

# Linking Genetic Variation to Important Phenotypes:

SNPs, CNVs, GWAS, and eQTLs

BMI/CS 776

[www.biostat.wisc.edu/bmi776/](http://www.biostat.wisc.edu/bmi776/)

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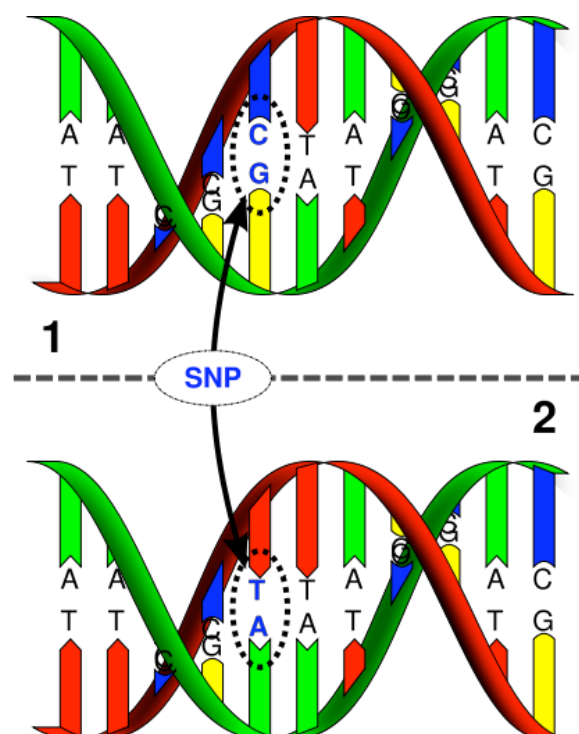
## 1. Understanding Human Genetic Variation

- the “human genome” was determined by sequencing DNA from a small number of individuals
- the HapMap project looked at polymorphisms in 270 individuals
- the 1000 Genomes project is sequencing the genomes of 1000 individuals from diverse populations

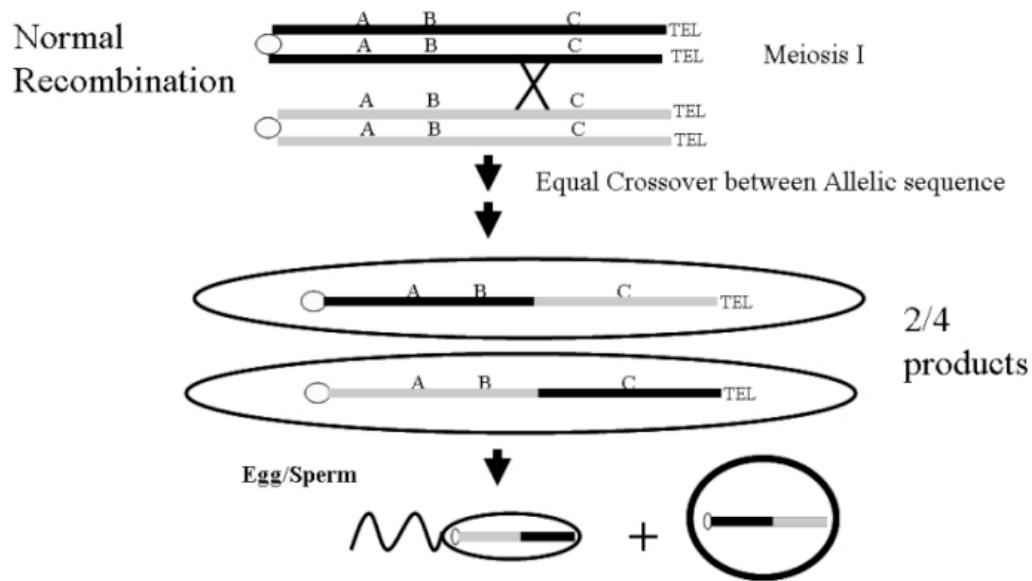
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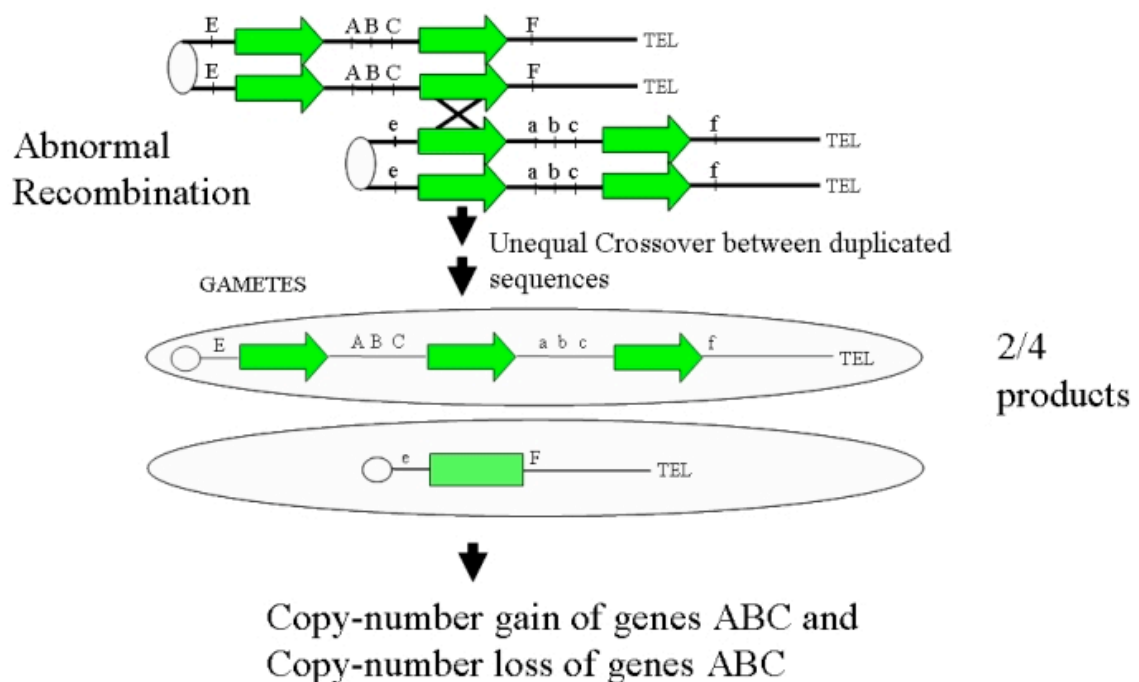
## Genetic Variation: Single Nucleotide Polymorphisms (SNPs)



## Genetic Variation: Copy Number Variants (CNVs)

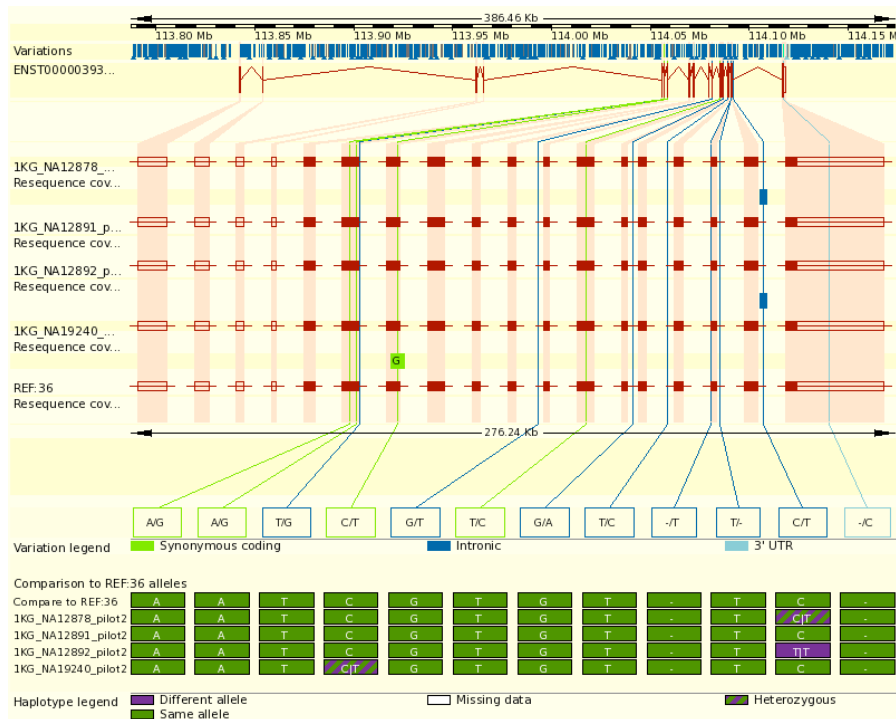


## Genetic Variation: Copy Number Variants (CNVs)



# 1000 Genomes Project

project goal: produce a catalog of human variation down to variants that occur at 1% frequency or less over the genome

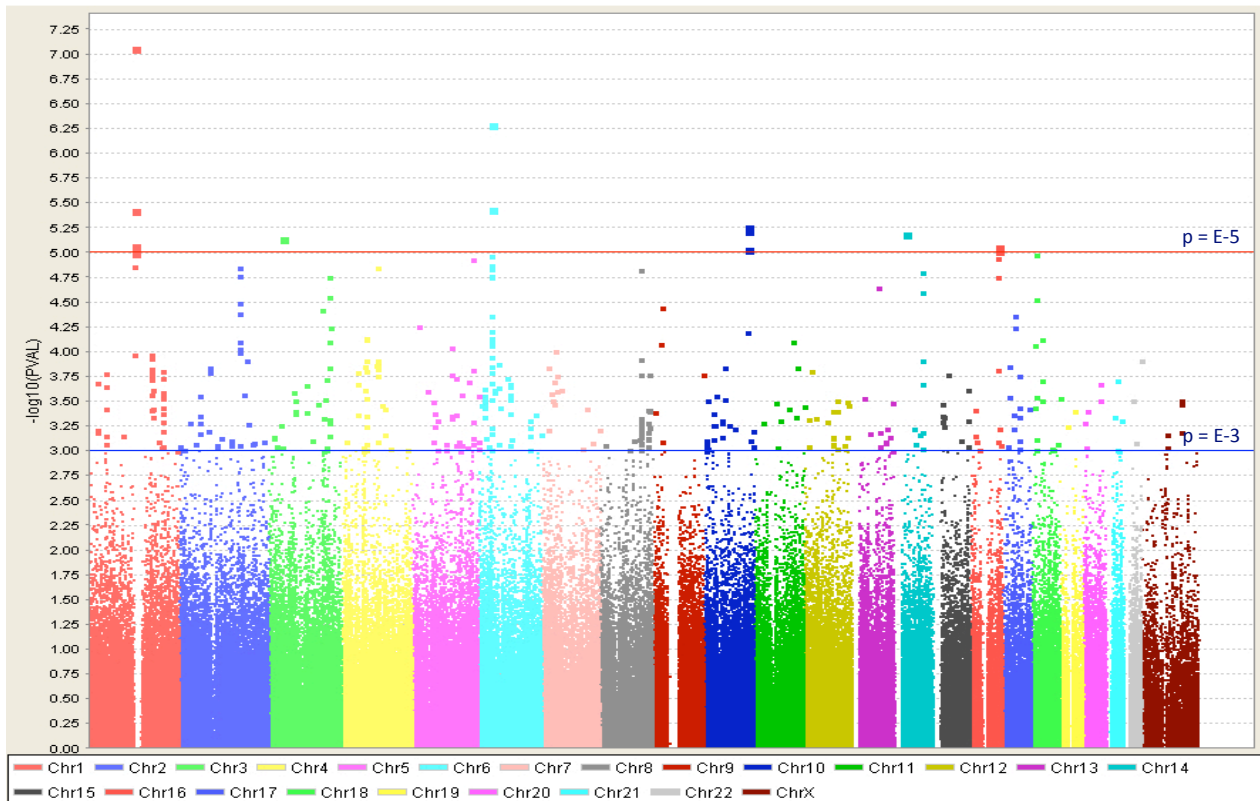


## 2. Understanding Associations Between Genetic Variation and Disease

*genome wide association study (GWAS)*

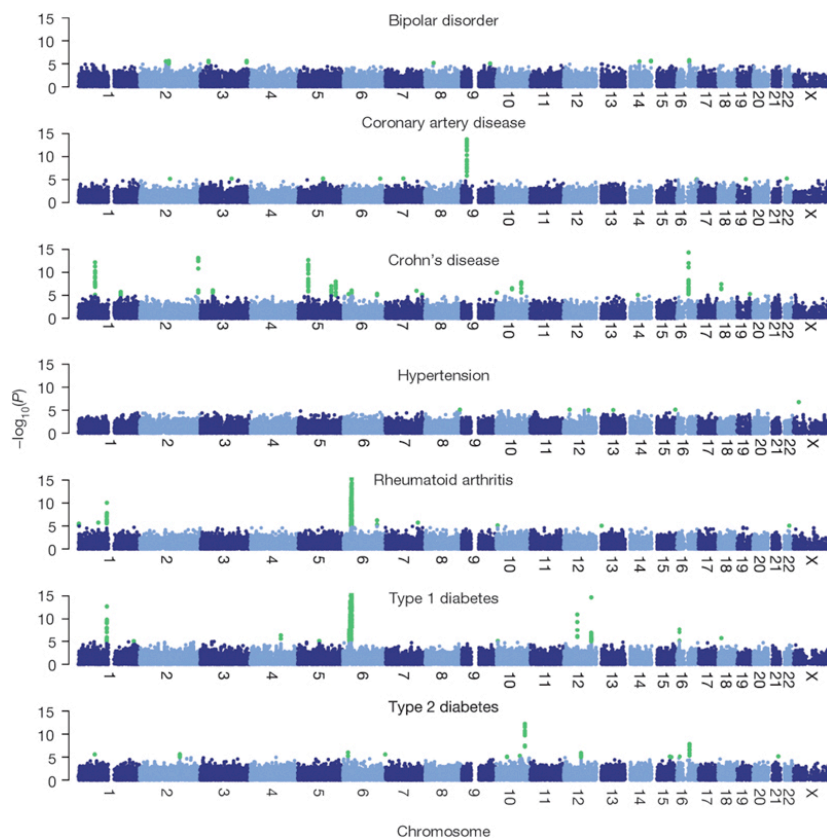
- gather some population of individuals
- genotype each individual at polymorphic markers (usually SNPs)
- test association between *state* at marker and some variable of interest (say disease)
- adjust for multiple comparisons

## Type 2 Diabetes Results: 386,731 markers



Type 2 diabetes association P values by chromosome (386,731 markers). The x-axis is the genomic position by chromosome 1-22 and X (by color), and the y-axis is the negative base 10 logarithm of the P value.

## Wellcome Trust GWAS

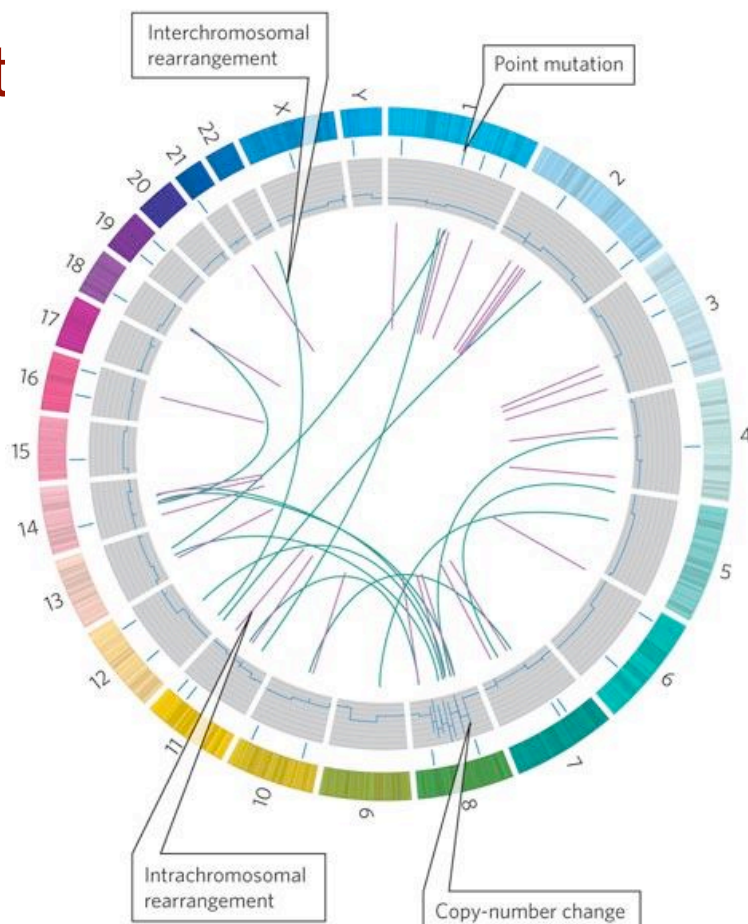


## 2. Understanding Associations Between Genetic Variation and Disease

### International Cancer Genome Consortium

- includes the NIH's *Cancer Genome Atlas*
- sequencing DNA from 500 tumor samples for each of 50 different cancers
- goal is to distinguish *drivers* (mutations that cause and accelerate cancers) from *passengers* (mutations that are byproducts of cancer's growth)

### A Circos Plot



# Some Cancer Genomes

## LUNG CANCER

Cancer: small-cell lung carcinoma

- Sequenced: full genome
- Source: NCI-H209 cell line
- Point mutations: 22,910
- Point mutations in gene regions: 134
- Genomic rearrangements: 58
- Copy-number changes: 334



## SKIN CANCER

Cancer: metastatic melanoma

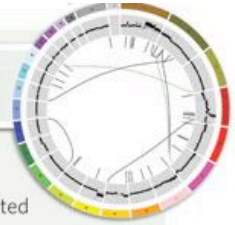
- Sequenced: full genome
- Source: COLO-829 cell line
- Point mutations: 33,345
- Point mutations in gene regions: 292
- Genomic rearrangements: 51
- Copy-number changes: 41



## BREAST CANCER

Cancer: basal-like breast cancer

- Sequenced: full genome
- Source: primary tumour, brain metastasis, and tumours transplanted into mice
- Point mutations: 27,173 in primary, 51,710 in metastasis and 109,078 in transplant
- Point mutations in gene regions: 200 in primary, 225 in metastasis, 328 in transplant
- Genomic rearrangements: 34
- Copy-number changes: 155 in primary, 101 in metastasis, 97 in transplant



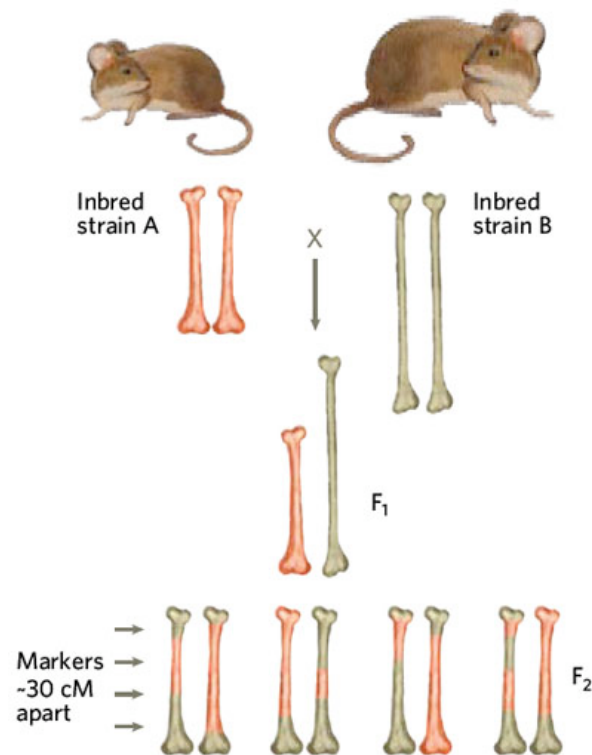
## 3. Understanding Associations Between Genetic Variation and Complex Phenotypes

quantitative trait loci (QTL) mapping

- gather some population of individuals
- genotype each individual at polymorphic markers
- map quantitative trait(s) of interest to chromosomal locations that seem to explain variation in trait

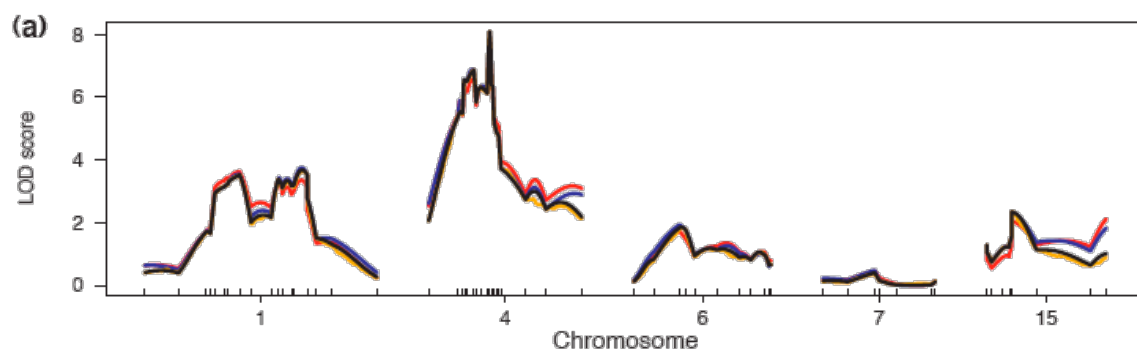


# QTL Mapping Example



# QTL Mapping Example

- QTL mapping of mouse blood pressure, heart rate [Sugiyama et al., Broman et al.]



$$\text{LOD}(q) = \log_{10} \frac{P(q | \text{QTL at } m)}{P(q | \text{no QTL at } m)}$$

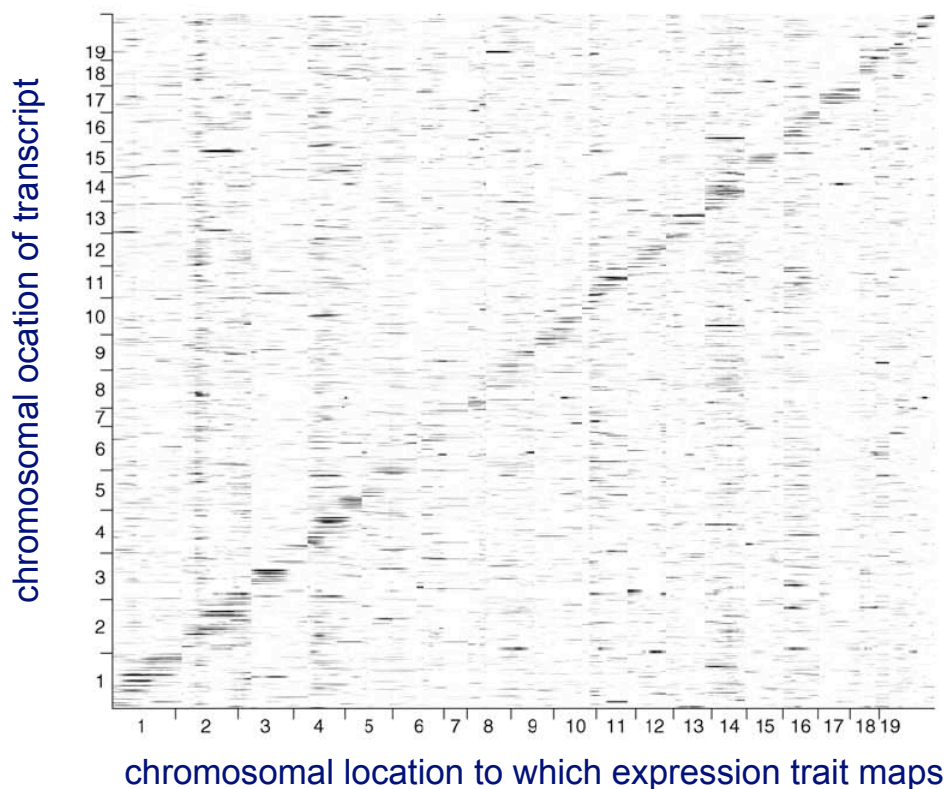


# QTL Mapping

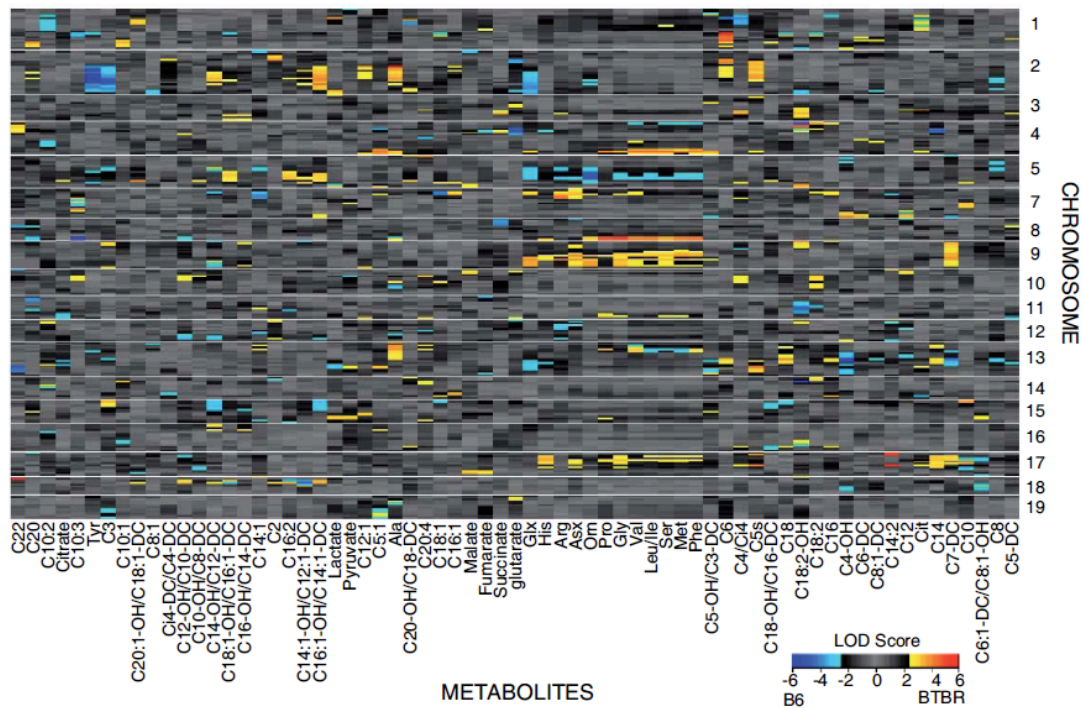
- QTL mapping can be done for large-scale quantitative data sets
  - expression QTL: traits are expression levels of various genes
  - metabolic QTL: traits are metabolite levels
- case study: uncovering the genetic/metabolic basis of diabetes (Attie lab at UW)



## *Expression* QTL (eQTL) Mapping



# Metabolite QTL (mQTL) Mapping



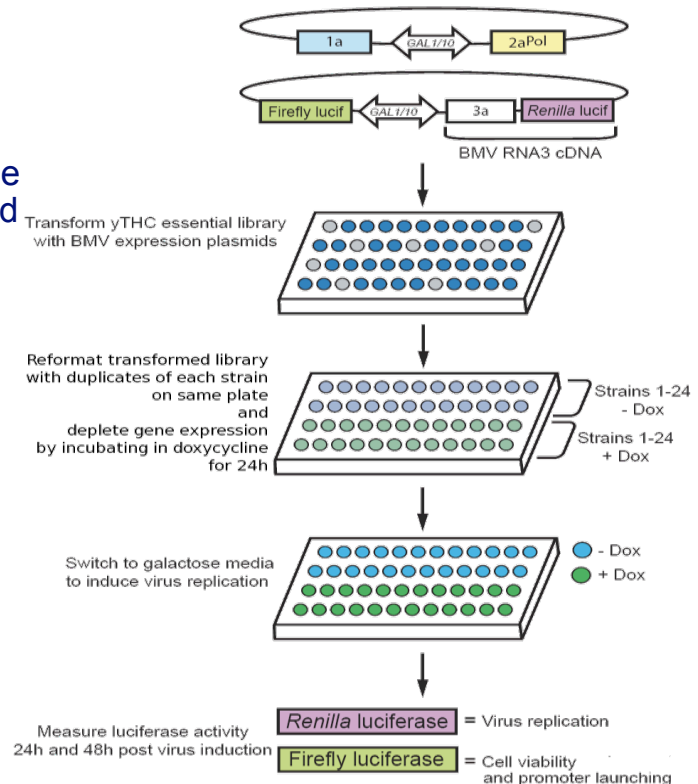
## 4. Understanding Associations Between Genetics and Phenotypes via High-Throughput Manipulations

can measure response of interest by suppressing/mutating genes on a genome-wide scale

- mutant libraries (i.e. knock out each gene)
- siRNA libraries (i.e. suppress the transcripts of each gene)

# Inferring Models of Host-Virus Interactions

- Brome mosaic virus (BMV) is a positive-strand RNA virus whose replication, gene expression and virion formation has been recapitulated in yeast
- The Ahlquist lab at UW has identified host genes affecting BMV replication via a genome-wide screen with the yeast deletion library

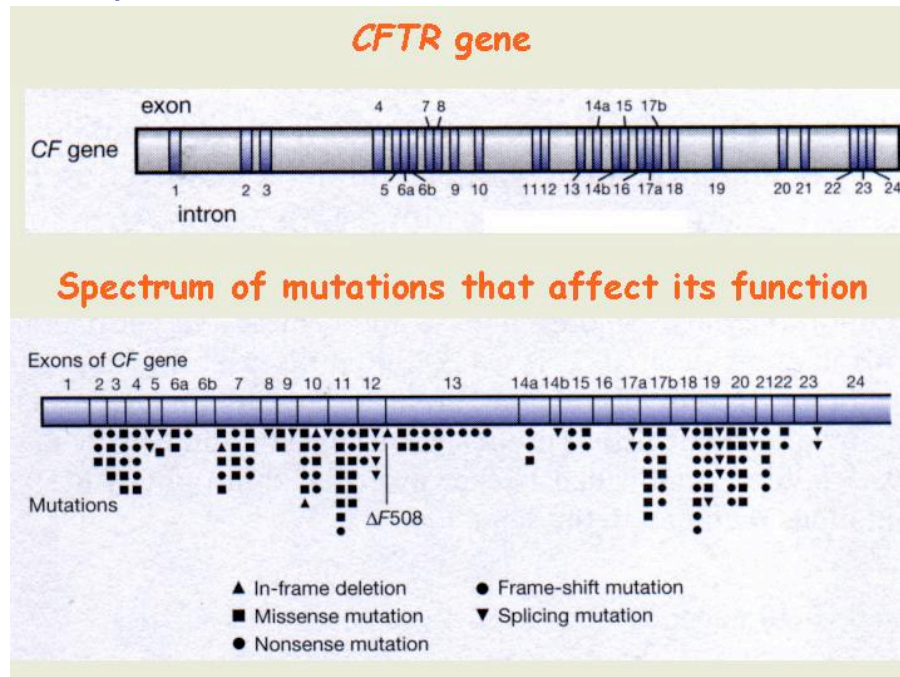


## Computational Problems

- assembly and alignment of thousands of genomes
- identifying functional roles of markers of interest (which genes/pathways does a mutation affect and how?)
- identifying interactions in multi-allelic diseases (which combinations of mutations lead to a disease state?)
- identifying genetic/environmental interactions that lead to disease
- inferring network models that exploit all sources of evidence: genotype, expression, metabolic

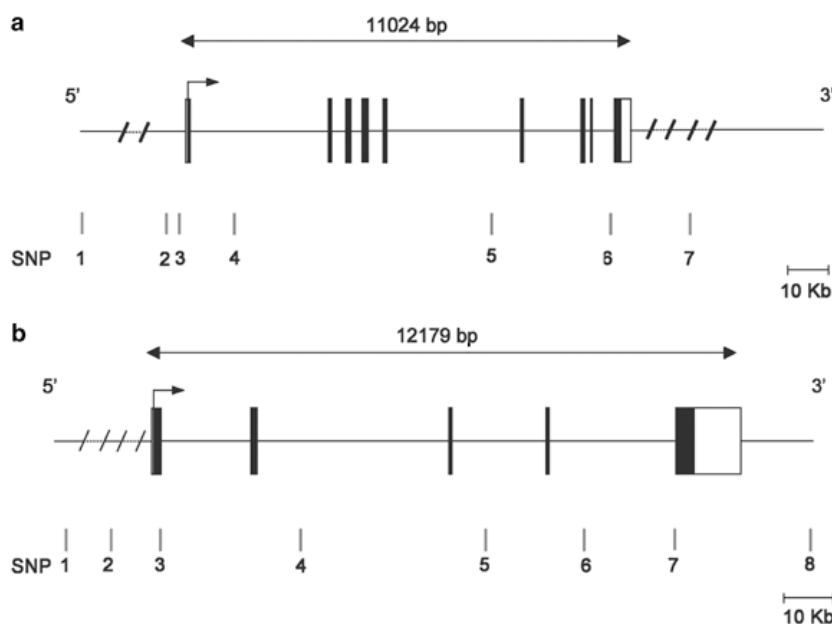
# Is determining association the end of the story?

a simple case: CFTR



## Many measured SNPs not in coding regions

- genes encoding CD40 and CD40L with relative positions of the SNPs studied



# Expression QTL (eQTL) Mapping

