RESEARCH STATEMENT

My research interest lies in both statistical methodology and application on genomics and bioinformatics. Nowadays, large amount of genomic data are publicly available and integrating these datasets provides unprecedented opportunities to reveal disease mechanisms. I have worked on horizontal omics meta-analysis (combine multiple cohorts of the same type of omics data) and vertical omics integrative analysis (combine multi-level omics data of the same patient cohort), which will help increase statistical power, interpretability, reproducibility and understanding of disease mechanism. In terms of statistical methodology, I am particularly interested in modeling and optimization for high-dimensional data, Bayesian methods, graphical models and statistical computing, which are capable of accommodating the high-dimensional nature of genomic data. In terms of genomics and bioinformatics application, I have collaborated with biologists in fields of cancer and psychiatry to analyze a broad range of genomic data (e.g. microarray and sequencing data), which motivates me to develop practical methodology and user-friendly softwares. However, genomics and bioinformatics are fast-evolving field and I am open minded to expand my research towards new data types (e.g. imaging data, single cell data) in the future. My long term goal is to bridge bioinformatics and statistics/machine learning.

Below is a highlight of my past and on-going research, as well as future research plan.

1 Statistical Methodology

I am particularly interested in data integration, modeling and optimization for high-dimensional data, Bayesian methods, graphical models and statistical computing. Note that some of these areas can potentially overlap. I am not restricted to these areas I have explored. If I encounter other meaningful and challenging problems, I am definitely eager to learn or collaborate with other researchers.

1.1 Data integration

• Past and on-going research:

Due to rapid development of high-throughput experimental techniques and dropping prices, many transcriptomic datasets have been generated and accumulated in the public domain (e.g. TCGA, GEO, SRA). Single cohort/data type may suffer from small sample size issue. A natural question is how to combine these complex data and increase statistical power, interpretation and reproducibility. This includes two directions: horizontal meta-analysis and vertical integrative analysis. Horizontal meta-analysis aims to combine genomic data of same type from multiple cohorts. I have worked on several meta-analytical methodologies including disease subtype discovery[1], candidate marker detection[2], differential co-expression network detection[4] and dimensional reduction[5]. Vertical integrative analysis combines multi-omics data (e.g. gene expression, CNV, genotyping, methylation) of the same cohort. I have developed a disease subtype discovery algorithm integrating multi-level omics data with prior biological information[3]. These methods will help better characterize a complex disease and develop towards personalized medicine.

• Future direction:

- 1. I have worked on horizontal meta-analysis and vertical integrative analysis respectively. A natural extension is towards **two-way integration** by combining horizontal meta-analysis and vertical integrative analysis.
- 2. More and more **epigenetic**, **single cell** and **neuroimaging data** are becoming available to help understand diseases with new insight. Integrating these data with genomic data is a potential future direction.

1.2 Modeling and optimization for high dimensional data

• Past and on-going research:

High-throughput data (e.g. genomic data) has more than 20,000 genes in human being but only tens or hundreds of samples. This large p and small n problem brings statistical challenges to reveal disease mechanism behind the big data. I am particularly interested in modeling high-dimensional data and solving related

optimization problem with various forms of **regularizations**. I have used a **lasso penalty** on clustering problems to formalize a statistical objective and perform optimization[1]. In another clustering problem, which required incorporating prior knowledge, I proposed **sparse overlapping group lasso** and used **alternating direction method of multipliers (ADMM)** to solve the challenging optimization problem[3]. By these models and techniques, we can discover the intrinsic information behind the high dimensional data.

• Future direction:

- I have worked on lasso, group lasso and overlapping group lasso problems. Other regularization techniques such as penalization on precision matrix or low rank penalty are also appealing to genomic applications.
- 2. High dimensional problems are challenging from perspectives of both optimization and theory. I have worked extensively on high-dimensional optimization problems. **High-dimensional theory** is also interesting and challenging to me, which is essential for a sound methodology with theoretical guarantees.

1.3 Bayesian methods and graphical models

• Past and on-going research:

Bayesian methods and graphical models are very flexible to model and reflect the biological generative process and its complex dependent structure. I have worked on a **Bayesian non-parametric** approach to combine summary statistics from multiple cohorts to perform meta-analysis[2]. I am working on **Bayesian variable selection** problems[6] with prior knowledge of multi-layer overlapping groups. Bayesian approaches and graphical models are also growing fields themselves and I expect these techniques will play an important role in genomics and bioinformatics.

• Future direction:

- 1. I have produced methodologies on high-dimensional clustering problem using frequentist approaches. I am very interested in high-dimensional or clustering problems from the Bayesian perspective.
- 2. Part of my thesis is about methods for single cell data. I proposed a **conditional random field** model for fast single cell imputation and I'll continue exploring that.

2 Bioinformatics Application

As a biostatistician, an important job and privilege is to work with local biologists and play with their own data. This is exciting for me because I can help biologists towards innovating scientific findings, and their data can motivate me to develop relevant statistical methodology as well.

• Past and on-going research:

I have been mainly involved in **cancer** and **psychiatry** diseases research. For cancer research, I have worked on disease subtypes of breast cancer[11], DNA methylation of parity-induced mice[8], copy number variation of prostate cancer[9], and fusion transcript discovery[7]. For psychiatry diseases, I have worked on schizophrenia, bipolar disorder and major depressive disorder in human pyramidal neurons[10] and parvalbumin neurons[12]. Currently, I am also working on other aspects of psychiatry (e.g. Induced pluripotent stem cell [13], sex related depression effect and circadian pattern).

• Future direction:

- 1. Collaboration is always an essential part for statistician/biostatistician. I will definitely seek for opportunities to work with local biologists in my future tenure-track research environment, which will in turn motivate statistical methodology development.
- 2. I have worked extensively on microarray data and sequencing data. But I understand that this is a fast moving field. I am ready to learn newly developed data types, as technique advances. At this stage, I am exposed to single cell data (single cell methylation and expression), which is part of my thesis.

References

- [1] **Zhiguang Huo**, Ying Ding, Silvia Liu, Steffi Oesterreich, and George Tseng. Meta-Analytic Framework for Sparse K-Means to Identify Disease Subtypes in Multiple Transcriptomic Studies. *Journal of the American Statistical Association*, 111, no. 513 (2016): 27-42.
- [2] **Zhiguang Huo**, Chi Song, George C. Tseng. (2016) Bayesian latent hierarchical model for transcriptomic meta-analysis to detect biomarkers with clustered meta-patterns of differential expression signals. Submitted to *Annals of Applied Statistics* (under second round of review).
- [3] **Zhiguang Huo**, George C. Tseng. (2016) Integrative Sparse K-means for disease subtype discovery using multi-level omics data. Submitted to Annals of Applied Statistics (under second round of review).
- [4] Li Zhu, Ying Ding, Cho-Yi Chen, Lin Wang, **Zhiguang Huo**, SungHwan Kim, Christos Sotiriou, Steffi Oesterreich and George C. Tseng. (2016) MetaDCN: meta-analysis framework for differential coexpression network detection with an application in breast cancer. *Bioinformatics* (accepted).
- [5] SungHwan Kim, Dongwan Kang, Zhiguang Huo, Yongseok Park, George C. Tseng. (2016) Meta-analytic principal component analysis. Submitted.
- [6] Li Zhu, **Zhiguang Huo**, Tianzhou Ma and George Tseng. Bayesian indicator variable selection model with multi-layer overlapping groups. (in preparation).
- [7] Silvia Liu, Wei-Hsiang Tsai, Ying Ding, Rui Chen, Zhou Fang, Zhiguang Huo, SungHwan Kim, Tianzhou Ma, Ting-Yu Chang, Nolan Michael Priedigkeit, Adrian V. Lee, Jianhua Luo, Hsei-Wei Wang, I-Fang Chung, George C. Tseng. (2015). Comprehensive evaluation of fusion transcript detection algorithms and a meta-caller to combine top performing methods in paired-end RNA-seq data. Nucleic Acids Research, 10.1093/nar/gkv1234.
- [8] Tiffany A. Katz, Serena G. Liao, Vincent J. Palmieri, Robert K. Dearth, Thushangi Pathiraja, Zhiguang Huo, Patricia Shaw, Sarah Small, Nancy E. Davidson, David G. Peters, George C. Tseng, Steffi Oesterreich, Adrian V. Lee. (2015) Targeted DNA Methylation Screen in the Mouse Mammary Genome Reveals a Parity-Induced Hypermethylation of IGF1R That Persists Long after Parturition. Cancer Prevention Research 8, no. 10 (2015): 1000-1009.
- [9] Yan P. Yu, Silvia Liu, Zhiguang Huo, Amantha Martin, Joel B. Nelson, George C. Tseng and Jian-Hua Luo. (2015) Genomic copy number variations in the genomes of leukocytes predict prostate cancer clinical outcomes. *PloS one*, 10(8):e0135982.
- [10] Dominique Arion, **Zhiguang Huo**, John F. Enwright, John P. Corradi, George Tseng and David A. Lewis. Transcriptome alterations in prefrontal pyramidal neurons distinguish schizophrenia from bipolar and major depressive disorders. Submitted to *Biological Psychiatry*, (under second round of review).
- [11] Oesterreich, S., Katz, T.A., Logan, G., Levine, K., Nagle, A., **Huo, Z.**, Tseng, G.C., Rui, H., Lee, A.V. and Butler, L.M., 2016. Abstract PD2-08: Potential role of prolactin signaling in development and growth of the lobular subtype of breast cancer. *Cancer Research*, 76(4 Supplement), pp.PD2-08.
- [12] John Enwright, Dominique Arion, **Zhiguang Huo**, George Tseng and David A. Lewis. Transcriptome alterations in layer 3 parvalbumin neurons in the dorsolateral prefrontal cortex in schizophrenia differ from those in layer 3 pyramidal cells. (in preparation).
- [13] Logan, R.W., Ozburn, A.R., **Huo, Z.**, Zhu, X., Fitzgerald, E., Arey, R.N., Jarpe, M., Tseng, G. and McClung, C.A. (2017) Valproic acid targets HDAC2 to normalize mania-like behaviors in mice. In preparation.