

eQTLs, genetics of other genomic variations

VJ Carey

CSAMA 2013

Overview

- Some recent literature
- A view of the workflow for cis-eQTL
- Prospects for generalization

2013

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Integrative Modeling of eQTLs and Cis-Regulatory Elements Suggests Mechanisms Underlying Cell Type Specificity of eQTLs

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Abstract

Genetic variants in cis-regulatory elements or trans-acting regulators frequently influence the quantity and spatiotemporal distribution of gene transcription. Recent interest in expression quantitative trait locus (eQTL) mapping has paralleled the adoption of genome-wide association studies (GWAS) for the analysis of complex traits and disease in humans. Under the hypothesis that many GWAS associations tag non-coding SNPs with small effects, and that these SNPs exert phenotypic control by modifying gene expression, it has become common to interpret GWAS associations using eQTL data. To fully exploit the mechanistic interpretability of eQTL-GWAS comparisons, an improved understanding of the genetic architecture and causal mechanisms of cell type specificity of eQTLs is required. We address this need by performing an eQTL analysis in three parts: first we identified eQTLs from eleven studies on seven cell types; then we integrated eQTL data with cis-regulatory element (CRE) data

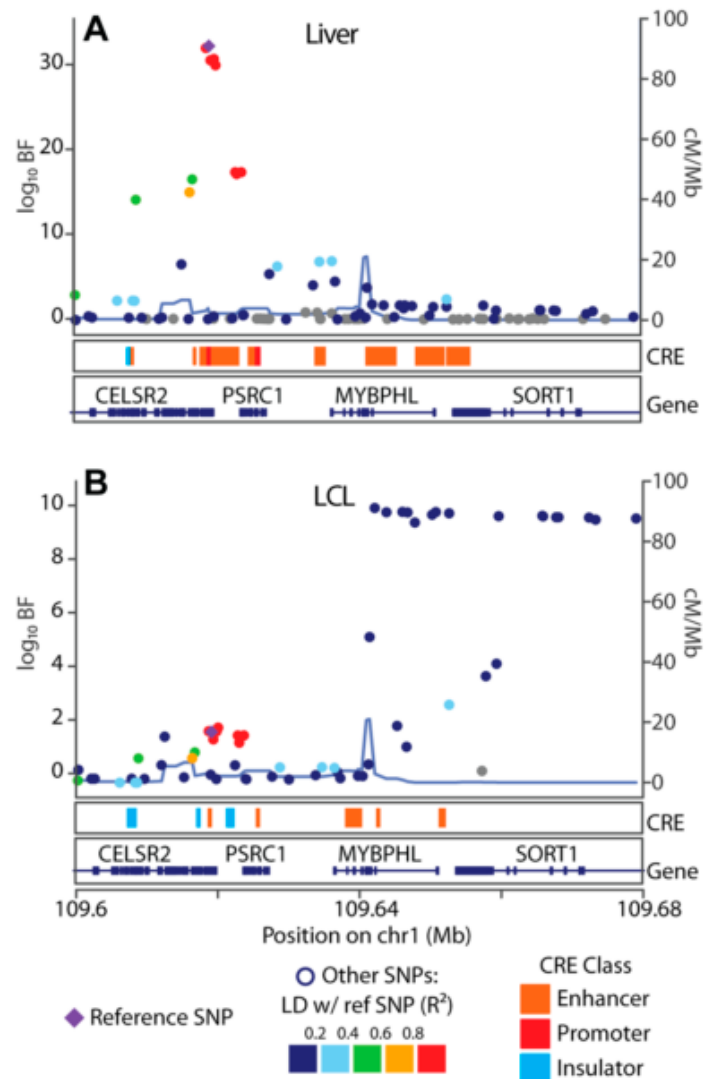


Figure 6. SORT1 eQTL illustrates mechanisms underlying cell specificity of eQTLs. Associations between (A) UChicago_liver and (B) CAP_LCL SORT1 expression and cis-linked SNPs (left y-axis; log₁₀ BF), plotted as points by SNP genomic coordinates (x-axis). Blue line

Query: What are the Bioconductor resources that could be used to develop these statistics and Visualizations?

2014

Common Genetic Variants Modulate Pathogen-Sensing Responses in Human Dendritic Cells

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Little is known about how human genetic variation affects the responses to environmental stimuli

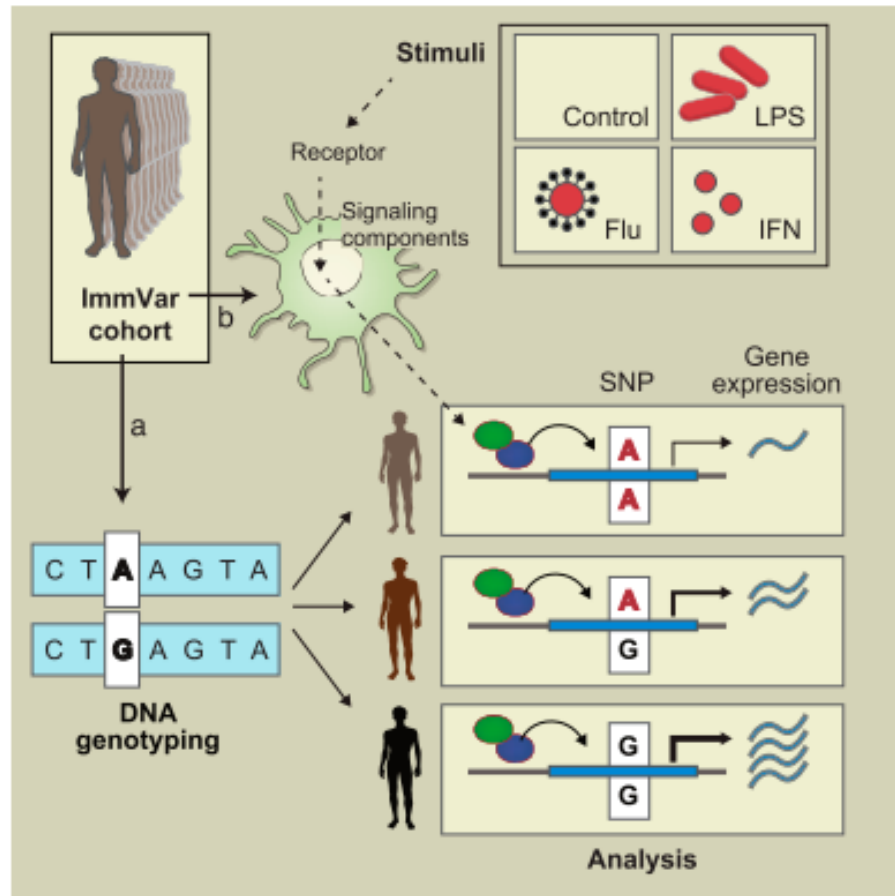
cleus to induce the expression of immune genes, including interferon- β (IFN- β) secretion that engages the type I IFN response pathway to induce the expression of hundreds of antiviral effectors. Genetic studies have associated common variants near many genes in these pathways with risk of different inflammatory diseases (11, 12). DCs also play a critical role in the pathologic immune responses underlying inflammatory diseases (11–13), also reflected in recent genome-wide association studies (GWAS) of several diseases (14–17), especially inflammatory bowel disease (14).

Results

Assessing the Impact of Genetic Variation on Pathogen Sensing in Primary Human DCs

We developed an integrated experimental and

“reQTL”: ‘response to stimulus’



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Research Article

B. P. Fairfax *et al.*, *Science* **343**, 1246949 (2014).

Perspective

P. K. Gregersen, *Science* **343**, 1087 (2014).

Identifying the genetic basis of variability in the host response to pathogens. A cohort of 534 individuals donated blood for (a) genotyping of common DNA variants and (b) isolation of immune DCs. DCs were stimulated with viral and bacterial components, and the variability in individuals' gene expression responses was mapped to specific DNA variants, which were then shown to affect binding of particular transcription factors.

Data flow for reQTL

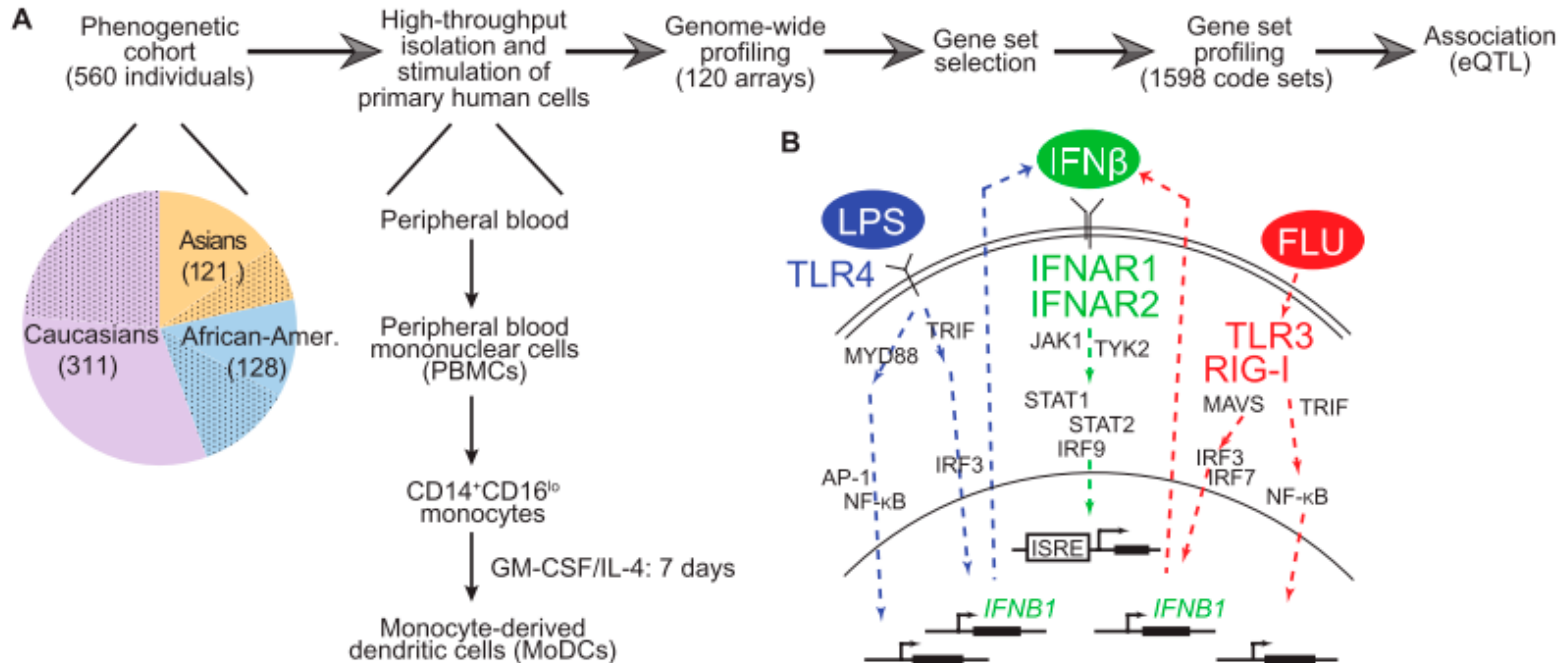
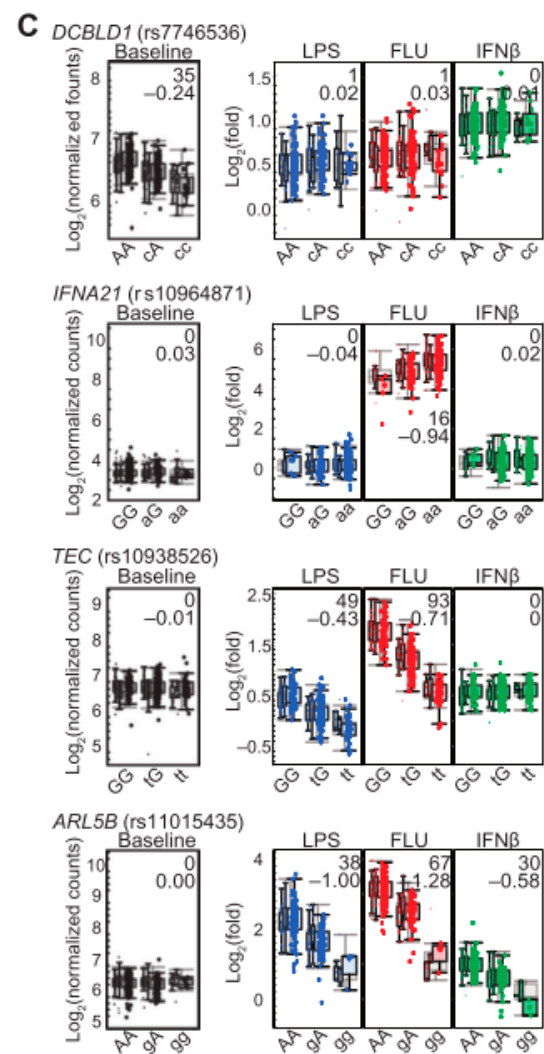
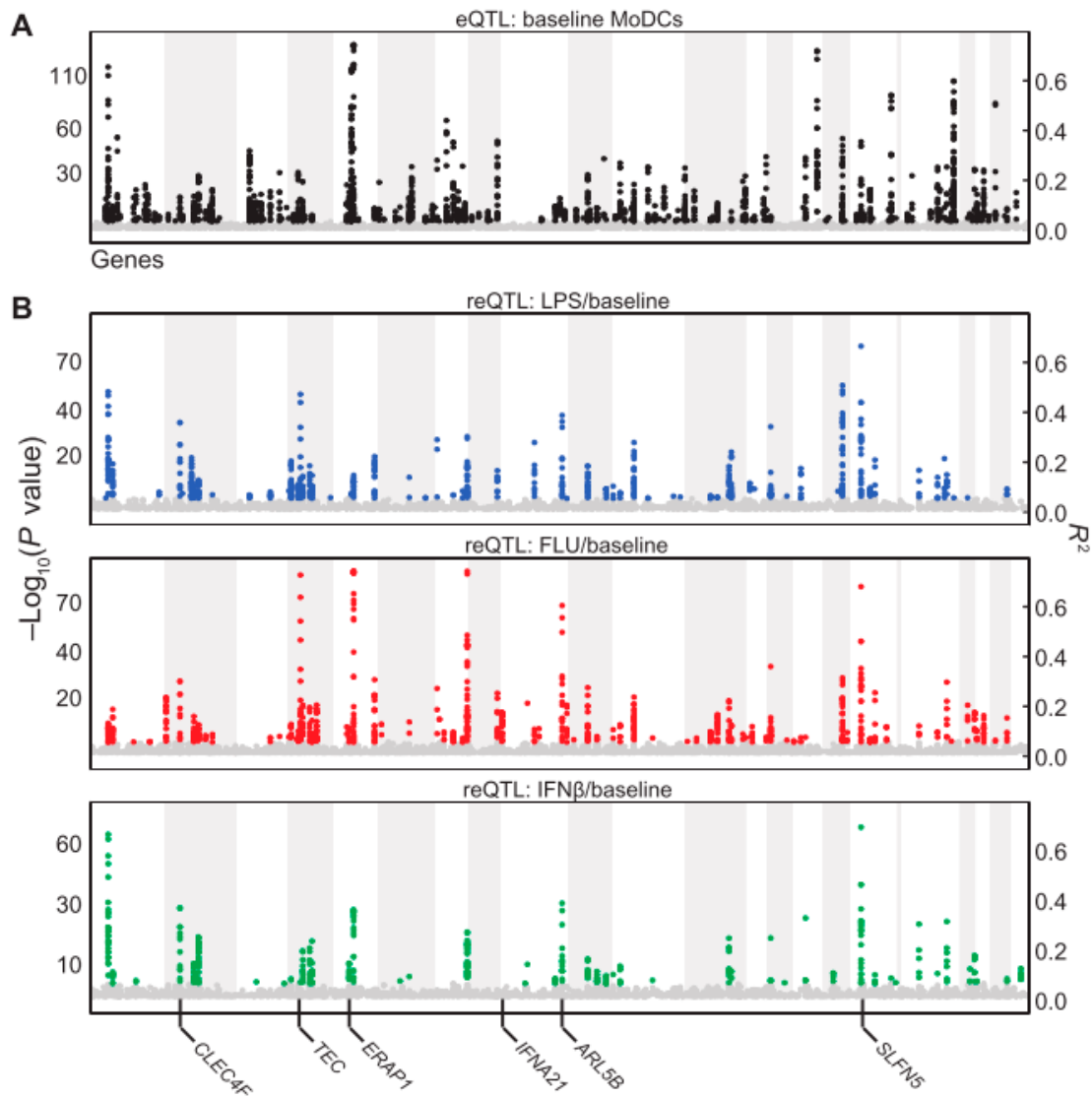


Fig. 1. A strategy to identify gene-by-environment interactions in the innate immune responses of primary human DCs. (A) Strategy used to identify baseline and response eQTLs and reQTLs, consisting of five steps: (i) high-throughput isolation and stimulation of primary human MoDCs from 560 healthy individuals (dotted slices, male; solid-colored slices, female) collected as part of the PhenoGenetic cohort; (ii) whole-genome gene expression measurements in a subset of the cohort; (iii) selection of signature gene set, consisting of regulators and regulated genes; (iv) digital multiplex gene expression measurements of signature genes in the entire cohort; and (v) mapping of genetic variation to

expression variation. GM-CSF, granulocyte-macrophage colony-stimulating factor; IL-4, interleukin-4. **(B)** Model of innate immune pathways activated by three stimuli demonstrating their downstream relationships. LPS from *E. coli* engages the TLR4 receptor; IFN- β engages the heterodimeric IFNAR receptor; influenza A/PR8 (Δ NS1) ("FLU") engages the cytoplasmic TLR3 and RIG-I receptors. Receptor engagement activates signal transduction cascades that regulate expression of inflammatory genes, IFNs, and IFN-stimulated genes. IFNAR activation also occurs during LPS and FLU stimulations because LPS and FLU both induce IFN production, leading to activation of ISREs. JAK1, Janus kinase 1.



How can we do the basic computations with Bioconductor?

- <http://bioconductor.org/help/workflows/eQTL/>