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Bayesian Network Modeling and Inference of GWAS Catalog

A thesis submitted in partial fulfillment of the requirements for the degree of Master of Science in Computer Science

by

Qiuping Pan Huaqiao University Bachelor of Network Engineering, 2009

> May 2018 University of Arkansas

This thesis is approved for recommendation to the Graduate Council.

Xintao Wu, Ph.D. Thesis Director

Wing Ning Li, Ph.D. Committee member Qinghua Li, Ph.D. Committee member

#### Abstract

Genome-wide association studies (GWASs) have received an increasing attention to understand genotype-phenotype relationships. The Bayesian network has been proposed as a powerful tool for modeling single-nucleotide polymorphism (SNP)-trait associations due to its advantage in addressing the high computational complex and high dimensional problems. Most current works learn the interactions among genotypes and phenotypes from the raw genotype data. However, due to the privacy issue, genotype information is sensitive and should be handled by complying with specific restrictions. In this work, we aim to build Bayesian networks from publicly released GWAS statistics to explicitly reveal the conditional dependency between SNPs and traits.

First, we focus on building a Bayesian network for modeling the SNP-categorical trait relationships. We construct a three-layered Bayesian network explicitly revealing the conditional dependency between SNPs and categorical traits from GWAS statistics. We then formulate inference problems based on the dependency relationship captured in the Bayesian network. Empirical evaluations show the effectiveness of our methods.

Second, we focus on modeling the SNP-quantitative trait relationships. Existing methods in the literature can only deal with categorical traits. We address this limitation by leveraging the Conditional Linear Gaussian (CLG) Bayesian network, which can handle a mixture of discrete and continuous variables. A two-layered CLG Bayesian network is built where the SNPs are represented as discrete variables in one layer and quantitative traits are represented as continuous variables in another layer. Efficient inference methods are then derived based on the constructed network. The experimental results demonstrate the effectiveness of our methods.

Finally, we present STIP, a web-based SNP-trait inference platform capable of a variety of inference tasks, such as trait inference given SNP genotypes and genotype inference given traits. The current version of STIP provides three services which are SNP-trait inference, Top-k trait prediction and GWAS catalog exploration.

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### **List of Published Papers**

[1]. Lu Zhang, Qiuping Pan, Xintao Wu, and Xinhua Shi. Building Bayesian Networks from GWAS statistics based on Independence of Causal Influence. In Bioinformatics and Biomedicine (BIBM), 2016 IEEE International Conference on (2016), IEEE. (Chapter 3)

[2]. Lu Zhang, Qiuping Pan, Xintao Wu, and Xinhua Shi. Bayesian Network Construction and Genotype-phenotype Inference using GWAS Statistics. IEEE/ACM Transactions on Computational Biology and Bioinformatics 99 (2017). (Chapter 3)

[3]. Lu Zhang, Qiuping Pan, Xintao Wu. Modeling SNP and Quantitative Trait Association from GWAS Catalog Using CLG Bayesian Network. In Bioinformatics and Biomedicine (BIBM), 2017 IEEEInternational Conference on (2017), IEEE. (Chapter 4)

[4]. Qiuping Pan, Lu Zhang, and Xintao Wu. Stip: A SNP-Trait Inference Platform. In Bioinformatics and Biomedicine (BIBM), 2017 IEEE International Conference on(2017), IEEE. (Chapter 5)

#### 1. Introduction

#### 1.1. Overview

Genome-wide association studies (GWASs) have received increasing attention due to the rapid decrease of genotyping costs and promising potential in genetic diagnostics, drug development, and personalized medicine. GWAS examines a genome-wide set of genetic variants in different individuals to see if any single nucleotide polymorphism (SNP) is associated with a phenotype/trait. It has been shown that genetic risk factors exist in many common, complex diseases such as schizophrenia and type II diabetes.

High-density genotyping microarrays, and recently next-generation sequencing technologies, have been utilized to identify common genetic variants that predispose an individual to diseases. Genotype data at the individual level are sensitive and thus are usually under controlled access with specific restrictions. For example, HIPAA (Health Insurance Portability and Accountability Act of 1996) provides data privacy and security provisions for protecting medical information in United States. It was shown that only 30-80 out of 30 million SNPs are needed to uniquely identify an individual [31]. Therefore, in addition to the HIPPA privacy rule, the USA Genetic Information Nondiscrimination Act of 2008 (GINA) requires data analyst follow severe privacy rules on acquiring individual's genetic tests and prohibits discrimination against employees or applicants because of genetic information. Hence, genotype profiles of participants are authorized accessible only if the confidentiality agreements are signed.

Meanwhile, in biomedical community, publicly available experimental data is in need because of replication or reanalyses by other researchers. Therefore, most of the GWAS statistics and SNP-trait associations are reported in the genetic literature. To capture such information, the GWAS catalog [57] collects and publicly releases literature-derived GWAS statistics. By extracting information from the literature, the GWAS catalog collects publication information, study cohort information such as sample description, country of recruitment and subject ancestry, and SNP-trait

association information including SNP identifier (i.e. RSID), risk allele frequency, odds ratio, p-value, etc.

Bayesian networks have been proved to be effective in modeling the causal relationships between SNPs and associated traits [12, 24, 60]. Unfortunately, raw data are required in these studies, which are not available in GWAS catalog. To tackle this limitation, some works aim to build Bayesian networks using the GWAS statistics. [15, 16] is employed to capture the joint effect of multiple SNPs on a trait. Consequently, the Bayesian network can be completely specified using the GWAS statistics only. Our previous work in [56] showed that the released GWAS statistics can be used to build a two-layered Bayesian network for inference. The proposed method only uses the statistics released in the GWAS catalog. However, this work suffers from significant limitations. For example, the constructed Bayesian network contains only the nodes representing traits and nodes representing SNP alleles. Thus, it cannot directly characterize the associations between the traits and the genotypes which are the combinations of two alleles. Meanwhile, the orientation of the arcs are pointing from trait nodes to SNP nodes, which contradicts to the fact that in GWAS researchers usually treat traits as the dependent variables and SNPs as the independent variables.

In this thesis, we develop methods to conduct Bayesian networks by utilizing GWAS statistics, which explicitly reveal the conditional dependency between SNPs and traits. Based on the built Bayesian networks, we further conduct SNP-trait inference tasks and study whether and to what extend exploiting the GWAS statistics can be used for inferring private information about a human individual.

First, we propose a Bayesian network to model the SNP and categorical trait associations from the GWAS catalog. The proposed Bayesian network is composed of three layers, the genotype layer, the allele layer, and the trait layer from top to bottom. Edges only go from an upper layer to a lower layer, and all edges among nodes within the same layer are prohibited. We then derive SNP categorical trait inference formulations based on the Bayesian network. The empirical re-

sults show the effectiveness of our proposed Bayesian network and also imply that meaningful private information can be inferred from public GWAS statistics.

Second, we propose a Conditional Linear Gaussian (CLG) Bayesian network to model the SNP and quantitative trait associations from the GWAS catalog. Existing methods in the literature can only deal with categorical traits. We address this limitation by leveraging the CLG Bayesian network, which can handle a mixture of discrete and continuous variables. A two-layered CLG Bayesian network is built where the SNPs are represented as discrete variables and quantitative traits are represented as continuous variables. We empirically evaluate the construction and inference methods and perform a case study to evaluate how much individual information would be disclosed using the constructed network. The results demonstrate the effectiveness of our methods.

Finally, we develop a web-based SNP-trait inference platform, called STIP. STIP is capable of a variety of inference tasks, such as trait inference given SNP genotypes and genotype inference given traits. The core of STIP is two Bayesian networks which model the SNP-categorical trait associations and SNP-quantitative trait associations, respectively. The inference tasks are based on the dependency relationship captured in the Bayesian networks. The current version of STIP provides three services which are SNP-trait inference, Top-k trait prediction and GWAS catalog exploration.

#### **1.2.** Contributions

The contributions of this thesis are as follows.

First, we build a classic three-layered Bayesian network from the GWAS statistics to explicitly reveal the conditional dependency between SNPs and categorical traits. The constructed Bayesian network can be used to conduct general SNP-categorical trait inference tasks.

Second, we address the problem that the classic Bayesian network can only deal with categor-

ical traits and propose a Conditional Linear Gaussian (CLG) Bayesian network to handle the quantitative traits. We build a two-layered CLG Bayesian network where SNPs are represented as discrete variables and quantitative traits are represented as continuous variables. The efficient SNP-quantitative trait inference methods are then derived based on the CLG Bayesian network.

Third, based on the dependency relationship captured in the Bayesian networks, we demonstrate the possibility of privacy breaching of individuals who are even not participants of a GWAS with only using GWAS statistics, and appropriate privacy protection mechanisms need to be developed to protect genetic privacy not only of GWAS participants but also regular individuals.

Finally, we develop STIP, a web-based platform that aims to aid common users in SNP-trait inference based on the proposed Bayesian networks and GWAS catalog exploration.

#### 1.3. Organization of the Thesis

The rest of this thesis is organized as follows. Chapter 2 briefly reviews the GWAS and GWAS catalog, and then introduces the Bayesian networks adopted in this thesis. Chapter 3 introduces a three-layered Bayesian network to model the associations between SNPs and categorical traits. Chapter 4 presents a Conditional Linear Gaussian (CLG) Bayesian network to model the SNP-quantitative trait associations. Chapter 6 concludes the paper and describes the future work.

#### 2. Background

#### 2.1. GWAS and GWAS Catalog

#### 2.1.1. Genome-Wide Association Study (GWAS)

In genetics, a genome-wide association study (GWAS or GWA study), also known as whole genome association study, is to examine the differences of genetic variants between two groups of people to identify the genetic risk factors that associated with a trait. Typically, GWA studies focus on associations between single nucleotide polymorphisms (SNPs) and major human diseases such as obesity, diabetes, and Parkinson's disease. SNP is the unit of genetic variation. In genetic studies, SNPs are considered as makers of a genomic region, with a small proportion of them having a great impact on common, complex diseases. Each SNP typically has two alleles, and each allele is assigned a value from the set {A, C, G, T}. The less common allele in the whole population is named as minor allele. In contrast, the more common allele is major allele. Meanwhile, a minor and a major allele frequency are assigned to each SNP. Furthermore, each individual carries a pair of alleles inherited from both parents and the genotype refers to the two alleles an individual has for a particular SNP. The genotype that contains two major alleles is homozygous minor, and the genotype that contains two minor alleles is homozygous.

All participants in a GWAS are genotyped for assaying SNPs using chip-based microarray technology. Two primary platforms used in most GWASs are Illumina and Affymetrix. Depending on different genotyping platforms, the number of SNPs may be varied from tens of thousands to tens of millions. For both categorical traits and quantitative traits, the association test between one single SNP and the trait is necessary but differing in methods. To be specific, contingency table methods or logistic regression are generally used to analyze Dichotomous case/control traits, whereas quantitative traits are generally analyzed using generalized linear model (GLM)[3].

#### • GWAS on categorical traits

For categorical traits, GWASs are usually conducted in a case-control setting, where cases are individuals with the trait under investigation and controls are matched individuals without the trait. Each individual is genotyped by microarrays or sequencing platforms. Dependent on genotyping platform, the number of SNPs genotyped in a GWAS setting typically ranges from tens of thousands to tens of millions. In a GWAS framework, we assume we study biallelic SNPs. Each biallelic SNP has two possible nucleotide variations in this base position, referred to as alleles (e.g., A/G). The allele that is more frequent in the case group comparing with the control group is called the risk allele (e.g., A), and the other one is called the non-risk allele (e.g., G). Each individual carries a pair of alleles inherited from both parents and the genotype refers to the two alleles an individual has for a particular SNP. The genotype that contains two risk alleles is called the homozygote for risk allele (e.g., AA), the genotype that contains two non-risk alleles is called the homozygote for non-risk allele (e.g., GG), and the genotype that contains one risk allele and one non-risk allele is called the heterozygote (e.g., AG).

Table 2.1: The genotype frequency

Table 2.2:	The allele	frequency
------------	------------	-----------

	AA	AG	GG	Total		A	G	Total
Cases	$r_0$	$r_1$	$r_2$	R	Cases	$2r_0 + r_1$	$r_1 + 2r_2$	2R
Controls	$s_0$	$s_1$	$s_2$	S	Controls	$2s_0 + s_1$	$s_1 + 2s_2$	2 <i>S</i>
Total	$n_0$	$n_1$	$n_2$	N	Total	$2n_0+n_1$	$n_1 + 2n_2$	2N

A GWAS is then to assess the difference of the frequency of alleles in the case and control groups. The typical process of a GWAS is described as follows. First, a genotype profile dataset is generated by genotyping the individuals in the case group and the control group. For each SNP, the genotype frequency is counted over the two groups to obtain a  $3 \times 2$  contingency table, as shown in Table 2.1. Here,  $r_0$  denotes the number of individuals in the case group with genotype AA and so forth. Then, the genotype frequency is transformed into the allele frequency represented by a  $2 \times 2$  contingency table as shown in Table 2.2. To be specific, each homozygote for risk/non-risk allele is counted as 2 copies of risk/non-risk alleles, and each heterozygote is counted as 1 risk

allele and 1 non-risk allele. After that, statistical tests such as chi-square test and Fisher's exact test, are performed on the allele contingency table to investigate whether there is an association between the SNP and the trait. In addition to a *p*-value indicating the significance of the association, the GWAS also reports odds ratios that measure the difference of frequency of an allele in the case versus control group. Specifically, the odds ratio is defined as the ratio between the proportion of individuals with a specific allele in the case group, and the proportion of individuals with the same allele in the control group. If the odds ratio is larger than 1, it indicates that the risk allele is more frequent in the case group than it is in the control group. Finally, the trait and its significantly associated SNPs are reported, along with the risk allele type and corresponding statistics (odds ratios, *p*-values, etc.).

#### • GWAS on quantitative traits

For quantitative traits, association tests are performed using the generalized linear model approaches. The allele that is positively associated with an increase in the trait is called the risk allele, and the other one is called the non-risk allele.

For a trait T and a SNP S, the association test basically asks to fit the linear regression

$$t = \beta_0 + \beta s + \varepsilon,$$

where *t* denotes the value of *T* and  $s = \{0, 1, 2\}$  represents the genotype of *S*. Parameters  $\beta_0$  and  $\beta$  are estimated using the least squares estimation. The statistical power of the association test is obtained using the Analysis of Variance (ANOVA). Specifically, consider the *n* individuals involved in the study, where the mean of *T* of all individuals is given by  $\bar{t}$ . For a particular individual *k*, denote his value of trait *T* by  $t_k$ , and denote his predicted value of *T* by  $\hat{t}_k$ . Then, the regression sum of squares (*SSR*) is defined as:

$$SSR = \sum_{k=1}^{n} \left( \hat{t}_k - \bar{t} \right)^2, \qquad (2.1)$$

Trait	SNP-risk allele	RAF in control	p-value	Odds ratio	Sample description	
Prostate cancer	rs4793529-T	0.22	2.00E-13	0.28	1146 European ancestry individuals	
Prostate cancer	rs1447295-A	0.71	6.00E-18	0.51	1146 European ancestry individuals	

Figure 2.1: Examples of categorical traits in the GWAS catalog.

and the error sum of squares (SSE) is defined as:

$$SSE = \sum_{k=1}^{n} (t_k - \hat{t}_k)^2.$$
 (2.2)

The ratio  $F^* = \frac{SSR/1}{SSE/(n-2)}$  is used to test hypotheses  $H_0 : \beta = 0$  versus  $H_a : \beta \neq 0$ . Since  $F^*$  is known to follow the *F* distribution with 1 numerator degree of freedom and n - 2 denominator degrees of freedom, i.e.,  $F^* \sim F(1, n - 2)$ , the *p*-value can be determined through an *F*-test.

#### 2.1.2. GWAS Catalog

For both quantitative and dichotomous trait analysis, SNP-trait associations and related statistics are published in genetic literature. The GWAS catalog[33] manually collects all published genome-wide association studies with assaying more than 100,000 SNPs and all SNP-trait associations with p-values  $< 1.0 \times 10^{-5}$ . The GWAS catalog data can be downloaded as a tsv file from its official website <sup>1</sup>. The latest version of GWAS catalog, which is released at July 4, 2017, contains 3020 publications and 37868 distinct SNP-trait associations. To be specific, there are 2014 traits and 31939 associated SNPs. For categorical (quantitative) traits, each record of the file indicates an association from a study, which contains the information like the trait, the SNP, the risk allele, risk allele frequency in controls, *p*-value, and odds ratio, as shown in Figure 2.1. For quantitative traits, each record contains the trait, the SNP, the risk allele, the sample size, the coefficient  $\beta$  and *p*-value of F-test, as shown in Figure 2.2.

<sup>&</sup>lt;sup>1</sup>https://www.ebi.ac.uk/gwas/

Trait	SNP-risk allele	Beta	p-value	Sample description	
Body mass index	rs7708584-A	0.021	5.00E-14	37956 African American individuals	
Mean platelet volume	rs7896518-A	5.18	2.00E-12	16388 African Amercan individuals	
Fibrinogen level	rs1976714-T	0.006	2.00E-08	120246 European ancestry individuals	

Figure 2.2: Examples of quantitative traits in the GWAS catalog.

#### 2.2. Bayesian Network

Bayesian networks are widely used for reasoning under uncertainty and its representation rigorously describes probabilistic relationships among variables of interest [9, 14, 22]. A Bayesian network G = (V, E) is a Directed Acyclic Graph (DAG), where the vertices (or nodes) in V corresponding to the variables and the edges (or links) in E represent the dependence relationships among the variables. The dependence/independence relationships are graphically encoded by the presence or absence of direct connections between pairs of variables. Hence a Bayesian network shows the (in)dependencies between the variables qualitatively, by means of the edges, and quantitatively, by means of conditional probability distributions which specify the relationships. In general, a Bayesian network represents the joint probability distribution by specifying a set of conditional independence assumptions together with sets of local conditional probabilities. An edge in the network represents the assertion that an variable is conditionally independent of its nondescendants in the network given its immediate predecessors. A conditional probability table is given for each variable, describing the probability distribution for that variable given the values of its immediate predecessors. Formally, for each variable  $X_i \in V$ , we have a family of conditional probability distributions  $P(X_i | Par(X_i))$ , where  $Par(X_i)$  represents the parent set of the variable  $X_i$  in G. From these conditional distributions we can compute the joint probability for any desired assignment of values  $\langle x_1, x_2, \cdots, x_n \rangle$  to the tuple of network variables  $X_1, X_2, \cdots, X_n$ by the formula:

$$P(x_1, x_2, \cdots, x_n) = \prod_{i=1}^n P(x_i | Par(X_i))$$
(2.3)

Note the values of  $P(x_i|Par(X_i))$  are precisely the values stored in the conditional probability table associated with variable  $X_i$ . Bayesian networks can be used to perform efficiently reasoning

tasks. There are several algorithms (including exact inference methods and approximate inference methods) to compute the posterior probability for any variable given the observed values of the other variables in the graph [23].

#### 2.2.1. Independence of Causal Influence

We describe the models of independence of causal influence that are widely used in building a Bayesian network. Consider a set of independent variables  $\mathbf{A} = \{A_1, \dots, A_m\}$  and a dependent variable *C*. In our context, we assume *C* is a binary variable. The CPT  $P(C|\mathbf{A})$  that exhibits ICI is defined as follows. First, each independent variable  $A_j$  is connected with a hidden variable  $X_j$ , which represents the "effective value" of  $A_j$  on *C*. The connection between  $A_j$  and  $X_j$  can be defined via various stochastic or deterministic functions. Then, the resulting hidden variables  $X_j$ s are combined using certain deterministic function  $f(\cdot)$ . Usually, in order to be a decomposable function,  $f(\cdot)$  is required to be associative and commutative. Besides, an additional hidden variable  $X_0$  is added to represent background knowledge, resulting a combination function  $X = f(X_0, X_1, \dots, X_m)$ . Finally, another stochastic or deterministic function is applied to X to obtain the value of *C*. The structure of the general formulation of the ICI models is shown in Figure 2.3. In general, learning an ICI model requires the raw data for estimating parameters in the presence of hidden variables.

In the following, we introduce the Noisy-Or model, one best known example of the ICI models. The Noisy-Or model can be considered as a generalization of the deterministic Or relation since it is an ICI model where the combination function is the Or function. In this model, each hidden variable  $X_j$  is a binary variable taking values of 0 and 1. The connection between each pair of  $A_j$ and  $X_j$  is defined as the following probabilistic distribution:

for each j, 
$$P(X_j = 0 | A_j = a_j) = \begin{cases} 1 & \text{if } a_j = 0, \\ \theta_j(a_j) & \text{otherwise}. \end{cases}$$



Figure 2.3: The ICI model

where  $\theta_j(a_j)$  is called the noise parameter representing the probability that the presence of  $A_j$ (i.e.,  $A_j \neq 0$ ) would be effective if the occurrence of *C* is true (i.e., C = 1). It is also defined that

$$P(X_0 = 0) = \theta_0,$$

which is called a leak probability that allows *C* to occur when all the  $A_j$ s are absent. Then,  $f(\cdot)$  is defined as the deterministic Or function that takes all  $X_j$ s as the input, i.e.,

$$f(X_0 = x_0, X_1 = x_1, \cdots, X_m = x_m) = x_0 \lor x_1 \lor \cdots \lor x_m.$$

Finally, *C* directly takes the value of the output of  $f(\cdot)$ . Straightforwardly, *C* equals 0 if and only if all  $X_j$ s take the value of 0. Thus, the probability of C = 0 given  $\mathbf{A} = \mathbf{a}$  is calculated by

$$P(C = 0 | \mathbf{A} = \mathbf{a}) = P(X_0 = 0) \prod_{j:a_j \neq 0} P(X_j = 0 | A_j = a_j)$$
$$= \theta_0 \prod_{j:a_j \neq 0} \theta_j(a_j).$$

By defining an indicator function

$$\mathbb{1}(a_j) = \begin{cases} 0 & \text{if } a_j = 0\\ 1 & \text{otherwise} \end{cases}$$

the above probability can be rewritten more compactly as

$$P(C = 0 | \mathbf{A} = \mathbf{a}) = \theta_0 \prod_{j=1}^m \theta_j(a_j)^{\mathbb{1}(a_j)}.$$
 (2.4)

To learn the Noisy-Or model, assume that we are given a dataset  $\mathcal{D} = \{\cdots, \mathbf{d}^l, \cdots\}$ , where each tuple  $\mathbf{d}^l = \{c^l, \mathbf{a}^l\}$  represents the values of *C* and **A**. The objective function is typically formalized as maximizing the log-likelihood of the model given the observed data, i.e.,  $\sum_{l=1}^{|\mathcal{D}|} \log P(\{C, \mathbf{A}\} = \mathbf{d}^l)$ . Following the procedure in [54], the Noisy-Or model can be learned using an EM algorithm [38]. The EM algorithm with the derived formulas is described as below:

1) The E-step is to compute the expected marginal count  $n(X_0 = 0)$  and  $n(X_j = 0, A_j = a_j)$  $(a_j \neq 0)$  given data  $\mathcal{D}$ :

$$n(X_0 = 0) = \sum_{l=1}^{|\mathcal{D}|} P(X_0 = 0 | \mathbf{d}^l),$$
  
$$n(X_j = 0, A_j = a_j) = \sum_{l=1}^{|\mathcal{D}|} P(X_j = 0, A_j = a_j | \mathbf{d}^l),$$

and for each tuple  $\mathbf{d}^l$ ,

$$P(X_j = 0, A_j = a_j | \mathbf{d}^l) = \begin{cases} P(X_j = 0 | \mathbf{d}^l) & \text{if } a_j^l = a_j \\ 0 & \text{otherwise.} \end{cases}$$

The above updated probabilities are computed as follows.

$$P(X_0 = 0 | \mathbf{d}^l) = \begin{cases} 1 & \text{if } c^l = 0, \\ \frac{1}{z} \cdot \left( \hat{\theta}_0 - \hat{\theta}_0 \prod_{i=1}^m \hat{\theta}_i(a_i^l)^{\mathbb{1}(a_i^l)} \right) & \text{if } c^l = 1, \end{cases}$$

$$P(X_{j} = 0|\mathbf{d}^{l}) = \begin{cases} 1 & \text{if } c^{l} = 0, \\ \frac{1}{z} \cdot \left(\hat{\theta}_{j}(a_{j}^{l}) - \hat{\theta}_{0} \prod_{i=1}^{m} \hat{\theta}_{i}(a_{i}^{l})^{\mathbb{1}(a_{i}^{l})} \right) & \text{if } c^{l} = 1, \end{cases}$$

where  $z = 1 - \hat{\theta}_0 \prod_{i=1}^m \hat{\theta}_i (a_i^l)^{\mathbb{1}(a_i^l)}$  is the normalization constant.

2) For the M-step, the parameters are updated as follows.

$$\hat{\theta}_0^* = \frac{n(X_0 = 0)}{n}$$
 and  $\hat{\theta}_j^*(a_j) = \frac{n(X_j = 0, A_j = a_j)}{n(A_j = a_j)}$ .

#### 2.2.2. CLG Bayesian Network

Traditional Bayesian network can only deal with the situations where the variables are all discrete or all continuous. To tackle this limitation, the CLG Bayesian network [27] is proposed to handle the mixture of discrete and continuous variables. In a CLG Bayesian network, all variables X are partitioned into a set of discrete variables  $X_{\Delta}$  and a set of continuous variables  $X_{\Gamma}$ . For discrete variables, it assumes that they only have discrete parents. A conditional probability distribution P(x|pa(X)) is defined for each discrete variable  $X \in X_{\Delta}$  to specify the conditional probability of X = x given all its parents pa(X). For each continuous variable  $X \in X_{\Gamma}$ , a CLG distribution P(x|pa(X)) is defined conditional on each value assignment of all its parents pa(X). Denote the discrete parents by  $pa_{\Delta}(X) \subseteq X_{\Delta}$ , and the continuous parents by  $pa_{\Gamma}(X) \subseteq X_{\Gamma}$ . The CLG distribution of *X* is specified as

$$P(x|pa(X)) = P(x|pa_{\Delta}(X) = \mathbf{i}, pa_{\Gamma}(X) = \mathbf{z})$$
  
=  $\mathcal{N}(x; a(\mathbf{i}) + b(\mathbf{i})^{\mathsf{T}}\mathbf{z}, c(\mathbf{i})),$  (2.5)

where  $a(\cdot)$  is a table of real numbers (one for each value assignment **i**),  $b(\cdot)$  is a table of  $|pa_{\Gamma}(X)|$ dimensional vectors (one for each value assignment **i**), and  $c(\cdot)$  is a table of non-negative real numbers (one for each value assignment **i**). Equation (2.5) shows that P(x|pa(X)) is a Gaussian distribution with mean  $\mu_{x|pa(X)} = a(\mathbf{i}) + b(\mathbf{i})^{\mathsf{T}}\mathbf{z}$  and variance  $\sigma_{x|pa(X)}^2 = c(\mathbf{i})$ .

Based on above definitions, the joint distribution over all variables in X is given by

$$P(X) = \prod_{X \in X_{\Delta}} P(x|pa(X)) \cdot \prod_{X \in X_{\Gamma}} P(x|pa(X)).$$
(2.6)

For any subset of variables  $\mathbf{X} \subseteq \mathcal{X}$ , the marginal distribution over  $\mathbf{X}$  is given by

$$P(\mathbf{x}) = \sum_{X_{\Delta} \setminus \mathbf{X}} \prod_{X \in X_{\Delta}} P(x|pa(X)) \cdot \int_{X_{\Gamma} \setminus \mathbf{X}} \prod_{X \in X_{\Gamma}} P(x|pa(X)) dx.$$
(2.7)

#### 3. Modeling SNP and Categorical Trait Association Using Bayesian Network

#### 3.1. Introduction

Genome-wide association studies (GWASs) have received intensive attention due to the rapid decrease of genotyping costs and promising potential in genetic diagnostics. GWASs typically focus on associations between single-nucleotide polymorphisms (SNPs) and human traits including common diseases. It has been shown that many common diseases such as various cancer types, have genetic disposition factors.

Facilitated by GWAS, modeling of SNP-trait associations using machine learning and data mining methods has been studied to aid the understanding of the interactions among genotypes and phenotypes. In particular, the Bayesian network has been proposed as a powerful tool for modeling SNP-trait associations due to its advantage in addressing the high computational complex and high dimensional problems [12, 24, 60]. However, most of the existing methods require raw genotypes of SNPs and such information is not publically available. This is because genotype data is usually classified as sensitive and should be handled by complying with specific restrictions.

On the other hand, the experimental results are publicly available so that the data can be compared with other studies or reanalyzed by other researchers. As a result, most of the GWAS statistics and SNP-trait associations are publicly accessible. The GWAS catalog [17, 57] is a database that collects and publicly releases literature-derived GWAS statistics, including pair-wise SNPtrait associations, risk allele frequency,  $\beta$ , *p*-value, etc.

In this chapter, we propose to construct a Bayesian network explicitly revealing the conditional dependency between SNPs and categorical traits from the GWAS statistics. In order to utilize the GWAS statistics, the constructed network is composed of three layers, the genotype layer, the allele layer, and the trait layer. Edges only go from an upper layer to a lower layer, and all edges among nodes within the same layer are prohibited.

The key challenge in specifying the Bayesian network is that, when the dependent variable (i.e., trait) has associations with multiple independent variables (i.e., SNPs), the Bayesian network needs to specify the conditional probability table (CPT) of the trait conditional on every value combination of its associated SNPs. However, GWAS statistics only provide the information for each trait-SNP association pair. The information about epistatic interactions among multiple SNPs that bring about joint effect on a trait is rather limited. Additionally, complex traits are commonly associated with many SNPs. Therefore, it is a combinatorial problem for specifying CPTs because the number of the conditional probability distribution values in the CPT is exponential to the number of SNPs associated with a trait.

To deal with this issue, we propose to adopt the models of independence of causal influences (ICI), a family of models which are widely used in building Bayesian networks [15, 16]. The ICI models assume that, when there are multiple parent variables, the causal mechanism of each parent variable is mutually independent. Hence, the combined influence of multiple parents is decomposable into a series of independent influence of each parent variable. Thus, an ICI model enables us to specify the CPT of a variable given its parents in terms of an associative and commutative operator on the contribution of each parent. The learning process of an ICI model generally requires raw data in order to find the parameters that make the model fit the data best [40, 54]. In this study, we investigate a scenario that the raw data (genotypes) are unknown and only GWAS statistics are available. This makes it challenging to build an ICI model for constructing a Bayesian network from only statistics. In order to do this, we derive a formulation based on the Noisy-Or model [26], one best known example of the ICI models, that can be used to specify the CPT from the released GWAS statistics where the underlying genotypes can be unknown. We prove that, the specified CPT is accurate as long as the individual-level genotype profile follows the Noisy-Or model. Then, we empirically evaluate the fitness of the Noisy-Or model to validate the proposed method.

As applications of the constructed Bayesian network, we propose three inference problems: 1)

trait inference given SNP genotype that aims to infer the probability of a target developing certain traits when the target's genotype profile is given; 2) genotype inference given trait that aims to infer the probability of a target having a certain genotype profile when some traits of the target are given; and 3) trait inference given trait that aims to infer the probability of having a new trait given known traits of the target. We study efficient inference methods to solve these problems using the reconstructed Bayesian network. To evaluate the derived inference methods, we simulate three scenarios. In the first scenario, we assume that an individual has taken a genetic test and wants to infer his/her probability of having some sensitive trait (e.g., disease) based on the genotype profile. For example, companies like Family Tree DNA, 23andMe, and Ancestry offer genotyping and analyzing service for various SNPs and traits. In the second scenario, we assume that an attacker such as an outsider has access to an anonymized genotype profile database which contains the target individual's record and aims to identify the target individual's record from the anonymized dataset. In the third scenario, we also assume the attacker knows some traits of the target individual. The attacker aims to derive new traits. For example, private traits and attributes of individuals can be predictable from easily accessible digital records of behavior such as Facebook Likes [28]. Other patient social networks and online communities like 'patientlikeme.com' provide a platform for users (mostly patients) to connect with others who have the same disease or condition and share their own experiences. Online publishing platform such as openSNP [10] also allows customers to share and publish their genotype and phenotype profiles. We evaluate how the derived inference methods perform in these scenarios.

The contributions of our study are as follows. 1) We apply the classic Bayesian network approach [9, 14, 22] to build a three-layered Bayesian network from the released GWAS statistics. The constructed Bayesian network explicitly reveals the conditional dependency between SNPs and traits, and can be used to compute the probability distribution for any subset of network variables given the values or distributions for any subset of the remaining variables. 2) We formulate three inference problems based on the dependency relationship captured in the Bayesian network and develop efficient formulas and algorithms to infer the posterior probabilities. 3) We conduct em-

pirical evaluations and the results show the effectiveness of our proposed methods, implying that meaningful private information can be inferred from public GWAS statistics on both participants and non-participants of GWAS. Our results imply that privacy protection mechanisms may need to be developed to protect genetic privacy of both GWAS participants and the general population.

#### 3.2. Learning Bayesian Network from GWAS Statistics

In this section, we elaborate how to build a three-layered Bayesian network. In general, we extract summary statistics of risk alleles from the GWAS catalog [57], build a three-layered Bayesian network from the aforementioned GWAS catalog, and prove the derived formula based on the Noisy-Or model for constructing a Bayesian network from GWAS statistics. The constructed Bayesian network, which explicitly captures the conditional dependency between SNPs and their associated traits, will be used as background knowledge for inference.

#### 3.2.1. Knowledge from GWAS Catalog

We use the information publicly available from the GWAS catalog [57] to construct the Bayesian network. As illustrated in Figure 2.1, such information includes trait/disease name, the associated SNPs and corresponding risk allele type, the risk allele frequency in control group, and statistics (e.g., odds ratio and p-value) in the association test of each SNP. Specifically, we extract the following data from the GWAS catalog: a trait set  $\mathcal{T}$ , which contains *m* traits, and an SNP set S, which contains *n* SNPs. For each specific trait  $T_k \in \mathcal{T}$ , we have a subset of associated SNPs  $\mathbf{S}_k$ . For each associated SNP  $S_{kj} \in \mathbf{S}_k$ , we can extract its corresponding risk allele type  $(r_{kj})$  associated trait  $T_k$ , the odds ratio  $O_{kj}$  of the association test, and the risk allele frequency in the control group  $f_{kj}^t(r)$ .

Though not directly given in the GWAS catalog, the risk allele frequency in the case group can be derived from the corresponding odds ratio and the risk allele frequency in the control group. For

an SNP  $S_{kj}$  associated with a trait  $T_k$ , its odds ratio is

$$O_{kj} = \frac{f_{kj}^c(r)(1 - f_{kj}^t(r))}{f_{kj}^t(r)(1 - f_{kj}^c(r))}.$$
(3.1)

With the released values of the odds ratio  $(O_{kj})$  and the risk allele frequency in the control group  $f_{kj}^t(r)$ , the risk allele frequency in the case group  $f_{kj}^c(r)$  can be derived as

$$f_{kj}^{c}(r) = \frac{O_{kj} \cdot f_{kj}^{t}(r)}{O_{kj} \cdot f_{kj}^{t}(r) + 1 - f_{kj}^{t}(r)}.$$
(3.2)

In summary, the background knowledge that an attacker can obtain from the GWAS catalog [57] includes: a trait set  $\mathcal{T}$ , an SNP set  $\mathcal{S}$ , the risk allele type  $(r_{kj})$ , the odds ratio  $O_{kj}$ , and the risk allele frequency in the control group  $f_{kj}^t(r)$  and in the case group  $f_{kj}^c(r)$  for each pair of trait and its associated SNPs.

#### 3.2.2. Three-layered Bayesian Network Construction

To construct a Bayesian network to represent the conditional dependencies between traits and SNPs, we treat each trait  $T_k \in \mathcal{T}$  as a binary random variable taking values in the set {1, 0}. Here, value 1 stands for the presence of the trait of a participant and value 0 stands for the absence. For each SNP  $S_j \in S$ , its allele and genotype are represented as two different random variables. We denote  $S_j$ 's allele by  $S_j^a$  taking values in {1, 0}, where 1 stands for that the SNP has the risk allele and 0 otherwise; denote  $S_j$ 's genotype by  $S_j^g$  taking values in {0, 1, 2}, where 0 represents the homozygote for non-risk allele, 2 represents the homozygote for risk allele, and 1 represents the heterozygote. Similarly, for a set of SNPs **S**, the set of their alleles are denoted by **S**<sup>a</sup>, and the set of their genotypes are denoted by **S**<sup>g</sup>.

We construct the Bayesian network with background knowledge shown in Section 2.2.1. The constructed network is composed of three layers, from top to bottom, the SNP genotype layer, the SNP allele layer, and the trait layer, based on the procedure of GWAS. Edges only go from an



Figure 3.1: A three-layered Bayesian network of traits and associated SNPs

upper layer to a lower layer, as shown in Figure 3.1. For each SNP  $S_j$ , two nodes  $S_j^g$  and  $S_j^a$  are at the top two layers respectively to denote its genotype and allele. The edge is pointing from  $S_j^g$ to  $S_j^a$  to represent the transformation of the genotype frequency to the allele frequency. For each trait  $T_k$ , there is a node at the bottom level of the network. If an SNP  $S_{kj}$  is associated with a trait  $T_k$  in the GWAS catalog, then an edge is added pointing from  $S_{jk}^a$  to  $T_k$  to represent this SNP-trait pair. Under the context of GWAS catalog analysis, we cannot acquire the SNP-SNP correlation or the trait-trait association. Thus, we prohibit the edges among SNP genotype nodes, the edges among SNP allele nodes, and the edges among trait nodes.

The next step to completely specify a Bayesian network is to determine the CPT stored at each node. We aim to accomplish all specifications by using only the background knowledge obtained from the GWAS catalog plus some prior information. Firstly, we need to acquire the prior probability  $P(S_j^g)$  of each SNP genotype  $S_j^g$  at the top level of the network. Since the comprehensive knowledge of the frequency of every SNP in a population is limited, we first estimate the allele prior probability  $P(S_j^a)$ , and then estimate  $P(S_j^g)$  using the Hardy-Weinberg principle [6]. It is straightforward to estimate  $P(S_j^a)$  as follows.

$$P(S_i^a = s_j) = P(S_i^a = s_j | T = 0)P(T = 0) + P(S_j^a = s_j | T = 1)P(T = 1).$$

By the Hardy-Weinberg principle,  $P(S_j^g)$  is estimated as

$$P(S_{j}^{g} = s_{j}) = \begin{cases} P(S_{j}^{a} = 1)^{2} & s_{j} = 2, \\ P(S_{j}^{a} = 0)^{2} & s_{j} = 0, \\ 2P(S_{j}^{a} = 1)P(S_{j}^{a} = 0) & s_{j} = 1. \end{cases}$$

Secondly, we need to specify the conditional probability  $P(S_j^a|S_j^g)$  for each SNP, which represents how the genotype frequency is transformed into the allele frequency in GWAS. For the typical procedure as shown in Section 2.1.1, we can directly define  $P(S_j^a|S_j^g)$  as

$$P(S_{j}^{a} = s_{1}|S_{j}^{g} = s_{2}) = \begin{cases} 1 & 2s_{1} = s_{2}, \\ 0.5 & s_{2} = 1, \\ 0 & \text{otherwise.} \end{cases}$$
(3.3)

Note that  $P(S_j^a|S_j^g)$  typically represents the assumption of the genetic effect in the data. The definition in Equation (3.3) is known as the additive model, which means that 2 copies of risk alleles impose twice genetic effect of a single risk allele on the trait. Our model can be easily extend to represent other assumptions. For example, to represent the dominant model where having one or more risk alleles imposes the same increased risk compared to the homozygote for non-risk allele, we can transform the heterozygote completely into the risk allele in Equation (3.3).

Finally, we need to specify the CPT of each trait  $T_k$  given its associated SNPs  $S_k$  which represents the SNP-trait association. It is challenging to estimate the combined effect of multiple independent variables on a dependent variable, especially when the raw data is not available. We compute  $P(T_k = 0 | S_k^a = s^a)$  as given by Equation (3.4) which is derived from the Noisy-Or model presented in the Section 3.2.3. We prove that, the computation in Equation (3.4) is accurate as long as the genotype profile follows the Noisy-Or model.

$$P(T_{k} = 0 | \mathbf{S}_{k}^{a} = \mathbf{s}^{a}) = \frac{P(T_{k} = 0) \prod_{S_{kj} \in \mathbf{S}_{k}} P(S_{kj}^{a} = s_{j}^{a} | T = 0)}{\prod_{S_{kj} \in \mathbf{S}_{k}} \sum_{S_{kj}^{g}} P(s_{kj}^{g}) P(s_{kj}^{a} | s_{kj}^{g})}.$$
(3.4)

As can be seen, the knowledge required for accomplish all above specifications only includes: 1) conditional probability  $P(S^a|T)$ , and 2) prior probability P(T). The former can be estimated from the allele frequencies  $f^t(\cdot)$  and  $f^c(\cdot)$  according to the maximum likelihood estimate, and the latter can be acquired from literature or internet.

#### 3.2.3. Modeling SNP-Trait Associations

This subsection derives the CPT specification formulation shown in Equation (3.4). Specifically, given a trait *T* and its associated SNP S, we assume that a Noisy-Or model holds for conditional probability of *T* given S's genotype  $S^g$ , i.e.,  $P(T|S^g)$ . This assumption will later be empirically validated using raw genotype data. Then, we derive Equation (3.4) from the obtained model.

**Lemma 1.** Let  $P(T|\mathbf{S}^g)$  follow the Noisy-Or model. Then for  $\mathbf{S}^g$  we have

$$P(\mathbf{S}^{g} = \mathbf{s}|T = 0) = \prod_{j=1}^{m} P(S_{j}^{g} = s_{j}|T = 0).$$

*Proof.* Conditional probability  $P(\mathbf{S}^g = \mathbf{s}|T = 0)$  can be written as

$$P(\mathbf{S}^{g} = \mathbf{s}|T = 0) = \frac{P(\mathbf{S}^{g} = \mathbf{s})}{P(T = 0)}P(T = 0|\mathbf{S}^{g} = \mathbf{s})$$
$$= \frac{\prod_{j=1}^{m} P(S_{j}^{g} = s_{j})}{P(T = 0)}P(T = 0|\mathbf{S}^{g} = \mathbf{s}),$$

$$P(S_j^g = s_j | T = 0) = \frac{P(S_j^g = s_j)}{P(T = 0)} P(T = 0 | S_j^g = s_j).$$

Thus we have

$$\frac{P(\mathbf{S}^g = \mathbf{s}|T = 0)}{\prod_{j=1}^m P(S_j^g = s_j|T = 0)} = \frac{P(T = 0|\mathbf{S}^g = \mathbf{s})P(T = 0)^{m-1}}{\prod_{j=1}^m P(T = 0|S_j^g = s_j)}.$$
(3.5)

According to the formula of total probability and the formulation of the Noisy-Or model (2.4), we have

$$P(T = 0) = \sum_{\mathbf{s}} P(\mathbf{S}^{g} = \mathbf{s}) P(T = 0 | \mathbf{S}^{g} = \mathbf{s})$$
$$= \sum_{\mathbf{s}} \left( \theta_{0} \prod_{j=1}^{m} P(S_{j}^{g} = s_{j}) \theta_{j}^{\mathbb{1}(s_{j})} \right) = \theta_{0} \prod_{j=1}^{m} \left( \sum_{s_{j}} P(S_{j}^{g} = s_{j}) \theta_{j}^{\mathbb{1}(s_{j})} \right)$$

and similarly

$$P(T = 0|S_j^g = s_j) = \theta_0 \sum_{\mathbf{s} \setminus \{s_j\}} \left( \prod_{i \neq j} P(S_i^g = s_i) P(T = 0|\mathbf{S}^g = \mathbf{s}) \right)$$
$$= \theta_0 \theta_j^{\mathbb{I}(s_j)} \prod_{i \neq j} \left( \sum_{s_i} P(S_i^g = s_i) \theta_i^{\mathbb{I}(s_i)} \right).$$

Then, it follows that

$$\frac{P(T=0)^m}{\prod_{j=1}^m P(T=0|S_j^g=s_j)} = \prod_{j=1}^m \frac{\sum_{s_j} P(S_j^g=s_j) \theta_j^{\mathbb{I}^{(s_j)}}}{\theta_j^{\mathbb{I}^{(s_j)}}}$$
$$= \frac{\theta_0 \prod_{j=1}^m \sum_{s_j} P(S_j^g=s_j) \theta_j^{\mathbb{I}^{(s_j)}}}{\theta_0 \prod_{j=1}^m \theta_j^{\mathbb{I}^{(s_j)}}} = \frac{P(T=0)}{P(T=0|\mathbf{S}^g=\mathbf{s})},$$

which indicates that Equation (3.5) equals 1. Hence, the lemma is proved.

**Lemma 2.** Let  $P(T|\mathbf{S}^g)$  follows the Noisy-Or model. Then for  $\mathbf{S}^a$  we also have

$$P(\mathbf{S}^{a} = \mathbf{s}|T = 0) = \prod_{j=1}^{m} P(S_{j}^{a} = s_{j}|T = 0).$$

*Proof.* We first derive the relationship between  $P(\mathbf{S}^a = \mathbf{s}|T = 0)$  and  $P(\mathbf{S}^g = \mathbf{s}|T = 0)$ . For each SNP  $S_j \in \mathbf{S}$ , suppose that the contingency table is as shown in Table 2.2. Then, the risk allele frequency in the case group is given by

$$\frac{2r_0+r_1}{2R}=r_0+\frac{1}{2}r_1.$$

Similarly we can compute other allele frequencies. In general, the connection between the allele frequency and the genotype frequency is given in Equation (3.6), where  $s = \{0, 1\}$  and  $t = \{0, 1\}$ .

$$P(S_{j}^{a} = s|T = t) = P(S_{j}^{g} = 2s|T = t) + \frac{1}{2}P(S_{j}^{g} = 1|T = t).$$
(3.6)

We extend this equation to a combination of multiple SNPs. Given an allele combination, if a genotype combination contains the corresponding homozygotes only, its contribution to the allele frequency is 1. If there are *j* heterozygotes in the genotype combination, the contribution to the allele frequency is  $\frac{1}{2^j}$ . To obtain the general formulation, given an allele combination **s** (e.g., (A,T)), we denote the genotype combination that contains the corresponding homozygotes only as 2**s** (e.g., (AA,TT)). We consider all of the possible genotype combinations that replace *j* homozygotes in 2**s** with heterozygotes, and denote this set by  $\pi_j(2\mathbf{s})$ . Note that  $\pi_0(2\mathbf{s}) = 2\mathbf{s}$ . Thus, the allele frequency of **s** is given by

$$P(\mathbf{S}^{a} = \mathbf{s}|T = t) = \sum_{j=0}^{m} \frac{1}{2^{j}} \sum_{\mathbf{s}' \in \pi_{j}(2\mathbf{s})} P(\mathbf{S}^{g} = \mathbf{s}'|T = t).$$

According to Equation (3.6), we have

$$\prod_{j=1}^{m} P(S_{j}^{a} = s_{j}|T = 0)$$
  
= 
$$\prod_{j=1}^{m} \left( P(S_{j}^{g} = 2s_{j}|T = 0) + \frac{1}{2}P(S_{j}^{g} = 1|T = 0) \right)$$
  
= 
$$\sum_{j=0}^{m} \frac{1}{2^{j}} \sum_{s' \in \pi_{j}(2s)} \prod_{s'_{j} \in s'} P(S_{j}^{g} = s'_{j}|T = 0).$$

According to Lemma 1,

$$P(\mathbf{S}^{g} = \mathbf{s}' | T = 0) = \prod_{s'_{j} \in \mathbf{s}'} P(S_{j}^{g} = s'_{j} | T = 0).$$

Combining the above three equations, we have

$$P(\mathbf{S}^{a} = \mathbf{s}|T = 0) = \prod_{j=1}^{m} P(S_{j}^{a} = s_{j}|T = 0).$$

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**Theorem 1.** Let  $P(T|\mathbf{S}^g)$  follow the Noisy-Or model. Then we have

$$P(T = 0 | \mathbf{S}^{a} = \mathbf{s}) = \frac{P(T = 0) \prod_{j=1}^{m} P(S_{j}^{a} = s_{j} | T = 0)}{\prod_{j=1}^{m} P(S_{j}^{a} = s_{j})}.$$

*Proof.* It directly follows Lemma 2 that

$$P(T = 0|\mathbf{S}^{a} = \mathbf{s}) = \frac{P(T = 0)P(\mathbf{S}^{a} = \mathbf{s}|T = 0)}{P(\mathbf{S}^{a} = \mathbf{s})}$$
$$= \frac{P(T = 0)\prod_{j=1}^{m} P(S_{j}^{a} = s_{j}|T = 0)}{\prod_{j=1}^{m} P(S_{j}^{a} = s_{j})}$$
$$= \frac{P(T = 0)\prod_{j=1}^{m} P(S_{j}^{a} = s_{j}|T = 0)}{\prod_{j=1}^{m} \sum_{s_{j}^{s}} P(s_{j}^{s})P(s_{j}^{a}|s_{j}^{s})}.$$

#### **3.3. Inference Based on the Constructed Bayesian Network**

With the three-layered Bayesian network constructed from the GWAS catalog, we can calculate the joint probability for any desired assignment of values to variable sets  $S^g$  of SNPs S and traits T, which reflects the relationship among genotypes and traits. We first develop the general formula for any inference on the constructed Bayesian network. Then we consider three specific inference problems, namely trait inference given SNP genotype, genotype inference given trait, and trait inference given trait. Finally, we present a typical application using the derived inference methods.

#### 3.3.1. General Inference Formula

**Theorem 2.** The joint probability for any value assignment to  $S^g$  of  $S \subseteq S$ ,  $T \subseteq T$ , i.e.,  $P(s^g, t)$ , is given by

$$P(\mathbf{s}^g, \mathbf{t}) = \prod_{S_j \in \mathbf{S}_1} P(s_j^g) \sum_{\mathbf{S}_2^a, \mathbf{S}_3^g, \mathbf{S}_3^a} \Big( \prod_{S_j \in \mathbf{S}_2 \cup \mathbf{S}_3} P(s_j^g) P(s_j^a | s_j^g) \prod_{T_k \in \mathbf{T}} P(t_k | Par(T_k)) \Big),$$

where  $S_1$  denotes the SNPs in S but not associated with T,  $S_2$  denotes the SNPs in S and also associated with T,  $S_3$  denotes the SNPs associated with T but not in S. Note that  $\sum_X f(x)$  means to sum up all f(x) going through all value assignments to attributes X.

*Proof.* The joint probability can be written as

$$P(\mathbf{s}^g, \mathbf{t}) = \sum_{\mathbf{S}^a, \bar{\mathbf{S}}^g, \bar{\mathbf{S}}^a, \bar{\mathbf{T}}} P(\mathbf{s}^g, \bar{\mathbf{s}}^g, \mathbf{s}^a, \bar{\mathbf{s}}^a, \mathbf{t}, \bar{\mathbf{t}}),$$

where  $\bar{\mathbf{S}} = S \setminus \mathbf{S}$  and  $\bar{\mathbf{T}} = \mathcal{T} \setminus \mathbf{T}$ .

According to the Markov property, the joint probability can be factorized as

$$P(\mathbf{s},\mathbf{t}) = \sum_{\mathbf{S}^a, \bar{\mathbf{S}}^g, \bar{\mathbf{S}}^a, \bar{\mathbf{T}}} \Big( \prod_{S_j \in \mathbf{S} \cup \bar{\mathbf{S}}} P(s_j^g) P(s_j^a | s_j^g) \prod_{T_k \in \mathbf{T}} P(t_k | Par(T_k)) \prod_{T_l \in \bar{\mathbf{T}}} P(t_l | Par(T_l)) \Big)$$

which follows that

$$P(\mathbf{s}^{g}, \mathbf{t}) = \sum_{\mathbf{s}^{a}, \mathbf{\bar{S}}^{g}, \mathbf{\bar{S}}^{a}S_{j} \in \mathbf{S} \cup \mathbf{\bar{S}}} \left( \prod_{s_{j} \in \mathbf{S} \cup \mathbf{\bar{S}}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \prod_{T_{k} \in \mathbf{T}} P(t_{k} | Par(T_{k})) \sum_{\mathbf{\bar{T}}} \prod_{T_{l} \in \mathbf{\bar{T}}} P(t_{l} | Par(T_{l})) \right)$$
$$= \sum_{\mathbf{s}^{a}, \mathbf{\bar{S}}^{g}, \mathbf{\bar{S}}^{a}} \left( \prod_{s_{j} \in \mathbf{S} \cup \mathbf{\bar{S}}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \prod_{T_{k} \in \mathbf{T}} P(t_{k} | Par(T_{k})) \right).$$

Then, we divide S into four disjoint subsets:  $S_1$  denotes the SNPs in S but not associated with T,  $S_2$  denotes the SNPs in S and also associated with T,  $S_3$  denotes the SNPs associated with T but not in S, and  $S_4$  denotes all the other SNPs. Thus,  $S = S_1 \cup S_2$ ,  $\bar{S} = S_3 \cup S_4$ , and  $Par(T_k)$  for  $T_k \in T$
only involves SNPs in  $S_2$  and  $S_3$ . It follows that

$$\begin{split} P(\mathbf{s}^{g}, \mathbf{t}) &= \sum_{\mathbf{S}^{a}, \tilde{\mathbf{S}}^{g}, \tilde{\mathbf{S}}^{g}} \left( \prod_{s_{j} \in \mathbf{S} \cup \mathbf{S}_{3}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \prod_{s_{j} \in \mathbf{S}_{4}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \prod_{T_{k} \in \mathbf{T}} P(t_{k} | Par(T_{k})) \right) \\ &= \sum_{\mathbf{S}^{a}, \mathbf{S}^{g}_{3}, \mathbf{S}^{g}_{3} \in \mathbf{S} \cup \mathbf{S}_{3}} \left( \prod_{s_{j} \in \mathbf{S} \cup \mathbf{S}_{3}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \prod_{T_{k} \in \mathbf{T}} P(t_{k} | Par(T_{k})) \sum_{\mathbf{S}^{g}_{4}, \mathbf{S}^{g}_{4}} \prod_{s_{j} \in \mathbf{S}_{4}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \right) \\ &= \sum_{\mathbf{S}^{a}, \mathbf{S}^{g}_{3}, \mathbf{S}^{g}_{3} \leq j \in \mathbf{S} \cup \mathbf{S}_{3}} \left( \prod_{s_{j} \in \mathbf{S} \cup \mathbf{S}_{3}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \prod_{T_{k} \in \mathbf{T}} P(t_{k} | Par(T_{k})) \right) \\ &= \sum_{\mathbf{S}^{a}, \mathbf{S}^{g}_{3}, \mathbf{S}^{g}_{3} \leq j \in \mathbf{S}_{2} \cup \mathbf{S}_{3}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \prod_{s_{j} \in \mathbf{S}_{1}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \prod_{T_{k} \in \mathbf{T}} P(t_{k} | Par(T_{k})) \right) \\ &= \sum_{\mathbf{S}^{a}, \mathbf{S}^{g}_{3}, \mathbf{S}^{g}_{3} \leq j \in \mathbf{S}_{2} \cup \mathbf{S}_{3}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \prod_{s_{j} \in \mathbf{S}_{1}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \prod_{T_{k} \in \mathbf{T}} P(t_{k} | Par(T_{k})) \sum_{\mathbf{S}^{a}_{1} \in \mathbf{T}} P(t_{k} | Par(T_{k})) \right) \\ &= \sum_{\mathbf{S}^{a}_{2}, \mathbf{S}^{g}_{3}, \mathbf{S}^{g}_{3} \leq j \in \mathbf{S}_{2} \cup \mathbf{S}_{3}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \prod_{T_{k} \in \mathbf{T}} P(t_{k} | Par(T_{k})) \prod_{S_{j} \in \mathbf{S}_{1}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \right) \\ &= \sum_{\mathbf{S}^{a}_{2}, \mathbf{S}^{g}_{3}, \mathbf{S}^{g}_{3} \leq j \in \mathbf{S}_{2} \cup \mathbf{S}_{3}} \left( \prod_{s_{j} \in \mathbf{S}_{2} \cup \mathbf{S}_{3}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \prod_{T_{k} \in \mathbf{T}} P(t_{k} | Par(T_{k})) \prod_{S_{j} \in \mathbf{S}_{1}} P(s_{j}^{g}) \right) \\ &= \prod_{S_{j} \in \mathbf{S}_{1}} P(s_{j}^{g}) \sum_{\mathbf{S}^{a}_{2}, \mathbf{S}^{g}_{3}, \mathbf{S}^{g}_{3} \leq j \in \mathbf{S}_{2} \cup \mathbf{S}_{3}} \left( \prod_{s_{j} \in \mathbf{S}_{2} \cup \mathbf{S}_{3}} P(s_{j}^{g}) P(s_{j}^{g} | s_{j}^{g}) \prod_{T_{k} \in \mathbf{T}} P(t_{k} | Par(T_{k})) \right) . \end{split}$$

Note that in Theorem 2, we apply marginalization to sum out 'irrelevant' variables so that we do not need to involve all variables in our summation to calculate  $P(\mathbf{s}^g, \mathbf{t})$ . As a result, the computation only involves variables in  $\mathbf{T}$ ,  $\mathbf{S}_1$ ,  $\mathbf{S}_2$  and  $\mathbf{S}_3$ .

Additionally, we can calculate the conditional joint probability for any *desired* assignment of values to variable sets  $\mathbf{S}_x^g$ ,  $\mathbf{T}_x$  given the *observed* assignment of variable sets  $\mathbf{S}_y^g$ ,  $\mathbf{T}_y$  following Theorem 3. Note that  $\mathbf{S}_x^g$  and  $\mathbf{S}_y^g$  denote the set of SNP genotypes; while  $\mathbf{T}_x$ ,  $\mathbf{T}_y$  denote the set of traits.

**Theorem 3.** The probability for any desired assignment of values  $\mathbf{s}_x^g, \mathbf{t}_x$  to variables in  $\mathbf{S}_x^g, \mathbf{T}_x$  given the (observed) assignment of values  $\mathbf{s}_y^g, \mathbf{t}_y$  to variables in  $\mathbf{S}_y^g, \mathbf{T}_y$  can be directly derived

$$P(\mathbf{s}_x^g, \mathbf{t}_x | \mathbf{s}_y^g, \mathbf{t}_y) = \frac{P(\mathbf{s}_x^g, \mathbf{t}_x, \mathbf{s}_y^g, \mathbf{t}_y)}{P(\mathbf{s}_y^g, \mathbf{t}_y)}$$
(3.7)

where the joint probability  $P(\mathbf{s}_x^g, \mathbf{t}_x, \mathbf{s}_y^g, \mathbf{t}_y)$  and  $P(\mathbf{s}_y^g, \mathbf{t}_y)$  can be calculated following Theorem 2.

A given Bayesian network can be used to derive the posterior probability distribution of one or more variables in the network given the values observed for other variables in the network. Theorem 2 and Theorem 3 show the simple and brute-force formulae, which have exponential time complexity and are not computationally tractable. Researchers have developed various efficient exact inference algorithms that take advantage of independence relationships represented in a Bayesian network, and stochastic approximation algorithms to estimate exact inference results when exact inference is prohibitively time consuming [41].

## 3.3.2. Trait Inference Given SNP Genotype

We assume that we have been given the genotype profile of the target and aim to derive the probability that the target has a specific trait using the constructed Bayesian network. The probability of the prevalence of a specific trait, which is retrievable from literature or internet, is used as the prior probability that the target has the specific trait. We then calculate the posterior probability of the target having the trait by inferring from the target's genotypes. Formally, we represent the genotypes of a target v as a vector,  $\mathbf{s}_{v}^{g} = (s_{v1}^{g}, s_{v2}^{g}, \dots, s_{vn}^{g})$ , with each entry  $s_{vj}^{g}$  denoting the genotype of SNP *j*.

**Definition 1.** The problem of trait inference given SNP genotype, aims to learn the posteriori probability  $P(t|\mathbf{s}_{v}^{g})$  that the target has a specific trait T given the target's genotype profile  $\mathbf{s}_{v}^{g}$  using the constructed Bayesian network.

The posteriori probability  $P(t|\mathbf{s}_v^g)$  can be calculated following Equation (3.7), specifically with  $\mathbf{s}_x^g = \emptyset$ ,  $\mathbf{t}_y = \emptyset$ ,  $\mathbf{t}_x = \{t\}$ , and  $\mathbf{s}_y^g = \mathbf{s}_v^g$ . In Lemma 3, we show our simplified formula where the calculation only involves SNPs that are associated with trait *T*.

**Lemma 3.** The posteriori probability  $P(t|\mathbf{s}_v^g)$  can be calculated as:

$$P(t|\mathbf{s}_{v}^{g}) = \sum_{\mathbf{Q}^{a}} \Big( \prod_{S_{j} \in \mathbf{Q}} P(s_{j}^{a}|s_{v_{j}}^{g}) P(t|\mathbf{q}^{a}) \Big),$$
(3.8)

where  $\mathbf{Q}$  denotes the SNPs that are associated with trait T.

*Proof.* Denote by **Q** the SNPs that are associated with trait *T*. We have  $P(t|\mathbf{s}_v^g) = \frac{P(t,\mathbf{s}_v^g)}{P(\mathbf{s}_v^g)}$  and apply Theorem 2 to compute  $P(t, \mathbf{s}_v^g)$ . Note that  $\mathbf{S}_1 = \bar{\mathbf{Q}}$ ,  $\mathbf{S}_2 = \mathbf{Q}$ , and  $\mathbf{S}_3 = \emptyset$ . Thus, we have

$$P(t, \mathbf{s}_{v}^{g}) = \prod_{S_{j} \in \bar{\mathbf{Q}}} P(s_{vj}^{g}) \sum_{\mathbf{Q}^{a}} \Big( \prod_{S_{j} \in \mathbf{Q}} P(s_{vj}^{g}) P(s_{j}^{a} | s_{vj}^{g}) P(t | \mathbf{q}^{a}) \Big).$$

Therefore, it results that

$$P(t|\mathbf{s}_{v}^{g}) = \frac{P(t, \mathbf{s}_{v}^{g})}{P(\mathbf{s}_{v}^{g})}$$

$$= \frac{\prod_{S_{j} \in \bar{\mathbf{Q}}} P(s_{vj}^{g}) \sum_{\mathbf{Q}^{a}} \left(\prod_{S_{j} \in \mathbf{Q}} P(s_{vj}^{g}) P(s_{j}^{a}|s_{vj}^{g}) P(t|\mathbf{q}^{a})\right)}{\prod_{S_{j} \in \mathbf{S}} P(s_{vj}^{g})}$$

$$= \frac{\prod_{S_{j} \in \bar{\mathbf{Q}}} P(s_{vj}^{g}) \prod_{S_{j} \in \mathbf{Q}} P(s_{vj}^{g}) \sum_{\mathbf{Q}^{a}} \left(\prod_{S_{j} \in \mathbf{Q}} P(s_{j}^{a}|s_{vj}^{g}) P(t|\mathbf{q}^{a})\right)}{\prod_{S_{j} \in \mathbf{S}} P(s_{vj}^{g})}$$

$$= \sum_{\mathbf{Q}^{a}} \left(\prod_{S_{j} \in \mathbf{Q}} P(s_{j}^{a}|s_{vj}^{g}) P(t|\mathbf{q}^{a})\right).$$

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		_

Specifically, according to Equation (3.4), we have

$$P(T = 0|\mathbf{s}_{v}^{g}) = P(T = 0) \sum_{\mathbf{Q}^{a}} \Big( \prod_{s_{j} \in \mathbf{Q}} \frac{P(s_{j}^{a}|s_{v_{j}}^{g})P(s_{j}^{a}|T = 0)}{\sum_{s_{j}^{g}} P(s_{j}^{g})P(s_{j}^{a}|s_{j}^{g})} \Big),$$

which shows how the prior probability is updated to obtain the posteriori probability. Note that  $P(T = 1|\mathbf{s}_{v}^{g}) = 1 - P(T = 0|\mathbf{s}_{v}^{g})$  as  $P(T = 1|\mathbf{s}_{v}^{g})$  is often of interest to users.

Lemma 3 implies that, instead of conducting inference based on the whole Bayesian network G, we can simply identify the subgraph G' that contains all associated SNPs of the target trait T, and then calculate the posterior probability following Equation (3.8).

Trait inference can help an individual discovers the risk of having a certain disease based on

his/her genotype profile. If the genotype profile of an individual has been stolen, then it introduces genetic privacy concerns since the genotypes can be used to infer private trait information of the target by attackers.

#### **3.3.3.** Genotype Inference Given Trait

In this problem, we aim to acquire the probability that an individual has specific genotypes for a set of SNPs given his/her associated trait information, with the Bayesian network constructed. Formally, we denote by  $\mathbf{s}_i^g = (s_{i1}^g, s_{i2}^g, \dots, s_{in}^g)$  an arbitrary genotype profile. A subset of a target's trait  $\mathbf{T}_v$  with its value assignment  $\mathbf{t}_v$  is given.

**Definition 2.** The problem of genotype inference given trait aims to learn the posteriori probability  $P(\mathbf{s}_i^g | \mathbf{t}_v)$  that the target has a genotype profile  $\mathbf{s}_i^g$  given the target's traits  $\mathbf{t}_v$  using the constructed Bayesian network.

**Lemma 4.** The posterior probability  $P(\mathbf{s}_i^g | \mathbf{t}_v)$  is

$$P(\mathbf{s}_{i}^{g}|\mathbf{t}_{v}) = \frac{\prod_{S_{j}\in\mathbf{Q}}P(s_{ij}^{g})\sum_{\mathbf{Q}^{a}}\left(\prod_{S_{j}\in\mathbf{Q}}P(s_{ij}^{g})P(s_{j}^{a}|s_{ij}^{g})\prod_{T_{k}\in\mathbf{T}_{v}}P(t_{k}|Pa(T_{k}))\right)}{\sum_{\mathbf{Q}^{g},\mathbf{Q}^{a}}\left(\prod_{S_{j}\in\mathbf{Q}}P(s_{j}^{g})P(s_{j}^{a}|s_{j}^{g})\prod_{T_{k}\in\mathbf{T}_{v}}P(t_{k}|Pa(T_{k}))\right)},$$

where **Q** denotes the SNPs that are associated with traits in  $\mathbf{T}_{v}$ , and  $P(t_k|Pa(T_k))$  is computed according to Equation (3.4).

*Proof.* We have  $P(\mathbf{s}_i^g | \mathbf{t}_v) = \frac{P(\mathbf{s}_i^g, \mathbf{t}_v)}{P(\mathbf{t}_v)}$  and apply Theorem 2 to compute the probabilities. For  $P(\mathbf{s}_i^g, \mathbf{t}_v)$ , similar to the proof to Lemma 3, we obtain

$$P(\mathbf{s}_i^g, \mathbf{t}_v) = \prod_{S_j \in \mathbf{Q}} P(s_{ij}^g) \sum_{\mathbf{Q}^a} \Big( \prod_{S_j \in \mathbf{Q}} P(s_{ij}^g) P(s_j^a | s_{ij}^g) \prod_{T_k \in \mathbf{T}_v} P(t_k | Pa(T_k)) \Big),$$

where **Q** denotes the SNPs that are associated with traits in  $\mathbf{T}_{v}$ . For  $P(\mathbf{t}_{v})$ , when applying Theo-

rem 2, note that  $\mathbf{S} = \emptyset$  and  $\mathbf{\bar{S}} = \mathbf{Q}$ . Thus we have

$$P(\mathbf{t}_{v}) = \sum_{\mathbf{Q}^{g}, \mathbf{Q}^{a}} \Big( \prod_{s_{j} \in \mathbf{Q}} P(s_{j}^{g}) P(s_{j}^{a} | s_{j}^{g}) \prod_{T_{k} \in \mathbf{T}_{v}} P(t_{k} | Pa(T_{k})) \Big).$$

3.3.4. Trait Inference Given Trait

A straightforward extension to the above two inferences is that, we can also infer other trait information of the target individual. Assume that we are given some of the target's traits  $\mathbf{t}_{v}$ . Then Lemma 5 gives the probability that the target has a new trait  $T_{new}$ .

**Lemma 5.** The probability that the target has a new trait  $T_{new}$  given some of the target's traits  $\mathbf{t}_v$  can be derived as

$$P(t_{new}|\mathbf{t}_v) = \sum_{\mathbf{Q}^g} P(t_{new}|\mathbf{q}^g) P(\mathbf{q}^g|\mathbf{t}_v),$$

where  $\mathbf{Q}$  is the set of SNPs associated with  $t_{new}$  and  $\mathbf{t}_{v}$ .

The proof is straightforward by applying the *d*-separation criterion [43]. We can see that  $P(t_{new}|\mathbf{q}^g)$  can be derived following Lemma 3, and  $P(\mathbf{q}^g|\mathbf{t}_v)$  can be derived following Lemma 4.



Figure 3.2: Identity attack

#### **3.3.5.** Application: Identity Attack

We present an attack that aims to infer the probability of a record in an anonymized genotype database that belongs to a target, when some traits of the target are available. As shown in Figure 3.2, assume that an attacker has access to an anonymized genotype dataset  $\mathcal{R}$  that contains the target's genotype record  $\mathbf{s}_{v}^{g}$ . The attacker also knows a subset of traits  $\mathbf{t}_{v}$  the target has. Then the attacker can learn the posteriori probability  $P(\mathbf{s}_{i}^{g} == \mathbf{s}_{v}^{g} | \mathbf{t}_{v})$  that each genotype record  $\mathbf{s}_{i}^{g}$  in the database corresponds to the target, as shown in Lemma 6. As a result, the attacker may be able to identify the target's record from the anonymized dataset.

**Lemma 6.** The posteriori probability that the genotype record  $\mathbf{s}_i^g$  corresponds to the target given *his trait*  $\mathbf{t}_v$  *is given by* 

$$P(\mathbf{s}_i^g == \mathbf{s}_v^g | \mathbf{t}_v) = \frac{P(\mathbf{s}_v^g | \mathbf{t}_v)}{\sum_{i=1}^{|\mathcal{R}|} P(\mathbf{s}_i^g | \mathbf{t}_v)}.$$

### **3.4. Further Extensions**

In this paper we treat SNPs as they are mutually independent since the SNP-SNP correlations cannot be obtained from the GWAS catalog. However, in some situations the SNP-SNP correlations may be available, e.g., being provided by some large-scale biomedical studies. In this subsection, we briefly discuss how to integrate the SNP-SNP correlations into our model.



Figure 3.3: An example network where  $S_1$  and  $S_2$  are correlated.

When the SNP-SNP correlations are available, we assume that in addition to the allele frequency in the case and control groups, we also know the joint genotype frequency of the correlated SNPs. Then, a straightforward extension of our model can be given as follows. For two or more correlated SNPs, we cluster their corresponding nodes in the genotype layer as a single super node. The super node represents the combination of the SNP genotypes, and takes value as the crossproduct of the sets of values of the genotypes. There is an edge pointing from the super node to each corresponding allele node. Note that the clustered Bayesian network represents the same joint probability distribution as the original Bayesian network.

Figure 3.3 shows an example, where SNPs  $S_1$  and  $S_2$  are correlated. Thus, we cluster nodes  $S_1^g, S_2^g$  as a single node  $S_{1,2}^g$ , i.e.,  $S_{1,2}^g = S_1^g \times S_2^g$ . Node  $S_{1,2}^g$  has two emanating edges pointing to  $S_1^a$  and  $S_2^a$  respectively. Denoting the value combination  $(s_1^g, s_2^g)$  by  $s_{1,2}^g$ , according to Equation (2.3), the joint probability of  $P(s_{1,2}^g, s_3^g, t_1, t_2)$  in the clustered Bayesian network is given by

$$P(s_{1,2}^g, s_3^g, t_1, t_2) = \sum_{s_1^a, s_2^a, s_3^a} P(s_{1,2}^g) P(s_3^g) P(s_1^a | s_{1,2}^g) P(s_2^a | s_{1,2}^g) P(s_3^a | s_3^g) P(t_1 | s_1^a, s_2^a) P(t_2 | s_2^a, s_3^a).$$
(3.9)

In Equation (3.9),  $P(s_{1,2}^g) = P(s_1^g, s_2^g)$  is assumed to be given representing the known SNP-SNP correlation. For  $P(s_1^a|s_{1,2}^g)$  (resp.  $P(s_2^a|s_{1,2}^g)$ ), as shown in Section 3.2.2 it represents for SNP  $S_1$  (resp.  $S_2$ ) how the genetic effect of the genotype is obtained from the genetic effects of its two alleles, hence has no connection with other SNPs. So, we have  $P(s_1^a|s_{1,2}^g) = P(s_1^a|s_1^g)$  and  $P(s_2^a|s_{1,2}^g) = P(s_1^a|s_2^g)$ . For  $P(t_2|s_2^a, s_3^a)$ , it can be accurately computed using Theorem 1 since  $S_{1,2}^g$  and  $S_3^g$  are independent. The only issue of exactly computing Equation (3.9) lies in the computing of  $P(t_1|s_1^a, s_2^a)$ . Since  $P(t_1|s_1^a, s_2^a)$  can be written as  $\frac{P(t_1)}{P(s_1^a, s_2^a)}P(s_1^a, s_2^a|t_1)$ , and we can easily obtain that  $P(s_1^a, s_2^a) = \sum_{s_{1,2}^g} P(s_1^a|s_1^g)P(s_2^a|s_2^g)P(s_{1,2}^g)$ , we focus on the computing of  $P(s_1^a, s_2^a|t_1)$ .

If  $P(s_1^a, s_2^a|t_1)$  is also given, then Equation (3.9) can be exactly computed. If not, we can estimate

 $P(s_1^a, s_2^a|t_1)$  as follows. We have

$$P(s_1^a, s_2^a) - P(s_1^a)P(s_2^a) = \sum_{T_1} P(s_1^a, s_2^a | t_1)P(t_1) - \sum_{T_1} P(s_1^a | t_1)P(t_1) \sum_{T_1} P(s_2^a | t_1)P(t_1).$$

Usually,  $P(T_1 = 0)$  is much larger than  $P(T_1 = 1)$ . Thus, by approximating  $\frac{P(T_1=1)}{P(T_1=0)}$  and  $\frac{P(T_1=1)}{\sqrt{P(T_1=0)}}$  by zero, it follows that

$$P(s_1^a, s_2^a) - P(s_1^a)P(s_2^a) \approx P(T_1 = 0) \left( P(s_1^a, s_2^a | T_1 = 0) - P(s_1^a | T_1 = 0)P(s_2^a | T_1 = 0) P(T_1 = 0) \right),$$

which leads to

$$P(s_1^a, s_2^a | T_1 = 0) \approx \frac{P(s_1^a, s_2^a) - P(s_1^a)P(s_2^a)}{P(T_1 = 0)} + P(s_1^a | T_1 = 0)P(s_2^a | T_1 = 0)P(T_1 = 0)$$

It should be noted that, the above extension cannot deal with the situation where the SNP-SNP correlations have overlaps, e.g., in Figure 3.3  $S_2$  is correlated with both  $S_1$  and  $S_3$  but the correlation among the three SNPs are not available. In this case, we can resort to the factor graph model [32] to represent the SNP-SNP correlations. We leave the detailed study to the future work.

# 3.5. Experiments

We first validate the Noisy-Or model in Section 3.5.1. Then we construct the Bayesian network from the GWAS catalog in Section 3.5.2. The inference methods and their applications are evaluated in Sections 3.5.3 and 3.5.4.

## 3.5.1. Noisy-Or Model Validation

To evaluate the fitness of the Noisy-Or model in modeling the SNP-trait association, we use raw data from openSNP [10] where more than two thousand users over the world share their genotype profiles and trait information. The genotype file contains the results of the genetic test taken by

each user. Each line in the file corresponds to one SNP with its identifier (rsid), its location on the reference human genome and alleles provided. Besides, users also contribute their phenotypes to openSNP, such as what the color of their eyes, whether they have astigmatism, or whether they are suffering from irritable bowel syndrome.

### Data Setup

In the experiments, we use openSNP of version 20151231. The genetic test results provided by users are taken from different genetic screening services. We focus on the genotyping files from 23andMe, Ancestry and FamilyTreeDNA. The data from these services account for more than 99% of the whole dataset. Among the 341 traits from the original data, there are 129 binary traits, 136 non-binary categorical traits, 39 numeric traits and 14 traits with unknown values. In align with GWAS case-control settings, we focus on the 129 binary traits to evaluate our models.

The data in openSNP is highly sparse and contains a mass of missing values due to various genetic testing platforms and varying willingness of individuals to share their traits. To ensure that the statistic tests in the model construction are meaningful, we further filter the data as follows. For each trait, we extract the individuals that belong to the control group and the case group. If the number of individuals contained in both groups for a trait is less than 10, we exclude this trait from our experiment. As a result, we obtain 71 traits satisfying the requirement. Then, following a typical GWAS procedure [52], from all associated SNPs for each trait, we remove the SNPs with: 1) low minor allele frequency (i.e., <1%); 2) call rate less than 90%; and 3) the number of records containing the risk allele less than 10. After that, we discard the traits with no associated SNPs left after filtering. Finally, we obtain a dataset which contains 23 traits and 256,845 SNPs.

### Results

To build the Bayesian network, we extract for each trait the associated SNPs along with risk allele types, risk allele frequencies and odds ratios. For each SNP, the allele frequencies in the case group and the control group and odds ratios are computed. If the odds ratio is larger than 1, the

	CULD		CULD
Traits	SNPs	Traits	SNPs
Eve with Blue Halo	rs6913354	Irritable Bowel Syndrome	rs8039023
Lyc with Dide Halo	rs10460585	Initiable Bower Syndrome	rs2948814
	rs1239925		rs3923767
Hair on Fingers	rs11715867		rs2531864
	rs2302025	Do You Grind Your Teeth	rs2042279
	rs1496496		rs12094507
	rs4619		rs9809185
	rs7235392		rs2409764
ADIID	rs664510		rs12564559
	rs1910236		rs10882959
	rs6922476	-	rs6601522
	rs747644	Enjoy Driving A Car	rs1002399
	rs1466410	Linjoy Driving A Car	rs6993841
Actionation	rs11680053	-	rs958648
Asuginausin	rs12358733		rs3808513
	rs1400390		rs6601518
	rs10508470		rs357281

Table 3.1: SNP-Trait associations

corresponding allele is considered as the risk allele. Then, we perform the Fisher's exact test of independence to test whether the association between the trait and the SNP is significant. The threshold of the *p*-value is set as  $4 \times 10^{-5}$ . We discard the traits with zero associated SNP, as well as the traits with only one associated SNP as they have no effect in testing ICI model. As a result, we obtain 7 traits and 34 associated SNPs for building the Bayesian network, as shown in Table 3.1.

Trait	Chi-square	df	p-Value	RMSEA
Eye with Blue Halo	6.73	4	0.15	0.10
Hair on Fingers	14.46	14	0.41	0.02
Irritable Bowel Syndrome	5.24	4	0.26	0.05
ADHD	55.32	53	0.38	0.02
Astigmatism	132.55	123	0.26	0.02
Do You Grind Your Teeth	50.13	49	0.42	0.02
Enjoy Driving A Car	96.33	98	0.52	NA

Table 3.2: The chi-square value, degree of freedom (df), p-value, RMSEA of the Noisy-or model

We then evaluate the fitness of the Noisy-Or model. For each trait, we predict the observed number of individuals with a specific trait and specific SNP genotypes, i.e.,  $n(T, \mathbf{S}^g)$ , by computing the predicted value as  $\hat{n}(T, \mathbf{S}^g) = P(T|\mathbf{S}^g)n(\mathbf{S}^g)$ , where  $n(\mathbf{S}^g)$  is the observed total number of individuals with the SNP genotypes. Since the data is highly sparse, when computing the chi-square value we only sum up the cells where  $n(S^g)$  does not equal to 0. We then compute the *p*-value to show the significance. The degree of freedom is computed as "total number of predictions - the number of non-zero  $n(S^g)$  - the number of model parameters". The null hypothesis  $H_0$  assumes that there is no relationship between the data and the model. Thus, the model is not rejected if *p*-value >0.05. In addition, we further compute the Root Mean Square Error of Approximation (RMSEA) values [34] which is an absolute measure of fit, to show the degree of the fitness. The RMSEA values are categorized into four levels: close fit (.00 - .05), fair fit (.05 - .08), mediocre fit (.08-.10) and poor fit (over .10). Note that RMSEA is applicable only when the chi-square value is larger than the degree of freedom (df), and is labelled as 'NA' otherwise. The results are shown in Table 3.2. As can be seen, the Noisy-Or model is accepted for all traits according to the *p*-values, which indicates the model is a good fit. The values of RMSEA show a close fit in general. Therefore, we validate the use of the Noisy-Or model in modeling SNP-trait association.

## 3.5.2. Bayesian Network Construction

With the justified Noisy-Or model for constructing a Bayesian network, we set out to construct a Bayesian network captured in GWAS statistics. Specifically, we construct a Bayesian network using data extracted from the online GWAS catalog [57] as of Feb 25th, 2016. This version of the GWAS catalog includes 2,347 publications and 23,152 records (SNP-trait pairs) about 17,781 SNPs associated with 1,457 traits. Publications included in such a catalog are limited to those attempted to assay at least 100,000 SNPs in the initial stage. SNP-trait pairs listed are limited to those with *p*-values less than  $10^{-5}$ . For each record, the odds ratio or beta coefficient is provided to indicate the association of the trait-SNP pair, depending on whether the trait is categorical (e.g., some disease) or numerical (e.g., height). The two values are contained in the same field in the dataset.

In this chapter, we target for categorical variables only. Thus, we focus on a subset of data published as the interactive diagram by the GWAS catalog, where an additional attribute "orType" is used to clearly indicate whether the odds ratio is provided. This subset of data includes 5,047

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records with 791 traits associated with 4,250 SNPs, and SNP-trait pairs are limited to those with p-values less than  $5 \times 10^{-8}$ . We extract the records with the odds ratio provided. As a result, we obtain 2,325 records with 266 traits associated 2,177 SNPs. Among these SNPs, there are 1,941 SNPs associated with a single trait, 122 SNPs associated with two traits, and at the most, one SNPs associated with 7 traits. Finally, we build a knowledge database for all extracted traits and associated SNPs including the risk allele type, risk allele frequency in the control group, and the odds ratio.

Based on the knowledge database, we build the Bayesian network according to Section 3.2.2. Particularly, to acquire the prior probability (prevalence) of each trait, we classify all the traits into 17 categories (e.g., immune system disease, nervous system disease), and retrieve the average prevalence of each category from the Wikipedia [58]. We use the average prevalence of a category as the prior probability of each trait belonging to the category. Our constructed Bayesian network can be refined by assigning the accurate prior probability for each trait when available.

#### 3.5.3. Simulated Scenario: Trait Inference

We evaluate the constructed Bayesian network using two simulated scenarios. In the first scenario, we infer the probability of an individual of having a trait given his/her genotype profile using the constructed Bayesian network. We use the genotype profiles in the 1000 Genomes Project [50] and extract a dataset referred to as 'CEU' for our experiment. It consists of 99 HapMap individuals from Utah residents with Northern and Western European ancestry (CEU) in the 1000 Genomes Project, which are treated as targets of trait inference in this study.

For each CEU individual *v*, we compute his/her posterior probability  $P(T_k = 1|\mathbf{s}_v^g)$  of having each trait  $T_k$  given the SNP genotype profile  $\mathbf{s}_v^g$  according to Lemma 3. Then, we compute the relative difference *rd* between the prior probability and the posterior probability of each trait, i.e.,  $rd = \frac{P(T_k=1|\mathbf{s}_v^g)-P(T_k=1)}{P(T_k=1)}$ , and rank the traits according to the *rd* for each individual. Figure 3.4 shows for top-3 and bottom-3 traits of each individual. A total of 24 traits are included as illustrated in



Figure 3.4: Top-3 and bottom-3 traits of each CEU individual

Figure 3.4, each of which is represented as a row. Each column shows the top traits of an individual, where the green and red dots represent the traits with the most positive and negative *rd* respectively.

Index	Trait	SNP-risk allele	$f_{kj}^t(r)$	$O_{kj}$	$f_{kj}^c(r)$	$P(t_k)$
1	Type 1 diabetes	rs9272346-G	0.13	8.3	0.55	0.25
		rs2647044-A	0.61	5.49	0.90	0.23
2	Behcet's disease	rs17482078-T	0.02	4.56	0.09	0.04
		rs11924265-C	0.02	3.99	0.08	
3	Crohn's disease	rs76418789-G	0.93	2.06	0.97	0.26
		rs2066847-G	0.06	2.27	0.13	
4	Fuchs's corneal dystrophy	rs613872-G	0.15	5.47	0.49	0.09
5	Freckles	rs1805007-T	0.05	4.37	0.19	0.05
6	Celiac disease	rs2187668 T	0.26	6.23	0.68	0.26
7	Immunoglobulin A	152107000-1	0.13	2.53	0.27	0.05

Table 3.3: Trait-SNP association

Tables 3.3 and 3.4 show the information of a snapshot of the constructed Bayesian network and the computed posterior probabilities. There are 7 traits and 9 SNPs. In Table 3.3, the risk allele type, risk allele in the control group and the odds ratio of each each SNP-trait pair are shown in Columns 3-5. Note that SNP rs2187668 is associated with two traits. The calculated risk allele frequency in the case group for each SNP-trait is shown in Column 6. Note that there is a big gap between the risk allele frequency in the case group and that in the control group. The prior probability (prevalence) of each trait is shown in Column 7. In Table 3.4, each index corresponds to the trait with the same index in Table 3.3. Columns  $s_{\nu}^{g}$  and *Count* respectively show the genotypes of

the associated SNPs and the number of individuals who have the genotypes. As before, 0 denotes the genotype of two non-risk alleles, 2 denotes the genotype of two risk alleles, and 1 denotes the genotype of one risk allele and one non-risk allele. Column  $P(t|\mathbf{s}_v^g)$  shows the posterior probability of one individual has a trait given his SNP genotype profile. The last column *rd* shows the relative difference between the prior probability and the posterior probability of each trait. As can be seen, all the posterior probabilities are significantly different from the corresponding prior probability of having a trait. In general, the posterior probability of a trait is larger if the individual has more risk alleles.Hence, the constructed Bayesian network is useful to infer new trait information. We also observe that, when there are multiple associated SNPs, the effect of each SNP can be different. For example, Trait 1 is associated with two SNPs. The posterior probability when the genotypes are (0, 1) is larger than that when the genotypes are (2, 0), implying that the second SNP has greater effect than the first one.

Table 3.4: Posterior probability of certain trait considering associated SNPs

Index	$\mathbf{s}_v^g$	Count	$P(t \mathbf{s}_v^g)$	rd	Index	$\mathbf{s}_v^g$	Count	$P(t \mathbf{s}_v^g)$	rd
	(0,0)	28	0.149	-0.403		(0,2,2)	89	0.349	0.341
	(1,0)	30	0.198	-0.208	3	(1,2,2)	7	0.367	0.411
1	(2,0)	22	0.247	-0.012		(2,2,2)	3	0.385	0.481
	(0,1)	10	0.269	0.078	4	(0)	66	0.056	-0.379
	(1,1)	6	0.311	0.245		(1)	27	0.150	0.670
	(2,1)	3	0.353	0.413		(2)	6	0.245	1.718
	(0)	55	0.037	-0.064	5	(0)	75	0.043	-0.138
2	(1)	36	0.094	1.351		(1)	23	0.104	1.076
	(2)	8	0.151	2.766		(2)	1	0.164	2.289
6	(0)	80	0.129	-0.501	7	(0)	80	0.024	-0.511
0	(1)	19	0.305	0.174	] /	(1)	19	0.108	1.162

## 3.5.4. Simulated Scenario: Identity Inference

In this scenario, we evaluate whether a target individual can be identified from an anonymized genotype database by an attacker given some traits of the target individual using the Bayesian network. For comparison we also include Humbert's de-anonymizing method proposed in [20]. This method also aims to identify the genotypes that correspond to the given traits, making use of the single SNP-single trait correlation. The difference lies in that this method rely upon some invalidated independence assumption, whereas our method is based on the independence of causal

influence, which is shown to have good fitness in modeling the SNP-trait associations in Section 6.1. We compare the identification accuracy of the Humbert's method to our method.

Trait	SNP-risk allele	$O_{kj}$
Exfoliation glaucoma	rs893818-A	20.94
Externation gradeonna	rs3825942-G	20.1
	rs11697186-A	33.33
Response to hepatitis C treatment	rs8099917-G	27.1
	rs6139030-T	25
Blue vs. brown eyes	rs1667394-T	29.43
Skin pigmentation	rs1834640-G	12.5

Table 3.5: Trait-SNP pairs

We consider the 7 trait-SNPs pairs listed in Table 3.5 whose odds ratios are larger than 10. The 'CEU' dataset is used to serve as the anonymized genotype database. To simulate an attack, we first designate a target individual whose traits  $\mathbf{t}_v$  and genotypes  $\mathbf{s}_v^g$  are known. Then we blend the genotype profile of the target into the 'CEU' dataset (containing the genotype records of the 99 unrelated CEU individuals), and attempt to re-identify it assuming that the attacker only knows the target's traits  $\mathbf{t}_v$ . To define the target, we assume that the target has all the traits, i.e., the target has  $T_k = 1$  for each trait  $T_k$ . We then randomly generate the genotype record for the target individual. The generating strategy is that for each SNP  $S_{kj}$  associated with one trait  $T_k$ , we generate  $S_{kj}^g = s_{kj}^g$  with the probability  $P(S_{kj}^g = s_{kj}^g|T_k = 1)$ . In this way we simulate a scenario where the target is randomly selected from the case group characterized by GWAS statistics. Finally, we calculate the probability that the generated record is correctly identified as belonging to the target individual, given the background trait information, according to Lemma 6. We also compare the identification capability with different amount of background knowledge, i.e., with the size of trait set  $\mathbf{t}_v$  ranging from one to four.

We run this whole process 10,000 times for each trait set. Figure 3.5a shows the average value of the resulted probabilities. As shown in Figure 3.5a, the green line is the baseline representing the probability 1/100 (100 = 99 'CEU' individuals + 1 target) that the generated record is inferred as belonging to the target individual without any background knowledge. The blue line represents the inferred probability based on the Bayesian network, and the red line represents the inferred



Figure 3.5: (a) Average Probability of Identification; (b) Probability Distribution of Identification

probability using the Humbert's method. The first points in the blue and red lines represent the results given the first trait (according to the trait index in Table 3.5) of the target. Similarly, the second points represent the results given both the first two traits of the target, and so on. The bar at each point shows the standard deviation of the resulting probabilities of 10,000 times of test.

We can see that in general, the probabilities of correctly identifying the target individual of both methods increase as the background knowledge increases, and the identification probability our method is significantly larger than that of the situation without any background knowledge (i.e., 0.01). Comparing the two methods, our method consistently outperforms the Humbert's method (the p-value of the t-test is 0.005). In addition, the identification probability given only one trait of our method is even larger than that given all four traits of the Humbert's method, showing that our method significantly improves the identification accuracy over the Humbert's method.

Figure 3.5b shows the distribution of the inference probability among the 10,000 times of identifications of our method. As the amount of background traits increases, the peaks of the process count would be located at positions with larger identifying probabilities. This indicates that in general, the more background knowledge we have, the more probably that the target individual's record is correctly identified. On the other hand, multiple peaks in each line represent different identifying probabilities due to different combinations of background traits, as well as different possible genotype records being randomly generated.

As an alternative method of defining the target, we leverage the openSNP users as they share both of their trait and genotype profiles online. By blending the profile of an openSNP user into the 'CEU' dataset and re-identifying it, we evaluate the risk of privacy leak of the openSNP users although their profiles are anonymized. One issue here is that the sets of traits and SNPs contained in the GWAS catalog and those contained in openSNP are not identical. In order to perform the attack, we select the target individuals from openSNP who have reported the traits and SNPs which are also contained in the GWAS catalog and openSNP. Among all the identified traits and SNPs, we further require that the odds ratio of the trait-SNP pair to be larger than 2 so that the effect of the SNP on the trait is significant. We have 3 traits and 7 associated SNPs satisfying the requirement, which are shown in Table 3.6. Then, we select the openSNP users who have reported traits. As a result, we obtain a total of 101 openSNP users who are considered as targets in the experiment.

Index	Trait	SNP-risk allele	$f_{kj}^t(r)$	$O_{kj}$	$f_{kj}^c(r)$	$P(t_k)$	
1	Rheumatoid arthritis	rs6457617-T	0.49	2.36	0.69	0.01	
		rs9275406-T	0.17	2.1	0.30	0.01	
2	Hypertriglyceridemia	rs964184-G	0.14	3.28	0.35	0.30	
3		rs3129889-G	0.2	2.97	0.43		
	Multiple colorosis	rs3129934-T	4-T 0.1 2.34	2.34	0.21	0.01	
	Multiple scierosis	rs3135388-A	0.22	2.75	0.44		
		rs9271366-G	0.15	2.78	0.33		

Table 3.6: Trait-SNP association

We compute the probability for each target to be correctly identified from the database. The results for all targets are shown in Figure 3.6a. As can be seen, nearly half (51/101 for our method and 41/101 for the Humbert's method) of the targets have the probability of identification higher than 0.01. In this case, it shows that there is no obvious risk for openSNP users. This is probably due to the difference between the openSNP users and the population represented by the GWAS catalog. However, as shown in Figure 3.6b, if we confine the targets to those who have at least reported the trait of "Hypertriglyceridemia", they have higher chances to be more accurately iden-



Figure 3.6: Probability of identification: (a) all targets; (b) targets with hypertriglyceridemia

tified (19/30 for our method and 13/30 for the Humbert's method). These results show that, for certain openSNP users there are higher risk of privacy leak. Under what circumstance the open-SNP users may face higher risk of privacy leak is worthy of further study. Comparing the two methods, it can be seen that our method still outperforms the Humbert's method in term of the identification accuracy.

# 3.6. Related Work

The detection of SNP-trait associations by building Bayesian networks has been studied in biomedical fields, where a Bayesian network is used to address the high computationally complex and high dimensional problems. In [24] the authors used a score-based Bayesian network structure learning algorithm to detect epistasis or interactions among SNPs. In [12], the same problem is addressed by using a new information-based score and a branch-and-bound search algorithm to discover the structure of the Bayesian network. As an extension to the work of [24], a recent study [60] proposed an exhaustive search on a Bayesian network to detect high order associations of SNPs with traits, without requiring marginal effects on low dimensional datasets. All of the related work aforementioned requires a raw genotype dataset to construct a Bayesian network. Our work is novel in that we build a Bayesian network from the publicly released GWAS statistics where the underlying genotypes are not publicly available. Our method is based on the models of Independence of Causal Influence. ICI is proposed to overcome the problem of specifying a large number of conditional probability distributions in the CPT for a node with multiple parents in the Bayesian network. Examples of widely used ICI models include Noisy-Or, Noisy-Max, Linear-Gaussian, etc. [16]. The Noisy-Max model is equivalent to the Noisy-Or model in our situation where each hidden variable  $X_j$  is binary. The Linear-Gaussian model is proposed for modeling numeric variables. Therefore, these two models are not discussed in this chapter.

Genetic privacy has also been actively studied in the literature (refer to survey papers [7, 39, 48]). For example, Homer et al. developed a method that can identify whether a target with some known SNPs comes from a population with known allele frequency [18]. It attracted more and more attention on the privacy disclosure of the public dissemination of the genotype-related data and aggregate statistics from the genome-wide association studies (GWAS) [21, 36, 46, 47, 53, 55, 64]. Another work [11] showed that full identities of personal genomes can be exposed via surname inference from recreational genetic genealogy databases followed by Internet searches. They considered a scenario in which the genomic data are available with the target's year of birth and state of residency, two identifiers that are not protected by HIPAA. In our previous work [56], we also studied whether and to what extent the unperturbed GWAS statistics can be exploited by attackers to breach the privacy of regular individuals who are not GWAS participants. Two attacks, namely trait inference attack and identify inference attack were formalized based on the 2-layer Bayesian network inference and empirically evaluated. In [49], the authors developed a likelihood-ratio test that uses allele presence or absence responses from a Web service called beacon to derive whether a target individual genome is present in the database. In [45], the authors proposed practical strategies including obscuration and access control for reducing re-identification risks in beacons. In [20], the authors studied the use of phenotypic traits to reidentify users in anonymized genomic databases such as OpenSNP and 23andMe and demonstrated that the privacy risks due to genotype-phenotype associations. In [19], the authors proposed to build a Bayesian network to represent the genotype and phenotype dependencies among

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family members, so that the genotype of a family member can be inferred from the genotypes and phenotypes of his relatives. When the correlation among genotypes are considered, the authors further adopted the factor graph instead of the Bayesian network to represent the familial dependencies.

Several research works [8, 25] have been conducted for the safe release of aggregate GWAS statistics without compromising a participant's privacy. Their ideas were based on differential privacy [5]. Differential privacy is defined as a paradigm of post-processing the output and provides guarantees against arbitrary attacks. A differentially private algorithm provides an assurance that the output cannot be exploited by the attacker to derive whether or not any individual's record is included. The privacy parameter  $\epsilon$  controls the amount by which the distributions induced by two neighboring data sets may differ (smaller values enforce a stronger privacy guarantee). A general method for achieving differential privacy for a query f is to compute the sum of the true output and random noise generated from a Laplace distribution. The magnitude of the noise distribution is determined by the sensitivity of the query and the privacy parameter specified by the data owner. The sensitivity of a computation bounds the possible change in the computation output over any two neighboring data sets (differing at most one record). For example, the sensitivity values of chi-square statistic and p-value were derived in [8]. For those statistics with large sensitivity values (e.g., the sensitivity of odds ratio is infinity), the authors in [8] adapted the idea of releasing the most significant patterns together with their frequencies in the context of frequent pattern mining [1] to release K most significant SNPs. In [25], the authors developed distancescore based privacy preserving algorithms for computing the number and location of SNPs that are significantly associated with the trait, the significance of any statistical test between a given SNP and the trait, correlation between SNPs, and the block structure of correlations. In [51], the authors developed methods for releasing differentially private  $\chi^2$ -statistics in GWAS while guaranteeing membership privacy in adversarial settings [30].

### 3.7. Summary

In this chapter, we first studied the construction of Bayesian networks from publicly released GWAS catalog. We employed the models of independence of causal influences (ICI) which assume that the causal mechanism of each parent variable is mutually independent. We derived a formulation from the Noisy-Or model, one of the ICI models, to specify the CPT using GWAS statistics, and developed a Bayesian Network construction algorithm based on the CPT specification formulation. We proved that, the specified CPT is accurate as long as the underlying individual-level genotype and phenotype profile data follows the Noisy-Or model. In the experiments, we empirically validated the fitness of the Noisy-Or model. Then, we developed three inference problems based on the constructed Bayesian network, namely trait inference given SNP genotype, genotype inference given trait, and trait inference given trait. We developed efficient formulas and algorithms to infer posterior probabilities. Finally, we empirically evaluated the derived inference methods for two applications. In the first application, we showed that significant amount of knowledge regarding traits can be inferred from the genotype profiles. In the second application, we showed that the probability of an individual to be identified from an anonymized genotype database is increasing given some traits of the individual.

Part of the work in this chapter is published in the Proceedings of the 2016 IEEE International Conference on Bioinformatics and Biomedicine [63] and IEEE/ACM Transactions on Computational Biology and Bioinformatics [62].

#### 4. Modeling SNP and Quantitative Trait Association Using CLG Bayesian Network

## 4.1. Introduction

In Chapter 3, we introduce a traditional Bayesian network to deal with the categorical traits. In the GWAS catalog, more than half of the traits are quantitative. The association between SNPs and these quantitative traits cannot be modeled using the previous method due to the mixture of discrete (SNPs and categorical traits) and continuous (quantitative traits) variables.

In this chapter, we address this problem by leveraging the Conditional Linear Gaussian (CLG) Bayesian network [27]. In a CLG Bayesian network, the variables are partitioned into two sets, a set of discrete variables and a set of continuous variables. For each discrete variable, it is associated with a CPT similar to the traditional Bayesian network. For each continuous variable, there is a CLG distribution conditional on each value assignment of its parents. We proposed the method to build the CLG Bayesian network from the GWAS catalog, focusing on the specification of the CLG distribution for quantitative traits. We then develop efficient inference methods based on the constructed network. For simplicity, we only consider quantitative traits in this chapter, but our model can be easily extended to the mixture of categorical and quantitative traits. We discuss the extensions of our model in Section 4.4.

We empirically evaluate the construction and inference methods. We show that useful information regarding the traits and SNPs can be inferred using the constructed network. A case study is also performed to evaluate how much individual information can be disclosed. This can help evaluate the potential risk due to the statistics released in GWAS catalog [20, 56]. We assume an anonymized genotype profile database and a target whose genotype profile is known to be contained in the database. An attacker knows some of traits of the target and aims to identify the target's record from the database. The results show that the attacker can increase his success likelihood by exploiting the traits of the target.

The rest of this chapter is organized as follows. Section 4.2 describes the construction of the CLG

Bayesian network from the GWAS catalog. Followed by that, Section 4.3 presents the inference methods based on the constructed network, and Section 4.4 presents several extensions. The experimental setup and results are discussed in Section 4.5. Section 4.6 summarizes the related work and Section 4.7 concludes this chapter.

#### 4.2. Learning CLG Bayesian Network from GWAS Statistics

In this section, we study the construction of a CLG Bayesian network from the GWAS catalog to model the SNP-quantitative trait association. We first identify the statistical knowledge that can be extracted and derived from the GWAS catalog. Then we show how to learn the CLG Bayesian network using the acquired statistics only. The challenge is to specify all parameters of the CLG distributions as shown in Equation (2.5) without access to the raw data. We make use of the assumption of ICI to facilitate the specification. The ICI assumption has already been shown to have good fitness in modeling SNP-trait associations [63]. Finally, we develop the methods to completely specify the CLG Bayesian network.

### 4.2.1. Knowledge from GWAS catalog

Consider a set of traits  $\mathcal{T}$  and a set of SNPs S in the GWAS catalog. Each trait  $T \in \mathcal{T}$  is a continuous variable whose mean is denoted by  $\mu$  and variance is denoted by  $\sigma^2$ . Assume that the trait is identified to be associated with a subset of m SNPs  $\mathbf{S} = \{S_1, \dots, S_m\} \subseteq S$ . For each SNP  $S_i \in \mathbf{S}$ , denote its genotype as  $s_i = \{0, 1, 2\}$ , where 0 represents two non-risk alleles, 1 represents one non-risk allele and one risk allele, and 2 represents two risk alleles. The association test for trait T and SNP  $S_i$  is performed over a simple of n individuals by fitting the linear regression  $t = \beta_0 + \beta s_i + \varepsilon$ . The values of n,  $\beta$  and the p-value of the ANOVA can be directly extracted from the GWAS catalog.

In addition to the above directly released statistics, we can also derive variance  $\sigma^2$  and parameter  $\beta_0$ , assuming that the values of mean  $\mu$  and the population genotype frequency  $P(s_i)$  can be acquired from the literature or Internet, as shown in Lemma 7. **Lemma 7.** The values of variance  $\sigma^2$  and intercept  $\beta_0$  can be derived as

$$\beta_0 = \mu - \beta \sum_{s_i = \{0, 1, 2\}} P(s_i) s_i,$$

$$\sigma^{2} = \frac{(n-2+F^{*})\left(n_{0}(\beta_{0}-\mu)^{2} + n_{1}(\beta_{0}+\beta_{1}-\mu)^{2} + n_{2}(\beta_{0}+2\beta_{1}-\mu)^{2}\right)}{nF^{*}}.$$

Proof. Straightforwardly we have the following estimation

$$\mu = \mathbb{E}[\hat{T}] = \sum_{s_i = \{0, 1, 2\}} p(s_i)\hat{t} = \sum_{s_i = \{0, 1, 2\}} p(s_i)(\beta_0 + \beta s_i).$$

Thus,  $\beta_0$  can be derived as

$$\beta_0 = \mu - \beta \sum_{s_i = \{0, 1, 2\}} p(s_i) s_i.$$

On the other hand, since the *p*-value is obtained through an *F*-test, ratio  $F^*$  can be recovered using the *p*-value and the F(1, n - 1) distribution. According to definition of SSR, we have

$$SSR = \sum_{k=1}^{n} (\hat{t}_k - \mu)^2 = n_0 (\beta_0 - \mu)^2 + n_1 (\beta_0 + \beta_1 - \mu)^2 + n_2 (\beta_0 + 2\beta_1 - \mu)^2,$$

where  $n_{s_i} = np(s_i)$  is the number of individuals with genotype  $s_i$ . Since  $F^* = \frac{SSR/1}{SSE/(n-2)}$ , we have  $SSE = (n-2)\frac{SSR}{F^*}$ . Meanwhile, the total sum of squares (SSTO) is defined as

$$SSTO = \sum_{k=1}^{n} (t_k - \mu)^2,$$

which can be estimated by  $n\sigma^2$ . In ANOVA, we have

$$SSTO = SSR + SSE.$$

As a result, we obtain

$$\sigma^{2} = \frac{(n-2+F^{*})\left(n_{0}(\beta_{0}-\mu)^{2}+n_{1}(\beta_{0}+\beta_{1}-\mu)^{2}+n_{2}(\beta_{0}+2\beta_{1}-\mu)^{2}\right)}{nF^{*}}.$$

Hence, the lemma is proved. We summarize the acquired knowledge from the GWAS catalog in Lemma 8.

**Lemma 8.** The knowledge that can be obtained from the GWAS catalog includes: a trait set  $\mathcal{T}$  and a SNP set S; for each trait T, its variance  $\sigma^2$  and associated SNPs S; and for each SNP-trait pair, the values of coefficients  $\beta_0$  and  $\beta$ .

4.2.2. CLG Bayesian Network Construction



Figure 4.1: The two-layered CLG Bayesian network

To construct the CLG Bayesian network using the knowledge in Lemma 8, the first step is to build the network structure based on the SNP-trait associations. We adopt a method similar to that in [63] to build a two-layered network, with all SNPs being nodes in the first layer, and all traits being nodes in the second layer. If a SNP is associated with a trait, then we add an edge pointing from the SNP to the trait. Since the SNP-SNP association cannot be acquired from the GWAS catalog, we assume no edge among SNPs, which means that all SNPs are mutually independent in the network. Similarly, we also assume no edge among traits, which means that there is no direct correlation between traits. However, two traits can still be correlated via SNPs if they share the same SNPs. We will discuss how to relax these assumptions in Section 4.4. As a result, we obtain a special CLG Bayesian network where the continuous variables (traits) only have discrete parents (SNPs). The structure of the network is illustrated in Figure 4.1.

The next step is to specify the conditional probability distribution for each SNP, and the CLG distribution for each trait. For a SNP *S*, the conditional probability distribution is directly specified by the genotype frequency p(s), which can be acquired from the literature or Internet as discussed in the previous subsection. For a trait *T*, following Equation (2.5), the CLG distribution given *T*'s associated SNPs **S** is given by

$$P(t|\mathbf{s}) = \mathcal{N}(t; a(\mathbf{s}), c(\mathbf{s})). \tag{4.1}$$

Note that the above equation does not include parameters  $b(\cdot)$  since the trait has no continuous parent.

Parameters  $a(\cdot)$  and  $c(\cdot)$  capture the joint effect of all associated SNPs on the trait. However, the GWAS statistics shown in Lemma 8 only provide the association between *T* and each of its associated SNP. In addition, the number of parameters in  $a(\cdot)$  and  $c(\cdot)$  is exponential to the size of **S**, making them impractical to be specified in a direct manner. Since it has already been shown in [63] that the assumption of ICI has a good fitness in modeling the SNP-trait association, we also adopt the ICI assumption to facilitate the parameter estimation in CLG distribution specification.

ICI assumes that, when there are multiple parents, the causal influence of each parent on the child is mutually independent, so that the combined influence of all parents is decomposable into a number of independent influence of each parent. The most commonly used ICI model when the child is a continuous variable is the Noisy-Add model [13], which represents the combined effect of all parents as a linear combination of the effect of each parent. According to the Noisy-Add model, the mean of T can be represented by a linear combination of association SNPs **S**, i.e.,  $a(\mathbf{s}) = a_0 + \sum_{i=1}^m a_i s_i$ . For *T*'s variance  $c(\cdot)$ , we simply assume that it is a constant *c* for all different **s**, i.e.,  $c(\mathbf{s}) = c$ . As a result, we denote the CLG distribution by a Gaussian distribution with mean  $\mu_{t|\mathbf{s}} = a_0 + \sum_{i=1}^m a_i s_i$  and variance  $\sigma_{t|\mathbf{s}}^2 = c$ . Parameters  $a_i$  ( $i = 0, \dots, m$ ) and *c* can be estimated from the GWAS statistics according to Lemma 9.

**Lemma 9.** Given GWAS statistics in Lemma 8, we generate following data points  $(y, x_1, \dots, x_m)$  in space  $\mathbb{R}^{m+1}$ :

$$\forall S_j \in \mathbf{S}, s_j = \{0, 1, 2\}, \qquad \begin{pmatrix} y = \beta_0 + \beta s_j \\ x_j = s_j \\ \forall i \neq j, \ x_i = \sum_{S_i} P(s_i) s_i \end{pmatrix}$$

Parameters  $a_i$  ( $i = 0, \dots, m$ ) can be estimated using the linear regression

$$y = a_0 + a_1 x_1 + \dots + a_m x_m + \varepsilon,$$

which aims to fit with above data points, and parameter c can be estimated by formula

$$\frac{1}{m}\sum_{j=1}^{m}\left(\sigma^{2}-\sum_{S_{j}}P(s_{j})\left(A(s_{j})+\mu_{t|s_{j}}^{2}-2\mu_{t|s_{j}}\mu+\mu^{2}\right)\right),$$

where  $A(s_j) = \sum_{\mathbf{S} \setminus \{S_j\}} \prod_{S_i \in \mathbf{S} \setminus \{S_j\}} P(s_i) \left( \mu_{t|\mathbf{s}}^2 - 2\mu_{t|s_j} \mu_{t|\mathbf{s}} + \mu_{t|s_j}^2 \right), \ \mu_{t|s_j} = \beta_0 + \beta s_j \ and \ \mu_{t|\mathbf{s}} = a_0 + \sum_{i=1}^m a_i s_i.$ 

*Proof.* Consider the conditional distribution of *T* given each parental SNP *S*<sub>j</sub>, and denote its mean and variance by  $\mu_{t|s_j}$  and  $\sigma_{t|s_j}^2$ . These values can be estimated using the CLG distribution, and meanwhile, they can also be estimated using the GWAS statistics. The goal is to find the parameters  $a_i$  and *c* that best fit the estimations from the CLG distribution to the estimations from the GWAS statistics.

For  $\mu_{t|s_j}$ , consider the conditional distribution  $P(t|s_j)$  obtained by marginalizing the CLG distribution, i.e.,

$$P(t|s_j) = \sum_{\mathbf{S} \setminus \{S_j\}} P(\mathbf{s} \setminus \{s_j\}) P(t|\mathbf{s}) = \sum_{\mathbf{S} \setminus \{S_j\}} \prod_{S_i \in \mathbf{S} \setminus \{S_j\}} P(s_i) P(t|\mathbf{s}).$$

Straightforwardly,  $\mu_{t|s_j}$  can be estimated from the mean of the above distribution, given by

$$\mathbb{E}[T|s_j] = \sum_{\mathbf{S} \setminus \{S_j\}} \prod_{S_i \in \mathbf{S} \setminus \{S_j\}} P(s_i) \left( a_0 + \sum_{i=1}^m a_i s_i \right),$$

which equals to

$$a_0 + s_j a_j + \sum_{i \neq j} \left( \sum_{S_i} P(s_i) s_i \right) a_i.$$

On the other hand, the estimation of  $\mu_{t|s_j}$  from the GWAS statistics is given by  $\beta_0 + \beta s_j$ . Thus, we consider the following linear regression:

$$y = a_0 + a_1 x_1 + \dots + a_m x_m + \varepsilon.$$

For each value assignment  $s_j$  of each SNP  $S_j$ , we have a data point  $(y, x_1, \dots, x_m)$  where  $y = \beta_0 + \beta s_j$ ,  $x_j = s_j$ , and  $x_i = \sum_{s_i} P(s_i) s_i$  for  $i \neq j$ . As a result, we have 3m data points in space  $\mathbb{R}^{m+1}$ , which can be used to train the above linear regression model and learn the parameters  $a_i$   $(i = 0, \dots, m)$ .

For  $\sigma_{t|s_j}^2$ , the estimation can be obtained from the variance of distribution  $P(t|s_j)$ , which is given by

$$\mathbb{E}[(T - \mu_{t|s_j})^2 | s_j] = \sum_{\mathbf{S} \setminus \{S_j\}} P(\mathbf{s} \setminus \{s_j\}) \mathbb{E}[(T - \mu_{t|s_j})^2 | s_j, \mathbf{s} \setminus \{s_j\}]$$
$$= \sum_{\mathbf{S} \setminus \{S_j\}} \prod_{s_i \in \mathbf{S} \setminus \{S_j\}} P(s_i) \mathbb{E}[(T - \mu_{t|s_j})^2 | \mathbf{s}].$$

We have

$$\mathbb{E}[(T - \mu_{t|s_j})^2 |\mathbf{s}] = \mathbb{E}[T^2 - 2T\mu_{t|s_j} + \mu_{t|s_j}^2 |\mathbf{s}]$$
  
=  $\mathbb{E}[T^2 |\mathbf{s}] - 2\mu_{t|s_j}\mu_{t|\mathbf{s}} + \mu_{t|s_j}^2,$  (4.2)

as well as

$$c = \mathbb{E}[(T - \mu_{t|\mathbf{s}}])^2 |\mathbf{s}] = \mathbb{E}[T^2 - 2T\mu_{t|\mathbf{s}} + \mu_{t|\mathbf{s}}^2 |\mathbf{s}]$$
  
=  $\mathbb{E}[T^2 |\mathbf{s}] - \mu_{t|\mathbf{s}}^2,$  (4.3)

where  $\mu_{t|s_j} = \beta_0 + \beta s_j$  and  $\mu_{t|s} = a_0 + \sum_{i=1}^m a_i s_i$ . Combining Equations (4.2) and (4.3), we have

$$\mathbb{E}[(T - \mu_{t|s_j})^2 | \mathbf{s}] = c + \mu_{t|\mathbf{s}}^2 - 2\mu_{t|s_j}\mu_{t|\mathbf{s}} + \mu_{t|s_j}^2.$$

As a result, we have

$$\mathbb{E}[(T - \mu_{t|s_{j}})^{2}|s_{j}] = \sum_{\mathbf{S} \setminus \{S_{j}\}} \prod_{S_{i} \in \mathbf{S} \setminus \{S_{j}\}} P(s_{i}) \left(c + \mu_{t|\mathbf{s}}^{2} - 2\mu_{t|s_{j}}\mu_{t|\mathbf{s}} + \mu_{t|s_{j}}^{2}\right)$$

$$= c + \sum_{\mathbf{S} \setminus \{S_{j}\}} \prod_{S_{i} \in \mathbf{S} \setminus \{S_{j}\}} P(s_{i}) \left(\mu_{t|\mathbf{s}}^{2} - 2\mu_{t|s_{j}}\mu_{t|\mathbf{s}} + \mu_{t|s_{j}}^{2}\right) = c + A(s_{j}).$$
(4.4)

Using similar strategy, we can also obtain

$$\mathbb{E}[(T-\mu)^2] = \sum_{s_j} p(s_j) \left( \mathbb{E}[(T-\mu_{t|s_j})^2 | s_j] + \mu_{t|s_j}^2 - 2\mu_{t|s_j}\mu + \mu^2 \right).$$
(4.5)

Combining Equations (4.4) and (4.5), we have

$$c = \mathbb{E}[(T-\mu)^2] - \sum_{s_j} P(s_j) \left( A(s_j) + \mu_{t|s_j}^2 - 2\mu_{t|s_j}\mu + \mu^2 \right).$$

The value of  $\mathbb{E}[(T - \mu)^2]$  is directly estimated by  $\sigma^2$ . Thus, the value of c is estimated by

$$\sigma^2 - \sum_{s_j} P(s_j) \left( A(s_j) + \mu_{t|s_j}^2 - 2\mu_{t|s_j} \mu + \mu^2 \right).$$

Since for each  $S_j$  we have the above estimation, the value of c that best fits the estimation from the CLG distribution and the estimation from the GWAS is the average of above estimation over

all SNPs in **S**, given by

$$\frac{1}{m} \sum_{j=1}^{m} \left( \sigma^2 - \sum_{S_j} P(s_j) \left( A(s_j) + \mu_{t|s_j}^2 - 2\mu_{t|s_j} \mu + \mu^2 \right) \right).$$

Based on the above analysis, the CLG distribution is specified as shown in the following theorem.

**Theorem 4.** The CLG distribution of trait T given its associated SNPs  $\mathbf{S} = \{S_1, \dots, S_m\}$  is denoted by a Gaussian distribution with mean  $\mu_{t|\mathbf{s}} = a_0 + \sum_{i=1}^m a_i s_i$  and variance  $\sigma_{t|\mathbf{s}}^2 = c$ , i.e.,

$$P(t|\mathbf{s}) = \mathcal{N}(t; a_0 + \sum_{i=1}^m a_i s_i, c),$$

where parameters  $a_i$  ( $i = 0, \dots, m$ ) and c are estimated according to Lemma 9.

### 4.3. Inference Based on the Constructed Bayesian Network

Using the CLG Bayesian network constructed from the GWAS catalog, we can calculate the conditional probability density for any desired value assignment to sets of SNPs  $S_x$  and traits  $T_x$ , given the observed value assignment to the disjoint sets of SNPs  $S_y$  and traits  $T_y$ . In the following we first develop the theorem to calculate the joint probability density for any value assignment to the SNPs and traits, as shown in Theorem 5.

**Theorem 5.** The joint probability density for any value assignment to subsets  $S \subseteq S$ ,  $T \subseteq T$  is given by

$$P(\mathbf{s},\mathbf{t}) = \sum_{\mathbf{S}_3} \prod_{s_j \in \mathbf{S}_1 \cup \mathbf{S}_2 \cup \mathbf{S}_3} P(s_j) \cdot \mathcal{N}(\mathbf{t};\boldsymbol{\mu},\boldsymbol{\Sigma}),$$

where  $S_1$  denotes the SNPs in S but not associated with T,  $S_2$  denotes the SNPs in S and are also associated with T,  $S_3$  denotes the SNPs associated with T but not in S;  $\mathcal{N}(t; \mu, \Sigma)$  is a multivariate Gaussian distribution with mean vector  $\mu$  and covariance matrix  $\Sigma$ , where

$$\boldsymbol{\mu} = \begin{pmatrix} \mu_{T_1|pa(T_1)} \\ \vdots \\ \mu_{T_n|pa(T_n)} \end{pmatrix}, \quad \boldsymbol{\Sigma} = \begin{pmatrix} \sigma_{T_1|pa(T_1)}^2 & \cdots & 0 \\ \vdots & \vdots & \ddots \\ 0 & \cdots & \sigma_{T_n|pa(T_n)}^2 \end{pmatrix}.$$

*Proof.* Divide S into four disjoint subsets:  $S_1, S_2, S_3$  denote the SNPs as described in the theorem, and  $S_4$  denotes all the other SNPs. Note that  $S = S_1 \cup S_2$ , and  $\bar{S} = S \setminus S = S_3 \cup S_4$ . Divide  $\mathcal{T}$ into two subsets: **T** and its complement  $\bar{T}$ . According to Equation (2.6), we have

$$P(\mathcal{S},\mathcal{T}) = P(\mathbf{s}_1,\mathbf{s}_2,\mathbf{s}_3,\mathbf{s}_4,\mathbf{t},\mathbf{\bar{t}}) = \prod_{s_j\in\mathcal{S}} P(s_j)\cdot\prod_{T_i\in\mathcal{T}} P(t_i|pa(T_i)).$$

According to Equation (2.7), joint probability density  $P(\mathbf{s}, \mathbf{t})$  is given by

$$P(\mathbf{s},\mathbf{t}) = \sum_{\bar{\mathbf{s}}} \prod_{S_j \in S} P(s_j) \cdot \int_{\bar{\mathbf{T}}} \prod_{T_i \in \mathcal{T}} P(t_i | pa(T_i)).$$

It can be straightforwardly derived that

$$\sum_{\mathbf{S}} \prod_{S_j \in S} P(s_j) = \sum_{\mathbf{S}_3, \mathbf{S}_4} \prod_{S_j \in \mathbf{S}_1 \cup \mathbf{S}_2 \cup \mathbf{S}_3 \cup \mathbf{S}_4} P(s_j)$$
$$= \sum_{\mathbf{S}_3} \prod_{S_j \in \mathbf{S}_1 \cup \mathbf{S}_2 \cup \mathbf{S}_3} P(s_j) \sum_{\mathbf{S}_4} \prod_{S_j \in \mathbf{S}_4} P(s_j) = \sum_{\mathbf{S}_3} \prod_{S_j \in \mathbf{S}_1 \cup \mathbf{S}_2 \cup \mathbf{S}_3} P(s_j),$$

and

$$\begin{split} \int_{\bar{\mathbf{T}}} \prod_{T_i \in \mathcal{T}} P(t_i | pa(T_i)) &= \prod_{T_i \in \bar{\mathbf{T}}} P(t_i | pa(T_i)) \int_{\bar{\mathbf{T}}} \prod_{T_i \in \bar{\mathbf{T}}} P(t_i | pa(T_i)) \\ &= \prod_{T_i \in \bar{\mathbf{T}}} P(t_i | pa(T_i)). \end{split}$$

Due to the Markov condition, CLG distributions  $p(t_i|pa(T_i))$  are mutually independent. This means that the product is also a Gaussian distribution, where the mean vector and covariance ma-

Based on Theorem 5, we provide the general inference formula as shown in Theorem 6.

**Theorem 6.** The probability for any desired value assignment to subsets  $S_x$ ,  $T_x$ , given the observed value assignment to disjoint subsets  $S_y$ ,  $T_y$ , is given by

$$P(\mathbf{s}_x, \mathbf{t}_x | \mathbf{s}_y, \mathbf{t}_y) = \frac{P(\mathbf{s}_x, \mathbf{s}_y, \mathbf{t}_x, \mathbf{t}_y)}{P(\mathbf{s}_y, \mathbf{t}_y)},$$

where  $P(\mathbf{s}_x, \mathbf{s}_y, \mathbf{t}_x, \mathbf{t}_y)$  and  $P(\mathbf{s}_y, \mathbf{t}_y)$  are calculated according to Theorem 5.

There can be many applications of the inference. One typical application is to evaluate the health risk, implied by a subset of traits, given the genotype profile and possibly a subset of other traits of the user. This means to calculate posterior distribution  $P(\mathbf{t}_x|\mathbf{s}_y)$  or  $P(\mathbf{t}_x|\mathbf{t}_y, \mathbf{s}_y)$ . For example, the bone mineral density is known to be associated with several SNPs. Hence, we can predict the bone mineral density in a certain period of lifetime based on the SNPs and some other traits such as physical activities the target has. Another application can be to infer the genotype profile of a user given an incomplete genotype record, i.e., to calculate  $P(\mathbf{s}_x|\mathbf{s}_y)$ . By doing so, the user can infer the genotype profile rather than doing another costly genotype scan. As another example, the CLG Bayesian network can be used to infer the genotype profile of a target based on the traits the target has, i.e., to calculate  $P(\mathbf{s}_x|\mathbf{t}_y)$ . This inference can be used to evaluate the potential privacy risk due to the statistics released in the GWAS catalog. We will evaluate the third application in the experiments.

## 4.4. Further Extensions

So far, we make several assumptions in order to simplify the representation and ensure that the CLG Bayesian network can be built from the GWAS catalog only. In this section, we briefly discuss the relaxation of the assumptions and the extensions of our model to these relaxed situations.

First, since the GWAS catalog does not include the correlations between SNPs, we assume no edge among SNPs in the network. This assumption can be relaxed if additional knowledge is available from other sources. For example, assume that we know the joint genotype frequency of all correlated SNPs, and we also know the direction of the influence between each pair of correlated SNPs. Then we can easily incorporate the SNP-SNP correlations into our CLG Bayesian network. For each correlated SNP pair, we simply add an edge between the two SNPs according to the direction of the influence. The CPTs of SNPs can be specified based on the joint genotype frequencies. On the other hand, if only SNP-SNP correlations without the direction exist, we can merge the correlated SNPs as a single super node, representing and taking values from the combination of the SNP genotypes. We add an edge pointing from the super node to each associated traits. After that, similar to Theorems 5 and 6 all inferences can be conducted.

Second, we assume no edge among traits. The assumption may also be relaxed with additional knowledge, where the network structure can be adapted based on trait-trait correlations. The only limitation here is that, we don't allow the edge to point from a quantitative trait to a categorical trait, since the CLG Bayesian network assumes that discrete variables only have discrete parents. As a result, we obtain a general CLG Bayesian network where each continuous variable can have both discrete and continuous parents. In this case, the general belief propagation algorithms such as [4, 29, 35] can be used for performing inference on the network.

## 4.5. Experiments

In this section, we first discuss our experimental setup and the CLG Bayesian network construction in Section 4.5.1. The inference methods are evaluated in Section 4.5.2. And finally, we conduct a case study to evaluate the individual information that can be inferred using the constructed network.

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### 4.5.1. CLG Bayesian Network Construction

We use data from the GWAS catalog of 5/24/2017. This version of the GWAS catalog includes 38,037 records (SNP-trait association pairs) extracted from 2,468 publications. Out these records, there are a total of 28,943 SNPs associated with a total of 1,864 traits. For each record, the risk allele type, the *p*-value, the sample description, and the odds ratio or  $\beta$  coefficient are provided. Note that the odds ratio or the  $\beta$  coefficient is provided in the same field, depending on whether the trait is categorical or quantitative.

In the experiments, we only consider the quantitative traits and focus on a subset of data used as an interactive diagram (https://www.ebi.ac.uk/gwas/diagram) published by the GWAS catalog. In this subset of data, an additional field "orType" is used to clearly indicate whether the odds ratio is provided and hence indicate whether the trait is categorical or quantitative. This subset of data includes 5,893 records which contains 809 traits and 4,643 SNPs. We extract the records with "orType = false", i.e., the trait is quantitative. As a result, we obtain 484 traits associated with 2,768 SNPs which are contained in 3,557 records. Finally, we build a knowledge database about the traits and the associated SNPs, including the risk allele type,  $\beta$  coefficient, the *p*-value, and the sample description for each trait-SNP association pair.

Index	Trait	$a_0$	с	SNP-risk allele	$a_i$	$p(s_i)$
1	PCA3 expression level	15.599	26.661	rs10993994-T	1.250	{0.23, 0.50, 0.27}
2	Head circumference	47.307	0.535	rs7980687-A	0.074	{0.67, 0.30, 0.03}
2	ricau circulificience			rs1042725-T	-0.065	{0.17, 0.49, 0.34}
3	Fat body mass	24.621	0.576	rs6567160-C	0.090	{0.60, 0.35, 0.05}
4	Iron levels	18.794	0.013	rs228916-T	-0.086	{0.00, 0.14, 0.86}
5 T	Thyroid-stimulating hormone levels	2.047	0.684	rs12126655-G	0.138	$\{0.45, 0.44, 0.11\}$
				rs11026407-C	-0.125	{0.31, 0.49, 0.20}

Table 4.1: Snapshot of CLG Bayesian network

Based on the knowledge database, the CLG Bayesian network has been built according to Section 4.2. Table 4.1 shows the information and statistics of a snapshot of the network. There are 5 traits and 7 associated SNPs in the snapshot, reported in 4 GWAS publications. For each trait, parameters  $a_0$  and c are shown in Columns 3 and 4. Column 5 shows each associated SNP of the trait,

along with the parameter  $a_i$  and the genotype frequencies  $P(s_i) = \{P(s_i = 0), P(s_i = 1), P(s_i = 2)\}$  as shown in Columns 6 and 7.



## 4.5.2. Inference Evaluation

Figure 4.2: Posteriori probability densities of traits.

We evaluate the inference methods by considering two situations. In the first situation, we calculate the CLG distribution of each trait given all SNPs, i.e., P(t|S) for each trait T, which is equivalent to P(t|pa(T)). The results for the traits in Table 4.1 are shown in Figure 4.2, where each subfigure corresponds to the trait with the same index in the table. In the subfigure, each dashed line represents the CLG distribution given one value assignment to the associated SNPs. We also plot the marginal distribution of the trait by marginalizing all CLG distributions, represented by the solid line. It can be seen that, for traits such as PCA3 expression level and Iron level, there are obvious differences in the marginal distribution and the CLG distributions, while for traits such as Fat body mass, the difference is very small. The difference in the distributions shows the difference in the strength of the influence of the associated SNPs on the trait.

In the second situation, we calculate the genotype frequencies of a subset of SNPs given the values of a subset of traits, i.e.,  $P(\mathbf{s}|\mathbf{t})$ . We set the values of traits to three levels: in the first level, all

traits are set to their means; in the second level, all traits are set to their means plus their standard deviations; and in the third level, all traits are set to their means plus two times of their standard deviations. The results for the traits and SNPs in Table 4.1 are shown in Figure 4.3, where each subfigure corresponds to one trait setting level. We can see that the distribution of each SNP changes with the increase of the traits correspondingly to the sign of  $a_i$ .

### 4.5.3. Application: Identity Inference

In this subsection, we consider an application where the CLG Bayesian network is used by an attacker in identifying the identity information of a target individual from a genotype database. Specifically, the attacker has access to an anonymized genotype database  $\mathcal{R}$  which contains the target's genotype record  $\mathbf{s}_{v}$ . The attacker also knows a subset of traits  $\mathbf{t}_{v}$  the target has. Then, the attacker can learn the posteriori probability of each genotype record  $\mathbf{s}_{i}$  in the database given the traits, i.e.,  $P(\mathbf{s}_{i}|\mathbf{t}_{v})$ , and use them to improve the identification success likelihood. Specifically, the improved probability that the target's record  $\mathbf{s}_{v}$  is identified is given by  $\frac{P(\mathbf{s}_{v}|\mathbf{t}_{v})}{\sum_{i=1}^{|\mathbf{r}|} P(\mathbf{s}_{i}|\mathbf{t}_{v})}$ . We evaluate the performance of the attack by comparing the identification probability with that of the random guess which uses no background knowledge.

We extract a genotype profile dataset from the 1000 Genomes Project [50] referred to as 'CEU', which is used to serve as the anonymized genotype database. The 'CEU' dataset consists of 99



Figure 4.3: Posteriori probability distribution of SNPs.
HapMap individuals from Utah residents with Northern and Western European ancestry. For the target, we manually generate his genotype and phenotype profiles at random. We assume that the target has the minor traits, which means that the value of each trait is less than  $\mu - \sigma$  or larger than  $\mu + \sigma$ . Thus, for each trait *T*, we randomly generate its value *t* according to the density *P*(*t*) with the constraint  $t < \mu - \sigma$  or  $t > \mu + \sigma$ . After all traits **t** are generated, for each SNP *S*, we randomly generate its value *s* according to the probability *P*(*s*|**t**). In this way, we simulate a scenario where the target is randomly picked from the population with minor traits. After that, we blend the genotype profile of the target into the 'CEU' dataset which contains the genotype records of 99 individuals, and attempt to re-identify it with the target's traits just like what an attacker does. We run this process 1,000 times, and evaluate the average probability of the target's record to be identified. We also compare the probability with different amount of knowledge, i.e., with the size of trait set **t**<sub>v</sub> increasing from one to five.



Figure 4.4: Probability of identification.

The results are shown in Figure 4.4. The green dashed line is the baseline representing the identification probability 1/100 without any background knowledge. The red solid line represents the average identification probability of our method along with the 15% and 85% percentiles. As can be seen, the identification probability generally increases as the background knowledge increases. The average identification probability of our method is larger than that of the random guess without any background knowledge, implying the potential risk in privacy leak due to the statistics released in GWAS catalog.

## 4.6. Related Work

Detecting and analyzing SNP-trait associations by building Bayesian networks has been studied in biomedical fields. In [24] the authors used a score-based Bayesian network structure learning algorithm to detect epistasis or interactions among SNPs. In [12], the same problem was addressed by using a new information-based score and a branch-and-bound search algorithm to discover the structure of the Bayesian network. As an extension to the work of [24], in [60] the authors proposed an exhaustive search on a Bayesian network to detect high order associations of SNPs with traits, without requiring marginal effects on low dimensional datasets. All of the related work aforementioned requires a raw genotype-phenotype dataset to construct a Bayesian network. However, the raw data required may not always be available. Different from above works, in [56, 63], the authors studied the construction of Bayesian network from the released GWAS statistics only. The two-layered Bayesian network was built and inference was conducted using the constructed network. However, these works can only deal with the categorical traits. Our work in this chapter uses the CLG Bayesian network to deal with the quantitative traits and is readily to be extended to the mixture of categorical and quantitative traits.

Similar to the work in [63], we adopt the assumption of Independence of Causal Influence in the specification of the CLG Bayesian network. The ICI is proposed to overcome the problem of specifying a large number of conditional probability distributions for a dependent node with multiple parents in the Bayesian network. Examples of widely used ICI models include Noisy-Or, Sigmoid, Noisy-Max, and Noisy-Add [16]. The first three models target the situation where the dependent node is a discrete variable, while the Noisy-Add model targets the situation where the dependent node is a continuous variable. In our case, only the trait can be the dependent node in the network. Thus, the Noisy-Add model is applied in this work.

# 4.7. Summary

In this chapter, we studied the exploration and modeling of associations between SNPs and quantitative traits from the GWAS catalog. The CLG Bayesian network which can deal with a mixture of discrete and continuous variables is employed. We developed the method of constructing the CLG Bayesian network only using the released statistics from the GWAS catalog, as well as inference methods for calculating posterior distributions. We empirically evaluate the construction and inference methods. The results show that important information can be inferred using the constructed network. The work in this chapter is published in the Proceedings of the 2017 IEEE International Conference on Bioinformatics and Biomedicine [61].

#### 5. STIP: A SNP-Trait Inference Platform

## 5.1. Introduction

In the previous two chapters, we have proposed methods to build Bayesian networks using only GWAS statistics instead of raw genotype data for characterizing SNP-trait associations and then developed efficient formulas and algorithms to conduct SNP-trait inference based on the Bayesian networks. In particular, we have proposed a three-layered traditional Bayesian network to model the SNP-categorical trait associations in Chapter 3 and a two-layered Conditional Linear Gaussian (CLG) Bayesian network to model the SNP-quantitative trait associations in Chapter 4.

In this chapter, we present STIP, a web-based platform that aims to aid common users in SNPtrait inference and GWAS catalog exploration. STIP is based on Bayesian networks proposed in Chapter 3 and 4 [61, 63]. Using STIP, users can easily infer SNPs/traits given their known traits/SNPs and explore the SNP-trait associations. In particular, STIP provides three services: 1) *SNP-trait inference* allows users to conduct any kind of SNP-trait inference task, such as trait inference given SNP genotypes or genotype inference given traits; 2) *Top-k trait prediction* shows the probabilities of having some traits based on the user's genotype profile; and 3) *GWAS catalog exploration* helps users explore the SNP-trait associations.

Currently, there are some tools for GWA studies, like PLINK[44], GCTA[59]. However, only a few systems have been developed to perform the SNP-trait inference. For example, hapassoc [2] is an R package for likelihood inference of trait associations with SNP haplotypes. CGBayesNets[37] is a MATLAB package that can build predictive models of a phenotype of interest using multimodal genomic data as possible predictors. The existing packages require raw genotypes of SNPs, yet such information is not publicly available. In this regard, all the data acquired by STIP is from the GWAS catalog which can be accessed directly online. Without using sensitive geno-type data, STIP can provide the SNP-trait inference service to public. Meanwhile, STIP can han-dle both categorical and quantitative traits reported in the GWAS catalog, while previous plat-

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forms only focus on one type of traits. Furthermore, unlike those packages that are designed for genomic researchers, STIP has user friendly interface for general end users.

# 5.2. STIP Overview

STIP is a web-based platform for SNP-trait inference and GWAS catalog exploration. Figure 5.1 shows the architecture of STIP. Users can visit our platform through the URL: http://csce. uark.edu/~xintaowu/STIP.htm. Currently, STIP supports three services: (1) SNP-trait inference allows users to infer SNPs/traits by providing their known SNPs/traits. (2) Top-k trait prediction helps users discover the risk of having a certain disease or identify significant changes of quantitative traits based on their genotype profile. (3) GWAS catalog exploration allows users to explore the SNP-trait associations especially when users find some unfamiliar SNPs or traits from the previous two services. The back end of our platform is two Bayesian networks which are derived from the GWAS catalog. All inference-related services are based on the constructed Bayesian networks.



Figure 5.1: The architecture of STIP

# 5.2.1. Bayesian Network Construction

There are two types of traits in the GWAS catalog, categorical traits  $\mathcal{T}^c$  and quantitative traits  $\mathcal{T}^q$ . For each type of trait, we construct a Bayesian network to model the relationships among traits and SNPs.

## The construction of the three-layered Bayesian network for modeling SNP-categorical traits

For categorical traits, a three-layered Bayesian network is constructed from the GWAS catalog, which is designed to represent the conditional dependencies between categorical traits and SNPs. The Bayesian network is consisted of three layers, the SNP genotype layer, the SNP allele layer, and the categorical trait layer, from top to bottom. Please refer to Chapter 3 for detailed descriptions.

To implement the Bayesian network, we adopt a subset of traits used as an interactive diagram published by the GWAS catalog (https://www.ebi.ac.uk/gwas/diagram). In this subset of data, an additional field "orType" is used to clearly indicate whether the odds ratio is provided. If the value of "orType" is "True", it indicates the trait is categorical. Otherwise, the trait is quantitative.

After obtaining a categorical trait set  $\mathcal{T}^c$ , we then extract a set of SNPs S given those traits. For each specific trait  $T_k^c \in \mathcal{T}^c$ , we have a subset of associated SNPs  $S_k$ . For each associated SNP  $S_{kj} \in S_k$ , we can extract its corresponding risk allele type  $r_{kj}$  associated trait  $T_k^c$ , the odds ratio of the association test, and the risk allele frequency in the control group. If there exist multiple entries for one SNP-trait pair, we only keep one SNP-trait association entry with the smallest pvalue. We further delete entries which do not contain risk allele, risk allele frequency in control group, or odds ratio because those statistics are required in the three-layered Bayesian network construction. To acquire the prior probability of each categorical trait, we classify all the categorical traits into 17 categories (e.g., immune system disease, nervous system disease, etc.), and retrieve the average prevalence of each category from the Wikipedia. We use the average prevalence of a category as the prior probability of each trait belonging to the category. Our threelayered Bayesian network can be refined by assigning the accurate prior probability for each trait when available.

The three layer Bayesian network is constructed from the GWAS catalog. Because the GWAS

catalog extracts information from literature, one SNP-trait association may exist in the GWAS catalog with multiple entries each of which may be submitted by one GWAS study and may have different statistics values. For these duplicated entries, we only keep one SNP-trait association entry with the smallest *p*-value. We further delete entries which do not contain risk allele, risk allele frequency in control group, or odds ratio because those statistics are required in the three layer Bayesian network construction.

Currently, STIP contains 265 categorical traits and their 2319 associated SNPs. Thus, the SNP genotype layer, the SNP allele layer, and the trait layer in the three-layered Bayesian network contain 2319, 2319, and 265 nodes, respectively. In particular, since each SNP has 2 alleles (e.g., A/G) and 3 genotypes (e.g., AA/AG/GG), each node in the SNP genotype layer has 3 values and each node in the SNP allele layer has 2 values. STIP saves the statistics of each SNP-categorical trait association in a JSON file. For each SNP-categorical trait association, the allele prior probabilities, the genotype prior probabilities, the risk allele frequency of case and control groups and the trait prior probability are included in the JSON file. The other conditional probabilities for SNP-trait inference can be easily derived from the existing information when a user submits a specific inference request. Figure 5.2a shows a code snippet about a SNP-categorical trait association in the JSON file. In this example, the trait "Chronic kidney disease" is associated with one SNP "rs6066043". The allele and the genotype prior probabilities of that SNP are included in this SNP-trait association. If a SNP is associated with multiple traits, those prior probabilities will be included in all the related SNP-trait associations. This redundancy design improves the runtime speed of the SNP-trait inference because the associated SNPs and their statistics can be directly acquired by given a trait.

# The construction of the CLG Bayesian network for modeling SNP-quantitative traits

We build a CLG Bayesian network for modeling SNP-quantitative trait associations. The CLG Bayesian network contains two layers that are SNP genotype layer and quantitative trait layer. In the CLG Bayesian network, the continuous variables (traits) only have discrete parents (SNPs).

Please refer to Chapter 4 for detailed descriptions.

Similar to the previous subsection, to implement the CLG Bayesian network, we obtain the following data from the GWAS catalog: a trait set  $\mathcal{T}^q$  and a SNP set  $\mathcal{S}$ ; for each trait  $T_k^q \in \mathcal{T}^q$ , its associated SNPs  $\mathbf{S}_k$ ; and for each SNP-trait pair, the coefficient  $\beta$ , *p*-value, and the sample size of the specific study.

Currently, STIP contains 484 quantitative traits and their 2768 associated SNPs. Thus, the CLG Bayesian network contains 484 nodes in the trait layer and 2768 nodes in the SNP genotype layer. We use another JSON file to store the information of each SNP-quantitative trait association. For each SNP-quantitative trait association, the genotype prior probabilities, trait distribution ( $\mu$  and  $\sigma$ ), and CLG distribution (coefficients and  $\sigma$ ) are included in the JSON file. Figure 5.2b shows a code snippet about a SNP-quantitative trait association in the JSON file. In this example, the trait "PCA3 expression level" is associated with one SNP "rs10993994".





(b) SNP-quantitative trait

Figure 5.2: The information of SNP-trait associations stored in JSON files.

#### 5.2.2. Services Provided by STIP

**SNP-trait inference:** The SNP-trait inference is the core service provided by STIP, which reports the posterior probability of having any specific SNPs or traits given the already known SNPs or traits. For example, we can calculate the posterior probability that a user has specific genotypes for a set of SNPs given the user's trait information.

In principle, we calculate the posterior probability for any *desired* assignment of values to variable sets  $\mathbf{S}_x^g$ ,  $\mathbf{T}_x$  given the observed assignment of variable sets  $\mathbf{S}_y^g$ ,  $\mathbf{T}_y$ , as shown in Equation (5.1). Note that  $\mathbf{S}_x^g$  and  $\mathbf{S}_y^g$  denote the set of SNP genotypes; while  $\mathbf{T}_x$ ,  $\mathbf{T}_y$  denote the set of traits.

$$P(\mathbf{s}_x^g, \mathbf{t}_x | \mathbf{s}_y^g, \mathbf{t}_y) = \frac{P(\mathbf{s}_x^g, \mathbf{t}_x, \mathbf{s}_y^g, \mathbf{t}_y)}{P(\mathbf{s}_y^g, \mathbf{t}_y)}.$$
(5.1)

Equation (5.1) indicates that we can derive the posterior probability distribution of one or more variables in the Bayesian network given the values observed for other variables in the network. Meanwhile, Equation (5.1) shows that to conduct inference, we first need to calculate the joint probability for any desired assignment of values to variable sets  $S^g$  of SNPs S and traits T, which reflects the relationship among SNPs and traits.

For categorical traits, the joint probability for any value assignment to  $S^g$  of  $S \subseteq S$ ,  $T^c \subseteq T^c$ , i.e.,  $P(s^g, t^c)$ , based on the three-layered Bayesian network is defined as

$$P(\mathbf{s}^{g},\mathbf{t}^{c}) = \prod_{S_{j}\in\mathbf{S}_{1}} P(s_{j}^{g}) \sum_{\mathbf{S}_{2}^{a},\mathbf{S}_{3}^{g},\mathbf{S}_{3}^{a}} \Big(\prod_{S_{j}\in\mathbf{S}_{2}\cup\mathbf{S}_{3}} P(s_{j}^{g})P(s_{j}^{a}|s_{j}^{g}) \prod_{T_{k}^{c}\in\mathbf{T}^{c}} P(t_{k}^{c}|pa(T_{k}^{c}))\Big),$$
(5.2)

where  $S_1$  denotes the SNPs in S but not associated with  $T^c$ ,  $S_2$  denotes the SNPs in S and also associated with  $T^c$ ,  $S_3$  denotes the SNPs associated with  $T^c$  but not in S.

For quantitative traits, the joint probability density for any value assignment to subsets  $S^g$  of  $S \subseteq$ 

 $\mathcal{S}, \mathbf{T}^q \subseteq \mathcal{T}^q$  is given by

$$P(\mathbf{s}^{g}, \mathbf{t}^{q}) = \sum_{\mathbf{S}_{3}} \prod_{S_{j}^{g} \in \mathbf{S}_{1} \cup \mathbf{S}_{2} \cup \mathbf{S}_{3}} P(S_{j}^{g}) \cdot \mathcal{N}(\mathbf{t}^{q}; \boldsymbol{\mu}, \boldsymbol{\Sigma}),$$
(5.3)

where  $S_1$  denotes the SNPs in S but not associated with  $T^q$ ;  $S_2$  denotes the SNPs in S and are also associated with  $T^q$ ;  $S_3$  denotes the SNPs associated with  $T^q$  but not in S;  $\mathcal{N}(t; \mu, \Sigma)$  is a multivariate Gaussian distribution with mean vector  $\mu$  and covariance matrix  $\Sigma$ , where

$$\boldsymbol{\mu} = \begin{pmatrix} \mu_{t_1^q \mid pa(T_1^q)} \\ \vdots \\ \mu_{t_n^q \mid pa(T_n^q)} \end{pmatrix}, \quad \boldsymbol{\Sigma} = \begin{pmatrix} \sigma_{t_1^q \mid pa(T_1^q)}^2 & \cdots & 0 \\ \vdots & \vdots & \ddots \\ 0 & \cdots & \sigma_{t_n^q \mid pa(T_n^q)}^2 \end{pmatrix}.$$

All terms in Equations (5.2) and (5.3) are completely specified in the two Bayesian networks.

**Top-k trait prediction:** The top-k trait prediction is a special case of SNP-trait inference, i.e., trait inference given SNP genotypes.

For categorical traits, this service lists the top-k traits of a user that have the most significant increase from the trait prior probabilities to the trait posterior probabilities by inferring from the user's genotype profile. Formally we represent the genotypes of a user as  $\mathbf{s}^g = (s_1^g, s_2^g, \dots, s_n^g)$ , with each entry  $s_j^g$  indicating the genotype of SNP *j*. The trait inference given SNP genotypes aims to learn the posterior probability  $P(t^c = 1|\mathbf{s}^g)$  that a user has a specific trait  $t^c$  given the target's genotype profile  $\mathbf{s}^g$  using the Bayesian network. The posterior probability  $P(t^c = 1|\mathbf{s}^g)$  can be calculated based on Equation (5.1) with  $\mathbf{s}_x^g = \emptyset$ ,  $\mathbf{t}_y^c = \emptyset$ ,  $\mathbf{t}_x^c = \{t\}$ , and  $\mathbf{s}_y^g = \mathbf{s}^g$ . After calculating the posterior probability  $P(t^c = 1|\mathbf{s}^g)$ , we further compute the change value for each trait  $L_{t^c} = P(t^c = 1|\mathbf{s}^g) - P(t^c = 1)$  and show traits with the top-k largest  $L_{t^c}$ .

*For quantitative traits*, this service lists the top-k traits of a user that have the most significant change from the mean value of the trait  $\mathbb{E}(t^q)$  to the mean value of the trait given the user's genotype profile  $\mathbb{E}(t^q | \mathbf{s}^g)$  based on the CLG Bayesian network. The mean value of the trait can be ac-

quired from the Internet or literature. The posterior mean value of the trait is defined as

$$\mathbb{E}[t^{q}|\mathbf{s}^{g}] = a_{0} + \sum_{i=1}^{m} a_{i}s_{i},$$
(5.4)

where  $a_i$   $(i = 0; \dots; m)$  are the CLG distribution coefficients of the associated SNPs given the trait  $t^q$ . After calculating  $\mathbb{E}(t^q | \mathbf{s}^g)$ , we further compute the change value for each trait  $L_{t^q} =$  $|\mathbb{E}(t^q | \mathbf{s}^g) - \mathbb{E}(t^q)|$  and show traits with the top-k largest  $L_{t^q}$ .

Figure 5.3 shows the procedures of using SNP-trait inference and Top-k trait prediction services. For SNP-trait inference, the known SNPs/traits are first specified and then STIP will report the probabilities of unknown SNPs/traits based on the Bayesian network. For Top-k trait prediction, users first submit their genetic file. STIP will report the probabilities or mean values of traits given the genotype profile.

**GWAS catalog exploration:** The third service provided by STIP is the GWAS catalog exploration. Although the GWAS catalog website allows users to search SNPs and traits, the search results are organized by the studies and do not clearly show the SNP-trait associations. Unlike the exploration provided by the GWAS catalog website, STIP provides trait-SNP and SNP-trait association explorations which allow users to choose a trait (SNP) and then show the associated SNPs (traits).



Figure 5.3: The procedures of SNP-trait inference and Top-k trait prediction

## 5.3. Demonstration

Since STIP is a web-based platform, users can visit STIP through an Internet browser. In this section, we illustrate the use of the three services provided by STIP.



Figure 5.4: SNP-categorical trait inference web UI with an example of trait inference given SNP genotypes and categorical traits

# 5.3.1. SNP-Trait Inference

Users can perform SNP-trait inferences on the "SNP-Categorical Trait Inference" and "SNP-Quantitative Trait Inference" pages. Users can first choose the known traits from the trait list and the known SNPs from the SNP list. Due to the large number of SNPs and traits, STIP provides search bars for both lists which allow users to search a specific SNP and trait. Users can choose multiple traits or SNPs at a time. Once users have selected the known traits and SNPs, they can further click the "Known Traits" and "Known SNPs" buttons. The selected traits and SNPs will be added to the "Known Traits ( $T_y$ )" and "Known SNPs ( $S_y$ )" lists on the right side of the page. In particular, for a categorical trait  $T_y^c \in \{Y, N\}$ , Y indicates a user with the trait and N indicates the user without the trait. For a quantitative trait, the default value of  $T_y^q \in (-\infty, \infty)$ , users can specify the range of the  $T_y^q$  based on their knowledge. Users can also specify the genotypes of each SNP. Then, users can further select the inference traits and SNPs by clicking the "Inference Traits" and "Inference SNPs". Finally, users can click the "Inference" button to get the result of the specified inference. Figure 5.4 shows an example of SNP-categorical trait inference. In this example, STIP reports the probability of a user with the trait "Psoriasis" by knowing that the user has two traits and three SNPs. The "SNP-Quantitative Trait Inference" has a similar UI which allows user to specify the range of a quantitative trait. Note that both pages provides general SNPtrait inference services which allow users to do any kind of inference without the need to specify all the lists on the right side of the pages. For example, if a user only knows some of the traits and wants to know the risk of having other traits, the user can just specify the "Known Traits ( $T_y$ )" and "Inference Traits ( $T_x$ )" and then runs the inference.

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op K Categorical Traits	Top K Quantitative Traits	All Traits		
	Results of top	K: 10 • traits with larg	gest probability changes	
Traits		Prior Probability	Posterior Probability	Difference
Celiac disease		0.26	0.3536	0.0936
Behcet's disease		0.04	0.094	0.054
Vascular dementia		0.3	0.3325	0.0325
Alopecia areata		0.09	0.1131	0.0231
Chronic kidney disease		0.09	0.1102	0.0202
Chronic hepatitis C infection		0.09	0.109	0.019
Galistones		0.26	0.275	0.015
Wegener's granulomatosis		0.04	0.0508	0.0108
Sagittal craniosynostosis		0.09	0.0987	0.0087
Nevirapine-induced rash		0.004	0.0100	0.0033

(a) Top-10 trait prediction

op K Trait Prediction					
Choose File usb_genotyping_file to: Submit O Currently only support the penotyping file from 23andble.					
Top K Categorical Traits Top K Quantitative Traits All Traits					
Celiac disease	Trait : Chronic kidney disease				
Vascular dementia	Prior Probability : 0.09 Posterior Probability : 0.1102				
Chronic kidney disease	Difference : 0.0202				
Chronic logantin C anfertion     Galinotos     Galinotos     Wengens's parallementosis     Sagintal emaioymotosis     Nervirguine noficoed rath     Epirobecin-induced logalopenia     Epirobecin-induced logalopenia     Engenda systeme	Associate 8399 : • m89804-0-:05				
Enteric fever     Testicular germ cell tumor     Non-alcoholic fanty liver disease     Lung cancer					



Figure 5.5: Top-k trait prediction web UIs with results of trait prediction given a user's genetic file

# 5.3.2. Top-K Trait Prediction

The "Top-K Trait Prediction" allows a user to submit his/her genetic file and reports the probabilities (mean values) of traits based on the genotype profile. Currently, STIP supports the genetic file provided by 23andme (https://www.23andme.com/). Figure 5.5 shows the result page after a user submits the genetic file. There are three tabs in the "Top-K Trait Prediction" page. The first one is the "Top-K Categorical Traits" tab (shown in Figure 5.5a) which shows top-k categorical traits with the largest increase values from trait prior probabilities to posterior probabilities given SNPs. The second one is the "Top-K Quantitative Traits" tab which shows top-k quantitative traits with the largest change values from trait prior mean values to posterior mean values. The last one is "All Traits" tab (shown in Figure 5.5b) which lists all the traits associated with the user's genotypes. For each categorical (quantitative) trait, STIP reports the trait prior and posterior probabilities (mean values), the change value between the trait prior and posterior probabilities (mean values), the associated SNPs, and the genotypes of the user.





(a) Exploring SNP-Trait associations

(b) Exploring Trait-SNP associations

Figure 5.6: GWAS catalog exploration web UIs

# 5.3.3. GWAS Catalog Exploration

STIP provides two separated pages for SNP-trait and trait-SNP association explorations. Figure 5.6a shows the SNP-trait association exploration where a user can explore the associated traits by providing a specific SNP. In this scenario, once a user chooses a SNP, the SNP-trait association page shows the number of associated traits and lists all the information about those traits. Figure 5.6b shows the trait-SNP association exploration where a user can explore the associated SNPs by specifying a trait.

# 5.4. Summary

We present STIP, a web-based platform that provides the SNP-trait inference and GWAS-catalog exploration. The SNP-trait inference is designed to infer the probabilities of unknown traits/SNPs by given known traits/SNPs. STIP also allows users to submit their genetic file and reports the risks of having specific traits based on their genotype profile. The GWAS-catalog exploration is designed to explore the SNP-trait associations. The work in this chapter is published in the Proceedings of the 2017 IEEE International Conference on Bioinformatics and Biomedicine [42].

## 6. Conclusions and Future Work

## 6.1. Conclusions

In this thesis, we studied the construction of the Bayesian networks from the public released GWAS catalog for modeling SNP and trait associations. Bayesian network has been proposed as a powerful tool for modeling SNP-trait associations. Most of existing works learn the Bayesian network from the raw data. However, due to privacy issue, genotype information is classified as sensitive which should be dealt with by complying with specific restrictions. To tackle this limitation, we proposed to build Bayesian networks from the GWAS catalog. The GWAS catalog is a database that collects and publicly releases literature-derived GWAS statistics, including pairwise SNP-trait associations, risk allele frequency, p-value, etc. In particular, since there are two types of traits (i.e., categorical trait and quantitative trait), we develop two types of Bayesian networks correspondingly to model the SNP-trait associations from the GWAS statistics.

First, we developed a method of building a classic three-layered Bayesian network using only GWAS statistics for characterizing SNP-categorical trait associations. In order to construct a Bayesian network from only statistics, we derived a formulation based on the Noisy-Or model, one best known example of the independence of causal influence (ICI) models, that can be used to specify the conditional probability table (CPT) from the released GWAS statistics where the underlying genotypes can be unknown. We theoretically and empirically validated the fitness of using the Noisy-Or model to specify CPT. Based on the constructed Bayesian network, we further conducted inference tasks including trait inference given SNP genotype, genotype inference given trait, and trait inference given trait. We showed that significant amount of knowledge regarding traits can be inferred from the genotype profiles and the probability of an individual to be identified from an anonymized genotype database was increasing given some traits of the individual.

Second, we employed a CLG Bayesian network which can deal with a mixture of discrete and

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continuous variables to model SNP-quantitative trait associations from GWAS statistics. Sepcifically, SNPs are represented as discrete variables, and the quantitative traits are represented as continuous variables. We then developed the inference methods to conduct SNP-quantitative trait inference tasks. The empirical results showed the effectiveness of our methods.

Finally, we developed STIP, a web-based platform that provided the SNP-trait inference and GWAS-catalog exploration. The SNP trait inference was designed to infer the probabilities of unknown traits/SNPs by given known traits/SNPs. STIP also allowed users to submit their genetic file and reported the risks of having specific traits based on their genotype profile.

## 6.2. Future Work

Our work can be further improved in the following directions:

Firstly, for simplicity we assume all traits are quantitative in the CLG Bayesian network. Since the CLG Bayesian network is able to deal with both continuous and categorical data, our model is readily to handle the mixture of categorical and quantitative traits. In the future, we plan to model the both categorical and quantitative traits in one unified CLG Bayesian network. In this case, besides the genotype frequencies and CLG distributions for quantitative traits, we also need to specify the CPTs for the categorical traits. This can be done by directly adopting the method proposed in [63]. After the network construction completes, all inferences can be performed similarly as shown in Theorems 5 and 6.

Second, we also plan to integrate the SNP-SNP correlations and trait-trait correlations into our model if additional knowledge is available from other sources.

Finally, we plan to develop methods to enable researchers to safely release aggregate GWAS data without compromising the anonymity of both GWAS participants and the general population.

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