

A PROBABILISTIC APPROACH TO DATA
INTEGRATION
IN BIOMEDICAL RESEARCH: THE IsBIG
EXPERIMENTS

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To my lovely daughters

Ila and Neha

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ABSTRACT

Vibha Anand

A PROBABILISTIC APPROACH TO DATA INTEGRATION IN BIOMEDICAL RESEARCH: THE IsBIG EXPERIMENTS

Biomedical research has produced vast amounts of new information in the last decade but has been slow to find its use in clinical applications. Data from disparate sources such as genetic studies and summary data from published literature have been amassed, but there is a significant gap, primarily due to a lack of normative methods, in combining such information for inference and knowledge discovery.

In this research using Bayesian Networks (BN), a probabilistic framework is built to address this gap. BN are a relatively new method of representing uncertain relationships among variables using probabilities and graph theory. Despite their computational complexity of inference, BN represent domain knowledge concisely. In this work, strategies using BN have been developed to incorporate a range of available information from both raw data sources and statistical and summary measures in a coherent framework. As an example of this framework, a prototype model (In-silico Bayesian Integration of GWAS or IsBIG) has been developed. IsBIG integrates summary and statistical measures from the NIH catalog of genome wide association studies (GWAS) and the database of human genome variations from the international HapMap project. IsBIG produces a map of disease to disease associations as inferred by genetic linkages in the population.

Quantitative evaluation of the IsBIG model shows correlation with empiric results from our Electronic Medical Record (EMR) – The Regenstrief Medical Record System

(RMRS). Only a small fraction of disease to disease associations in the population can be explained by the linking of a genetic variation to a disease association as studied in the GWAS. None the less, the model appears to have found novel associations among some diseases that are not described in the literature but are confirmed in our EMR. Thus, in conclusion, our results demonstrate the potential use of a probabilistic modeling approach for combining data from disparate sources for inference and knowledge discovery purposes in biomedical research.

Mathew J. Palakal, Ph.D., Chair

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Chapter 1 INTRODUCTION

Opportunities for Informatics in Biomedical Research

The past few years have witnessed major advances in the area of biomedical research due to rapid advances in technology like database management and the availability of open source software tools to name a few. Due to the latter, the research in this area has become increasingly collaborative and several major initiatives including the mapping of entire human genome have been successfully completed in the last decade.

[1]

Informatics, which is viewed as a science of information, is often studied as a branch of computer science and information technology relating to databases, ontology and software engineering and is primarily concerned with transformation of information by computation or communication; by machines or people. Health informatics or Biomedical informatics is an emerging discipline engaged in study, invention and implementation of *structures and algorithms* to improve understanding and management of medical information. The end objective of biomedical informatics is coalescing of data, knowledge and the tools necessary to apply the data and knowledge in the decision making process at the time and place that a decision needs to be made.

Thus, in the post genome era the role of biomedical informatics has shifted from managing and integrating genetic sequence databases to discovering knowledge from biomedical databases. More recently integrating this knowledge from disparate sources such as from biological databases and clinical data from electronic medical records (EMR) for applications such as personalized medicine has received an increasing amount of interest from both the National Institutes of Health (NIH) and individual researchers.

Challenges for Informatics in Biomedical research

However, in biomedical research, to make inferences based on data especially using traditional statistical methods, one requires a unified dataset, i.e. a dataset where all variables are measured on the same set of individuals. Furthermore, making new discoveries depends on having access to these original datasets. However, in the current research paradigm the variables of interest are being measured in separate studies and on different study populations. They are being stored in silos of specialized databases that do not relate to each other on an individual level. Therefore a significant gap exists in our ability to draw inference from these datasets in order to further our understanding of the outcomes of such research and its applicability for instance to clinical care.

For example, Figure 1-1 on the next page shows a model for a common disease like asthma that is known to have many causes. To gain a full understanding of the disease and its management, one has to account for all the causes –environmental, clinical, genetic, socio economic and demographic factors along with any sub clinical symptoms (phenotypes) that may be presented. Thus asthma presents as a common but a complex disease involving many risk factors. To apply cutting edge research to this disease in a clinical application, for example, on how environment or genes may affect an individual's disease status, one needs to integrate all such information in a coherent model and draw inference from it. Thus, finding methods to integrate information from disparate sources – biological databases, clinical databases, and published literature in a coherent model for purposes of prediction and eventually pre-emption of disease has become the goal of biomedical informatics researchers in this decade.

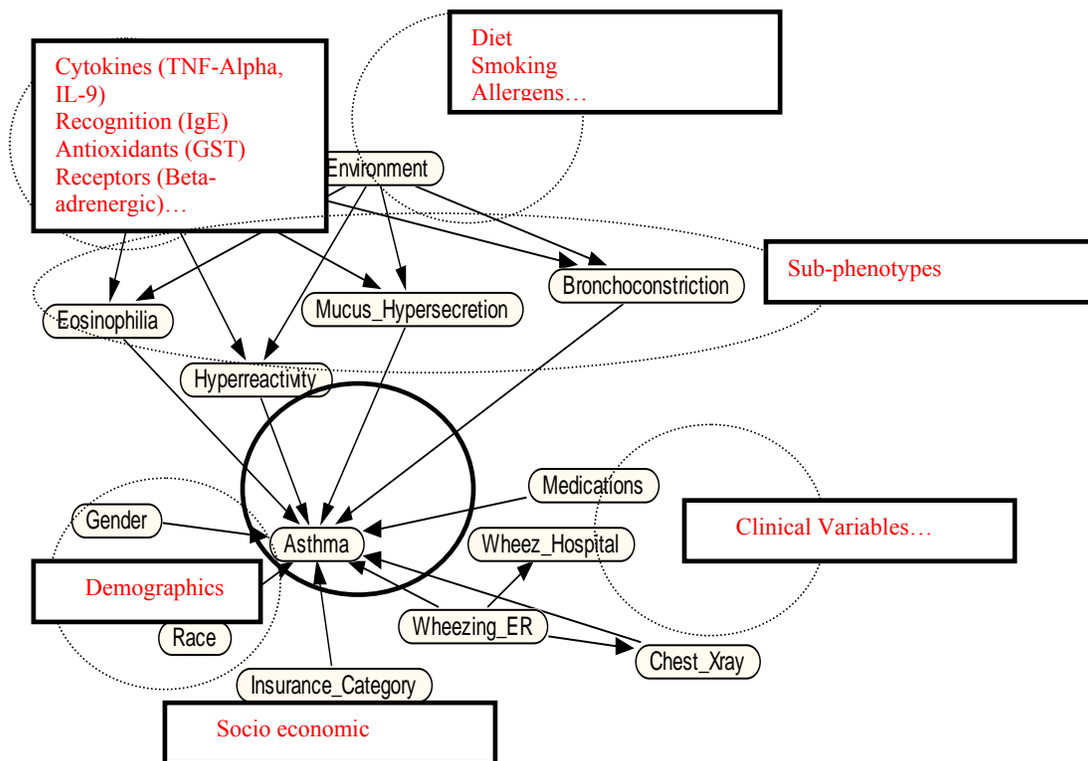


Figure 1-1 A complex dataset example

However, at present no such coherent model can be built because data collected from disparate study populations reside in silos of biomedical databases, with each of them focused on one of a number of causes, for example, how environmental factors like tobacco smoke exposure may affect asthma. Due to lacking unified datasets, our best hope of linking information is by using informatics tools that employ non-traditional statistical methods, for example, to combine information from available datasets such as EMR with sources such as summary or statistical measures from published literature.

In this work, we approach the integration problem with probabilistic modeling tools. We outline a methodology to move beyond the boundary of a dataset that is limited by a set of variables. Assuming independence of causal influences (ICI) among many causes that lead to a common effect, we strategically combine disparate sources of information with a Bayesian Network (BN) framework for identifying associations among the disparate datasets. Our approach uses available summary and statistical measures of correlations (r^2) and odds ratio (OR) from published literature when no unifying dataset is present to build a model that integrates information in a systematic and normative form for further knowledge discovery. Figure 1.2 outlines our conceptual model for data-information-knowledge discovery.

Figure 1-2 In-silico Bayesian Integration – Conceptual Model

We demonstrate our approach by integrating information from at least two disparate sources in a coherent model - 1) statistical correlation on genetic linkages (associations) between Single Nucleotide Polymorphisms (SNP) in the human genome, and 2) data from multiple genome wide association studies (GWAS) where the magnitude of the association between a SNP and a disease phenotype is measured as an odds ratio (OR) in each GWAS. SNPs and diseases are modeled as nodes in a BN and the edges that connect the nodes represent the strength of the relationship between nodes (i.e. SNP to SNP and SNP to disease). We demonstrate that as the effect of the SNP nodes from this model are averaged out, i.e. absorbed out, a disease to disease association map emerges as inferred from genetic linkages and discovered by the integration of these two disparate sources. We call the methodology In-silico Bayesian Integration and the model In-silico Bayesian Integration of GWAS (IsBIG).

Thus, IsBIG combines information from various GWAS in a coherent model which otherwise is not available from a unified dataset. IsBIG therefore also presents a qualitative and quantitative structure that can be used for further knowledge discovery, for example, for generating new hypotheses for future studies associating diseases as inferred by genetic linkages.

This thesis is organized into several major chapters. We first introduce the background of this research in Chapter 2. We describe existing methods for statistical analysis and modeling that have been employed for such research and their limitations. We then describe probabilistic modeling methods as a knowledge representation tool and give a brief literature review of their use in modeling healthcare data with emphasis on Bayesian networks.

In Chapter 3, we describe our data integration strategy using a series of experimental studies within the clinical domain. We learn BN from large clinical datasets and compare their performance with an expert's version to assess the feasibility of this modeling technique. When the data available are sparse, we use the causal independence assumption using the Noisy-OR formalism to learn the conditional probability distributions in our model BN. We apply the above to a feasibility study in the area of childhood asthma case finding from our electronic medical record (EMR), the Regenstrief Medical Record System (RMRS), [2] and find that the results are comparable to an expert's model in real world datasets. To model domain causal relationships, over and above causal independence, we test a recently published algorithm – Recursive Noisy OR (RNOR) and evaluate it with our previous childhood asthma case finding application. We find no statistically significant differences between the RNOR and causal independence approaches with this real world dataset. Therefore we stick to using the causal independence approach as a data integration strategy for successive experiments.

In Chapter 4, we extend beyond our clinical domain to apply the causal independence assumption to an experimental study where data from our EMR for childhood asthma is integrated with statistical and summary data published in one study of asthma, linking a genotype and environmental tobacco smoke exposure to the risk of the disease. We develop this approach into a formal method – In-silico Bayesian Integration and demonstrate its applicability to generate a phenotype to phenotype map (IsBIG) from statistical and summary data on diseases and / or traits linked to Single Nucleotide Polymorphisms (SNPs) found in Genome Wide Association Studies (GWAS).

In Chapter 5 we empirically evaluate the IsBIG model using data derived from our EMR, the Regenstrief Medical Record System (RMRS) [2] and literature search.

In Chapter 6, we summarize our work, discuss limitations of our approach as a knowledge representation tool and data integration strategy and outline some possible future directions.

Chapter 2 **BACKGROUND**

This chapter introduces the background of our research, including a brief introduction on Bayesian Networks and their comparison to other statistical and machine learning techniques and their applicability to our research as a knowledge representation tool for building a probabilistic framework for biomedical research.

Bayesian Networks (BN)

Bayesian networks are a modeling and inference tool for problems involving uncertainty. They have been shown to represent domain knowledge with natural perception of cause and effect. [3] A BN is a graphical model that both represents a qualitative structure and encodes quantitative parameters of the structure by defining a unique probability distribution. Because of their concise representation and their ability for belief propagation; BN have been widely used in many real world problems, [4] for example, in modeling probabilistic relationships in medical diagnoses. [5]

Computational Methods of BN

A Bayesian network is represented as a directed acyclic graph (DAG). The nodes within the DAG of BN denote relevant entities or random variables and the directed edges denote probabilistic relationships among them. For example, the DAG in Figure 2-1 below models a structure encoding relationships between History of Smoking (H), Lung Cancer (L), Bronchitis (B), Fatigue (F) and Chest X-ray results (C), as described in [6]. The numerical values of these relationships are encoded as a joint probability distribution (JPD) over a set of these random variables.

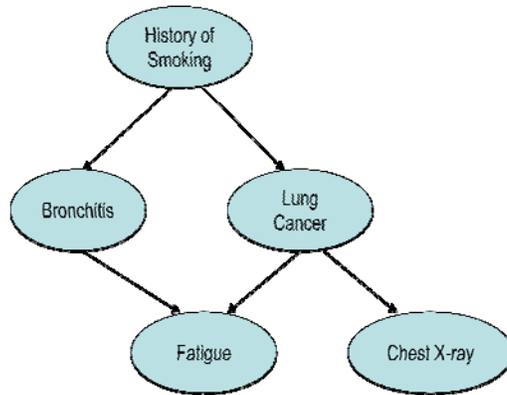


Figure 2-1 A Directed Acyclic Graph (DAG)

In probability theory, the notation $P(X / Y)$ denotes the *conditional probability* of a variable X *given* (denoted by symbol “[/]”) another variable Y . Two variables X and Y are independent if the probability of X given Y is the same as the probability of X occurring alone (i.e. $P(X / Y) = P(X)$) and vice versa, and when both events are known to occur with a certain probability, i.e. $P(X) \neq 0$ and $P(Y) \neq 0$. However there may be times when two variables are not independent by themselves but independent when *conditioned* upon a third variable, say Z , i.e., X and Y are *conditionally independent* given Z . A variable X is conditionally independent of Y given Z if

$$P(X | Z) = P(X | Z, Y) \text{ when } P(X | Z) > 0 \text{ and } P(Y | Z) > 0$$

i.e. if Z is given, the probability of X will not be affected by the discovery of Y . [3] At the core of BN is this notion of *conditional independence*. For example, in the example above in Figure 2-1, the node Bronchitis (B) is conditionally independent of nodes Lung cancer (L) and Chest X-ray (C) given that we know about History of smoking (H). Table 2-1 below gives other conditional independencies in this DAG.

Table 2-1 Conditional Independencies in Figure 2-1

Node	Parent	Conditional Independence
C	(L)	(C), (H, B, F) (L)
B	(H)	(B), (L, C) (H)
F	(B, L)	(F), (H, C) (B, L)
L	(H)	(L), (B) (H)

Another notion that BN model encodes is that of the *Markov condition*, also called the *Markov independence assumption*. This assumption says that each variable is conditionally independent of the set of all its non-descendants given the set of all its parents [3, 6], for example, Fatigue (F) is independent of History of Smoking (H) and Chest X-ray (C) given that we know about Bronchitis (B) and Lung Cancer (L).

Under these two assumptions, i.e. conditional independence and Markov assumption, the factorization theorem as described by Pearl encodes a unique probability distribution for a graph G which is described by the following equation (1) [3]

$$P(X_1, \dots, X_n) = \prod_{i=1}^n P(X_i | Pa_i^G) \quad (1)$$

Where Pa_i^G are the parent nodes of the variables, X_i , in G . Equation (1) is called the *chain rule for Bayesian Networks*. [3] As an example, the graph structure G in Figure 2-1 of a BN encodes independence assumptions while the conditional probability distributions (CPD), of the form $P(X_i / Pa_i)$ where Pa_i are parents of X_i , provide the quantitative parameters for the joint probability distribution (JPD) of the BN represented by this graph G . The JPD of the DAG shown in Figure 2-1 can be calculated as follows by equation (2) as follows –

$$P(f, c, b, l, h) = P(f|b, l) * P(c|l) * P(b|h) * P(l/h) * P(h) \quad (2)$$

In a BN, the DAG defines the *structure* and the CPD values for each variable are called the *parameters*. *Inference* refers to the query for finding the probability distribution score of a node (node in question) given values of a subset of nodes (instantiated nodes) in the DAG. For example in Figure 2-1, if we know a patient has history of smoking and positive chest x-ray, we may be interested in finding the probability of that patient having lung cancer, i.e. $(P(l | h, c))$ and having bronchitis, i.e. $P(b | h, c)$. Exact inference is a non-deterministic polynomial time hard (*NP-hard*) problem [7]. Algorithms developed earlier such as message passing [3] in DAG, Symbolic Probabilistic Inference (SPI) [8], arc reversal / node reduction operations [9-10], and the Junction Tree algorithm, [11-12] all have NP hard computational complexity in a multiply connected DAG and can become intractable for inference [6] in large networks. Following this, Cooper [7] obtained a result that the problem of determining the conditional probabilities is tractable in multiply connected networks and belongs to the class of problems that is P – complete if the remaining variables in a BN are restricted to having no more than two states per node and no more than two parents per node but with no restriction on number of children per node, given that certain variables are instantiated. Therefore, approximate inference algorithms such as stochastic simulation and deterministic search [13], finding the most probable explanation also called abductive inference methods [7] have been developed by many researchers in the field.

Besides the development of approximate algorithms for inference, approximate algorithms to learn the structure and parameters of BN from data have been developed as well. When the variable X or its parents are discrete valued (i.e. binary or multinomial),

to learn the CPD of a single variable, a beta density function or a dirichlet density function is used. In case of binary variables, a beta density function is used; in the case of multinomial variables, a dirichlet density function is used. [6] Unlike the case of discrete variables, when variable X or its parents are real valued, linear Gaussian conditional densities [14] or other appropriate density functions [6] are used to represent the underlying data for assessing CPD values.

In case of discrete variables a conditional probability table (CPT) is defined to represent the probability of X_i conditioned on each of its parents Pa_i . Together, the CPTs of all variables and the DAG define the JPD. For example, if the number of parents of a node denoted by Pa_i consists of K binary variables, the table (CPT) for the node defines 2^K rows of distributions. Therefore, while a full table form can describe any discrete conditional probability distribution (CPD), the number of parameters required grows exponentially in the number of parents Pa_i . [3] Therefore methods to reduce the complexity of parameter estimation for local CPTs have been developed. These methods all involve independence of causal influence assumption (ICI). Below, we describe these methods in detail.

Causal Independence and ICI models

A major difficulty in model building using Bayesian networks (BN) arises when numerical parameters to quantify them for conditional probability tables (CPT) are needed [15]. The complete CPT for a binary variable with n binary predecessors in a BN requires 2^n independent parameters [3]. Hence the number of parameters in a CPT grows exponentially with the number of parents and can become prohibitive for model building.

The BN however, does not constrain how a variable depends upon its parents; one interpretation is that the directed edges between parent and the child represent causal relationships [16]. Nonetheless, as shown previously by other researchers, there is some structure in the dependencies and probability functions of parents and child that can be exploited for knowledge acquisition and inference. The dependencies can be stated as rules [17], trees [18], multinets [19] or some form of binary operation that can be applied to values from each of the parent variables. *Independence of Causal Influence* (ICI) or *Causal Independence* [3, 20-21] is one such dependency and refers to the situation where multiple causes *independently* contribute to the common effect. An assumption of causal independence among the parent nodes that affect the child node greatly reduces the number of parameters required.

Noisy-OR Model

The Noisy-OR gate [3, 22], or distribution, is a member of the ICI family. [21, 23-24] The Noisy-OR model [3] makes this assumption and provides a logarithmic reduction in the number of parameters required relative to the CPT. This model has been shown to perform reasonably well in the field of medical diagnosis. [5] The word ‘noisy’ reflects the fact that the interaction among the cause(s) and the effect is not deterministic thus allowing the presence of the effect in presence or absence of any modeled causes. One can think of Noisy-OR as a probabilistic extension of the deterministic binary OR model. In practice, it is often impossible to capture all the possible causes for an effect. To address this issue and help the domain experts in the knowledge engineering process, Henrion proposed an extension of the Noisy-OR by introducing the concept of “*leak*” or

background probability [23]. Leak can be formally considered as one of the causes of the effect.

In Figure 2-2 below, the nodes X_i denote independent causes and Y is the common effect. The nodes U_i are called the inhibitor nodes [3] and encode individual effect (via their CPTs) of corresponding U_i on Y . A leaky Noisy-OR model can be described (Figure 2-2) using the following equations for several possible causes (X_1, X_2, \dots, X_n) of an effect Y under the two assumptions –

(a) Each of the causes X_i has a probability of producing the effect in the absence of all other causes and (b) each cause is sufficiently independent of the presence of other causes – i.e. –

$$p_i = P(y \mid x_i \text{ only}) = P(y \mid \overline{x_1}, \overline{x_2}, \dots, \overline{x_{i-1}}, \overline{x_{i+1}}, \dots, \overline{x_n}) \quad (3)$$

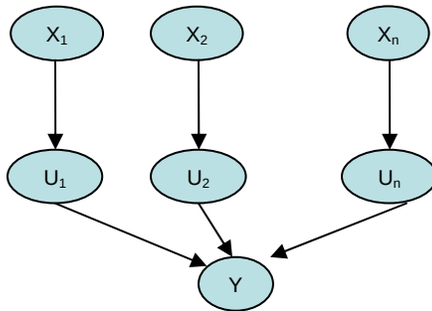


Figure 2-2 A Noisy-OR model

Using a deterministic mapping function such as Boolean OR, the CPT of Y defines how individual causes X_i interact to produce the effect Y . Therefore, the probability p_i as defined by equation (3) is also called the *link* probability and determines the causal strength between the cause, i , and the effect, Y , in the absence of all other

causes. It has been shown [3] that the probability of $Y = y$ given a subset X_p (x_i present), i.e. a set consisting of causes that are present, is given by the following equation (4) –

$$P(y/X_p) = 1 - \prod_{x_i \in X_p} (1 - p_i) \quad (4)$$

Under the assumption that causes produce the common effect independently, equation (4) can calculate the probability value for an effect solely based on the causal strength p_i of each cause to the effect. Therefore using the assumption of causal independence, the number of values required for CPT elicitation of effect Y reduces from *exponential to linear in number of causes*.

Further the *leak* probability p_0 which models un-modeled causes can be defined as

$$p_0 = P(y | x_1, x_2, \dots, x_i, \dots, x_{n-1}, x_n) \quad (5)$$

Let p' define the probability that Y is present when x_i is present and all other causes of Y including un-modeled causes (*leak*) are absent. From [3] for the leaky Noisy-OR model following is defined –

$$1 - p' = (1 - p_i) / (1 - p_0) \quad (6)$$

$$\therefore p_i = P(y/x_i \text{ only}) = p' + p_0 - p' * p_0 \quad (7)$$

The two ways of parameterization of CPD using Noisy-OR gate equation (4) is credited to Henrion [23] and Diez [22]. For calculating the link probability, the difference between the Henrion method and the Diez method lies in learning Noisy-OR parameters from equation 7. [15] The Henrion method seeks the p_i parameter where as the Diez method seeks the p' parameter.

Among the studies in the past, where the Noisy-OR gate model has been successfully applied are – reformulation of the rule based expert INTERNIST-1/QMR system into a probabilistic system by combining probabilities from disease profiles and hospital discharge statistics [5], deriving parameters from small data sets for converting from single-disorder to multiple-disorder liver disease diagnostic system (HEPAR-II) [15] and to an artificial domain for comparison of human expert’s judgment of parameters using the two ways of parameterization credited to Diez [22] and Henrion [23] in using [25] the Noisy-OR assumption. The result of this last study [25] claims that the Henrion method is better at providing Noisy-OR parameters from data [15] when the underlying distribution follows the Noisy-OR assumption, and the Diez method is better when human experts provide the parameters.

Thus the Noisy-OR model may be suitable for parameter estimation in large scale domains, such as medical diagnosis, where an observation such as a symptom can be triggered independently by a number of causes (diseases), or a number of causes can independently lead to a common complex disease. Similar to Noisy-OR, other forms of noisy deterministic functions (Noisy-AND, Noisy-MAX, Noisy-MIN, Noisy-ADD) [22-24, 26-27] have been defined and proposed for assessing values for CPD in a BN using the assumption of causal independence.

Other Related Models

While these models greatly reduce the complexity of parameter estimation in CPT and can serve as a good first approximation (Chapter 3) for modeling, the conditional probability distributions (CPD) in themselves do not account for interactions among causes that lead to the common effect. Although all of the above models take into account

the probability of an effect given each single cause as an input, the interactions defined among causes (by virtue of the noisy deterministic function) are considered to be synergistic or reinforcing. [28] Therefore, when multiple causes are present, the causes may reinforce each other (i.e. making the effect more likely to be present) or may undermine the impact of each other (i.e. the effect becomes less likely when more causes are present). As pointed by Xiang et al, [28] all of the above distributions (Noisy-AND, Noisy-MAX, Noisy-MIN, Noisy-ADD) can only express one type of causal interaction in a model, i.e. reinforcing.

To address the possibility of reinforcing interactions between causes, recently Lemmer and Gossink [29] proposed the Recursive Noisy-OR (RNOR) distribution which allows elicitation of probability parameters of the effect given subsets of causes as input. RNOR defines the concept of positive causality and how the dependent causes can work together as being either “synergistic” or “interfering.” The RNOR model can incorporate an expert provided probability distribution for an effect as well as a subset of values of causes given as input, wherever applicable and claims to be a generalization of Noisy-OR model. The RNOR model does not handle expert assertions of interference between causes. If an expert-provided subset of values implies inhibitions or interference, and the causes are undermining, RNOR can produce probability values that are greater than one. [29] Its application to problem domains such as medical diagnosis looks promising because of its ability to represent a subset of causes but it needs empiric evaluation.

More recently, Xiang and Jia have proposed a variation of this model. The non-impeding Noisy-AND tree (or NIN-AND tree) model [28] can represent both types of causal interactions among a set of causes, some of which can be reinforcing and others

undermining. However, this model also has limitations in expressing all possible causal interactions. For example, to model grouped causes that can selectively reinforce and grouped causes that can selectively undermine. Druzzdel et al. [30] have proposed yet another theoretical model – Probabilistic Independence of Causal Influences (PICI) as an extension of ICI models that leads to more expressive parametric models that are able to cope with a combination of positive and negative influences. To the best of our knowledge, RNOR distributions, PICI and NIN-AND tree models have not been studied extensively in real world problems but hold promise for medical domain applications.

BN Vs Other Methods

Our decision to use a BN framework, as opposed to other methods, for this research is best explained if we compare BN with some of the other statistical and computational methods available for analysis and model formation. BN belongs to a class of generative models. Generative models differ from discriminative models in that a generative model contains a full probability distribution of all variables, whereas a discriminative model provides a model only of the outcome variable(s) conditioned on the observed variables. Therefore, BN models are both diagnostic and predictive at the same time. They offer several advantages when compared to other computational methods for knowledge representation such as Neural Networks or even traditional statistical methods such as linear regression models for data analysis. We describe some of the differences briefly below.

Artificial Neural Network (ANN) models need large numbers of complete cases for training to be used in prediction and classification problems. They often overfit the data to the problem and, unlike BN, lack explanation capabilities. Therefore, they have

limited use in a domain like biomedical research. BN models on the other hand are probabilistic and can take personal (expert) beliefs (using a subjectivist approach) for model building – they are well suited to derive prior probability values from small sample sizes and are able to handle missing data values reasonably well while avoiding over fitting of data to the model. [4] They model cause and effect in a normative way [3] and therefore provide a framework for incorporating all available information in a systematic manner for both model and parameter estimation to produce a predictive distribution.

Linear regression models are non-parametric (distribution free) statistical models and can also be used for prediction and classification problems like ANN and BN. However, they do not handle missing data well in the input, and due to their susceptibility to noise in the data, their use for model building is limited, particularly in the biomedical domain where data are noisy. As with ANN, linear regression models lack normative explanation capabilities and also suffer from the “curse of dimensionality” i.e. an exponential increase in model space with addition of extra dimensions, as is the case in domains with large number of variables such as the biomedical domain.

Other methods of knowledge acquisition such as systematic review and meta-analysis aim to more precisely estimate the true “effect size” from a group of studies as opposed to the less precise estimates derived in a single study under a given set of assumptions. But these are not computational methods to synthesize a prediction model.

Therefore, despite the computational complexity of inference, BN methods offer several advantages over traditional methods of knowledge acquisition and representation. They have been shown to represent domain knowledge with conciseness and normative form. The product form of equation (1), i.e. the chain rule and the conditional

independence assumption makes the BN representation of the JPD compact. Their ability to handle missing data values and ability to calculate probability scores from small data sets is another reason to choose BN methods. Most of all, BN methods provide a strategic framework for our research to incorporate all available information in a systematic manner for both model and parameter estimation to produce a predictive distribution that can be used for inference and hypotheses generation. In this research we develop a methodology using the BN framework and summary and statistical measures from published studies for integration of information from disparate sources of data in the biomedical domain.

Challenges for Knowledge Representation and Inference in Biomedical Domain

To gain full understanding of the implications of the research thus far in the biomedical domain, we first need to understand how we could represent the existing information, derive knowledge and inference from it. In light of the data gathered from separate environmental and genetic studies and not on the same individual, a unifying model is desperately needed to represent such information perhaps in a computational model (In-silico). Such a model, once constructed could also be used for knowledge discovery or in clinical applications.

For instance, it is believed that both genetic and environmental risk factors have an important role to play in most common diseases. [31-32] A 2003 review article titled “Genomics as a Probe for Disease Biology” in the New England journal of Medicine highlights the importance of understanding genetics together with the pathology of a disease in order to unravel the underlying disease processes. [33] Asthma, which is best considered as a cluster of related disorders, [34] is one such common complex disease.

The prevalence of asthma has risen dramatically in the last few decades, [35] suggesting that environmental risk factors have a key role to play together with genetic factors in developing a risk of the disease [35-37] in early childhood. As is the case with asthma, there is also evidence suggesting the role of environmental and genetic factors for most other common diseases such as diabetes, obesity and heart disease. [38-39]

Recently, it has also been argued that the current classification of human disease has significant shortcomings as reflected in its lack of sensitivity in identifying pre-clinical disease and lack of specificity in defining disease unequivocally. [40] Therefore, it has been proposed that an approach using network principles and *linking phenotype or clinical data with the genotype and environmental data* associated with the risk of disease, can lead to more accurate identification and classification of disease diagnostic and treatment options. [40] For example, in the field of cancer biology, bioinformatics methods that integrate diverse data (clinical and genotype) in their analysis for predicting survival rates have achieved higher accuracy than use of clinical data alone, even when the data analyzed are from different sources. [41-42]

Given the example above, we believe that the challenges of knowledge representation and inference in this domain are three fold.

Existence of silos of datasets

First, data are being amassed in silos of biomedical databases. Currently, there are major initiatives underway by the National Institutes of Health (NIH) to address the rise in common diseases (like asthma and diabetes) by studying their genetic linkages and disease-environment interactions [43-47] in Genome Wide Association Studies (GWAS) and Environment Wide Association Studies (EWAS) respectively. Due to the interplay

of environment, lifestyle and small effects of many genes, researchers have focused on very different aspects, for example pharmacogenetics / pharmacogenomics, [37, 48-49] gene-environment interaction [47, 50] and clinical environmentally focused [51-52] studies. Despite their best efforts, researchers find it hard to conduct unbiased studies in well defined populations that have sufficient power to detect small effects attributed to genetic or environmental factors. [49, 53-54] Therefore, to date these studies are being conducted in sub populations and patient level data are being collected and stored within the individual institution's repository.

Access Rights

Second, due to lack of data sharing agreements among institutions and patient privacy concerns, [55-56] the data are not accessible in their raw form to outside entities like researchers in other institutions for any secondary analysis. The only publicly accessible results from these studies are published summary and statistical results. Thus, if any form of computational model needs to be developed to unify the information it most likely will have to use the published results.

Inference in Patient's Context

Third, there is a big gap in application of knowledge gained from biomedical research and its use in patient's context, for example from an EMR. Research that applies to clinical management of diseases and many rare disorders which are governed by straightforward Mendelian rules of inheritance have been known for some time. However, teasing out the genetic and environmental components for complex disorders such as diabetes, heart and lung disease, autoimmune and psychiatric disorders and their clinical management remains challenging [33] due to this application gap.

Therefore, there are many practical challenges for application of existing research in the biomedical domain. In the next chapter we discuss how, based on our background research, we can use BN for building a probabilistic framework for knowledge representation and inference in this domain. We specifically develop strategies using this framework to incorporate all available knowledge in a coherent model from various sources for example as presented in Figure 1-1 – environmental, genetic, demographic, socio-economic and clinical phenotypes.

Our hypothesis in building such a model is to 1) find associations across the domain that are not apparent in the silos of datasets and 2) confirm these associations by testing – a) against data from our EMR and b) by evaluating against what has been published in the literature so far. We are interested to know how much of explanation is provided by a subset of data, for example how well genetic associations can explain the risk of a complex disorder like asthma or diabetes.

To test our model we use Area under the Curve (AUC) of Receiver Operator Characteristics (ROC) curves as a performance measure. We describe the ROC performance measure below.

Receiver Operating Characteristic (ROC) Curve

A ROC curve is a plot of pairs of true positives (Sensitivity) vs. false positives (1 – Specificity) for various cut points of a binary classifier as its discrimination threshold is varied. The ROC curve has its roots in Signal Detection Theory from World War II and since then they have been extensively applied as an analysis tool in areas of medicine, radiology [57], and many other fields. [58]

Since ROC curve analysis is a non-parametric method and does not rely on the underlying distribution, their use as an analysis tool is particularly attractive to the machine learning community; especially for use as a model comparison tool to select an optimal model given the data. The area under the curve (AUC) of an ROC curve is equal to the probability that a classifier will rank a randomly chosen positive instance higher than a randomly chosen negative one [58] and therefore measures the performance of the model. A ROC with AUC of 0.5 score has no predictive value and is as good as chance.

A Bayesian approach to represent domain knowledge

Based on our background research on Bayesian Networks and the breadth of the challenges involved in building a coherent model in the biomedical domain, we evaluate the use of a probabilistic framework, combining BN fundamentals of conditional independence and the Markov condition to encode domain knowledge both qualitatively and quantitatively. We then evaluate the use of the Independence of Causal Influence (ICI) assumption as a potential data integration strategy. In this chapter, we describe our experiments in the clinical domain using data on childhood asthma from our EMR, the Regenstrief Medical Record System (RMRS) [2] and another independent test data source from our pediatric decision support system in practice – Child Health Improvement through Computer Automation (CHICA) system [59] described below.

In experiment 1, we compare performance of a data derived DAG to a domain expert’s model DAG by testing it on the same test datasets to evaluate the sensitivity of the BN function to the structure of the DAG with real world datasets.

In experiment 2, we test the validity of the Independence of Causal Influence (ICI) assumption in particular the Noisy-OR model using the same datasets from experiment 1. To model domain causal relationships, over and above causal independence, we test the validity of Recursive Noisy-OR (RNOR) rule.

Experiment 1: Bayesian Networks from Electronic Medical Records

Probabilistic Asthma Case Finding - A Pilot Study using the CHICA system [60]

Introduction

One of the most useful characteristics of BN is the ability to construct DAG models based on expert knowledge of causal relationships or entirely empirically, using large datasets. In fact, this feature makes BN ideally suited for our goal of merging information from different sources. However, the comparability of DAGs derived in these different ways has not been tested. In this series of experiments, we use Bayesian Networks as a strategy for modeling patients' clinical status with the goal of comparing two DAGs: the one developed by the domain expert with the one mined from data. A large retrospective cohort consisting of 16,187 children having wheezing prior to age two was mined from data to derive a DAG to predict asthma after age five. We compare the predictive power of this mined network with a domain expert's DAG using two test scenarios – (a) using a test dataset from our EMR and (b) using an independent dataset from our clinical decision support system (CDSS).

Methods

Our goal is to derive these BNs from data in our clinical data repository. To achieve this we considered two possibilities 1) use a clinical expert to define the nodes and arcs in the BN and train the resulting BN to derive parameters using data or 2) use data mining techniques to derive the BN structure and parameters from data. In this chapter we describe an experiment in which we compare these approaches in the domain of childhood asthma. Pediatric Asthma cases and controls were identified from RMRS and from the CHICA system [59] for an independent test set. CHICA is a Clinical

Decision Support System (CDSS) used in our Pediatric Primary Care (PCC) practice in conjunction with RMRS, [2] and we briefly describe it below.

CHICA Overview

The CHICA system went live on Nov. 5th, 2004 at the Pediatric Primary Care Center (PCC) of Wishard Hospital, Indianapolis, Indiana, and now has data from over 25,000 patients. The system provides decision support for well child care and management of common childhood problems. The user interface consists of scannable paper forms called adaptive turnaround documents (ATD). [61] Data collected on ATDs are used to generate questions to the patient and reminders to physicians at the point of care. CHICA uses a knowledge base encoded as Arden Syntax medical logic modules (MLM) [62] and patient data from the RMRS [2] and CHICA databases to generate dynamic content on the ATD forms. The MLMs are prioritized using a global priority scheme to address the most relevant questions and reminders on the ATD. [59, 63] The CHICA system electronically receives a record of all clinical observations from the RMRS database for every patient visit.

We analyzed data for all children over 5 years of age in our system. Children were classified as cases or controls based on the presence of an ICD-9 code for asthma (493.*) or more than two prescriptions of an asthma medication. From the filtered set we were able to extract the variables listed in Table 3-1 to get an “Asthma Status,” sex and race for each patient (ages 5 years or older) who had a visit to the PCC clinic. The CHICA system in its current state has been described in detail in previous manuscripts. [59, 64-66]

Model

We used Netica software [67] (Norsys Software Corporation, Vancouver BC, www.norsys.com) to construct BN for our expert model and WinMine toolkit [68] (<http://research.microsoft.com/~dmax/WinMine/Tooldoc.htm>) for mining a directed acyclic graph (DAG) from data. Netica allows network construction and parameter learning from data. The WinMine toolkit provides software for learning a DAG from data. Table 3-1 below lists the data variables used for modeling expert BN and data mining the DAG.

Data

To compare the two DAGs – expert BN and mined BN we compared the performance of the BNs on two datasets. First, the data from 16,187 cases from Regenstrief Medical Record Systems (RMRS) were split randomly into 2/3 of cases for a training set and 1/3 for a test set. For the second dataset, the CHICA system electronically receives a record of all clinical observations from the RMRS database for every patient visit. We filtered these observations and preprocessed them to extract the data variables listed in Table 3-1. *These data were collected for children ages 5 and above to predict childhood asthma.* At the time of the study, the CHICA system had data for 1984 cases. Table 3-2 lists the baseline characteristics of the datasets used in these experiments.

Table 3-1 Data Variables for Model – Experiment 1

Variable	Values
Race	White, Black, Hispanic, Other, Unknown
Sex	Male, Female
Eczema	True, False
Wheeze	ICD9 or clinic billing diagnosis before age 2 (True, False)
Asthma	ICD9 (493.*) or any clinic billing diagnosis after age 5 or at least 3 drugs from a specified list within 12 months after age 5 (True, False)
X-ray	Chest x-ray before age 2 (True, False)
Drug	Drugs from a specified list before (True, False)
Wz_hosp	Inpatient admission with hospital ICD9 as wheezing (True, False)
Wz_er	Any ER visit with billing ICD9 as wheezing (True, False)
Ins_cat	Insurance category - first available insurance in the same year of the first wheezing diagnosis (True, False)

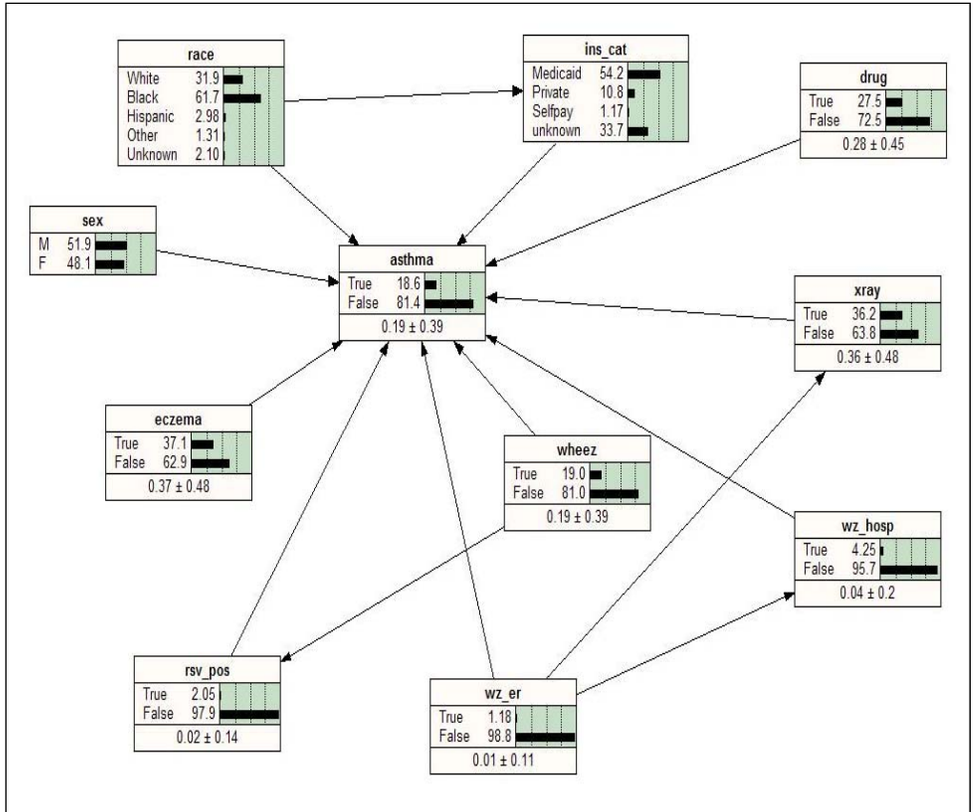


Figure 3-1 Expert's BN trained with data from RMRS

Table 3-2 Baseline characteristics of training and test sets

Variables	Training Set from RMRS (n = 11,000)		Test Set from RMRS (n = 5,187)		Test Set from CHICA (n = 1,984)		
	#	%	#	%	#	%	
Race	Hispanic	98	1%	373	7%	429	22%
	Unknown	115	1%	220	4%	24	1%
	Black	6859	62%	3116	60%	1156	58%
	White	3806	35%	1385	27%	327	16%
	Other	122	1%	93	2%	48	2%
Sex	Female	5357	49%	2503	48%	916	46%
	Male	5641	51%	2684	52%	1068	54%
Eczema	True	4021	37%	2021	75%	NA	NA
	False	6979	63%	3166	61%	NA	NA
Wheeze	True	1661	15%	1431	28%	187	9%
	False	9339	85%	3756	72%	1797	91%
Asthma	True	1561	14%	548	11%	536	27%
	False	9439	86%	4639	89%	1448	73%
X-ray	True	4015	37%	1900	37%	1529	77%
	False	6985	64%	3287	63%	455	23%
Drug	True	3013	27%	1488	29%	529	27%
	False	7987	73%	3699	71%	1455	73%
Wz_hosp	True	433	4%	247	5%	159	8%
	False	10567	96%	4940	95%	1825	92%
Wz_er	True	102	1%	98	2%	143	7%
	False	10898	99%	5089	98%	1841	93%
Ins_cat	Medicaid	4762	43%	4051	78%	1631	82%
	Unknown	5276	48%	182	4%	64	3%
	Private	844	8%	893	17%	160	8%
	Self-pay	118	1%	61	1%	0	0%
rsv_pos	True	244	2%	114	2%	17	1%
	False	10756	98%	5073	98%	1967	99%

Expert's Design of BN with training using Netica

Using the predictor data variables of Table 3-1 as nodes and the domain knowledge for joining them with arcs, the domain expert (SMD) created a BN as shown in Figure 3-1. This BN was trained with the training set and compiled using Netica software. In Figure 3-1 the BN shows marginal probabilities of each node with an asthma prior probability of 18.6%.

BN Derived using Data Mining Techniques

The training set from RMRS data was used to derive the DAG for this approach. The software from WinMine toolkit was used to preprocess the data from raw format (excel tab delimited) to WinMine XML format, which was then used for creating and editing a plan file to instruct the learning algorithm to model each predictor variable based on a) the role of each variable – *input* (used to predict other variables), *output* (predicted by other variables) and *input-output* (both predicted and used to predict) or *ignored* (not used); b) the model distribution used for each variable – specifies the tree versus the table representation and the local distribution of the variable, the representation chosen in this case is tree for discrete variables and the distribution chosen is binary multinomial to accommodate missing values; c) Model-as-binary information (*missing vs non-missing values for binary variable or one state vs all other states for discrete variables.*). Figure 3-2 shows the roles, the distributions used and model-as-binary information for each of the predictor variables for our model in the WinMine toolkit.

Variable	Role	Distribution	Model as binary
race	input-output	tree-multinomial	Black vs. Others
sex	input-output	tree-multinomial	None
asthma	input-output	tree-multinomial	0 vs. Others
eczema	input-output	tree-multinomial	None
xray	input-output	tree-multinomial	None
rsv_pos	input-output	tree-multinomial	0 vs. Others
drug_BF2	input-output	tree-multinomial	0 vs. Others
wz_hosp	input-output	tree-multinomial	0 vs. Others
wz_er	input-output	tree-multinomial	0 vs. Others
wheez	input-output	tree-multinomial	0 vs. Others
ins_cat	input-output	tree-multinomial	Medicaid/Wishard vs. Others

Figure 3-2 Plan file for deriving the WinMine model

We replicated the DAG derived by the WinMine software in Netica. This DAG is shown in Figure 3-3. We then trained this BN using the same training set as the expert BN and compiled it to get the prior probabilities for each node in the model. The asthma prior probability in this model was 13.9%.

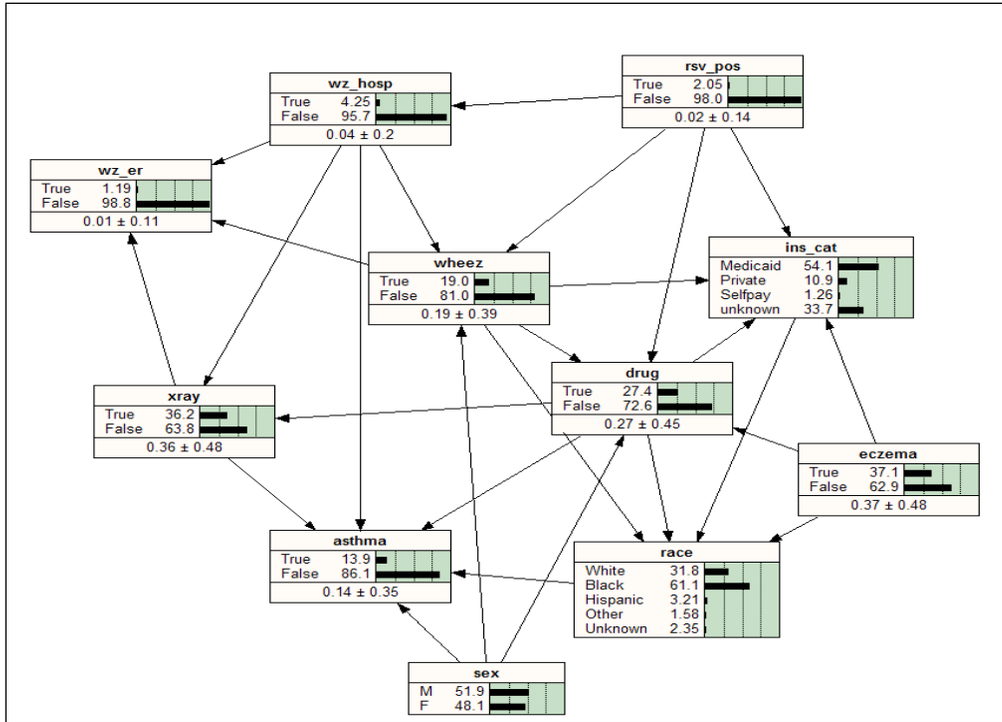


Figure 3-3 BN mined and trained with data from RMRS

Testing the Two Models

The two BN models were evaluated, first, using data from our test set from RMRS data (1/3 split from the large cohort study) and, second, using the CHICA data set derived from CHICA database.

Netica provides an interface to test the BN using a case file of test data. The node(s) of interest for prediction are treated as “unobserved nodes”. Asthma was used as an unobserved node in our tests. The software reports several measures for each unobserved node. We chose to use the quality of test results, which gives a performance measure in the form of a table for sensitivity, specificity, positive predictive value and negative predictive value.

We compared BNs using Receiver Operating Characteristics (ROC) curves [57]. The area under the curve was used as a measure of overall test performance. The ROC

curve was obtained by plotting pairs of true positive rate (sensitivity) and false positive rate (1 - specificity).

Results

We had 5188 cases in our RMRS test set and 2000 cases in the CHICA test set. Both the Expert and the Mined BN were tested using these sets, the results of which are listed below in Table 3-3, Figure 3-4 and Table 3-4, Figure 3-5 for RMRS and CHICA test sets respectively.

Using RMRS Test Set

Table 3-3 Operational Characteristics with RMRS test set

	(* Expert BN		+ Mined BN)	
Sensitivity (%)	Specificity (%)	Predictive (%)	Predict-Neg (%)	1 - specificity (%)
*84.77	*38.11	*16.28	*94.63	*61.89
*63.31	*68.4	*22.15	*92.92	*31.6
*50.07	*78.8	*25.12	*91.74	*21.2
*36.03	*88.05	*29.99	*90.65	*11.95
*15.76	*95.6	*33.71	*88.88	*4.4
*6.89	*96.82	*23.53	*87.98	*3.18
*1.99	*99.53	*37.5	*87.73	*0.47
+82.38	+44.51	+17.41	+94.68	+55.49
+53.91	+77.16	+25.11	+92.18	+22.84
+43.71	+83.97	+27.92	+91.31	+16.03
+26.09	+93.38	+35.88	+89.89	+6.62
+10.86	+98.34	+48.24	+88.6	+1.66
+0.93	+99.68	+29.17	+87.63	+0.32

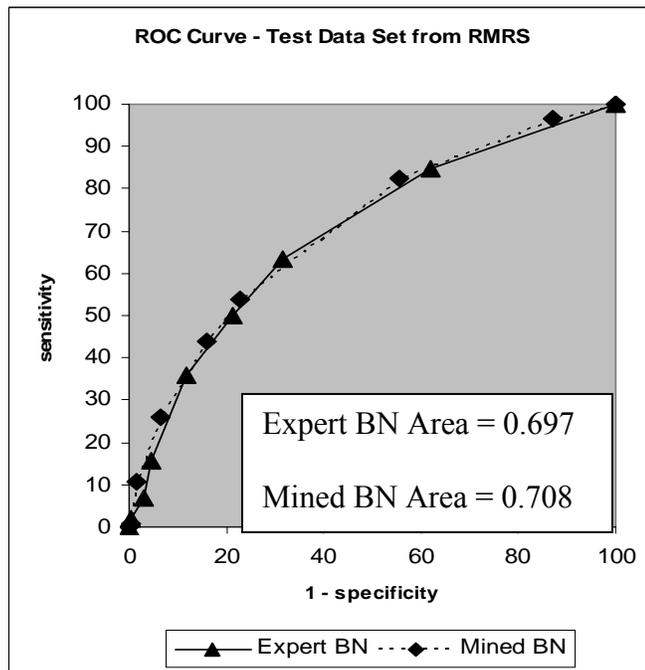


Figure 3-4 ROC curves using test data from RMRS

Using CHICA Test Set

Table 3-4 Operational Characteristics with CHICA test set

Sensitivity (%)	Specificity (%)	Predictive (%)	Predict-Neg (%)	1-Specificity (%)
*75.56	*38.05	*31.11	*80.79	*61.95
*56.53	*64.3	*36.95	*79.98	*35.7
*48.69	*73.69	*40.65	*79.51	*26.31
*30.04	*90.81	*54.76	*77.81	*9.19
*25.56	*93.99	*61.16	*77.33	*6.01
*1.31	*99.93	*87.5	*73.23	*0.07
+77.43	+27.97	+28.46	+77	+72.03
+44.59	+76.66	+41.42	+78.89	+23.34
+40.11	+80.18	+42.83	+78.34	+19.82
+26.87	+89.92	+49.66	+76.86	+10.08
+15.49	+97.24	+67.48	+75.66	+2.76
+5.97	+99.65	+86.49	+74.11	+0.35

* Expert BN + Mined BN

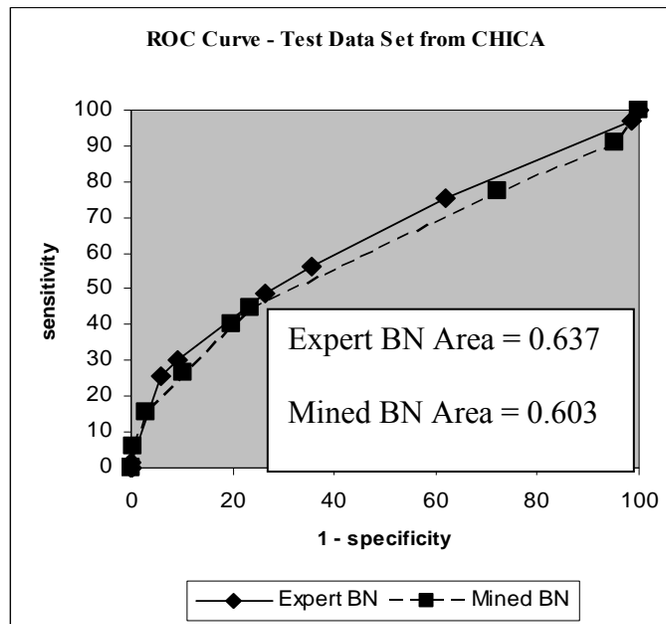


Figure 3-5 ROC curves using test data from CHICA

Discussion

The results of AUC for the ROC for both BNs using the same test set are comparable. Both the expert BN and the mined BN performed better with the test set from the RMRS data set when compared with the CHICA data set. We attribute degraded performance when testing with CHICA data set due to our less stringent inclusion criteria in the CHICA test set. For example, any chest x-ray observation will satisfy the inclusion criteria for CHICA data, where as for the RMRS test set only a chest x-ray finding before age 2 will satisfy the inclusion criteria.

Similar performance of each BN in each test scenario suggests that the mined BN has a predictive value similar to the DAG derived by the expert. Furthermore, the two compared BNs in this experiment were derived from two different data sources – a subjective model based on a clinical expert’s judgment and from data from our EMR. The data derived BN was as good as the subjective model suggesting the BN method presents a knowledge representation and inference tool where subjective decisions can be incorporated to approximate the domain knowledge.

Experiment 2: Strategies for Learning BN parameters

Probabilistic asthma case finding: A Noisy-OR reformulation [69]

An Empirical Validation of Recursive Noisy-OR (RNOR) Rule for Asthma Prediction [70]

Introduction

Development of a BN to represent the relationships between GWAS results and gene disequilibrium data requires assumptions about independent causal associations. As a preliminary evaluation, we wanted to evaluate the Noisy-OR and Recursive Noisy-OR formalisms by comparing the predictive power of BN developed using these methods to a “gold standard” BN trained on clinical data.

Noisy-OR: In combining disparate data sets in which one data set describes the relationship between some causes or risks and their consequences, and another data set describes the relationship between other causes or risks and the same consequences, there are no cases from which to infer the combined effect causes recorded in the different datasets. One approach to this challenge is to assume causal independence among predecessors (parents) of a given node. In this case, it may be reasonable to apply a Noisy-OR calculation [3] to estimate the probability of the child node given a particular combination of values for the parents [22-24]. By assuming these variables have independent causal effects, the Noisy-OR allows us to assign posterior probabilities conditioned on causes from these different sets. However, the validity of the independence assumption is rarely tested. We wanted to test this assumption by applying the Noisy-OR to combinations of conditioning variables for which we knew the joint probability distributions.

RNOR: To test the Noisy-OR model against a “gold standard,” we need a BN trained on a data set that represents the joint probability distributions. Several algorithms have been developed for training BN by learning their conditional probability distributions (CPD) from such datasets. [27, 71] However, a challenge arises when the training data set has no cases representing a particular combination of values for variables that condition a particular CPD. This is a common problem in complex BN even when large training sets are available. [22] A common strategy in this situation is to assign a uniform (uninformed) distribution to the dependent variable, conditioned by this combination of variables. For example, when the probability of asthma is conditioned on the sex, race, insurance and past wheezing history of a patient, there may be no cases in the training data that are male, white, on Medicaid and with a positive history of wheezing. Under the uniform distribution strategy, the probability of asthma would have a 50-50 distribution.

The ideal strategy would retain posterior distributions for combinations of parent node values that exist in the training set while applying the Noisy-OR rule when there are no cases representing a combination of conditioning variables. In 2004, a potential solution to this problem was published by Lemmer and Gossink. [29] The Recursive Noisy-OR (RNOR) rule described by these authors was intended to incorporate expert estimates of probabilities conditioned on more than one node while applying the Noisy-OR rule when these higher order conditional probabilities were not available. We reasoned that the RNOR algorithm might be a successful strategy for training a BN from a data set that did not contain cases representing all combinations of variables conditioned on a given node. We hypothesized that this RNOR approach would produce

a BN with better predictive power than either a Noisy-OR formulated or traditionally trained BN. This chapter describes the development and evaluation of this strategy.

Methods

We constructed a BN in the domain of asthma prediction in children, using expert knowledge to derive a directed acyclic graph (DAG) and applied a commercially available software package to learn the CPDs (parameters) from a large clinical dataset. This empiric BN has been described before in Chapter 3 (Figure 3-1). [60] For this study, we reformulated the CPDs in our domain expert's BN using both Noisy-OR and the RNOR rule. Our empiric BN, Noisy-OR BN and RNOR BN were tested against two independent clinical data sets described below.

EMR Data and Variables

Clinical data for this study were derived from two datasets – RMRS and CHICA as described in experiment 1 in Tables 3-1 and 3-2.

Bayesian Network and Noisy-OR model

We took the Expert's BN from Figure 3-1 and reformulated it as a Noisy-OR model (Figure 3-6). The expert BN was trained with data to derive a CPT for each node. Since the Noisy-OR model inherently assumes binary causes (absent / present; true/false), we dichotomized the non-binary nodes i.e. *race and, insurance category*) into “true” and “false” condition by assigning “true” to the state that minimized the global leak (p_0) when all the other nodes are in a “false” state. Thus, because boys were more likely to have asthma, male sex was coded as the “true” state. Similarly, race = Black and ins_cat = Private were coded as “true” states. The marginal probabilities of all the causal nodes (*sex, eczema, wheeze, xray, drug, rsv_pos, wz_er, wz_hosp*) remain the same from our

expert BN. For the study, we only wished to compute the local CPT of the *node asthma* using the Noisy-OR parameters.

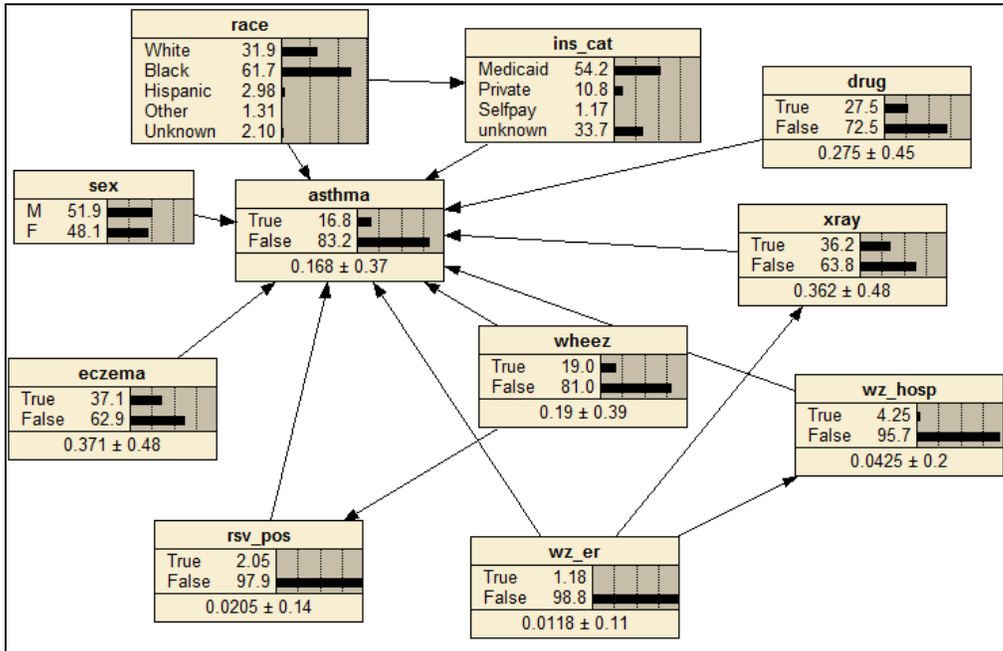


Figure 3-6 BN with Noisy-OR parameters

Obtaining Noisy-OR Parameters from Data

To derive the *leak* parameter for the network in Figure 3-6, we set all the nodes in the network that had an arc to the node *asthma* to a state *false*. The resulting posterior probability for the node *asthma* was our *leak* parameter, p_0 (0.014). Using this leak probability, we calculated the parameter p_i (the causal strength when no other cause is present) of each node (*node = True*) to the effect (*asthma*).

We used Netica to compute the posterior probability p_i of the effect given only one of the causes at a time. This is equivalent to eliciting the Henrion parameter which includes the leak parameter in the posterior probability. From there, we were able to derive the probability p' (Diez parameter) for each cause alone, using equation (7) in chapter 2.

Netica provides an interface to input Noisy-OR parameters for a given node and is able to calculate the conditional probability distribution for the node from these input parameters – effect node’s leak, individual link probabilities of causes. We used this feature to calculate the CPD for the *asthma* node. This Noisy-OR calculation is achieved with the function in Figure 3-7. The link parameters are listed in Table 3-5.

```

P (asthma | sex, eczema, wheez, drug, xray, rsv_pos, wz_er, wz_hosp, race, ins_cat) =
NoisyOrDist (asthma, 0.014, sex == M, 0.022, eczema, 0.048, wheez, 0.17, drug, 0.117,
xray,0.072, rsv_pos, 0.4, wz_er, 0.37, wz_hosp, 0.104, race == Black, 0.042, ins_cat ==
Private, 0.058)

```

Figure 3-7 Calculation of Noisy-OR parameters

The prior probability of the node asthma computed is slightly different in the two BNs - Noisy-OR reformulation vs. the Expert BN (16.8% Vs 18.6%).

Testing of Noisy-OR model

The two BN models – Expert (empiric) BN (Figure 3-1) and Noisy-OR Reformulation (Figure 3-6) were evaluated, first, using data from our test set from RMRS data (1/3 split, 5187 cases) and, second, using the CHICA data set derived from the prospectively collected CHICA database (1984 cases). Netica provides an interface to test the BN using a case file of test data. The node(s) of interest for prediction are treated as “unobserved nodes”. Asthma was used as an unobserved node in our tests. We compared the sensitivity, specificity, positive predictive value and negative predictive value of the Noisy-OR and the expert BN.

We compared the BNs using Receiver Operating Characteristic (ROC) curves. [57] The ROC curve was obtained by plotting pairs of true positive rate (sensitivity) and

false positive rate (1–specificity). The area under the curve (AUC) was used as a measure of overall test performance. AUCs were compared using the methods of Hanley, J.A. and B.J. McNeil. [72]

Results of Noisy-OR Reformulation

We had 5187 cases in our RMRS test set and 1984 cases in the CHICA test set. The empiric BN and the Noisy-OR reformulation of the empiric BN were tested using these sets. Results for these tests are plotted in Figures 3-8 and Figure 3-9 respectively.

When comparing the BN that utilized the Noisy-OR assumption to the BN with the fully data derived CPD (empiric BN), we saw a modest decrement in the AUC (0.697 vs. 0.726) when applied to the evaluation dataset from RMRS. When applied to the prospectively collected CHICA dataset, the decrement was about the same as before (0.612 vs. 0.637). However, neither BN (empiric or Noisy-OR reformulated) was particularly effective with the independent CHICA dataset.

Bayesian Network and Recursive Noisy-OR (RNOR) model

From experiment 1, the Expert (empiric) BN (Figure 3-1) had an 18.6% marginal probability of asthma. On examining the CPD of asthma in this empiric BN, we found the majority of its rows contained uninformed priors (i.e., uniformly distributed probability score – asthma = 50-50). We attributed this to the lack of cases satisfying the particular combination of causes in the training set (e.g., race = Other, Sex = male, Ins_cat = selfpay, Eczema = true, Wz_hosp = true). We hypothesized that by using the RNOR rule (described below) to calculate conditional probability for the CPD row value for asthma in such cases, the RNOR reformulated BN will outperform the empiric BN.

Recursive Noisy-OR Rule

In 2004, a rule for estimating complex probabilistic interactions was published. [29] The rule builds upon the Noisy-OR equation (4) in Chapter 2, to accommodate non-independent causes of an effect for calculating RNOR probability $p^R(x)$. Interested readers are encouraged to refer to the original paper [29]. The RNOR rule is a generalization of Noisy-OR and reduces to Noisy-OR in cases where $|x| = 1$, i.e. a subset of a single cause is provided by the expert $p^E(x)$. Furthermore, the rule preserves certain ratios (synergies and interference, see below), and the authors claim a major advantage of the rule is that it allows for arbitrary causally dependent subsets of probability scores to be incorporated in the estimation of $p^R(x)$, the probability score of effect.

The rule states that as long as the expert provided values for a subset of causes do not imply inhibition (i.e. abides positive causality, see synergy and interference below) it can be applied. Expert provided values implying inhibition among a set of causes will cause RNOR to produce probability scores greater than one, rendering the rule inapplicable. The RNOR rule is summarized in the following equation 1–

$$P^R(x) = \begin{cases} p^E(x), & \text{if expert provides information} \\ 1 - \prod_{i=0}^{n-1} \frac{(1 - p^R(x \setminus \{x_i\}))}{(1 - p^R(x \setminus \{x_i, x_{(i+1) \bmod n}\}))} & \\ \text{, otherwise} & \end{cases} \quad (1)$$

Synergy and Interference

To judge the rule for semantic correctness (i.e. the numbers produced make sense), the authors introduce the notion of positive causality. *Positive causality* refers to the idea that additional causes always increase the probability of achieving an effect.

Synergy of causes satisfies positive causality and produces an effect (probability) greater than the Noisy-OR calculation. *Interference* of causes also satisfies positive causality but produces an effect less than the Noisy-OR calculation. However, *inhibition*, in contrast to interference, violates positive causality and will produce probability scores greater than one. Therefore, the “information on the probability of an effect from a combination of causes provided by an expert (or derived from data) can be represented as a scalar multiple of the regular Noisy-OR.” This can be represented using equation (2) as defined in the original paper. [29]

$$1 - p^R(x) \equiv \delta(x)(1 - p^N(x)) \quad (2)$$

where $p^R(x)$ represents the probability from a RNOR estimation and $p^N(x)$ represents the standard Noisy-OR estimation. The factor $\delta(x)$ represents a scalar gain or attenuation coefficient between the two estimates. If $\delta(x)$ is less than one then it represents a biased amplifying coefficient of the probability and hence *synergy*. Conversely if $\delta(x)$ is greater than one then it represents a biased attenuating coefficient and hence *interference*. Finally, if it is equal to one then an independent product combination could hold implying causal independence.

Application of RNOR rule to the Asthma case finding BN

We took the expert’s BN from Figure 3-1 and calculated the CPD of the asthma node using the RNOR rule as follows.

Deriving link probabilities for RNOR rule

To apply the RNOR rule to the network for CPD computation of asthma, we individually calculated the link probabilities (p') of each cause of the effect (asthma) This was a two step process – (a) reduce the network by absorbing all nodes (see below

node absorption) except the node in question (e.g. eczema), looking up the “true” conditional probability score of the effect given the cause (i.e. $asthma = True$ given $eczema = True$), then looking up its “false” conditional probability score to find leak p_0 for this reduced network and (b) using equation (7) in Chapter 2 as before to find the Diez probability (p') in absence of all un-modeled causes including the leak calculated in (a). We repeated this two step process for all causal nodes for asthma thus first deriving the Henrion parameter (p_i) which includes the leak parameter in the posterior probability. From there, we derived the link probability p' (Diez parameter) for each cause alone. Table 3-5 lists the values for each link probability (p') to asthma.

Table 3-5 Link probability of each node to asthma

P(eczema=True)=0.048	P(drug= True) = 0.117
P(rsv_pos=True) = 0.4	P(wheez=True) = 0.17
P(wz_er=True) = 0.37	P(ins_cat=Private)= 0.058
P(wz_hosp=True)=0.104	P(race=Black)=0.042
P(xray=True) = 0.072	P(sex=Male) = 0.022

Node absorption

Node absorption is a network transform which removes nodes from a BN and makes any necessary adjustments to the resulting network. Also known as averaging out or “summing out a variable”, this transform leaves the full joint probability distribution of the remaining nodes unchanged.

Node absorption is part of the network transform for solving decision problems using influence diagrams (Bayesian networks with decision nodes) and is described in

detail in Shachter’s algorithm [9-10]. Shachter’s algorithm involves three simple reductions. First, any nodes that have no direct successors or “barren” nodes are removed as their value do not influence the successors and are irrelevant to the decision problem at hand, second, the propagation of the deterministic node, i.e. if any direct successor j of a node i has a CPD for which node i is a conditioning variable, that function is substituted in the distribution of j and in the process, node i is replaced as a conditional predecessor to node j by the conditional predecessors of the node i , i.e. node i is absorbed out. This may introduce new directed edges or links if not present between the predecessors of node i and node j . Third, arc reversal – if there is an arc or a directed edge from i to j , it is possible to transform the net into an arc from j to i instead and both i and j inherit each other’s conditional predecessors. The CPD for node i is found by summing out and the new CPD for node j is calculated from Bayes theorem.

Applying RNOR rule using link probabilities

Using the Java programming language and the Netica application programming interface, we programmed the RNOR algorithm as defined in [29]. Appendix A.8 details the code. The algorithm calculated the probability score of asthma in each row of the CPD using equation (1). An example combination – (s)ex = Male, (r)ace = White, (w)heez = True is given here –

$$P^R(x) = 1 - \frac{1 - P(r, w)}{1 - P(w)} \times \frac{1 - P(w, s)}{1 - P(s)} \times \frac{1 - P(s, r)}{1 - P(r)} \quad (3)$$

In the example above (equation 3), the RNOR algorithm calculated the row value for the CPD by first calculating each of the subsets, multiplying them and subtracting

from 1. For this experiment, the probability score for asthma for each combination of conditioning variables was calculated in three different ways–

Using RNOR rule

The algorithm recursively calculated the probability score for successive combinations; using the original scores, $p^E(x)$, from data learnt CPD wherever they existed (i.e., not 50-50), using equation (3). For example, if P (Wheez, Sex) was non-uniform (i.e., not 50-50) in the original data learnt CPD of asthma (from the empiric BN), it was used in successive calculations for higher order combinations. When the resulting probability was negative or exceeded 1, suggesting inhibition, the CPD row was left with 50-50.

Noisy-OR approach

Starting with link probabilities listed in Table 3-5, the algorithm recursively calculated a probability score of asthma for successive combinations of conditioning variables without using any values from the original data learnt CPD from the empiric BN. This essentially reduced the model to a Noisy-OR model.

Adaptive Noisy-OR approach

As with the RNOR calculations, the algorithm computed CPD values recursively using $p^E(x)$ where applicable. However, for those scores resulting in negative values, instead of leaving a uniform distribution, we used the Noisy-OR value, reasoning that it would be a better approximation than the 50-50 value in the CPD row. We termed this “*Adaptive Recursive Noisy-OR*” or *Adaptive RNOR (ARNOR)*.

BNs with probability tables calculated from these three methods were used in two separate tests, comparing their predictive abilities in the two test sets.

Testing the Models

The BN models were evaluated, first, using data from our original test set of RMRS data (1/3 split from the large cohort study 5187 cases) and, second, using the data set derived from the prospectively collected CHICA database (1982 cases), utilizing Netica's test interface. We compared the sensitivity, specificity, positive predictive value, and negative predictive value using Receiver Operating Characteristic (ROC) curves. [57] The ROC curve was obtained by plotting pairs of true positive rate (sensitivity) and false positive rate (1 - specificity). The area under the ROC curve (AUC) was used as a measure of overall test performance. AUCs were compared using the method of Hanley and McNeil [72] which specifically accounts for comparing ROC derived from same cases.

Results of RNOR rule Reformulation

We had 5187 cases in our RMRS test set and 1984 cases in the independent test set (CHICA). Results for the tests are shown in Figures 3-8 and 3-9 respectively. There were no statistically significant differences between the predictive ability of the RNOR or Adaptive RNOR and that of the empirically trained BN. Both RNOR and Adaptive RNOR had larger AUC than the Noisy-OR BN in both RMRS and CHICA datasets but the difference did not reach statistical significance (Figures 3-8 and 3-9 respectively and Table 3-6).

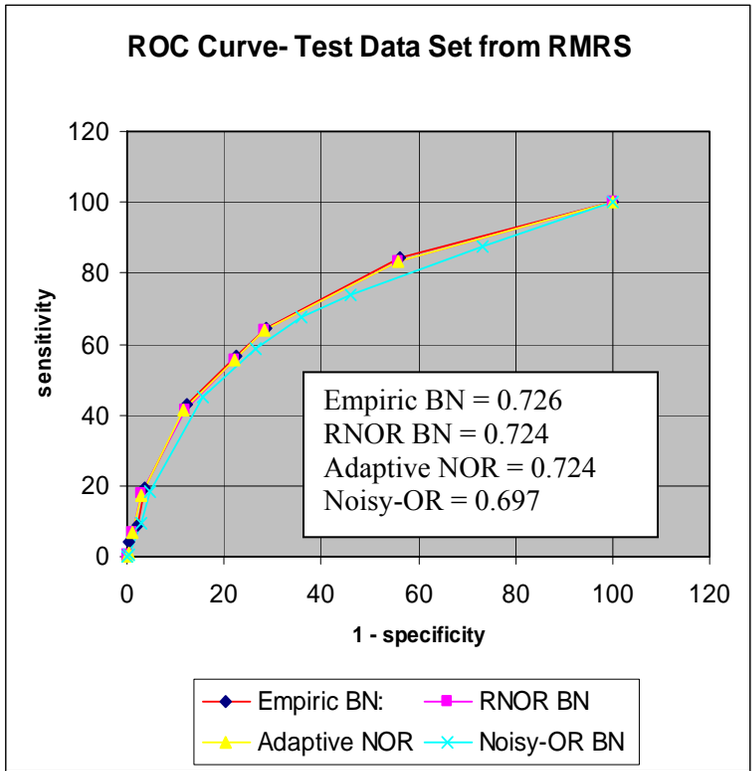


Figure 3-8 Evaluation using test data from RMRS

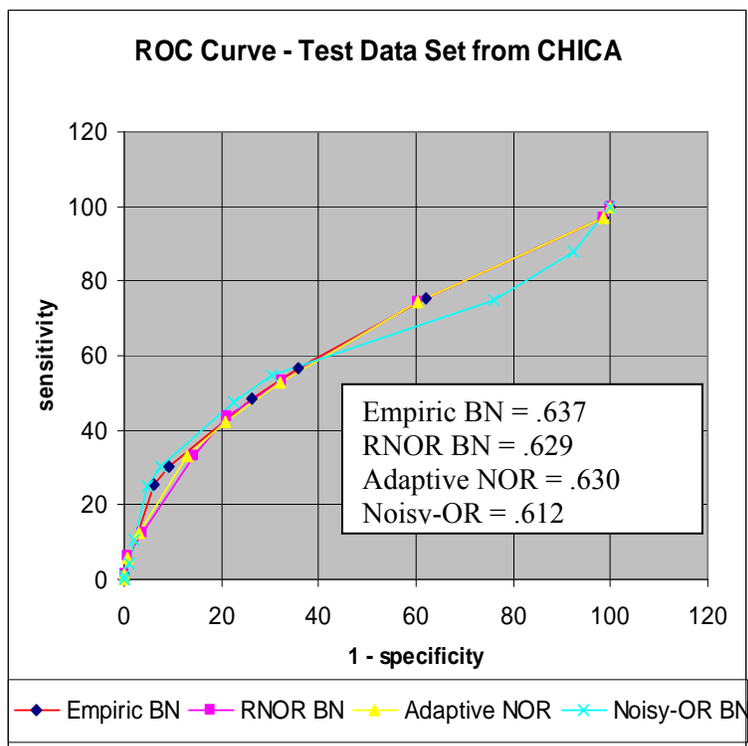


Figure 3-9 Evaluation using test data from CHICA

Table 3-6 Statistical comparison of models

Dataset	AUC1	AUC2	P-value
RMRS	Empiric BN	RNOR	0.956
RMRS	Empiric BN	ARNOR	0.968
CHICA	Empiric BN	RNOR	0.620
CHICA	Empiric BN	ARNOR	0.625
RMRS	Noisy-OR	RNOR	~1.0
RMRS	Noisy-OR	ARNOR	~1.0
CHICA	Noisy-OR	RNOR	~1.0
CHICA	Noisy-OR	ARNOR	0.851

Discussion

In case of Noisy-OR reformulation, the predictive power of the BN decayed significantly when applied to another dataset (CHICA), it did not decay as much when we made the Noisy-OR assumption in the original dataset. This is somewhat surprising since we wouldn't necessarily expect the predecessors to the asthma node (xray, drugs, wheeze) to be independent causes (or even causes at all). Nonetheless, the relatively small decrement in AUC suggests that the Noisy-OR may be a robust assumption to make in an array of situations, even if BNs trained in one setting may be less effective in another. We believe that the Noisy-OR formalism provides sound theoretic background for combining a number of causes leading to a common disease manifestation if the underlying distribution follows the assumptions.

In case of RNOR rule reformulation, we had anticipated that the RNOR would perform better than the empirically trained BN. The RNOR retains the higher level

posterior distributions that can be extracted from the data, but in the absence of cases in the data, the RNOR estimates posterior distributions that are more reasonable than a uniform distribution. This is especially true for a variable like *asthma* where the posterior probability would never be expected to reach 50%. We observed such a trend, but the differences were not statistically significant.

In our application of the RNOR we ignored any negative probability scores that are produced (as a result of recursive calculations in the algorithm). We believe these negative scores are produced where inhibition exists between subsets of dependent causes and the scores produced by the algorithm are greater than one. One such example is – *sex* = *Male*, *drug* = *True*, *xray* = *True* – inhibition between *drug* and *xray* (subset score=1.0034) and *sex* and *drug* (subset score = 1.03), though *x-ray* and *sex* subset has a score within bound (subset score = 0.969).

The implication is that the domain chosen for this study may not be an ideal application of the RNOR strategy but the RNOR algorithm can be used to detect inhibition in large datasets. Additionally, we have evaluated conditions in our dataset which render the RNOR rule inapplicable and discussed our use of Noisy-OR calculations in such situations.

For our domain, the Noisy-OR formalism produced results comparable to the empirically trained BN. Surprisingly, the RNOR did not contribute significantly more predictive power than the Noisy-OR. Therefore, we conclude that the Noisy-OR approach to combine information can serve as a satisfactory strategy for merging data in the IsBIG experiments described in the next chapter.

Using causal independence and statistical measures

In this chapter, we describe experiments that extend the use of causal independence assumption, using the Noisy-OR approach to link disparate sources in a normative form. In the absence of complete datasets of all the domain variables, we present a methodology using summary and statistical measures and the causal independence assumption.

In experiment 3, we apply the concept of linking disparate data sources for knowledge representation and inference with BN, again in the domain of childhood asthma. We combine data from our EMR (RMRS) with published data on the interaction of genotype and smoking to the risk of asthma using the causal independence assumption. Our aim is to leverage the BN representation and causal independence assumption beyond the use case of learning conditional probability distributions from independent causes. We use causal independence as a data integration strategy to learn and inference from disparate sources, for example, for testing genetic hypotheses in large clinical data sets from an EMR. We demonstrate this use case using an experimental study where data from RMRS and CHICA system are combined with statistical and summary data published in one study linking a particular genetic variant and an environmental variable (tobacco smoke exposure) to asthma in an integrated model. We evaluate the integrated model against our EMR data using a “goodness of fit” metric as a performance measure.

Our aim in experiment 4 is to integrate several published studies. For the purposes of this research, we choose the domain of genome wide association studies (GWAS) where findings link a genetic variation of Single Nucleotide Polymorphism (SNP) type

with a disease or a trait. However, as discussed in the background chapter (chapter 2), we have no publicly accessible primary data source for these studies to link with each other or with a data source like an EMR.

Therefore, in experiment 4, in the absence of any primary data source in the domain of genome wide association studies, we extend our methodology to integrate statistical measures of correlations and effect sizes from published studies to incorporate all available information in a data integration framework – In-silico Bayesian Integration as previously described in Figure 1-2. Using statistical measures of correlation, we learn conditional probability distributions of disparate BNs; these BNs are linked to each other by effect size of common nodes. The common nodes are then “absorbed” because they are hidden and not the primary variables of interest. The result of the transformation of absorbing nodes is that it preserves the joint probability distribution of the BN but may introduce new edges to preserve its quantitative structure. Therefore the integrated BN may find new relationships which otherwise may be hidden knowledge.

We believe this approach has three main advantages: 1) It can incorporate all available information across boundaries of individual datasets by either learning directly from data or assuming certain independencies in the dataset or from secondary sources such as summary and statistical measures. 2) Once a model is built, it can make inferences in context, for example, from patient data from an EMR. 3) It creates the capacity to keep adding more information as it becomes available using the same framework.

We hypothesize that this approach will discover hidden associations across silos of biomedical data, for example predictive distributions which are otherwise unknown.

We also believe that this approach can demonstrate the state of the current research by putting it in context with the patient data, for example, from our EMR. This would allow us to quantify from a domain like GWAS, for example, how much risk of common disease(s) is explained by genetic linkages. With this aim of linking disparate studies, we conduct experiment 4 detailed in following pages.

Experiment 3: Integrating Published and EMR Data

In-Silico Testing of Genotype-Phenotype Associations with Electronic Medical Records [73]

Introduction

In recent years, several published studies have reported associations of a given genetic polymorphism with a particular common complex disease. In separate clinical studies these associations are also stratified by various demographic, racial, ethnic and environmental factors. However, due to the challenges discussed in the Chapter 2, the two separate sources of information are almost never combined for use in clinical practice. Therefore, informatics methods are needed to integrate them with clinical data, for example, from an EMR to a) validate findings in larger populations and b) generate higher order hypotheses to study separately and c) for future use in data from an EMR for application in patient's context.

In this experiment, we developed a BN methodology to integrate summary and statistical data from full-text published biomedical literature with records identified from our EMR system RMRS to test genotype-phenotype associations in our clinical population. Here we report our results with the methodology in the domain of asthma risk.

With the advent of microarray technologies, clinical effects can be predicted based on functional effect(s) of a gene. [40] It has been shown that DNA mutations in coding regions effect the function or the efficiency of the protein that the gene encodes, which can lead to physiologic effects that are clinically relevant. [33] Therefore, strategies applied to genetic diseases like sickle cell anemia (a pure genetic disease) can

also be applied to common diseases like asthma, but the task is more complex due to gene-gene and gene-environment interactions. [33] However, for complex diseases like asthma, genetics studies can contribute to better healthcare outcomes in one or more ways.

A considerable number of studies point to a lack of consensus on asthma clinical subtypes. [40] Therefore a classification of asthma based on genetics and environment could provide more accurate clinical subtypes. [33, 40] Genetic classification can provide improved prognostic information including identification of patients who are at highest risk for severe life threatening episodes of asthma. In addition, a more detailed understanding of the pathophysiology of the disease can lead to a more precise definition of the environmental modifications which can most likely reduce the risk of asthma. Finally, such studies could lead to genetic testing to predict a patient's response to a drug or development of new drug therapies. [74]

We report our experimental findings for an asthma case finding application that integrates summary and statistical data derived from the full-text biomedical literature and RMRS. The published paper details the association of asthma with the Beta 2 Adrenergic Receptor (β 2AR, also referred to as ADRB2 genotype) gene polymorphism and cigarette Smoking. [50] Our methods utilize Bayesian Networks (BN) to integrate disparate data sources – a) summary data from literature b) pediatric data from RMRS and c) self reported smoke exposure data by families from CHICA system. We use a “goodness to fit” metric [75] as a measure to compare our integrated model with a clinical only model.

Methods

Pediatric Asthma cases and controls were identified from the RMRS and the CHICA system, [59] a Clinical Decision Support System (CDSS) used in our Pediatric Primary Care (PCC) practice in conjunction with RMRS. An overview of CHICA system is described in Chapter 3, experiment 1.

Data

The CHICA system electronically receives a record of all clinical observations from the RMRS database for every patient visit. For this study, we analyzed data for all children over 5 years of age in our system. Children were classified as cases or controls based on the presence of an ICD-9 code for asthma (493.*) or more than two prescriptions of an asthma medication. From the filtered set we were able to extract the variables listed in Table 4-1 to get an “Asthma Status,” sex and race for each patient (ages 5 years or older) who had a visit to the PCC clinic. We combined these data with the self reported data on environmental tobacco smoke exposure collected from the CHICA pre-screener form (PSF), a computer generated questionnaire that is given to the patient family to complete in the waiting room. The combined data set is used to build our clinical model using a Bayesian Network (Figure 4-1). Table 4-2 details the characteristics of the dataset used to build the model.

Table 4-1 Data Variables – Experiment 4

Variable	Values
Race	White, Black, Hispanic, Other, Unknown
Sex	Male, Female
Asthma	ICD9 (493.*) or any clinic billing diagnosis for asthma after age 5 (True, False)
Smoke Exposure	“Does anyone in [your child’s] home smoke” (yes, no)

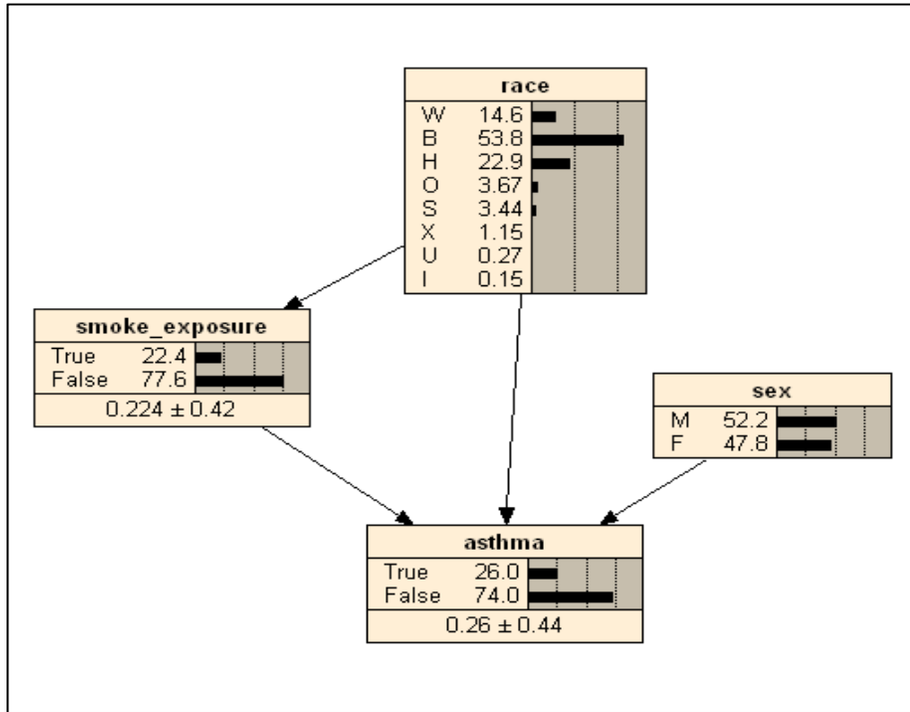


Figure 4-1 Clinical Model (with limited nodes)

Table 4-2 Baseline Characteristics – Clinical model

Variables	Training Set (n = 2609)		
	%	#	
Race	Hispanic (H)	597	23%
	Spanish (S)	89	3%
	Asian (X)	29	1%
	Islander (I)	3	< 1%
	Unknown (U)	6	0%
	Black (B)	1408	54%
	White (W)	382	15%
	Other (O)	95	4%
Sex	Female (F)	1246	48%
	Male (M)	1363	52%
Asthma	True (T)	666	26%
	False (F)	1943	75%
Smoke Exp	True (T)	579	22%
	False (F)	2030	78%

Bayesian Network for Clinical Model

We used Netica software [67] (www.norsys.com) to construct BN for our clinical model (Figure 4-1). Netica allows network construction and parameter learning from data. The model parameters were obtained using data from the CHICA database as specified in the data section. At the time of this experiment the CHICA system provided data on both asthma and smoke exposure status for about 2600 patients. The baseline characteristics of the cohort for the training set are shown in Table 4-2. The prior probability of asthma from this model is 26% (Figure 4-1).

Bayesian Network for the Genetic Model

We used a published case-control study – “*Association of Asthma with Beta 2-Adrenergic Receptor (β 2AR) Gene Polymorphism and Cigarette Smoking*” [50] to build a genetic model linking genotype to smoke exposure and asthma. The study reported an interaction between cigarette smoking and β 2AR-16 genotype. It showed a synergistic relationship between tobacco smoke exposure and the Arg-16 homozygous genotype with respect to asthma. When compared with Gly-16 homozygotes who never smoked, the smokers who were Arg-16 homozygotes had a significantly increased risk of asthma (Odds ratio = 7.81). We used summary data from this study, adjusted for our population’s asthma prevalence (26%), in our clinical model. Since the study reported 128 cases, our experiment required 364 controls (for a total of 492 subjects, resulting in a 26% prevalence). Therefore, we multiplied the number of controls in each genotype and smoking status group by a factor*** ($f = 2.68$) (Table 4-3). We constructed a BN using Netica software and learnt its parameters using the summary data in Table 4-3. This constituted our genomic BN (Figure 4-3).

**Table 4-3 Literature Summary data adjusted for Asthma prevalence
(From our EMR)**

β 2AR-16 genotype	Smoking Status	Cases (n)	Controls (n)	Adjusted Controls (Asthma prevalence = 26%) ***
GG	Never-smokers	16	28	75.04
AG	Never-smokers	43	52	139.36
AA	Never-smokers	30	33	88.44
GG	Ever-smokers	6	6	16.08
AG	Ever-smokers	11	12	32.16
AA	Ever-smokers	22	5	13.4
Total		128	136	364

$$0.26 = \frac{\# \text{ cases } (n = 128)}{\# \text{ cases } (n = 128) + \text{Total } \# \text{ controls } (? = 364)}$$

$$\text{factor } (f) = \frac{\text{Total } \# \text{ controls } (n = 364)}{\# \text{ controls } (n = 136)} = 2.68$$

The prevalence of β 2AR-16 allele frequencies in the genomic network from this cohort is Arg-16 (A) – 56.4% and Gly-16 (G) – 43.6% (Figure 4-4).

Integration of the two models

The β 2AR genotype can be considered a risk factor or hidden node affecting asthma for the clinical model; specifically accounting for the effect of smoke exposure. In order to integrate this genetic relationship into a clinical model, we needed an observable proxy for genotype status. For this experiment, race was considered as a surrogate for the β 2AR genotype in absence of actual genotype data for the patient.

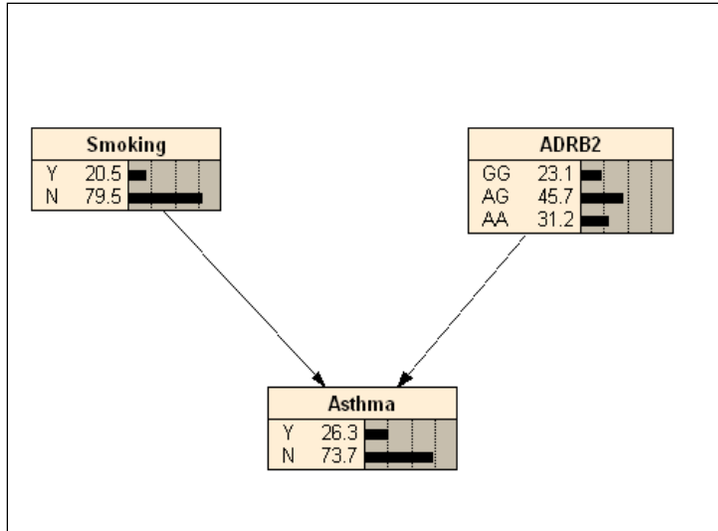


Figure 4-2 Genomic Model (Genotype)

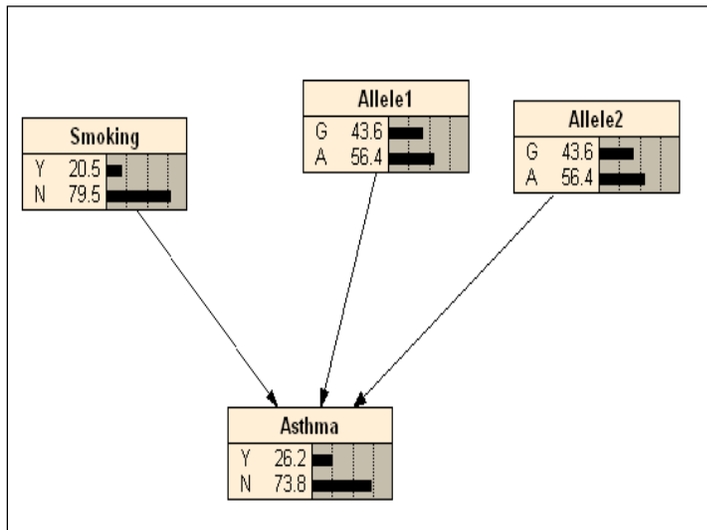


Figure 4-3 Genomic Model (Alleles)

Table 4-4 Allele Distribution by Race from public sources

Race	Allele A (%)	Allele G (%)
White (W)	34.0	66.0
Black (B)	45.0	55.0
Hispanic (H)	44.0	56.0
Other (O)*	50.0	50.0
Spanish (S)	44.0	56.0
Asians (X)	55.0	45.0
Unknown (U)*	50.0	50.0
Islander (I)*	50.0	50.0

*Uniform distribution assumed since no info found

Table 4-4 lists the distribution of alleles A and G by race for β 2AR genotype. We obtained this distribution from publicly available databases – ALFRED (<http://alfred.med.yale.edu>), PharmGKB (<http://www.pharmgkb.org>) and entrez SNP (<http://www.ncbi.nlm.nih.gov/snp>) databases. Using this observable proxy, we did three things to integrate as follows –

We first inserted the two explicit nodes for β 2AR alleles (Figure 4-3) in the clinical model (Figure 4-1) and used the distribution described in Table 4-4 to obtain the conditional probability distribution (CPD) of allele1 and allele2 given race.

Second, since the CPD of asthma in the genetic model reflects the findings that are reported in the presence or absence of smoke exposure in the study, we used the adjusted published distribution (i.e. Figure 4-3) to replace the CPD of the asthma node in our clinical model (Figure 4-1.). In the integrated model, we kept the CPD for smoke exposure node from the clinical model since this is our population of interest.

Third, we compiled the integrated model using Netica’s compilation tool. This operation creates a “junction tree” for “belief updating” and also calculates the full joint probability distribution (JPD) of the resulting network – our integrated model is shown in Figure 4-4.

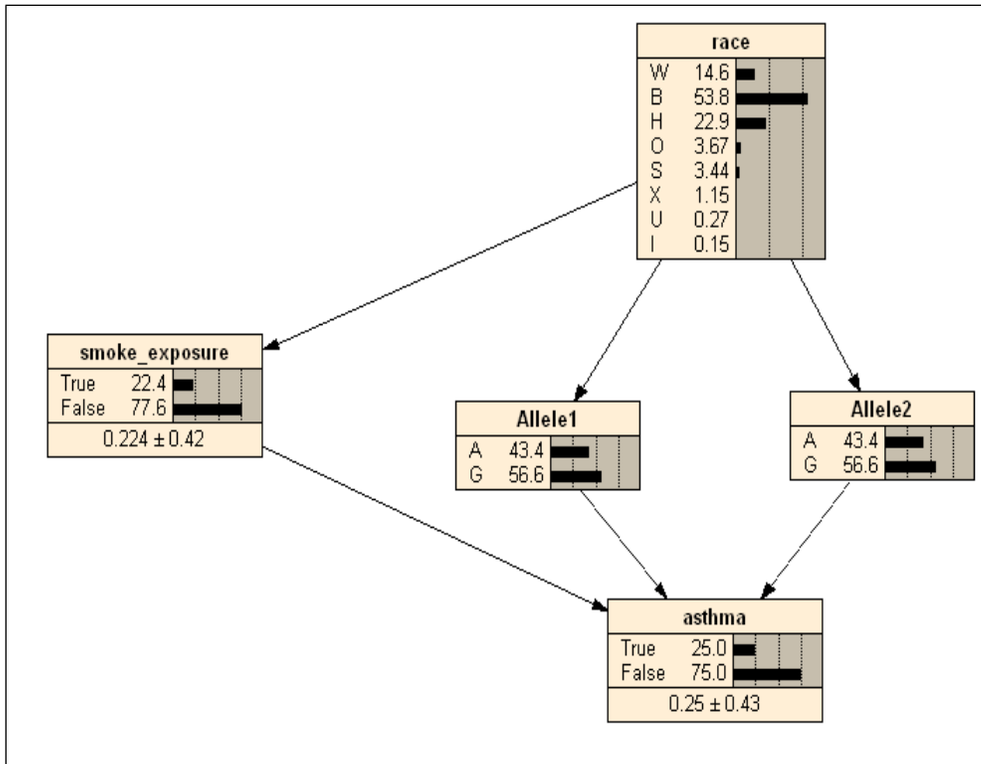


Figure 4-4 Integrated model – Clinical and Genomic

The resulting prior probabilities of β 2AR-16 alleles in this integrated model are – Arg-16 (A) – 43.4%, Gly-16 (G) – 56.6%. They are different from the prevalence found in the published study (A - 56.4% and G - 43.6%) and reflect the inferred distribution in our population based on racial distribution. The network also compiled a resulting prior probability of asthma (25%); quite close to the marginal prevalence in our clinical model (26%), suggesting that the relationship between race and asthma could be mediated by β 2AR-16 genotype. In order to compare the two models – clinical with integrated, we absorbed the allele nodes in the integrated model (Figure 4-5).

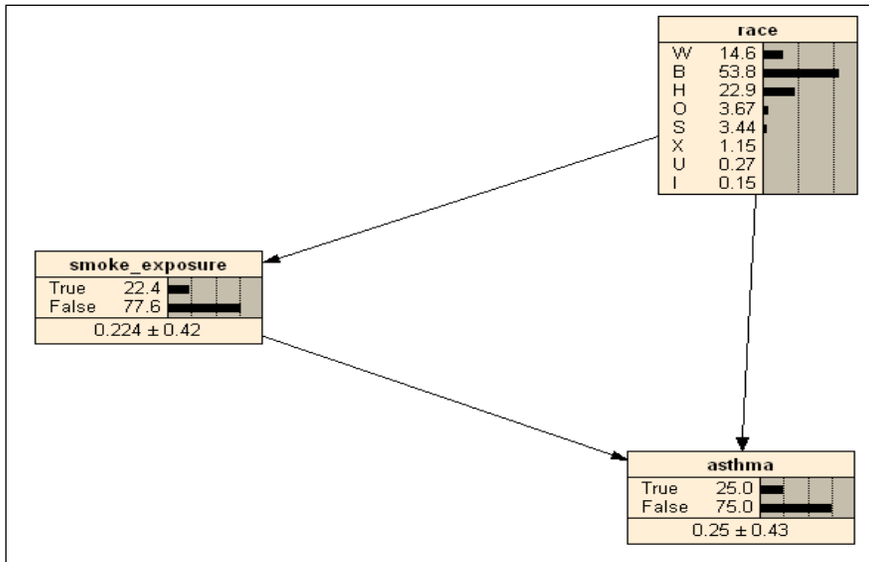


Figure 4-5 Model (with Allele nodes absorbed)

Also known as “averaging out” or “summing out a variable,” a node absorption transform leaves the full joint probability distribution of the remaining nodes unchanged and results in the final integrated model (Figure 4-5) that could be used for comparison.

Comparing the two models

We compared the CPD of asthma node for the clinical and integrated models to see how well the two distributions match. Our hypothesis was that associations between genotype and phenotype could explain, at least in part, the clinical findings that we observed in our population, specifically as the relationship between asthma and tobacco smoke exposure. Table 4-5 lists the distribution of the asthma node in both models.

Goodness of fit Metric

We used a “Goodness of fit” as the comparison metric. [75] This metric is based on the Chi-square distribution and the method of least squares. [75] The chi-square test compares the observed frequencies with the expected frequencies, giving the probability of observed differences under the null hypothesis (there is no difference). [76] The method of least squares is built on the hypothesis that the optimum description of a set of

data is one which minimizes the weighted sum of squares of deviations of data (y_i) from the fitting function $y(x_i)$. [75] The variance of the fit s^2 , which is an estimate of the variance of data σ^2 , characterizes this sum for $v = N - n - 1$ number of degrees of freedom, where n coefficients fit to N data. Variance of fit s^2 is also characterized by the statistic χ^2 as described in equation 1 below. Equation 2 describes the *reduced chi-square* χ^2_v

$$\chi^2 = \sum \left\{ \frac{1}{\sigma^2} [y_i - y(x_i)]^2 \right\} \quad (1)$$

$$\chi^2_v = \frac{\chi^2}{v} = \frac{s^2}{\sigma^2} \quad (2)$$

where σ^2 in equation 2 is the weighted average of individual variances. However, the estimated variance of fit s^2 is also a characteristic of both the spread of the data and the accuracy of the fit. [75] Therefore its definition as a ratio of the estimated variance s^2 to the parent variance σ^2 (times the number of degrees of freedom) from equation 2 makes it a convenient measure of the “goodness of fit.” If the fitting function is a good approximation to the parent function, the estimated variance s^2 should agree well with the parent variance σ^2 and the value of the reduced chi-square should be approximately unity $\chi^2_v = 1$. [75] Furthermore, the Q-Statistic defines the probability that a function $Q(\chi^2/v, v)$ for a set of deviations obtained by *randomly sampling N observations* from normal distribution would *exceed* the value for χ^2/v that was obtained by the fitting function. [75]

Results

Table 4-5 CPD of Asthma node in clinical and integrated models

Smoke Exposure	Race	Clinical Model $y(x_i)$	Integrated Model (y_i)	Chi Square
F	W	27.21	21.68	0.71
F	H	19.00	22.52	0.29
F	S	14.59	22.52	1.45
F	B	30.70	22.60	1.52
F	O	23.75	22.99	0.01
F	U	22.61	22.99	0.00
F	I	36.94	22.99	4.50
F	X	24.63	23.35	0.04
T	W	21.59	31.86	2.44
T	H	25.80	34.32	1.68
T	S	36.94	34.32	0.16
T	B	27.49	34.61	1.17
T	O	31.92	36.17	0.42
T	U	42.04	36.17	0.80
T	I	42.04	36.17	0.80
T	X	30.45	37.91	1.29
			Variance (σ^2) = 43.25	$X^2 = 17.28$
		v = # of degrees of freedom	$v = N - n - 1$ $= 16 - 2 - 1 = 13$	$X^2_v = X^2/v = 1.33$

We calculated the value of reduced chi-square $\chi^2_v = 1.33$, (p-value = 0.187) for 13 degrees of freedom ($N = 16$ for $n = 2$ coefficients- race and smoke exposure) and $Q(\chi^2/v, v)$ to be $0.10 < Q < 0.20$. A p-value of 0.187 suggests the two distributions for clinical and integrated models may be similar. Q is “small” ($0.10 < Q < 0.20$) therefore the fit is “poor”. Only 10–20 % of large number of trials would result in a chi-square value as large as observed. Therefore, β 2AR genotype (ADRB2 status), as inferred by race, explains only a fraction of the risk of asthma association with smoke exposure in our suggested integrated model.

Discussion

Our results are consistent with a minor contribution of the β 2AR gene to the association between asthma and smoke exposure observed in our EMR. One explanation is that the association between the genotype (ADRB2 or β 2AR) and the clinical condition (asthma) in the literature was reported with “active smoking” in an adult population. Our experiment models these associations for a pediatric population where the children are exposed to environmental tobacco smoke or “passive smoking.” Asthma is also thought to have a multigenic etiology. The incorporation of other associated genes into the model is likely to improve its predictive power. Finally, the data associating race with genotype is crude and incomplete. Therefore, race (the only surrogate available to us) is a poor surrogate for genotype. A better surrogate would be helpful.

Conclusion

We have developed and demonstrated a methodology for integrating a published data source with a primary data source – EMR. This methodology can be useful for testing genetic hypotheses in large clinical data sets. The approach is applicable to a wide range of genotypes that are associated with two or more clinically observed phenomena (in this case asthma and smoke exposure). We believe this approach is worth investigating for other common diseases like diabetes and obesity and their associations with genomic and environmental findings.

Experiment 4: Integrating Disparate Sources of Summary Data

In-silico Bayesian Integration of GWA Studies (IsBIG)

Introduction

Our aim for this research is to develop probabilistic methods to integrate disparate sources of data to form a coherent model even when no primary data are available using secondary sources of information such as published summary and statistical measures for knowledge discovery. In the last experiment, we demonstrated integration of genetic, clinical and environmental sources using both data and published data summaries. To demonstrate how this approach can be used to link secondary sources of data, we choose the domain of genome wide association studies (GWAS). Using the NIH compiled catalog of GWAS (www.genome.gov) and the database of human genome variations from the international HapMap project (www.hapmap.org), we combine information from these two secondary sources using BN framework as described in our previous experiments. We call this model – In Silico Bayesian Integration of GWAS or IsBIG. In this experiment we describe the methodology and report our preliminary results. In the next chapter we formally evaluate the IsBIG model using data from our EMR and published literature identified in the Pubmed database.

Genome Wide Association Studies (GWAS)

Genome Wide Association Studies (GWAS) have become the standard to report associations of a genetic variation type – Single Nucleotide Polymorphism (SNP) with a particular disease such as diabetes, heart and lung disease, autoimmune and psychiatric disorders. It is only in very recent years with the advent of microarray technology and the mapping of the variations of human genome (<http://www.hapmap.org>) [77-78] that the

tools to conduct genome wide scans for finding associations of gene to disease have become available. The first results of the genome wide scans started to appear in 2005 [79] and since have become increasingly sophisticated in the number of gene loci that they can address. Since the genome wide scans follow a specific methodology, they are now known as Genome Wide Association Studies (GWAS).

At the heart of any GWA study is a cohort of individuals with a known disease or trait status and a comparable control group without the disease or trait. Their whole genome is genotyped for known variations of Single Nucleotide Polymorphism (SNP). Such SNP variations have been cataloged among diverse populations in the international HapMap [78] project. The results from the GWAS are analyzed for strong statistically significant (below $p < 5 \times 10^{-8}$) associations between each SNP and the disease or trait status of interest.

Thus, GWAS differ from traditional genetic linkage studies of the past where a hypothesis driven candidate gene approach was being used. GWAS, to date, have amassed large datasets linking genotype to phenotype and have provided many useful insights [80] into certain diseases such as specific forms of cancer [81] and drug metabolism, for example warfarin. [82] However, they are resource intensive and require large sample sizes to detect even modest effect sizes, and yet their applications for defining new therapies or preventive measures are largely unknown. [83] This is mainly because the influence of the genetic variation to the phenotype is unclear and there is a need to point to the causal variants; to move beyond the process of gene identification. [84]

But as GWAS (often conducted by individual groups) reported ever more potential etiologic and functional implications for similar diseases or traits [85-88], there was a need to share the results beyond publication. Therefore, summary and statistical results from these studies have been cataloged as an online resource for future investigations at <http://www.genome.gov/gwastudies> [83] at NIH.

This catalog contains the following details on each study – population characteristics, initial sample size, sub population type, strength of statistical association as odds ratio (OR) or a beta coefficient (Beta), and the frequency of the risk allele in the study. [83] However, to the best of our knowledge no secondary use of this catalog has been reported.

In this catalog, the risk allele is the marker for the SNP that is found to be strongly associated with the disease or trait. Additionally, from previous knowledge, the reported SNPs in GWAS are known to follow *non-random patterns of association* between alleles from different markers. [89] This is also known as linkage disequilibrium (LD) [90], which we describe below.

Linkage Disequilibrium (LD)

LD is non-random association between SNP alleles from different markers, i.e. SNP loci on the same or different chromosomes. Different geographic populations have different allele frequencies, and therefore, LD differs between them. Thus the non-random pattern of allele associations varies by sub populations – for example European descent, Japanese ancestry, etc. [89, 91] LD is measured using correlation coefficient (r^2) [90] and varies between 0-1. Two alleles have a high LD score if the r^2 value between them is greater than 0.5 meaning that they are associated with each other in the population.

The international HapMap consortium (<http://www.hapmap.org>) [77] has developed a haplotype map of the human genome, the HapMap, describing common patterns of human DNA sequence variation. It has enabled LD data to be readily cataloged and available in sub populations. Thus LD data quantitatively associates SNPs with other SNPs and can be measured in a sub population using a tool such as SNP annotation and proxy search (SNAP).

The SNAP tool, [92] available from Broad Institute at (<http://www.broadinstitute.org/mpg/snap>) computes LD scores (r^2) between SNPs up to 500 kilo base pairs apart and takes as input a list of SNPs of interest, a threshold value for r^2 above which to search the database and the sub population of interest. The output of the tool is a paired list consisting of the input SNP and another SNP henceforth called “proxy” SNP that exists in a non-random association above the given threshold cutoff value of r^2 in the HapMap database for the desired sub population.

Disparate Sources of information

We reasoned if diseases can be linked with SNPs and SNPs with one another, then it should be possible to predict associations among diseases that could be mediated through linked genetic determinants. Predicting quantitatively the associations among diseases and traits that would be predicted by genetic patterns alone would allow us to undertake three novel studies: (1) Validate the associations among diseases and traits with electronic health record data (2) quantify the amount of variation among disease linkages that can be explained by currently cataloged GWAS and linkage studies, (e.g. heart disease and obesity) and (3) predict novel associations among diseases that could be tested in future genetic studies.

In this experiment, we build a computational model (IsBIG) to integrate the two disparate sources – 1) results from the studies cataloged in the NIH catalog [83] linking SNPs to diseases or traits and 2) HapMap (www.hapmap.org) data associating SNPs to other SNPs. With the IsBIG model, we hoped to find a disease map linking diseases to other diseases that would be similar to what one would find in a clinical database such as our EMR.

Methods

In-silico Bayesian Integration of GWAS (IsBIG) model

The IsBIG model was assembled in the following four steps. 1) Extract Gene-Disease associations from the NIH catalog for a sub population to form a model catalog; 2) for the SNPs listed in this model catalog, compute genome wide LD scores (r^2) from a comparable sub population from the HapMap database to form a SNP-SNP (proxy) association dataset; 3) for the SNPs linked by LD in the SNP-SNP (proxy) dataset, find diseases or traits linked to each other pair wise as a result of LD. This formed the SNP-SNP-Disease-Disease dataset for input to the IsBIG algorithm described in the following sections. 4) Using this input, the IsBIG algorithm computes a BN with two separate components in steps – a) SNP-SNP BN using LD (r^2) to calculate CPDs. b) SNP-Disease BN, using OR to calculate CPDs. Finally, the algorithm absorbs the SNP nodes (as described below) to compute the disease-disease BN or the output of disease map. The algorithm is coded using Java API and a commercially available software package for modeling Bayesian networks – Netica’s API (www.norsys.com) [67]. The details of each step are listed as follows.

Extract Gene-Disease Association

A copy of the GWAS catalog from the <http://www.genome.gov/gwastudies/> website was downloaded on 12-28-09. The GWAS catalog contains information from a wide range of GWAS studies associating specific SNPs to diseases or traits. From the variables included in the catalog we chose the variables listed in Table 4-6 for the experiment. Table 4-7 lists sample studies from the GWAS catalog. Appendix A.1 gives a summary of the studies from the GWAS catalog we used.

Table 4-6 Variables for Model Catalog

DISEASE/TRAIT	Disease or trait examined in study
SNPS	Strongest SNP
INITIAL SAMPLE SIZE	Sample size for Stage 1 of GWAS, population subtype
RISK ALLELE FREQUENCY	Reported risk allele frequency associated with strongest SNP
OR or BETA	Reported odds ratio or beta-coefficient associated with strongest SNP risk allele

Table 4-7 Sample Studies in GWAS catalog

Disease /Trait	Risk Allele	SNP	Risk Allele Freq	OR	Initial Sample Size
Type 1 diabetes	G	rs4900384	0.29	1.09	7,514 cases, 9,045 controls
Type 2 diabetes	C	rs4607103	0.76	1.09	4,549 cases, 5,579 controls
Multiple sclerosis	A	rs1335532	0.87	1.28	1,618 cases 3,413 European ancestry controls

We planned to evaluate the model disease map with data from our EMR, the RMRS [2]. Therefore we decided to use only those studies from the downloaded catalog

where the initial population originated from European ancestry. Thus, we filtered studies where the initial sample size variable did not contain population from European descent (i.e. Japanese, Chinese, Korean, Gambian and other sub populations in the NIH catalog). Where the sub population was not defined, we assumed it to be of European ancestry.

2564 GWAS studies consisting of 1708 unique SNPs from the GWAS catalog were downloaded (in Dec. 2009). Of these 807 GWAS studies (Appendix A.1) qualified as conducted in population with European ancestry. These studies contributed 182 unique disease traits and 850 unique SNPs in this sub population for our *Model Catalog*.

Gene (SNP) – Gene (SNP) Association

We extracted gene-gene associations from linkage disequilibrium data in the HapMap database. [77] Since we decided to use studies with European ancestry population, we used the HapMap CEU dataset (phase 3 release r2). Using the SNAP tool [92] we derived LD between SNPs reported in our *Model Catalog* and other SNPs present genome wide.

For our experiment, we used a LD score (r^2) cutoff threshold value of 0.3. This threshold was chosen empirically to capture *weaker associations between SNPs* and yet keep our SNP to SNP (proxy) dataset computationally tractable. Higher and lower values of cutoff threshold brought in fewer and more SNPs with stronger and weaker associations respectively. However, empirically, this threshold ($r^2 = 0.3$) provided an optimum set for this experiment. Table 4-8 gives a snippet of the output from the SNAP tool for our input SNP (e.g. rs2191566).

Table 4-8 Sample output from SNAP tool

SNP	SNP (proxy)	Distance	r^2
rs2191566	rs7255512	63311	0.616
rs2191566	rs8104605	77627	0.604
rs2191566	rs4803675	78327	0.561

Pair wise disease-disease associations using LD

When we searched in the HapMap phase 3 release 2 CEU dataset using the SNAP tool, the 850 unique SNPs were in LD ($r^2 \geq 0.3$) with more than 16,000 proxy SNPs. From these 16,879 proxy SNPs, we were only interested in the proxy SNPs that were also reported in our *Model Catalog* and their associated diseases or traits. Therefore, we imported the proxy SNP data produced by the SNAP tool and our *Model Catalog* in a database. From there, we were able to do association mining to obtain SNPs that were both in our model catalog and had a LD ≥ 0.3 with another SNP (proxy) in the *Model Catalog* and obtain its associated disease or trait.

This resulted in 397 unique SNP to SNP pair-wise associations and their associated disease or trait relationships (Appendix A.6) where LD between SNPs was < 1 (i.e. $r^2 \neq 1$). The lowest LD found in this association mining step was 0.302 and the highest LD was 0.983 between two SNPs. Association mining with LD data from HapMap provided us the ability to examine the model GWAS catalog beyond the pair wise single SNP to disease or trait association. Table 4-9 shows some associations found between diseases or traits using the LD scores of correlations (r^2) derived from HapMap.

Table 4-9 Pair wise associations from GWAS catalog (by association mining)

Disease or Trait associated with a SNP	Disease or Trait associated with a SNP (proxy)	r^2
Coronary Disease	Glioma	0.384
Rheumatoid arthritis	Inflammatory bowel disease	0.389
Celiac disease	Schizophrenia	0.400
Schizophrenia	Celiac disease	0.400
Primary biliary cirrhosis	Systemic lupus erythematosus	0.425
Type 1 diabetes	Rheumatoid arthritis	0.796

However, traditional statistical methods like correlation can only examine pair wise relationships, and additionally carry the burden of identifying the actual functional relationships (in this case between the SNP and the proxy SNP) and suffer from multiple testing issues as well. Therefore, for analyzing functional relationships between common diseases studied in GWAS, we need methods that handle complexity beyond the pair wise paradigm. This is where Bayesian methods are useful – they compute the probability of hypotheses rather than probability of committing an error [93] and model multiple random variables whose probability distributions can be factorized into smaller conditional probability distributions (CPD). [21]

We use this property of BN, to compute CPDs from available subsets of data, i.e. from statistical data reported in the GWAS and from LD measure of correlations from HapMap database, to form smaller BNs which, on absorbing the SNP nodes, combine in one large BN – our disease map. We describe the In-silico Bayesian Integration of GWAS (IsBIG) algorithm below.

In-Silico Bayesian Integration of GWAS (IsBIG) Algorithm (Figure 4-10)

We used the BN approach and causal independence assumption described in Chapter 2 to draw a DAG of the SNP to SNP and SNP to disease or trait relationship. The qualitative structure of this model is assembled from the input SNP-SNP-Disease-Disease dataset (formed from association mining as described above) as follows. The complete table is listed in Appendix A.6.

A directed edge connecting a SNP to another SNP (proxy) node is drawn programmatically (using Netica API) if the SNPs are correlated (LD correlation) in the input dataset. They assume a parent-child relationship from SNP to SNP (proxy). Similarly a directed edge connecting each SNP to one or more diseases or traits is drawn if they are correlated in the input GWAS data. Please note that in this dataset, there are several SNP to SNP links, and therefore, there are N way interactions because each SNP may be correlated with several others. Similarly, multiple SNPs can be correlated with a disease. These *multiply connected* SNPs present a problem in computing the conditional probability distributions (CPD) of the nodes that have more than one parent because the data sources only describe one to one association metrics. We address this problem of multiply connected SNPs with the method of partial correlations described below.

Figure 4-6 below depicts the DAG of the model GWAS catalog constructed for this study. In this figure $S_1, S_2 \dots S_n$ are the SNPs in the *Model Catalog*, they are in LD (r^2) with each other (from HapMap data) and have been shown to have strong associations in GWAS with their respective diseases or traits (e.g. S_2 to asthma). The odds ratio (OR) quantifies the strength of each of the relationships between the SNP

and the disease as derived from a GWAS. In this figure, SNP S_3 is multiply connected with S_1 and S_2 as parents and the disease Psoriasis has both SNPs S_4 and S_5 as parents. However, each of these represent separate BNs derived from individual pieces of information and combined into a single BN. The IsBIG strategy for deriving the parameters follows –

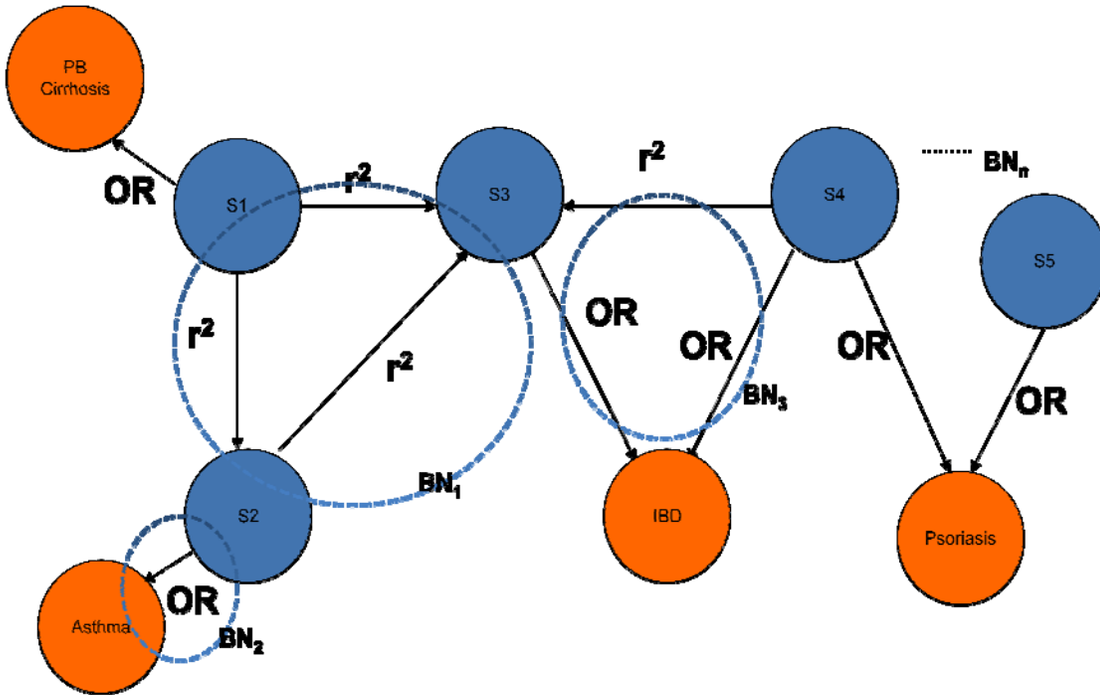


Figure 4-6 Model DAG of SNP-Proxy SNP-Disease

Computation of SNP – SNP BN Parameters

The task at hand is to compute the CPD of each SNP – we need to calculate the CPD of a SNP that has no parent (S_1), 1 parent (S_2) or many parents (S_3). As shown in Table 4-7, the GWAS catalog lists a risk allele (RA) frequency (RAF) for each SNP to disease or trait association. The normal allele (NA) frequency (NAF) is the complement of RAF, i.e. $-(1-RAF)$. Therefore, for a SNP that has no parent (e.g. S_1 in Figure 4-6),

the two cells in CPD of S_1 , i.e. $P(S_1 = RA)$ and $P(S_1 = NA)$ is simply the prevalence of the RA, i.e. RAF and its complement (1-RAF).

For a SNP like S_2 that has S_1 as a parent, i.e. $S_2 | S_1$, (Figure 4-6), the CPD can be constructed using the RAF and NAF of both SNPs and the correlation coefficient (r^2), the linkage disequilibrium measure between the SNPs. We need to compute the four cells for the CPD of S_2 , i.e. $P(S_2 = RA | S_1 = RA)$, $P(S_2 = NA | S_1 = RA)$, $P(S_2 = NA | S_1 = RA)$ and $P(S_2 = NA | S_1 = NA)$. By definition of LD (Table 4-10), if T_1 and T_2 are RAF for S_1 and S_2 from the catalog, the four cells in the CPD of S_2 given S_1 i.e. $S_2 | S_1$ can be computed by calculating the deviation measure (D) between the two SNPs. D is a measure of deviation from *linkage equilibrium which is a random association* of the two SNP alleles and is calculated by equation 1. [90].

$$D = r * \sqrt{T_1 * (1 - T_1) * T_2 * (1 - T_2)} \quad (1)$$

Table 4-10 CPD from Linkage Disequilibrium and Risk Allele Frequency

$P(S_2 S_1)$	Risk Allele (RA)	Normal Allele (NA)	Total
RA	$X = T_1 * T_2 + D$	$Y = T_1 * (1 - T_2) - D$	T_1
NA	$V = (1 - T_1) * T_2 - D$	$W = (1 - T_1) * (1 - T_2) + D$	$1 - T_1$
	T_2	$1 - T_2$	1

For multiply connected SNP nodes, e.g. S_3 in figure 4-6 which has 2 parents, S_1 and S_2 , we cannot use the simple approach as described above because of the SNP-SNP triangulations formed as in Figure 4-7. In such situations, we use partial correlations to first assess the strength of the relationship between SNP pairs as described below.

Computation of CPD for multiply connected SNP

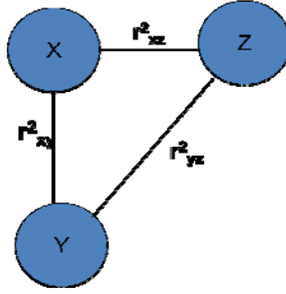


Figure 4-7 SNP-SNP Triangulations

Partial correlation quantifies the correlation between two variables x and y when conditioning on one or several other variables [94], for example z . The 0th order correlation is the regular correlation. The 1st order partial correlation between variables x , y and z in Figure 4-7 is given by equation 2 below. For example, $r_{xy \cdot z}$ is the correlation between parts of x and y that are uncorrelated with z .

$$r_{xy \cdot z} = \frac{r_{xy} - r_{xz}r_{yz}}{\sqrt{(1 - r^2_{xz})(1 - r^2_{yz})}} \quad (2)$$

For calculating the CPD of a multiply connected SNP (e.g. S_3 in Figure 4-6 which has 2 parents, S_1 and S_2), the 1st order partial correlations are computed, ignoring the directionality of the link to first assess whether the strength of the relationship between the pairs S_1 - S_2 , S_2 - S_3 and S_1 - S_3 are above a predetermined threshold value. If the partial correlation is below this threshold value (for example, 0.5), the link that has low partial correlation value in the S_1 - S_2 - S_3 triangulation is removed thus breaking the triangulation. If no links are removed, we use partial correlation coefficients ($r_{13 \cdot 2}$, $r_{23 \cdot 1}$) for S_1 - S_3 and S_2 - S_3 links to calculate two deviation measures (D) in equation, one for each parent, i.e. $P(S_3 | S_1)$ and $P(S_3 | S_2)$. Now we have two sets of CPDs (four cells) for S_3 , one from each

parent that we need to combine. Since these are independent linkages, i.e. S_1 - S_3 and S_2 - S_3 , we can combine them assuming causal independence using a Noisy-OR calculation as described in Chapter 3. We can apply the same calculations for more than two parents, however in our *Model Catalog*, the links were pruned and we rarely needed to calculate beyond the single deviation measure.

Thus, applying partial correlations to our scenario pruned the links between SNPs when there were triangulations of SNPs present and reduced the computational complexity of calculating the CPDs of multiply connected SNPs. Specifically, links that were weaker than a predetermined threshold value of partial correlation coefficient were removed. For our experiment, we used a threshold value of 0.2 for 1st order partial correlation. As we show later in the results, the threshold of partial correlations defined how sparsely or densely the resulting network connected.

Computation of SNP – Disease BN Parameters

For each SNP-disease association to be modeled from a GWAS, the only data that are available are the odds ratio (OR). Given a prior probability (aka disease prevalence) of a disease or trait and the strength of the relationship (OR) from the GWAS, a posterior probability (CPD) can be derived.

To derive a CPD from the prevalence $P(D_i)$, and an odds ratio, $P(D_i)$ must first be converted to the prior odds (O_{prior}) of the disease according to equation 3. From there, the posterior odds of the disease $O_{\text{posterior}}$ can be calculated using prior odds and the odds ratio reported in the GWA study (O_{gwa}) as in equation 4. [95]

$$O(\text{prior}) = \frac{P(D_i)}{1 - P(D_i)} \quad (3)$$

$$O(\text{posterior}) = O(\text{prior}) * O(\text{gwa}) \quad (4)$$

From posterior odds of disease (or trait) we can find the probability P_i of the disease given the SNP, i.e. $P(D_i | \text{SNP})$ by equation 5 below.

$$P_i = P(D_i | \text{SNP}) = \frac{O(\text{posterior})}{1 + O(\text{posterior})} \quad (5)$$

P_i is the link probability [27] of the SNP to the disease or trait node in absence of any other cause. For disease or trait nodes that have multiple links from SNPs (e.g. S_4 to IBD and Psoriasis in Figure 4-6), we assume causal independence of SNP to disease (as in the GWAS). Therefore, in those situations, we calculate the CPD of the disease given the SNPs, i.e. $P(D | S_1, S_2, \dots)$ using the Noisy-OR equation below (equation 6).

$$P(D / S_p) = 1 - \prod_{s_i \in S_p} (1 - p_i) \quad (6)$$

The disease or trait's prevalence $P(D_i)$ measure can either be specified by a domain expert or from data, for example, from an EMR. Since in this experiment we are developing the methodology, we arbitrarily assumed certain prevalence of common diseases. For a formal evaluation of the IsBIG model in the next chapter, we used prevalence measures from RMRS, our EMR.

Deriving Disease – Disease Map

We have constructed two components of a BN: SNP to SNP and SNP to disease from two separate sources of information. Our goal in the next chapter will be to evaluate the predictive power of this BN against clinical data from the RMRS. To do this, we wish to reduce the BN to a disease map by absorbing out the SNP nodes which are unobserved in clinical data. Node absorption is a network transform and is described in the experiments in Chapter 3.

Results

Our preliminary results show that from the data in our model catalog, the IsBIG model linked clinically related nodes of diseases or traits on absorbing the SNP nodes. For example, testicular germ cell tumor was linked to testicular cancer, and chronic obstructive pulmonary disease was linked to lung cancer, which was also linked to lung adenocarcinoma. Similarly, coronary disease was linked to early myocardial infarction. Some of the diseases the model linked are given in Table 4-11.

Table 4-11 Disease linkage patterns from GWAS catalog

Diet–Environment	Obesity, Type 2 diabetes
Autoimmune1	Type 1 diabetes, Celiac disease, Inflammatory bowel disease
Autoimmune2	Psoriasis, Crohn’s Disease, Inflammatory Bowel Disease Celiac Disease, Multiple Sclerosis
Cardiac	Coronary Disease, early Myocardial infarction, Intracranial Aneurysm, LDL cholesterol
Lung	Lung cancer, Lung adenocarcinoma, Chronic obstructive pulmonary disease
Cancer	Colorectal cancer, Prostate cancer
Traits	Blond vs Brown hair color, Skin sensitivity to sun, Freckles, Red vs non-red hair color, Melanoma

We conducted normative evaluation of the IsBIG model. We used two part criteria by varying the following network parameters – 1) LD threshold between SNPs and 2) Partial correlation threshold used for pruning the SNP links and their effect on the number of nodes and connectivity in the network.

Linkage Disequilibrium Threshold

LD threshold (r^2) was chosen as $r^2 \geq 0.1$, $r^2 \geq 0.3$, and $r^2 \geq 0.5$, corresponding to weak and strong associations between SNPs. Based on the LD threshold value used, we expected the number of SNPs brought in the model from the HapMap dataset to increase or decrease. As a result of this variation, the number of disease pairs that became part of the model changed as well. Table 4-12 below details the number of SNP-SNP pairs and disease pairs that entered the model based on the choice of LD threshold.

As can be seen in Table 4-12 (from Disease Pairs column) changing the network parameter for linkage disequilibrium (LD) threshold between SNPs changes the number of disease nodes that become part of the model. The number of disease nodes that entered the model varied inversely to the LD threshold value chosen.

Table 4-12 Effect of LD Threshold on network size

LD (r^2) Threshold	SNP-SNP Pairs From HapMap	Disease Pairs	High / Low LD
≥ 0.5	9416	134	0.982/0.505
≥ 0.3	16879	397	0.982/0.303
≥ 0.1	41485	518	0.982/0.101

Partial Correlation Threshold

Next we changed the partial correlation threshold values determining which links were pruned between the SNPs when the triangulations existed (Figure 4-7). We expected the network to become sparse when the partial correlation threshold was set high (≥ 0.5) and to become dense when it was set low (≥ 0.2). Figure 4-8 shows a sparsely connected DAG computed by IsBIG when 1st order partial correlation threshold is set at $r^2 = 0.8$. Figure 4-9 shows a computed DAG when 1st order partial correlation threshold is set at $r^2 = 0.2$ and when the SNP nodes are absorbed.

Discussion

We outlined a methodology that we call “In-silico Bayesian Integration” and applied it to the domain of GWAS to build a model – In-silico Bayesian Integration of GWAS or IsBIG. The IsBIG model is able to find relationships qualitatively and quantitatively between various diseases or traits as inferred by the linked genetic determinants of the disease or traits in the GWAS catalog. We made a few assumptions: 1) Where the population was not defined in the catalog, we assumed European descent. 2) The strength of association between the SNP and the disease or trait (given by odds ratio) is assumed for a single copy of the allele; the GWAS catalog does not detail about the haplotype effect. 3) We focused on GWAS with discrete outcomes for diseases or traits (i.e. absent / present). A majority of studies in the GWAS catalog list discrete outcomes as opposed to the studies that describe change in a continuous trait like diastolic blood pressure measured as mm Hg increase / decrease and where the strength of the association is reported as a beta coefficient. 4) Lastly, we assumed independent causal

independence among SNPs affecting a disease or trait. This allowed us to combine the individual effect using a Noisy-OR gate as described before.

Additionally, we made some empirical observations about the correlation thresholds. These thresholds were chosen to keep the computational model tractable – values below these thresholds (i.e. LD threshold value below 0.3 and partial correlation threshold below 0.2) resulted in many more SNP nodes which had weaker associations. However when they were included, the algorithm generated out of memory exceptions when trying to absorb many more nodes due to the physical memory addressing limitation of the Java Virtual Machine (2GB on a 32 bit operating system.) Exploring this methodology on systems with greater memory addressing capacity was beyond the scope of this exploratory research.

In Chapter 5 we evaluate our model quantitatively using data from our EMR, the RMRS to quantify the amount of variation among disease linkages that can be explained by studies currently cataloged in the domain of GWAS and to report any novel associations the model finds.

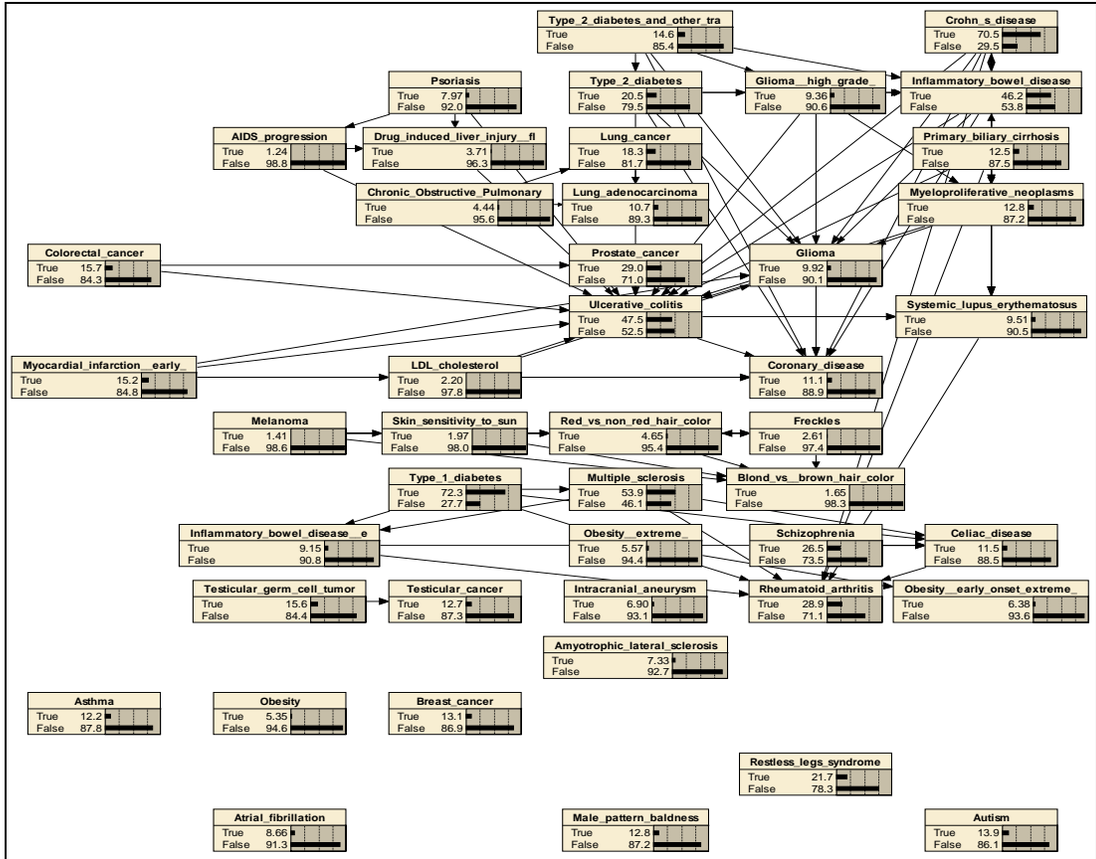


Figure 4-8 IsBIG DAG (SNP-SNP $r^2 = 0.3$, 1st order partial $r^2 = 0.8$)

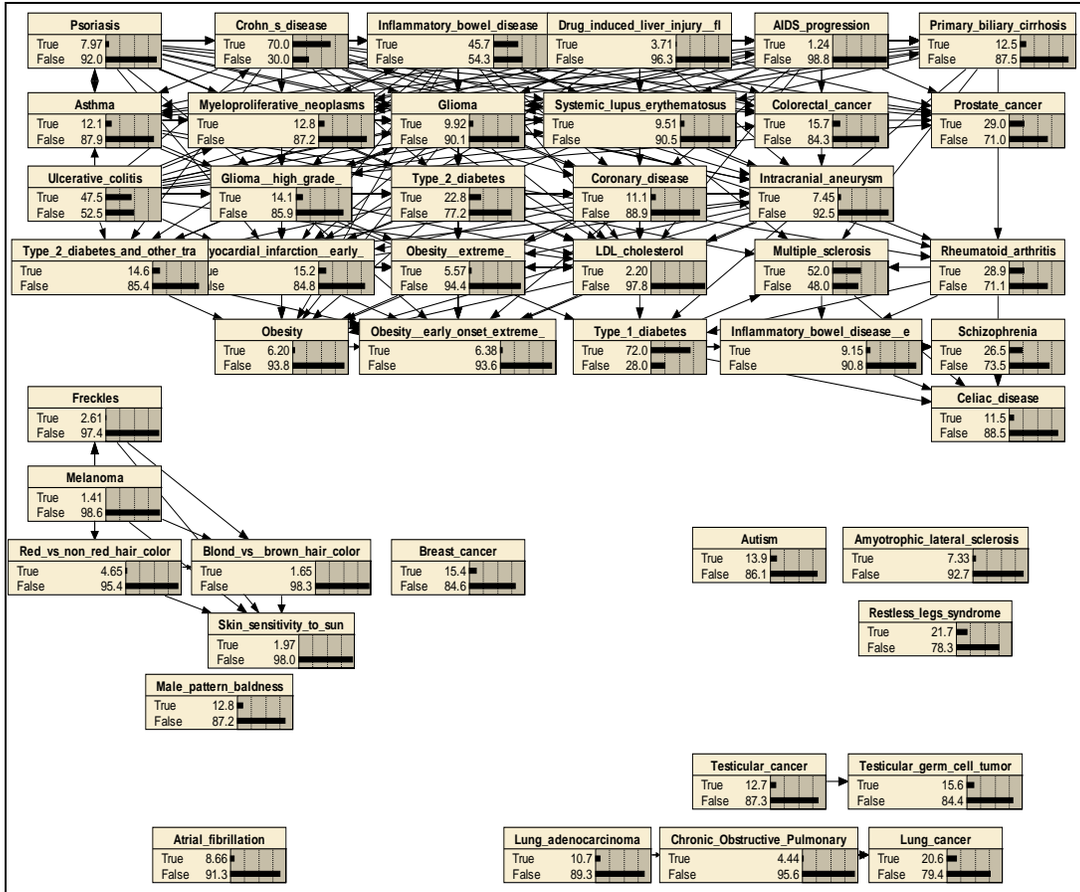


Figure 4-9 IsBIG DAG (SNP-SNP $r^2 = 0.3$, 1st order partial $r^2 = 0.2$)

Table 4-13 Effect of Partial LD Threshold on network connectivity

Partial correlation LD(r^2) Threshold	# of triangulations	# of cycles removed	# of edges removed	# of SNPs with > 1 parent
≥ 0.1	28	7	17	18
≥ 0.2	28	6	30	13
≥ 0.35	28	6	41	9

Figure 4-10 In-silico Bayesian Integration of GWAS Algorithm

Find unique SNP-Proxy Pairs order by descending r^2 value in set M.

Associate each SNP to its list of disease/traits and Proxy SNP to its list of disease trait in a hash map data structure D

DrawDAG(): For each SNP in M

Make SNP a parent node $p(i)$ and Proxy a child node $c(i)$, if the nodes do not already exist in DAG. Draw a link from $p(i)$ to $c(i)$

Triangulate(): For each SNP-Proxy pair in M

If SNP-Proxy have common child $c(i)$ - mark it as a triangulation in set T

RemoveCycles(): For each SNP-Proxy pair in M

If proxy SNP has a child node $c(i)$ that has the gwas SNP as a parent node $p(i)$, reverse the link between $c(i)$ and $p(i)$

Prune(): For each traingulation in T

Compute 1st order partial correlation of each pair. If 1st order partial correlation $>$ defined threshold, keep the link otherwise mark it for removal in set R. Remove all links in set R, this results in SNP map DAG

DrawDAG_DiseaseTraits(): For each SNP in M

Make SNP a parent node $p(i)$ and disease/trait a child node $c(i)$, if the nodes do not already exist in DAG. Draw a link from $p(i)$ to $c(i)$

Compute_CPD(): For each SNP-Proxy pair in M

Use known MAF of each SNP – Proxy duo from GWAS and LD (r^2) value from HapMap data to compute $P(S2/S1)$ as described using equation (1) and (2)

ComputeCPD_Disease (): For each SNP-Proxy pair in M and each SNP-Disease pair in D

Use equation (3)-(5) to compute $P(D/S_i)$. In cases where $P(D/S1,S2)$, use Noisy-OR calculations using equation (6)

CompileNet(): Netica builds a “junction tree” for fast belief updating.

AbsorbSNP(): Absorb out all the SNP nodes in the compiled DAG

WriteDAG(): Serialize the compiled and absorbed DAG

Chapter 5 VALIDATION AGAINST PRIMARY EMR DATA

Introduction

In this chapter we present our evaluation of the unified BN constructed using the In-silico Bayesian Integration methodology described in the previous chapter. In particular we evaluate the In-silico Bayesian Integration of GWAS, hence forth called the IsBIG model (or I-Model) with data from our EMR, the Regenstrief Medical Record System (RMRS). [2]

We evaluate the IsBIG model with the following hypotheses –

1) IsBIG can discover disease-disease associations that are valid. These associations can be confirmed and quantified meaningfully by testing against a) data from our EMR and b) by evaluating against what has been published in the literature.

2) IsBIG can discover novel disease-disease associations. These associations do not exist in the literature.

3) Genetic data can only explain a small fraction of the risk of the disease. IsBIG model can quantify the proportion of risk of disease that can be explained by linked genetic determinants.

Below we describe our methods to test the above three hypotheses. We believe this study is novel as it quantifies the degree of associations found among diseases determined by genetic linkages alone and compares it to a real world EMR.

Methods

For evaluation of disease-disease associations (hypothesis 1), we construct a mixed model (M-Model) where the structure of the IsBIG DAG remains the same but its conditional probability distributions (CPDs) are derived from our EMR, the RMRS. We

test the I-Model against this M-Model and PubMed searches. For evaluation of novel disease-disease associations (hypothesis 2), we evaluate I-Model against published literature in the PubMed database. To evaluate the proportion of the risk attributed to linked genetic determinants (hypothesis 3), we evaluate I-Model against a purely clinical model (C-Model) derived from RMRS. Below we describe each of these models but first we describe the dataset derived from RMRS that is used for this evaluation.

EMR Data for evaluation

With IRB approval we obtained de-identified data from the Regenstrief EMR (RMRS) for 169,711 individuals for 89 diseases. These diseases or proxies of these diseases were also listed in our *Model Catalog* described in the previous chapter. Extraction of these data was sought before the disease-disease relationships for I-Model were known. The data obtained for each patient were extracted from the last 15 years of the individual's medical record, and for each disease each individual was coded as a case or control using the ICD-9 diagnostic codes. The ICD-9 diagnostic codes were selected to match the diseases in the GWAS catalog by a Regenstrief data core expert physician. Thus, in this dataset the same individual can be a case for one or more diseases and a control for others. The dataset had no missing values and henceforth is referred to as the "Regenstrief dataset". We randomly split the Regenstrief dataset into 2/3 training set (112,829 records) and 1/3 test set (56,882 records) for our evaluation. Next, we describe construction of each of the models, i.e. *learning the parameters* of the M-Model and *deriving the DAG and parameters* of the C-Model from the Regenstrief training set.

I-Model Construction

The IsBIG model (I-model) was built using software developed by the author, using Java and Netica software [67] APIs. The software implements the IsBIG algorithm as described in Chapter 5 and produces a network file (.dne) suitable for representation and inference by Netica software. The Java code for the algorithm is included in Appendix A.9. The input for computing the model consisted of the following network parameters – LD threshold value of $r^2 \geq 0.3$ for SNP to SNP correlation and partial correlation threshold values = 0.2. These network parameters were chosen empirically to keep the computational model tractable and yet include weaker correlations of SNPs to test with a real world sub population using RMRS data. The pre-processed input (from the GWAS catalog and HapMap data) for the IsBIG algorithm is detailed in Appendix A.6, and its pre processing (association mining) is described in Chapter 5. The prevalence data for the modeled diseases was derived from our EMR. (Appendix A.7)

M-Model Construction

The mixed model retained the network structure that was derived from the IsBIG algorithm, i.e. the DAG structure of the I-Model. Using Netica's "case file" interface, the 2/3 training set was used to learn the conditional probability distributions (CPD) of each node (disease) in the I-Model DAG. Netica resets any CPDs to uninformed priors (50-50) for each of the nodes before a case file is incorporated. Thus, incorporating the training set transformed the IsBIG model (I-Model) to the mixed model (M-Model) with network structure derived from the GWAS and SNP data and conditional probability distributions learnt from the clinical data from RMRS. Similar to the I-Model, the M-Model was also evaluated for its discriminative power using the same test dataset from RMRS.

C-Model Construction

Using the Winmine toolkit [68] and the 2/3 split training set (same as above), we mined the structure of a DAG that represents the clinical model or the C-Model (Figure 5-1). In this DAG, 42 of the 89 diseases or traits from the initial dataset were connected by an edge with another disease or trait. The rest of the diseases were either disconnected or, because in the GWAS catalog they listed as a continuous trait, they could not be evaluated by the discrete Winmine algorithm.

Winmine uses a greedy algorithm to mine the best structure for the DAG, given the data. This mined DAG was then implemented using Netica [67] Bayesian Network software. The conditional probability distributions for this DAG were learnt from the same 2/3 split training set from which the structure was mined. Thus this model represented the DAG structure and parameters; both learnt from clinical data and became our clinical model (C-Model). Similar to the other two models, the C-Model was evaluated for its discriminative or predictive power using the same test dataset from RMRS.

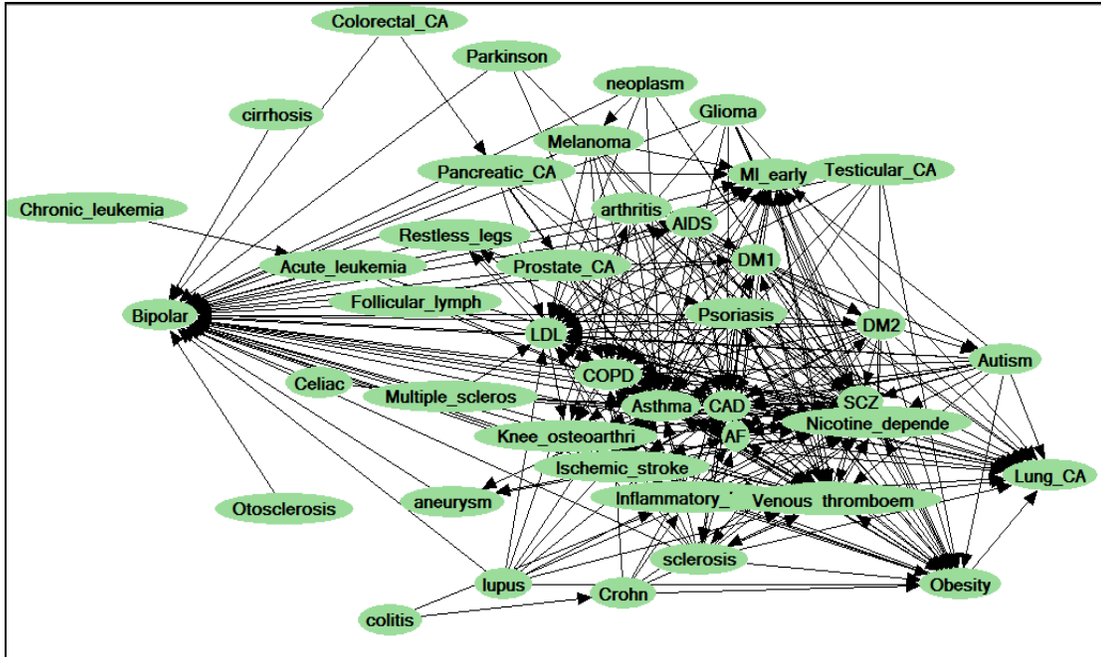


Figure 5-1 DAG Structure of C-Model learnt from RMRS Training set

Performance measure

We use Area Under the Curve (AUC) of Receiver Operating Characteristic (ROC) curves [57] as a performance measure for evaluating the discriminative power of the three BNs. The ROC curve performance measure is described before in Chapter 2. A ROC with AUC of 0.5 score has no predictive value and is as good as chance.

Computing Area under the Curve (AUC)

We used Netica’s “test interface” and the randomly split 1/3 test set from RMRS to evaluate the performance for discrimination of cases and controls for each disease by calculating the AUCs for each of the 42 diseases or the subset of them in each of the DAGs.

Testing for statistical significance

We did a test of hypothesis for each model constructed above as follows. For each disease node, we tested whether the BN had statistically significant predictive power (i.e.,

AUC significantly greater than 0.5) using the Hanley and McNeil method described in [96] for calculating the standard error from which z-scores can be calculated without distributional assumptions (i.e. based on the count of normal and abnormal cases in the test set). The results of these tests are listed in Appendix A.2 to A.4 and described below.

Additionally, to compare the predictive power of the M-Model and the C-Model, we used the Hanley McNeil method for ROC derived from *same cases* [72] to compare ROCs for the same disease. Please note that both these models differ in structure, but the conditional probability distributions were learnt from the same training set and they were tested using the same test set.

Evaluation of IsBIG Compared to Published Literature

To evaluate the I-Model, we evaluated direct pair wise associations (Appendix A.4) between diseases that the IsBIG algorithm found against – 1) PubMed database searches, 2) direct pair wise associations between diseases in C-Model, i.e. DAG derived from EMR and 3) against both of the above combined. We searched the PubMed database for articles linking the diseases found to be associated in the IsBIG DAG by searching for one disease as a keyword with a boolean AND condition to the second condition, for example – Primary Biliary Cirrhosis AND Crohn’s Disease. These searches were conducted in May 2010. The number of articles meeting these simple criteria was counted for our measure.

Results

I-Model Evaluation with Test set

29 of the 42 diseases (Appendix A.2) from Regenstrief EMR were included in the I-Model construction by the IsBIG software with the chosen network parameters

described above. With the 1/3 randomly split test set, the I-Model predicted 5 (17%) of the 29 diseases or traits with an area under the curve ($AUC > 0.5$, $p < 0.05$). In other words the network had the discriminative power to differentiate cases from controls from our EMR test set for 17% of diseases. The diseases that were predicted with statistical significance in I-Model are listed in Table 5-1 below and highlighted in Appendix A.2.

Table 5-1 Discriminative power in IsBIG (I-Model)

	Node	AUC	p-value
1	Coronary Disease	0.6856	0
2	Lung cancer	0.6263	0
3	LDL Cholesterol (Elevated)	0.5823	0
4	Type 2 Diabetes	0.5431	0
5	Obesity	0.5192	4.73E-09

M-Model Evaluation with Test set

As described before, the I-Model was parameterized for its 29 nodes using the 2/3 randomly split training set. This parameterization transformed the I-Model into the M-Model. Using the same 1/3 test set (as before), the M-Model predicted 12 (41%) of the 29 diseases or traits with an area under the curve ($AUC > 0.5$, $p < 0.05$). The diseases that were predicted with statistical significance in M-Model are listed in Table 5-2 below and highlighted in Appendix A.3.

Table 5-2 Predictable diseases in Parameterized IsBIG (M-Model)

	Node	AUC	p-value
1	Myocardia I Infarction Early	0.937	0
2	Coronary Disease	0.7318	0
3	Psoriasis	0.6343	0
4	Lung cancer	0.6263	0
5	Systemic Lupus Erythematosus	0.5947	2.08E-06
6	LDL Cholesterol Elevated	0.5945	0
7	Rheumatoid Arthritis	0.5871	0
8	Chronic Obstructive Pulmonary Disorder	0.533	1.2E-10

9	Asthma	0.5274	0
10	AIDS Progression	0.522	0.029809
11	Obesity	0.5192	4.73E-09
12	Type 1 Diabetes	0.5098	0.038912

C-Model Evaluation with Test set

We used the same test set (randomly 1/3 split from RMRS data) to evaluate the performance of the clinical model (C-Model). Please note that the structure of the C-Model DAG was independently derived from the 2/3 training set. With the same test set, the C-Model predicted 31 (74%) of 42 disease or trait nodes with an area under the curve (AUC > 0.5, $p < 0.05$). The diseases that were predicted with statistical significance in this model are listed and highlighted in Appendix A.4.

Comparing the I-Model to the M-Model

Since the only difference between the I-Model and the M-Model is the conditional probability distributions of each node; (the I-Model contains the genetic linkage representation where as the M-Model contains both the genetic and clinical representation, therefore we compared the statistical significance of the I-Model with the M-Model. This enabled us to evaluate our hypothesis that the associations discovered by IsBIG model can be validated in our EMR. Table 5-3 below gives the counts of the number of diseases that were statistically significant in each model.

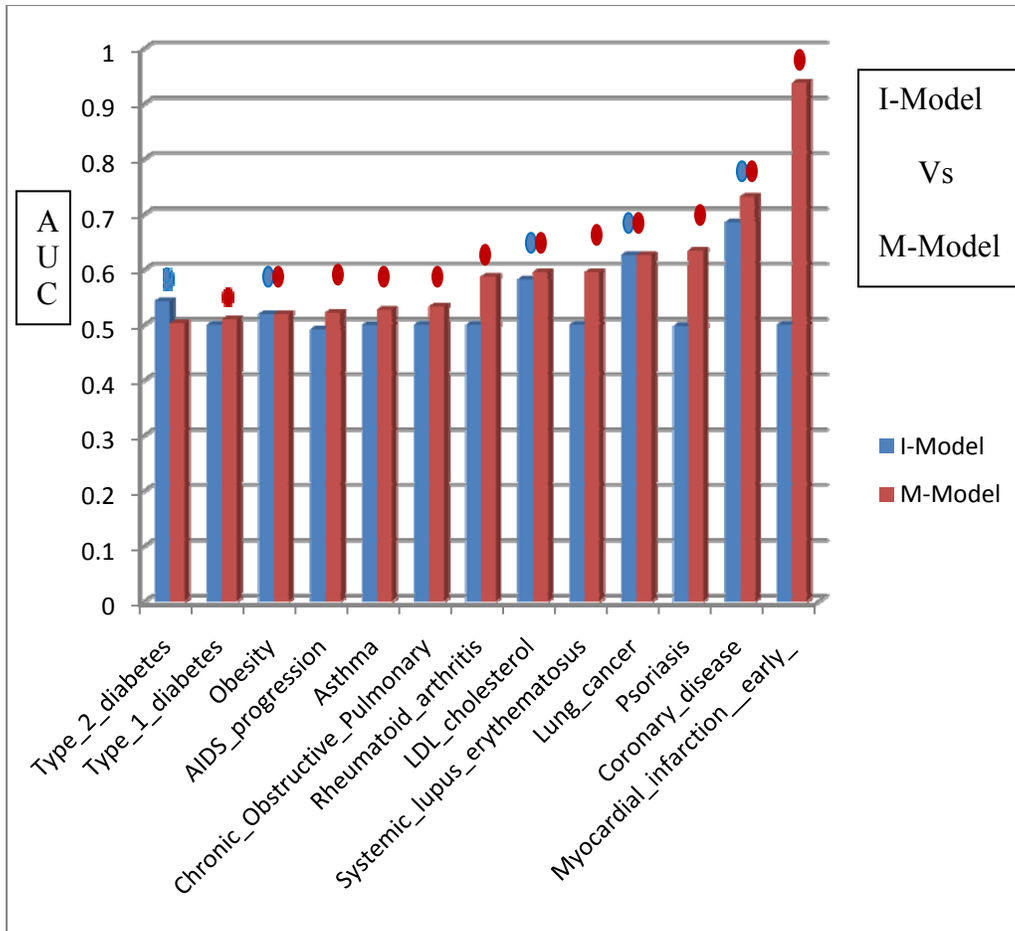


Figure 5-2 IsBIG Model performance, statistical significance denoted by ● ●

Table 5-3 Number of statistically significant diseases predicted by each model

		+	M-Model	-
I-Model	+	4	1	
	-	8	16	

Predicted diseases denoted by +

Of the 29 diseases, 5 were predictable by the I-Model and 12 were predictable by the M-Model (AUC > 0.5, p < 0.05). There were 4 diseases common to both the I-Model and the M-Model, i.e. the I-Model is 33% sensitive when compared to M-Model. There

were 17 diseases not predictable by the M-Model, of those, 16 were common to both the models i.e. I-Model is 94% specific when compared to the M-Model. The details of the disease nodes marked with a colored dot for statistical significance are in Figure 5-2 above.

Of the 4 diseases predictable by the I-Model and the M-Model, the networked performed the same for *Obesity and Lung Cancer* i.e. AUCs did not change on computation of conditional probability distributions (parameterization) of the nodes. For the other diseases, there was a modest gain (1 to 14%) in the discriminative power of the IsBIG DAG on parameterization with the EMR data i.e. in the M-Model (Figure 5-3).

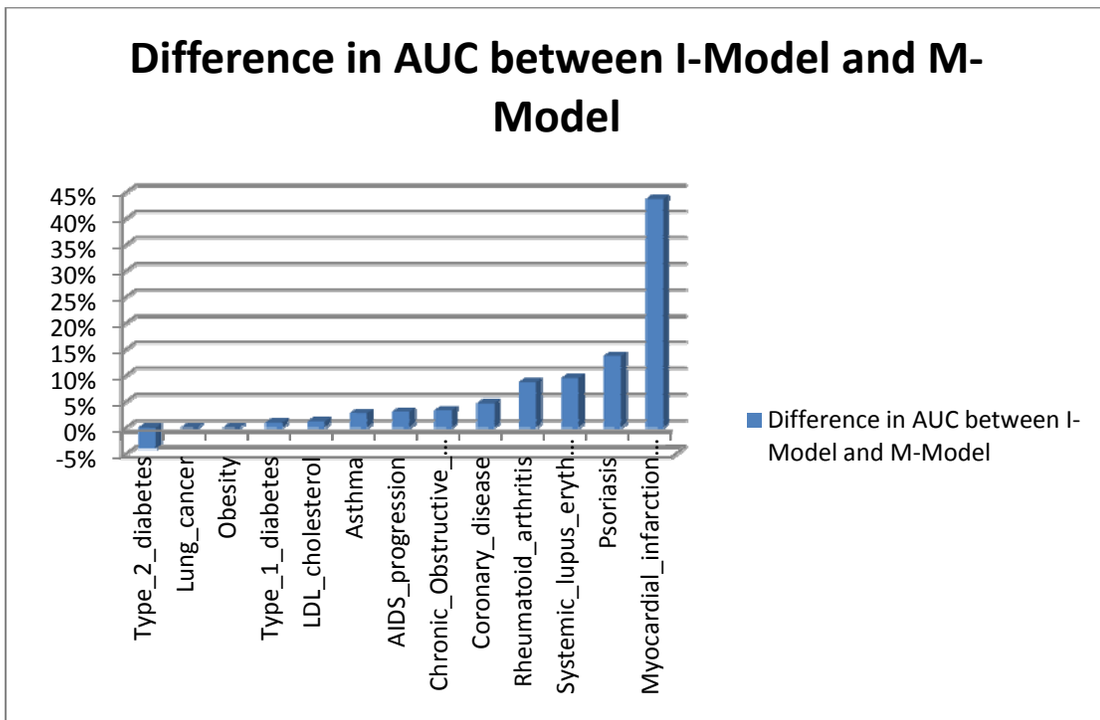


Figure 5-3 Change in IsBIG AUC on parameterization

There were a few exceptions though, most notably for early detection of *Myocardial infarction*. The M-Model’s discriminative power (AUC) increased by 44%, i.e. a relative jump of 87% when compared to the I-Model. However, surprisingly on the

other hand, the M-Model's discriminative power decreased for *Type 2 diabetes* by 4% on parameterization with the clinical data.

Compare the IsBIG Models (I-Model and M-Model) to the C-Model

As expected, overall the C-Model derived from clinical data from our EMR outperformed the IsBIG models (Figure 5-4). The C-Model was able to discriminate 31 diseases when compared to 5 and 12 diseases in the I-Model and the M-Model respectively. Furthermore, the AUCs of the C-Model were much bigger ($p < 0.05$) for all but one of the statistically significant diseases common in both the IsBIG models. The C-Model had the same AUC for Systemic Lupus Erythematosus (or SLE) as in the M-Model ($p = 0.4095$). Interestingly, the difference in AUCs for the early Myocardial infarction node between the M-Model and the C-Model was very small (~ 0.02) but it was statistically significant ($p = 5.3 \times 10^{-6}$), i.e. the discriminative powers of the two DAGs were very different.

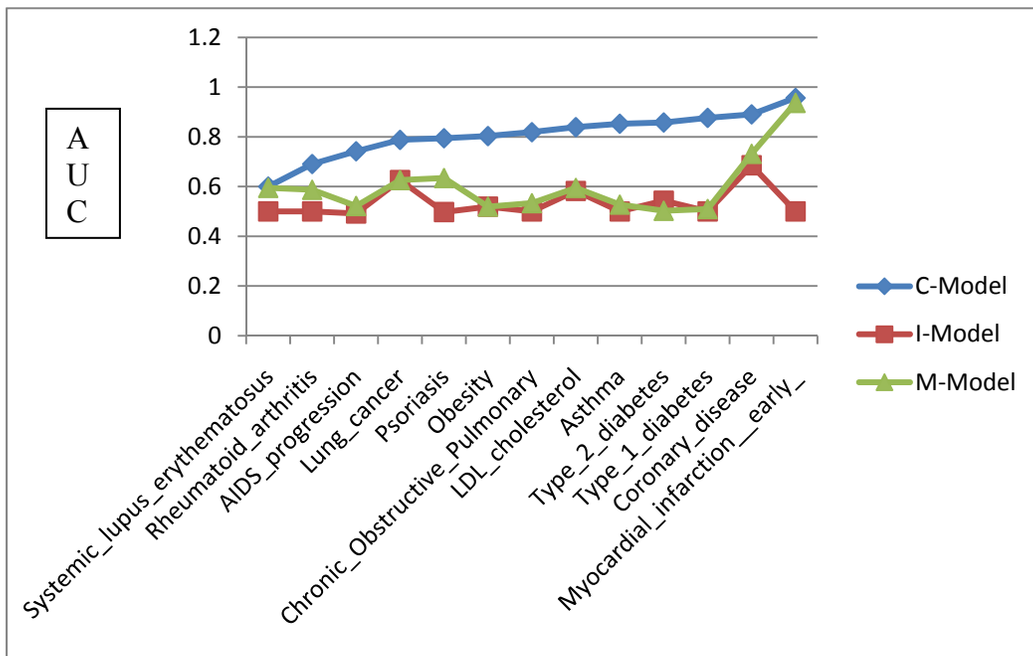


Figure 5-4 AUC comparison of models

Compare the I-Model against PubMed Literature searches

We evaluated 117 direct pair wise (parent-child) associations (Appendix A.4) in the I-Model against the PubMed database literature searches. Twenty (17%) of the pair wise associations in the I-Model had *no references* in the literature and can be possibly considered novel. At least 1 of these associations (Rheumatoid Arthritis with Systemic Lupus Erythematosus) was also found in direct pair wise associations in the C-Model (Table 5-4).

Table 5-4 Novel Associations in IsBIG Model (I-Model)

	Disease Node 1	Disease Node 2
***1	Rheumatoid arthritis	Systemic lupus erythematosus
2	AIDS progression	Crohn's disease
3	Glioma	Crohn's disease
4	Glioma	Primary biliary cirrhosis
5	Glioma	Myeloproliferative neoplasms
6	Intracranial aneurysm	Crohn's disease
7	Intracranial aneurysm	Inflammatory bowel disease
8	LDL cholesterol (Elevated)	Crohn's disease
9	Myeloproliferative neoplasms	Colorectal cancer
10	Myeloproliferative neoplasms	Crohn's disease
11	Myeloproliferative neoplasms	AIDS progression
12	Myeloproliferative neoplasms	Primary biliary cirrhosis
13	Myeloproliferative neoplasms	Asthma
14	Myocardial infarction early	Crohn's disease
15	Primary biliary cirrhosis	Crohn's disease
16	Primary biliary cirrhosis	Colorectal cancer
17	Primary biliary cirrhosis	Prostate cancer
18	Systemic lupus erythematosus	Crohn's disease
19	Systemic lupus erythematosus	Primary biliary cirrhosis
20	Systemic lupus erythematosus	Myeloproliferative neoplasms

*** Also found in clinical model (C-Model)

Compare I-Model pair wise associations to C-Model pair wise associations

Of the 117 direct pair wise disease associations in the I-Model, 27 (23%) associations were also found in direct pair wise disease associations in the C-Model Table 5-5 below lists these associations.

**Table 5-5 Associations common in IsBIG Model with C-Model
(with literature reference count)**

	Disease Node 1	Disease Node 2	PubMed Ref count
1	Rheumatoid arthritis	Systemic lupus erythematosus	0
2	Coronary disease	Crohn's disease	1
3	LDL cholesterol (Elevated)	AIDS progression	4
4	Coronary disease	Glioma	5
5	Asthma	AIDS progression	6
6	Myocardial infarction early	AIDS progression	9
7	LDL cholesterol (Elevated)	Glioma	11
8	LDL cholesterol (Elevated)	Inflammatory bowel disease	12
9	Glioma	Asthma	24
10	Crohn's disease	Ulcerative colitis	32
11	Inflammatory bowel disease	Crohn's disease	33
12	LDL cholesterol (Elevated)	Psoriasis	33
13	LDL cholesterol (Elevated)	Myocardial infarction early	35
14	Myocardial infarction early	Inflammatory bowel disease	38
15	Coronary disease	AIDS progression	50
16	Chronic Obstructive Pulmonary	Lung cancer	51
17	Coronary disease	Inflammatory bowel disease	51
18	Coronary disease	Psoriasis	66
19	LDL cholesterol (Elevated)	Coronary disease	160
20	Multiple sclerosis	Asthma	233
21	Asthma	Psoriasis	261
22	Inflammatory bowel disease	Ulcerative colitis	262
23	Asthma	Inflammatory bowel disease	271
24	Obesity	Type 2 diabetes	320
25	Rheumatoid arthritis	Asthma	894
26	Schizophrenia	Type 1 diabetes	931
27	Myocardial infarction early	Coronary disease	54128

Literature Ref count of pair wise associations in I-Model (also found in C-Model)

Of the 27 pair wise disease associations in the I-Model that were also found in the C-Model, 1 (4%) association had no literature reference, 5 (19%) had up to 10 references. These disease associations can be possibly considered as worth exploring further in future

studies. Additionally, 9 (33%) associations had only up to 50 references. The details are listed in Figure 5-4 below and Table 5-5 above.

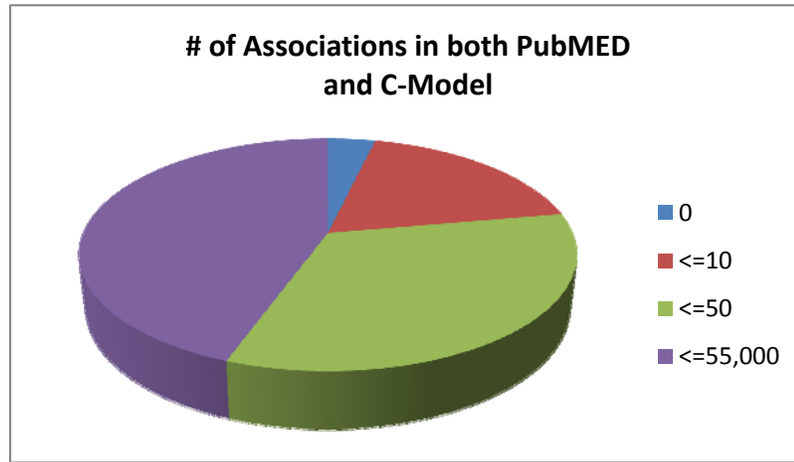


Figure 5-5 Reference count of IsBIG associations also found in C-Model

Discussion

We have described our IsBIG methodology and applied it to combine statistical correlations between SNPs and GWAS to generate a disease map of common diseases, the IsBIG model as inferred from genetic underpinnings. We evaluated the model against both raw clinical data and published literature.

Our results show that IsBIG discovered disease-disease associations are valid. We tested this by training the IsBIG DAG with RMRS data and found that IsBIG correctly found 33% of the diseases in the M-Model; and of the associations found by IsBIG, 80% of them were confirmed in the M-Model. This essentially means that for the 4 diseases or traits (*Coronary Disease, Elevated LDL Cholesterol, Lung Cancer and Obesity*) predictable by IsBIG, there is discriminative power in the model as inferred by genetic variations alone in the population. We also found that the IsBIG model has the same discriminative power for early detection of Myocardial Infarction as the same model trained on clinical data (i.e. M-Model).

IsBIG can discover novel disease-disease associations that have not been described. We compared disease-disease associations to the PubMed database and found that many had never been described. We believe these associations can be considered as hypotheses for future studies. Furthermore, we were able to confirm some of them in the clinical model (C-Model). At least one pair wise disease association (*Rheumatoid arthritis and Systemic lupus Erythematosus*) found by IsBIG had no reference in the literature and can be considered novel, and further, it can be confirmed by the C-Model derived from our EMR data.

Finally, our results show that probabilistic methods can extract data from disparate sources and combine them in a normative way such that it correlates with what we may empirically find in an EMR. Only a fraction of associations among diseases found in an EMR can be explained by genetics of SNP to disease linkage. We compared the AUCs of IsBIG and the C-Model and found that the C-Model AUCs were much bigger. Therefore, genetic data alone can only explain a small fraction of the risk of a disease.

Chapter 6 **DISCUSSION**

Summary of findings

In this work, we formed a methodology which we call In-silico Bayesian Integration, to integrate data from disparate sources using probabilistic modeling methods, specifically the Bayesian Network framework. We demonstrated that this methodology can combine information in a normative way and that the results can be validated against real world datasets from our EMR. Additionally, we demonstrated that the model built using this methodology discovers new knowledge that can be used as hypotheses for future studies.

As an example of our approach, we combined two disparate sources – the genetic linkages (associations) found between Single Nucleotide Polymorphisms (SNP) in the human genome and the effect size linking a SNP with a disease or a trait found in Genome Wide Association Studies (GWAS). We call this model In-silico Bayesian Integration of GWAS (or IsBIG). IsBIG produces a map of disease-disease associations as inferred from their genetic underpinnings.

Seventeen percent of disease-disease associations found by the IsBIG model are not described in the literature and can be considered novel, with at least one association confirmed in our EMR. Furthermore, our results show that IsBIG finds meaningful associations that can be empirically validated against our EMR. The IsBIG model is 33% sensitive, 94% specific and has a positive predictive value of 80% when we evaluate its discriminative power compared to our EMR data. We found from this evaluation that GWA studies in the domain of *Obesity, Type 2 Diabetes, elevated LDL Cholesterol trait, Lung cancer and Coronary Disease* have signal to detect effect of genetic variations to

disease likelihood in our clinical population and may have potential clinical applications. By generating a predictive distribution of disease-disease associations, the IsBIG model has also quantified that the simple linking of genetics to a disease only explains a fraction of the risk of disease in our EMR.

Summary of contributions

Applying Bayesian Networks to the real world task of integrating data from disparate sources required the use of causal independence assumptions (Independence of Causal Influence models, ICI) particularly the Noisy-OR model. Our empiric evaluations of the BN framework with ICI model using large datasets from our EMR produced comparable results to that of an expert. Therefore, in this research we have demonstrated that ICI models provide a successful strategy for computing conditional distributions of a large BN with many linked nodes in a tractable way. We believe the use of ICI models, particularly the use of Noisy-OR model seems to be robust under a broad range of conditions. To the best of our knowledge, this has not been demonstrated before, especially in the biomedical field.

Additionally, we have also shown that the use of Noisy-OR model provides a successful strategy for combining disparate sources, where individual conditional distributions can be computed from effect sizes and statistical correlations. We have demonstrated it by utilizing odds ratios and correlation and partial correlation coefficients as strategies to translate genetic data from published GWAS into conditional probability distributions for disease associations. This has not been demonstrated before to the best of our knowledge.

There is no analytic method, to the best of our knowledge, which computationally combines disparate source of both raw data and summary measures in a quantitative framework. Applications of text mining using ontology [97] and more recently applications of semantic web technology coupled with graph algorithms [98] have been used to find associations between biological entities from different sources of information. However, they rely on purely qualitative associations; i.e. subject-predicate-object relationships, for example, drug A may cause symptom B sometimes is represented by a triplet such as <A causes B>. However this representation does not account for the uncertainty (“may”) in the assertion. The knowledge that the relationship between A and B exists say 2% of the times is not represented in this framework. Such frameworks lay out a graph of the associations but do not produce predictive distributions of these associations. On the other hand, quantitative methods such as meta-analysis of studies are able to give more precise estimates of the effect size by pooling effect sizes from a number of studies under assumptions, but they are not a computational model that can be used for future inference. They also do not permit for additional information to be incorporated as it becomes available. Other quantitative methods, in particular data mining methods which are computational and can generate predictive distributions, require a large numbers of cases for model building, training and testing. These datasets seldom exist in the biomedical domain for use in hypotheses generation tools.

Therefore, due to the lack of these original datasets, in this work, we have developed a quantitative method using Bayesian Networks framework to combine both raw data and summary measures in a computational model. The method is able to discover new associations from disparate sources and is able to produce predictive

distributions for these associations. It is also able to incorporate new information as it becomes available. Our evaluation of this method shows correlation to large real world datasets.

On Use of Causal Independence Assumption (Noisy-OR)

The Bayesian network framework does not constrain how a variable depends upon its parents. One interpretation is that the directed edges or arcs represent causal relationships among the parent and the child. Thus the local structure encodes the dependencies and probability distributions of parent and child (conditioned upon parents). The probability distribution of the child node can be approximated assuming causal independence among the parents using a boolean function such as OR, but since the parent-child relationship is probabilistic, the relationship is “noisy”.

The underlying assumption in this situation is that each parent acts independently as a cause to the child (effect) with an independent mechanism of action, and each parent can sufficiently cause the effect. In the absence of any parent, there is no effect, unless we assume a leaky model.

Without the causal independence assumption, we had no way of combining the SNP’s effect size to a disease. These assumptions allowed us to combine the contribution of each existing cause (SNP) to the effect (disease) with fewer calculations than the calculations for full conditional probability distribution. Additionally they allowed the flexibility of tractably adding new causes in the future. It is these capabilities of Noisy-OR that we exploited in this research to combine information (data or statistical summaries) in a causal (normative) way.

Limitations

One of the limitations of our methodology is the approximation of the underlying data – we modeled only discrete variables. This assumption allowed us to compute conditional probability tables (CPTs) using a boolean function such as OR in the Noisy-OR calculations. For this research, the boolean function “OR” and discrete data fitted our assumption of the real world datasets but may not hold true for other datasets. In future studies, there may be disparate sets that may need combinations of other boolean functions such as AND or XOR, besides OR. Nonetheless, similar methodology can be applied in such situations by replacing the Noisy-OR calculations with the one that fits the datasets. For example, if we make the assumption that both genetics and environment lead to a disease status, we may want to model it using a Noisy AND. For modeling continuous variables, methods similar to Monte Carlo simulations [99] assuming a-priori probability distributions can be applied.

Another assumption in this study is that effect sizes or correlations are defined for one variable to the outcome variable. When using effect sizes (such as odds ratio) to link two different BNs to calculate CPTs, we are assuming the effect is attributable to a single variable. For example, for the IsBIG study, it is the odds ratio or strength of association of single allele to the disease or trait in the study. There may be situations, where this assumption does not hold true. The effect size could result from more than one variable. For example, two copies of a specific allele may produce more than an additive effect as is the case in recessive traits. In those situations, assuming causal independence between the two causal variables (for e.g. between alleles) and the outcome variable will not be justified.

Lastly, though the IsBIG study illustrated how disparate datasets, summary or statistical measures from different sources can be combined. However, the information in both the sources belonged to a similar population (i.e. European Ancestry). We propose that as a variable, such as the LD measure, varies by a sub-population, this should be accounted for in the model when disparate sources are combined.

Future Directions

We believe our methodology has been successfully applied in integrating sub domains, especially in the biomedical domain and has many practical applications. We envision that using this method secondary data sources such as summary and statistical data from published sources can be merged with data from primary sources such as electronic health records to provide a more normative and quantitative evaluation of the domain.

We are envisioning one such application of this method for dynamic prioritization of the reminder prompts in our CHICA system. [59] CHICA's static, global prioritization scheme limits the flexibility of the system by evaluating the predetermined prompts for primary care alerts from a set of guidelines for a specific age group. At present there are many such guidelines and recommendations. [100-104] We plan to use Bayesian networks (BN) as a strategy for modeling a patient's clinical status with the idea of calculating the expected value of making alternative recommendations to physicians in order to tailor prioritization to the patient, [63] using the research in this thesis.

APPENDICES

Appendix A.1 Summary of Studies from GWAS Catalog

Disease/Trait	Number of Studies
Type 2 diabetes	67
Type 1 diabetes	62
Crohn's disease	61
Bipolar disorder	43
Multiple sclerosis	41
Prostate cancer	37
Amyotrophic lateral sclerosis	28
Rheumatoid arthritis	24
Schizophrenia	24
Ulcerative colitis	19
Acute lymphoblastic leukemia (childhood)	17
Breast cancer	16
Lung cancer	16
Colorectal cancer	15
Parkinson's disease	14
Smoking behavior	14
Obesity (extreme)	13
Psoriasis	12
Response to treatment for acute lymphoblastic leukemia	12
Systemic lupus erythematosus	12
Coronary disease	11
Celiac disease	10
Myocardial infarction (early onset)	9
Nonsyndromic cleft lip with or without cleft palate	9
Inflammatory bowel disease	8
AIDS	7
Alzheimer's disease	7
Glioma	7
Response to citalopram treatment	7
Alcohol dependence	6
Chronic lymphocytic leukemia	6
Drug-induced liver injury (flucloxacillin)	6
Melanoma	6
Restless legs syndrome	6
Testicular germ cell tumor	6
Type 2 diabetes and other traits	6

Atrial fibrillation	5
Blond vs. brown hair color	5
Hypertension	5
Lung adenocarcinoma	5
Obesity	5
Primary biliary cirrhosis	5
Stroke	5
Blue vs. green eyes	4
Coronary artery disease	4
Freckles	4
Inflammatory bowel disease (early onset)	4
Intracranial aneurysm	4
Autism	3
Exercise (leisure time)	3
Glioma (high-grade)	3
Male-pattern baldness	3
Skin pigmentation	3
Skin sensitivity to sun	3
Alzheimer's disease (late onset)	2
Arthritis (juvenile idiopathic)	2
Asthma	2
Atrial fibrillation/atrial flutter	2
Basal cell carcinoma (cutaneous)	2
Chronic Obstructive Pulmonary Disease	2
Kawasaki disease	2
Knee osteoarthritis	2
Left ventricular mass	2
Nicotine dependence	2
Response to Hepatitis C treatment	2
Thyroid cancer	2
Urinary bladder cancer	2
Age-related macular degeneration	1
Age-related macular degeneration (wet)	1
AIDS progression	1
Asthma (childhood onset)	1
Atopic dermatitis	1
Atopy	1
Bladder cancer	1
Blue vs. brown eyes	1
Burning and freckling	1
Crohn's disease and Sarcoidosis (combined)	1

Diabetic nephropathy	1
Disease/Trait	1
End-stage renal disease	1
Essential tremor	1
Follicular lymphoma	1
Gallstones	1
Glaucoma (exfoliation)	1
Height	1
Ischemic stroke	1
Kidney stones	1
LDL cholesterol	1
Major depressive disorder	1
Myeloproliferative neoplasms	1
Myocardial infarction	1
Narcolepsy	1
Neuroblastoma	1
Neuroblastoma (high-risk)	1
Neuroticism	1
Obesity (early onset extreme)	1
Osteonecrosis of the jaw	1
Otosclerosis	1
Ovarian cancer	1
Pancreatic cancer	1
Parkinson's disease (familial)	1
Periodontitis	1
QT interval prolongation	1
Red vs. non-red hair color	1
Red vs. non-red hair color	1
Renal function and chronic kidney disease	1
Response to antipsychotic treatment	1
Response to statin therapy	1
Response to ximelagatran treatment	1
Testicular cancer	1
Venous thromboembolism	1
	807

Appendix A.2 AUC and p-values in IsBIG model (I-Model)

	Node	AUC	p-value
1	Coronary Disease	0.6856	0
2	Lung cancer	0.6263	0
3	LDL	0.5823	0
4	Type 2 Diabetes	0.5431	0
5	Obesity	0.5192	4.73E-09
6	AIDS Progression	0.4917	0.234547
7	Psoriasis	0.4976	0.351112
8	Asthma	0.4998	0.470214
9	MI_Early	0.5	0.5
10	SLE	0.5	0.5
11	Type 1 Diabetes	0.5	0.5
12	Prostate Cancer	0.5	0.5
13	RA	0.5	0.5
14	COPD	0.5	0.5
15	Ulcerative Colitis	0.5	0.5
16	Testicular cancer	0.5	0.5
17	Schizophrenia	0.5	0.5
19	Primary Biliary Cirrhosis	0.5	0.5
19	Myeloproliferative Neoplasms	0.5	0.5
20	Melanoma	0.5	0.5
21	Intracranial Aneurysm	0.5	0.5
22	Inflammatory Bowel Disease	0.5	0.5
23	Crohn's Disease	0.5	0.5
24	Colorectal cancer	0.5	0.5
25	Celiac Disease	0.5	0.5
26	Autism	0.5	0.5
27	Atrial Fibrillation	0.5	0.5
28	Multiple Sclerosis	0.5	0.5
29	Glioma	0.4703	0.13301

Appendix A.3 AUC and p-values in Mixed model (M-Model)

	Node	AUC	p-value
1	Myocardial Infarction Early	0.937	0
2	Coronary Disease	0.7318	0
3	Psoriasis	0.6343	0
4	Lung cancer	0.6263	0
5	Systemic Lupus Erythematosus	0.5947	2.08E-06
6	LDL Cholesterol Elevated	0.5945	0
7	Rheumatoid Arthritis	0.5871	0
8	Chronic Obstructive Pulmonary Disorder	0.533	1.2E-10
9	Asthma	0.5274	0
10	AIDS Progression	0.522	0.029809
11	Obesity	0.5192	4.73E-09
12	Type 1 Diabetes	0.5098	0.038912
13	Type 2 Diabetes	0.5025	0.30807
14	Prostate Cancer	0.5012	0.476679
15	Ulcerative Colitis	0.5	0.5
16	Testicular cancer	0.5	0.5
17	Schizophrenia	0.5	0.5
18	Primary Biliary Cirrhosis	0.5	0.5
19	Myeloproliferative Neoplasms	0.5	0.5
20	Melanoma	0.5	0.5
21	Intracranial Aneurysm	0.5	0.5
22	Inflammatory Bowel Disease	0.5	0.5
23	Crohn's Disease	0.5	0.5
24	Colorectal cancer	0.5	0.5
25	Celiac Disease	0.5	0.5
26	Autism	0.5	0.5
27	Atrial Fibrillation	0.5	0.5
28	Multiple Sclerosis	0.4987	0.479927
29	Glioma	0.4958	0.438703

Appendix A.4 AUC and p-value in Clinical Model (C-Model)

	Node	AUC	p-value
1	Crohn's Disease	0.9985	0
2	Ulcerative Colitis	0.9969	0
3	Myocardial Infarction Early	0.9568	0
4	Coronary Disease	0.8906	0
5	Type 1 Diabetes	0.8766	0
6	Schizophrenia	0.862	0
7	Type 2 Diabetes	0.8581	0
8	Atrial Fibrillation	0.8577	0
9	Asthma	0.853	0
10	LDL Cholesterol Elevated	0.8392	0
11	Bipolar	0.8268	0
12	Chronic Obstructive Pulmonary Disorder	0.8192	0
13	Obesity	0.8034	0
14	Nicotine Dependence	0.8013	0
15	Psoriasis	0.7947	0
16	Lung cancer	0.788	0
17	Breast cancer	0.7805	0
18	Ischemic Stroke	0.7603	0
19	Autism	0.7501	0
20	AIDS Progression	0.7423	0
21	Venous thromboembolism	0.7239	0
22	Colorectal Cancer	0.7051	0.001966
23	Knee osteoarthritis	0.7003	0
24	Rheumatoid Arthritis	0.6908	0
25	Prostate Cancer	0.6702	0
26	Glioma	0.6546	2.29E-08
27	Restless Leg Syndrome	0.6241	0.000132
28	Testicular cancer	0.6139	0.001118
29	Systemic Lupus Erythematosus	0.6	5.93E-07
30	Intracranial Aneurysm	0.5884	0.006792
31	Pancreatic Cancer	0.5757	0.005877
32	Follicular Lymphoma	0.5526	0.122542
33	Myeloproliferative Neoplasms	0.5433	0.111934
34	Melanoma	0.5268	0.173608
35	Acute Leukemia	0.5	0.5
36	Otosclerosis	0.5	0.5
37	Celiac Disease	0.5	0.5
38	Chronic Leukemia	0.5	0.5

39	IBD	0.5	0.5
40	Multiple Sclerosis	0.5	0.5
41	Parkinson	0.5	0.5
42	Primary Biliary Cirrhosis	0.5	0.5

Appendix A.5 Relationships evaluated in IsBIG Model (I-Model)

	Disease Node 1	Disease Node 2	Lit Ref count	RMRS Eval
1	Rheumatoid arthritis	Systemic lupus erythematosus	0	Yes
2	AIDS progression	Crohn s disease	0	No
3	Glioma	Crohn s disease	0	No
4	Glioma	Primary biliary cirrhosis	0	No
5	Glioma	Myeloproliferative neoplasms	0	No
6	Intracranial aneurysm	Crohn s disease	0	No
7	Intracranial aneurysm	Inflammatory bowel disease	0	No
8	LDL cholesterol (Elevated)	Crohn s disease	0	No
9	Myeloproliferative neoplasms	Colorectal cancer	0	No
10	Myeloproliferative neoplasms	Crohn s disease	0	No
11	Myeloproliferative neoplasms	AIDS progression	0	No
12	Myeloproliferative neoplasms	Primary biliary cirrhosis	0	No
13	Myeloproliferative neoplasms	Asthma	0	No
14	Myocardial infarction early	Crohn s disease	0	No
15	Primary biliary cirrhosis	Crohn s disease	0	No
16	Primary biliary cirrhosis	Colorectal cancer	0	No
17	Primary biliary cirrhosis	Prostat cancer	0	No
18	Systemic lupus erythematosus	Crohn s disease	0	No
19	Systemic lupus erythematosus	Primary biliary cirrhosis	0	No
20	Systemic lupus erythematosus	Myeloproliferative neoplasms	0	No
21	Coronary disease	Crohn's disease	1	Yes
22	Asthma	Crohn's disease	1	No
23	Intracranial aneurysm	Psoriasis	1	No
24	Intracranial aneurysm	AIDS progression	1	No
25	Myeloproliferative neoplasms	Psoriasis	1	No
26	Primary biliary cirrhosis	AIDS progression	1	No
27	Psoriasis	Crohn's disease	1	No
28	Glioma	AIDS progression	2	No
29	Myeloproliferative neoplasms	Prostat cancer	2	No
30	Myeloproliferative neoplasms	Ulcerative colitis	2	No
31	AIDS progression	Ulcerative colitis	3	No
32	Glioma	Psoriasis	3	No
33	Myeloproliferative neoplasms	Inflammatory bowel disease	3	No
34	LDL cholesterol (Elevated)	AIDS progression	4	Yes
35	Glioma	Ulcerative colitis	4	No
36	Glioma	Inflammatory bowel disease	4	No
37	Intracranial aneurysm	Ulcerative colitis	4	No

38	Rheumatoid arthritis	Crohn's disease	4	No
39	Coronary disease	Glioma	5	Yes
40	Colorectal cancer	Crohn's disease	5	No
41	Intracranial aneurysm	LDL cholesterol (Elevated)	5	No
42	Asthma	AIDS progression	6	Yes
43	Colorectal cancer	AIDS progression	6	No
44	Myocardial infarction early	Glioma	7	No
45	Systemic lupus erythematosus	AIDS progression	8	No
46	Myocardial infarction early	AIDS progression	9	Yes
47	Prostate cancer	AIDS progression	10	No
48	LDL cholesterol (Elevated)	Glioma	11	Yes
49	Asthma	Primary biliary cirrhosis	11	No
50	Colorectal cancer	Psoriasis	11	No
51	Glioma	Systemic lupus erythematosus	11	No
52	Intracranial aneurysm	Prostat cancer	11	No
53	LDL cholesterol (Elevated)	Ulcerative colitis	11	No
54	LDL cholesterol (Elevated)	Inflammatory bowel disease	12	Yes
55	AIDS progression	Psoriasis	12	No
56	AIDS progression	Inflammatory bowel disease	12	No
57	Prostate cancer	Psoriasis	15	No
58	Celiac disease	Type 1 diabetes	17	No
59	Multiple sclerosis	Systemic lupus erythematosus	17	No
60	Primary biliary cirrhosis	Psoriasis	19	No
61	Prostate cancer	Ulcerative colitis	23	No
62	Glioma	Asthma	24	Yes
63	Prostate cancer	Inflammatory bowel disease	28	No
64	Crohn's disease	Ulcerative colitis	32	Yes
65	Inflammatory bowel disease	Crohn's disease	33	Yes
66	LDL cholesterol (Elevated)	Psoriasis	33	Yes
67	Multiple sclerosis	Primary biliary cirrhosis	33	No
68	LDL cholesterol (Elevated)	Myocardial infarction early	35	Yes
69	Myocardial infarction early	Inflammatory bowel disease	38	Yes
70	Intracranial aneurysm	Colorectal cancer	41	No
71	Myocardial infarction early	Ulcerative colitis	47	No
72	Coronary disease	AIDS progression	50	Yes
73	Chronic Obstructive Pulmonary	Lung cancer	51	Yes
74	Coronary disease	Inflammatory bowel disease	51	Yes
75	Multiple sclerosis	Rheumatoid arthritis	58	No
76	Coronary disease	Psoriasis	66	Yes
77	Celiac disease	Asthma	66	No
78	Celiac disease	Multiple sclerosis	68	No

79	Glioma	Colorectal cancer	68	No
80	Schizophrenia	Celiac disease	71	No
81	Inflammatory bowel disease	Psoriasis	80	No
82	Myocardial infarction early	Psoriasis	82	No
83	Celiac disease	Primary biliary cirrhosis	86	No
84	Celiac disease	Systemic lupus erythematosus	96	No
85	Type 1 diabetes	Primary biliary cirrhosis	100	No
86	Intracranial aneurysm	Coronary disease	104	No
87	Intracranial aneurysm	Glioma	110	No
88	Intracranial aneurysm	Myocardial infarction early	117	No
89	Primary biliary cirrhosis	Inflammatory bowel disease	117	No
90	Intracranial aneurysm	Type 2 diabetes	137	No
91	Glioma	Prostat cancer	138	No
92	Primary biliary cirrhosis	Ulcerative colitis	145	No
93	Coronary disease	Ulcerative colitis	151	No
94	LDL cholesterol (Elevated)	Coronary disease	160	Yes
95	Intracranial aneurysm	Systemic lupus erythematosus	166	No
96	Celiac disease	Rheumatoid arthritis	178	No
97	Systemic lupus erythematosus	Inflammatory bowel disease	182	No
98	Rheumatoid arthritis	Primary biliary cirrhosis	184	No
99	Psoriasis	Ulcerative colitis	201	No
100	Systemic lupus erythematosus	Ulcerative colitis	207	No
101	Multiple sclerosis	Asthma	233	Yes
102	Asthma	Ulcerative colitis	238	No
103	Asthma	Psoriasis	261	Yes
104	Inflammatory bowel disease	Ulcerative colitis	262	Yes
105	Asthma	Inflammatory bowel disease	271	Yes
106	Obesity	Type 2 diabetes	320	Yes
107	Systemic lupus erythematosus	Psoriasis	521	No
108	Prostate cancer	Colorectal cancer	882	No
109	Rheumatoid arthritis	Asthma	894	Yes
110	Type 1 diabetes	Systemic lupus erythematosus	894	No
111	Schizophrenia	Type 1 diabetes	931	Yes
112	Colorectal cancer	Ulcerative colitis	1019	No
113	Type 1 diabetes	Multiple sclerosis	1032	No
114	Type 1 diabetes	Rheumatoid arthritis	1990	No
115	Intracranial aneurysm	Myeloproliferative neoplasms	4126	No
116	Colorectal cancer	Inflammatory bowel disease	37538	No
117	Myocardial infarction early	Coronary disease	54128	Yes

Appendix A.6 Input to the IsBIG Algorithm for constructing I-Model

	SNP ₁	SNP ₂	LD (r ²)	SNP ₁ OR	SNP ₂ OR	SNP ₁ RAF	SNP ₂ RAF
1	rs564398	rs1412829	0.983	1.13	1.42	0.56	0.39
2	rs11190140	rs10883365	0.983	1.2	1.18	0.48	0.48
3	rs11190140	rs10883365	0.983	1.2	1.2	0.48	0.48
4	rs10883365	rs11190140	0.983	1.18	1.2	0.48	0.48
5	rs10883365	rs11190140	0.983	1.2	1.2	0.48	0.48
6	rs1412829	rs564398	0.983	1.42	1.13	0.39	0.56
7	rs4263839	rs6478109	0.982	1.22	1.36	0.68	0.69
8	rs6478109	rs4263839	0.982	1.36	1.22	0.69	0.68
9	rs6074022	rs4810485	0.977	1.2	1.15	0.25	0.25
10	rs4810485	rs6074022	0.977	1.15	1.2	0.25	0.25
11	rs13277113	rs2736340	0.976	1.39	1.19	0.23	0.24
12	rs2736340	rs13277113	0.976	1.19	1.39	0.24	0.23
13	rs3135388	rs9271366	0.974	2.75	2.78	0.22	0.15
14	rs3135388	rs9271366	0.974	1.99	2.78	0.23	0.15
15	rs9271366	rs3135388	0.974	2.78	2.75	0.15	0.22
16	rs9271366	rs3135388	0.974	2.78	1.99	0.15	0.23
17	rs2981582	rs1219648	0.966	1.26	1.2	0.38	0.4
18	rs7931342	rs10896449	0.966	1.19	1.1	0.51	0.52
19	rs2241880	rs10210302	0.966	1.45	1.19	0.55	0.48
20	rs10210302	rs2241880	0.966	1.19	1.45	0.48	0.55
21	rs1219648	rs2981582	0.966	1.2	1.26	0.4	0.38
22	rs10896449	rs7931342	0.966	1.1	1.19	0.52	0.51
23	rs1335532	rs2300747	0.964	1.28	1.3	0.87	0.88
24	rs2300747	rs1335532	0.964	1.3	1.28	0.88	0.87
25	rs4506565	rs7901695	0.96	1.36	1.37	0.32	NR
26	rs7901695	rs4506565	0.96	1.37	1.36	NR	0.32
27	rs6931514	rs7756992	0.958	1.25	1.2	NR	0.26
28	rs7756992	rs6931514	0.958	1.2	1.25	0.26	NR
29	rs3197999	rs9858542	0.956	1.2	1.17	0.27	0.29
30	rs3197999	rs9858542	0.956	1.2	1.09	0.27	0.28
31	rs9858542	rs3197999	0.956	1.17	1.2	0.29	0.27
32	rs9858542	rs3197999	0.956	1.09	1.2	0.28	0.27
33	rs9941349	rs1121980	0.95	1.48	1.66	0.43	0.41
34	rs907092	rs2872507	0.95	1.29	1.12	0.45	0.47
35	rs2872507	rs907092	0.95	1.12	1.29	0.47	0.45
36	rs1121980	rs9941349	0.95	1.66	1.48	0.41	0.43

37	rs11755527	rs3757247	0.949	1.13	1.13	0.47	NR
38	rs3757247	rs11755527	0.949	1.13	1.13	NR	0.47
39	rs477515	rs2395185	0.948	1.38	1.52	0.69	0.67
40	rs2395185	rs477515	0.948	1.52	1.38	0.67	0.69
41	rs2903692	rs12708716	0.941	1.54	1.19	0.62	0.65
42	rs2903692	rs12708716	0.941	1.54	1.23	0.62	0.68
43	rs12708716	rs2903692	0.941	1.19	1.54	0.65	0.62
44	rs12708716	rs2903692	0.941	1.23	1.54	0.68	0.62
45	rs2201841	rs10889677	0.94	1.13	1.29	0.3	0.3
46	rs10889677	rs2201841	0.94	1.29	1.13	0.3	0.3
47	rs3764021	rs11052552	0.933	1.57	1.49	0.47	0.49
48	rs11052552	rs3764021	0.933	1.49	1.57	0.49	0.47
49	rs3828309	rs2241880	0.932	1.25	1.45	0.53	0.55
50	rs2241880	rs3828309	0.932	1.45	1.25	0.55	0.53
51	rs10210302	rs3828309	0.932	1.19	1.25	0.48	0.53
52	rs8050136	rs1421085	0.932	1.3	1.39	NR	0.4
53	rs8050136	rs1421085	0.932	1.17	1.39	0.38	0.4
54	rs8050136	rs1421085	0.932	1.23	1.39	0.4	0.4
55	rs8050136	rs1421085	0.932	1.15	1.39	NR	0.4
56	rs3828309	rs10210302	0.932	1.25	1.19	0.53	0.48
57	rs1421085	rs8050136	0.932	1.39	1.3	0.4	NR
58	rs1421085	rs8050136	0.932	1.39	1.17	0.4	0.38
59	rs1421085	rs8050136	0.932	1.39	1.23	0.4	0.4
60	rs1421085	rs8050136	0.932	1.39	1.15	0.4	NR
61	rs2814707	rs3849942	0.931	1.22	1.23	0.23	0.23
62	rs3849942	rs2814707	0.931	1.23	1.22	0.23	0.23
63	rs7903146	rs4506565	0.921	1.38	1.36	0.3	0.32
64	rs7903146	rs4506565	0.921	1.31	1.36	NR	0.32
65	rs7903146	rs4506565	0.921	1.49	1.36	NR	0.32
66	rs7903146	rs4506565	0.921	1.71	1.36	NR	0.32
67	rs7903146	rs4506565	0.921	1.34	1.36	0.18	0.32
68	rs7903146	rs4506565	0.921	1.38	1.36	0.26	0.32
69	rs7903146	rs4506565	0.921	1.65	1.36	0.3	0.32
70	rs7903146	rs4506565	0.921	1.37	1.36	NR	0.32
71	rs7903146	rs4506565	0.921	1.48	1.36	0.27	0.32
72	rs4506565	rs7903146	0.921	1.36	1.48	0.32	0.27
73	rs4506565	rs7903146	0.921	1.36	1.38	0.32	0.3
74	rs4506565	rs7903146	0.921	1.36	1.31	0.32	NR
75	rs4506565	rs7903146	0.921	1.36	1.49	0.32	NR
76	rs4506565	rs7903146	0.921	1.36	1.71	0.32	NR
77	rs4506565	rs7903146	0.921	1.36	1.34	0.32	0.18

78	rs4506565	rs7903146	0.921	1.36	1.38	0.32	0.26
79	rs4506565	rs7903146	0.921	1.36	1.65	0.32	0.3
80	rs4506565	rs7903146	0.921	1.36	1.37	0.32	NR
81	rs7901695	rs7903146	0.919	1.37	1.34	NR	0.18
82	rs7901695	rs7903146	0.919	1.37	1.38	NR	0.26
83	rs7901695	rs7903146	0.919	1.37	1.65	NR	0.3
84	rs7901695	rs7903146	0.919	1.37	1.37	NR	NR
85	rs7901695	rs7903146	0.919	1.37	1.48	NR	0.27
86	rs4474514	rs995030	0.919	3.07	2.55	NR	0.8
87	rs4474514	rs995030	0.919	3.07	2.69	NR	0.83
88	rs995030	rs4474514	0.919	2.55	3.07	0.8	NR
89	rs995030	rs4474514	0.919	2.69	3.07	0.83	NR
90	rs599839	rs646776	0.917	1.29	1.19	0.23	0.81
91	rs599839	rs646776	0.917	0.95	1.19	0.24	0.81
92	rs646776	rs599839	0.917	1.19	1.29	0.81	0.23
93	rs646776	rs599839	0.917	1.19	0.95	0.81	0.24
94	rs2981579	rs1219648	0.916	1.17	1.2	0.41	0.4
95	rs1219648	rs2981579	0.916	1.2	1.17	0.4	0.41
96	rs1000113	rs11747270	0.905	1.54	1.33	0.07	0.09
97	rs11209026	rs11465804	0.905	2.92	2.5	0.92	0.93
98	rs11209026	rs11465804	0.905	2.56	2.5	0.94	0.93
99	rs11209026	rs11465804	0.905	3.84	2.5	0.93	0.93
100	rs11209026	rs11465804	0.905	1.79	2.5	0.93	0.93
101	rs11465804	rs11209026	0.905	2.5	2.92	0.93	0.92
102	rs11465804	rs11209026	0.905	2.5	2.56	0.93	0.94
103	rs11465804	rs11209026	0.905	2.5	3.84	0.93	0.93
104	rs11465804	rs11209026	0.905	2.5	1.79	0.93	0.93
105	rs11747270	rs1000113	0.905	1.33	1.54	0.09	0.07
106	rs1121980	rs1421085	0.902	1.66	1.39	0.41	0.4
107	rs6983267	rs10505477	0.902	1.24	1.17	0.48	0.5
108	rs6983267	rs10505477	0.902	1.27	1.17	0.49	0.5
109	rs6983267	rs10505477	0.902	1.42	1.17	0.49	0.5
110	rs6983267	rs10505477	0.902	1.26	1.17	0.5	0.5
111	rs6983267	rs10505477	0.902	1.28	1.17	0.53	0.5
112	rs1421085	rs1121980	0.902	1.39	1.66	0.4	0.41
113	rs10505477	rs6983267	0.902	1.17	1.24	0.5	0.48
114	rs10505477	rs6983267	0.902	1.17	1.27	0.5	0.49
115	rs10505477	rs6983267	0.902	1.17	1.42	0.5	0.49
116	rs10505477	rs6983267	0.902	1.17	1.26	0.5	0.5
117	rs10505477	rs6983267	0.902	1.17	1.28	0.5	0.53
118	rs4977574	rs1333049	0.9	1.29	1.36	0.56	0.47

119	rs4977574	rs1333049	0.9	1.29	1.47	0.56	0.47
120	rs1333049	rs4977574	0.9	1.36	1.29	0.47	0.56
121	rs1333049	rs4977574	0.9	1.47	1.29	0.47	0.56
122	rs8034191	rs1051730	0.899	1.4	1.31	0.33	0.35
123	rs8034191	rs1051730	0.899	1.29	1.31	NR	0.35
124	rs8034191	rs1051730	0.899	1.3	1.31	0.34	0.35
125	rs8034191	rs1051730	0.899	1.3	1.31	NR	0.35
126	rs8034191	rs1051730	0.899	1.38	1.31	NR	0.35
127	rs8034191	rs1051730	0.899	1.29	1.35	NR	NR
128	rs8034191	rs1051730	0.899	1.3	1.35	0.34	NR
129	rs8034191	rs1051730	0.899	1.3	1.35	NR	NR
130	rs8034191	rs1051730	0.899	1.38	1.35	NR	NR
131	rs8034191	rs1051730	0.899	1.4	1.35	0.33	NR
132	rs1051730	rs8034191	0.899	1.31	1.4	0.35	0.33
133	rs1051730	rs8034191	0.899	1.35	1.4	NR	0.33
134	rs1051730	rs8034191	0.899	1.35	1.29	NR	NR
135	rs1051730	rs8034191	0.899	1.35	1.3	NR	0.34
136	rs1051730	rs8034191	0.899	1.35	1.3	NR	NR
137	rs1051730	rs8034191	0.899	1.35	1.38	NR	NR
138	rs1051730	rs8034191	0.899	1.31	1.29	0.35	NR
139	rs1051730	rs8034191	0.899	1.31	1.3	0.35	0.34
140	rs1051730	rs8034191	0.899	1.31	1.3	0.35	NR
141	rs1051730	rs8034191	0.899	1.31	1.38	0.35	NR
142	rs17221417	rs2076756	0.892	1.29	1.71	0.29	0.27
143	rs2076756	rs17221417	0.892	1.71	1.29	0.27	0.29
144	rs2981582	rs2981579	0.884	1.26	1.17	0.38	0.41
145	rs2981579	rs2981582	0.884	1.17	1.26	0.41	0.38
146	rs9941349	rs1421085	0.884	1.48	1.39	0.43	0.4
147	rs9941349	rs8050136	0.884	1.48	1.3	0.43	NR
148	rs9941349	rs8050136	0.884	1.48	1.17	0.43	0.38
149	rs9941349	rs8050136	0.884	1.48	1.23	0.43	0.4
150	rs9941349	rs8050136	0.884	1.48	1.15	0.43	NR
151	rs1421085	rs9941349	0.884	1.39	1.48	0.4	0.43
152	rs8050136	rs9941349	0.884	1.3	1.48	NR	0.43
153	rs8050136	rs9941349	0.884	1.17	1.48	0.38	0.43
154	rs8050136	rs9941349	0.884	1.23	1.48	0.4	0.43
155	rs8050136	rs9941349	0.884	1.15	1.48	NR	0.43
156	rs4975616	rs401681	0.882	1.15	1.15	NR	NR
157	rs401681	rs4975616	0.882	1.15	1.15	NR	NR
158	rs8050136	rs1121980	0.87	1.3	1.66	NR	0.41
159	rs8050136	rs1121980	0.87	1.17	1.66	0.38	0.41

160	rs8050136	rs1121980	0.87	1.23	1.66	0.4	0.41
161	rs8050136	rs1121980	0.87	1.15	1.66	NR	0.41
162	rs1121980	rs8050136	0.87	1.66	1.3	0.41	NR
163	rs1121980	rs8050136	0.87	1.66	1.17	0.41	0.38
164	rs1121980	rs8050136	0.87	1.66	1.23	0.41	0.4
165	rs1121980	rs8050136	0.87	1.66	1.15	0.41	NR
166	rs1701704	rs2292239	0.851	1.25	1.28	0.35	0.34
167	rs2292239	rs1701704	0.851	1.28	1.25	0.34	0.35
168	rs3135388	rs3129934	0.847	2.75	3.3	0.22	NR
169	rs3135388	rs3129934	0.847	1.99	3.3	0.23	NR
170	rs3129934	rs3135388	0.847	3.3	2.75	NR	0.22
171	rs3129934	rs3135388	0.847	3.3	1.99	NR	0.23
172	rs2943641	rs2943634	0.843	1.19	1.21	0.63	0.65
173	rs2943634	rs2943641	0.843	1.21	1.19	0.65	0.63
174	rs774359	rs2814707	0.831	1.19	1.22	0.25	0.23
175	rs2814707	rs774359	0.831	1.22	1.19	0.23	0.25
176	rs2872507	rs7216389	0.826	1.12	1.45	0.47	0.52
177	rs7216389	rs2872507	0.826	1.45	1.12	0.52	0.47
178	rs9271366	rs3129934	0.824	2.78	3.3	0.15	NR
179	rs3129934	rs9271366	0.824	3.3	2.78	NR	0.15
180	rs401681	rs31489	0.821	1.15	1.12	NR	0.59
181	rs31489	rs401681	0.821	1.12	1.15	0.59	NR
182	rs3849942	rs774359	0.811	1.23	1.19	0.23	0.25
183	rs4788084	rs8049439	0.811	1.09	1.14	0.42	0.37
184	rs8049439	rs4788084	0.811	1.14	1.09	0.37	0.42
185	rs774359	rs3849942	0.811	1.19	1.23	0.25	0.23
186	rs907092	rs7216389	0.808	1.29	1.45	0.45	0.52
187	rs7216389	rs907092	0.808	1.45	1.29	0.52	0.45
188	rs947474	rs4750316	0.796	1.1	1.14	0.19	0.2
189	rs4750316	rs947474	0.796	1.14	1.1	0.2	0.19
190	rs258322	rs1805007	0.783	1.67	2.34	0.09	0.08
191	rs258322	rs1805007	0.783	1.67	4.37	0.09	0.05
192	rs258322	rs1805007	0.783	1.67	12.47	0.09	NR
193	rs258322	rs1805007	0.783	1.67	2.94	0.09	0.06
194	rs1805007	rs258322	0.783	2.34	1.67	0.08	0.09
195	rs1805007	rs258322	0.783	4.37	1.67	0.05	0.09
196	rs1805007	rs258322	0.783	12.47	1.67	NR	0.09
197	rs1805007	rs258322	0.783	2.94	1.67	0.06	0.09
198	rs7193343	rs2106261	0.776	1.21	1.25	NR	0.174
199	rs2106261	rs7193343	0.776	1.25	1.21	0.174	NR
200	rs31489	rs4975616	0.74	1.12	1.15	0.59	NR

201	rs4975616	rs31489	0.74	1.15	1.12	NR	0.59
202	rs7501939	rs4430796	0.734	1.41	1.22	0.57	0.49
203	rs7501939	rs4430796	0.734	1.41	1.19	0.57	0.52
204	rs7501939	rs4430796	0.734	1.41	1.18	0.57	0.54
205	rs4430796	rs7501939	0.734	1.22	1.41	0.49	0.57
206	rs4430796	rs7501939	0.734	1.19	1.41	0.52	0.57
207	rs4430796	rs7501939	0.734	1.18	1.41	0.54	0.57
208	rs4977756	rs1412829	0.724	1.24	1.42	0.6	0.39
209	rs1412829	rs4977756	0.724	1.42	1.24	0.39	0.6
210	rs7756992	rs7754840	0.722	1.2	1.12	0.26	0.31
211	rs7756992	rs7754840	0.722	1.2	1.12	0.26	0.36
212	rs7754840	rs7756992	0.722	1.12	1.2	0.31	0.26
213	rs7754840	rs7756992	0.722	1.12	1.2	0.36	0.26
214	rs4712523	rs7756992	0.722	1.2	1.2	0.32	0.26
215	rs10946398	rs7756992	0.722	1.18	1.2	NR	0.26
216	rs10946398	rs7756992	0.722	1.16	1.2	0.32	0.26
217	rs7756992	rs4712523	0.722	1.2	1.2	0.26	0.32
218	rs7756992	rs10946398	0.722	1.2	1.18	0.26	NR
219	rs7756992	rs10946398	0.722	1.2	1.16	0.26	0.32
220	rs564398	rs4977756	0.71	1.13	1.24	0.56	0.6
221	rs4977756	rs564398	0.71	1.24	1.13	0.6	0.56
222	rs2201841	rs11805303	0.7	1.13	1.39	0.3	0.68
223	rs11805303	rs2201841	0.7	1.39	1.13	0.68	0.3
224	rs6931514	rs4712523	0.688	1.25	1.2	NR	0.32
225	rs6931514	rs10946398	0.688	1.25	1.18	NR	NR
226	rs6931514	rs10946398	0.688	1.25	1.16	NR	0.32
227	rs4712523	rs6931514	0.688	1.2	1.25	0.32	NR
228	rs10946398	rs6931514	0.688	1.18	1.25	NR	NR
229	rs10946398	rs6931514	0.688	1.16	1.25	0.32	NR
230	rs7754840	rs6931514	0.688	1.12	1.25	0.31	NR
231	rs7754840	rs6931514	0.688	1.12	1.25	0.36	NR
232	rs6931514	rs7754840	0.688	1.25	1.12	NR	0.31
233	rs6931514	rs7754840	0.688	1.25	1.12	NR	0.36
234	rs10758593	rs7020673	0.674	1.13	1.14	NR	0.5
235	rs2292239	rs11171739	0.674	1.28	1.34	0.34	0.42
236	rs7020673	rs10758593	0.674	1.14	1.13	0.5	NR
237	rs11171739	rs2292239	0.674	1.34	1.28	0.42	0.34
238	rs6932590	rs13194053	0.653	1.16	1.28	0.78	0.82
239	rs6932590	rs13194053	0.653	1.16	1.22	0.78	0.86
240	rs13194053	rs6932590	0.653	1.28	1.16	0.82	0.78
241	rs13194053	rs6932590	0.653	1.22	1.16	0.86	0.78

242	rs10889677	rs11805303	0.649	1.29	1.39	0.3	0.68
243	rs11805303	rs10889677	0.649	1.39	1.29	0.68	0.3
244	rs12722489	rs2104286	0.626	1.25	1.16	0.85	0.73
245	rs12722489	rs2104286	0.626	1.25	1.15	0.85	0.76
246	rs2104286	rs12722489	0.626	1.16	1.25	0.73	0.85
247	rs2104286	rs12722489	0.626	1.15	1.25	0.76	0.85
248	rs11171739	rs1701704	0.618	1.34	1.25	0.42	0.35
249	rs1701704	rs11171739	0.618	1.25	1.34	0.35	0.42
250	rs4598195	rs4730276	0.614	1.23	1.22	0.54	0.39
251	rs4730276	rs4598195	0.614	1.22	1.23	0.39	0.54
252	rs2076756	rs5743289	0.612	1.71	1.46	0.27	0.17
253	rs5743289	rs2076756	0.612	1.46	1.71	0.17	0.27
254	rs4977574	rs1333040	0.603	1.29	1.29	0.56	0.55
255	rs1333040	rs4977574	0.603	1.29	1.29	0.55	0.56
256	rs2180439	rs1160312	0.602	1.82	1.6	0.43	0.43
257	rs1160312	rs2180439	0.602	1.6	1.82	0.43	0.43
258	rs9888739	rs11574637	0.556	1.62	1.33	0.13	0.19
259	rs11574637	rs9888739	0.556	1.33	1.62	0.19	0.13
260	rs1333049	rs1333040	0.555	1.36	1.29	0.47	0.55
261	rs1333049	rs1333040	0.555	1.47	1.29	0.47	0.55
262	rs1333040	rs1333049	0.555	1.29	1.36	0.55	0.47
263	rs1333040	rs1333049	0.555	1.29	1.47	0.55	0.47
264	rs4730276	rs4730273	0.551	1.22	1.22	0.39	0.7
265	rs4730273	rs4730276	0.551	1.22	1.22	0.7	0.39
266	rs17221417	rs5743289	0.546	1.29	1.46	0.29	0.17
267	rs5743289	rs17221417	0.546	1.46	1.29	0.17	0.29
268	rs7014346	rs10505477	0.541	1.19	1.17	0.18	0.5
269	rs10505477	rs7014346	0.541	1.17	1.19	0.5	0.18
270	rs477515	rs9272346	0.512	1.38	5.49	0.69	0.61
271	rs9272346	rs477515	0.512	5.49	1.38	0.61	0.69
272	rs9465871	rs7756992	0.509	1.18	1.2	0.18	0.26
273	rs7756992	rs9465871	0.509	1.2	1.18	0.26	0.18
274	rs17696736	rs653178	0.505	1.22	1.21	0.42	0.48
275	rs17696736	rs653178	0.505	1.34	1.21	0.42	0.48
276	rs653178	rs17696736	0.505	1.21	1.22	0.48	0.42
277	rs653178	rs17696736	0.505	1.21	1.34	0.48	0.42
278	rs9272346	rs2395185	0.497	5.49	1.52	0.61	0.67
279	rs2395185	rs9272346	0.497	1.52	5.49	0.67	0.61
280	rs2412973	rs5753037	0.493	1.15	1.1	0.46	0.39
281	rs5753037	rs2412973	0.493	1.1	1.15	0.39	0.46
282	rs7014346	rs6983267	0.488	1.19	1.42	0.18	0.49

283	rs7014346	rs6983267	0.488	1.19	1.26	0.18	0.5
284	rs7014346	rs6983267	0.488	1.19	1.28	0.18	0.53
285	rs6983267	rs7014346	0.488	1.24	1.19	0.48	0.18
286	rs6983267	rs7014346	0.488	1.27	1.19	0.49	0.18
287	rs6983267	rs7014346	0.488	1.42	1.19	0.49	0.18
288	rs6983267	rs7014346	0.488	1.26	1.19	0.5	0.18
289	rs6983267	rs7014346	0.488	1.28	1.19	0.53	0.18
290	rs7014346	rs6983267	0.488	1.19	1.24	0.18	0.48
291	rs7014346	rs6983267	0.488	1.19	1.27	0.18	0.49
292	rs4763879	rs11052552	0.484	1.09	1.49	0.37	0.49
293	rs11052552	rs4763879	0.484	1.49	1.09	0.49	0.37
294	rs6897932	rs1445898	0.467	1.12	1.12	0.75	0.55
295	rs6897932	rs1445898	0.467	1.18	1.12	0.75	0.55
296	rs6897932	rs1445898	0.467	1.12	1.12	0.71	0.55
297	rs1445898	rs6897932	0.467	1.12	1.12	0.55	0.75
298	rs1445898	rs6897932	0.467	1.12	1.18	0.55	0.75
299	rs1445898	rs6897932	0.467	1.12	1.12	0.55	0.71
300	rs4763879	rs3764021	0.451	1.09	1.57	0.37	0.47
301	rs3764021	rs4763879	0.451	1.57	1.09	0.47	0.37
302	rs10038113	rs4307059	0.45	1.33	1.19	0.59	0.61
303	rs4307059	rs10038113	0.45	1.19	1.33	0.61	0.59
304	rs4977756	rs4977574	0.446	1.24	1.29	0.6	0.56
305	rs4977574	rs4977756	0.446	1.29	1.24	0.56	0.6
306	rs9296249	rs3923809	0.442	1.67	1.9	0.76	0.66
307	rs3923809	rs9296249	0.442	1.9	1.67	0.66	0.76
308	rs10484554	rs2395029	0.432	2.8	3.47	0.15	0.03
309	rs12191877	rs2395029	0.432	2.64	3.47	0.15	0.03
310	rs12191877	rs2395029	0.432	2.64	45	0.15	0.05
311	rs10484554	rs2395029	0.432	2.8	45	0.15	0.05
312	rs10484554	rs2395029	0.432	2.8	4.1	0.15	0.03
313	rs12191877	rs2395029	0.432	2.64	4.1	0.15	0.03
314	rs2395029	rs10484554	0.432	3.47	2.8	0.03	0.15
315	rs2395029	rs10484554	0.432	45	2.8	0.05	0.15
316	rs2395029	rs10484554	0.432	4.1	2.8	0.03	0.15
317	rs2395029	rs12191877	0.432	4.1	2.64	0.03	0.15
318	rs10974944	rs10758669	0.477	3.1	1.12	NR	0.35
319	rs10758669	rs10974944	0.477	1.12	3.1	0.35	NR
320	rs9465871	rs6931514	0.473	1.18	1.25	0.18	NR
321	rs6931514	rs9465871	0.473	1.25	1.18	NR	0.18
322	rs6897932	rs1445898	0.467	1.12	1.12	0.75	0.55
323	rs6897932	rs1445898	0.467	1.18	1.12	0.75	0.55

324	rs6897932	rs1445898	0.467	1.12	1.12	0.71	0.55
325	rs1445898	rs6897932	0.467	1.12	1.12	0.55	0.75
326	rs1445898	rs6897932	0.467	1.12	1.18	0.55	0.75
327	rs1445898	rs6897932	0.467	1.12	1.12	0.55	0.71
328	rs4763879	rs3764021	0.451	1.09	1.57	0.37	0.47
329	rs3764021	rs4763879	0.451	1.57	1.09	0.47	0.37
330	rs10038113	rs4307059	0.45	1.33	1.19	0.59	0.61
331	rs4307059	rs10038113	0.45	1.19	1.33	0.61	0.59
332	rs4977756	rs4977574	0.446	1.24	1.29	0.6	0.56
333	rs4977574	rs4977756	0.446	1.29	1.24	0.56	0.6
334	rs9296249	rs3923809	0.442	1.67	1.9	0.76	0.66
335	rs3923809	rs9296249	0.442	1.9	1.67	0.66	0.76
336	rs10484554	rs2395029	0.432	2.8	3.47	0.15	0.03
337	rs12191877	rs2395029	0.432	2.64	3.47	0.15	0.03
338	rs12191877	rs2395029	0.432	2.64	45	0.15	0.05
339	rs10484554	rs2395029	0.432	2.8	45	0.15	0.05
340	rs10484554	rs2395029	0.432	2.8	4.1	0.15	0.03
341	rs12191877	rs2395029	0.432	2.64	4.1	0.15	0.03
342	rs2395029	rs10484554	0.432	3.47	2.8	0.03	0.15
343	rs2395029	rs10484554	0.432	45	2.8	0.05	0.15
344	rs2395029	rs10484554	0.432	4.1	2.8	0.03	0.15
345	rs2395029	rs12191877	0.432	4.1	2.64	0.03	0.15
346	rs2395029	rs12191877	0.432	3.47	2.64	0.03	0.15
347	rs2395029	rs12191877	0.432	45	2.64	0.05	0.15
348	rs6010620	rs2315008	0.428	1.28	1.36	0.23	0.69
349	rs6010620	rs2315008	0.428	1.52	1.36	0.77	0.69
350	rs2315008	rs6010620	0.428	1.36	1.28	0.69	0.23
351	rs2315008	rs6010620	0.428	1.36	1.52	0.69	0.77
352	rs10488631	rs12537284	0.426	1.52	1.54	NR	0.13
353	rs12537284	rs10488631	0.426	1.54	1.52	0.13	NR
354	rs2187668	rs9272219	0.4	7.04	1.14	0.14	0.72
355	rs9272219	rs2187668	0.4	1.14	7.04	0.72	0.14
356	rs660895	rs477515	0.389	3.62	1.38	0.21	0.69
357	rs477515	rs660895	0.389	1.38	3.62	0.69	0.21
358	rs2158836	rs4598195	0.388	1.21	1.23	0.35	0.54
359	rs4598195	rs2158836	0.388	1.23	1.21	0.54	0.35
360	rs9465871	rs10946398	0.385	1.18	1.18	0.18	NR
361	rs9465871	rs10946398	0.385	1.18	1.16	0.18	0.32
362	rs9465871	rs4712523	0.385	1.18	1.2	0.18	0.32
363	rs9465871	rs7754840	0.385	1.18	1.12	0.18	0.31
364	rs9465871	rs7754840	0.385	1.18	1.12	0.18	0.36

365	rs7754840	rs9465871	0.385	1.12	1.18	0.31	0.18
366	rs7754840	rs9465871	0.385	1.12	1.18	0.36	0.18
367	rs4712523	rs9465871	0.385	1.2	1.18	0.32	0.18
368	rs10946398	rs9465871	0.385	1.18	1.18	NR	0.18
369	rs10946398	rs9465871	0.385	1.16	1.18	0.32	0.18
370	rs1333049	rs4977756	0.384	1.36	1.24	0.47	0.6
371	rs1333049	rs4977756	0.384	1.47	1.24	0.47	0.6
372	rs4977756	rs1333049	0.384	1.24	1.36	0.6	0.47
373	rs4977756	rs1333049	0.384	1.24	1.47	0.6	0.47
374	rs660895	rs2395185	0.381	3.62	1.52	0.21	0.67
375	rs2395185	rs660895	0.381	1.52	3.62	0.67	0.21
376	rs4730273	rs4598195	0.374	1.22	1.23	0.7	0.54
377	rs4598195	rs4730273	0.374	1.23	1.22	0.54	0.7
378	rs9929218	rs1728785	0.368	1.1	1.17	0.29	0.76
379	rs1728785	rs9929218	0.368	1.17	1.1	0.76	0.29
380	rs17594526	rs9960767	0.35	1.44	1.23	0.03	0.06
381	rs9960767	rs17594526	0.35	1.23	1.44	0.06	0.03
382	rs660895	rs6457617	0.344	3.62	2.36	0.21	0.49
383	rs660895	rs6457620	0.344	3.62	2.55	0.21	0.5
384	rs6457617	rs660895	0.344	2.36	3.62	0.49	0.21
385	rs6457620	rs660895	0.344	2.55	3.62	0.5	0.21
386	rs9272346	rs9271366	0.331	5.49	2.78	0.61	0.15
387	rs9271366	rs9272346	0.331	2.78	5.49	0.15	0.61
388	rs3135388	rs9272346	0.322	2.75	5.49	0.22	0.61
389	rs3135388	rs9272346	0.322	1.99	5.49	0.23	0.61
390	rs9272346	rs3135388	0.322	5.49	2.75	0.61	0.22
391	rs9272346	rs3135388	0.322	5.49	1.99	0.61	0.23
392	rs9272346	rs3129934	0.315	5.49	3.3	0.61	NR
393	rs3129934	rs9272346	0.315	3.3	5.49	NR	0.61
394	rs11805303	rs7517847	0.314	1.39	1.61	0.68	0.56
395	rs7517847	rs11805303	0.314	1.61	1.39	0.56	0.68
396	rs11228565	rs10896449	0.303	1.23	1.1	0.2	0.52
397	rs10896449	rs11228565	0.303	1.1	1.23	0.52	0.2

NR – Not Reported – assume RAF = 0.5

Disease /Trait associated with each SNP (from above)

	SNP ₁ - Disease	SNP ₂ - Disease
1	Type 2 diabetes	Glioma (high-grade)
2	Crohn's disease	Crohn's disease
3	Crohn's disease	Crohn's disease
4	Crohn's disease	Crohn's disease

5	Crohn's disease	Crohn's disease
6	Glioma (high-grade)	Type 2 diabetes
7	Crohn's disease	Inflammatory bowel disease
8	Inflammatory bowel disease	Crohn's disease
9	Multiple sclerosis	Rheumatoid arthritis
10	Rheumatoid arthritis	Multiple sclerosis
11	Systemic lupus erythematosus	Rheumatoid arthritis
12	Rheumatoid arthritis	Systemic lupus erythematosus
13	Multiple sclerosis	Multiple sclerosis
14	Multiple sclerosis	Multiple sclerosis
15	Multiple sclerosis	Multiple sclerosis
16	Multiple sclerosis	Multiple sclerosis
17	Breast cancer	Breast cancer
18	Prostate cancer	Prostate cancer
19	Crohn's disease	Crohn's disease
20	Crohn's disease	Crohn's disease
21	Breast cancer	Breast cancer
22	Prostate cancer	Prostate cancer
23	Multiple sclerosis	Multiple sclerosis
24	Multiple sclerosis	Multiple sclerosis
25	Type 2 diabetes	Type 2 diabetes
26	Type 2 diabetes	Type 2 diabetes
27	Type 2 diabetes	Type 2 diabetes
28	Type 2 diabetes	Type 2 diabetes
29	Crohn's disease	Crohn's disease
30	Crohn's disease	Crohn's disease
31	Crohn's disease	Crohn's disease
32	Crohn's disease	Crohn's disease
33	Obesity (extreme)	Obesity (early onset extreme)
34	Primary biliary cirrhosis	Crohn's disease
35	Crohn's disease	Primary biliary cirrhosis
36	Obesity (early onset extreme)	Obesity (extreme)
37	Type 1 diabetes	Type 1 diabetes
38	Type 1 diabetes	Type 1 diabetes
39	Inflammatory bowel disease	Ulcerative colitis

40	Ulcerative colitis	Inflammatory bowel disease
41	Type 1 diabetes	Type 1 diabetes
42	Type 1 diabetes	Type 1 diabetes
43	Type 1 diabetes	Type 1 diabetes
44	Type 1 diabetes	Type 1 diabetes
45	Psoriasis	Ulcerative colitis
46	Ulcerative colitis	Psoriasis
47	Type 1 diabetes	Type 1 diabetes
48	Type 1 diabetes	Type 1 diabetes
49	Crohn's disease	Crohn's disease
50	Crohn's disease	Crohn's disease
51	Crohn's disease	Crohn's disease
52	Type 2 diabetes	Obesity
53	Type 2 diabetes	Obesity
54	Type 2 diabetes	Obesity
55	Type 2 diabetes	Obesity
56	Crohn's disease	Crohn's disease
57	Obesity	Type 2 diabetes
58	Obesity	Type 2 diabetes
59	Obesity	Type 2 diabetes
60	Obesity	Type 2 diabetes
61	Amyotrophic lateral sclerosis	Amyotrophic lateral sclerosis
62	Amyotrophic lateral sclerosis	Amyotrophic lateral sclerosis
63	Type 2 diabetes	Type 2 diabetes
64	Type 2 diabetes	Type 2 diabetes
65	Type 2 diabetes	Type 2 diabetes
66	Type 2 diabetes	Type 2 diabetes
67	Type 2 diabetes	Type 2 diabetes
68	Type 2 diabetes	Type 2 diabetes
69	Type 2 diabetes	Type 2 diabetes
70	Type 2 diabetes	Type 2 diabetes
71	Type 2 diabetes and other traits	Type 2 diabetes
72	Type 2 diabetes	Type 2 diabetes and other traits
73	Type 2 diabetes	Type 2 diabetes
74	Type 2 diabetes	Type 2 diabetes
75	Type 2 diabetes	Type 2 diabetes
76	Type 2 diabetes	Type 2 diabetes

77	Type 2 diabetes	Type 2 diabetes
78	Type 2 diabetes	Type 2 diabetes
79	Type 2 diabetes	Type 2 diabetes
80	Type 2 diabetes	Type 2 diabetes
81	Type 2 diabetes	Type 2 diabetes
82	Type 2 diabetes	Type 2 diabetes
83	Type 2 diabetes	Type 2 diabetes
84	Type 2 diabetes	Type 2 diabetes
85	Type 2 diabetes	Type 2 diabetes and other traits
86	Testicular cancer	Testicular germ cell tumor
87	Testicular cancer	Testicular germ cell tumor
88	Testicular germ cell tumor	Testicular cancer
89	Testicular germ cell tumor	Testicular cancer
90	Coronary disease	Myocardial infarction (early onset)
91	LDL cholesterol	Myocardial infarction (early onset)
92	Myocardial infarction (early onset)	Coronary disease
93	Myocardial infarction (early onset)	LDL cholesterol
94	Breast cancer	Breast cancer
95	Breast cancer	Breast cancer
96	Crohn's disease	Crohn's disease
97	Crohn's disease	Crohn's disease
98	Inflammatory bowel disease	Crohn's disease
99	Inflammatory bowel disease	Crohn's disease
100	Ulcerative colitis	Crohn's disease
101	Crohn's disease	Crohn's disease
102	Crohn's disease	Inflammatory bowel disease
103	Crohn's disease	Inflammatory bowel disease
104	Crohn's disease	Ulcerative colitis
105	Crohn's disease	Crohn's disease
106	Obesity (early onset extreme)	Obesity
107	Colorectal cancer	Colorectal cancer
108	Colorectal cancer	Colorectal cancer

109	Prostate cancer	Colorectal cancer
110	Prostate cancer	Colorectal cancer
111	Prostate cancer	Colorectal cancer
112	Obesity	Obesity (early onset extreme)
113	Colorectal cancer	Colorectal cancer
114	Colorectal cancer	Colorectal cancer
115	Colorectal cancer	Prostate cancer
116	Colorectal cancer	Prostate cancer
117	Colorectal cancer	Prostate cancer
118	Myocardial infarction (early onset)	Coronary disease
119	Myocardial infarction (early onset)	Coronary disease
120	Coronary disease	Myocardial infarction (early onset)
121	Coronary disease	Myocardial infarction (early onset)
122	Chronic Obstructive Pulmonary Disease	Lung adenocarcinoma
123	Lung cancer	Lung adenocarcinoma
124	Lung cancer	Lung adenocarcinoma
125	Lung cancer	Lung adenocarcinoma
126	Lung cancer	Lung adenocarcinoma
127	Lung cancer	Lung cancer
128	Lung cancer	Lung cancer
129	Lung cancer	Lung cancer
130	Lung cancer	Lung cancer
131	Chronic Obstructive Pulmonary Disease	Lung cancer
132	Lung adenocarcinoma	Chronic Obstructive Pulmonary Disease
133	Lung cancer	Chronic Obstructive Pulmonary Disease
134	Lung cancer	Lung cancer
135	Lung cancer	Lung cancer
136	Lung cancer	Lung cancer
137	Lung cancer	Lung cancer
138	Lung adenocarcinoma	Lung cancer
139	Lung adenocarcinoma	Lung cancer
140	Lung adenocarcinoma	Lung cancer
141	Lung adenocarcinoma	Lung cancer
142	Crohn's disease	Crohn's disease

143	Crohn's disease	Crohn's disease
144	Breast cancer	Breast cancer
145	Breast cancer	Breast cancer
146	Obesity (extreme)	Obesity
147	Obesity (extreme)	Type 2 diabetes
148	Obesity (extreme)	Type 2 diabetes
149	Obesity (extreme)	Type 2 diabetes
150	Obesity (extreme)	Type 2 diabetes
151	Obesity	Obesity (extreme)
152	Type 2 diabetes	Obesity (extreme)
153	Type 2 diabetes	Obesity (extreme)
154	Type 2 diabetes	Obesity (extreme)
155	Type 2 diabetes	Obesity (extreme)
156	Lung cancer	Lung cancer
157	Lung cancer	Lung cancer
158	Type 2 diabetes	Obesity (early onset extreme)
159	Type 2 diabetes	Obesity (early onset extreme)
160	Type 2 diabetes	Obesity (early onset extreme)
161	Type 2 diabetes	Obesity (early onset extreme)
162	Obesity (early onset extreme)	Type 2 diabetes
163	Obesity (early onset extreme)	Type 2 diabetes
164	Obesity (early onset extreme)	Type 2 diabetes
165	Obesity (early onset extreme)	Type 2 diabetes
166	Type 1 diabetes	Type 1 diabetes
167	Type 1 diabetes	Type 1 diabetes
168	Multiple sclerosis	Multiple sclerosis
169	Multiple sclerosis	Multiple sclerosis
170	Multiple sclerosis	Multiple sclerosis
171	Multiple sclerosis	Multiple sclerosis
172	Type 2 diabetes and other traits	Coronary disease
173	Coronary disease	Type 2 diabetes and other traits
174	Amyotrophic lateral sclerosis	Amyotrophic lateral sclerosis

175	Amyotrophic lateral sclerosis	Amyotrophic lateral sclerosis
176	Crohn's disease	Asthma
177	Asthma	Crohn's disease
178	Multiple sclerosis	Multiple sclerosis
179	Multiple sclerosis	Multiple sclerosis
180	Lung cancer	Lung adenocarcinoma
181	Lung adenocarcinoma	Lung cancer
182	Amyotrophic lateral sclerosis	Amyotrophic lateral sclerosis
183	Type 1 diabetes	Inflammatory bowel disease (early onset)
184	Inflammatory bowel disease (early onset)	Type 1 diabetes
185	Amyotrophic lateral sclerosis	Amyotrophic lateral sclerosis
186	Primary biliary cirrhosis	Asthma
187	Asthma	Primary biliary cirrhosis
188	Type 1 diabetes	Rheumatoid arthritis
189	Rheumatoid arthritis	Type 1 diabetes
190	Melanoma	Blond vs. brown hair color
191	Melanoma	Freckles
192	Melanoma	Red vs non-red hair color
193	Melanoma	Skin sensitivity to sun
194	Blond vs. brown hair color	Melanoma
195	Freckles	Melanoma
196	Red vs non-red hair color	Melanoma
197	Skin sensitivity to sun	Melanoma
198	Atrial fibrillation	Atrial fibrillation
199	Atrial fibrillation	Atrial fibrillation
200	Lung adenocarcinoma	Lung cancer
201	Lung cancer	Lung adenocarcinoma
202	Prostate cancer	Prostate cancer
203	Prostate cancer	Prostate cancer
204	Prostate cancer	Prostate cancer
205	Prostate cancer	Prostate cancer
206	Prostate cancer	Prostate cancer
207	Prostate cancer	Prostate cancer
208	Glioma	Glioma (high-grade)
209	Glioma (high-grade)	Glioma

210	Type 2 diabetes	Type 2 diabetes
211	Type 2 diabetes	Type 2 diabetes
212	Type 2 diabetes	Type 2 diabetes
213	Type 2 diabetes	Type 2 diabetes
214	Type 2 diabetes and other traits	Type 2 diabetes
215	Type 2 diabetes	Type 2 diabetes
216	Type 2 diabetes	Type 2 diabetes
217	Type 2 diabetes	Type 2 diabetes and other traits
218	Type 2 diabetes	Type 2 diabetes
219	Type 2 diabetes	Type 2 diabetes
220	Type 2 diabetes	Glioma
221	Glioma	Type 2 diabetes
222	Psoriasis	Crohn's disease
223	Crohn's disease	Psoriasis
224	Type 2 diabetes	Type 2 diabetes and other traits
225	Type 2 diabetes	Type 2 diabetes
226	Type 2 diabetes	Type 2 diabetes
227	Type 2 diabetes and other traits	Type 2 diabetes
228	Type 2 diabetes	Type 2 diabetes
229	Type 2 diabetes	Type 2 diabetes
230	Type 2 diabetes	Type 2 diabetes
231	Type 2 diabetes	Type 2 diabetes
232	Type 2 diabetes	Type 2 diabetes
233	Type 2 diabetes	Type 2 diabetes
234	Type 1 diabetes	Type 1 diabetes
235	Type 1 diabetes	Type 1 diabetes
236	Type 1 diabetes	Type 1 diabetes
237	Type 1 diabetes	Type 1 diabetes
238	Schizophrenia	Schizophrenia
239	Schizophrenia	Schizophrenia
240	Schizophrenia	Schizophrenia
241	Schizophrenia	Schizophrenia
242	Ulcerative colitis	Crohn's disease
243	Crohn's disease	Ulcerative colitis
244	Multiple sclerosis	Multiple sclerosis
245	Multiple sclerosis	Multiple sclerosis
246	Multiple sclerosis	Multiple sclerosis
247	Multiple sclerosis	Multiple sclerosis

248	Type 1 diabetes	Type 1 diabetes
249	Type 1 diabetes	Type 1 diabetes
250	Ulcerative colitis	Ulcerative colitis
251	Ulcerative colitis	Ulcerative colitis
252	Crohn's disease	Inflammatory bowel disease
253	Inflammatory bowel disease	Crohn's disease
254	Myocardial infarction (early onset)	Intracranial aneurysm
255	Intracranial aneurysm	Myocardial infarction (early onset)
256	Male-pattern baldness	Male-pattern baldness
257	Male-pattern baldness	Male-pattern baldness
258	Systemic lupus erythematosus	Systemic lupus erythematosus
259	Systemic lupus erythematosus	Systemic lupus erythematosus
260	Coronary disease	Intracranial aneurysm
261	Coronary disease	Intracranial aneurysm
262	Intracranial aneurysm	Coronary disease
263	Intracranial aneurysm	Coronary disease
264	Ulcerative colitis	Ulcerative colitis
265	Ulcerative colitis	Ulcerative colitis
266	Crohn's disease	Inflammatory bowel disease
267	Inflammatory bowel disease	Crohn's disease
268	Colorectal cancer	Colorectal cancer
269	Colorectal cancer	Colorectal cancer
270	Inflammatory bowel disease	Type 1 diabetes
271	Type 1 diabetes	Inflammatory bowel disease
272	Type 2 diabetes	Type 2 diabetes
273	Type 2 diabetes	Type 2 diabetes
274	Type 1 diabetes	Celiac disease
275	Type 1 diabetes	Celiac disease
276	Celiac disease	Type 1 diabetes
277	Celiac disease	Type 1 diabetes
278	Type 1 diabetes	Ulcerative colitis
279	Ulcerative colitis	Type 1 diabetes
280	Inflammatory bowel	Type 1 diabetes

	disease (early onset)	
281	Type 1 diabetes	Inflammatory bowel disease (early onset)
282	Colorectal cancer	Prostate cancer
283	Colorectal cancer	Prostate cancer
284	Colorectal cancer	Prostate cancer
285	Colorectal cancer	Colorectal cancer
286	Colorectal cancer	Colorectal cancer
287	Prostate cancer	Colorectal cancer
288	Prostate cancer	Colorectal cancer
289	Prostate cancer	Colorectal cancer
290	Colorectal cancer	Colorectal cancer
291	Colorectal cancer	Colorectal cancer
292	Type 1 diabetes	Type 1 diabetes
293	Type 1 diabetes	Type 1 diabetes
294	Multiple sclerosis	Type 1 diabetes
295	Multiple sclerosis	Type 1 diabetes
296	Type 1 diabetes	Type 1 diabetes
297	Type 1 diabetes	Multiple sclerosis
298	Type 1 diabetes	Multiple sclerosis
299	Type 1 diabetes	Type 1 diabetes
300	Type 1 diabetes	Type 1 diabetes
301	Type 1 diabetes	Type 1 diabetes
302	Autism	Autism
303	Autism	Autism
304	Glioma	Myocardial infarction (early onset)
305	Myocardial infarction (early onset)	Glioma
306	Restless legs syndrome	Restless legs syndrome
307	Restless legs syndrome	Restless legs syndrome
308	Psoriasis	AIDS progression
309	Psoriasis	AIDS progression
310	Psoriasis	Drug-induced liver injury (flucloxacillin)
311	Psoriasis	Drug-induced liver injury (flucloxacillin)
312	Psoriasis	Psoriasis
313	Psoriasis	Psoriasis
314	AIDS progression	Psoriasis
315	Drug-induced liver injury (flucloxacillin)	Psoriasis

316	Psoriasis	Psoriasis
317	Psoriasis	Psoriasis
318	Myeloproliferative neoplasms	Crohn's disease
319	Crohn's disease	Myeloproliferative neoplasms
320	Type 2 diabetes	Type 2 diabetes
321	Type 2 diabetes	Type 2 diabetes
322	Multiple sclerosis	Type 1 diabetes
323	Multiple sclerosis	Type 1 diabetes
324	Type 1 diabetes	Type 1 diabetes
325	Type 1 diabetes	Multiple sclerosis
326	Type 1 diabetes	Multiple sclerosis
327	Type 1 diabetes	Type 1 diabetes
328	Type 1 diabetes	Type 1 diabetes
329	Type 1 diabetes	Type 1 diabetes
330	Autism	Autism
331	Autism	Autism
332	Glioma	Myocardial infarction (early onset)
333	Myocardial infarction (early onset)	Glioma
334	Restless legs syndrome	Restless legs syndrome
335	Restless legs syndrome	Restless legs syndrome
336	Psoriasis	AIDS progression
337	Psoriasis	AIDS progression
338	Psoriasis	Drug-induced liver injury (flucloxacillin)
339	Psoriasis	Drug-induced liver injury (flucloxacillin)
340	Psoriasis	Psoriasis
341	Psoriasis	Psoriasis
342	AIDS progression	Psoriasis
343	Drug-induced liver injury (flucloxacillin)	Psoriasis
344	Psoriasis	Psoriasis
345	Psoriasis	Psoriasis
346	AIDS progression	Psoriasis
347	Drug-induced liver injury (flucloxacillin)	Psoriasis
348	Glioma	Inflammatory bowel disease
349	Glioma (high-grade)	Inflammatory bowel

		disease
350	Inflammatory bowel disease	Glioma
351	Inflammatory bowel disease	Glioma (high-grade)
352	Primary biliary cirrhosis	Systemic lupus erythematosus
353	Systemic lupus erythematosus	Primary biliary cirrhosis
354	Celiac disease	Schizophrenia
355	Schizophrenia	Celiac disease
356	Rheumatoid arthritis	Inflammatory bowel disease
357	Inflammatory bowel disease	Rheumatoid arthritis
358	Ulcerative colitis	Ulcerative colitis
359	Ulcerative colitis	Ulcerative colitis
360	Type 2 diabetes	Type 2 diabetes
361	Type 2 diabetes	Type 2 diabetes
362	Type 2 diabetes	Type 2 diabetes and other traits
363	Type 2 diabetes	Type 2 diabetes
364	Type 2 diabetes	Type 2 diabetes
365	Type 2 diabetes	Type 2 diabetes
366	Type 2 diabetes	Type 2 diabetes
367	Type 2 diabetes and other traits	Type 2 diabetes
368	Type 2 diabetes	Type 2 diabetes
369	Type 2 diabetes	Type 2 diabetes
370	Coronary disease	Glioma
371	Coronary disease	Glioma
372	Glioma	Coronary disease
373	Glioma	Coronary disease
374	Rheumatoid arthritis	Ulcerative colitis
375	Ulcerative colitis	Rheumatoid arthritis
376	Ulcerative colitis	Ulcerative colitis
377	Ulcerative colitis	Ulcerative colitis
378	Colorectal cancer	Ulcerative colitis
379	Ulcerative colitis	Colorectal cancer
380	Schizophrenia	Schizophrenia
381	Schizophrenia	Schizophrenia
382	Rheumatoid arthritis	Rheumatoid arthritis
383	Rheumatoid arthritis	Rheumatoid arthritis

384	Rheumatoid arthritis	Rheumatoid arthritis
385	Rheumatoid arthritis	Rheumatoid arthritis
386	Type 1 diabetes	Multiple sclerosis
387	Multiple sclerosis	Type 1 diabetes
388	Multiple sclerosis	Type 1 diabetes
389	Multiple sclerosis	Type 1 diabetes
390	Type 1 diabetes	Multiple sclerosis
391	Type 1 diabetes	Multiple sclerosis
392	Type 1 diabetes	Multiple sclerosis
393	Multiple sclerosis	Type 1 diabetes
394	Crohn's disease	Inflammatory bowel disease
395	Inflammatory bowel disease	Crohn's disease
396	Prostate cancer	Prostate cancer
397	Prostate cancer	Prostate cancer

Appendix A.7 Disease prevalence from RMRS data

Disease / Trait (listed in GWAS)	Prevalence (%) in RMRS
Acute_lymphoblastic_leukemia_	0.0002
AIDS_progression	0.01
Amyotrophic_lateral_sclerosis	0.0006
Asthma	0.29
Atrial_fibrillation	0.0374
Atrial_fibrillation_atrial_fl	0.0374
Autism	0.0004
Biochemical_measures	0.01
Bipolar_disorder	0.19
Black_vs__red_hair_color	0.01
Blond_vs__brown_hair_color	0.01
Body_mass_index	0.01
Breast_cancer	0.017
C_reactive_protein	0.1
Celiac_disease	0.0004
Cholesterol__total	0.01
Chronic_lymphocytic_leukemia	0.00001
Chronic_Obstructive_Pulmonary	0.06
Colorectal_cancer	0.0004
Coronary_disease	0.13
Crohn_s_disease	0.0064
Cutaneous_nevi	0.01
Diastolic_blood_pressure	0.1
Drug_induced_liver_injury__fl	0.01
F_cell_distribution	0.01
Folate_pathway_vitamin_levels	0.1
Follicular_lymphoma	0.01
Freckles	0.01
Glioma	0.0002
Glioma__high_grade_	0.00002
HDL_cholesterol	0.01
Height	0.1
Hematocrit	0.01
Hemoglobin	0.1
Hemoglobin_levels	0.1
Inflammatory_bowel_disease	0.01
Inflammatory_bowel_disease__e	0.01
Intracranial_aneurysm	0.0012

Ischemic_stroke	0.024
Knee_osteoarthritis	0.027
LDL_cholesterol	0.18
Lung_adenocarcinoma	0.00015
Lung_cancer	0.015
Male_pattern_baldness	0.01
Mean_corpuscular_hemoglobin	0.01
Mean_corpuscular_volume	0.01
Melanoma	0.002
Menarche_age_at_onset_	0.1
Multiple_sclerosis	0.0024
Myeloproliferative_neoplasms	0.0014
Myocardial_infarction__early_	0.04
Nicotine_dependence	0.23
Nonsyndromic_cleft_lip_with_o	0.01
Obesity	0.16
Obesity__early_onset_extreme_	0.0016
Obesity_extreme_	0.0016
Obesity_related_traits	0.0016
Other_metabolic_traits	0.01
Otosclerosis	0.0035
Pancreatic_cancer	0.0018
Parkinson_s_disease	0.00025
Plasma_eosinophil_count	0.01
Plasma_level_of_vitamin_B12	0.01
Plasma_levels_of_liver_enzyme	0.01
Primary_biliary_cirrhosis	0.00035
Prostate_cancer	0.0037
Psoriasis	0.04
Pulmonary_function_measures	0.01
Quantitative_traits	0.01
Recombination_rate__females_	0.01
Recombination_rate__males_	0.01
Red_vs_non_red_hair_color	0.5
Restless_legs_syndrome	0.01
Rheumatoid_arthritis	0.01
Schizophrenia	0.067
Serum_iron_concentration	0.01
Serum_markers_of_iron_status	0.01
Skin_sensitivity_to_sun	0.01
Systemic_lupus_erythematosus	0.01

Systolic_blood_pressure	0.01
Testicular_cancer	0.01
Testicular_germ_cell_tumor	0.01
Triglycerides	0.01
Type_1_diabetes	0.05
Type_2_diabetes	0.064
Type_2_diabetes_and_other_tra	0.0006
Ulcerative_colitis	0.0035
Venous_thromboembolism	0.02
Waist_circumference_and_relat	0.01
Weight	0.01

Appendix A.8 Java code for RNOR Subroutine

```
/****** RNOR Algorithm *****/
public Float calculaterNORTrue(NodeList nList, int[] pStates, Node
nodeForRNOR)
{
    Float retValTrue = 0.0f;
    Node n1, n2;
    Float f1,f2,probScoreTrue;
    NodeList subtractList, numeratorList, denominatorList;

    float[] cptRow = {0.0f, 0.0f};
    float [] vecp = {0.0f, 0.0f};

    try {
        if(nList.size() == 1)
        {
            n1 = (Node) nList.get(0);
            vecp = nodeForRNOR.getCPTable(pStates, null);
            if(vecp[0] == 0.5)
            {
                retValTrue = linkProbsTrue.get(n1.getName());
            }
            else
            {
                retValTrue = vecp[0];
            }
            linkProbsTrue.put(nList.toString(),
retValTrue); // also save it for further calculations
            printProbsTrue.put(getParentStateName(pStates),
retValTrue);
        }
        else if(nList.size() == 2)
        {
            n1 = (Node) nList.get(0);
            n2 = (Node) nList.get(1);
            f1 = linkProbsTrue.get(n1.getName());
            f2 = linkProbsTrue.get(n2.getName());

            vecp = nodeForRNOR.getCPTable(pStates, null);
            if(vecp[0] == 0.5)
            {
                retValTrue = 1 - ( (1 - f1)* (1 -
f2)) ;
            }
            else
            {
                retValTrue = vecp[0];
            }

            linkProbsTrue.put(nList.toString(), retValTrue);
            // also save it for further calculations
            printProbsTrue.put(getParentStateName(pStates),
retValTrue);
        }
    }
}
```

```

else
{
    subtractList = new NodeList(tempNet);
    Node nodeToSubtract, nodeToSubtractPlusOne;
    ArrayList<Float> resultProbs = new
    ArrayList<Float>();
    ArrayList<Float> resultProbsTrue = new
    ArrayList<Float>();

    for(int i=0; i< nList.size(); i++)
    {
        nodeToSubtract = nList.getNode(i);
        subtractList.clear();
        // Numerator
        numeratorList = new NodeList(nList);
        subtractList.add((Node) nodeToSubtract);
        setSubtract(numeratorList, subtractList);
        //After the call numeratorSet = nList - subtractList

        // Denominator
        subtractList.clear();
        denominatorList = new NodeList(nList);
        subtractList.add((Node) nodeToSubtract);
        if((nList.size() - i) == 1)
        // Are we at the end of the list, then the plus
        one subtract node needs to be wrapped
        {
            nodeToSubtractPlusOne = nList.getNode(0);
        }
        else
        {
            nodeToSubtractPlusOne =
            nList.getNode(i+1);
        }
        subtractList.add((Node) nodeToSubtractPlusOne);
        setSubtract(denominatorList, subtractList);
        //After the call denominatorSet = nList -
        subtractList

        probScoreTrue =
        calculateTrueProbScore(numeratorList,
        denominatorList, pStates, nodeForRNOR);
        resultProbsTrue.add(probScoreTrue);
    }

    // Now multiply individual scores and subtract from 1
    retValTrue = resultProbsTrue.get(0);
    for(int i=1; i < resultProbsTrue.size(); i++)
    {
        retValTrue = retValTrue*resultProbsTrue.get(i);
    }
    retValTrue = 1 - retValTrue;

    if(retValTrue == 0)

```

```

        {
            retValTrue = 0.0f;
        }
        else if(retValTrue == 1)
        {
            retValTrue = 1.0f;
        }

        //if(retValTrue >= 0.0f && retValTrue <= 1.0f)

    {
        if(!linkProbsTrue.containsKey(nList.toString())) // we store it
        for future calculations if needed
        {
            linkProbsTrue.put(nList.toString(), retValTrue); // also save
            it for further calculations

            printProbsTrue.put(getParentStateName(pStates), retValTrue);
        }
    }
    //else
    if(retValTrue < 0.0f || retValTrue > 1.0f)
    {
        try {
            writerError.write("Combination -VE or +VE for:" +
            getParentStateName(pStates) + ":" + retValTrue.toString());
            writerError.write("\n");

            for(int i=0; i < resultProbsTrue.size(); i++)
            {
                writerError.write("val_" + i + ": " +
                resultProbsTrue.get(i));
                writerError.write("\n");
            }
        } catch (IOException e) {

            // TODO Auto-generated catch block
            try {
                writer.write(e.toString());
            } catch (IOException e1) {
                // TODO Auto-generated catch block
                e1.printStackTrace();
            }

            e.printStackTrace();
        }

    }

} catch (NeticaException e) {
    // TODO Auto-generated catch block
    e.printStackTrace();
}

return retValTrue;
}

```

Appendix A.9 Java code for IsBIG Subroutine

```

public void processRequest (String filename_org, String filename_other,
String filename_trait_prevalance, Double threshold)
{

    logBuf = ""; //resets log

    assocFileName = filename_org;
    outputFileName = filename_other;
    initTraitPrevalance(filename_trait_prevalance);
    partialAssociationElement.setThreshold(threshold);

    try{
        //-- Do Netica stuff
        try {
            if (env == null) {
                env = Environ.getDefaultEnviron();
                if (env == null) {
                    String errMsg = initSession();
                    if (env == null) {
                        System.out.println( errMsg );
                        return; //no point in continuing
                    }
                }
            }
        }
    } catch (Exception e) {
        System.out.println( e.getMessage() );
    }

    totalSNPs = 0;
    totalTraits = 0;
    parseAssocFile(assocFileName);
    if(printDebug)
    {
        System.out.println("/*****
        *****/");
        System.out.println("Total Duplicate Associations Found for input:
        " + myAssociations.getDuplicateAssociation());
        System.out.println("Total Associations Found: " +
        myAssociations.getTotalAssociation());
        Iterator<Map.Entry<String, associationElement>> assocIterator =
        myAssociations.getIterator();
        while(assocIterator.hasNext())
        {
            System.out.println(assocIterator.next().getKey());
        }
        System.out.println("/*****
        *****/");
    }

    drawDAG();
    System.out.println("Writing to File Total SNPs: " + totalSNPs);

    String fn = outputFileName.substring(0,
    outputFileName.indexOf("."));

```

```

Streamer os1 = new Streamer(fn + "_SNP.dne" );
tempNet.write(os1);
triangulate();
prune();

System.out.println("Writing to File Pruned SNPs Network: ");

Streamer os2 = new Streamer(fn + "_PrunedSNP.dne" );
tempNet.write(os2);

Iterator<snp> snpListIterator= snpListByRSquare.iterator();
if(printDebug)
{
    while(snpListIterator.hasNext())
    {
        System.out.println(snpListIterator.next().getName());
    }
    System.out.println("/*****
    *****/");
}
drawDAG_traits();
System.out.println("Total Traits: " + totalTraits);
Iterator<String> traitListIterator= traitList.iterator();
if(printDebug)
{
    while(traitListIterator.hasNext())
    {
        System.out.println(traitListIterator.next());
    }
    System.out.println("/*****
    *****/");
}
removeCyclesBeforeCPT();
computeCPT();
computeCPTofTraits();

boolean compiled = false;
while(!compiled)
{
    try{

        tempNet.compile();
        compiled = true;
    } catch(NeticaException e)
    {
        if(e.getMessage().contains("is a cycle (containing link)")
        {
            String [] msg = e.getMessage().split("->");
            String [] n = msg[0].split("link");
            String [] msgg = msg[1].split("\n");
            String c_name = msgg[0].trim();
            String n2 = c_name.substring(0,c_name.length()-1);
            Node child = (Node) tempNet.getNode(n2);
            Node parent = (Node) tempNet.getNode(n[1].trim());
            removeCycles(parent, child);
        }
        else if(e.getMessage().contains("doesn't have a CPT table"))

```

```

        {
            String [] msg = e.getMessage().split("node");
            String [] n = msg[0].split("rs");
            String [] msgg = msg[1].split("\n");
            String c_name =
                msgg[0].substring(2,msgg[0].indexOf("doesn't")-2);

            Node Odd_n = (Node) tempNet.getNode(c_name);
            int size = Odd_n.getParents().size();
            float[] cptRow = new float [size*2*2];
            for(int i=0; i<cptRow.length; i++)
            {
                cptRow[i] = 0.5f;
            }
            Odd_n.setCPTTable(cptRow);
        }
        else
        {
            System.out.println(e.getMessage());
        }
    }
}

double size = tempNet.sizeCompiled();
System.out.println("Total compiled size: " +
    Double.toString(size));
double memSize = tempNet.getEnviron().getMemoryUsageLimit();
System.out.println("Total memory size: " +
    Double.toString(memSize));
Streamer os = new Streamer(outputFileName);
tempNet.write(os);

Net netToAbsorb = new Net(new Streamer (outputFileName));

//Absorb nodes
NodeList nodes = new NodeList (netToAbsorb);
Iterator<snp> snpNodeListIterator=
snpListByRSquare.iterator();
while(snpNodeListIterator.hasNext())
{
    snp thisSNP = snpNodeListIterator.next();
    String name = thisSNP.getName();

    nodes.add(netToAbsorb.getNode(name));
}
netToAbsorb.absorbNodes (nodes);

Streamer os_mod = new Streamer(fn + "_absorbed.dne");
netToAbsorb.write(os_mod);
netToAbsorb.finalize();

tempNet.finalize();
tempNet = null;
}
catch (Exception e) {

    System.out.println( e.getMessage() );
}
}

```

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CURRICULUM VITAE

Vibha Anand

Education

Ph.D. in Health Informatics (2005 – 2010)

Indiana University, Indianapolis, Indiana, USA

M.S. in Computer Sciences (1992 – 1994)

University of New Haven, West Haven, Connecticut, USA

B.E. in Electrical and Electronics Engineering (1982 – 1986)

Motilal Nehru National Institute of Technology, India

Research, Academic, and Professional Experience

Ph.D. student, Indiana University-Purdue University, Indianapolis (2005 – 2010)

- Develop models for medical decision support using probabilistic methods and data mining
- In-Silico Bayesian Integration of GWA Studies (IsBIG)
- An Empirical Validation of Recursive Noisy OR (RNOR) Rule for Asthma
- Prediction
- In-Silico Testing of Genotype-Phenotype Associations with Electronic Medical Records
- Probabilistic Asthma Case Finding – A Noisy-OR Reformulation
- Probabilistic Asthma Case Finding – A Pilot Study using the CHICA system

Research Associate Faculty, Pediatrics, IU School of Medicine (Oct 2008 – Dec 2010)

Lead Systems Analyst (May 2002 – Oct 2008)

- Direct Children's Health Informatics Research and Development Lab (CHIRDL) for software development. Manage a team of developers.
- Medical informatics integration at point of care, Medical informatics applications, Use of healthcare standards
- Collaborating with multi-disciplinary team in randomized controlled trials using pediatric decision support system (DSS), The Child Health Improvement through Computer Automation system (CHICA)
- Design, development and integration of open source electronic medical record systems (OpenMRS – <http://www.openmrs.org>) into second generation of CHICA system.
- Lead the design and development of the first generation CHICA system for Pediatric care and a DSS for Arthritis use in geriatric care.
- Developed an Arden Syntax parser using open source tool (ANTLR) for translating Arden Syntax Medical logic modules based clinical guideline rules into JAVA code
- Design and development of Decision Support System for Indiana Statewide Newborn Screening Alert System (using INPC network)

Senior Engineer, Hippo Inc., New Haven, CT (Oct 2000 – Mar 2002)

- Designed and developed Voice over IP (VoIP) protocol based software
- Integrated a third party SIP stack in the firmware

Consultant Engineer, Advanced Decisions Inc., Shelton, CT (Jan 1998 – Oct 2000)

- Design and develop software systems for advanced computer telephony interfaces
- Design and develop software systems for business logic in advanced postal meters

Lead Engineer, Executone Information Systems, Milford, CT (Jul 1993 – Jan 1998)

- Design and develop software system for Private Branch Exchanges (PBX)

Senior Engineer, Bull HN Info Systems, Melbourne, Australia, (Jun 1989 – Sep 1992)

- Design and develop software system for Electronic Funds Transfer Point of Sale systems (EFTPOS)

Engineer Altos / UMCL / Uptron India Ltd, India, (Aug 1986 – Mar 1989)

- Design and develop hardware and firmware for personal computers

Memberships

- American Medical Informatics Association (AMIA) – Member of Knowledge Discovery and Data Mining Working Group

Publications

- Anand V, Downs SM (2010) An Empirical Validation of Recursive Noisy-OR (RNOR) Rule for Asthma Prediction, AMIA Annual Fall Symposium 2010, In press
- Downs SM, Anand V, Dugan TM, Carroll AE (2010) You Can Lead a Horse to Water: Physician's Responses to Clinical Reminders, AMIA Annual Fall Symposium 2010, In press
- Anand, V., et al., Tailoring Interface for Spanish Language: A Case Study with CHICA System, in Proceedings of the 1st International Conference on Human Centered Design: Held as Part of HCI International 2009. 2009, Springer-Verlag: San Diego, CA. p. 398-407.
- Anand V, Downs SM (2008) Probabilistic Asthma Case Finding: A Noisy-OR Reformulation, AMIA Annual Fall Symposium 2008,6-10,Published
- Grannis S, Biondich PG, Downs SM, Sheley ME, Anand V, Egg J (2008) Leveraging Open-Source Matching Tools and Health Information Exchange to Improve Newborn Screening Follow-up, Public Health Information Network Conference (PHIN) 2008
- Anand V, Downs SM (2007) Probabilistic Asthma Case Finding - A Pilot Study using the CHICA system. Medinfo, 2007. 12(Pt 1): p. 292,Published

- Downs SM, Biondich PG, Anand V, Zore MM, Carroll AE (2006) Using Arden syntax and adaptive turnaround documents to evaluate clinical guidelines,, AMIA Fall Symposium, AMIA, 214-8, Published
- Downs SM, Carroll AE, Anand V, Biondich PG (2005) Human and system errors, using adaptive turnaround documents to capture data in a busy practice, AMIA Fall Symposium, AMIA, 211-215, Published
- Biondich PG, Downs SM, Anand V, Carroll AE (2005) Automating the recognition and prioritization of needed preventive services: early results from the CHICA system, AMIA Fall Symposium, AMIA, 51-55, Published
- Anand V (2004) The CHICA RuleBuilder: A practical tool to author Arden Syntax, Medinfo, IMIA, Published
- Anand V, Biondich PG, Downs SM (2004) Child Health Improvement through Computer Automation: The CHICA System, MedInfo, IMIA, 187-191, Published
- Biondich PG, Anand V, Downs SM, McDonald CJ (2003) Using Adaptive Turnaround Documents to Electronically Acquire Structured Data in Clinical Settings, AMIA fall symposium, AMIA, 56-60, Published

Skills

Software Design and Development

- Languages - C#, JAVA, 'C', SQL, Assembly, C++, Visual Basic, Visual C++, Pascal, Basic, Prolog, Lisp, VBScript, XML, Description Logic language
- Software Libraries / Framework - MFC, ATL, STL, MVC – Spring Framework
- Source code control – Visual Source Safe, CVS, SCCS, Subversion (SVN)
- Databases - Microsoft SQL server, MySQL, Access, Btrieve
- OS - UNIX tools - SED, AWK, YACC, 'C' shell, K-shell, SCCS, CVS
- Network Communication protocols / tools - HDLC, DDCMP, NETBIOS, TCP/UDP/IP, XNS, Windows RPC Windows Sockets, BSD Sockets, SIP, RTP, TSAPI, CSTA, Dialogic CT-Connect, Cisco ICR. Protocol Analyzer, HP In circuit emulators
- Healthcare standards – HL7, Arden Syntax, LOINC, SNOMED, UMLS, MeSH, DICOM

Modeling

- UML Modeling, ER Diagrams, ANTLR (www.antlr.org) tool
- Bayesian Networks