 2000.
 - Drummond, McGuire (eds.). Economic Eval. in Health Care Programs. 2001.
 - Levin and McEwan. Cost-Effectiveness Analysis. 2nd ed., 2001.
 - Parmigiani. Modelling in Medical Decision Making. 2002.
 - Haddix et al. Prevention Effectiveness. 2nd ed., 2003.
 - Fox-Rushby and Cairns. Economic Evaluation. 2005.
 - Briggs et al. Decision Modelling for Health Economic Evaluation, 2006.
 - Alemi and Gustafson. Decision Analysis for Healthcare Managers, 2006.
 - Arnold. Pharmacoeconomics: From Theory to Practice. 2009.
 - Kassirer et al. Learning Clinical Reasoning. 2nd ed., 2010.
 - Mushlin and Greene. Decision Making in Medicine. 3rd ed., 2010.

(cont'd)

IDs in the literature on MDM (2/3)

- Books that mention decision trees but do <u>not</u> mention IDs (cont.)
 - Gray et al. Applied Methods of CEA in Health Care, 2011.
 - Alfaro-LeFevre. Critical Thinking, Clinical Reasoning... 5th ed., 2013.
 - Morris et al. Economic Analysis in Healthcare. 2nd ed., 2012.
 - Rascati. Essentials of Pharmacoeconomics. 2nd ed., 2013.
 - Sox et al. Medical Decision Making. Latest ed., 2013.
 - Hunink et al. Decision Making in Health and Medicine. 2nd ed., 2014.
 - Drummond et al. Methods for the Economic Evaluation of... 4th ed. 2015.
 - Edlin et al. Cost Effectiveness Modelling for HTA... 2015.
 - Neumann et al. Cost-Effectiveness in Health and Medicine. 2016
 - Caro et al. Discrete Event Simulation for HTA. 2016

One book that mentioned IDs

 Muennig. Designing and Conducting Cost-Effectiveness Analyses in Medicine and Health Care. 2002, page 242:

"An influence diagram (also known as a tornado diagram) ..."

The 2nd edition (2007) and the 3rd (2016) do not mention them.

IDs in the literature on MDM (3/3)

- Three books that <u>describe</u> IDs
 - Chapman and Sonnenberg (eds.). Decision Making in Health Care. 2000 (5 pages out of 421, in a chapter authored by Mark Roberts)
 - Schwartz and Bergus. Medical Decision Making. A Physician's Guide. 2008.
 (2 pages out of 230)
 - Kattan. Encyclopedia of Medical Decision Making. 2009 (4 pages out of 1200+).
- Summary of the informal survey of books on MDM and EBM
 - 26 books published after 1984
 - All of them explain DTs but only 3 describe IDs, very briefly.
- Some books on medical informatics mention IDs:
 - Shortliffe and Cimino. Biomedical Informatics. 4th ed., 2013 (2.5 pages out of 991).
 - Kalet. Principles of Biomedical Informatics. 2nd ed., 2013 (3 pages out of 708).
- Why are IDs so little known in health sciences after 35+ years?

Limitations of IDs

- 1. The "reasoning" of an ID is not easy to understand
- 2. The evaluation returns large policy tables
- 3. IDs can only model symmetric problems
 - IDs require a total ordering of the decisions
 - IDs cannot represent incompatibilities between values
 - Non-standard versions of IDs partially solve this problem, but none of the alternatives was completely satisfactory.
- 4. Algorithms could only evaluate unicriterion IDs
 - They could not perform cost-effectiveness analysis
- 5. Temporal reasoning was not possible with IDs
 - Dynamic IDs are computationally unfeasible.

Solutions we have proposed

- 1. Explanation in influence diagrams
 - showing the posterior probabilities and expected values
 - introduction of evidence
 - hypothetical reasoning (what if) by means of imposed policies
- 2. Synthesizing the optimal intervention
 - in the form of a compact tree
- 3. Decision analysis networks (DANs)
 - an alternative to IDs for asymmetric decision problems.
- 4. Cost-effectiveness analysis with IDs
- 5. Markov influence diagrams
 - including cost-effectiveness analysis

5. Multicriteria decision making

5.1. Analysis with two criteria: cost and effectiveness

An example with costs and effectiveness

- Two therapies
 - Effectiveness (QALY)

	No therapy	Therapy 1	Therapy 2
Disease present	1.2	4.0	6.5
Disease absent	10	9.9	9.3

- ➤ Therapy 1 cost = 20,000 €
- ➤ Therapy 2 cost = 70,000 €

Questions:

- What therapy to apply when the disease is present
- What therapy to apply when the disease is absent
- Problem: how to compare health and money

5.2. Combining cost and effectiveness into a single criterion

Net benefit

◆ Net monetary benefit:

$$NMB = \lambda \cdot E - C$$

- E = effectiveness, usually measured in QALYs (utility)
- C = cost, in monetary units (€, £, \$...)
- λ = willingness to pay = cost-effectiveness threshold
- λ is usually measured in \$/QALY, €/QALY, £/QALY...
- > It converts effectiveness into monetary units
- It is specific for each decision maker
- When comparing two or more interventions/strategies, which one is more beneficial?
 - It may depend on λ

NMB as a function of λ

♦ If $\lambda = 6,000 \in /QALY$:

NMB	No therapy	Therapy 1	Therapy 2
Disease present	7,200 €	4,000 €	–31,000€
Disease absent	60,000€	39,400€	–14,200 €

♦ If $\lambda = 15,000$ €/QALY:

NMB	No therapy	Therapy 1	Therapy 2
Disease present	18,000€	40,000€	27,500 €
Disease absent	150,000€	128,500 €	69,500 €

♦ If $\lambda = 30,000$ €/QALY:

NMB	No therapy	Therapy 1	Therapy 2
Disease present	36,000 €	100,000€	125,000€
Disease absent	300,000€	277,000 €	209,000€

Problem: difficult to estimate the WTP

 \bullet λ is different for each decision maker:

```
• USA $50,000-100,000 / QALY
```

• UK £20,000-30,000 / QALY

Spain, Italy ~ €30,000/QALY

Norway ~ €70,000/QALY

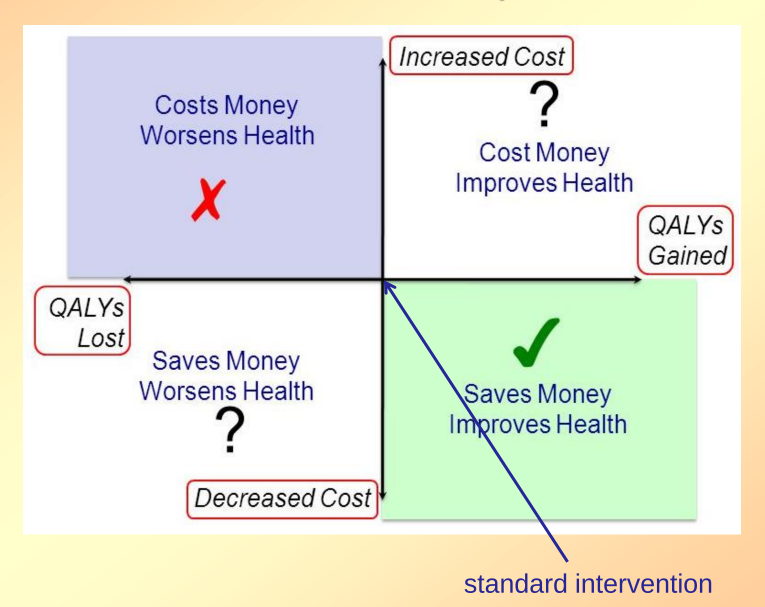
WHO ~3 × (annual per capita GDP) / DALY

- In some countries the range of variation is very wide.
- How to estimate it?
 - 1. Shadow threshold: what interventions are covered in a country
 - 2. Econometric methods
 - No consensus among health economists
- lacktriangle What value of λ should we use in our analyses?
- Solution (partial solution): cost-effectiveness analysis

5.3. Cost-effectiveness analysis

5.3.1. Deterministic CEA

Cost-effectiveness plane



An example with costs and effectiveness

- Two therapies
 - Effectiveness (QALY)

	No therapy	Therapy 1	Therapy 2
Disease present	1.2	4.0	6.5
Disease absent	10	9.9	9.3

- ➤ Therapy 1 cost = 20,000 €
- ➤ Therapy 2 cost = 70,000 €

Questions:

- What therapy to apply when the disease is present
- What therapy to apply when the disease is absent
- Problem: cost-effectiveness analysis

Incremental cost-effectiveness ratio (ICER)

One intervention is more effective but more expensive

$$NMB_{1} = \lambda \times E_{1} - C_{1}$$

$$NMB_{2} = \lambda \times E_{2} - C_{2}$$

$$NHB_{1} > NHB_{2} \Leftrightarrow \frac{C_{2} - C_{1}}{E_{2} - E_{1}} < \lambda$$

Def.: Incremental cost-effectiveness ratio (ICER)

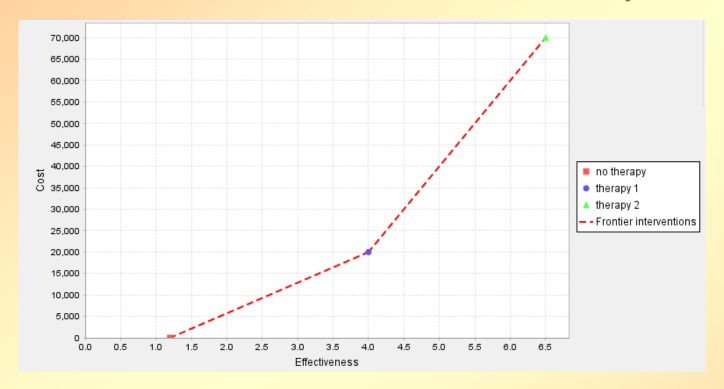
ICER _{2,1} =
$$\frac{C_2 - C_1}{E_2 - E_1}$$

Conclusion

$$NHB_2 > NHB_1 \Leftrightarrow ICER_{2,1} < \lambda$$

 \bullet λ , the WTP, determines which option is more beneficial

When we know that the disease is present



Interval for λ	Cost	Effect.	Best therapy
(0, 7,143)	0	1.2	no-therapy
(7,143, 13,208)	20.000	4.0	therapy-1
(13,208, +∞)	70.000	6.5	therapy-2

Cost-effectiveness partition (CEP)

- Elements of a CEP
 - > n-1 thresholds
 - > n costs
 - > n effectiveness values
 - > n interventions
- ightharpoonup Example with n = 3

Interval for λ	Cost	Effect.	Best therapy
(0, 7,142)	0	1.2	no-therapy
(7,142, 13,208)	20.000	4.0	therapy-1
(13,208, +∞)	70.000	6.5	therapy-2



CEPs are the basis of our algorithms for CEA with PGMs.

An example with costs and effectiveness

- Two therapies
 - Effectiveness (QALY)

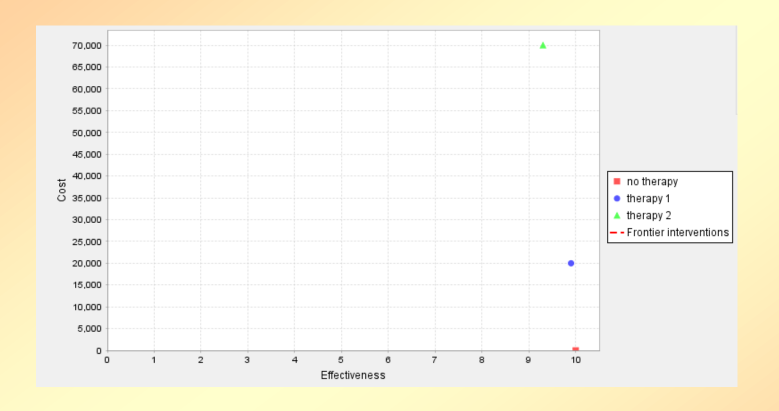
	No therapy	Therapy 1	Therapy 2
Disease present	1.2	4.0	6.5
Disease absent	10	9.9	9.3

- ➤ Therapy 1 cost = 20,000 €
- ➤ Therapy 2 cost = 70,000 €

Questions:

- What therapy to apply when the disease is present
- What therapy to apply when the disease is absent
- Problem: cost-effectiveness analysis

When we know that the disease is absent



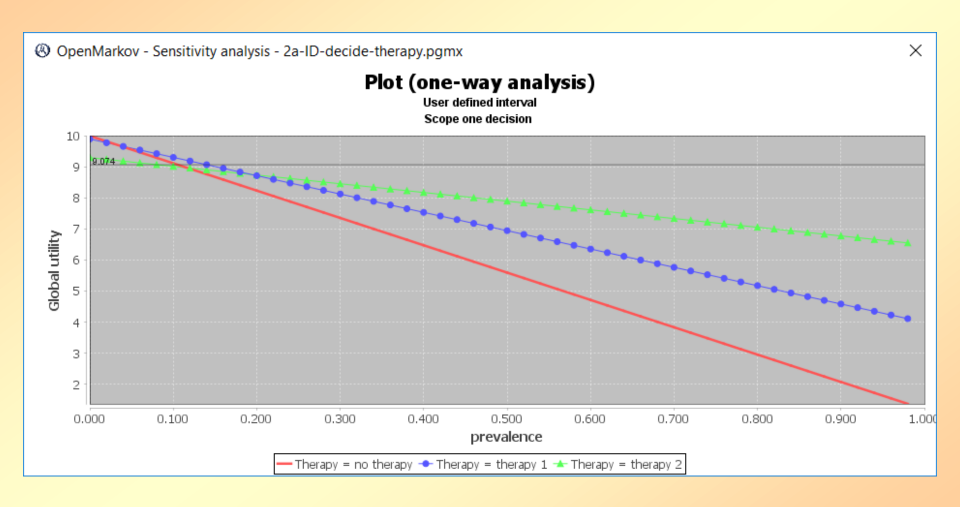
Interval for λ	Cost	Effect.	Best therapy
$(0, +\infty)$	0	10	no-therapy

5.3.2. CEA with uncertain outcomes

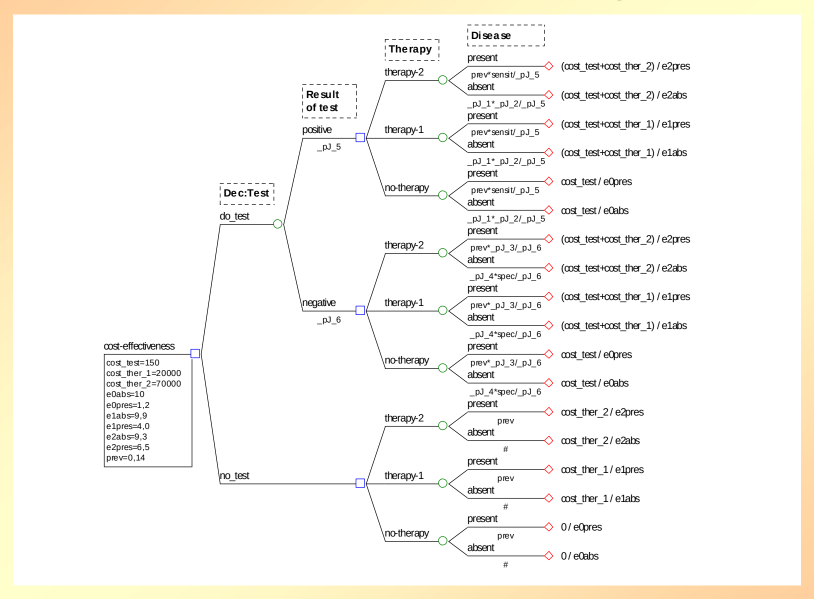
Example with uncertain outcomes: cost-effectiveness of a test

- The costs and effectiveness of the two therapies are the same as in the previous example
- But there is uncertainty (probabilities):
 - prevalence of the disease: 0.14
 - > test: sensitivity 0.90 specificity 0.93
- ◆ Also the test has a cost: 150 €
- Questions:
 - When is the test <u>cost-effective</u>? = What is its <u>ICER</u>?
 - \triangleright What is the most beneficial therapy for each value of λ ?

Effectiveness as a function of prevalence



A decision tree for this example



Problem: the standard algorithm only works for the unicriterion case

A warning and a (rudimentary) solution

"Embedded, or downstream, decision nodes are not useful in cost-effectiveness analysis because the optimal branch cannot be determined when folding back the tree without an explicit decision rule for comparing costs and consequences.

Cost-effectiveness analyses can be performed with a decision tree that has <u>one decision node at the root</u>. The branches of the initial decision node represent <u>all of the strategies</u> that are to be compared."

Kuntz and Weinstein [2001]

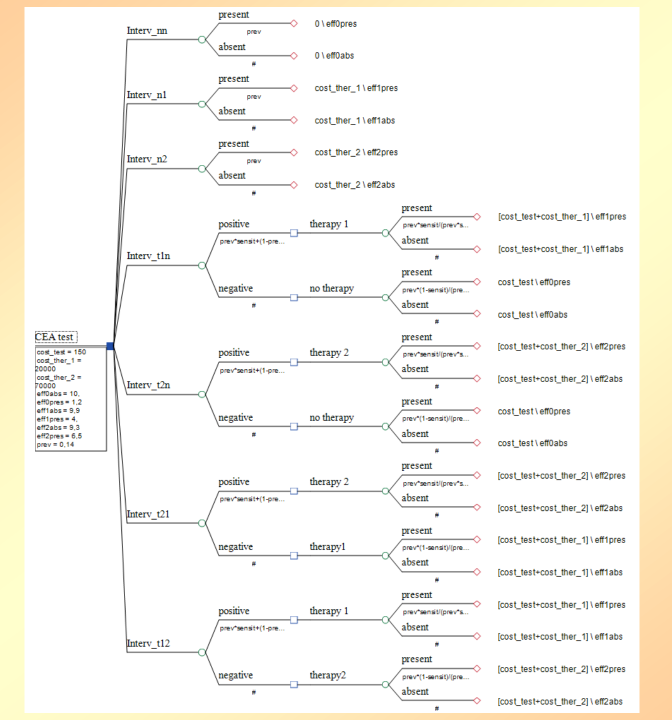
How many strategies for this example?

Without testing

- No therapy in any case
- Always therapy 1
- Always therapy 2

With testing

- ➤ If positive, no therapy; if negative, no therapy.
- ➤ If positive, no therapy; if negative, therapy 1.
- ➤ If positive, no therapy; if negative, therapy 2.
- If positive, therapy 1; if negative, no therapy.
- ➤ If positive, therapy 1; if negative, therapy 1.
- If positive, therapy 1; if negative, therapy 2.
- If positive, therapy 2; if negative, no therapy.
- If positive, therapy 2; if negative, therapy 1.
- ➤ If positive, therapy 2; if negative, therapy 2.



Author's personal copy

PharmacoEconomics (2014) 32:1141–1145 DOI 10.1007/s40273-014-0195-1

RESEARCH LETTER

The Problem of Embedded Decision Nodes in Cost-Effectiveness Decision Trees

Manuel Arias · Francisco Javier Díez

Published online: 31 July 2014

© Springer International Publishing Switzerland 2014

1 Introduction

Cost-effectiveness analysis (CEA) is increasingly used to inform health policies. Decision trees are the standard method for decision analysis in non-temporal domains. A decision node that is not the root of the tree is said to be embedded.

All books on medical decision analysis discuss both CEA *and* decision trees [1–11], but few explain how to conduct a CEA *with* decision trees [1, 2, 10, 11], and only

build a decision tree with one decision node at the root, which represents all the strategies to be evaluated, as proposed by Kuntz and Weinstein; the other is to apply the algorithm presented in Arias and Díez [13].

As a case study, we consider the common problem of finding the incremental cost-effectiveness ratio (ICER) of a test:

Example 1 For a disease with a prevalence of 0.14, there are two possible therapies, the effectiveness of which depends on whether or not the disease is present, as shown

CISIAD

Español | English

Home

Members

Research

- Areas
- Projects
- Contracts
- Seminars
- Colleagues

Publications

- Books
- Papers
- Conferences and workshops
- Technical reports
- Theses

Postgraduate courses (in Spanish)

- Medicina (modular)
- Master AI
- Doctorado

News

Links

Contact

Technical Report

M. Arias and F. J. Díez. **Cost-effectiveness analysis with sequential decisions.** Technical Report CISIAD-11-01, UNED, Madrid, 2011.

26 pages. PDF (859 KB), zip version (827 KB), BibTeX entry.

Abstract

In this paper we present a new method for performing cost-efectiveness analysis of problems that involve multiple decisions and probabilistic outcomes. This problem has been ignored by most of the literature on medical decision making, and the few solutions proposed so far are either wrong or unfeasible except for very small problems. The method proposed in this paper consists of building a decision tree with several decision nodes and evaluating it with a modified roll-back algorithm that operates with partitions of intervals.

Decision trees

See the technical report for an explanation of these examples.

- natural tree (WinDM)
- natural tree (TreeAge Pro)
- <u>all-strategies tree</u> (<u>TreeAge Pro</u>)

Additional information

- Slides presented at SMDM-2007.
- Cost-effectiveness analysis in OpenMarkov.

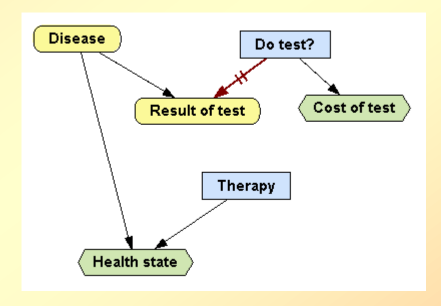
CISIAD. Research Center on Intelligent Decision-Support Systems. UNED. Madrid, Spain.

5.3.3. CEA with IDs and DANs

Influence diagram

Result of test Cost of test Therapy Health state

DAN



- The same structure as in the unicriterion case
- but now we have two criteria: cost and effectiveness

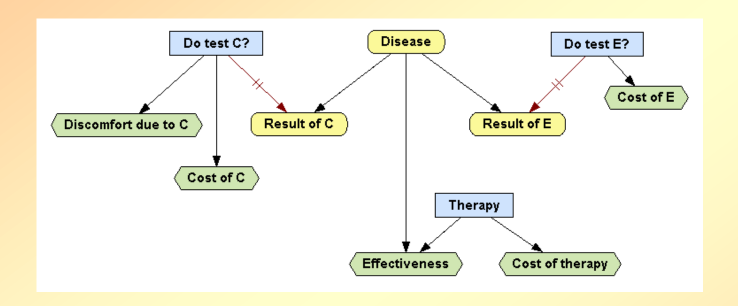
An example with two tests

- ◆ The same disease and therapies as in the previous example.
- ◆ Test E (the same as in the previous example)
 - > sensitivity: 0.90
 - > specificity: 0.93
 - > cost: 150 €
- ◆ Test C
 - > sensitivity: 0.78
 - > specificity: 0.91
 - > cost: 18 €
 - discomfort: 0.001 QALY
- \bullet What is the optimal policy (for each value of λ)?

It is a difficult problem

- Impossible to solve this problem with an ID
 - IDs require a total ordering of the decisions
 - The trick of using two IDs does not work in this case because it does not return all the ICERs
- ◆ Difficult to build a decision tree with embedded dec. nodes
 - It would have 90 leaves
 - Computing the probability of each scenario is cumbersome
- Much more difficult to build a decision tree without embedded decision nodes
 - Finding the possible interventions is a daunting task

... that can be easily solved with a DAN



λ _{inf} (€/QALY)	λ _{sup} (€/QALY)	cost (€)	effect. (QALY)	policy
0	7,747	0	8.768	do nothing
7,747	21,385	2,120	9.046	do test C; if positive {do test E; if positive, therapy 1}
21,385	24,090	7,305	9.284	do test C; if positive {do test E; if positive, therapy 2}
24,090	74,131	9,062	9.357	do test E; if positive {do test C; if negative, therapy 1; if positive, therapy 2}
74,131	112,564	10,735	9.380	do both tests; if both are positive, therapy 2; if only one is positive, therapy 1
112,564	+∞	14,857	9.416	do test E; if positive, therapy 2; if negative {do test C; if positive, therapy 1}

Original Articles

Cost-effectiveness Analysis with Influence Diagrams*

M. Arias; F. J. Díez

Department of Artificial Intelligence, UNED, Madrid, Spain

Keywords

Cost-benefit analysis, cost-effectiveness analysis, decision trees, influence diagrams

Summary

Background: Cost-effectiveness analysis (CEA) is used increasingly in medicine to determine whether the health benefit of an intervention is worth the economic cost. Decision trees, the standard decision modeling technique for non-temporal domains, can only perform CEA for very small problems.

Objective: To develop a method for CEA in problems involving several dozen variables.

Methods: We explain how to build influence diagrams (IDs) that explicitly represent cost and effectiveness. We propose an algorithm for evaluating cost-effectiveness IDs directly, is a without available of the decision of t

Results: The evaluation of an ID returns a set of intervals for the willingness to pay — separated by cost-effectiveness thresholds — and, for each interval, the cost, the effectiveness, and the optimal intervention. The algorithm that evaluates the ID directly is in general much more efficient than the brute-force method, which is in turn more efficient than the expansion of an equivalent decision tree. Using OpenMarkov, an open-source software tool that implements this algorithm, we have been able to perform CEAs on several IDs whose equivalent decision trees contain millions of branches.

Conclusion: IDs can perform CEA on large problems that cannot be analyzed with decision trees.

units divided by cost units; for example, in dollars per death avoided or euros per quality-adjusted life year (QALY) [4]. As the willingness to pay is different for each decision maker, CEA must consider all its possible values. The result of the analysis is usually a set of intervals for λ , each one having an optimal intervention.

When the consequences of the interventions are not deterministic, it is necessary to model the probability of each outcome. Decision trees are the tool used most frequently for this task, especially in medicine [5]. Their main drawback is that their size grows exponentially with the number of variables^b. In the medical literature, trees usually have 3 or 4 variables and between 6 and 10 leaf nodes. A tree of 5 variables typically contains around 20 leaf nodes,









Cost-effectiveness analysis with decision analysis networks

Manuel Arias Manuel Luque Jorge Pérez-Martín Francisco Javier Díez

Universidad Nacional de Educación a Distancia (UNED), Madrid, Spain

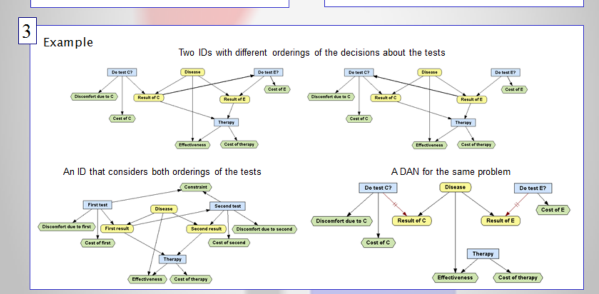
I | Elements of a DAN

- ◆ Structural information: graph
- > three types of nodes: chance, decision, value
- > links connecting nodes; usually represent causality
- ◆ Quantitative information
- > probabilities: prevalence, sensitivities, specificities...
- > value functions:
 - · effectiveness (life years, QALYs...)
 - economic costs (in \$, €, £...)

The same as in influence diagrams.

Representing the flow of information

- ◆ In influence diagrams (IDs)
- > information links
- > temporal-order links between decisions
- > requisite (by definition): a total ordering of the decisions
- ♦ In DΔNs
- > always observed variables
- > revelation links
- > the decisions may be partially ordered
- > the evaluation algorithm will determine the optimal order



4

Result of evaluating the DAN

The optimal policy depends on λ, the willingness to pay: 5 ICER thresholds ⇒ 6 intervals

Hands-on exercise 4

Exercise: Optimal stratety for two tests

Test	sensitivity	specificity	discomfort	cost
Α	0.60	0.92	0.0003 QALY	\$100
В	0.80	0.91	0.0001 QALY	\$200

Disease ->	absent	present
therapy	38 QALY	30 QALY
no therapy	40 QALY	20 QALY

cost of therapy = \$7,000

- ◆ The same probabilities and effectiveness as in exercise 3
- but now we are also considering economic costs.
- Question: What is the most beneficial strategy?

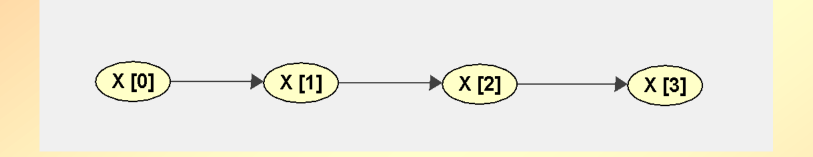
6. Temporal models

Temporal PGMs

- Markov models
 - The future is independent of the past given the present
 - "Markov models do not have memory"
 - Key concept: state
 - Types of models: Markov chains, HMMs, MDPs, POMDPs, DBNs, MIDs, DLIMIDs...
- ◆ Temporal non-Markov models
 - The future is **not** determined by the current state
 - for example, birth occurs around 9 months after conception
 - An type of non-Markov model: event networks
 - Galán, Aguado, Díez, Mira. NasoNet: Modelling the spread of nasopharyngeal cancer with temporal Bayesian networks. Al in Med, 2002.

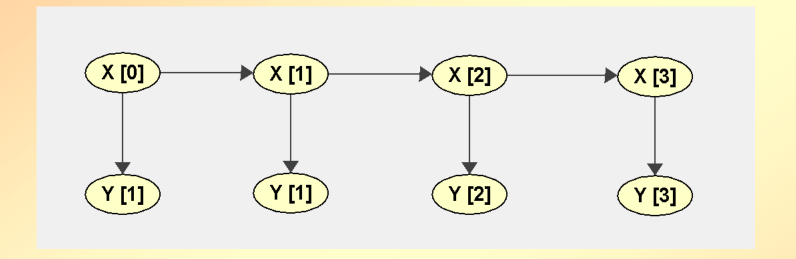
6.1. Types of Markov models

Markov chain



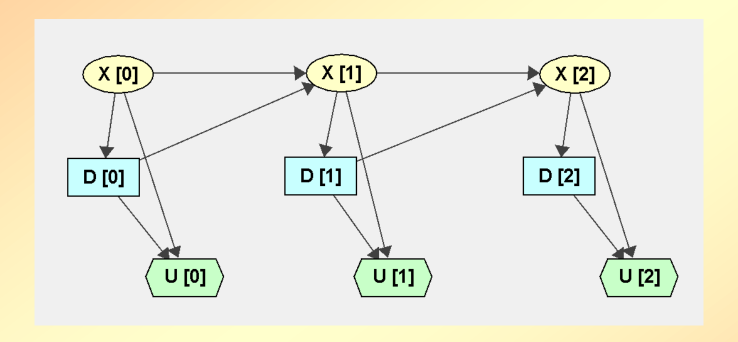
- One variable that evolves over time
- lacktriangle Transition probabilities: $P(x_{i+1}|x_i)$

Hidden Markov model (HMM)



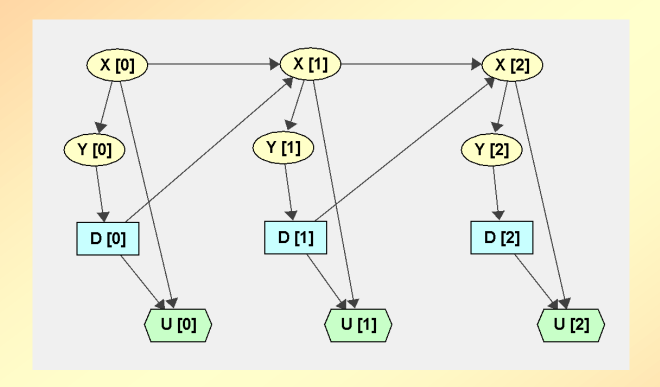
- ◆ Observed variable: *Y*
- ♦ Non-observed (hidden) variable: X
- lacktriangle Transition probabilities: $P(x_{i+1}|x_i)$
- lacktriangle Probability of each observation: $P(y_i|x_i)$

Markov decision process (MDP)



- ◆ Observed variable: X
- ◆ Decision: D
- lacktriangle Transition probabilities: $P(x_{i+1}|x_i)$
- lacktriangle Reward: $U(x_i, d_i)$

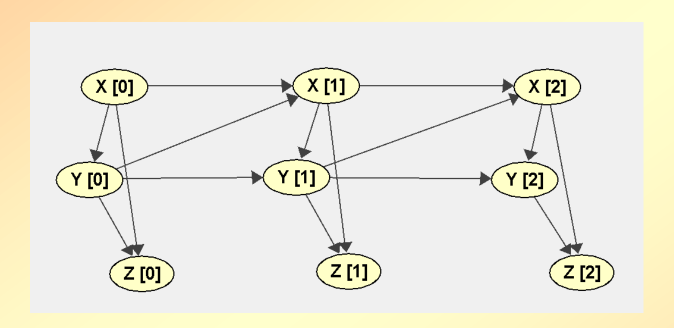
Partially observable MDP (POMDP)



- ◆ Hidden variable: X
- Observed variable : Y
- Decision: D

- Observation prob.: $P(y_i|x_i)$
- ♦ Transition prob.: $P(x_{i+1}|x_i)$
- lacktriangle Reward: $U(x_i, d_i)$

Dynamic Bayesian network (DBN)

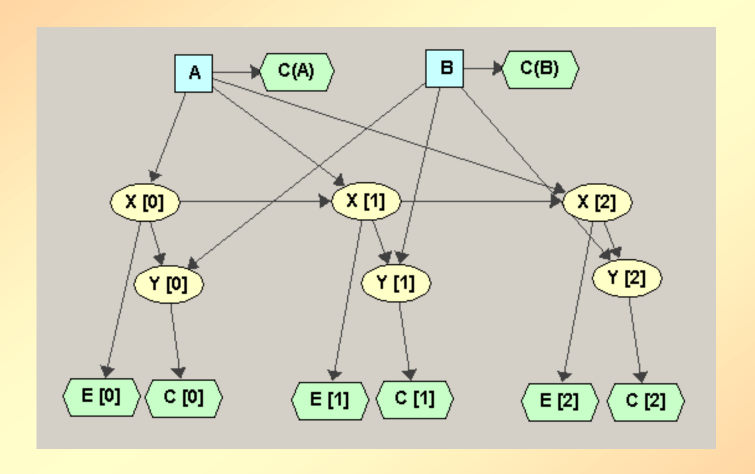


- Markov chain or hidden Markov model:
 - one variable, X
 - one conditional probability: $P(x_{i+1}|x_i)$
- Dynamic Bayesian network:
 - several variables, $\{X, Y, Z...\}$
 - factored probability: $P(y_i|x_i)$, $P(z_i|x_i, y_i)$, $P(x_{i+1}|x_i, y_i)$...

Factored extensions of Markov models

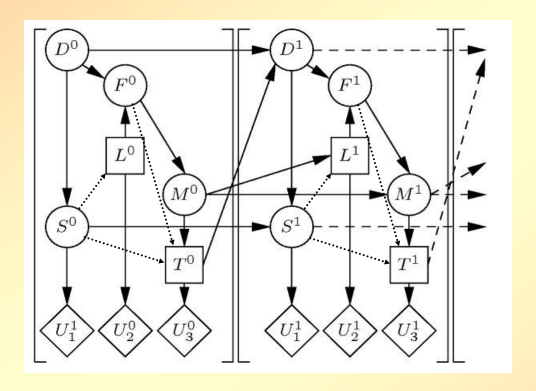
Flat model	Factored model	
Markov chain	Dynamic Bayesian network [Dean and Kanazawa, 1989]	
Hidden Markov model		
Markov decision process (MDP)	Factored MDP [Boutilier et al., 1995, 2000]	
Partially-observable MDP (POMDP)	Factored POMDP [Boutilier and Poole, 1996]	

Markov influence diagrams



Can be used for cost-effectiveness analysis

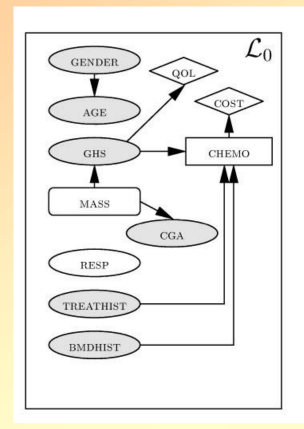
Dynamic limited-memory IDs (DLIMIDs)

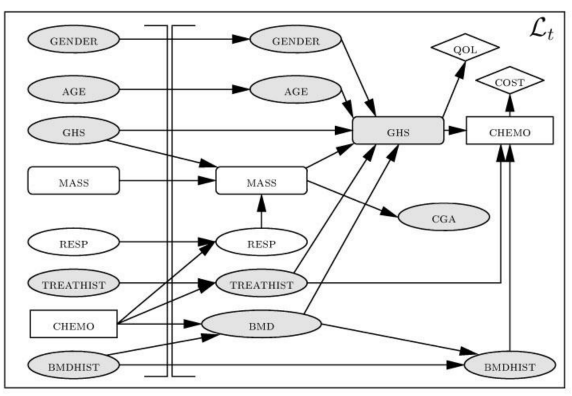


Differences wrt POMDPs

- Several decisions in each time slice.
- Limited memory: the decision maker only knows the observations made in the current time slice and in the previous one
- Memory variables summarize the past.

A DLIMID for a carcinoid tumors





➤ Therapy selection for high-grade carcinoid tumors (van Gerven et al., 2007)

MDPs in Medicine: Opportunities and Challenges

F. J. Díez M. A. Palacios M. Arias Dept. Artificial Intelligence. UNED Madrid, Spain

Abstract

In the last three decades hundreds of Markov models have been built for medical applications, but most of them fall under the paradigm of what we call *simple Markov models* (SMMs). Markov decision processes (MDPs) are much more powerful as a decision analysis tool, but they are ignored in medical decision analysis books and the number of medical applications based on them is still very small. In this paper we compare both types of models and discuss the challenges that MDPs must overcome before they can be widely accepted in medicine. We present a software tool, Open-Markov, that addresses those challenges and has been used to build a Markov model for analyzing the cost-effectiveness of the HPV vaccine.

1 Introduction

Markov models were introduced in the beginning of the 20th century by the Russian mathematician Andrei Andreyevich Markov [1906]. In the three decades passed since the pioneering work of Beck and Pauker [1983], hundreds of the emergence of partially observable Markov decision processes (POMDPs) [Åström, 1965], in which the state of the system is not directly observable, but there is a variable that correlates probabilistically with it. POMDPs were developed in the field of automatic control as an extension of MDPs, but currently most of the research about them is carried out in artificial intelligence (AI), again as a tool for planning, especially in robotics [Ghallab et al., 2004]. The main contribution of AI to this field comes from the area of probabilistic graphical models: Bayesian networks [Pearl, 1988] led to the development of dynamic Bayesian networks [Dean and Kanazawa, 1989], which generalize Markov chains and hidden Markov models [Murphy, 2002]. The idea of using several variables to represent the state of the system, instead of only one, led to factored MDPs [Boutilier et al., 1995; 2000] and factored POMDPs [Boutilier and Poole, 1996], which can model efficiently many problems that were unmanageable with flat (i.e., non-factored) representations; correspondingly, there are new algorithms that can solve problems several orders of magnitude bigger than in the recent past [Hoey et al., 1999; Poupart, 2005; Spaan and Vlassis, 2005].

In the rest of the paper, we use the acronym MDPs to refer to both fully observable and partially observable models

6.2. Markov influence diagrams

Markov Influence Diagrams: A Graphical Tool for Cost-Effectiveness Analysis

Francisco J. Díez, PhD, Mar Yebra, MEng, Iñigo Bermejo, PhD, Miguel A. Palacios-Alonso, MSc, Manuel Arias Calleja, PhD, Manuel Luque, PhD, Jorge Pérez-Martín, MEng

Markov influence diagrams (MIDs) are a new type of probabilistic graphical model that extends influence diagrams in the same way that Markov decision trees extend decision trees. They have been designed to build state-transition models, mainly in medicine, and perform costeffectiveness analyses. Using a causal graph that may contain several variables per cycle, MIDs can model various patient characteristics without multiplying the number of states; in particular, they can represent the history of the patient without using tunnel states. OpenMarkov, an open-source tool, allows the decision analyst to build and evaluate MIDs—including cost-effectiveness analysis and

several types of deterministic and probabilistic sensitivity analysis—with a graphical user interface, without writing any code. This way, MIDs can be used to easily build and evaluate complex models whose implementation as spreadsheets or decision trees would be cumbersome or unfeasible in practice. Furthermore, many problems that previously required discrete event simulation can be solved with MIDs; i.e., within the paradigm of state-transition models, in which many health economists feel more comfortable. Key words: Markov models; influence diagrams; cost-effectiveness analysis; outcomes research. (Med Decis Making XXXX; XX:xx-xx)

Hands-on exercise 5

5. Cost-effectiveness analysis with a Markov model

Exercise

- A disease D may be latent or active. When it is latent, the probability of becoming active in the
 next month is 11% and that of dying is 2%. When the disease is active, the monthly probability of
 dying is 15%. Quality of life is 0.9 for latent and 0.7 for active disease. The cost of the standard
 therapy is £150/month when the disease is latent and £2,500/month when it is active.
- There is a new therapy, with a cost of £950/month, which slows down the progression of the
 disease, so that the monthly probability of becoming active reduces to 8%, without affecting the
 probability of death. Unfortunately, this therapy has no effect when the disease is already active.
- With an annual discount rate of 3.5% for both cost and effectiveness, what is the ICER of the new therapy?
- Is it cost-effective for a willingness-to-pay of £20,000 per QALY?

6.2.1. Example: Chancellor's model for HIV

Case study: HIV/AIDS

(Chancellor et al., 1997)

ORIGINAL RESEARCH ARTICLE

Pharmacoeconomics 1997 Jul 12 (1) 54-66 1170-7690/97/0007-0054/\$06.50/0

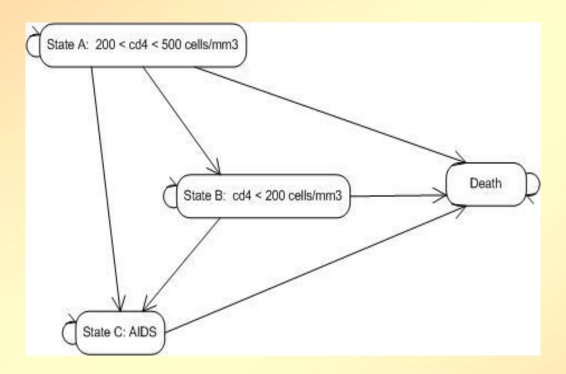
© Adis International Limited All rights reserved

Modelling the Cost Effectiveness of Lamivudine/Zidovudine Combination Therapy in HIV Infection

Jeremy V. Chancellor, Andrew M. Hill, Caroline A. Sabin, Kit N. Simpson and Mike Youle⁵

- Glaxo Wellcome UK Ltd, Uxbridge, Middlesex, England
- 2 Glaxo Wellcome Research and Development Ltd, Greenford, Middlesex, England
- 3 Department of Primary Care and Population Sciences, Royal Free Hospital, London, England
- 4 University of North Carolina, Chapel Hill, North Carolina, USA
- 5 HIV/GUM Research Unit, Chelsea and Westminster Hospital, London, England

◆ State-transition diagram: 4 states

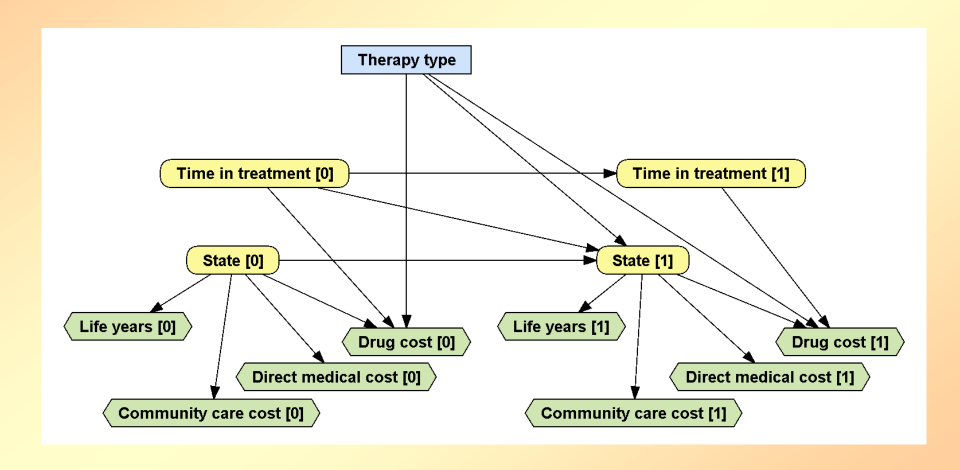


◆ Two therapies:

- monotherapy: AZT only
- > combined therapy: AZT + lamivudine for 2 years; then only AZT

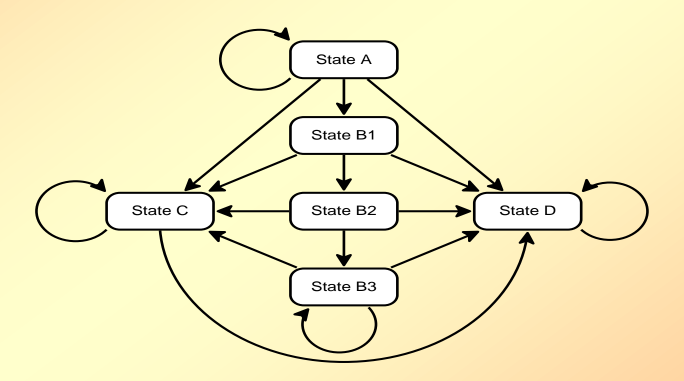
A MID version of the HIV model

[Chancellor et al., 1997]



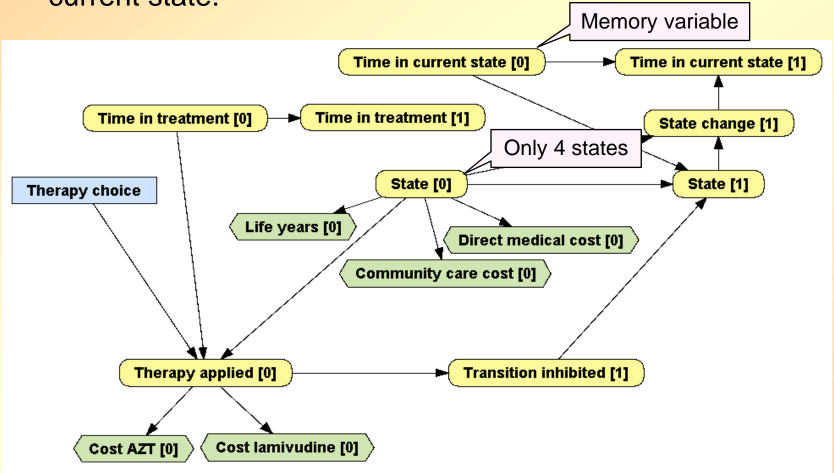
Representing the patient history (1)

- Transition probabilities that depend on the time spent in current state:
 - State-transition model with tunnel states



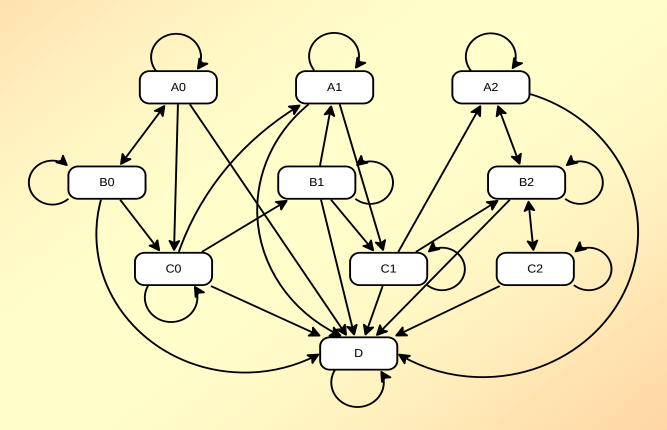
Representing the patient history (1)

◆ Transition probabilities that depend on the time spent in current state:



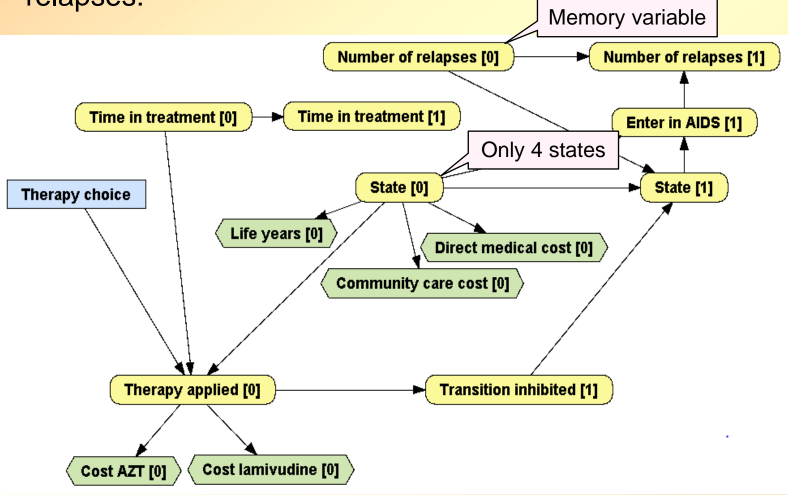
Representing the patient history (2)

Transition probabilities that depend on the number of relapses:



Representing the patient history (2)

Transition probabilities that depend on the number of relapses:





Case study: Hip replacement

(Briggs et al., 2004)

ARTICLE

Appl Health Econ Health Policy 2004; 3 (2): 79-89 1175-5652/04/0002-0079/\$31.00/0

2004 Adis Data Information BV. All rights reserved.

The Use of Probabilistic Decision Models in Technology Assessment

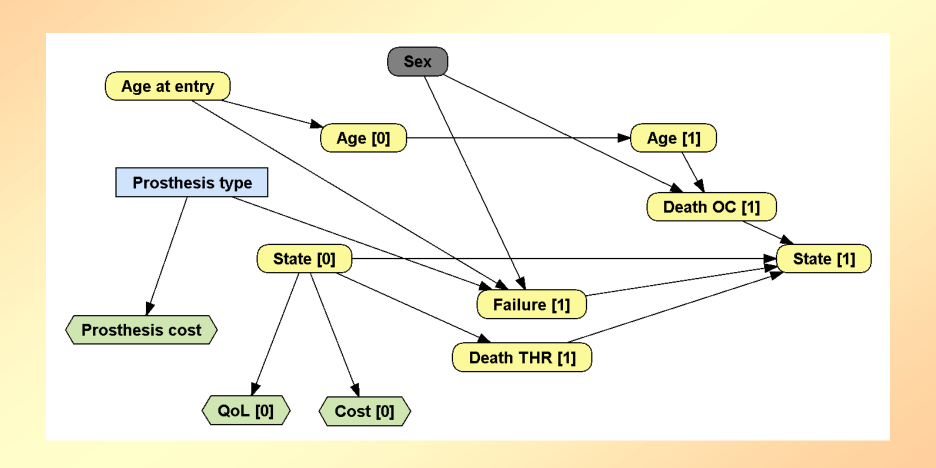
The Case of Total Hip Replacement

Andrew Briggs, Mark Sculpher, Jill Dawson, Ray Fitzpatrick, David Murray and Henrik Malchau

- 1 Health Economics Research Centre, Department of Public Health, University of Oxford, Old Road Campus, Headington, Oxford, UK
- 2 Centre for Health Economics, University of York, Heslington, York, UK
- 3 School of Health and Social Care, Oxford Brookes University, Oxford, UK
- 4 Department of Public Health, University of Oxford, Old Road Campus, Headington, Oxford, UK
- 5 Nuffield Orthopaedic Centre, Headington, Oxford, UK
- 6 Department of Orthopaedics, Massachusetts General Hospital, Boston, USA

A MID version of the hip replacement model

[Briggs et al., 2004]



Case study: HPV vaccine

(Insinga et al., 2009)

BMC Infectious Diseases



Research article

Open Access

Epidemiologic natural history and clinical management of Human Papillomavirus (HPV) Disease: a critical and systematic review of the literature in the development of an HPV dynamic transmission model Ralph P Insinga*, Erik J Dasbach and Elamin H Elbasha

Address: Department of Health Economic Statistics, Merck Research Laboratories, North Wales, PA, USA

Email: Ralph P Insinga* - ralph_insinga@merck.com; Erik J Dasbach - erik_dasbach@merck.com; Elamin H Elbasha - elamin_elbasha@merck.com

* Corresponding author

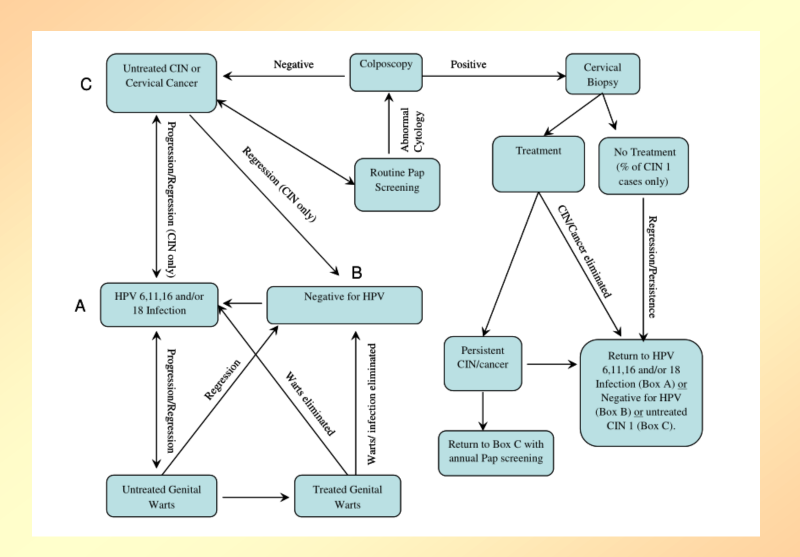
Published: 29 July 2009

BMC Infectious Diseases 2009, 9:119 doi:10.1186/1471-2334-9-119

This article is available from: http://www.biomedcentral.com/1471-2334/9/119

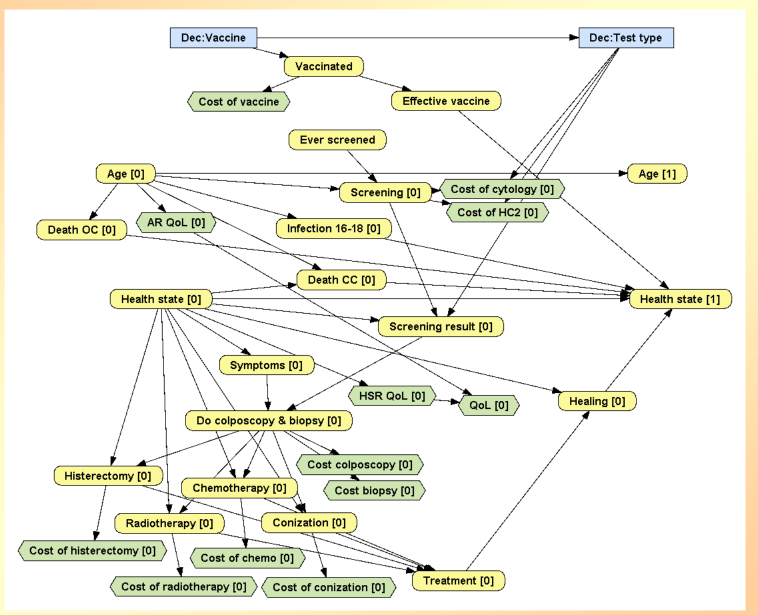
Received: 13 May 2008

Accepted: 29 July 2009



A MID version of the HPV vaccination model

[Callejo et al., 2010]



Content of one of the Excel cells for this model:

=VLOOKUP(\$C5; Variables!\$A\$4:\$H\$21;8;TRUE)*(((BI5+BJ5)+BK5*u CIN1+SUM(BL5:BP5)*uCIN2_3+(BQ5+BR5)*uLCC+(BS5+BT5)*uRCC+(BU5+BV5)*uDCC)+((BI4+BJ4)+BK4*uCIN1+SUM(BL4:BP4)*uCIN2_3+(BQ4+BR4)*uLCC+(BS4+BT4)*uRCC+(BU4+BV4)*uDCC)*VLOOKUP(\$C4; Variables!\$A\$4:\$H\$21;2;TRUE)+(BQ4+BR4)*uLCC*VLOOKUP(\$C4; Variables!\$A\$4:\$H\$21;4;TRUE)+(BS4+BT4)*uRCC*VLOOKUP(\$C4; Variables!\$A\$4:\$H\$21;5;TRUE)+(BU4+BV4)*uDCC*VLOOKUP(\$C4; Variables!\$A\$4:\$H\$21;5;TRUE)+(BU4+BV4)*uDCC*VLOOKUP(\$C4; Variables!\$A\$4:\$H\$21;5;TRUE)+(BU4+BV4)*uDCC*VLOOKUP(\$C4; Variables!\$A\$4:\$H\$21;2;TRUE))

Case study: AIDS in Africa

(Ryan et al., 2009)

The cost-effectiveness of cotrimoxazole prophylaxis in HIV-infected children in Zambia

Máirín Ryan^a, Susan Griffin^b, Bona Chitah^c, A. Sarah Walker^d, Veronica Mulenga^e, Donald Kalolo^e, Neil Hawkins^b, Concepta Merry^a, Michael G. Barry^a, Chifumbe Chintu^e, Mark J. Sculpher^b and Diana M. Gibb^d

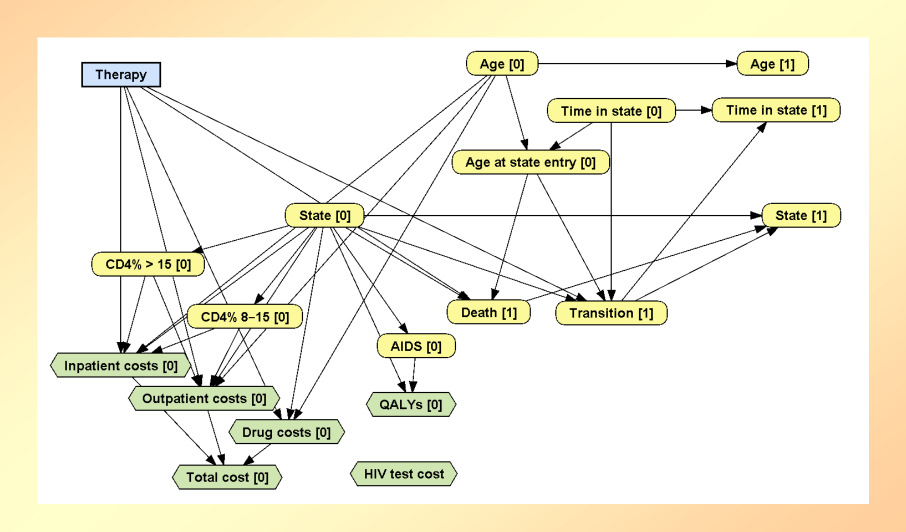
Objective: To assess the cost-effectiveness of cotrimoxazole prophylaxis in HIV-infected children in Zambia, as implementation at the local health centre level has yet to be undertaken in many resource-limited countries despite recommendations in recent updated World Health Organization (WHO) guidelines.

Design: A probabilistic decision analytical model of HIV/AIDS progression in children based on the CD4 cell percentage (CD4%) was populated with data from the placebocontrolled Children with HIV Antibiotic Prophylaxis trial that had reported a 43% reduction in mortality with cotrimoxazole prophylaxis in HIV-infected children aged 1–14 years.

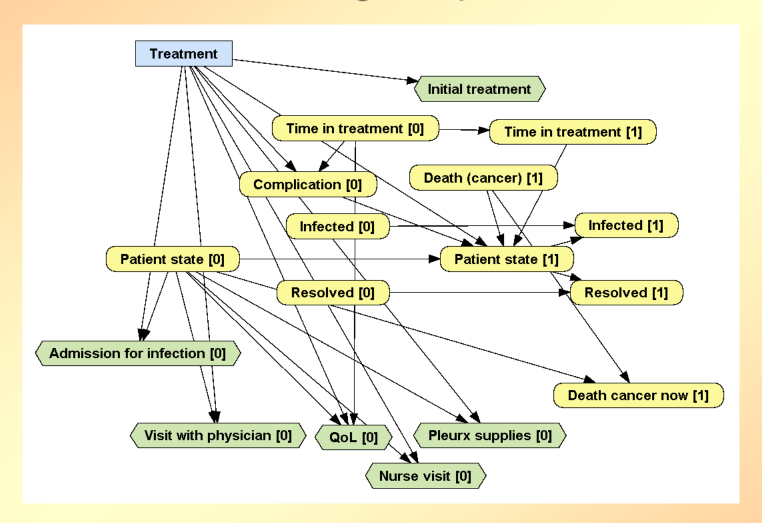
Methods: Unit costs (US\$ in 2006) were measured at University Teaching Hospital, Lusaka. Cost-effectiveness expressed as cost per life-year saved, cost per quality adjusted life-year (QALY) saved, cost per disability adjusted life-year (DALY) averted was calculated across a number of different scenarios at tertiany and primary healthcare centres.

A MID version of the CHAP model

[Ryan et al., 2008]

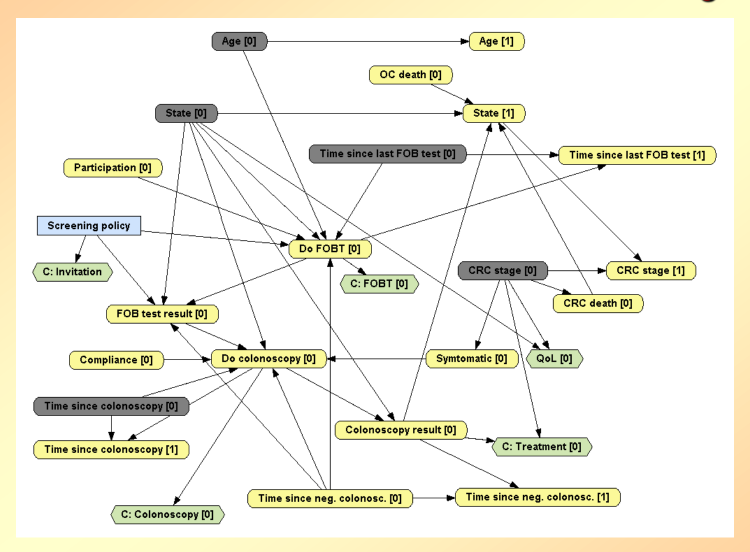


Our model for malignant pleural effusion

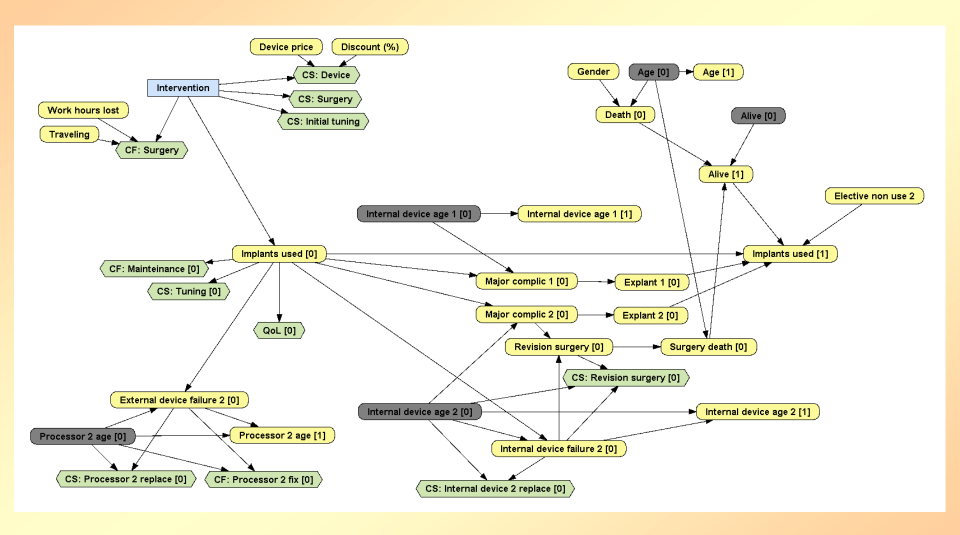


Meeting of the Society for Medical Decision Making (SMDM 2015), St. Louis, October 2015.

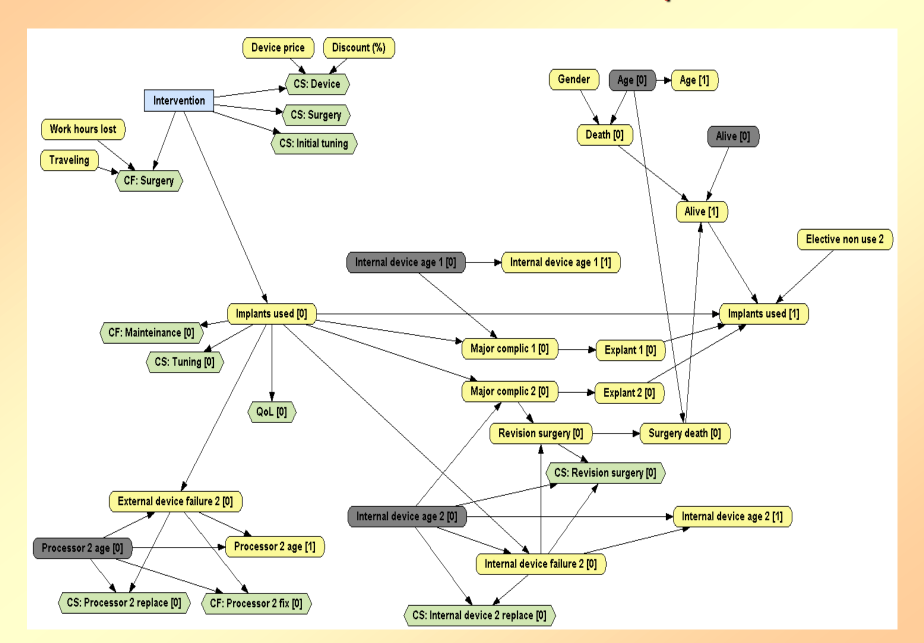
Our model for colorectal cancer screening

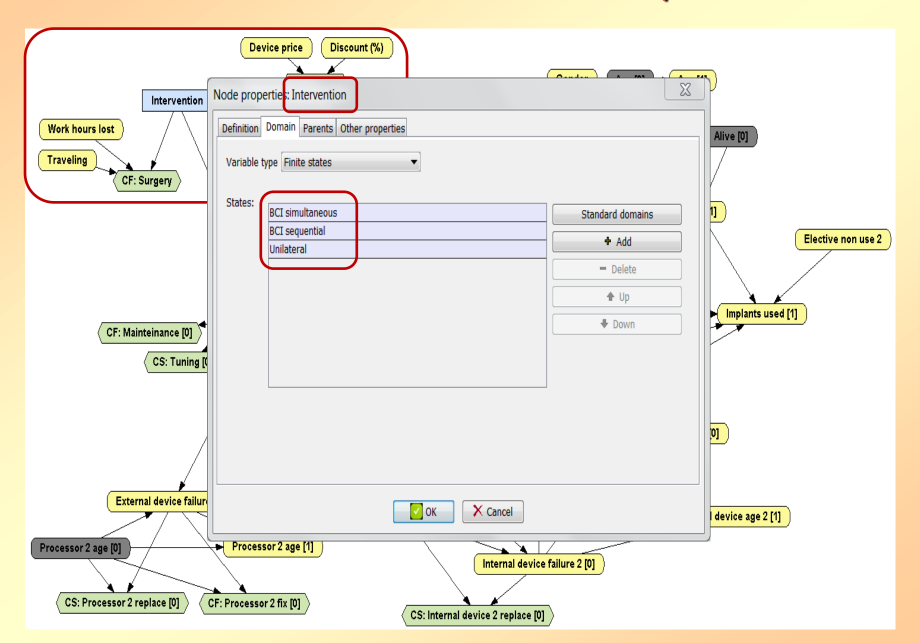


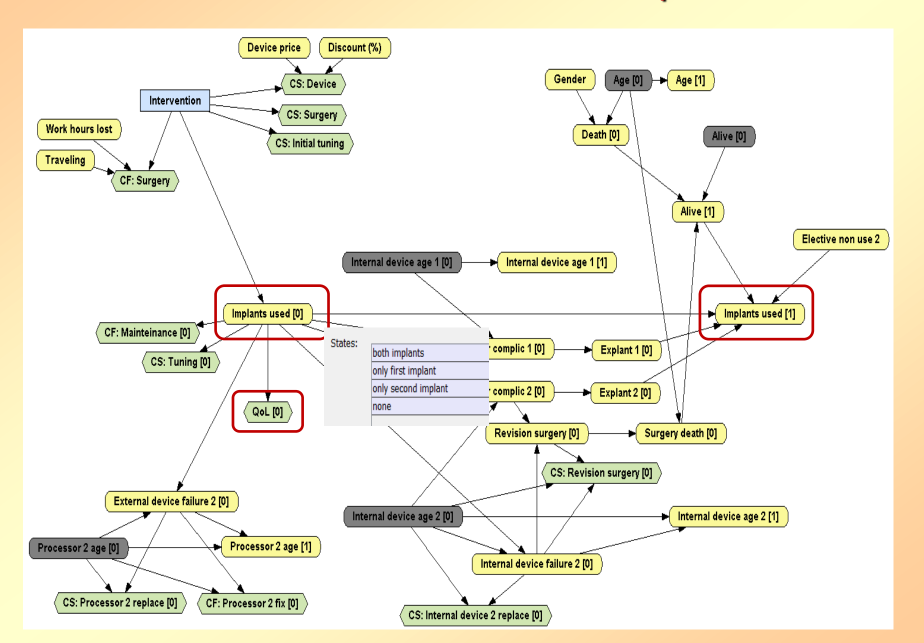
European Conference of the Society for Medical Decision Making, London, UK, June 2015.

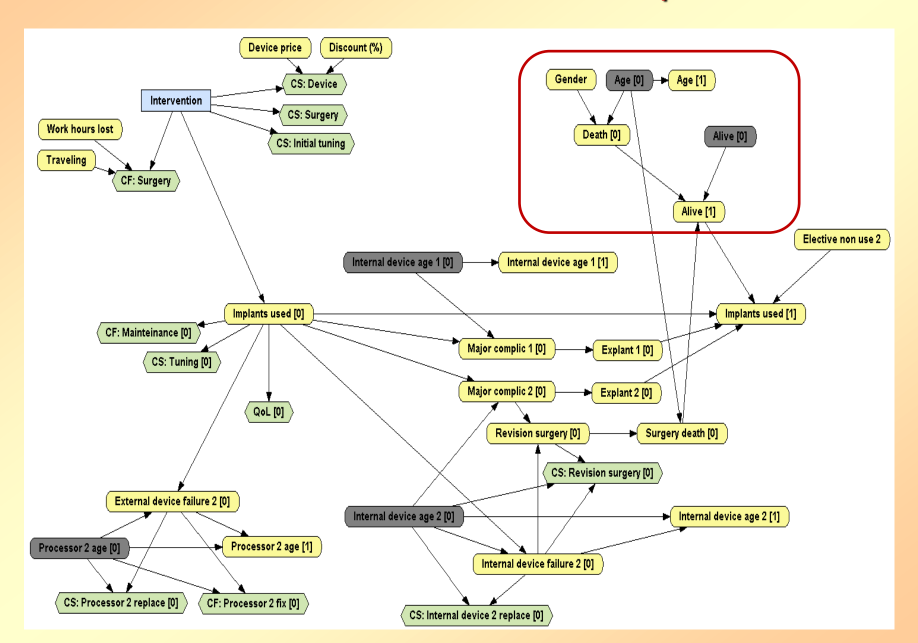


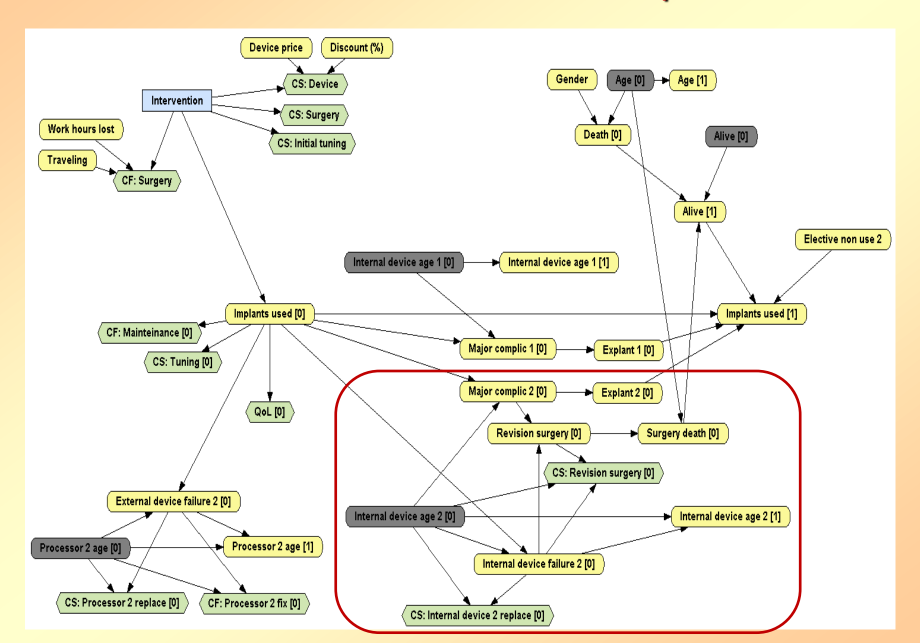
Cochlear Implant Symposium, Washington DC, October 2015.

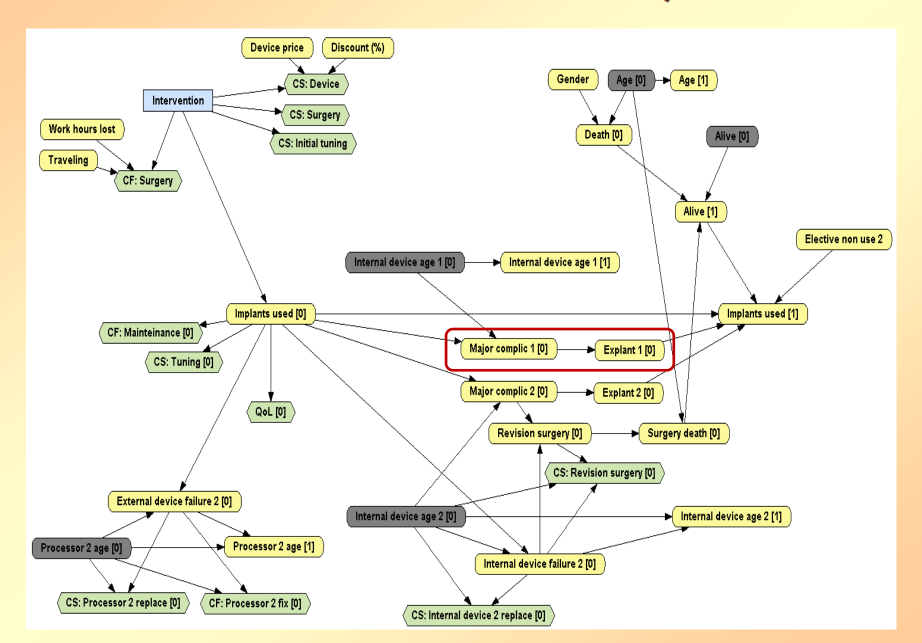


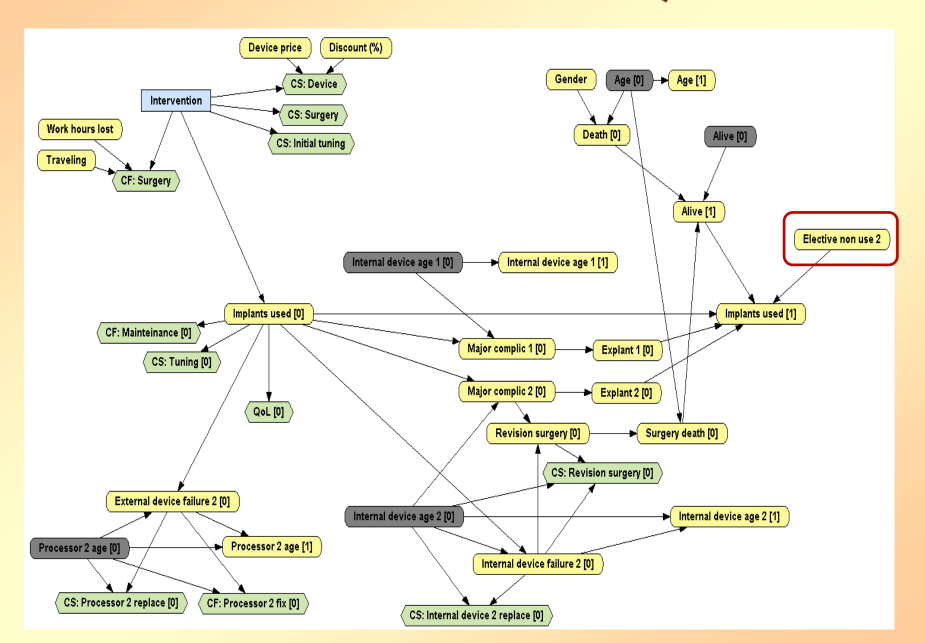


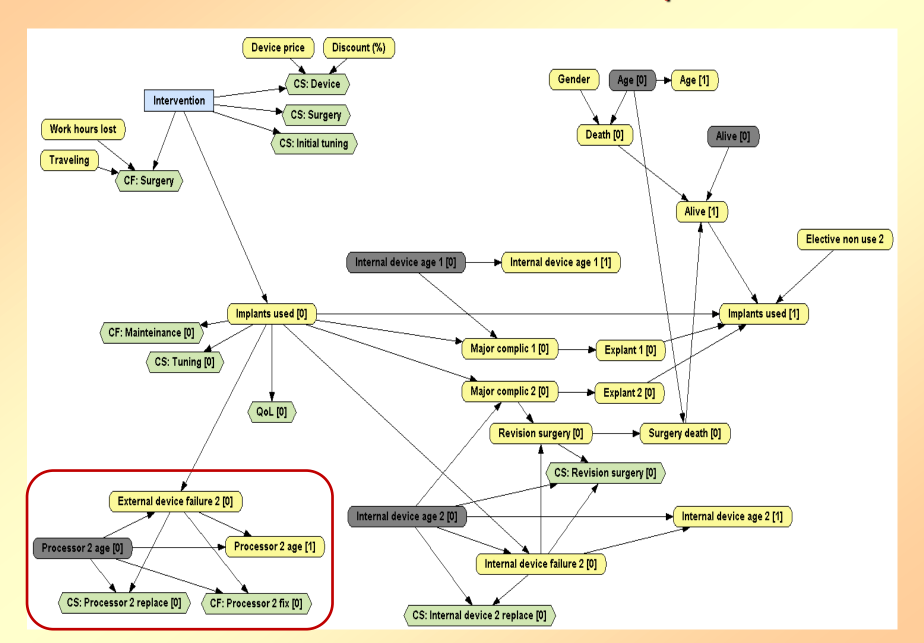






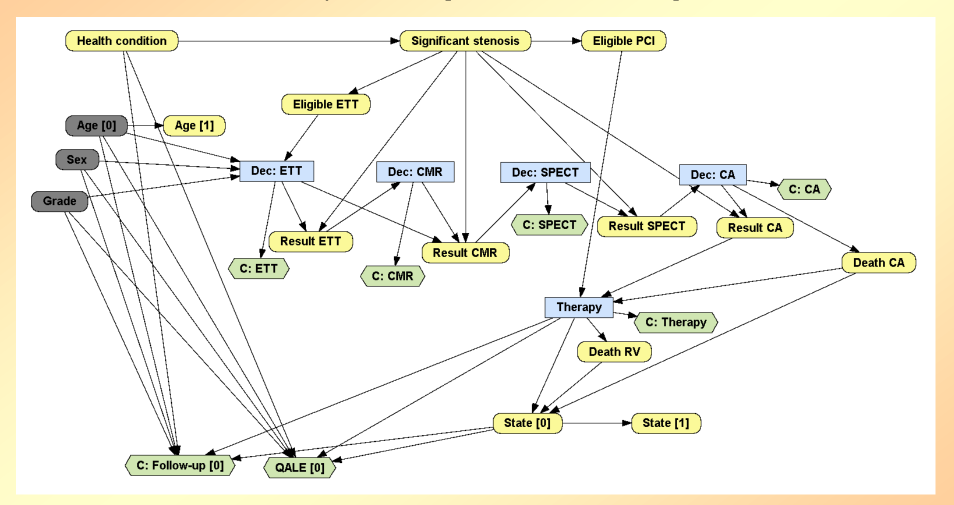






A MID with several decisions

Adapted from [Walker et al., 2013]



- This model evaluates all the possible interventions.
- It can cope with heterogeneity: sex, age, grade.

ORIGINAL ARTICLE

Cost-effectiveness of cardiovascular magnetic resonance in the diagnosis of coronary heart disease: an economic evaluation using data from the CE-MARC study

Simon Walker, ¹ François Girardin, ^{1,2,3} Claire McKenna, ¹ Stephen G Ball, ⁴ Jane Nixon, ⁵ Sven Plein, ⁴ John P Greenwood, ⁴ Mark Sculpher ¹

► Additional material is published online only. To view please visit the journal online (http://dx.doi.org/10.1136/ heartjnl-2013-303624).

Centre for Health Economics, University of York, York, UK ²Medical Direction, Geneva University Hospitals, Geneva, Switzerland 3Division of Clinical Pharmacology and Toxicology, Geneva University Hospitals, Geneva, Switzerland 4Multidisciplinary Cardiovascular Research Centre and Leeds Institute of Genetics, Health and Therapeutics, University of Leeds, Leeds, UK ⁵Clinical Trials Research Unit, University of Leeds, Leeds, UK

Correspondence to

Simon Walker, Centre for Health Economics, University of York, Alcuin A Block, Heslington, York YO10 5DD, UK; simon.walker@york.ac.uk

Received 10 January 2013 Revised 15 March 2013 Accepted 17 March 2013

ABSTRACT

Objective To evaluate the cost-effectiveness of diagnostic strategies for coronary heart disease (CHD) derived from the CE-MARC study.

Design Cost-effectiveness analysis using a decision analytic model to compare eight strategies for the diagnosis of CHD.

Setting Secondary care out-patients (Cardiology Department).

Patients Patients referred to cardiologists for the further evaluation of symptoms thought to be angina pectoris.

Interventions Eight different strategies were considered, including different combinations of exercise treadmill testing (ETT), single-photon emission CT (SPECT), cardiovascular magnetic resonance (CMR) and coronary angiography (CA).

Main outcome measures Costs expressed as UK sterling in 2010–2011 prices and health outcomes in quality-adjusted life-years (QALYs). The time horizon was 50 years.

Results Based on the characteristics of patients in the CE-MARC study, only two strategies appear potentially cost-effective for diagnosis of CHD, both including CMR. The choice is between two strategies: one in which CMR follows a positive or inconclusive ETT, followed by CA if CMR is positive or inconclusive (Strategy 3 in the model); and the other where CMR is followed by CA if

INTRODUCTION

Coronary heart disease (CHD) is a leading cause of death and disability worldwide. In the UK, over 2 million people are living with CHD and, in 2007, it was estimated to account for over 94 000 deaths, of which over 31 000 were considered premature.¹

A variety of investigations may be used to diagnose CHD and identify patients who require coronary revascularisation; all these tests, however, have their limitations. Increasingly, non-invasive imaging has replaced exercise treadmill testing (ETT), with single-photon emission CT (SPECT) being the most commonly used test for myocardial ischaemia worldwide.² Cardiovascular magnetic resonance (CMR) imaging is increasingly used for the diagnosis of CHD as a result of its safety (no ionising radiation), high spatial resolution and ability to assess multiple aspects of CHD pathology in both the stable and unstable clinical settings.^{3–8}

The diagnosis of CHD has no direct health benefit in itself; instead, any improved accuracy in diagnosis should result in more appropriate treatment which can confer health benefits on patients. The optimal management of patients with CHD continues to be debated, but options include medical therapy, percutaneous coronary intervention (PCI) or coronary artery bypass grafting (CABG). Many patients with

Model structure

To conduct the economic evaluation a decision analytic model was developed. For the initial diagnosis a decision tree allocates patients to the appropriate diagnostic group. The prognostic implications of being in one of these groups are then quantified using three distinct Markov models. An example of the decision tree for Strategy 2 (ETT, followed by CA if ETT is positive or inconclusive) is shown in figure 1.

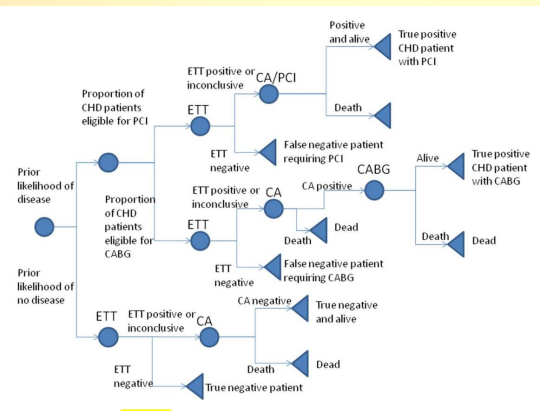


Figure 1 Structure of decision tree using Strategy 2 as an example. CA, coronary angiography; CABG, coronary artery bypass grafting; CHD, coronary heart disease; ETT, exercise treadmill testing; PCI, percutaneous coronary intervention.

CrossMark

REVIEW ARTICLE

A Comparison of Four Software Programs for Implementing Decision Analytic Cost-Effectiveness Models

Chase Hollman¹ · Mike Paulden^{1,2} · Petros Pechlivanoglou^{3,4,5} · Christopher McCabe¹

Published online: 9 May 2017

© Springer International Publishing Switzerland 2017

Abstract The volume and technical complexity of both academic and commercial research using decision analytic modelling has increased rapidly over the last two decades. The range of software programs used for their implementation has also increased, but it remains true that a small number of programs account for the vast majority of cost-effectiveness modelling work. We report a comparison of four software programs: TreeAge Pro, Microsoft Excel, R and MATLAB. Our focus is on software commonly used for building Markov models and decision trees to conduct cohort simulations, given their predominance in the published literature around cost-effectiveness modelling. Our comparison uses three qualitative criteria as proposed by Eddy et al.: "transparency and validation", "learning curve" and "capability". In addition, we introduce the quantitative criterion of processing speed. We also consider the cost

sion of this implementary

Electronic supplementary material The online version of this article (doi:10.1007/s40273-017-0510-8) contains supplementary material, which is available to authorized users.

of each program to academic users and commercial users. We rank the programs based on each of these criteria. We find that, whilst Microsoft Excel and Tree-Age Pro are good programs for educational purposes and for producing the types of analyses typically required by health technology assessment agencies, the efficiency and transparency advantages of programming languages such as MATLAB and R become increasingly valuable when more complex analyses are required.

Key Points for Decision Makers

Microsoft Excel and TreeAge Pro are good programs for implementing the types of cost-effectiveness analyses commonly required by health technology assessment bodies.

MATLAB and R are particularly valuable for implementing more complex decision analytic models and computationally demanding analyses, such as expected value of perfect parameter information (EVPPI), due to their processing speed and transparency.

Hands-on exercise 5

6.2.3. Within-cycle corrections in Markov models

Theoretical Foundations and Practical Applications of Within-Cycle Correction Methods

Elamin H. Elbasha, PhD, Jagpreet Chhatwal, PhD

Background. Modeling guidelines recommend applying a half-cycle correction (HCC) to outcomes from discretetime state-transition models (DTSTMs). However, there is still no consensus on why and how to perform the correction. The objective was to provide theoretical foundations for HCC and to compare (both mathematically and numerically) the performance of different correction methods in reducing errors in outcomes from DTSTMs. Methods. We defined 7 methods from the field of numerical integration: Riemann sum of rectangles (left, midpoint, right), trapezoids, life-table, and Simpson's 1/3rd and 3/8th rules. We applied these methods to a standard 3-state disease progression Markov chain to evaluate the costeffectiveness of a hypothetical intervention. We solved the discrete- and continuous-time (our gold standard) versions of the model analytically and derived expressions for various outcomes including discounted qualityadjusted life-years, discounted costs, and incremental cost-effectiveness ratios. Results. The standard HCC

method gave the same results as the trapezoidal rule and life-table method. We found situations where applying the standard HCC can do more harm than good. Compared with the gold standard, all correction methods resulted in approximation errors. Contrary to conventional wisdom, the errors need not cancel each other out or become insignificant when incremental outcomes are calculated. We found that a wrong decision can be made with a less accurate method. The performance of each correction method vastly improved when a shorter cycle length was selected; Simpson's 1/3rd rule was the fastest method to converge to the gold standard. Conclusion. Cumulative outcomes in DTSTMs are prone to errors that can be reduced with more accurate methods like Simpson's rules. We clarified several misconceptions and provided recommendations and algorithms for practical implementation of these methods. Key words: state-transition models; discrete time; continuous time; half-cycle correction; numerical integration. (Med Decis Making XXXX;XX:XX-XX)

Within-cycle correction methods

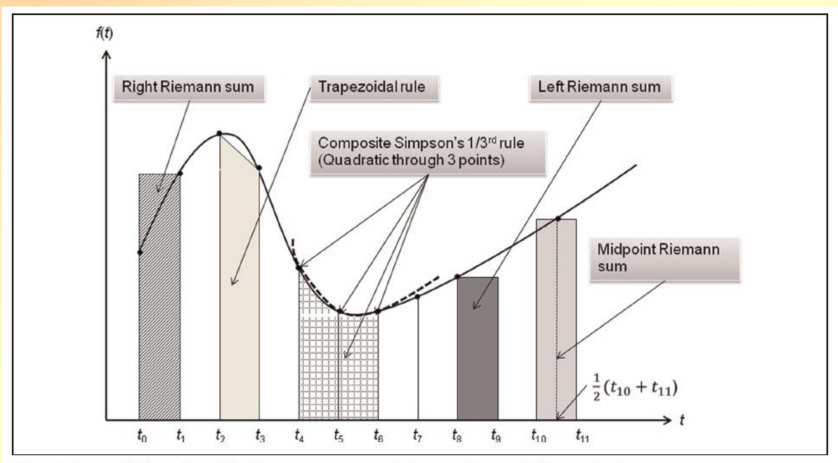


Figure 1 Geometric illustration of the Riemann sums, trapezoidal rule, and composite Simpson's rules.

Med Decis Making. 2019 May;39(4):414-420. doi: 10.1177/0272989X19837974. Epub 2019 Mar 28.

Evaluation of Markov Models with Discontinuities.

Pérez-Martín J¹, Bermejo I, Díez FJ¹.

Author information

1 Department of Artificial Intelligence, Universidad Nacional de Educación a Distancia (UNED), Madrid, Spain (JP, FJD), and School of Health and Related Research, University of Sheffield, Sheffield, South Yorkshire, UK.

Abstract

Background. Several methods, such as the half-cycle correction and the life-table method, were developed to attenuate the error introduced in Markov models by the discretization of time. Elbasha and Chhatwal have proposed alternative "corrections" based on numerical integration techniques. They present an example whose results suggest that the trapezoidal rule, which is equivalent to the half-cycle correction, is not as accurate as Simpson's 1/3 and 3/8 rules. However, they did not take into consideration the impact of discontinuities. Objective. To propose a method for evaluating Markov models with discontinuities. Design. Applying the trapezoidal rule, we derive a method that consists of adjusting the model by setting the cost at each point of discontinuity to the mean of the left and right limits of the cost function. We then take from the literature a model with a cycle length of 1 year and a discontinuity on the cost function and compare our method with other "corrections" using as the gold standard an equivalent model with a cycle length of 1 day. Results. As expected, for this model, the life-table method is more accurate than assuming that transitions occur at the beginning or the end of cycles. The application of numerical integration techniques without taking into account the discontinuity causes large errors. The model with averaged cost values yields very small errors, especially for the trapezoidal and the 1/3 Simpson rules. Conclusion. In the case of discontinuities, we recommend applying the trapezoidal rule on an averaged model because this method has a mathematical justification, and in our empirical evaluation, it was more accurate than the sophisticated 3/8 Simpson rule.

KEYWORDS: Markov models; discontinuities; half-cycle correction; state-transition models; within-cycle correction

PMID: 30920897 DOI: 10.1177/0272989X19837974

6.2.3. MIDs vs. other types of models

Advantages of MIDs for CEA

For model builders

- No programming is required, not even for sensitivity analysis
- The construction of the model is much faster and easier.
- It is possible to accomplish each phase (structure, numeric parameters, deterministic analysis, sensitivity analysis) without thinking of the next one
- Debugging consists only of refining the knowledge contained in the model: it is not necessary to debug formulas and macros.
- For the recipients of the model (agencies: NICE, etc.)
 - Just by observing the graph it is possible to find out the basic structure of the model its main hypotheses.
 - It is not necessary to check that the code (formulas, macros...) is correct.

Comparison of MIDs with other techniques

- ♦ MIDs vs. spreadsheets (Excel)
 - no need to write any formulas nor VisualBasic macros
 - no need to multiply the number of states
 - difficult to write functions of parameters
- MIDs vs. Markov decision trees
 - ➤ much more compact ⇒ possible to build much larger models
 - no need to add tracking variables (microsimulation)
- ♦ MIDs vs. a programming language (R, C++, MATLAB...)
 - > no need to write any code, not even for sensitivity analysis
 - but programming languages are much more flexible
- MIDs vs. discrete event simulation
 - cohort propagation (exact algorithm) is often much faster
- MIDs vs. all the others: may contain several decisions.

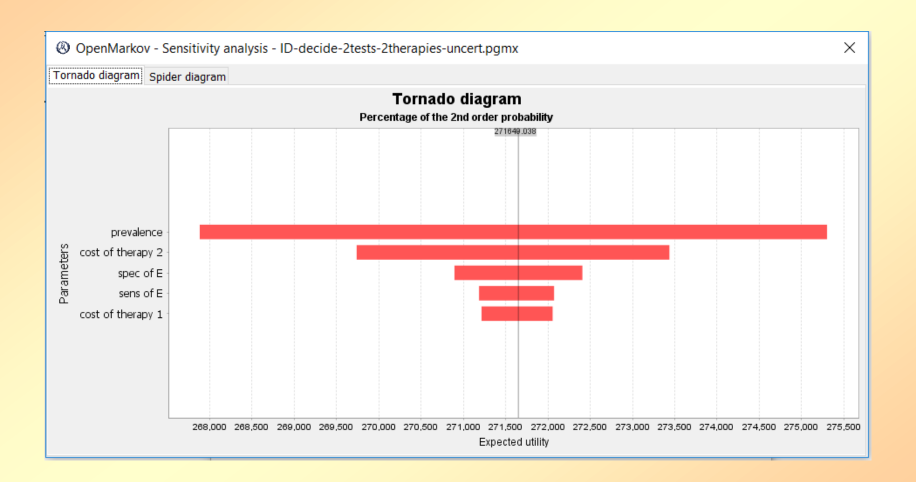
7. Sensitivity analysis

Types of sensitivity analysis

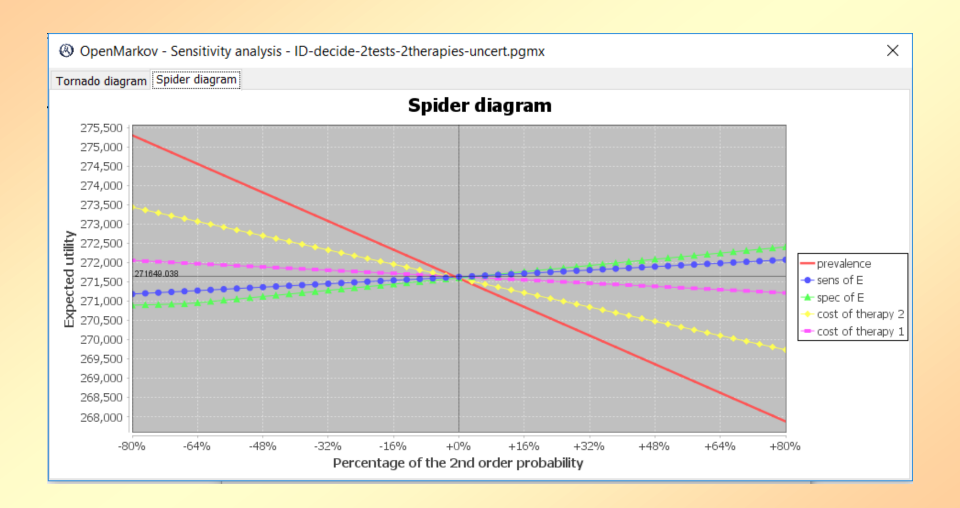
- Two main types
 - structural (qualitative)
 - parametric (quantitative)
- Depending on the effect analyzed
 - analysis of utility
 - analysis of decisions / policies
- Depending on how many parameters are varied
 - one-way analysis
 - n-way analysis (independent or join analysis)
- Depending on how the parameters are varied
 - range (interval)
 - probability distribution
 - look for thresholds (changes in policies)

7.1. Unicriterion sensitivity analysis

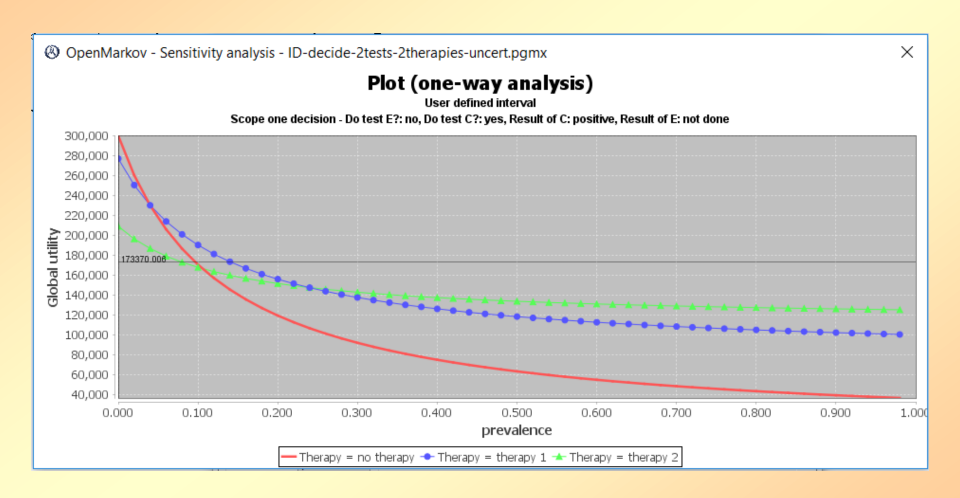
Tornado diagram



Spider diagram



Plot (one-way sensitivity analysis)



7.2. Cost-effectiveness sensitivity analysis

Scatter plot



Acceptability curve



Some sensitivity analysis options

	unicriterion	cost-effectiveness
deterministic	 tornado / spider diagram (global) plot (global / for a decision) map (global / for a decision) 	C.E. spider diagram (global)
probabilistic	acceptability (for a decision)EVPI (global)	 scatter plot + acceptability curve (for a decision) EVPI curve (global)

8. Overview of software tools

Software tools for CEA

REVIEW OF SOFTWARE FOR DECISION MODELLING.

DECISION SUPPORT UNIT

Jon Tosh and Allan Wailoo

Health Economics and Decision Science, School of Health and Related Research,

University of Sheffield

Table 1 - Software used for NICE Technology Appraisals

Software	Respondents that used this software		Number of	Number of	Number of	
			TAGs	Manufacturers	Consultancies	
	n	%				
MS Excel	28	100%	6	14	8	
TreeAge Pro	16	57%	6	7	3	
WinBUGS	6	21%	1	2	3	
R	5	18%	1	2	2	
Arena	3	11%	0	2	1	
SAS	3	11%	0	1	2	
Crystal Ball	2	7%	1	0	1	
Simu8	2	7%	1	0	1	
STATA	1	4%	1	0	0	
RevMAN	1	4%	1	0	0	
Borland	1	4%	1	0	0	
Delphi						
S-PLUS	1	4%	1	0	0	
@risk	1	4%	0	0	1	
STELLA	0	0%	0	0	0	
Witness	0	0%	0	0	0	

Software Packages for Graphical Models

Written by Kevin Murphy.

Last updated 16 June 2014.

(Thanks to Alex Gorban for helping me with the switch to Google Sheets.)

Review articles

- List of GM code at MLOSS
- Click here for a short article I wrote for the ISBA (International Society for Bayesian Analysis) Newsletter, December 2007, sumarizing some of the packages below.
- Click here for a more detailed discussion of some of these packages written by Ann Nicholson and Kevin Korb in 2004.
- Click here for a French version of my comparison table (not necessarily up-to-date).

What do the headers in the table mean?

- Src = source code included? (N=no) If so, what language?
- Cts = are continuous (latent) nodes supported? G = (conditionally) Gaussians nodes supported analytically, Cs = continuous nodes supported by sampling, Cd = continuous nodes supported by discretization, Cx = continuous nodes supported by some unspecified method, D = only discrete nodes supported.
- GUI = Graphical User Interface included?
- · Learns parameters?
- Learns structure? CI = means uses conditional independency tests
- Utility = utility and decision nodes (i.e., influence diagrams) supported?
- Free? 0 = free (although possibly only for academic use). \$ = commercial software (although most have free versions which are restricted in various ways, e.g., the model size is limite or models cannot be saved, or there is no API.)
- Undir? What kind of graphs are supported? U = only undirected graphs, D = only directed graphs, UD = both undirected and directed, CG = chain graphs (mixed directed/undirected).
- Inference = which inference algorithm is used? jtree = junction tree, varelim = variable (bucket) elimination, MH = Metropols Hastings, G = Gibbs sampling, IS = importance sampling sampling = some other Monte Carlo method, polytree = Pearl's algorithm restricted to a graph with no cycles, VMP = variational message passing, EP = expectation propagation, SL = the program is designed for structure learning from completely observed data, not state estimation
- Comments. If in "quotes", I am quoting the authors at their request.

If you want your package to be listed, please fill out this form.

	Name	Authors	Src	Cts	GUI	Params	Struct	Utility	Free	Undir	Inference	Comments	
	<												
	<u>AgenaRisk</u>	Agena	N	Сх	Υ	Υ	N	N	\$	D	JTree	Simulation by Dynamic discretisation	
	<u>Analytica</u>	Lumina	N	G	Y	N	N	Υ	\$	D	sampling	spread sheet compatible	
	B-course	U. Helsinki	N	Cd	Y	Υ	Υ	N	0	D	?	Runs on their server: view results us	
	<u>Banjo</u>	Hartemink	Java	Cd	N	N	Υ	N	0	D	none	structure learning of static or dynami	
_[Daggiot	II Heleinki	CII	_	NI	V	M	NI.	0	n	MIT	Caparatas C.L. for MCMC /No long.	

Open-source tools for PGMs

	Weka	JavaBayes	Elvira	BNT	Riso	UnBBayes	OpenMarkov	BayesLine	PNL	BNJ	OBP
Start	1993	1996	1997	1999	2000	2000	2002	2003	2003	2004	2006
Stopped		2001	2010	2007	2004	2014		2003	2005	2004	2007
Programming language	Java	Java	Java	Matlab	Java	Java	Java	Java	C++	Java	Python
License	GPL	GPL	?	GPL	GPL	GPL	GPL	LGPL	IOSL	GPL	GPL
Bayesian networks	yes	yes	yes	yes	yes	yes	yes	yes	yes	yes	yes
Influence diagrams	no	no	yes	yes	no	yes	yes	no	no	no	no
Dynamic/Markov models	no	no	no	yes	no	no	yes	no	no	no	no
User manuals	yes	yes	yes	yes	yes	yes	yes	no	no	yes	yes
Developer manuals	yes	no	no	no	no	yes	yes	no	no	no	no
Users list/forum	yes	no	no	yes	yes	yes	yes	yes	no	yes	yes
Developers list/forum	yes	no	yes	yes	yes	yes	yes	yes	no	yes	yes
Source HTML docs	yes	yes	yes	no	yes	yes	yes	yes	no	no	no
Version control	yes	no	yes	no	yes	yes	yes	yes	no	yes	yes
Bug tracker	yes	yes	no	no	yes	yes	yes	yes	no	yes	yes

- Only <u>BNT</u> and <u>OpenMarkov</u> can represent <u>Markov models</u>.
- Among the tools having a GUI for editing PGMs, only
 Weka and OpenMarkov are still under active development.

Software tools for PGMs

Type of PGM	Application	Software tools			
Bayesian networks	Diagnosis	Many tools			
Dayesian networks	Statistical inference	BUGS, Stan			
Influence diagrams	Decision analysis for unicriterion problems	Many tools			
G	Cost-effectiv. analysis	OpenMarkov			
Decision analysis networks (DANs)	Decision analysis with partially-ordered decisions	OpenMarkov			
MIDs	Cost-effectiv. analysis in temporal problems	OpenMarkov			
Markov decision processes	Planning (e.g., robotics)	Several tools			

OpenMarkov. Main features

- ◆ Main advantage: open source
 - Free
 - Users can adapt it to their needs
 - Software engineering tools: JUnit, maven, mercurial (bitbucket), nexus, bugtracker, etc.

Strengths

- Written in Java: portability (Windows, linux, MacOS...)
- Many types of models, potentials, etc.
- Algorithms not available in any other package
 - CEA with IDs
 - interactive learning
- Very active: new features are continuously added
- Support for users and developers: wiki, lists, mail...
- Well-documented format for encoding networks: ProbModelXML.

OpenMarkov. Limitations

- Main weakness
 - Still a prototype: needs debugging
- Other weaknesses
 - Written in Java: relatively slow (in some cases)
 - No on-line help, documentation still poor
 - Support is limited, due to scarcity of human resources.

8. Conclusions

Conclusions

- ♦ BNs overcame the limitations of the naïve Bayes method.
- ◆ IDs have several advantages over decision trees, but also have serious limitations for medical decision making.
- DANs are similar to IDs, but more suitable for asymmetric decision problems, especially partially ordered decisions.
- ♦ It is possible to do cost-effectiveness analysis with IDs.
- and also with Markov IDs (MIDs) if all decisions are atemporal.
- There are other types of Markov PGMs having one or more decisions per cycle: MDPs, POMDPs, DLIMIDs...

How to bring PGMs from artificial intelligence into medical decision making

- Develop powerful user-friendly software tools
- Dissemination
 - Seminars, short courses...
 - Tutorials and textbooks written in the language of clinicians, epidemiologists and health economists

Research

- New methods for the representation of knowledge
- New algorithms for CEA, sensitivity analysis...
- Discrete event simulation with PGMs

Thank you very much for your attention!

- ◆ Links
 - www.cisiad.uned.es
 - www.OpenMarkov.org
 - www.ProbModelXML.org/networks
- ◆ Contact: fjdiez@dia.uned.es