Bayesian network internals Inference algorithms, time series & distributed learning with Big Data

Dr John Sandiford, CTO Bayes Server



Contents

- Introduction
- What is a Bayesian network?
- What is inference?
- Probability
- Bayesian network inference
- Inference with time series
- Distributed parameter learning



Introduction



Profile

linkedin.com/in/johnsandiford

- PhD Imperial College Bayesian networks
- Machine learning 15 years
 - Implementation
 - Application
 - Numerous techniques
- Algorithm programming even longer
 - Scala , C#, Java, C++
- Graduate scheme mathematician (BAE Systems)
- Artificial Intelligence / ML research program 8 years (GE/USAF)
- BP trading & risk analytics big data + machine learning
- Also: NYSE stock exchange, hedge fund, actuarial consultancy, international newspaper



What is a Bayesian network?

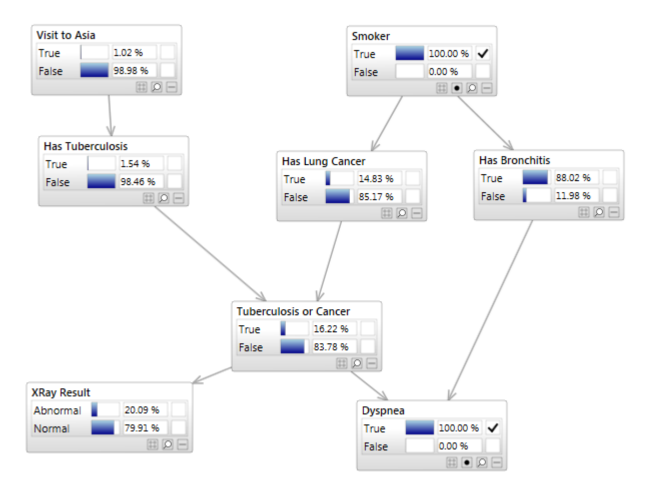


What is a Bayesian network?

- DAG directed acyclic graph
- Nodes, links, probability distributions
- Each node requires a probability distribution conditioned on its parents (if any)

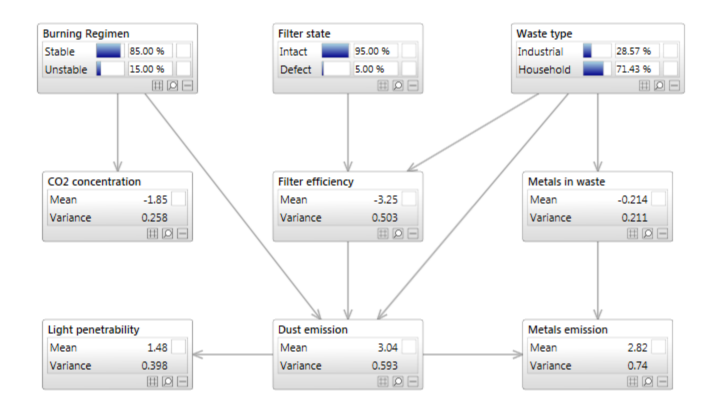


Example – Asia network



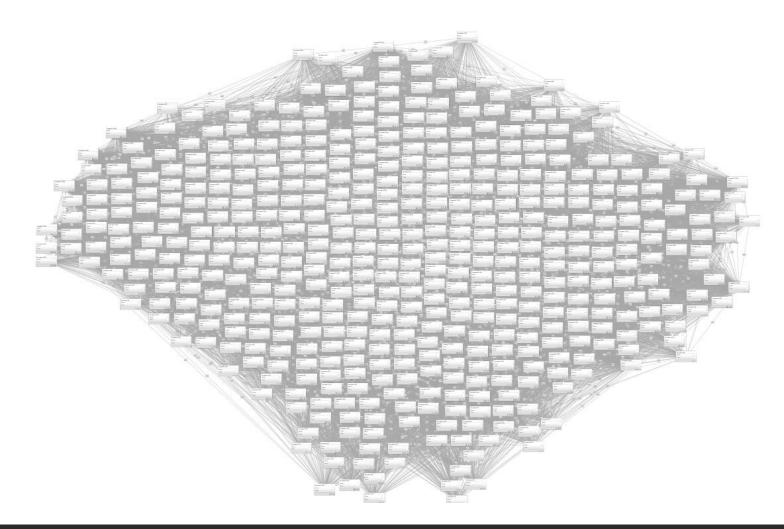


Example – Waste network



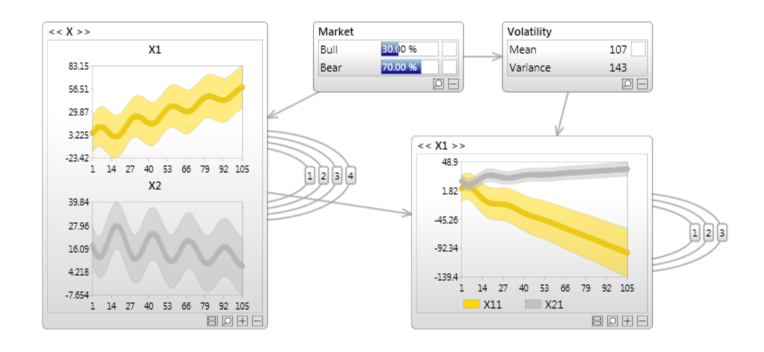


Example – the bat (40,000 links)



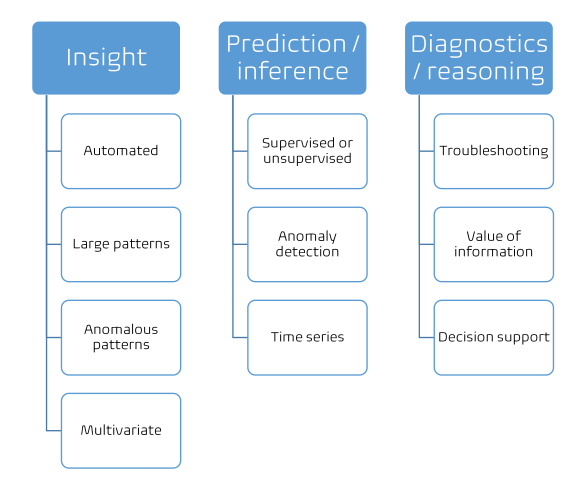


Example – static & temporal





Insight, prediction & diagnostics





What is inference?



Inference

- Asking a question given things you already know
- Encompasses prediction, reasoning & diagnostics
- Given a number of symptoms, which diseases are most likely?
- How likely is it that a component will fail, given the current state of the system?
- Given recent behaviour of 2 stock, how will they behave together for the next 5 time steps?
- Handles missing data



Exact & approximate

- Exact inference
 - Applicable to a large range of problems, but not all
 - May not be possible when combinations/paths get large
 - Correct answer subject to rounding errors
- Approximate inference
 - Wider class of problems
 - Deterministic / non deterministic
 - No guarantee of correct answer



Exact inference

- We will discuss exact inference
- Many concepts apply to both



Probability



Probability notation

- P(A)
- P(A|B) Conditional probability (probability of A given B)
- P(A,B) Joint probability (probability of A and B)
- P(Head | Tail)
 - variables on the left are referred to as head, and variables on the right are referred to as tail
- P(A,B) = P(A|B)P(B) = P(B|A)P(A) =>
 - P(A|B) = P(B|A)P(A) / P(B)
 - This is Bayes theorem
 - Used during inference



Joint probability

- E.g. P(Raining, Windy)
- Sums to 1

| Raining | Windy = False | Windy = True |
|---------|---------------|--------------|
| False | 0.64 | 0.16 |
| True | 0.1 | 0.1 |



Marginalization

P(Raining, Windy)

| Raining | Windy = False | Windy = True | Sum |
|---------|---------------|--------------|-----|
| False | 0.64 | 0.16 | 0.8 |
| True | 0.1 | 0.1 | 0.2 |



P(Raining)

| Raining = False | Raining = True |
|-----------------|----------------|
| 0.8 | 0.2 |

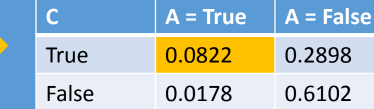
For discrete variables we sum, whereas for continuous variables we integrate

Marginalization – multiple variables

P(A,B,C,D)

| В | С | D | A = True | A = False |
|-------|-------|-------|----------|-----------|
| True | True | True | 0.0036 | 0.0054 |
| True | True | False | 0.0098 | 0.0252 |
| True | False | True | 0.0024 | 0.0486 |
| True | False | False | 0.0042 | 0.1008 |
| False | True | True | 0.0256 | 0.0864 |
| False | True | False | 0.0432 | 0.1728 |
| False | False | True | 0.0064 | 0.2016 |
| False | False | False | 0.0048 | 0.2592 |





Marginal probability P(A,C) created by marginalizing B and D from the joint probability P(A,B,C,D)

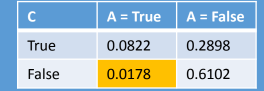


Multiplication

 $P(B,D \mid A, C)$

| В | С | D | A = True | A = False |
|-------|-------|-------|----------|-----------|
| True | True | True | 0.0438 | 0.0186 |
| True | True | False | 0.1192 | 0.0870 |
| True | False | True | 0.1348 | 0.0796 |
| True | False | False | 0.2360 | 0.1652 |
| False | True | True | 0.3114 | 0.2981 |
| False | True | False | 0.5255 | 0.5963 |
| False | False | True | 0.3596 | 0.3304 |
| False | False | False | 0.2697 | 0.4248 |





P(A,B,C,D)

| В | С | D | A = True | A = False |
|-------|-------|-------|----------|-----------|
| True | True | True | 0.0036 | 0.0054 |
| True | True | False | 0.0098 | 0.0252 |
| True | False | True | 0.0024 | 0.0486 |
| True | False | False | 0.0042 | 0.1008 |
| False | True | True | 0.0256 | 0.0864 |
| False | True | False | 0.0432 | 0.1728 |
| False | False | True | 0.0064 | 0.2016 |
| False | False | False | 0.0048 | 0.2592 |



Instantiation – (evidence)

P(A=False,B,C,D)

| В | С | D | A = True | A = False |
|-------|-------|-------|----------|-----------|
| True | True | True | 0.0036 | 0.0 |
| True | True | False | 0.0098 | 0.0 |
| True | False | True | 0.0024 | 0.0 |
| True | False | False | 0.0042 | 0.0 |
| False | True | True | 0.0256 | 0.0 |
| False | True | False | 0.0432 | 0.0 |
| False | False | True | 0.0064 | 0.0 |
| False | False | False | 0.0048 | 0.0 |



P(B, C, D)

| С | D | B=True | B=False |
|-------|-------|--------|---------|
| True | True | 0.0036 | 0.0256 |
| True | False | 0.0098 | 0.0432 |
| False | True | 0.0024 | 0.0064 |
| False | False | 0.0042 | 0.0048 |

Bayesian network inference



Joint probability – Bayesian network

- If we multiply all the distributions of a Bayesian network together, we get the joint distribution over all variables
- What can we do with the joint?
- Any evidence e is information we know (e.g. D=True)

$$P(\mathbf{X}, \mathbf{e}) = \sum_{\mathbf{U} \setminus \mathbf{x}} P(\mathbf{U}, \mathbf{e}) = \sum_{\mathbf{U} \setminus \mathbf{x}} \prod_{i} P(\mathbf{U}_{i} | pa(\mathbf{U}_{i})) \mathbf{e}$$

U = universe of variablesX = variables being predictede= evidence on any variables



Just use the joint over all variables?

- We could perform the same tasks if memory and time were not an issue.
- The problem?
 - Exponential increases in size with discrete variables
- The answer?
 - Bayesian network inference



Distributive law

$$if A \notin \mathbf{X}, A \in \mathbf{Y}, then \sum_{A} \phi_{\mathbf{X}} \phi_{\mathbf{Y}} = \phi_{\mathbf{X}} \sum_{A} \phi_{\mathbf{Y}}$$

This simply means that if we want to marginalize out the variable A we can perform the calculations on the subset of distributions that contain A

φ is a probability distribution over the variables in the subscript



Consider calculating P(A|D=True)



$$P(A|\mathbf{e}) \propto \sum_{B,C,D} P(A)P(B|A)P(C|B)P(D|C)e_D$$
 Distributive law

$$P(A|\mathbf{e}) \propto P(A) \sum_{B} P(B|A) \sum_{C} P(C|B) \sum_{D} P(D|C) e_{D}$$



Elimination order

- The order in which marginalization is performed is called an elimination order.
- Many different possible orders
- NP hard
- A number of algorithms exist to determine orderings that work well in practise
 - E.g. pick the variable(s) that result in the smallest distribution to be marginalized at each step
 - Multiple variables can be eliminated at each step.

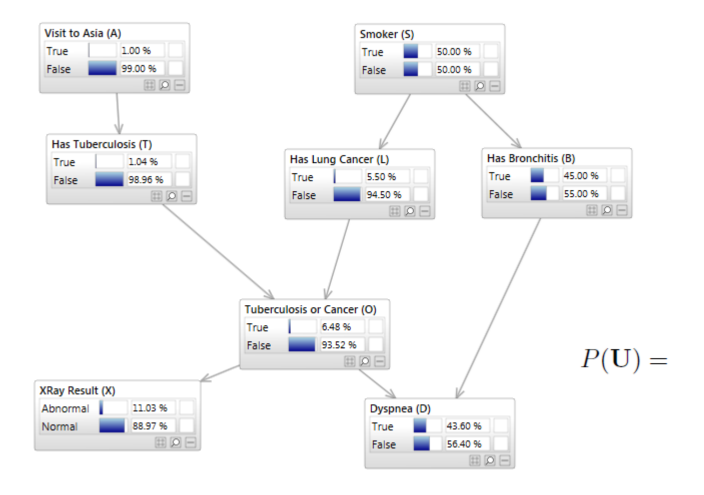
$$P(A|\mathbf{e}) \propto P(A) \sum_{B} P(B|A) \sum_{C} P(C|B) \sum_{D} P(D|C) e_{D}$$

Junction trees

- What if we want to predict all variables, not just A?
- We could use the previous procedure known as Variable Elimination multiple times.
- Or we can use a junction tree (join tree)
 - Simply the tree formed by eliminating all variables in the same way as before



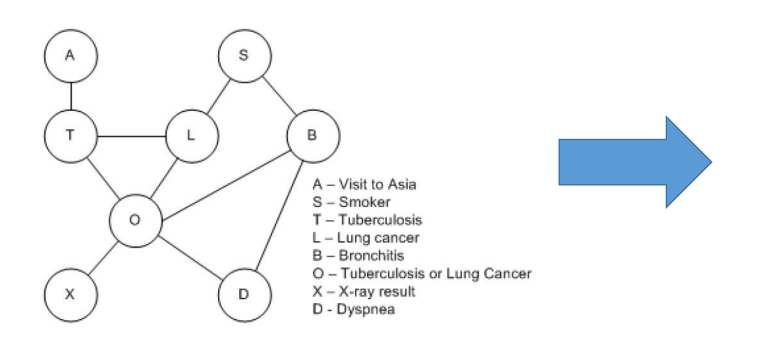
Asia network

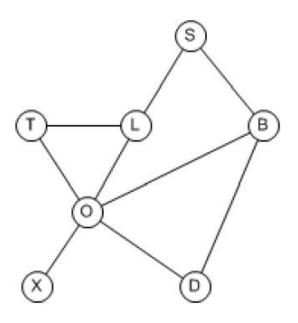


| Node | Distribution |
|------|--------------|
| A | P(A) |
| T | P(T A) |
| S | P(S) |
| L | P(L S) |
| B | P(B S) |
| O | P(O T,L) |
| X | P(X O) |
| D | P(D O,B) |

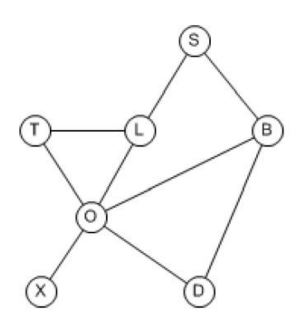
$$P(A)P(S)P(T|A)P(L|S) \times$$

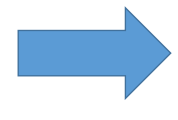
$$P(B|S)P(O|T,L)(X|O)P(D|B,O)$$

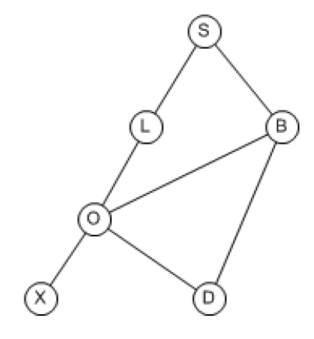




Domain graph after elimination of A

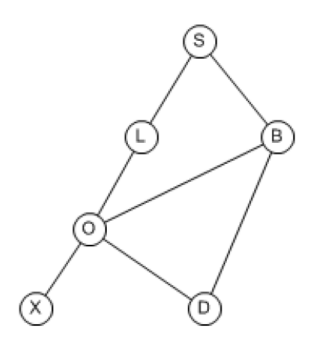




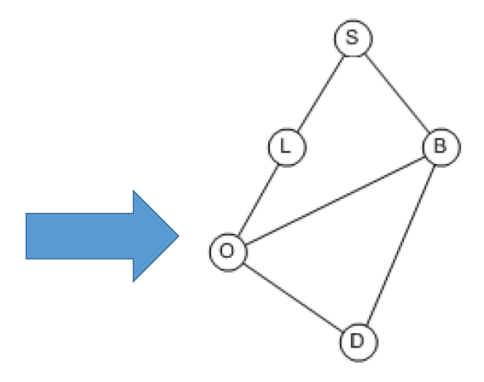


Domain graph after elimination of A

Domain graph after elimination of ${\cal T}$

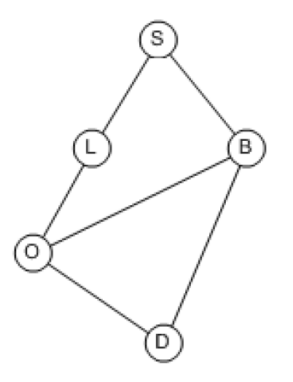


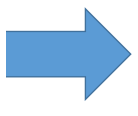


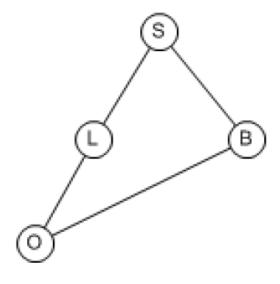


Domain graph after elimination of X



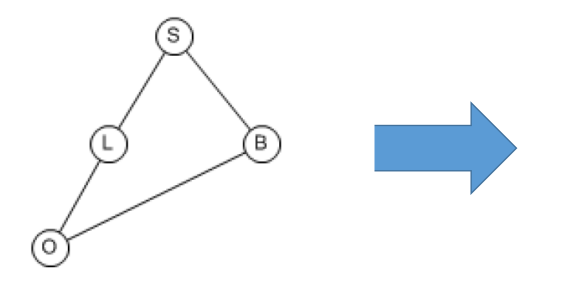


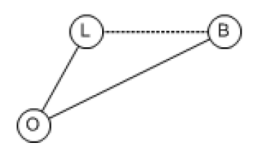




Domain graph after elimination of D

Domain graph after elimination of X



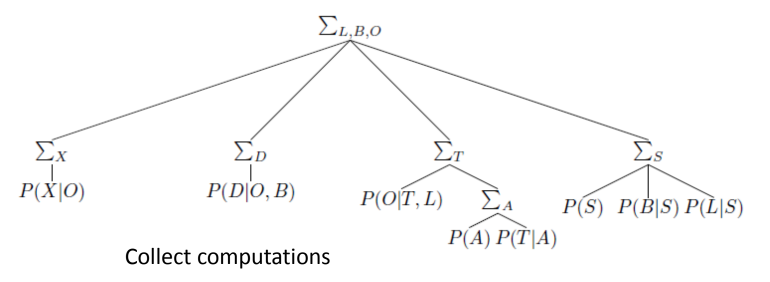


7.: Domain graph after elimination of S. The dotted line is a required fill-in.

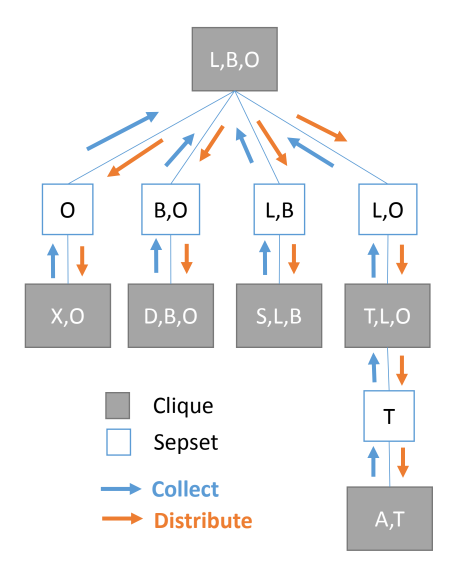
Domain graph after elimination of D

The complete elimination order is... {A}, {T}, {X}, {D}, {S}, {L,B,O}

Junction trees

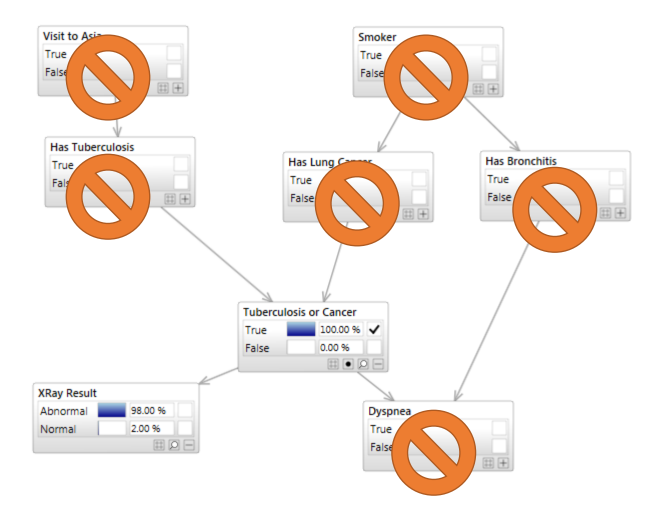


- Collect towards the root {L,B,O} similar to variable elimination
- Distribute from the root {L,B,O} back to the leaves allows us to calculate all marginals– P(A), P(X), P(B), P(L) etc...





Relevance – Bayes ball algorithm

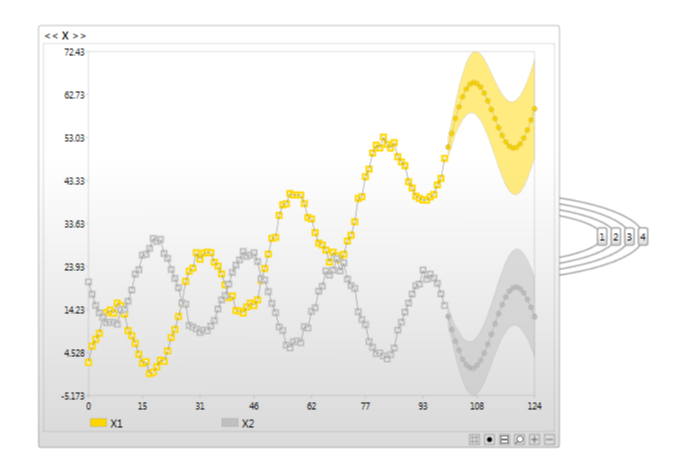




Inference with time series -Dynamic Bayesian networks

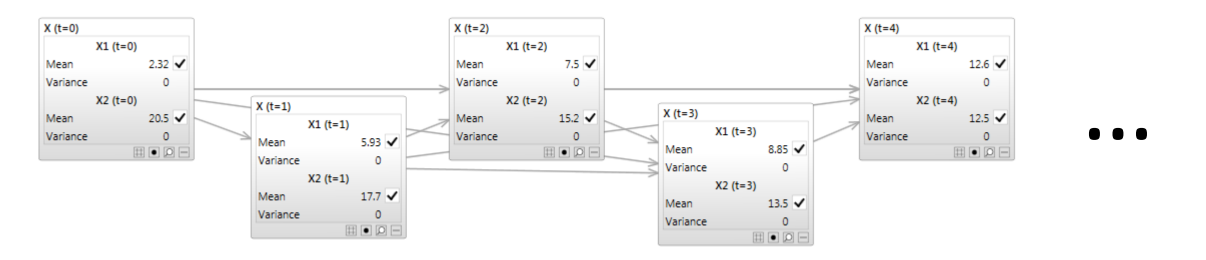


Dynamic Bayesian networks





Unrolling



We could unroll, and use standard methods



Distributions that understand time

| | | X1[t] | X2[t] |
|---|--------------------|----------------|----------------|
| • | Intercept | 3.076214646583 | -1.58979124120 |
| | Covariance (X1[t]) | 4.142028922619 | -1.63113437658 |
| | Covariance (X2[t]) | -1.63113437658 | 2.023002098810 |
| | Weight (X1[t-1]) | 0.995368300968 | -0.00816950459 |
| | Weight (X2[t-1]) | 0.026861977953 | 0.942548514594 |

Note that X1 appears in the same distribution twice, but at different times

Time aware distributions

- Marginalization & multiplication is well defined
- We can use all the existing algorithms
- We can construct queries like...
- P(X1@t=4)
 - Returns probabilities for discrete, mean & variance for continuous
- P(X1@t=4, X2@t=4)
 - Joint time series prediction (funnel)
- P(X1@t=2, X1@t=3)
 - Across different times
- P(A, X1@t=2)
 - Mixed static & temporal
- Log-likelihood of a multivariate time series
 - Anomaly detection



Distributed parameter learning



Different types of scalability

Data size

Big data?

Connectivity

(discrete -> exponential)

Network size,

Rephil > 1M nodes

Inference (distributed)



Apache Spark

RDD Objects

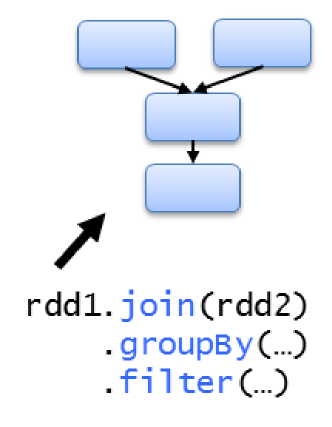
Spark SQL Spark Streaming

MLlib (machine learning) GraphX (graph)

Apache Spark









Apache Spark

- RDD (Resilient distributed dataset)
- Can be in memory
- Code automatically converted to DAG execution engine
- Serialization of variables



Distributed architecture

- 1. Distribute Bayesian network to worker nodes
- 2. Calculate the sufficient statistics per partition
 - This often requires an inference algorithm per thread/partition
- 3. Aggregate the sufficient statistics (reduce)
- 4. Update the Bayesian network based on the new statistics
- 5. Return to 1 until convergence



Distributed parameter learning

Node 1

| Х | Υ | X | Υ | х | Υ |
|------|------|------|------|------|-----|
| 10.2 | 12.5 | 3.4 | 3.2 | 2.0 | 4.0 |
| 14.5 | 3.2 | 6.6 | 1.6 | 8.1 | 3.4 |
| 15.6 | 8.2 | 5.5 | 4.3 | 2.2 | 7.7 |
| 9.2 | 12.2 | 12.4 | 8.9 | 15.1 | 1.2 |
| 15.8 | 9.2 | -1.1 | -2.4 | 4.6 | 4.5 |
| 4.5 | 2.1 | 4.5 | 4.2 | 2.4 | 1.9 |
| | | | | | |

 \sum stats \sum stats

Node 2

| X | Υ | Х | Υ | X | Υ |
|------|------|------|------|------|-----|
| 10.2 | 12.5 | 3.4 | 3.2 | 2.0 | 4.0 |
| 14.5 | 3.2 | 6.6 | 1.6 | 8.1 | 3.4 |
| 15.6 | 8.2 | 5.5 | 4.3 | 2.2 | 7.7 |
| 9.2 | 12.2 | 12.4 | 8.9 | 15.1 | 1.2 |
| 15.8 | 9.2 | -1.1 | -2.4 | 4.6 | 4.5 |
| 4.5 | 2.1 | 4.5 | 4.2 | 2.4 | 1.9 |
| | | | | | |

∑ stats

Node 3

| х | Υ | Х | Υ |
|------|------|------|------|
| 10.2 | 12.5 | 3.4 | 3.2 |
| 14.5 | 3.2 | 6.6 | 1.6 |
| 15.6 | 8.2 | 5.5 | 4.3 |
| 9.2 | 12.2 | 12.4 | 8.9 |
| 15.8 | 9.2 | -1.1 | -2.4 |
| 4.5 | 2.1 | 4.5 | 4.2 |
| | | | |

∑ stats

∑ stats

Σ stats Σ stats

∑ stats

Σ stats Σ stats Σ stats Σ stats Σ stats Σ stats



∑ stats

Distributed parameter learning





Example – distributed learning

```
val sc = new SparkContext(conf)
// hard code some test data. Normally you would read data from your cluster.
val data = createRDD.cache()
// A network could be loaded from a file or stream
// we create it manually here to keep the example self contained
val network = createNetwork
val parameterLearningOptions = new ParameterLearningOptions
// Bayes Server supports multi-threaded learning
// which we want to turn off as Spark takes care of this
parameterLearningOptions.setMaximumConcurrency(1)
/// parameterLearningOptions.setMaximumIterations(...) // this can be useful to limit the number of iterations
val config = new MemoryNameValues // we could also use broadcast variables
val output = ParameterLearning.learnDistributed(network, parameterLearningOptions,
 new BayesSparkDistributer[Seq[(Double, Double)]](
    data.
   config,
    (ctx, iterator) => new TimeSeriesEvidenceReader(ctx.getNetwork, iterator)
```



Distributed time series prediction

```
// make some time series predictions into the future
val predictions = Prediction.predict[TimeSeries](
  network.
  testData,
  Seq(
    PredictVariable("X1", Some(PredictTime(5, Absolute))), PredictVariance("X1", Some(PredictTime(5, Absolute))),
    PredictVariable("X2", Some(PredictTime(5, Absolute))), PredictVariance("X2", Some(PredictTime(5, Absolute))),
    PredictVariable("X1", Some(PredictTime(6, Absolute))), PredictVariance("X1", Some(PredictTime(6, Absolute))),
    PredictVariable("X2", Some(PredictTime(6, Absolute))), PredictVariance("X2", Some(PredictTime(6, Absolute))),
    PredictLogLikelihood() // this value can be used for Time Series anomaly detection
  (network, iterator) => new TimeSeriesReader(network, iterator))
predictions.foreach(println)
```



Thankyou

