

Research II allocation proposal (Multi-tissue methods and GTEx analysis)

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Research goals and potential impact

Variation in gene expression plays a crucial role in the etiology of complex human disease. Understanding the genetic factors that underpin the quantitative levels of gene expression (known as expression quantitative trait loci, or eQTL) provides intermediate insight into biological basis for disease associations identified in genetic mapping studies such as GWAS [1]. To date, most efforts on eQTL discovery have focused on their contribution to gene expression within a single type of cell or tissue. Although the importance of shared eQTL between human cells and tissues has been well acknowledged [3], multi-cell and multi-tissue analyses remain challenging due to the lack of adequate data source and analytic tools that are both statistically powerful and computationally efficient.

Recently, the Genotype Tissue Expression (GTEx) Project, a renowned global research effort aiming at understanding the role of regulatory variants in multiple tissues, have completed its Pilot Phase [2]. Our group, being part of the Statistical Genetics Committee in GTEx, had previously developed a statistical framework that improved power to detect eQTL sharing among multiple tissues [4]. However, a critical drawback of the framework is that its computational time is proportional to 2^R (R is the number of tissues); consequently it becomes intractable to even consider over 10 tissues jointly. On the other hand, however, there is an immediate urge for being capable of performing analyses involving large number of tissues, as we are faced with the challenge of analyzing the current phase of GTEx which has data on 44 tissues.

Recently we have been devising two independent statistical approaches to jointly analyze large number of tissues. Once these new methods are fully developed, they will be applied to analyzing GTEx project for discovering novel patterns of shared gene regulations in humans. Furthermore, our methods can potentially be applied to generic problems involving assessment of genetic factors across multiple data sources, a type of data integration problem of general interest in the emerging field of genetic data science.

Results from previous allocations and anticipated tasks for future allocations

Our previous statistical framework for joint tissue analysis had been applied to the GTEx pilot phase involving 9 tissues. Computations were performed on the Midway cluster. The result have contributed to the *Science* paper published in May, 2015 [2].

For large tissue problem posed by the current GTEx data we are developing two approaches to model, respectively, the qualitative and quantitative heterogeneity among tissues using lower dimensional spaces of probability measures, thus avoiding a growth in computation from 2^9 to 2^{44} . We are currently finishing up the theoretical portion and a proof-of-concept computer implementation of the new methods. However development of high quality software for the new methods are still in progress and we are yet to perform comprehensive evaluation and comparison of the two approaches in simulation studies, as well as to complete genome-wide joint eQTL mapping in 44 GTEx tissues.

In preparing for analysis-ready data-sets, we have developed an efficient data management framework based on HDF5 file system. Approximately 50 billion lines of text files have been processed to create the

data infrastructure for GTEx release in April, 2015. We are expecting a complete update of source data, and consequently the need to rebuild the data system, in October, 2015.

Estimate of requested resources

Based on SU consumption in the method development phase last quarter (70,000 SUs) we estimate a usage of 550,000 SUs and 1TB of storage upon completion of the project in the next 12 months. The SU requested will be dedicated to mainly 3 tasks: data processing, methods development and comparison and GTEx analysis.

For data-set per tissue it requires 450 SUs (16–20 CPUs on one node with 1GB memory per CPU) to perform genome-wide single tissue analysis and organizing the outcome into HDF5 system. It requires another 100 SUs (1 CPU on one node with 100 - 200GB memory) to merge and compute summary statistics from output across all tissues. Since the input information for the two methods we are developing are partially different, an additional 20% SUs is required to create companion data-sets to provide information specifically required by different methods. In sum the data processing step have consumed 24,000 SUs from previous allocation. The data to be released in October 2015 is a milestone update in GTEx project, approximately twice in size compared to current release. We therefore estimate a requirement of 48,000 SUs for the upcoming data processing task. Our data is highly compressed (in `bz2` format with level 9 compression), currently taking about 300GB in space. We expect 600GB storage requirement for the October release.

For the quantitative model we are developing, it requires 200 SUs (16–20 CPUs on one node) to fit the model on 50,000 data points, which is about 0.05% of the total data set. In continued development of the method we will experiment with 0.05% data to improve model parameter configurations (SUs required can be negligible). The ultimate goal is to apply the model on 50% data points which we believe will provide good representation for the entire data. The expected resource usage is 200,000 SUs.

For the qualitative model we need 1% data points to provide reliable experimental results. Depending on the desired accuracy of convergence in our algorithm it requires 1,000 – 3,000 SUs (16 – 20 CPUs on one node with 200G memory total) to complete the paralleled portion of computation involved. Computational implementation of the remaining part of algorithm is not yet paralleled, but we estimate a requirement of 2,000 - 6,000 SUs in conducting it. In tuning the model and implementing its parallel version, we expect non-trivial SU consumption (about 20,000 SUs) on experiments. Once we apply the model to 50% data points we expect usage of 170,000 – 470,000 SUs. Thus we request 300,000 SUs for data analysis under this model.

Substantial amount of output data for each gene and regulatory variant pair across genome will be generated in the joint GTEx data analysis. Given that intermediate results are constantly removed as the analyses move on, we ultimately expect about 300GB for storing the genome-wide summary statistics.

References

- [1] Frank W. Albert and Leonid Kruglyak. “The role of regulatory variation in complex traits and disease”. In: *Nature Reviews Genetics* 16.4 (Feb. 2015), pp. 197–212. ISSN: 1471-0056. DOI: 10.1038/nrg3891. URL: <http://dx.doi.org/10.1038/nrg3891>.
- [2] K. G. Ardlie et al. “The Genotype-Tissue Expression (GTEx) pilot analysis: Multitissue gene regulation in humans”. In: *Science* 348.6235 (May 2015), pp. 648–660. ISSN: 0036-8075. DOI: 10.1126/science.1262110. URL: <http://www.sciencemag.org/content/348/6235/648.abstract>.

- [3] Antigone S Dimas et al. “Common regulatory variation impacts gene expression in a cell type-dependent manner.” In: *Science (New York, N.Y.)* 325.5945 (Sept. 2009), pp. 1246–50. ISSN: 1095-9203. DOI: 10.1126/science.1174148. URL: <http://www.sciencemag.org/content/325/5945/1246.long>.
- [4] Timothée Flutre, Xiaoquan Wen, Jonathan Pritchard, and Matthew Stephens. “A Statistical Framework for Joint eQTL Analysis in Multiple Tissues”. In: *PLoS Genetics* 9.5 (2013), pp. 1–8. ISSN: 15537390. DOI: 10.1371/journal.pgen.1003486. arXiv: [arXiv:1212.4786v1](https://arxiv.org/abs/1212.4786v1).