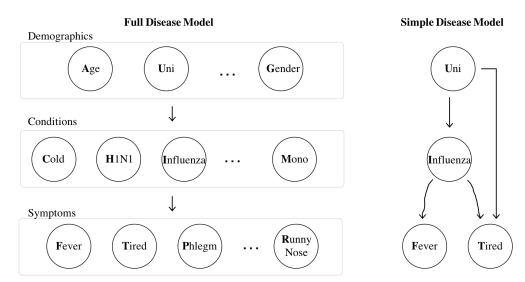
Bayesian Networks

At this point in the reader we have developed tools for analytically solving for probabilities. We can calculate the likelihood of random variables taking on values, even if they are interacting with other random variables (which we have called multi-variate models, or we say the random variables are jointly distributed). We have also started to study samples and sampling.

Consider the WebMD Symptom Checker. WebMD has built a probabilistic model with random variables which roughly fall under three categories: symptoms, risk factors and diseases. For any combination of observed symptoms and risk factors, they can calculate the probability of any disease. For example, they can calculate the probability that I have influenza given that I am a 21-year-old female who has a fever and who is tired: P(I=1|A=21,G=1,T=1,F=1). Or they could calculate the probability that I have a cold given that I am a 30-year-old with a runny nose: P(C=1|A=30,R=1). At first blush this might not seem difficult. But as we dig deeper we will realize just how hard it is. There are two challenges: (1) Modelling: sufficiently specifying the probabilistic model and (2) Inference: calculating any desired probability.

Bayesian Networks

Before we jump into how to solve probability (aka inference) questions, let's take a moment to go over how an expert doctor could specify the relationship between so many random variables. Ideally we could have our expert sit down and specify the entire "joint distribution" (see the first lecture on multi-variable models). She could do so either by writing a single equation that relates all the variables (which is as impossible as it sounds), or she could come up with a joint distribution table where she specifies the probability of any possible combination of assignments to variables. It turns out that is not feasible either. Why? Imagine there are N=100 binary random variables in our WebMD model. Our expert doctor would have to specify a probability for each of the $2^N>10^{30}$ combinations of assignments to those variables, which is approaching the number of atoms in the universe. Thankfully, there is a better way. We can simplify our task if we know the "generative" process that creates a joint assignment. Based on the generative process we can make a data structure known as a **Bayesian Network**. Here are two networks of random variables for diseases:



For diseases the flow of influence is directed. The states of "demographic" random variables influence whether someone has particular "conditions", which influence whether someone shows particular "symptoms". On the right is a simple model with only four random variables. Though this is a less interesting model it is easier to understand when first learning Bayesian Networks. Being in university (binary) influences whether or not someone has influenza (binary). Having influenza influences whether or not someone has a fever (binary) and the state of university and influenza influences whether or not someone feels tired (also binary).

In a Bayesian Network an arrow from random variable X to random variable Y articulates our assumption that X directly influences the likelihood of Y. We say that X is a *parent* of Y. To fully define the Bayesian network we must provide a way to compute the probability of each random variable (X_i) conditioned on knowing the value of all their parents: $P(X_i = k | \text{Parents of } X_i \text{ take on specified values})$. Here is a concrete example of what needs to be defined for the simple disease model. Recall that each of the random variables is binary:

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P(\text{Uni}=1)=0.8 \\ P(\text{Influenza}=1|\text{Uni}=1)=0.2 \\ P(\text{Influenza}=1|\text{Uni}=0)=0.1 \\ P(\text{Tired}=1|\text{Uni}=0,\text{Influenza}=0)=0.1 \\ P(\text{Tired}=1|\text{Uni}=1,\text{Influenza}=0)=0.8 \\ P(\text{Tired}=1|\text{Uni}=1,\text{Influenza}=1)=0.9 \\ P(\text{Tired}=1|\text{Uni}=1,\text{Influenza}=1)=0.9 \\ P(\text{Tired}=1|\text{Uni}=1,\text{Influenza}=1)=1.0 \\ P(\text{Uni}=1,\text{Influenza}=1)=1.0 \\ P(\text{Uni}=1,\text{Influenza}=1)=1.0 \\ P(
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Let's put this in programming terms. All that we need to do in order to code up a Bayesian network is to define a function: **getProbXi(i, k, parents)** which returns the probability that X_i (the random var with index **i)** takes on the value **k** given a value for each of the parents of X_i encoded by the list **parents**: $P(X_i = x_i|V)$ alues of parents of X_i)

Deeper understanding: The reason that a Bayes Net is so useful is that the "joint" probability can be expressed in exponentially less space as the product of the probabilities of each random variable conditioned on its parents! Without loss of generality, let X_i refer to the ith random variable (such that if X_i is a parent of X_j then i < j):

$$P(\mathrm{Joint}) = P(X_1 = x_1, \dots, X_n = x_n) = \prod_i P(X_i = x_i | \mathrm{Values \ of \ parents \ of} \ X_i)$$

What assumptions are implicit in a Bayes Net? Using the chain rule we can decompose the **exact** joint probability for n random variables. To make the following math easier to digest I am going to use x_i as shorthand for the event that $X_i = x_i$:

$$P(x_1,\ldots,x_n) = \prod_i P(x_i|x_{i-1},\ldots,x_1)$$

By looking at the difference in the two equations, we can see that a Bayes Net is assuming that

$$P(x_i|x_{i-1},\ldots,x_1) = P(x_i|\text{Values of parents of }X_i)$$

This is a conditional independence statement. It is saying that once you know the value of the parents of a variable in your network, X_i , any further information about non-descendents will not change your belief in X_i . Formally we say that X_i is conditionally independent of its non-descendents, given its parents. What is a non-descendent again? In a graph, a descendent of X_i is anything which is in the subtree that starts at X_i . Everything else is a non-descendent. Non-descendents include the "ancestor" nodes of X_i as well as nodes which are totally unconnected to X_i . When designing Bayes Nets you don't have to think about this assumption directly. It turns out to be a naturally good assumption if the arrows between your nodes follow a causal path.

Designing a Bayes Net

There are several steps to designing a Bayes Net.

- 1. Choose your random variables, and make them nodes.
- 2. Add edges, often based off your assumptions about which nodes directly cause which others.
- 3. Define $P(X_i = x_i | \text{Values of parents of } X_i)$ for all nodes.

As you might have guessed, we can do step (2) and (3) by hand, or, we can have computers try and perform those tasks based on data. The first task is called "structure learning" and the second is an instance of "machine learning." There are fully autonomous solutions to structure learning—but they only work well if you have a massive amount of data. Alternatively people will often compute a statistic called correlation between all pairs of random variables to help in the art form of designing a Bayes Net.

In the next part of the reader we are going to talk about how we could learn $P(X_i = x_i | \text{Values of parents of } X_i)$ from data. For now let's start with the (reasonable) assumption that an expert can write down these functions in equations or as python: **getProbXi**.

Next Steps

Great! We have a feasible way to define a large network of random variables. First challenge complete. We haven't talked about continuous or multinomial random variables in Bayes Nets. None of the theory changes: the expert will just have to define **getProbXi** to handle more values of **k** than 0 or 1.

A Bayesian network is not very interesting to us unless we can use it to solve different conditional probability questions. How can we perform "inference" on a network as complex as a Bayesian network?