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Statistical Methods in Integrative Genomics

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Abstract

Statistical methods in integrative genomics aim to answer important biology questions by jointly analyzing multiple types of genomic data (vertical integration) or aggregating the same type of data across multiple studies (horizontal integration). In this article, we introduce different types of genomic data and data resources, and then review statistical methods of integrative genomics, with emphasis on the motivation and rationale of these methods. We conclude with some summary points and future research directions.

Keywords

genomics; integrative genomics; horizontal data integration; vertical data integration

1. INTRODUCTION

It is an exciting time to work on statistical methods for genomic problems. The rapid development of high-throughput techniques allows researchers to collect large amounts of genomic data, which can answer more biological questions and enable the development of more effective therapeutic strategies for human diseases. Since multiple types of genomic data are often available within and across studies, the integrated analysis of genomic data has becomes popular. One may integrate the same type of genomic data across multiple studies (horizontal integration), or integrate different types of genomic data in the same set of samples (vertical integration). We will review both horizontal and vertical integration studies, while putting more emphasis on the latter. Before discussing statistical methods, we will give a brief review of different types of genomic data as well as resources on where to

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1.1. Different Types of Genomic Data

1.1.1. DNA—A common genomic analysis is to study DNA features from germline (normal) tissue and tumor tissue separately, as they often have very different characteristics. DNA variants from germline tissue include single nucleotide polymorphisms (SNPs), indels (short insertion or deletion), copy number variation (CNV), and other structural changes such as translocations. SNP arrays can provide high confidence genotype estimates because the underlying genotypes belong to one of three classes: AA, AB, and BB, where A and B indicate the two alleles of a SNP. Most SNP arrays are designed to target common variants (e.g. those DNA variants that occur in more than 1% of individuals in a population). Both Array CGH and SNP array can measure CNVs. While array CGH can only measure total copy number, SNP arrays can measure the copy number in each of two homologous chromosomes, i.e., allele-specific copy number (Wang et al. 2007; Sun et al. 2009). Recently, high-throughput sequencing, including whole genome sequencing or exome sequencing (exome-seq) has been used to study DNA variants. With sufficient read-depth, sequencing data can provide more accurate estimates of SNP genotypes and copy number calls. In addition, sequencing data can detect rare mutations (i.e., the mutations with low population frequencies) that are usually not captured by arrays (Nielsen et al. 2011; Mills et al. 2011).

In cancer studies, we are often interested in somatic DNA mutations that occur in tumor tissues but not found in the germline. Somatic point mutations, including single nucleotide changes and indels, are often rare. It is likely that two cancer patients share few or no somatic point mutations across the whole exonic regions. In this sense, cancer may be better considered as a collection of rare diseases rather than one disease. Due to such rareness, somatic point mutations are usually detected by sequencing. A somatic copy number aberration (SCNA) often occupies a relatively long genomic region (e.g., one-third of a chromosome may be deleted or amplified), and can be relatively common. SCNAs can be studied by either array CGH, SNP array, or by high throughput sequencing. Studying somatic DNA mutations (either point mutations or SCNAs) is challenging because tumor samples are often composed of a mixture of tumor and normal cells (e.g., the normal cells from connective tissues or blood vessels) and tumor cells may have more than or less than 2 copies of DNA on average. These two issues are known as purity and ploidy issues. Unknown purity and ploidy affect each other and should be estimated together (Van Loo et al. 2010; Carter et al. 2012). In addition, recent sequencing studies have revealed that tumor cell populations may be composed of several subclones. Some somatic mutations may only occur in one or some of the subclones, and thus have low allele frequencies. It is challenging to distinguish such mutations from sequencing errors (Ding et al. 2014).

 1.1.2. Epigenetic Marks—Normal cells within the human body share almost identical DNA, with sporadic somatic mutations contributing a small amount of variation between cells. Despite this similarity, different types of cells are observed to have dramatically different sizes, shapes, and/or functions. Such cell-type-specific traits are often maintained

by epigenetic marks, which are modifications on DNA molecules or proteins that can be passed to daughter cells during mitosis. The term "epi" means "over, outside of, around" in Greek. Although epigenetic marks do not change the DNA sequence itself, they may also be inheritable, and their role in the etiology of human diseases are increasingly recognized (Jiang, Bressler & Beaudet 2004). We will introduce three types of epigenetic marks: open chromatin regions, histone modifications, and DNA methylation.

Within a cell nucleus, DNA is packed around multiple proteins called histones. This complex of DNA and proteins is referred to as chromatin. Chromatin usually takes a condensed form so that the packed DNA sequence is not accessible by other proteins, such as regulatory transcription factors. Open chromatin regions, where previously packed DNA sequence is loosened and exposed, often harbor active regulatory elements bound to DNA. Open chromatin regions can be detected by DNase I hypersensitive sites (DHSs) sequencing (DNase-seq), where DNA sequences on DHSs are captured and then located by high-throughput sequencing techniques (Figure 1) (Song & Crawford 2010).

Histone modifications include different types of chemical modifications (e.g., methylation, acetylation, or phosphorylation) on different amino acids of histone proteins. Chromatin immunopreciptation (ChIP) followed by microarray (ChIP-chip) or sequencing (ChIPseq) are popular choices to capture DNA sequence associated with modified histones (Figure 1). The ChIP step enriches for such DNA sequences, and the following microarray or sequencing step determines their likely genomic location. Each type of histone modification may occur on short/long genomic regions, and is associated certain biological features e.g., active promoters or genes with suppressed expression (ENCODE Consortium 2012; Rashid, Sun & Ibrahim 2014).

DNA methylation usually refers to the addition of a methyl group to cytosine residues within CpG dinucleotides. In the human genome, there are approximately 28 million CpG sites, which are not uniformly distributed. Clusters of CpG sites $(a.k.a. CpG$ islands) tend to occur on gene promotors (Stirzaker et al. 2014). DNA methylation on promoter regions usually represses gene expression; in contrast, DNA methylation in genic or exonic regions is often positively associated with gene expression. Popular techniques to measure DNA methylation including array-based methods (e.g., Infinium HumanMethylation450 Bead-Chip (HM450)), whole genome bisulfite sequencing (WGBS), and reduced representation bisulfite sequencing (RRBS) (Figure 1). From the HM450 array, two measurements are obtained for a CpG locus, reflecting methylation (M) and unmethylation (U) signals, respectively. A commonly used measurement of methylation is referred to as beta-value, which equals to $M/(M + U)$ (See Figure 1 for examples). Using WGBS, one can count the number of sequence reads with methylated or unmethylated CpG's, where methylated CpG's are marked by bisulfite transformation. Although RRBS covers less than 5% of CpG's genomewide $(\sim 1$ million of the 28 million CpG sites), its coverage is enriched for CpG's at promotor regions (~ 0.5 million of 2 million CpG sites on promoters) (Stirzaker et al. 2014).

1.1.3. RNA—Three types of RNA molecules are commonly encountered in genomic data: messenger RNA (mRNA) which encode proteins, and two non-coding RNAs with regulatory roles: microRNA (miRNA) and long non-coding RNA (lncRNA). In fact, the field

has also gradually recognized miRNA as one epigenetic machinery (Malumbres 2013). Expression (of any type of RNA) has traditionally been studied by different types of microarrays, where the expression of one gene/RNA may be measured by one or more microarray probes. In recent years, RNA-seq has been replacing microarrays to become the major platform of transcriptomic studies. Compared with microarrays, RNA-seq provides more accurate estimates of gene expression, allows de novo discovery of transcripts, and delivers new information such as allele-specific expression and RNA isoform-specific expression. Recent studies have systematically evaluated different RNA-seq protocols and paved the way for future large scale RNA-seq studies (Kratz & Carninci 2014). Using RNA-seq data, the expression of one gene could be quantified by the number of RNA-seq fragments mapped to this gene, after correcting for read-depth and gene length.

1.1.4. Protein—Proteins perform many fundamental functions within living organisms and understanding their abundance or activity is biologically important. The protein expression is, however, often less studied, mostly because amino acids do not form double helix structure as in nucleotides and the amplification and hybridization techniques used in microarray and sequencing for DNA and RNA are not conveniently applicable to proteins.. The activity of a protein may depend on a specific set of post-translational modifications (PTM). There are over 200 types of PTMs that may occur in multiple positions of a protein, and thus the combinations of such PTMs lead to an enormous number of protein states that cannot be handled by any current technology. A particular form of PTM, phosphorylation, has been better studied because phosphorylated proteins (phosphoproteins) play important roles in signaling pathways and assays are available to measure phosphoprotein abundance in large scale (Terfve et al. 2012). Existing techniques for proteomics study can be classified into two classes. One is antibody-based array and the other is mass spectrometry (MS). The Cancer Genome Atlas (TCGA) project has measured expression more than 100 proteins or phosphoproteins across thousands of cancer patients using an antibody-based array named reverse phase protein array (RPPA). Traditional gene expression array measures genome-wide expression of one sample on an array. In contrast, each spot in a RPPA corresponds to a sample, and a RPPA measures the expression of one protein or phosphoprotein across all the samples spotted on this array. Therefore the output of RPPA is comparable for one protein across all the samples, but in general not comparable for multiple proteins of one sample. Proteomics is a fast-growing field. Several large scale proteomics projects are ongoing, e.g., The Clinical Proteomic Tumor Analysis Consortium (Ellis et al. 2013).

1.2. Genomic Data Resources

There are huge amount of publicly available genomic data deposited in different databases (Table 1). The results of Genome-Wide Association Studies (GWAS), including DNA genotype and phenotype data, are often deposited at dbGAP, which is hosted by National Center for Biotechnology Information (NCBI). Since genotyping information can theoretically trace to patient identity, one needs to complete a secure access application through dbGAP to protect privacy of patients. Gene expression and epigenetic data are often deposited at NCBI GEO (Gene Expression Omnibus) or ArrayExpress. The NCBI SRA

There are also grsowing data resources from large consortium projects. A widely cited example is The Cancer Genome Atlas (TCGA). The TCGA data portal allows users to directly download open access data, which includes de-identified data of clinical and demographic features, mRNA or microRNA expression, copy number alterations, DNA methylation, and protein or phosphoprotein abundance (Figure 2). The primary sequence data and genotype data belong to controlled access data, which can be downloaded from CGHub (Cancer Genomics Hub). The International Cancer Genome Consortium (ICGC) is another large consortium that also collects genomic data from different types of cancers. A few other notable genomic data resources include the Roadmap Epigenomics Project, which focuses on genome-wide epigenetic marks; Genotype-Tissue Expression project (GTEx), which produces RNA-seq data from different human tissues, and ENCODE (Encyclopedia Of DNA Elements) project, which aims to study all functional elements in the human genome sequence.

Although lots of datasets are freely available for academic use, effort is needed to become familiar with different data resources and make correct use of them. Many datasets are publicly available but are not well-annotated, making them difficult to use. Databases with standardized uploading protocols are typically easier to use. For example, GEO adopts the MIAME standard and has volunteer personnel to constantly check data quality when new datasets are uploaded. Furthermore, sequencing or genotyping data of human samples often involve issues regarding privacy and legal consent, and thus their datasets need protection through protocols such as dbGAP. The administrative burden to access such data is usually not negligible and should be considered when using these datasets.

1.3. Genomic Annotation Databases

Genomic annotations, such as locations and functions of genomic features, are valuable knowledge to assist analysis of any genomic study. Due to limitations of space, we only provide a brief review of selected annotation databases (Table 2). Arguably, the most important annotation for most genomic studies is the reference genome. The most recent release of human reference genome is GRCh38.p2 (released on December 8, 2014 by Genome Reference Consortium), which is the second patch release for the GRCh38 reference assembly. Reference genomes can be accessed online at Ensembl or the UCSC Genome Browser, among other online locations. At the DNA level, NCBI dbSNP provides a comprehensive annotation of known SNPs, taken from various sequencing/genotyping projects such as the 1000 Genomes Project. Current version of dbSNP (Human build 142) has a total of 112 million reference SNPs (refSNPs). Gene structure annotation includes the location of a gene and its exons, as well as its transcripts (i.e., RNA isoforms). Ensembl's Genebuild pipeline automatically annotates genes based on existing evidence of mRNA and proteins in public scientific databases (Curwen et al. 2004). The GENCODE annotation combines the automatic annotation from Ensembl and manual annotation from the HAVANA (Human and Vertebrate Analysis and Annotation) team (Harrow et al. 2012). The functional annotation of each gene is incorporated by many gene-centered databases

such as NCBI Entrez Gene database. Gene Ontology (GO) database provide standardized ontology terms for gene functions in three categories: biological process, molecular function, and cellular component (Ashburner et al. 2000). There are also many databases for pathway annotations such as KEGG and NCI Pathway Interaction Database. Pathway Commons provide a centralized location to store pathway information from multiple databases. Many annotation databases can be conveniently found in the "Annotation" category of Bioconductor, a comprehensive collection of bioinformatics tools on the R language platform. There are also numerous useful databases to systematically catalog existing biological findings such as GWAS Catalog (disease association findings), COSMIC (mutations and gene translocation), miRanda (miRNA target genes), Genomics of Drug Sensitivity in Cancer (GDSC, for drug response in cancer), MIPS (protein-protein interactions), and Transfac (transcription factor binding motifs), just to name a few.

2. Horizontal Data Integration

With rapidly accumulated GWAS, gene expression and methylation studies, and often limited sample size in each study, applications and development of meta-analysis methods to increase statistical power and achieve a consensus conclusion have significantly grown and evolved in the past decade. The ultimate goal is usually to improve detection of differentially expressed genes, disease associated SNPs or differentially methylated sites. Due to the large^p-small-n nature of omics datasets (also partly because Microsoft Excel could only accommodate at most 256 columns in the 90's), samples are usually arranged on the columns and gene features (SNPs, gene symbols or methylation sites) are on the rows, a reversed convention than general statistical practices. As a result, when multiple GWAS or transcriptomic studies are combined for meta-analysis, the datasets are laid out horizontally with gene features matched on the rows. As a result, such multi-study data integration is often called "horizontal genomic meta-analysis" and is the focus of this section. In contrast, when multiple omics datasets of the same cohort of samples are combined, the datasets are aligned vertically with samples matched on the columns. The data integration is called "vertical genomic integrative analysis" (Section 3). For horizontal meta-analysis, interested readers may refer to the following publications for details: GWAS meta-analysis review papers (Thompson, Attia & Minelli 2011; Begum et al. 2012; Evangelou & Ioannidis 2013), microarray meta-analysis review papers (Ramasamy et al. 2008; Tseng, Ghosh & Feingold 2012) and relevant comparative studies (Wang et al. 2013; Chang et al. 2013). Below we mainly focus on GWAS and transcriptomic meta-analysis to illustrate the basic principles and common issues, as well as discussing related challenges and opportunities.

2.1. Data collection and preprocessing

Researchers first determine a systematic search and inclusion/exclusion criteria to identify, extract, annotate and prepare datasets for meta-analysis. This process may involve special data management consideration and tedious preprocessing protocol. In GWAS metaanalysis, for example, raw genotyping data are usually not allowed to share without patient consent. The GWAS meta-analysis consortium usually needs to develop a rigorous data exchange protocol to determine sharing of clinical information and summary statistics (e.g. effect size and its standard deviation) for millions of SNPs for meta-analysis. For

transcriptomic meta-analysis, determining whether studies have similar underlying biological comparison suitable for meta-analysis is critical. After data preparation, methods may be applied to ensure quality control for meta-analysis (Kang et al. 2012).

2.2. Statistical methods for meta-analysis

Many traditional meta-analysis methods have been applied to genomic applications. These include two major categories: combine p-values and combine effect sizes. In the first category, Fisher's method and Stouffer's method are probably the most popular. Methods taking the minimum and maximum p-values have been used. In the second category, fixed, random or mixed effects models are popular. In transcriptomic meta-analysis, nonparametric methods based on ranks have also been developed (Hong et al. 2006).

2.3. Targeted biological objectives and underlying hypothesis setting

An important prerequisite decision behind genomic meta-analysis is to determine the targeted biological objective and the corresponding hypothesis setting. Tseng, Ghosh & Feingold (2012) demonstrated two hypothesis settings (HS_A and HS_B) to detect biomarkers differentially expressed (or SNPs associated to disease) in "all studies" or "one or more studies", respectively. Although HS_A is more often the desired biological objective, HS_B can be considered when study heterogeneity is expected and of research interest (e.g. when studies utilize different tissues; see next paragraph). These two hypothesis settings are closely related to traditional union-intersection test (UIT) and intersection-union test (IUT), and choosing a hypothesis setting affects the selection of a suitable meta-analysis method. For example, Fisher's method combines p-values by summation of log-transformed pvalues. One sufficiently small p-value is enough to generate statistical significance and thus, the method is for HS_B (IUT). Chang et al. (2013) conducted a comprehensive comparative study to compare different methods for transcriptomic meta-analysis according to different hypothesis settings. Song $\&$ Tseng (2014) discussed a robust hypothesis setting to relax HS_A from the stringent requirement of differential expression in "all studies" to "most studies" and proposed a solution by order statistics.

2.4. Cross-study heterogeneity

Genomic studies often contain heterogeneity across studies due to different cohorts, experimental protocols, platforms or tissues used to generate the data. Although the main purpose of meta-analysis is to combine consensus information to improve statistical power, the heterogeneities across studies are also often of importance. For example, if different tissues are applied in different studies, tissue-specific biomarkers are expected and of concern. In the HS_B (IUT) hypothesis setting described above, adaptively weighted concept (a.k.a. subset-based approach) and meta-lasso approach have recently been developed to identify gene-specific subset of studies that contain differential expression (Li 2011; Li et al. 2014) or disease association information (Bhattacharjee et al. 2012; Han & Eskin 2012). The result characterizes both homogeneous and heterogeneous signals across studies.

2.5. Horizontal meta-analysis for purposes other than biomarker detection

The discussion so far focuses on meta-analysis of multiple genomic datasets to improve biomarker detection. Beyond biomarker detection, the concept of horizontal meta-analysis can be extended to virtually any statistical learning area that has been developed and applied to a single high-throughput experimental dataset. In transcriptomic analysis, for example, gene set analysis (a.k.a. pathway analysis) is a popular and powerful tool to characterize biological pathways associated with the disease or condition contrast (Khatri, Sirota & Butte 2012; Newton & Wang 2015). The methods can be extended towards a meta-analytic setting to improve statistical power and to reach a more consensus conclusion (Shen & Tseng 2010). Other statistical learning areas such as dimension reduction, clustering, classification and network analysis may also take advantage of combining information from multiple studies to improve performance and leave many open and challenging opportunities in the field.

3. Vertical Data Integration

With vertical integration, we are concerned with multiple data types *on the same set of* samples. Following the fundamental principle of systems biology that biological mechanisms are built upon multiple molecular phenomena acting at different levels, we aim to gain understanding of complex phenotypic traits by *jointly analyzing* different layers of genomic information. Vertical integration tasks are directly linked to the type of biological questions that are posed, and the methods used are correspondingly and extremely varied. Main distinguishing characteristics are whether the biological question is focused on predictive, regressive (supervised) or exploratory (unsupervised) aims, and how prior biological information is utilized within the statistical analysis. Most approaches to vertical integration are model-based and we shall focus our review on these. None model-based approaches will be briefly discussed when we describe examples of analysis strategies aimed at answering specific biological questions (see Section 3.4).

3.1. Integrative clustering

Many diseases exhibit substantial heterogeneity with respect to biological characteristics and clinical outcomes. Motivated by the known influence of genetic aberrations (germline and somatic) and the accessibility of tumor samples, early work focused almost exclusively on cancer and on using simple hierarchical clustering of gene expression profiles to uncover cancer subtypes. The landmark papers by Golub et al. (1999) on leukemia and by Perou et al. (2000) on breast cancer were followed by numerous studies endeavoring to describe molecular subtypes for a large variety of cancers, with only a few studies related to noncancer pathologies, such as myopathies (Greenberg et al. 2002) or auto-immune diseases (Lee et al. 2011). Cancer genomes exhibit considerable heterogeneity with abnormalities occurring in different genes among different individuals, posing a great challenge to identify those genes with functional importance and therapeutic implications. Hence, to go beyond a straightforward catalog and to provide deeper biological insight and clinical significance, it became apparent that additional biological information should be incorporated into the clustering process; and new robust clustering strategies that would integrate simultaneously diverse genomic characteristics were warranted.

3.1.1. iCluster—An important step in this direction was made by Shen, Olshen & Ladanyi (2009) who proposed **iCluster**, an integrative clustering approach to infer latent subtypes based on multiple genomic data types measured on the same samples. They achieve this by specifying a latent model for each data type X_j (where each dataset is row centered):

$$
\mathbf{X}_j = \mathbf{W}_j \mathbf{Z} + \varepsilon_j \quad j = 1, \dots, m \quad (1)
$$

where $\mathbf{Z} = (\mathbf{z}_1, ..., \mathbf{z}_{N-1})$ are the latent subtypes *common* to the *m* data types and \mathbf{W}_j are the coefficient the latent subspaces. The independent error terms ($\varepsilon_1, \ldots, \varepsilon_m$), with diagonal covariance matrix, represent the residual variance in each dataset after accounting for the correlation between data types. In order to derive a computationally efficient procedure for evaluating the likelihood in (1), a Gaussian latent variable model representation, based on a continuous parametrization **Z*** of **Z**, is used. An EM iterative algorithm is employed to derive the reduced representation $\hat{E}(\mathbf{Z}^*|\mathbf{X})$. Additional lasso-type penalties are imposed on the factor scores **W** to obtain a sparse solution and pinpoint important features contributing to the clustering. Finally, the class indicators **Z** are recovered using a K-means procedure on $\hat{E}(Z^*|X)$ to derive N clusters. Model choice for the lasso penalty and the number of clusters is based on an empirical separability criterion.

This approach, preceded by a number of filtering steps, was used in the high profile METABRIC paper (Curtis et al. 2012) to derive a novel classification of breast cancer patients into clinically meaningful subgroups. Information from inherited variants (CNVs and SNPs) and acquired somatic copy number aberration (SCNAs) was integrated to define these subgroups. While **iCluster** was originally formulated for clustering continuous data types, **iCluster+** (Mo et al. 2013) extends the framework to cope with both discrete and continuous data, replacing the linear formulation in (1) by a generalized linear one.

 3.1.2. Bayesian integrative clustering approaches—The METABRIC paper and its potential importance for clinical management of breast cancer opened the door to numerous studies of tumor heterogeneity. It also stimulated the development of alternative clustering approaches, aiming to exploit the power of Bayesian mixture models to increase the flexibility of integrative clustering. Besides being able to use different types of data (discrete and continuous) and to include a natural assessment of uncertainty provided by the use of Bayesian Dirichlet multinomial models as underlying structure, it was also considered important to allow for the possibility of not assuming the same clustering on all data types. Instead, the Bayesian formulations aim to find related clustering structures across the data types. Two main approaches have been taken to model cluster dependence, consisting in either relating clusters or in uncovering common and specific cluster patterns between the data types.

Building on the work of Savage et al. (2010) and Yuan, Savage & Markowetz (2011) for integrative clustering of two data types, Kirk et al. (2012) proposed **MDI** (Multi Dataset Integration). Denoting the observed data for gene i in data type k by X_{ik} , where $i = 1, ..., n$

and $k = 1, ..., K$. A Dirichlet-Multinomial allocation model (DMA) for each data type is specified:

- each gene *i* is classified into one of N components (N fixed, same for each dataset, components may be empty) with allocation probabilities given by $P(z_{ik} = j) = \pi_{jk}$ for $j = 1, ..., N;$
- $\frac{1}{2}$ in each dataset k , a mixture model is specified using appropriate parametric densities, f_k , involving parameters Θ_k ;
- association parameters φ_{km} θ are introduced to control the strength of association between pairs (k, m) of datasets:

$$
P(z_{i1},...,z_{iK}) \propto \prod_{k=1}^{K} \pi_{z_{ik}k} \prod_{k=1}^{K-1} \prod_{m=k+1}^{K} (1 + \phi_{km} I_{[z_{ik}=z_{im}]}).
$$

where π_{Zik} is the allocation probability of gene *i* to the component Z_{ik} in data type *k*.

Estimation proceeds by stochastic simulation using Gibbs sampling, exploiting natural conjugacy in the model formulation. As clusters are allowed to be empty, N should be sufficiently large, with $N = n/2$ a practical recommended choice. If φ_{km} is large, then groups of co-clustering genes in dataset k will be encouraged to have the same 'label' in dataset m . Interpretation of these parameters and the associated posterior probabilities for a sample *i* to be 'fused' across the datasets, i.e., to have the same label in a subset of data types, allow a rich interpretation of the posterior output.

Lock & Dunson (2013) propose **BCC** (Bayesian Consensus Clustering) which aims to simultaneously uncover *source-specific* clusters for each data type and a *common* clustering pattern for all data types. Such a decomposition, in line with a tradition of hierarchical modeling of several data sources in epidemiology into common and specific patterns (see for example Knorr-Held & Best (2001) or Ancelet et al. (2012)), makes stronger structural assumptions than **MDI**. As in **MDI**, **BCC** uses a fixed number N of clusters for each data type. **BCC** considers an overall 'consensus clustering' C, with corresponding latent allocation z_i and links the cluster labels z_{ik} in the different data types to the consensus clustering C through a dependence function ν:

$$
P(z_{ik}=j|z_i)=\nu(j,z_i,\alpha_k)=\begin{cases} \alpha_k & \text{if } z_i=z_{ik} \\ (1-\alpha_k)/(N-1) & \text{otherwise} \end{cases}
$$

where $a_k \in [1/N, 1]$ controls the level of 'adherence' of data type k to overall clustering and the function ν aligns the cluster labels in the different data types. Similar to **MDI**, estimation of **BCC** is implemented through Gibbs sampling. By using a formulation and parametrisation that increases linearly with the number of clusters rather than in a quadratic fashion as for **MDI**, the **BCC** algorithm is more scalable to a large number of data sources and samples. On the other hand, the appropriateness of the basic assumption of the existence of a consensus clustering has to be evaluated for each case study. Both **MDI** and **BCC** use

datasets from The Cancer Genome Atlas (TCGA) to illustrate performance and interpretability of the clustering patterns uncovered, integrating up to four TGCA data sources.

It is clear that flexible clustering approaches, which exploit jointly several genomics levels, plays an important role in integrative genomics. A natural extension of such approaches would be to incorporate additional outcome data, i.e., to use a joint model of features and response in a semi-supervised manner, rather than proceed sequentially with clustering first, then by linking clusters with survival outcome as presented in the METABRIC paper. In the genetic epidemiology context, Papathomas et al. (2012) used a joint clustering of genes and lung cancer outcomes to explore potential for gene-gene interactions. They adopt a nonparametric Bayesian approach referred to as profile regression (Molitor et al. 2010), which also allows the selection of the important features that drive the clustering. Integration of additional structure in the data, i.e., spatial organization, into the formulation of integrative clustering models, would also be of great interest, as it may provide additional interpretability of the clusters (see Pettit et al. (2014)). Such extension would be particularly relevant in view of the recent developments of single-cell technologies.

3.2. Integrative regression

Integrative clustering addresses vertical integration for unsupervised tasks. For supervised problems, regression approaches are ubiquitously employed. When regression tasks involve many more features than samples, the so-called large p small n paradigm, additional structure or constraints are needed in order to derive useful solutions. A large body of work has been developed along the lines of penalized regressions, which produce shrinkage estimates of the regression coefficients, the most common being ℓ_2 or ℓ_1 penalties corresponding to ridge or lasso regressions, respectively. Penalized regression approaches for high dimensional data and their numerous extensions have been thoroughly reviewed in the article by Bühlmann, Kalisch & Meier (2014) in ARSIA Volume 1. In this section, we will cover in more details Bayesian approaches, which combine variable selection for high dimensional problems with integrative genomics tasks, than their penalised regression counterparts.

 3.2.1. Including prior information into variable selection—Given a set of ^p covariates, $\{X_j, j = 1, ..., p\}$, let us consider the regression model of outcome y on the set of covariates:

$$
y=X\beta+\varepsilon
$$
, $\varepsilon \sim N(0, \sigma_{\varepsilon}^2 I_n)$, (2)

where $y = (y_1, ..., y_n)^T$, $X_{n \times p}$ is the covariate matrix, *n* the number of samples with $p \gg n$, and $\beta = (\beta_1, ..., \beta_p)^T$ is the vector of regression coefficients (response and covariates are assumed centered).

Bayesian variable selection methods typically include binary variable selection indicators γ_i indicating if variable X_j is included or not $(\gamma_j = 1 \text{ if } \beta_j \quad 0 \text{ and } \gamma_j = 0 \text{ if } \beta_j = 0)$ and aim to explore the vast set of 2^p possible models corresponding to $\gamma = (\gamma_1, ..., \gamma_p)$. Alternatively,

spike and slab priors for the regression coefficients have also been considered (Ishwaran & Rao 2005). Focussing for now on conditional formulations,(2) reduces to:

$$
y|\gamma = X_{\gamma}\beta_{\gamma} + \varepsilon
$$
, $\varepsilon \sim N(0, \sigma_{\varepsilon}^2 I_n)$ (3)

where β_{γ} is the vector of non-zero coefficients, X_{γ} is the $n \times p_{\gamma}$ reduced design matrix with columns corresponding to $\gamma_j = 1$ and p_γ is the overall number of non zero coefficients.

Full Bayesian inference require prior specification for the regression coefficients. In order to explore the vast model space efficiently, conjugate priors for the regression coefficients β_{ν} are commonly adopted, so that the regression coefficients can be integrated out. Both independent priors (Hans, Dobra $&$ West 2007) and Zellner g-prior structure with a hyperprior on g have been used (Bottolo & Richardson 2010). A range of efficient stochastic algorithms to explore the vast model space have been proposed, e.g. the Stochastic Shotgun (SSS) sampler (Hans, Dobra & West 2007), some inspired from population Monte Carlo, e.g. the Evolutionary Stochastic Search (ESS) sampler (Bottolo & Richardson 2010; Bottolo et al. 2011a).

The prior model of the binary indicators has direct influence on the sparsity of the model space. Under an exchangeability assumption, it can be tuned to encompass prior assumptions on the mean and variability of the overall expected number of selected covariates (Bottolo & Richardson 2010). On the other hand, specific external information W_i might be available on each of the covariates X_j , information that could make the selection of X_j more or less likely. Such a situation arises, for example, in genetic association studies where additional functional characterizations of the SNPs in terms of genomic regions or functional annotation might be relevant. To integrate such information in a flexible manner, a natural extension of (3) is to specify a hierarchical model for $\{\gamma_i\}$ and use a probit link for linking the underlying probabilities to the external information:

$$
\gamma_j \sim B(\pi_j) \quad \pi_j | \alpha \sim \Phi(\alpha_0 + \mathbf{W}_j \alpha_1), j = 1, \dots, p. \tag{4}
$$

If $\alpha_1 \simeq 0$ then the model (4) is equivalent to a standard exchangeable prior on the selection indicators. Estimating (α_0, α_1) together with (3) allows quantifying the influence of the external information. Quintana & Conti (2013) propose such an extension and illustrate its benefits in a genetic association study of smoking cessation involving 121 SNPs. For each SNP, they integrate external information on gene regions and on a quantitative association with a nicotine metabolite ratio. Integrative regression can also be used when building directed networks, as discussed in Section 3.3.

Besides quantitative information, structural and distributional information can also be integrated in a variable selection framework to improve inference. For example, Stingo et al. (2011) include prior information on gene networks to better select discriminatory variables. They model the joint distribution of the binary selection indicators $\{\gamma_i\}$ as a Markov Random Field (MRF):

$$
P(\gamma_j | d, f, k \in N_j) \hspace{-1mm} = \hspace{-1mm} \frac{\exp(\gamma_j(d\hspace{-0.1mm}+\hspace{-0.1mm}f \hspace{-0.1mm}\sum_{k \in N_j} \hspace{-0.1mm}\gamma_k))}{1\hspace{-0.1mm}+\hspace{-0.1mm}\exp(\gamma_j(d\hspace{-0.1mm}+\hspace{-0.1mm}f \hspace{-0.1mm}\sum_{k \in N_j} \hspace{-0.1mm}\gamma_k))}.
$$

where N_j is the set of direct neighbors of variable *j* is a preset graph, e.g., extracted from KEGG database; *d* controls the sparsity of the model and *f* controls the strength of 'spatial' structure, with both d and f fixed.

The Bayesian approaches discussed above for integrating information into the variable selection have analog in the penalized regression context. Group lasso (Yuan & Lin 2006) allows groups or network of genes to be viewed as additional information to tailor the penalization. These ideas were refined in the genomic context by Pan, Xie & Shen (2010) who used a group penalty based on KEGG pathway information to predict survival of glioblastoma patients. To incorporate external information provided by additional sources of data, Bergersen, Glad & Lyng (2011) propose a weighted lasso approach, where the weights are inversely proportional to a quantitative function linking external information, response and covariates. An additional tuning parameter controlling the relative strength of all the weights is calibrated through cross validation. This exible approach allow a variety of external information to be straightforwardly incorporated into the analysis, e.g. copy number alterations when looking for prognostic gene expression signatures, and was shown to improve predictive ability.

 3.2.2. Multiple response model—Faced with a set of correlated responses or related phenotypes, performing joint regression analysis of these responses is another way of borrowing information to increase sensitivity. Multiple response models extend the single outcome regression (3) to multidimensional responses $Y(n \times \ell)$ where ℓ is the number of responses, by considering the residual variance-covariance between the responses $\Sigma(\ell \times \ell)$. The likelihood (2) becomes:

$$
Y|\gamma = X_{\gamma}B_{\gamma} + \varepsilon, \quad \varepsilon \sim \mathcal{N}(I_n, \Sigma) \quad (5)
$$

where as before $X_{\gamma}(n \times p_{\gamma})$ and $B_{\gamma}(p_{\gamma} \times \ell)$ now represents the matrix of regression coefficients for the selected variables. In (5), it is assumed that the predictors have an effect (possibly different) on multiple outcomes at once and the borrowing of information is effected through the correlation between the responses. The 2^p model search task remains as previously and Bayesian algorithms can be extended straightforwardly to multiple response models.

This approach was used by Bottolo et al. (2013) to analyze groups of correlated lipid phenotypes. Inspired by the known structure of HDL and LDL cholesterol pathways, different combinations of lipid biomarkers (Trigyceride, HDL and LDL cholesterol, APOA1 and APOB) were analyzed following (5) and regressed on a genome wide set of 273,675 SNPs derived from Affymetrix Genome-Wide Human SNP arrays 6.0 (tagged $r^2 > 0.8$) To cope with the challenging computational task of performing model exploration on this large

set of SNPs, observed on 3175 individuals, a GPU (Graphics Processing Units) version of the ESS algorithm (GUESS) was developed. Synthetic measures of evidence such as a list of "top models", together with estimate of their posterior probability and Bayes factors (BF) against the null model, as well as marginal posterior probability of inclusion for each SNP (using model averaging) rescaled to be comparable across different combination, were derived. The results provided new insight into the genetic control of lipid pathways, refining some of the previous GWAS results (Bottolo et al. 2013).

3.2.3. Joint regression of two -omics datasets and eQTL models—Many

biological questions can be expressed under the generic framework of performing the joint regression analysis of two or more different types of genomics datasets. In this section, we focus on analyses where a large number of responses is regressed on a very large number of predictors. Multiple response models described in Section 3.2.2, which assume that a set of predictors affect all the responses at once, are not adapted to analyses involving a large number of responses. A canonical example of genomic studies involving a large number of responses are the so-called eQTL studies, which investigate the genetic control of expression by regressing expression profiles on DNA variants (Figure 3). Other examples are mQTL studies, which link DNA variations to metabolite synthesis (Marttinen et al. 2014) or studies investigating the influence of SCNAs on tumor gene expression. Flexible ways of borrowing information between the high dimensional phenotypes are required to increase power. In other words, rather than testing the association between each pair (marker \times expression) separately and subsequently face a huge multiplicity adjustment, the high dimensional set of responses, e.g. gene expression, is treated as related outcomes. The statistical aims are thus expanded to not only uncover the multivariate association of each (expression) response with a large number of features, e.g., genetic markers, but to also highlight the features that are associated with many responses. Finding key control points associated with the the expression of many genes, sometimes called '*hot spots*', is an important step towards a better understanding of biological pathways.

Different approaches have been proposed for discovering regression links between a large number q of responses $(y_k, 1 \mid k \mid q)$ and a large set of predictors X in a way that exploits the relatedness of the responses. Penalized approaches use structured regularization to account for the correlation of the responses. For example, Peng et al. (2008) propose a combination of ℓ_1 and ℓ_2 penalties to encourage the detection of master regulators, while Kim (2012) use a tree-guided lasso to account for the relationship between the genes. An early Bayesian approach is the Mixture Over Markers (MOM) method (Kendziorski et al. 2006), which associates each response with any of the p predictors (or none of them) via a mixture model, so each response is associated with at most one marker, a workable but restrictive assumption. Stochastic partition approaches where the responses are partitioned into disjoint subsets that have a *similar dependence* on a subset of covariates, have also been implemented in eQTL analyses, making strong assumptions on the commonality of effects within the blocks (Monni 2009); see also Zhang et al. (2010) for an application of Bayesian partitioning to find pleiotropic and epistatic eQTL modules.

Bayesian approaches combining high dimensional variable selection for each response with a hierarchical structure on the selection indicators have the benefit of being fully multivariate while retaining scalability. The key quantities that are involved in such models are:

- **•** the latent binary vectors $\gamma_k = (\gamma_{k1}, ..., \gamma_{kj}, ..., \gamma_{kp})^T$ for each regression of y_k on X , where each indicator has a Bernoulli prior $p(\gamma_{kj}|\omega_{kj}) = \omega_{kj}^{\gamma_{kj}}(1-\omega_{kj})^{1-\gamma_{kj}}$
- **•** $\Gamma = (\gamma_{kj} 1 \quad k \quad q, 1 \quad j \quad p)$, the $(q \times p)$ matrix of selection indicators
- **•** a hierarchical structure for the matrix of prior probabilities for Γ

$$
\Omega = \begin{bmatrix}\n\omega_{11} & \cdots & \omega_{1j} & \cdots & \omega_{1p} \\
\vdots & \ddots & \vdots & \ddots & \vdots \\
\omega_{k1} & \cdots & \omega_{kj} & \cdots & \omega_{kp} \\
\vdots & \ddots & \vdots & \ddots & \vdots \\
\omega_{q1} & \cdots & \omega_{qj} & \cdots & \omega_{qp}\n\end{bmatrix}
$$

that facilitates sparsity control in each regression as well as the borrowing of information across responses to highlight important predictors common to several responses.

Different prior structures for ω_{kj} have been proposed. Bottolo et al. (2011b) introduce a multiplicative parametrization: $\omega_{kj} = \omega_k \times \rho_j$ with $\omega_k \sim Beta(a_{\omega_k}, b_{\omega_k}), \rho_j \sim Gam(c_{\rho_j}, d_{\rho_j}),$ subject to the constraint: $0 \quad \omega_{kj} \quad 1$. In this choice of parametrization, ρ_j captures the 'propensity' for predictor *j* to influence several outcomes at the same time, while ω_k controls the sparsity of each regression. Scott-Boyer et al. (2011) propose a mixture model for ω_{ki} with an atom at zero to reduce the false discovery rate, a Beta distribution for the second mixture component, and a SNP specific mixture weight: $\omega_{kj} = p_j \delta_0 (\omega_{kj}) + (1 - p_j) Beta(a_j)$ b_j)(ω_{kj}). Both approaches are implemented by MCMC. The choice of a g prior structure for the regression coefficients in Bottolo et al. (2011b) allows the latter to be fully integrated out and to use a hierarchical extension of their ESS sampler to traverse the model space, while the prior structure defined by Scott-Boyer and implemented in their algorithm iBMQ (Imholte et al. 2013) requires joint updating of the variable selection and regression coefficients. Despite efficient MCMC implementation, fully Bayesian joint eQTL analysis strategy are nevertheless quite demanding in terms of computational time and would typically need to be run in parallel on each chromosome on a few thousands genes only. Ultra fast implementation of a linear model, which tests the association of each SNP with each transcript, as implemented in Shabalin (2012) could be used as a pre-selection step.

Both Bottolo et al. (2011b) and Scott-Boyer et al. (2011) make the simplifying assumption of no residual dependence between the responses conditional on the selected model. Adding a model of the residual structure to the previous set-up, in a framework akin to Seemingly Unrelated Regressions, has been proposed by Bhadra & Mallick (2013). The additional computational complexity is severe and such approaches will not easily scale up for large q. To encompass additional nuisance correlation in a computationally feasible manner, Stegle et al. (2010) and Fusi, Stegle & Lawrence (2012) propose a variational Bayesian approach

which accounts for additional known and hidden sources of variation using an computationally tractable latent factor approach. The factors can be used as covariates in standard eQTL mapping, or can be interpreted as corresponding to transcription factor activations if additional biological information is provided to the model. The efficient software PEER (Stegle et al. 2012), which is designed to uncover such hidden factors, has been used to re-analyze several eQTL studies where an increase of power for eQTL detection is shown. In the same spirit, Bayesian reduced rank regression has been used by Marttinen et al. (2014) to analyze gene-metabolome associations, account for known factors and combine information over multiple SNPs and phenotypes.

3.3. Graphical model

Graphical models or biological networks are powerful tools to describe the relationships of different biological entities. Commonly used biological networks include proteinprotein interaction networks, co-expression networks, transcription regulation networks, signaling pathway, and etc. (Figure 4). Many statistical methods have been developed to construct or to exploit the knowledge from such networks to analyze genomic data. We will focus on three approaches in the subsection. First, we use gene expression quantitative trait loci (eQTLs) to construct a Directed Acyclic Graph (DAG) for gene expression. Second, we integrate miRNA expression, gene expression data, annotation data to infer the miRNA-gene regulations. Third, we will review graphical model approaches for pathway activity estimation.

 3.3.1. eQTL-guided DAG construction—A co-expression network is usually an undirected graph. However, there are many situations where a directed graph is desirable, for example, to infer the consequence of a perturbation by a drug. DAG models have been used to construct directed graphs using gene expression data (Neto et al. 2008, 2010; Hageman et al. 2011; Bühlmann, Kalisch & Meier 2014). In such a DAG, each vertex represents a gene and each edge represents a direct causal relation between two genes. For example, an edge $g_1 \rightarrow g_2$ implies that perturbation of g_1 alters g_2 while changes on g_2 leaves g_1 unaffected. It is well known that interventions or perturbations are needed to infer causal relations. However, a huge number of interventions on gene expression (e.g., gene knock out) are needed to infer causal relations of thousands of genes, which are not feasible yet. The eQTLs of gene expression provide natural perturbations to the expression of a large number of genes. It can be considered as a randomized experiments on DNA genotype (i.e., Mendelian Randomization (Smith 2007; Sheehan et al. 2008)), and the design of experiment (i.e., interventions on DNA genotype rather than gene expression) is also consistent with our intuition that DNA genotype affects gene expression rather than vice versa (Chen et al. 2007). The genetic variants of eQTL studies may be genotype, copy number variations, or other DNA variants. Without loss of generality, we assume such genetic variants is DNA genotype in the following.

The example in Figure $5(A)$ illustrates a situation where eQTLs can help to estimate edge directions in a DAG. Consider genes g_2 and g_3 , which are co-expressed, and thus there is an undirected edge $g_2 - g_3$ in the graph. Without external data, we cannot distinguish $g_2 \rightarrow g_3$ and $g_2 \leftarrow g_3$ because two DAGs encode the same dependence assumption and have the same

likelihood: $L(g_2 \to g_3) = f(g_3|g_2)f(g_2) = f(g_2, g_3) = f(g_2|g_3)f(g_3) = L(g_3 \to g_2)$. If we know that g_2 has an eQTL, denoted by c_2 . Then the partially directed graph is $c_2 \rightarrow g_2 - g_3$, and the possible DAG is $c_2 \rightarrow g_2 \rightarrow g_3$ or $c_2 \rightarrow g_2 \leftarrow g_3$. These two graphs can be distinguished because they encode different conditional independence assumptions. $c_2 \rightarrow g_2 \rightarrow g_3$ implies $c_2 \perp g_3|g_2$ and $c_2 \rightarrow g_2 \leftarrow g_3$ implies $c_2 \sim g_3|g_2$, and thus have different likelihoods. To understand the reason that $c_2 \sim g_3|g_2$, one may consider an example "rain \rightarrow wet grass \leftarrow sprinkler", where given the event that grass being wet, the two parent vertices rain and sprinkler are dependent.

To use eQTLs to derive causal gene expression network, we also need to separate direct and indirect eQTL effects. Using the example in the previous paragraph and assuming the causal relation is $c_2 \rightarrow g_2 \rightarrow g_3$, then c_2 may appear to be an eQTL for both g_2 and g_3 . We need to know that c_2 directly affects g_2 , but indirectly affects g_3 for the purpose of DAG estimation. Such information can be obtained by separating cis -eQTL and $trans$ -eQTL using RNA-seq data (Sun 2012; Sun & Hu 2013). All of the cis-eQTLs directly influence their target genes and a trans-eQTL may be inuencing its target's expression directly or indirectly. Therefore, it is desirable to use only cis-eQTLs for DAG construction.

Neto et al. (2008) developed the QTL Directed Dependency Graph (QDG) method and implemented in the R package qtlnet. The QDG method was originally designed to study the relations of multiple phenotypes given their QTLs, though it can be applied for eQTL studies as well. The QDG method assumes "multiple QTLs associated with these traits had previously been determined". It has the following steps. (1) Construct a DAG skeleton from the PC algorithm, which is a popular algorithm for DAG skeleton construction (Spirtes, Glymour & Scheines 2000). (2) Distinguish QTLs with direct and indirect effect. (3) Orient each edge by LOD score, which is the log_{10} likelihood ratio for the edge $Y_i \rightarrow Y_j$ versus Y_j $\rightarrow Y_i$ given all the vertices (either phenotype or DNA genotype) connected to Y_i or Y_j . (4) Randomly choosing an order of all the edges, and then following this order, sequentially update the directions of the edges using the LOD score conditioning on the vertices that are parents of Y_i or Y_j . (5) Repeat step (4) for 1000 times and choose the graph with the highest score, which could be a likelihood-based measure of goodness of fit.

In a later paper, Neto et al. (2010) developed a new method named QTLnet, which jointly estimates the graphical structure of the phenotypes and the underlying genetic architecture. This method would be computationally too demanding to study genome-wide eQTL data with tens of thousands genes and millions of SNPs. In addition, the genetic architecture of human gene expression is relatively simple, with the vast majority of the eQTLs being local eQTLs. Therefore it may be a reasonable approximation to assume the genetic architecture only involves local eQTLs and then estimate eQTLs before DAG estimation. In contrast to QDG, which reports the most likely graph, the QTLnet approach reports graph structure based on Bayesian model averaging. In other words, the posterior probability of edge $Y_i \rightarrow$ Y_j is the summation of the posterior probabilities of the graphs that have the edge $Y_i \rightarrow Y_j$.

Another type of approach for graphical model estimation is structural equation models (SEM) that permit both cyclic and acyclic graphs. Li et al. (2006) employed a score-based model selection method. Logsdon & Mezey (2010) estimated network skeleton by applying

an adaptive lasso regression for each gene expression trait, and then transformed the skeleton into a DAG or a Directed Cyclic Graph (DCG) based on eQTL perturbations. Cai, Bazerque & Giannakis (2013) extended the work of Logsdon & Mezey (2010) by providing the adaptive lasso a set of initial parameter estimates from penalized regressions using the LASSO penalty.

 3.3.2. Construction of miRNA regulation network—Recent studies have shown that microRNA (miRNA), a class of short non-coding RNA molecule (21–24 nucleotides), may play an important role in transcriptional and post-transcriptional regulation of gene expression (Pasquinelli 2012). The human genome may encode over 1,000 miRNAs, which may target more than half of human transcripts. One miRNA's sequence may match the complementary sequences of one or more mRNAs, and thus this miRNA can bind these base-paired mRNAs, which leads to mRNA degradation or represses the translational process. Therefore, over-expression of a miRNAs usually reduces the expression of its targets. In plants, a miRNA is often perfectly or almost perfectly matched with its targets. However, animal miRNAs typically exhibit only partial complementarity to their mRNA targets. A "seed region" of about 6–8 nucleotides in length at the 5' end of an animal miRNA is thought to be an important determinant of target specificity (Pasquinelli 2012). Many computational approaches have been developed to predict miRNA targets based on sequencing similarity. However, these methods have limited accuracy due to the relatively low target specificity based on sequence data alone. Motivated by this problem, several methods have been developed to integrate gene expression, miRNA expression as well as miRNA target annotation based on sequence similarity to infer the miRNA regulatory network (Muniategui et al. 2013). Here we briefly review a Bayesian graphical model approach (Stingo et al. 2010).

Denote the expression of G genes by $\mathbf{Y} = (Y_1, ..., Y_G)$, and the expression of M miRNAs by $X = (X_1, ..., X_M)$. Stingo et al. (2010) construct a DAG for these $G + M$ variables where the only allowable edges are those of the form of $X_i \to Y_k$ where $i = 1, ..., M$ and $j = 1, ..., G$. In other words, they assume that $X_i \perp X_j$ for any $i, j = 1, ..., M$ and $i \neq j$, and $Y_k \perp Y_j \perp X$ for any k, $l = 1, ..., G$ and k l (Figure 5(B)). Since the marginal distribution of **X** does not affect the estimation of regulatory relations, the assumption $X_i \perp X_j$ is a reasonable choice to simplify the computation. When sample size is large enough, one may further relax the assumption $Y_k \perp Y_j \mathbf{X}$, though imposing this assumption may be the best one can do given a limited sample size and/or computational power. An additional assumption is that all the edges must point from X_i to Y_j , which is justifiable by the regulatory role of miRNA. Give such an underlying DAG, the problem reduces to G regression problems where the g -th problem aims to select those miRNAs that regulate the g-th gene.

Stingo et al. (2010) assumes a linear model with Gaussian errors such as

$$
Y_g = -\sum_{m=1}^{M} X_m \beta_{gm} + \varepsilon_g, \tag{6}
$$

where ε_g 's are i.i.d. $\mathcal{N}(0, \sigma_g)$, and the negative sign in front of the term $\sum_{m=1}^{M} X_m \beta_{gm}$ indicates that miRNAs repress gene expression. The prior distribution for the regression coefficients $β_{gm}$'s are

$$
\pi(\beta_{gm}|\sigma_g, r_{gm}) = r_{gm} \text{Gam}(1, c\sigma_g) + (1 - r_{gm})I_{(\beta_{gm} = 0)}, \quad (7)
$$

where *Gam*() indicates a gamma distribution, $I_{(\beta_{gm}=0)}$ is an indicator function, $r_{gm} = 1$ if the m-th miRNA regulates the g-th gene and $r_{gm} = 0$ otherwise. Then the key part of the method, which is to incorporate the annotation based on sequencing similarity, is implemented as the prior distribution for r_{gm} :

$$
\log\left(\frac{P(r_{gm}=1)}{1 - P(r_{gm}=1)}\right) = \eta + \sum_{u=1}^{U} s_{gm}^u \tau_u, \tag{8}
$$

where s_{qm}^u is a score describing the degree of confidence of that the *m*-th miRNA regulates the g-th gene. The regression coefficients τ_u 's are additional set of parameters with at hyper prior set as a gamma distribution $Gam(a_t, b_t)$. Then Stingo et al. (2010) designed an MCMC approach to sample all the parameters, among them r_{gm} 's are of primary interest because they indicate whether the m -th miRNA regulate the g -th gene.

3.3.3. Inference of each gene's contribution to the activity of a pathway—

Most human diseases are complex diseases (e.g., diabetes or cancer) that are associated with mutations or perturbations of multiple genes. Such complex diseases may be better described at the pathway level. For example, two patients may have different sets of mutations but each may modify the activity of the same pathway. Therefore, to study the importance of a gene that belongs to a known pathway, one may quantify the change of pathway activity by turning on or off this gene. As shown in Figure 4 (C) and (D), the contribution of a gene to a pathway (e.g., a transcriptional regulation pathway or a signaling pathway) should be measured by its protein activity, which could be the abundance of an active form, e.g., a specific phosphoprotein, conditioning on the states of other proteins within the pathway. These quantities are often latent variables that cannot be directly measured in genome-scale using current techniques. PARADIGM (PAthway Recognition Algorithm using Data Integration on Genomic Models) is popular computational method that addresses this challenge by integrating different types of genomic data and pathway annotation.

In the PARADIGM model, Vaske et al. (2010) assume the pathway information is known. Considering a pathway as a graph, a vertex of this graph can be a protein-coding gene, a protein complex, a gene family, an abstract processes (e.g., apoptosis), or other biological entities. Each vertex has three states: activated, nominal, or deactivated relative to a control level and is encoded as 1, 0 or −1, respectively. Therefore the graph is a factor graph, and such a simplified assumption of three states greatly reduces the difficulty of model estimation. Each edge of this graph has a sign, indicating whether the parent vertex

has a positive or negative influence on the child vertex. PARADIGM includes four entities for the *j*-th protein-coding gene: DNA copy number (c_j) , mRNA expression (g_j) , protein abundance (p_j) , and protein activity (a_j) , see Figure 6A for an example of three protein coding genes. Directed edges with positive signs are introduced as $c_j \rightarrow g_j \rightarrow p_j \rightarrow a_j$. The relation between protein coding genes are introduced based on pathway annotation. For example, if activated protein 1 induces the activity of protein 2, a directed edge $a_1 \rightarrow a_2$ with positive sign is added. If activated protein 2 represses the expression of gene 3, a directed edge $a_2 \rightarrow g_3$ with negative sign is added (Figure 6A).

The graph allows one to compute the expected state of the *i*-th vertex, denoted by μ_i , given its parents. Assuming the parents contribute additively, $\mu_i = sign(\sum_{j \in Pa_i} \mu_j \beta_{ji})$, where Pa_i denotes the parent set of the *i*-th vertex, and $\beta_{ji} = 1$ or -1 is the sign of the edge *j* 1 *i*. PARADIGM also allows the contribution from all the parents to be summarized by an "AND" or "OR" operation (Figure 6B). We use X_i to denote the underlying state of the *i*-th vertex. If X_i is unobserved, it follows a categorical distribution across the three classes such that $P(X_i = a) = 1 - \varepsilon$ if $a = \mu_i$ and $P(X_i = a) = \varepsilon/2$ otherwise, where ε is a small value, e.g., ε $= 0.001$. For some of the vertices, the values of X_i 's are observed (assuming no measurement error) and the purpose of PARADIGM method is to infer the state of unobserved X_i 's. They employ an EM algorithm to infer such hidden states. Finally, an IPA (Integrated Pathway Activity) is estimated for each biological entity. Note that the name IPA may be misleading. It does not estimate pathway activity per se. Instead, it calculates how much each entity contributes to the pathway activity. Specifically, let $\ell(i, a) = \log [P(D|X_i =$ a)/ $P(D|X_i \mid a)$], which is the log likelihood ratio comparing the situation $X_i = a$ versus. X_i ^a. Then

$$
IPA(i) {=} \left\{ \begin{array}{ll} \ell(i,1) & \text{if} \ \operatorname{argmax}_a \ell(i,a) {=} 1 \\ -\ell(i,-1) & \text{if} \ \operatorname{argmax}_a \ell(i,a) {=} -1 \\ 0 & \text{otherwise.} \end{array} \right.
$$

In other words, $IPA(i)$ is the signed log-likelihood ratio if the most likely state is 1 or −1, and $IPA(i)$ is 0 otherwise. Vaske et al. (2010) further demonstrated that clustering IPAs may reveal meaningful clusters that cannot be identified by clustering each type of genomic data directly.

3.4. Other methods for integrative genomics

In the previous sections, we have highlighted some recent work for integrative genomics. Since integrative genomics is a very active research area with huge amount of literature, we cannot give an exhaustive review of all work in this area. In the following, we summarize some methods that aim to answer some specific biological questions.

 3.4.1. Phenotype association/prediction—Xiong et al. (2012) proposed a method named Gene Set Association Analysis (GSAA) to identify disease-associated gene sets. They first assessed the association between a phenotype and the expression and genotype of a gene separately. Then they combined the z-statistics of differential expression and genotype association using Fisher's method for each gene, and used the combined gene-

specific test statistic for gene set enrichment analysis (Newton & Wang 2015). Instead of analyzing each type of data separately, Tyekucheva et al. (2011) first summarized each type of genomic data at the gene level, and then studied the association between a gene and a phenotype using one regression model. In this approach, the phenotype was used as the response variable and different types of genomic data were used as covariates. They scored this gene using a test statistic for the null hypothesis that the regression coefficients for all the genomic data are 0's. Then this test statistic was used for gene set enrichment analysis.

Another question that is often of interest is whether one type of genomic data mediates the effect of the other type of genomic data on phenotype. For example, whether gene expression mediate the effect of DNA genotype on disease outcomes. Huang, VanderWeele & Lin (2014) used mediation analysis to address this question, and provided quantification of SNP genotype's direct effect and indirect effect (mediated by gene expression) on disease outcomes.

In addition to association testing, genomic data can be used for the prediction of phenotypes. It is not an unusual situation that many genomic features may have relatively small effects on the phenotype, so that there is not enough power to identify such genomic features by association testing. However, one may still be able to perform predictions without selecting phenotype-associated genomic features. An example is OmicKriging (Wheeler et al. 2014). Kriging is a well-known geo-statistical method for prediction of spatially measured outcomes, by making prediction using observations from nearby locations (Cressie 1993). In OmicKriging, Wheeler et al. (2014) assumes the similarity matrix of phenotype data across

all samples is $\Sigma = \sum_{j=1}^{d_j} \sum_{j=1}^{d_j} \sum_{j=1}^{d_j} \sum_{j=1}^{d_j} \sum_{j=1}^{d_j}$ where S_j is the similarity matrix for the *j*-th type of genomic data and *I* is an identify matrix to capture variance due to environmental factors. Given such phenotype similarity derived from genomic data, one can easily make predictions on phenotypes. For example, the phenotype of a testing sample could be a weighted average of the phenotypes of training samples, where the weights are the similarities between this testing sample and all the training samples. Wheeler et al. (2014) showed that their method could provide good prediction of several phenotypes after combining multiple types of genomic data.

 3.4.2. Gene expression regulation modules—Several integrative genomic methods have been developed to identify gene expression regulation modules. Sun, Yu & Li (2007) developed a method to detect modules where a local eQTL modifiles the expression of a gene, which modifiles the activity of a transcription factor (TF), and the TF in turn regulates the expression of a group of genes. They integrated TF binding site data and gene expression data to infer latent TF activities and then used genotype data, gene expression and estimated TF activity to build regulation modules. Akavia et al. (2010) proposed a method named CONEXIC (copy number and expression in cancer) to detect modules where copy number affects the expression of a driver gene, which in turn regulates the expression of a group of genes.

 3.4.3. Study functional consequence of somatic mutations by integrating somatic mutation data and gene-gene interaction annotations—Because somatic

point mutations tend to be rare, it is difficult to assess their effects directly. Several methods have been developed to borrow information of known gene-gene interactions (e.g., proteinprotein interactions or regulation relations) to study the functional consequence of somatic point mutations. The method of DriverNet (Bashashati et al. 2012) seeks to study the consequence of somatic mutations on gene expression by connecting genes A and B such that A has a somatic mutation, B has extreme gene expression, and A and B are connected by known gene-gene interaction(s).

Driver mutations that increase the survival advantages of tumor cells often occur together with a number of passenger mutations in cancer patients. Many methods have been developed to find such drivers by identifying recurrently mutated genes. Several recent works have shown that exploiting a "mutual exclusive pattern of somatic mutations" may help to identify a set of driver mutations. MEMo (Ciriello et al. 2012) selects a group of recurrently mutated genes that are close in the gene-gene interaction graph and having mutually exclusive mutations. The focus on genes that are close in the gene-gene interaction graph is partly due to the high computational cost of exhaustive search. However, recently Leiserson et al. (2013) developed a computationally efficient approach to select multiple groups of genes such that the genes within groups have mutually exclusive mutations and good coverage (i.e., most patients have mutations in at least one gene), while without relying on gene-gene interaction information.

HotNet (Vandin, Upfal & Raphael 2011) identifiles significantly altered subnetworks in an interaction network by a network diffusion approach, which can be understood as a random walk on a gene-gene interaction graph. In other words, a somatic mutation in gene A may also affect gene A's neighbors in the interaction graph. After network diffusion, HotNet evaluates the frequency that a subnetwork being altered across a number of patients and find those subnetworks that are recurrently altered. Network diffusion provides a networksmoothed version of the consequence of somatic mutations. Hofree et al. (2013) propose to cluster the network-smoothed mutation profiles by non-negative matrix factorization. TieDIE (Paull et al. 2013) use network diffusion to identify pathways linking somatic mutations and transcriptional regulation pathways.

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Glossary

Array CGH

comparative genomic hybridization (CGH) on DNA microarray to compare the copy number of two DNA samples.

Read depth

the (average) number of times a nucleotide is covered by sequencing process.

Somatic mutation

a DNA mutation that is not inherited from a parent and not passed to offspring. In contrast, a germline mutation is an inheritable DNA mutation occurred in the germ cells (i.e., sperm and eggs).

Tumor purity

the percentage of tumor cells within a tumor sample.

Ploidy

the number of sets of chromosomes within a cell. A normal human cell is diploid, i.e., with 2 sets of chromosomes.

Tumor subclone

All the tumor cells within one subclone are descended from the same cell and have the same set of somatic mutations.

Histone modification

methylation, acetylation, and phosphorylation of certain amino acids of histone proteins.

DNA methylation

an epigenetic mark of DNA sequence by adding a methyl group (CH3) to DNA nucleotides.

DNase I hypersensitive sites

genomic loci that are sensitive to cleavage by the DNase I enzyme. Such sensitivity implies the DNA sequence is loosened and exposed instead of taking a condensed form.

ChIP-chip/ChIP-seq

Chromatin immunoprecipitation (ChIP) followed by microarray (chip) or sequencing.

Bisulfite sequencing

sequencing DNA after bisulfite treatment that converts un-methylated cytosine to uracil while leaving methylated cytosine unaffected.

Microarray probe

a fragment of DNA sequence located in spot of a microarray.

Phosphorylation

the addition of a phosphate ($PO₄^{3–}$) group to a protein other organic molecule.

Mass spectrometry

a technique that measures the amount of analytes (e.g., protein peptides) by their mass-tocharge ratios.

Reverse phase protein array

a protein array designed to measure the expression of one protein across multiple samples.

1000 Genome Project

An international collaboration to produce an extensive public catalog of human genetic variation.

Protein-protein nteraction

physical interaction of two or more proteins for various biological function, e.g. signal transduction.

Co-expression network

a network where two genes are connected if their expression are dependent, either with or without conditioning on the expression of other genes.

Transcription regulation network

a network where an edge indicates a transcriptional regulation. The regulators can be transcription factors, or other molecules, such as microRNAs.

Signaling pathway

a pathway for signal transduction. It starts with activation of a specific receptor by extracellular molecules. Then the receptor triggers a series of events, such phosphorylation of proteins, and these events lead to certain cell response.

cis-eQTL

A cis-eQTL is located on the same chromosome as its target gene and inuences the gene expression in an allele-specific manner.

trans-eQTL

A trans-eQTL of a gene can be located anywhere in the genome and it inuences the gene expression of both alleles to the same extent.

DAG skeleton

an undirected graph that is constructed by removing the directions of all the edges within a DAG.

Protein complex

a group of two or more proteins that are physically associated and its function may require each individual protein to be active.

Gene family

a collection of genes in which any single gene is sufficient to perform a specific function.

Apoptosis

programmed cell death.

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- **1.** All models have their explicit and implicit assumptions. Model fitting to the underlying data structure largely determines success of omics data analysis.
- **2.** In data integration, certain biological mechanisms and prior knowledge are often known across different omics data. Proper modeling of such prior knowledge is crucial to enhance statistical power and identify biologically interpretable results.
- **3.** Incorporating prior biological information using Bayesian hierarchical modelling is a very powerful way for data integration.
- **4.** Integration of multiple omics data involves ultra-high dimensional problems. Feature selection and its related model selection problem is a major topic when developing a novel method.
- **5.** Network-based methods are effective approaches to integrate multiple types of data and biological knowledge.

Figure 1.

Different types of genomic data and the platforms to measure these genomic data.

Figure 2.

A screenshot of a spreadsheet shown in the TCGA data portal web page when querying available data from breast cancer patients. The column of this spreadsheet corresponds to a combination of data types and platforms. For each platform, there could be data from tiers 1 to 3. Usually raw data belong to tier 1, and processed data belong to tier 2 or 3. One type of genomic data may be collected on multiple platforms. For example, gene expression are measured by both microarray and RNA-seq and two types of DNA methylation arrays (JHU-USC HumanMethylation27 and JHU-USC HumanMethylation450) have been used. A platform name often starts with the institute that processes those tumor samples using that particular platform. For example, BI Genome_Wide_SNP_6 means Affymetrix 6.0 array from Broad Institute. Each row of this spreadsheet corresponds to a patient. The meaning of the patient ID can be found at<https://wiki.nci.nih.gov/display/TCGA/TCGA+Barcode>

Figure 3.

An illustration of the first genome-wide eQTL study (Brem et al. 2002) conducted in yeast sergeants (offsprings) from a cross of two yeast strains, denoted by RM and BY. Because yeast has haploid genome, the genotype data in these yeast sergeants are binary. The figure in the right panel illustrate the association between the expression of one gene LEU1 and the genotype of one SNP around gene LEU2.

Figure 4.

Examples of biological networks (A) Protein-protein interaction. The edges are detected from experiment for physical interaction. (B) Co-expression network. The edges are inferred from the expression data of a group of samples, as dichotomized correlation or partial correlation matrix. (C) Gene regulatory network. The Octagons indicate protein activity and the ovals indicates gene expression. The dash line between g_3 and a_3 indicates a_3 is the activity of the protein that is encoded by g_3 . (D) Signaling pathway. EGFR (epidermal growth factor receptor) is a receptor located at cell surface that can be activated by epidermal growth factor. The symbol^{−P} arounds a protein indicates a phosphorylation of the corresponding protein.

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Figure 5.

Statistical approaches to use graphical model for integrative genomics. (A) To infer directed co-expression network with the aid of DNA variation information. (B) To infer the regulatory relation between miRNA and gene expression.

Figure 6.

The PARADIGM method of integrative genomics (Vaske et al. 2010). (A) The graphical model of underlying states. $a_1 \rightarrow a_2$ indicates activated protein 1 regulates the activity of protein 2. $a_2 \rightarrow g_3$ indicates activated protein 2 regulates the expression of g_3 . (B) Examples of "AND" and "OR" relations. (C) Adding observed data into the graph. Here observed data, like the underlying state, has three values, −1, 0, or 1. Having observed data in certain vertices mean that the underlying states of those vertices are known.

Table 1

Genomic data resources

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Table 2

Annotation databases

