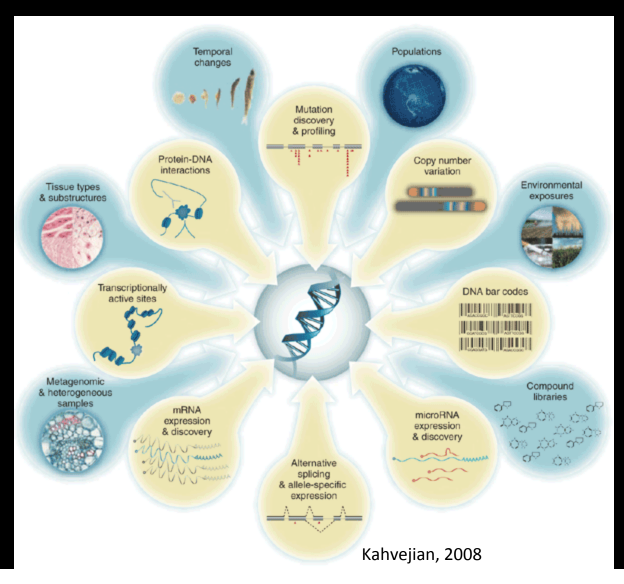
## Use of Bayesian Statistical Techniques in High-throughput Sequencing

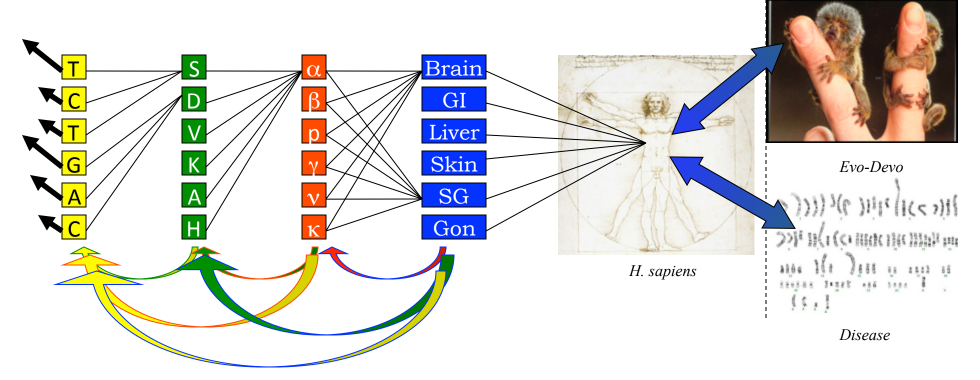
```
catttqtattatccctcttccta*caaacacactqtccqca ACGCACTCTCCATTGTTACTGCAGATTTCTGAACT
AATTTCATTTGTATTATCCCTCTTCCTA*CAAACACACTGTCCGCAG cqqaqtctccattqttactqcagatttctgaa GTTTTCTTTCCTGCAGTAAGCATCCAT
 TTGTATTATCCCTCTTCCTA*CAAACACACTGTCCGCAGACGCAC CTCCATTGTTACTGCAGATTTCTGAACTGTTTTCTTTCCTGCAGTAAGCATCCATG
  catttqtattatccctcttccta*caaacactqtccqcaqacqcac CTCCATTGTTACTGCAGATTTCTGAACTGTTTTCT
  ATTTGTATTATCCCTCTTCCTA*CAAACACACTGTCCGCAGACGCACTCTC ttgttactgcagatttctgaactgttttcttt tgcagtaagcatccatgtcttcact
                                               ntaactttttctttactgcagtaaacatccatgtcttcact
  ttgtattatccctcttccta*caaacacactgtccg agacgcactctccattgttactgcagatttct
  CAT tgtattatccctcttccta*caaacacactgtc gcagacgcactctccattgttactgcagattt gaactgttttctttcctgcagtaagcatcca
     qtattatccctcttccta*caaacacactqtcc caqacqcactctccattqttactqcaqatttc aactqttttctttcctqcaqtaaqcatccatq
      tattatccctcttccta*caaacacactgtccg agacgcactctccattgctactgcagatttct
  CATTTGTATTATCCCTCTTCCTA*CAAACACACTGTCCGCAGACGCACTCTCCAT ttactgcagatttctgaactgt
        TATCCCTCTTCCTA*CAAACACACTGTCCGCAGACGCACTCTCCATTGT ctgcagatttctgaactgttttctttcctgca aagcatccatgtc
  Assistant Professor
                The Institute for Computational Biomedicine,
               Department of Physiology and Biophysics at
                 Weill Cornell Medical College, and the
    Tri-I Program (Cornell, MSKCC, Rockefeller) on Comp. Bio. & Medicine
                            February 21st, 2012
```

Since DNA defines the biochemical recipe for the genesis of organisms, sequencing allows us to create molecular portraits of development and disease at single-base resolution.



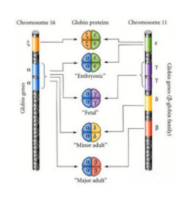


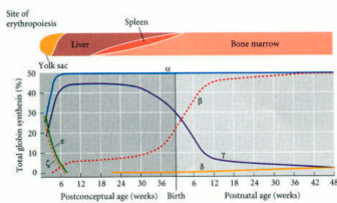
## Understanding the genome's mutation, selection, and/or drift



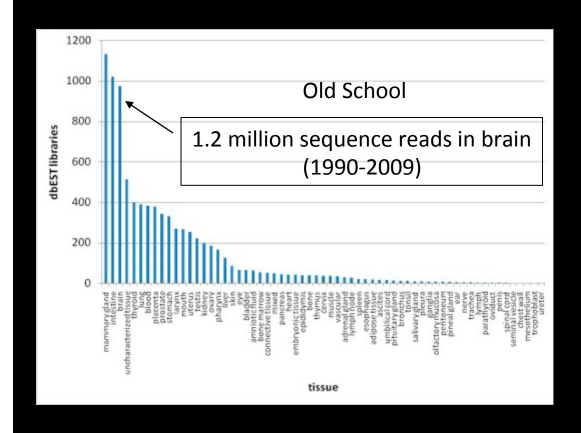
How can we understand these interactions?

Integrated, spatiotemporal molecular profiling.





## What erudition do we have now on the functional elements?



- Currently limited amount of EST info at NCBI
- ➤ EST data is expensive, timeconsuming (cloning), and exhibits 3' bias.
- Much EST and cDNA data is for whole brains, and few libraries exist with region-specific data.

## New School: One run of a NGS machine = billions of sequence reads in days

# Description/Discussion of the Various Technologies

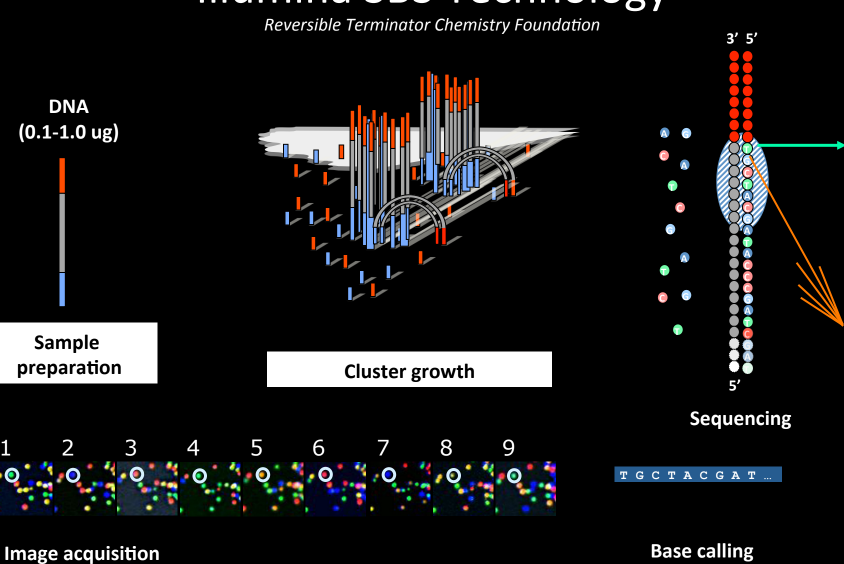
- The goal of the Archon X prize in Genomics is to enable a \$1,000 genome,
- Currently at \$3,000-\$50,000
- Certain platforms are better suited for certain tasks:
  - Counting applications (ChIP-Seq, RNA-Seq) need more reads
  - De novo assembly work needs longer reads
  - Whole genome re-sequencing requires lower errors rate and high processivity

#### But, there are many options:

Platform	Library/ template preparation	NGS chemistry	Read length (bases)	Run time (days)	Gb per run	Machine cost (US\$)	Pros	Cons	Biological applications	Refs
Roche/454's GS FLX Titanium	Frag, MP/ emPCR	PS	330*	0.35	0.45	500,000	Longer reads improve mapping in repetitive regions; fast run times	High reagent cost; high error rates in homo- polymer repeats	Bacterial and insect genome de novo assemblies; medium scale (<3 Mb) exome capture; 16S in metagenomics	D. Muzny, pers. comm.
Illumina/ Solexa's GA <sub>II</sub>	Frag, MP/ solid-phase	RTs	75 or 100	4 <sup>‡</sup> , 9 <sup>§</sup>	18‡, 35 <sup>§</sup>	540,000	Currently the most widely used platform in the field	Low multiplexing capability of samples	Variant discovery by whole-genome resequencing or whole-exome capture; gene discovery in metagenomics	D. Muzny, pers. comm.
Life/APG's SOLiD 3	Frag, MP/ emPCR	Cleavable probe SBL	50	7 <sup>‡</sup> , 14 <sup>§</sup>	30 <sup>‡</sup> , 50 <sup>§</sup>	595,000	Two-base encoding provides inherent error correction	Long run times	Variant discovery by whole-genome resequencing or whole-exome capture; gene discovery in metagenomics	D. Muzny, pers. comm.
Polonator G.007	MP only/ emPCR	Non- cleavable probe SBL	26	5 <sup>§</sup>	12 <sup>§</sup>	170,000	Least expensive platform; open source to adapt alternative NGS chemistries	Users are required to maintain and quality control reagents; shortest NGS read lengths	Bacterial genome resequencing for variant discovery	J. Edwards, pers. comm.
Helicos BioSciences HeliScope	Frag, MP/ single molecule	RTs	32*	8 <sup>‡</sup>	37‡	999,000	Non-bias representation of templates for genome and seq-based applications	High error rates compared with other reversible terminator chemistries	Seq-based methods	91
Pacific Biosciences (target release: 2010)	Frag only/ single molecule	Real-time	964*	N/A	N/A	N/A	Has the greatest potential for reads exceeding 1 kb	Highest error rates compared with other NGS chemistries	Full-length transcriptome sequencing; complements other resequencing efforts in discovering large structural variants and haplotype blocks	S. Turner, pers. comm.

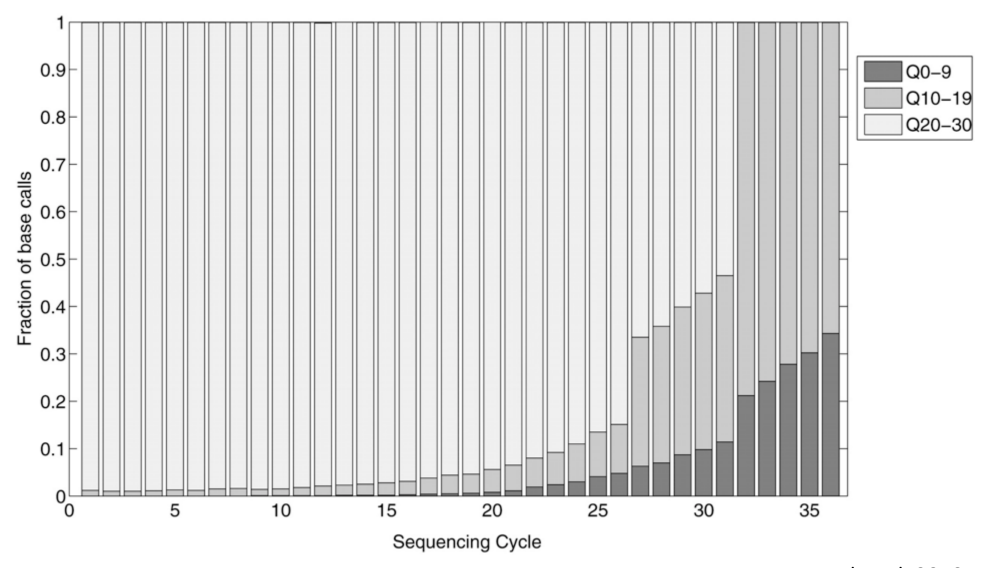
Michael Metzker, 2010

#### Illumina SBS Technology



http://www.illumina.com/technology/sequencing\_technology.ilmn

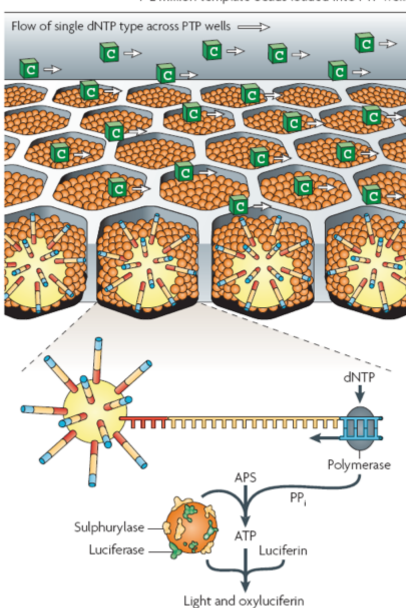
## **Quality Scores vary**

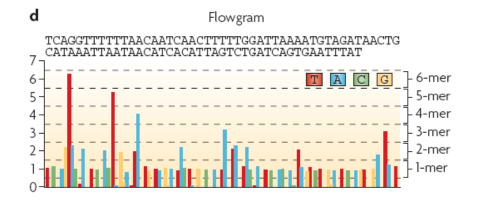


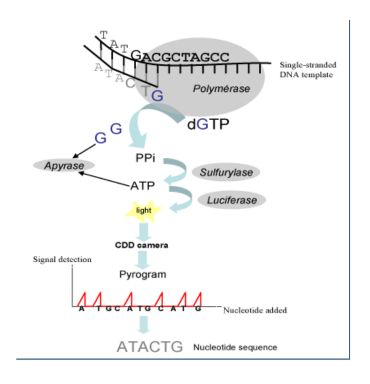
#### Pyrosequencing

#### Roche/454 — Pyrosequencing

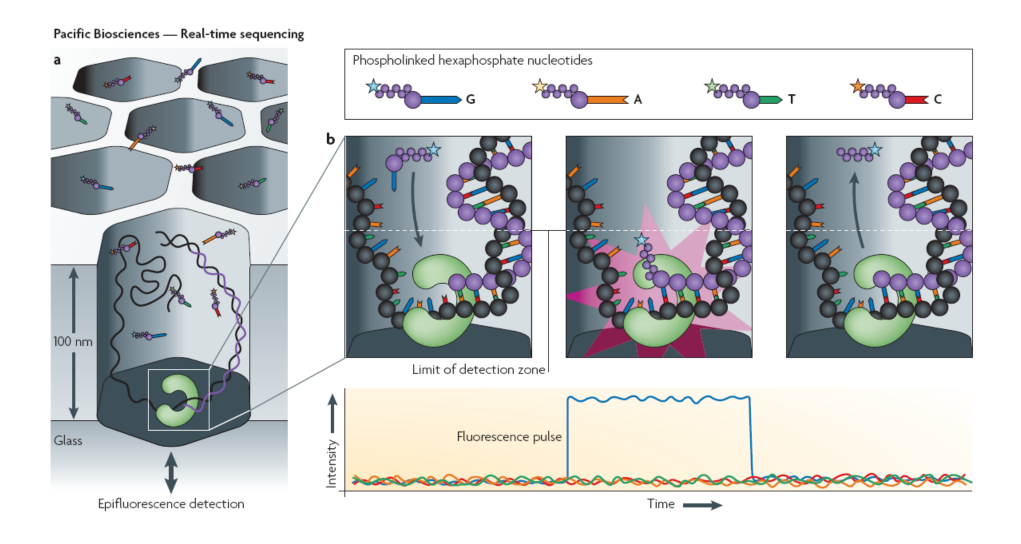
1-2 million template beads loaded into PTP wells







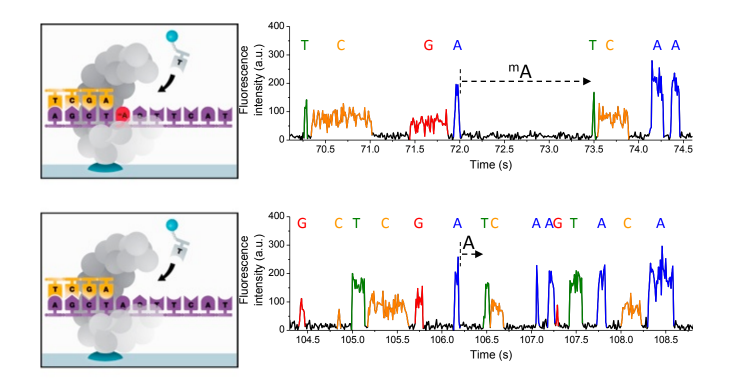
#### Single Molecule Real-Time



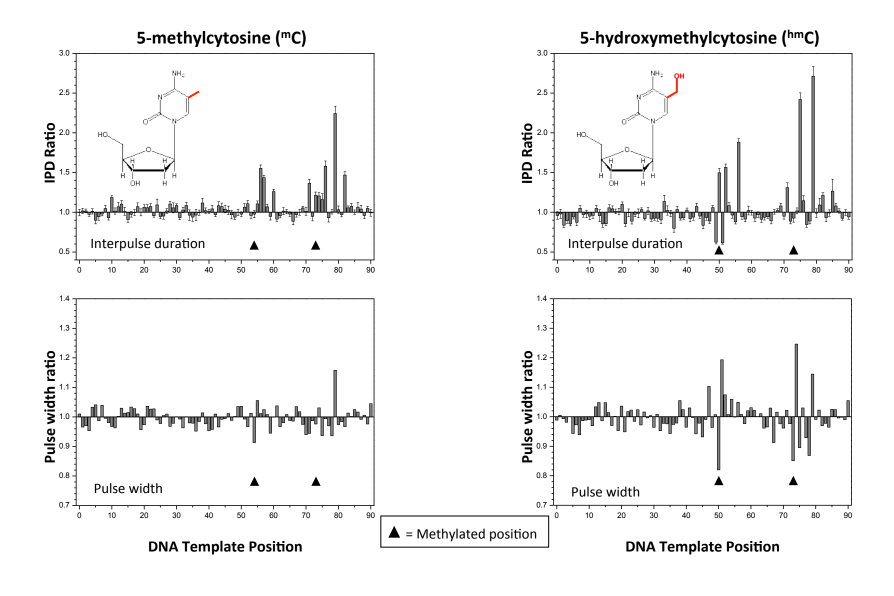
#### Direct Detection of Methylation

Approach: Kinetic detection of methylated bases during SMRT DNA sequencing

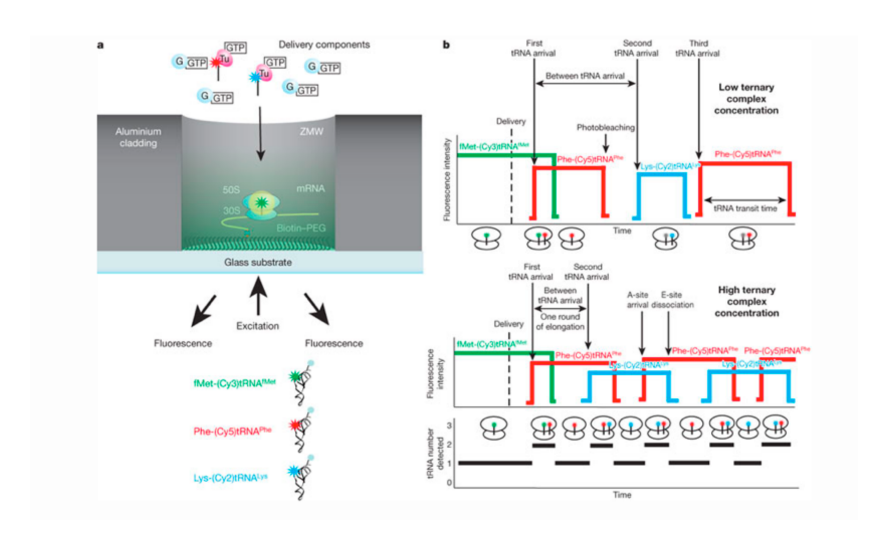
Example: N<sup>6</sup>-methyladenosine (<sup>m</sup>A)



#### detect other base modifications

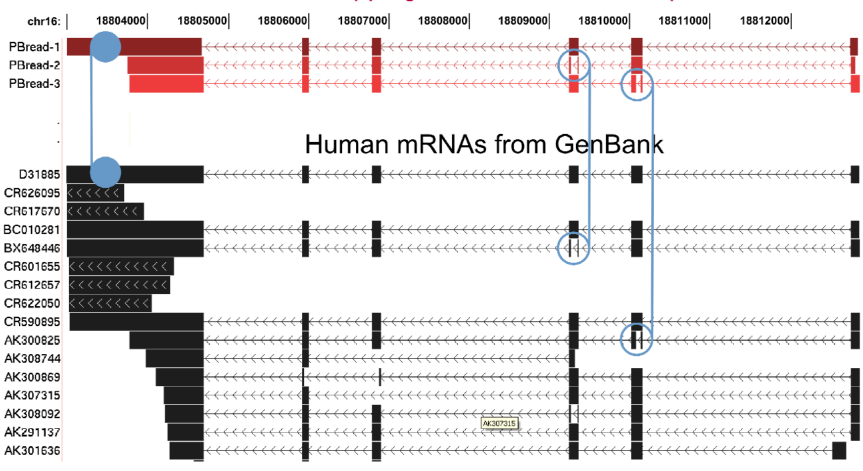


#### Watch Translation in Real-Time

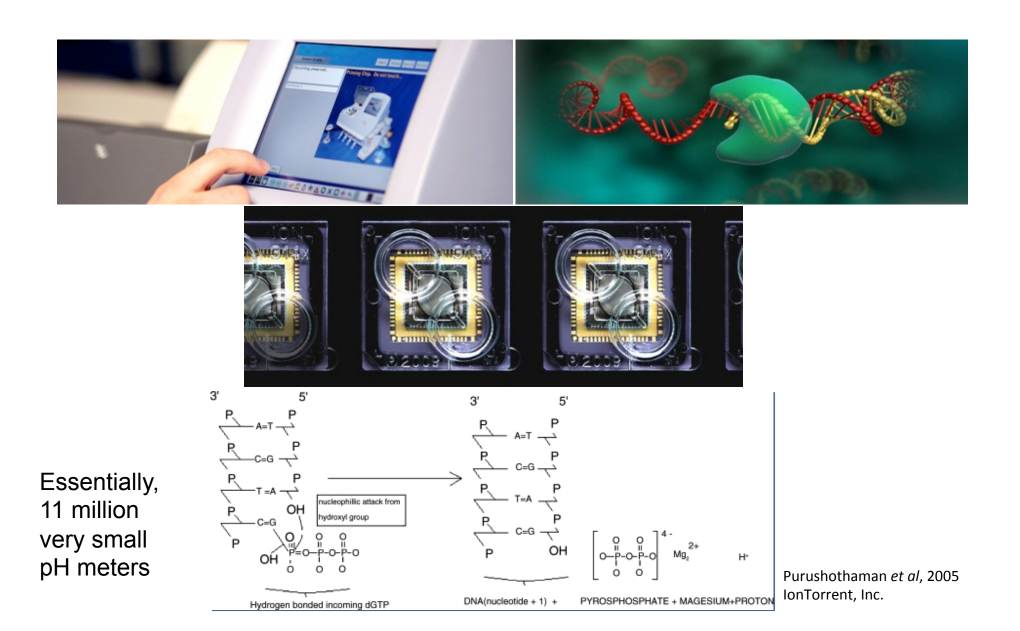


#### Full-length cDNA sequencing

#### PacBio Reads Mapping to ARL6IP1 mRNA Splice Variants

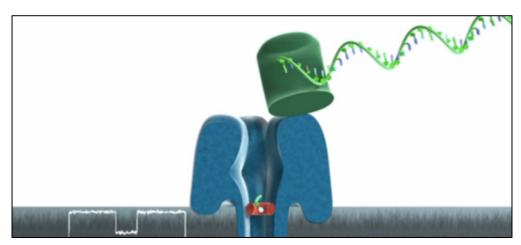


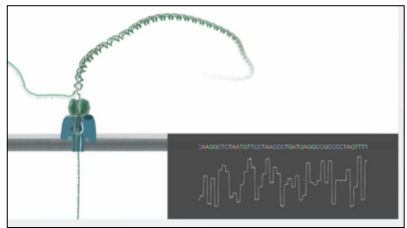
"Post-Light," Semi-Conductor Sequencing





#### **DNA Sequencing**

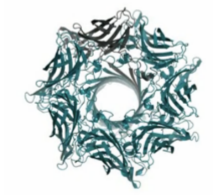




Exonuclease-Seq

Strand-Seq



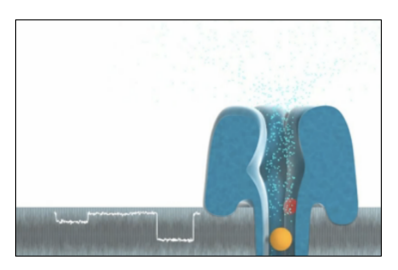




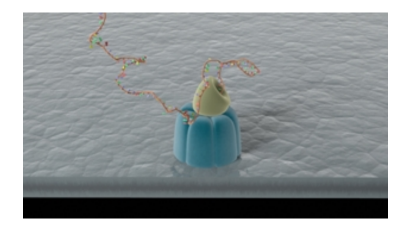
MinION GridION



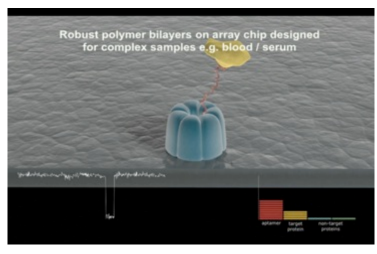
#### Other (Maybe Killer) Apps



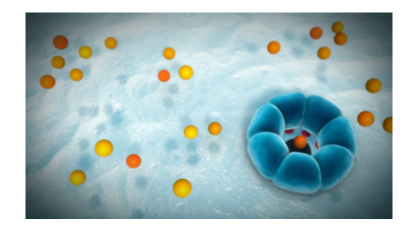
Analyte



**Direct RNA Sequencing** 



**Protein Aptamer** 

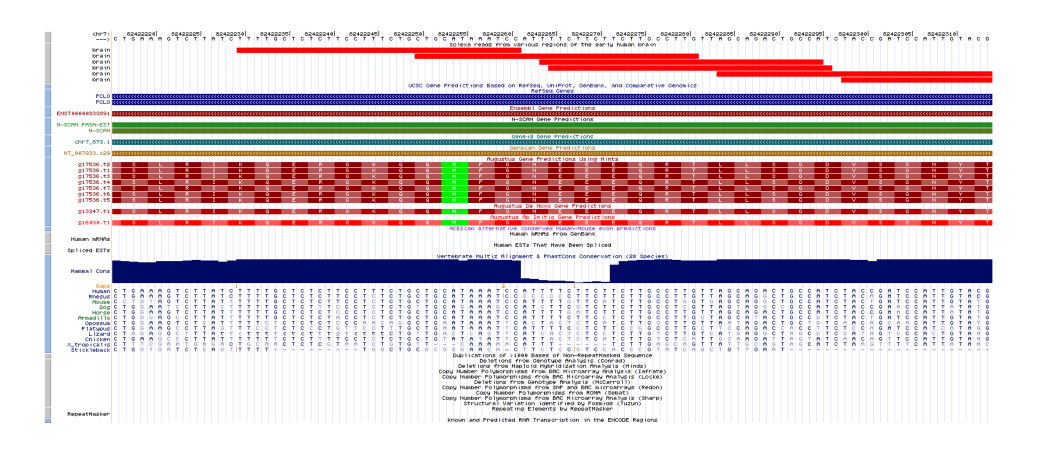


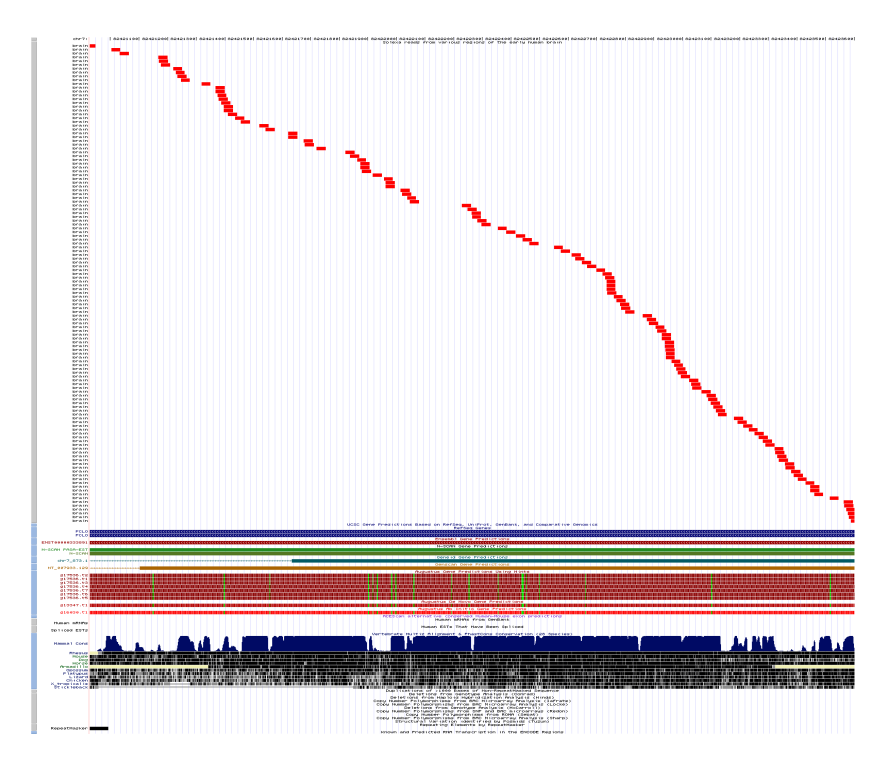
Small molecule

# Each Platform has various sources of noise, and thus Error

- De-Phasing
  - Lagging strand dephasing from incomplete extension
  - Leading strand dephasing from over-extension
- Dark Nucleotides
- Polymerase errors (10<sup>-5</sup> to 10<sup>-7</sup>)
- Platform-specific errors
  - Illumina more likely to have error after 'G'
  - PCR-based methods miss GC- and AT-rich regions

#### Alignment to the genome





## Analyzing High-Resolution Data

- Bayesian Methods
- Hidden Markov Models
- Permutation Testing
- Circular Binary Segmentation
- Seed-seeking
- Least Squares Regression
- Democratic Voting

#### Prior

- The prior function Pr (H) gives the probability of different possible values of the quantity of interest <u>before</u> the data are considered that is, it represents the state of knowledge <u>prior</u> to the data.
- Prior may be broad or flat if we have few data (non-informative prior), or peaked if we have more information (informative prior).

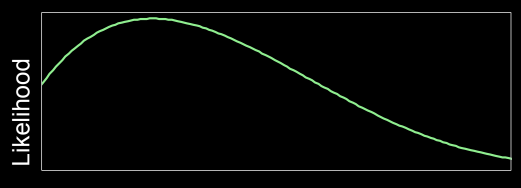


Unknown quantity of interest

(Population size, length at sexual maturity, haplotype frequency, model parameter)

#### Likelihood

- The likelihood function Pr (data | H) gives the probability of obtaining the data, given different possible values of the unknown quantity of interest (the "hypothesis" H).
- The likelihood is calculated using a statistical model, which represents the process that produced the data. The likelihood function connects the parameters of the model to the data.

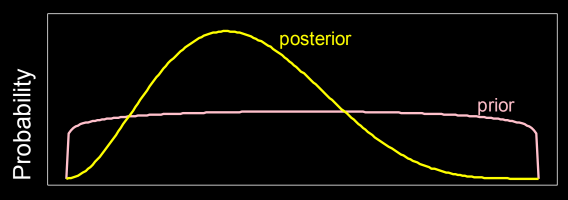


Unknown quantity of interest

(Population size, length at sexual maturity, haplotype frequency, model parameter)

#### **Posterior**

- The posterior function Pr (H | data) gives the probability of different possible values of the quantity of interest <u>after</u> the data are considered that is, it represents the state of knowledge <u>posterior</u> to the data.
- The posterior is a combination of the prior (what we knew before) and the likelihood (what the data told us).
- The difference between the prior and the posterior indicates how much we learned from the data.



Unknown quantity of interest

(Population size, length at sexual maturity, haplotype frequency, model parameter)

#### Paradigm for Bayesian inference

posterior likelihood x prior

new state information x current state of knowledge from new data of knowledge

Thus, in Bayesian reasoning, new data update the current state of knowledge through Bayes' Theorem. The result is a new state of knowledge represented by the posterior.

#### Bayes' Theorem

(H) The Hypothesis (the unknown) and 
$$x = \text{data}$$

$$p \text{ (H | data)} = \frac{p \text{ (data | H)} p \text{ (H)}}{p \text{ (data)}}$$

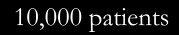
posterior = likelihood x prior

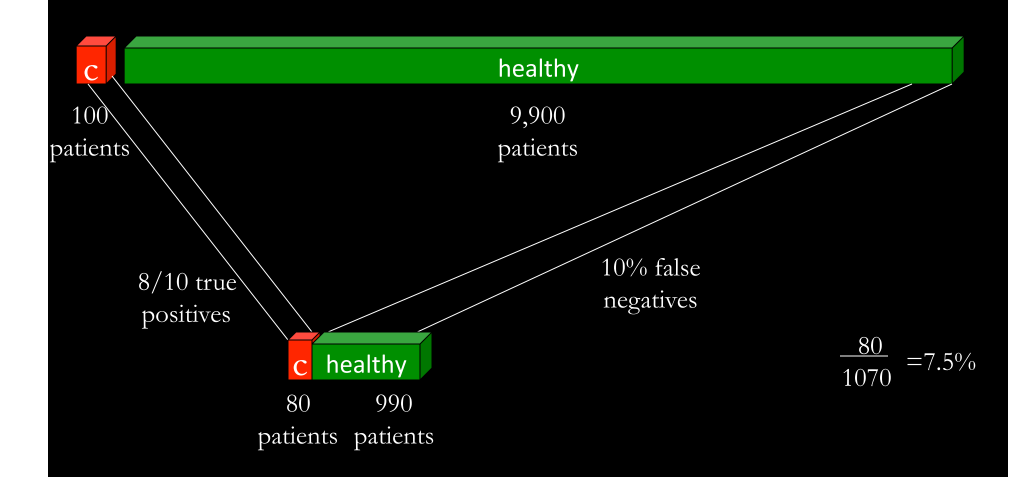
#### 2 fundamental Bayesian concepts:

- 1. Things that are unknown are represented by probability distributions.
- 2. Things that are known (data) are used to improve the knowledge of unknowns through Bayes' Theorem.

## Bayesian view of cancer

- 1% of women at 40 yrs have breast cancer (Prior)
- Mammography diagnoses 8/10 correctly (true positive rate, false negative rate)
- 10% of mammographies are false positives
- If you get a positive result, what are your odds of having breast cancer?





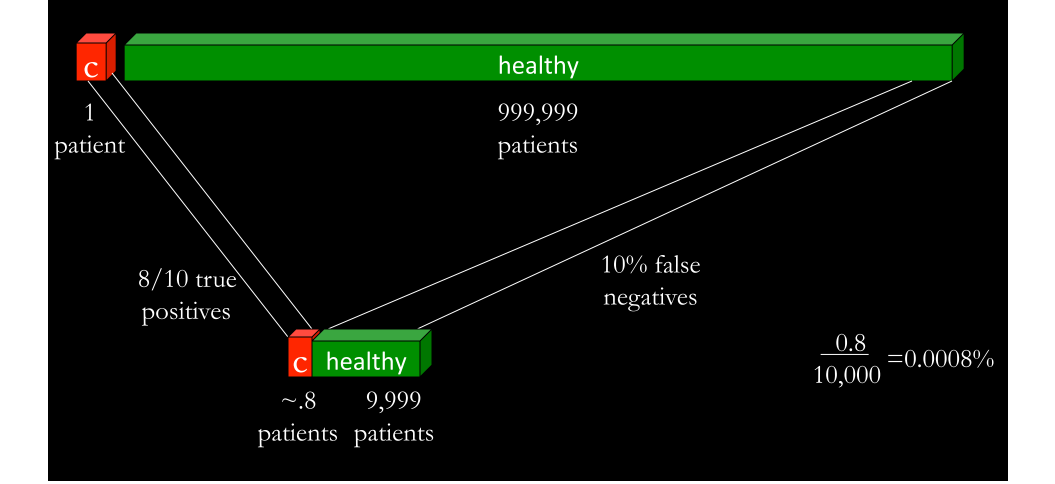
# Bayes Theorem depends on the prior probability (pr(A)):

$$\Pr(A|B) = \frac{\Pr(B|A) \Pr(A)}{\Pr(B)}.$$

A = have cancer B = positive result

$$\frac{.8*.01}{.8*.01 + .1*.99} = 7.5\%$$

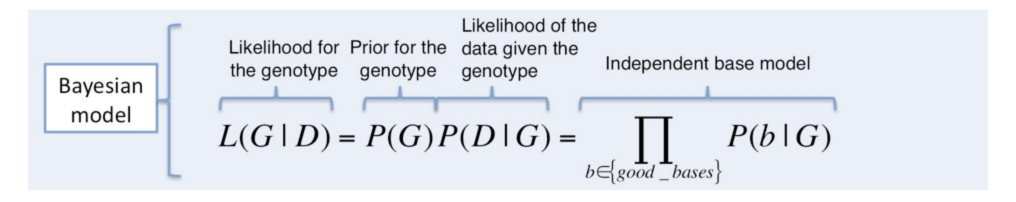




Q. What is the Bayesian Conspiracy?	
that controls publication, grants, tenure, a be accepted into the Bayesian Conspiracy or college, and gradually work your way up	onal, interdisciplinary, and shadowy group of scientists and the illicit traffic in grad students. The best way to is to join the Campus Crusade for Bayes in high school p to the inner circles. It is rumored that at the upper e silent figures known only as the Bayes Council.

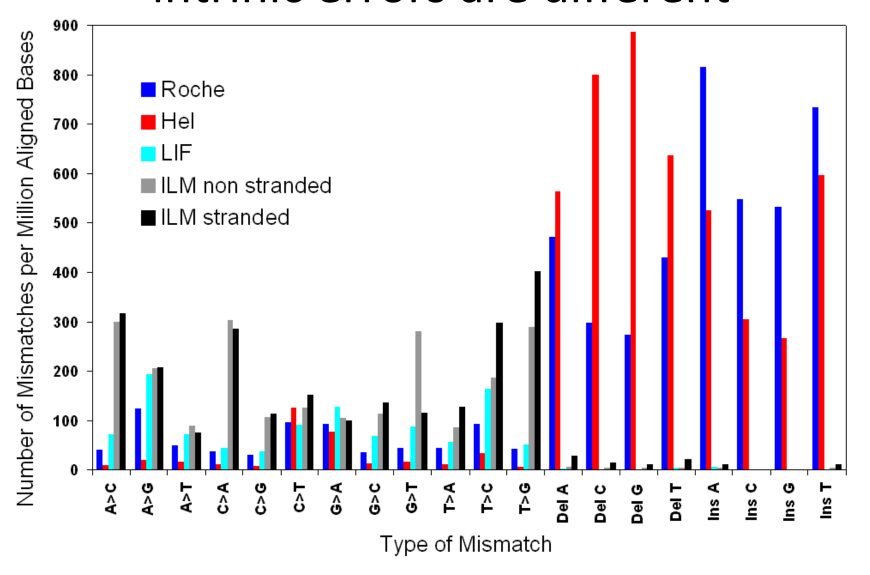
### Applications of Bayes to DNA-Seq

#### GATK single sample genotype likelihoods

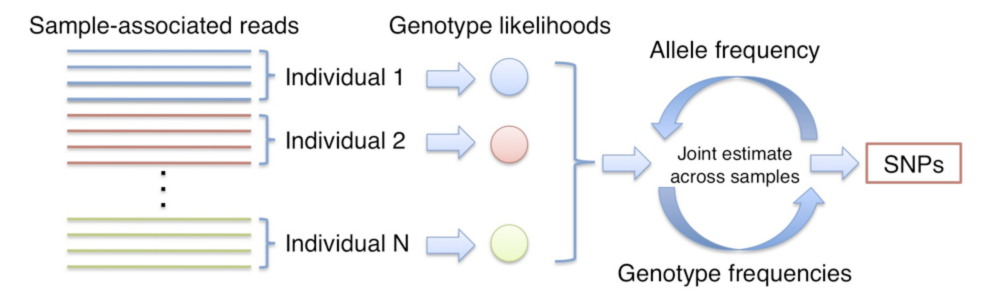


- Priors applied during multi-sample calculation; P(G) = 1
- Likelihood of data computed using pileup of bases and associated quality scores at given locus
- Only "good bases" are included: those satisfying minimum base quality, mapping read quality, pair mapping quality, NQS
- P(b | G) uses a platform-specific confusion matrix
- L(G|D) computed for all 10 genotypes

## Each Platform is slightly different, and so intrinic errors are different



## The Broad Unified Genotyper SNP caller multiplesample allele frequency and genotype estimates



 This approach allows us to combine weak single sample calls to discover variation among samples with high confidence

## SNIP-Seq SNP calling

For each potential variant site in the sequenced region: Set the base quality value for each base call to the Illumina quality value

For 
$$k = 1, 2,...$$

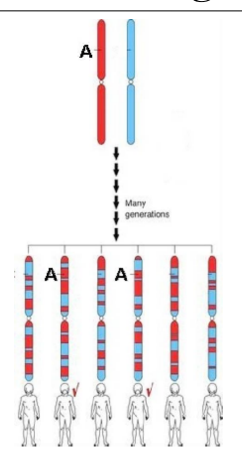
- a. Sample a genotype for each individual from the posterior distribution using a heterozygote prior of 0.001.
- b. Recalibrate the quality score for each base call using genotypes for all individuals.

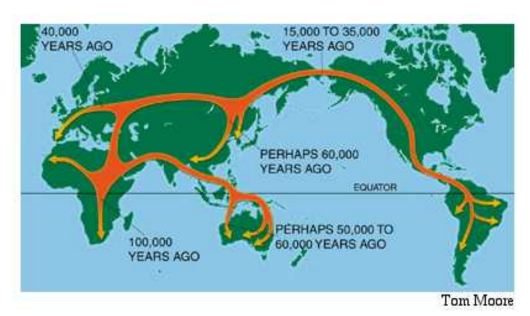
If the genotype of any individual is different from the reference, identify position as a SNP.

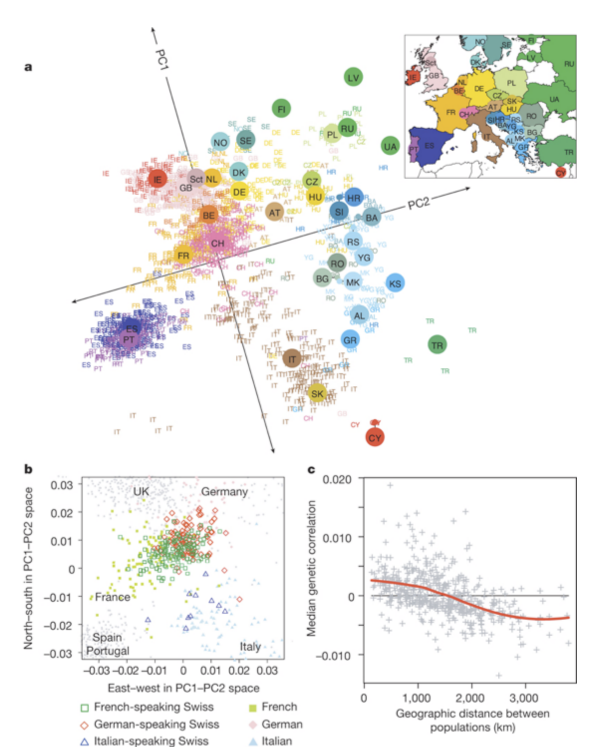
Sample a genotype for each individual from the posterior distribution computed using a heterozygote prior of 0.2.

## Population-based models can overcome systematic errors, but...

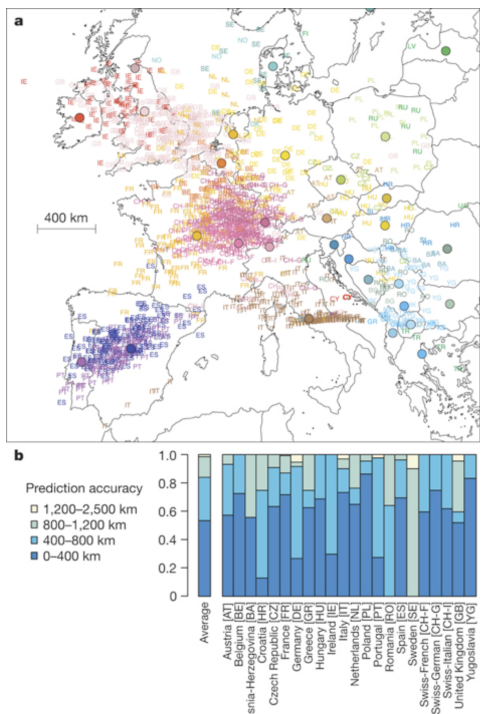
# Population Stratification is from the migration patterns of haplotypes throughout human history







Genes mirror geography within Europe Novembre et al, 2008



Genes mirror geography within Europe Novembre et al, 2008

## Applications of Bayes to RNA-Seq

## Microarrays vs. RNA-Seq





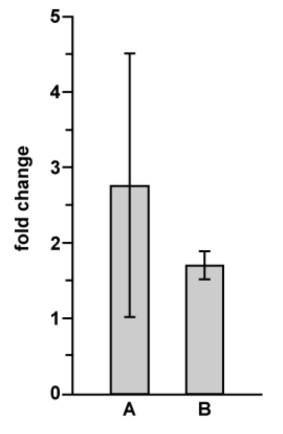


RNA-Seq: An assessment of technical reproducibility and comparison with gene expression arrays

Marioni and Mason et. al, Genome Research, 2008

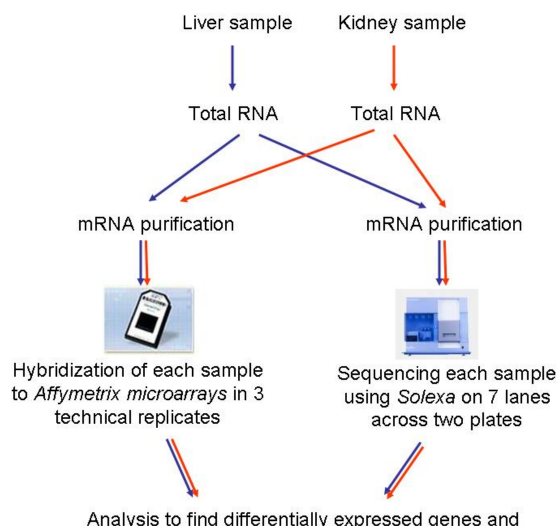
## Data Analysis: What genes are differentially expressed?

- Early days—fold change cutoffs (e.g., 2x difference or better)
- not a very satisfying approach:
  - -doesn't take into account variance
  - -misses any small changes



Here, "A" has a fold change >2.5, but varies greatly between replicate experiments. "B" has a fold change of only 1.75, but changes reliably each time the experiment is performed.

## Experimental Design: Liver vs. Kidney



Analysis to find differentially expressed genes and comparison between technologies

## Metric for RNA-Seq Expression

#### **RPKM:**

Reads per Kilobase per Million Reads

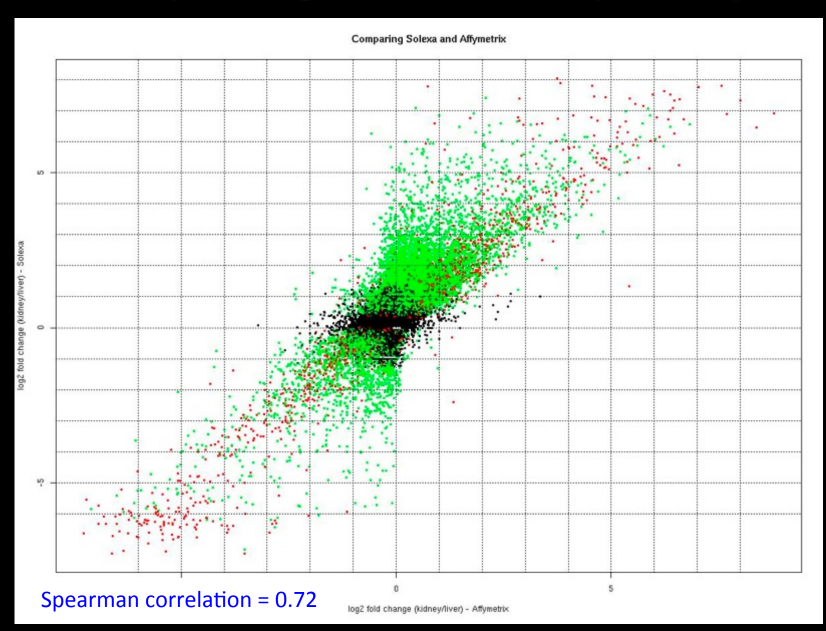
Normalizes for (1) gene size and (2) sequencing depth

(~0.1-1 transcript/cell)

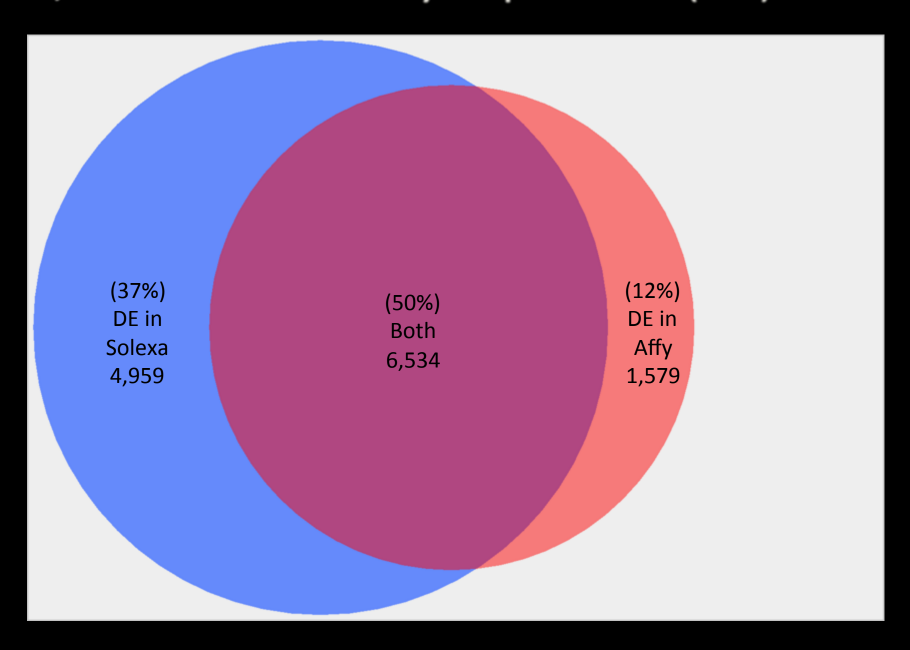
$$\text{RPKM} = \frac{N \ reads}{1 \ gene} \times \frac{1 \ \text{gene}}{B \ bp} \times \frac{1000 \ \text{bp}}{1 Kb} \times \frac{1 \ Million \ reads}{Y \ total \ reads}$$

Y = (exons, introns, intergenic reads)

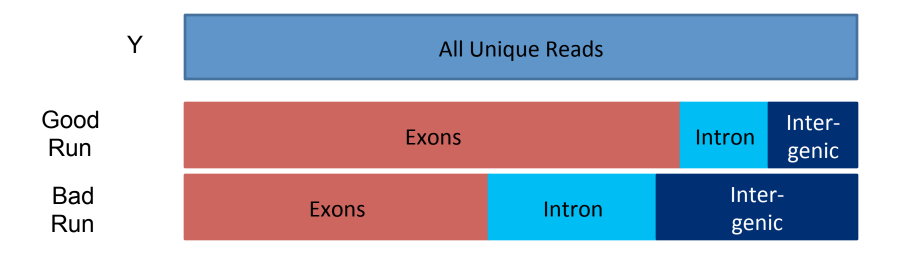
## Comparing GA and Affy arrays



## 13,072 Differentially Expressed (DE) Genes



## Bias is introduced if these ratios are not kept:



## Normalization is needed

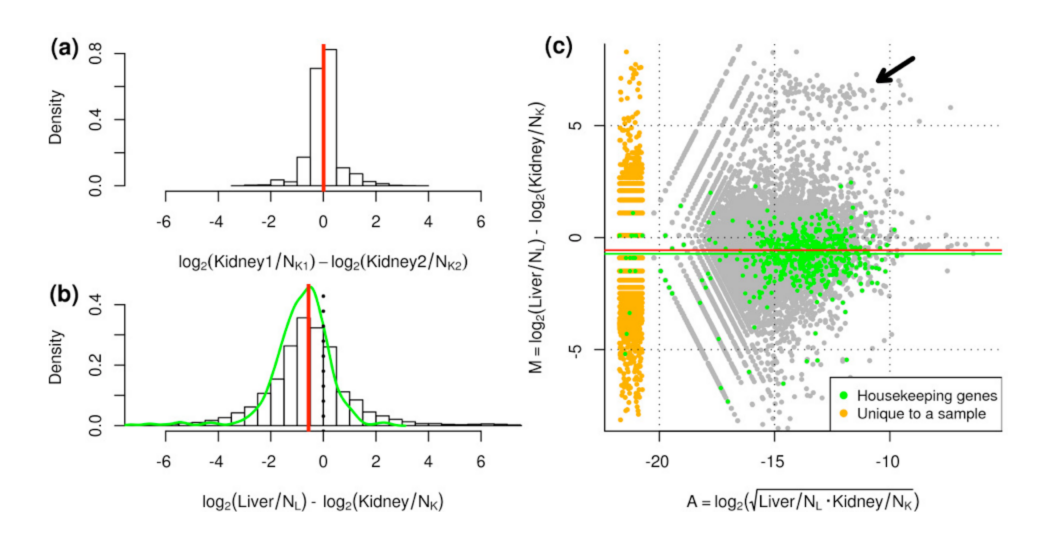
Define  $Y_{gk}$  as the observed count for gene g in library k summarized from the raw reads,  $\mu_{gk}$  as the true and unknown expression level (number of transcripts),  $L_g$  as the length of gene g and  $N_k$  as total number of reads for library k. We can model the expected value of  $Y_{gk}$  as:

$$E[Y_{gk}] = \frac{\mu g k^L g}{S_k} N_k$$
where  $S_k = \sum_{g=1}^G \mu_{gk} L_g$ ;

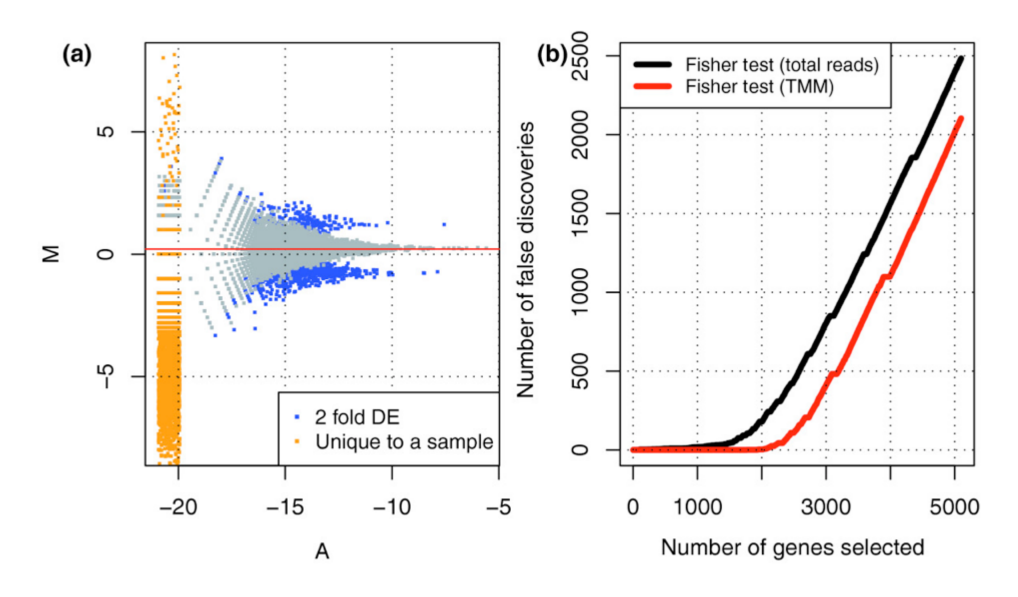
 $S_k$  represents the total RNA output of a sample. The problem underlying the analysis of RNA-seq data is that while  $N_k$  is known,  $S_k$  is unknown and can vary drastically from sample to sample, depending on the RNA composition.

$$M_g = \log_2 \frac{Y_{gk}/N_k}{Y_{gk'}/N_{k'}}$$

#### Normalization is needed



## Normalization is needed





## MISO / Probabilistic analysis and design of RNA-Seq experiments for identifying mRNA isoform regulation

Home | Paper | Software | Documentation | Datasets | Contact

About MISO

#### **About MISO**

MISO (Mixture of Isoforms) is a probabilistic framework that quantitates the expression level of alternatively spliced genes from RNA-Seq data, and identifies differentially regulated isoforms or exons across samples. By modeling the generative process by which reads are produced from isoforms in RNA-Seq, the MISO model uses Bayesian inference to compute the probability that a read originated from a particular isoform.

MISO uses the inferred assignment of reads to isoforms to quantitate the abundances of the underlying set of alternative mRNA isoforms. Confidence intervals over estimates can be obtained, which quantify the reliability of the estimates.

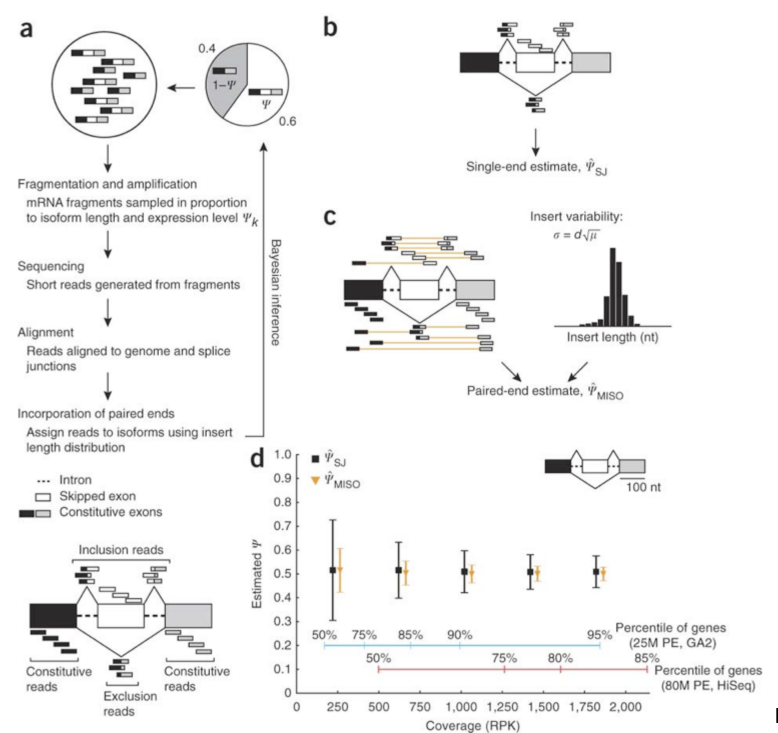






#### Contact

Department of Biology MIT 31 Ames Street, 68-271A Cambridge, MA 02139-4307



Katz et al, 2010

## Coverage Requirements: How many lanes/plates/wells?

### Depends on:

1.Read Length

2. Size of Transcriptome

3. Complexity of Transcriptome

4. Complexity of Tissue

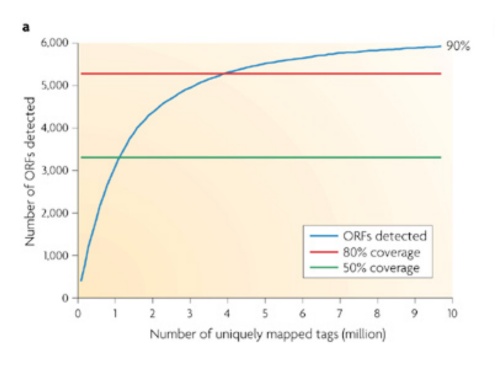
5.Biological Variance

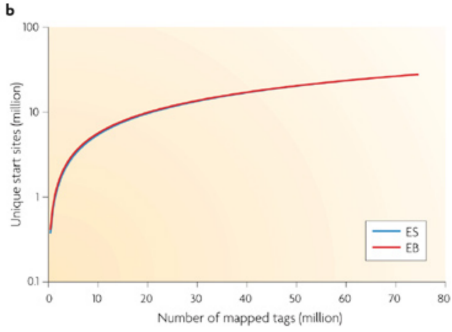
6.Errors (random and systematic)

## Plateau of Information Starts @ ~500Mb

Number of lanes		Differentially expressed	Overlap with genes Correlation of fold o		old changes	
compared		genes	called from the array	between Solexa and the array		
One vs One		5670	4208	0.67		
Two vs Two		7994	5340	0.70		
Three vs Three		9482	5909	0.71		
Four vs Four		10580	6278	0.72		
Five vs Five		11493	6534	0.73		
	I					
				Kidney		
Liv		•		Ridirey		
	No gen	es Percentage		No genes	Percentage	
Five Lanes	2008	0 100	Five Lanes	20921	100	
Four Lanes	1969	5 97.9	Four Lanes	20552	98.2	
Three Lanes	1917	0 95.5	Three Lanes	20064	96.0	
Two Lanes	1839	0 91.6	Two Lanes	19355	92.5	
One Lane	1697	3 84.5	One Lane	18080	86.4	

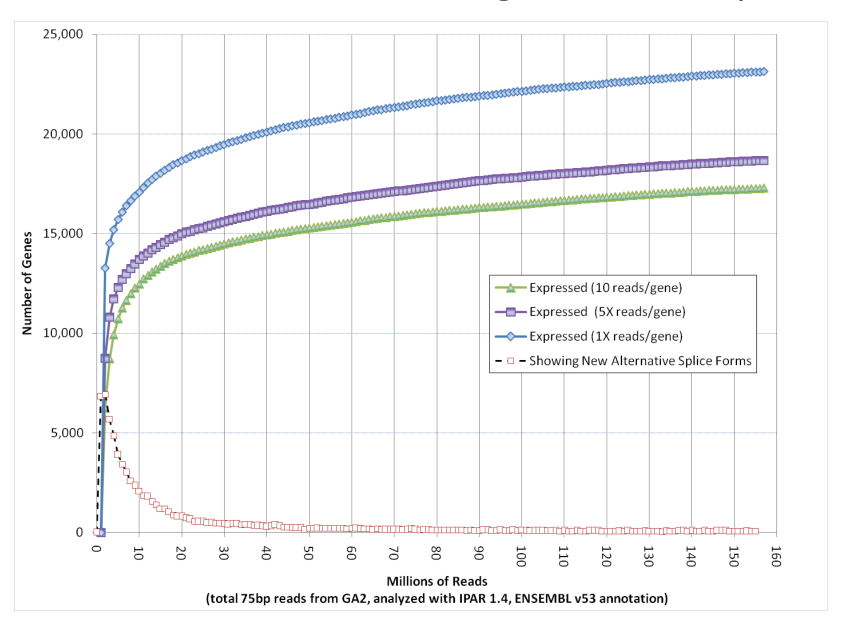
## Coverage Requirements





Nature Reviews | Genetics

### No current visible end of gene discovery



### How many replicates do I need?

Calculation of the number of replicates depends on:

- 1. An estimate of  $\sigma^2$  obtained from previous experiments.
  - 2. The size of the difference ( $\delta$ ) to be detected.
- 3. The assurance with which it is desired to detect the difference (i.e., Power of the test =  $1-\beta$ ).
- 4. The level of significance to be used in the actual experiment (i.e., Type I error).

  5. The test required, whether a one-tail or two-tail test.

To determine the number of replicates use the following formula:

$$\#reps = 2\left(Z_{\alpha/2} + Z_{\beta}\right)\left(\frac{\sigma}{\delta}\right)^{2}$$

where:  $Z\alpha/2$  is associated with the Type I error (two-tailed) Z $\beta$  is associated with the Type II error  $\delta$  is the true difference to be detected, and  $\sigma$  is the known variance obtained from previous experiments

## Bayes in Chip-seq too!

Research article

Highly accessed Open Access

#### BayesPeak: Bayesian analysis of ChIP-seq data

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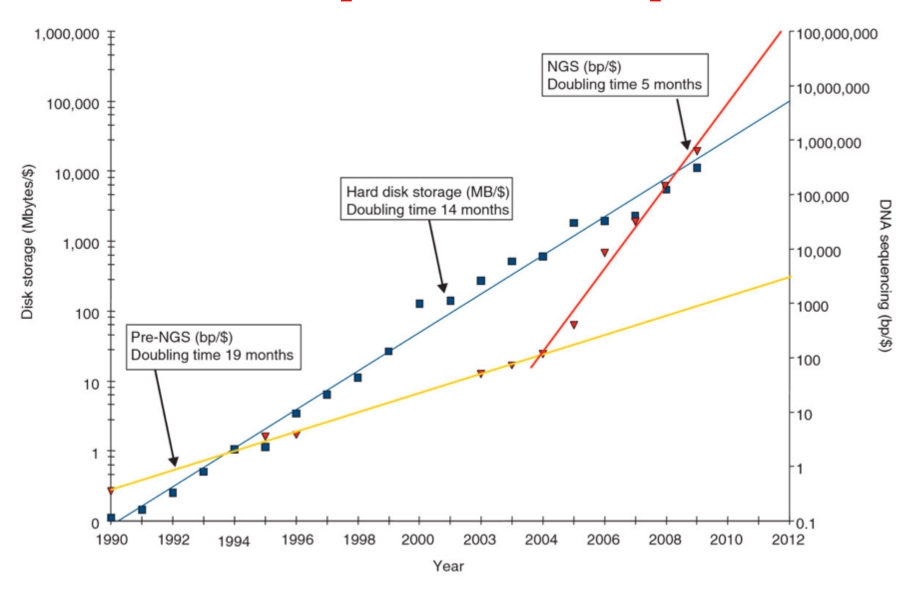
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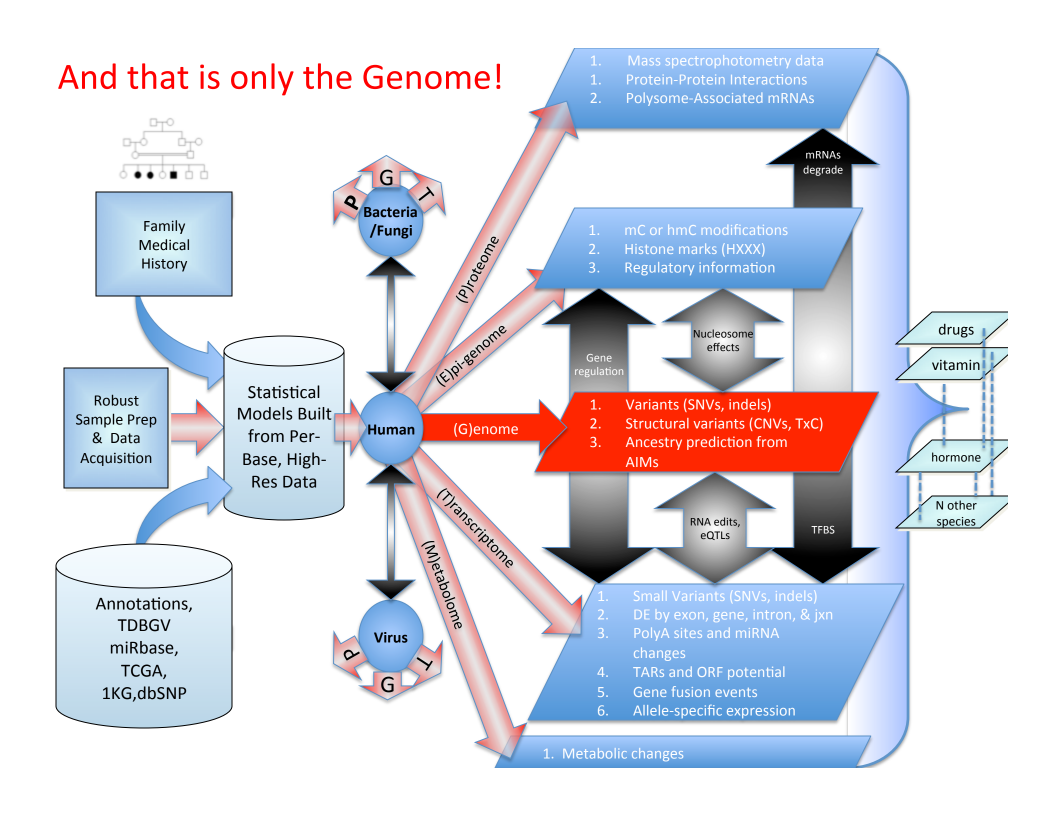
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## What's the problem with Bayesian statistics? (according to non-Bayesians)

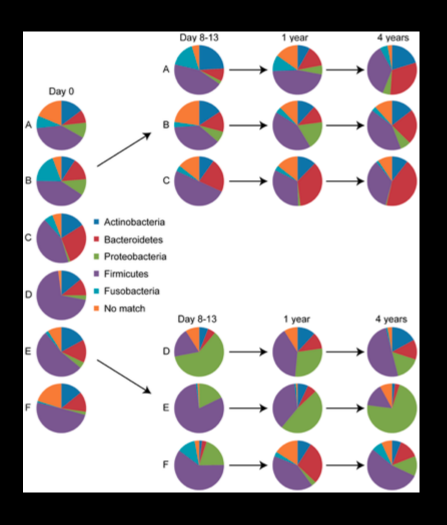
- Priors can introduce subjective judgment into data analysis.
- 2. Priors affect the result. Different people can get different answers from the same data.
- It's too hard. There are no simple point-andclick programs.

## This requires a lot of space





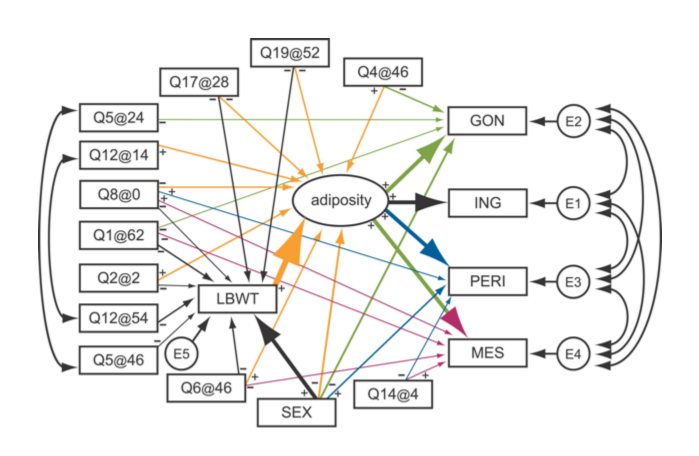
Meta-genomic phenotypes can persist for years, and "passenger genomes" can be a phenotype, as well as their distributions.



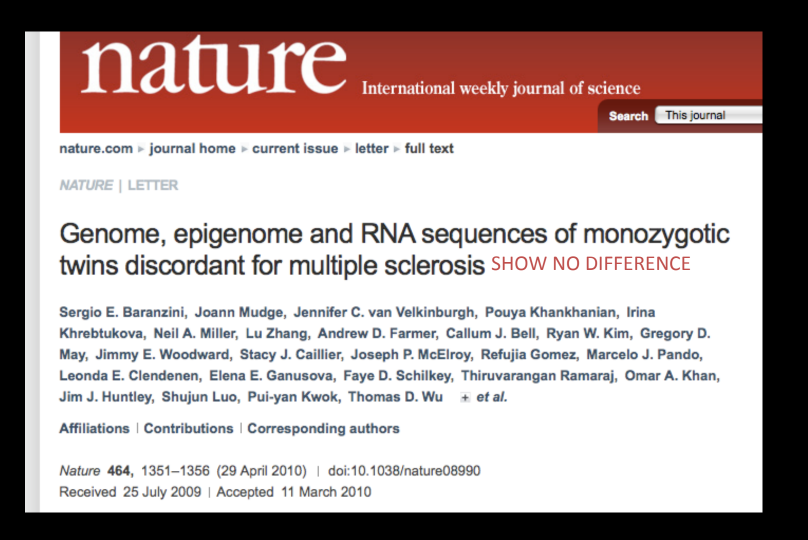
Normal throat

Throat + antibiotics

## Risk factors for diseases usually involve many genes and pathways



## There are other factors than these!



Systems biology requires spatiotemporal monitoring of the genome, epigenome, transcriptome, proteome, metabolome, and the environment, to see the interactome