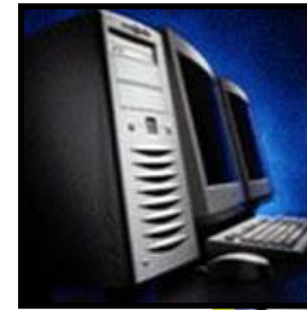
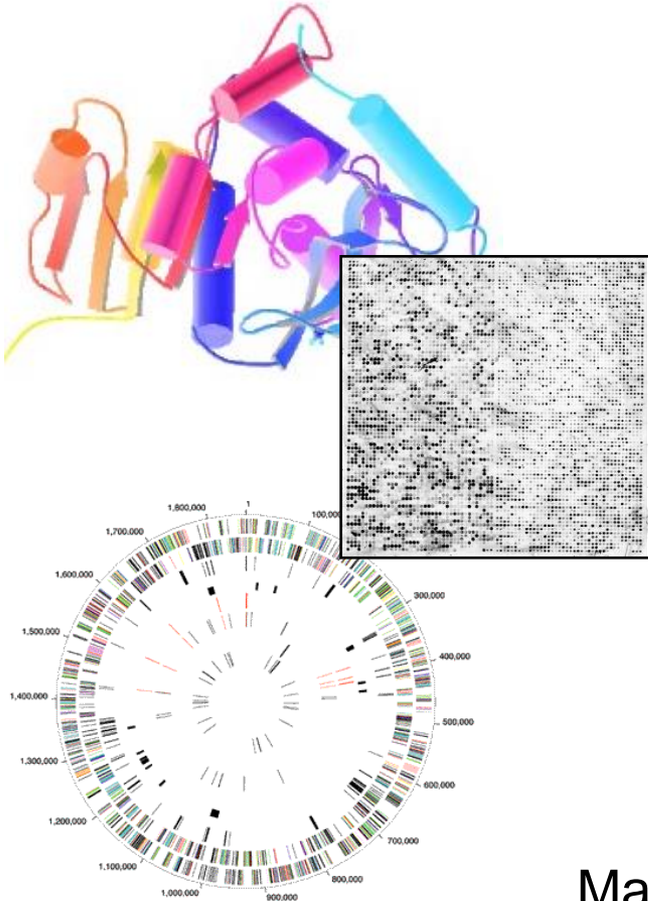


Biomed. Data Sci. Personal Genomes Intro.

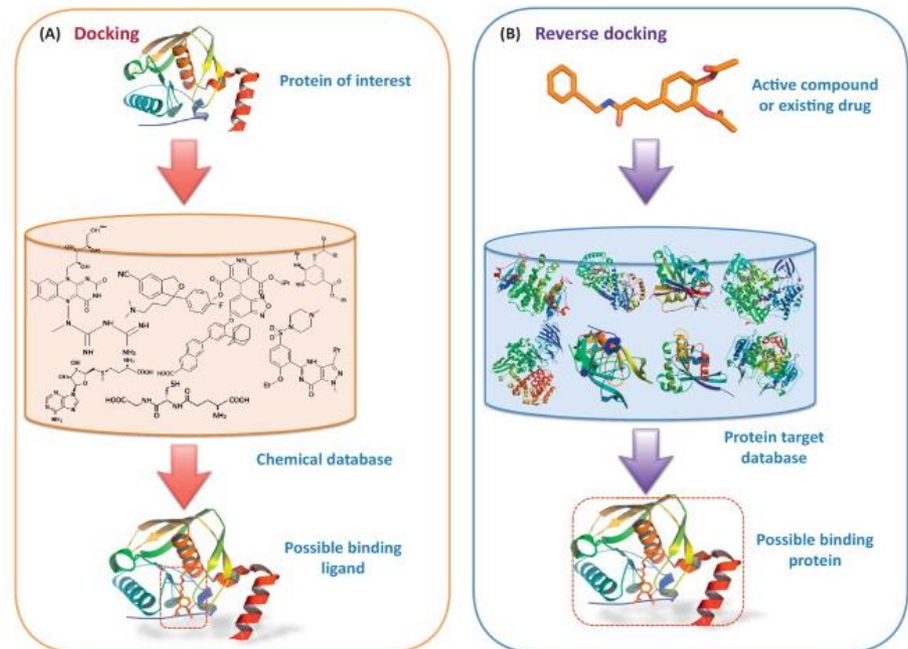
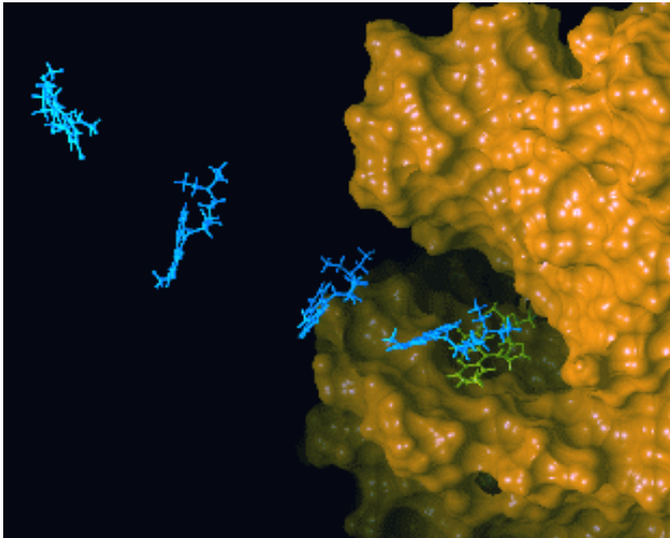


Mark Gerstein, Yale University
GersteinLab.org/courses/452

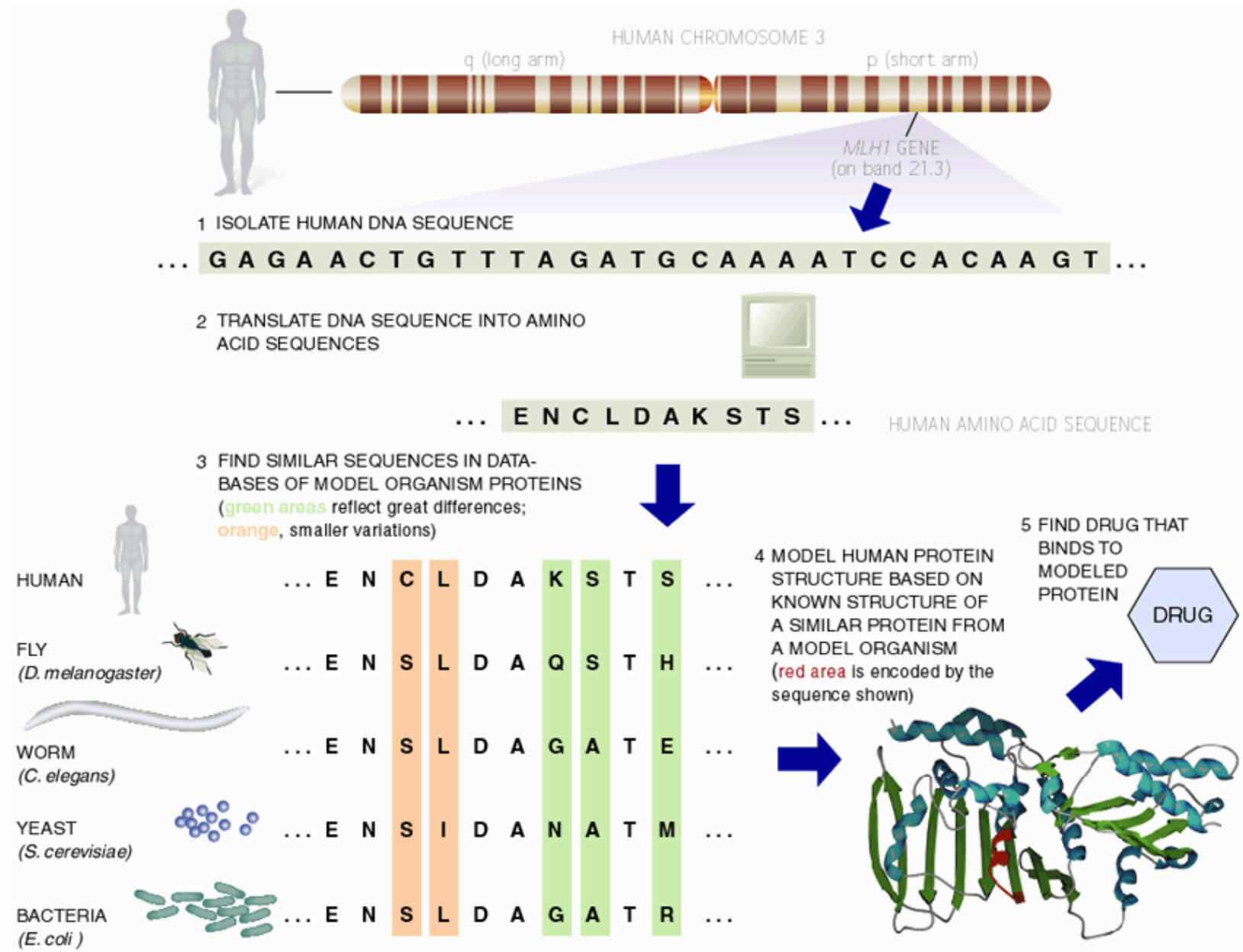
(Last edit in spring '22, this is 22i2a which has a slight edit on slide 5 relative to l2a)

Major Application I: Designing Drugs from Structural Targets

- Understanding how structures bind other molecules
- Designing inhibitors using docking, structure modeling
- *In silico* screens of chemical and protein databases

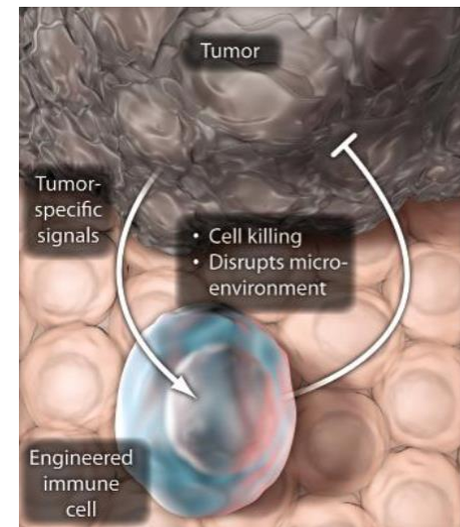
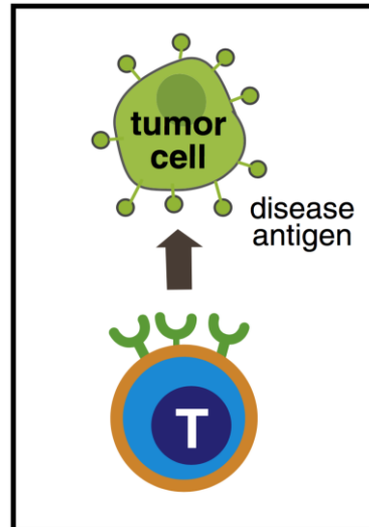
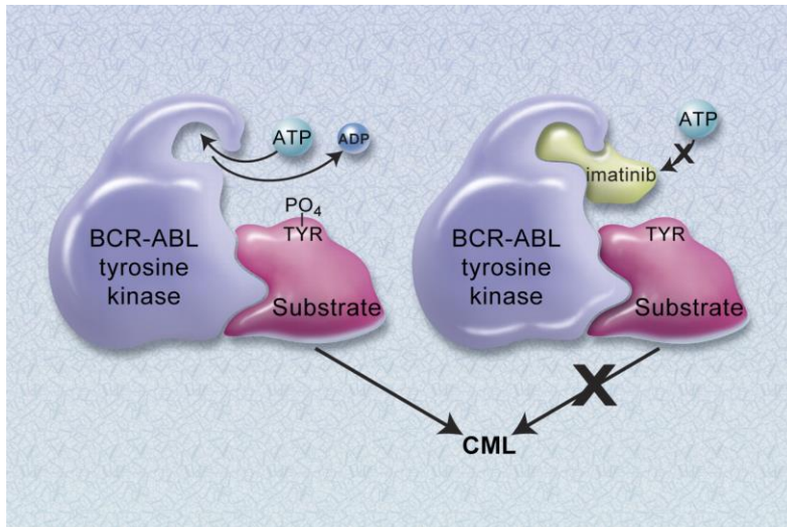


Major Application II: Finding Homologs, to Find Experimentally Tractable Gene Targets



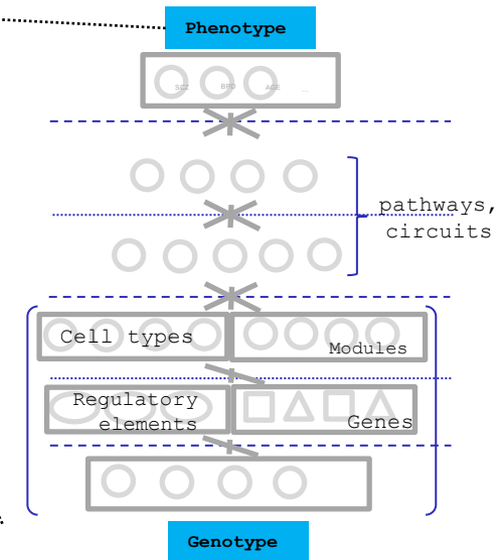
Major Application III: Customizing treatment in oncology

- Identifying disease causing mutations in individual patients
- Designing targeted therapeutics
 - e.g. BCR-abl and Gleevec
 - Cancer immunotherapies targeting neoantigens



Major Application IV: Finding molecular mechanisms & drug targets for diseases we know little about (Neuro-psychiatric Diseases)

Disease	Heritability*	Molecular Mechanisms
Schizophrenia	81%	-
Bipolar disorder	70%	-
Alzheimer's disease	58 - 79%	Apolipoprotein E (APOE), Tau
Hypertension	30%	Renin–angiotensin–aldosterone
Heart disease	34-53%	Atherosclerosis, VCAM-1
Stroke	32%	Reactive oxygen species (ROS), Ischemia
Type-2 diabetes	26%	Insulin resistance
Breast Cancer	25-56%	BRCA, PTEN



Many psychiatric conditions are highly heritable

Schizophrenia: up to 80%

But we don't understand basic molecular mechanisms underpinning this association
(in contrast to many other diseases such as cancer & heart disease)

Moreover, current models substantially underestimate heritability using genetic data

Schizophrenia : ~25%

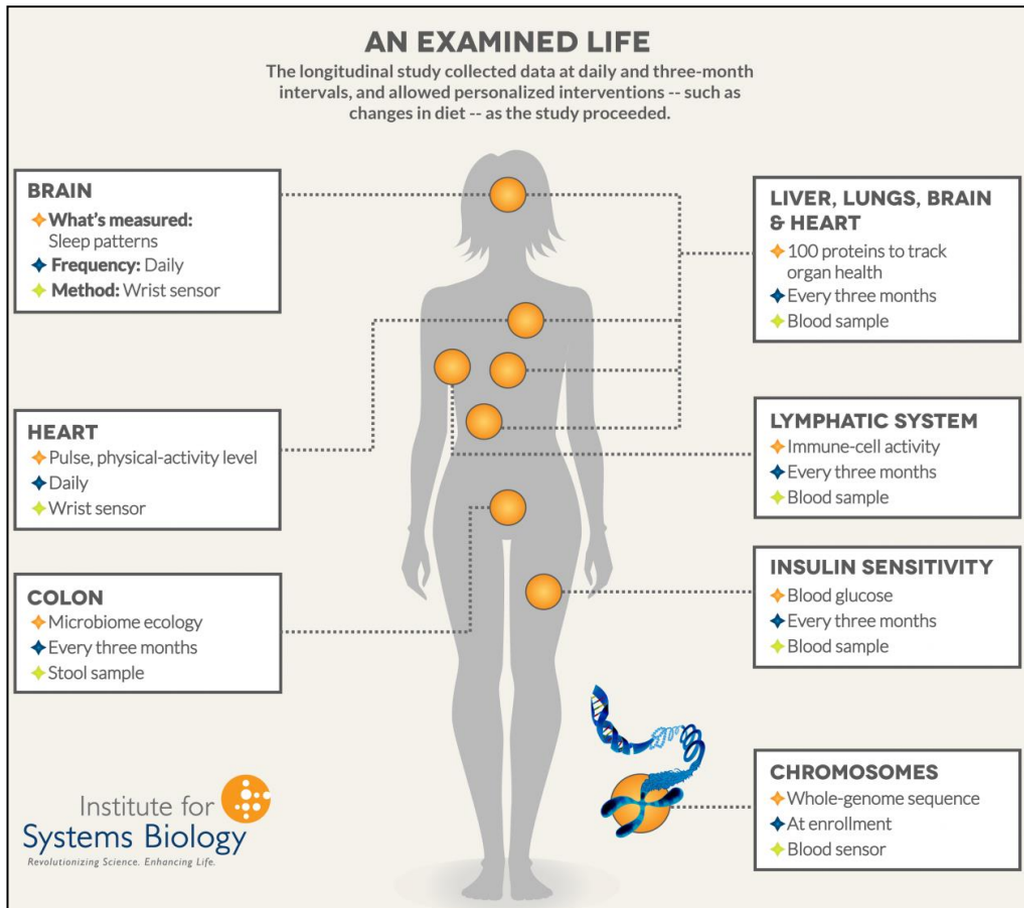
Thus, interested in developing predictive models of psychiatric traits which:

Use observations at intermediate (molecular levels) levels to inform latent structure.

Use the predictive features of these “molecular endo phenotypes” to begin to suggest actors involved in mechanism

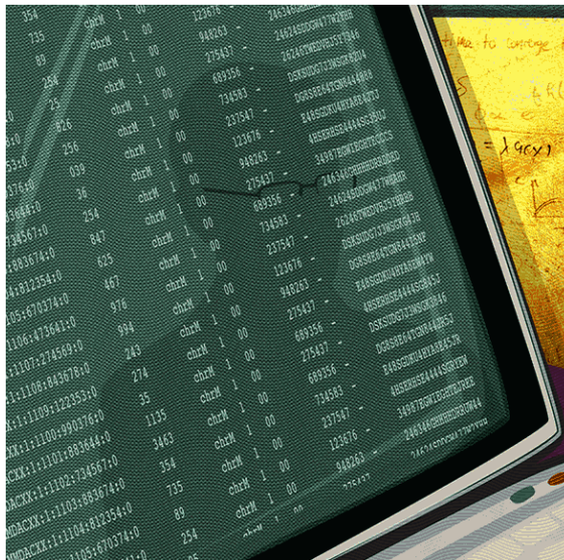
Major Application V: Holistic Personal Genome Characterization, in Normal Individuals

- Mental disease & cancer are two extremes with respect to genomics (CEN, 92: 26)
 - Many other conditions in between, often involving interaction with the environment
- Pers. Genome Characterization
 - Identify mutations in personal genomes (SNPs, SVs, &c)
 - Estimate phenotypic (deleterious or protective) impact of variants.
 - Compare one person to wider population.
- Track changes over time & consider interaction w/ environment
 - Transcriptome studies
 - Longitudinal health studies (e.g. 100K wellness project, Framingham Heart Study)



Analyzing Carl Zimmer's genome

CARL ZIMMER'S GAME OF GENOMES SEASON 1



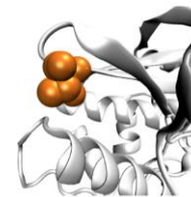
SNV

AAGCT → ACGCT

Protein Structure



Wild-type



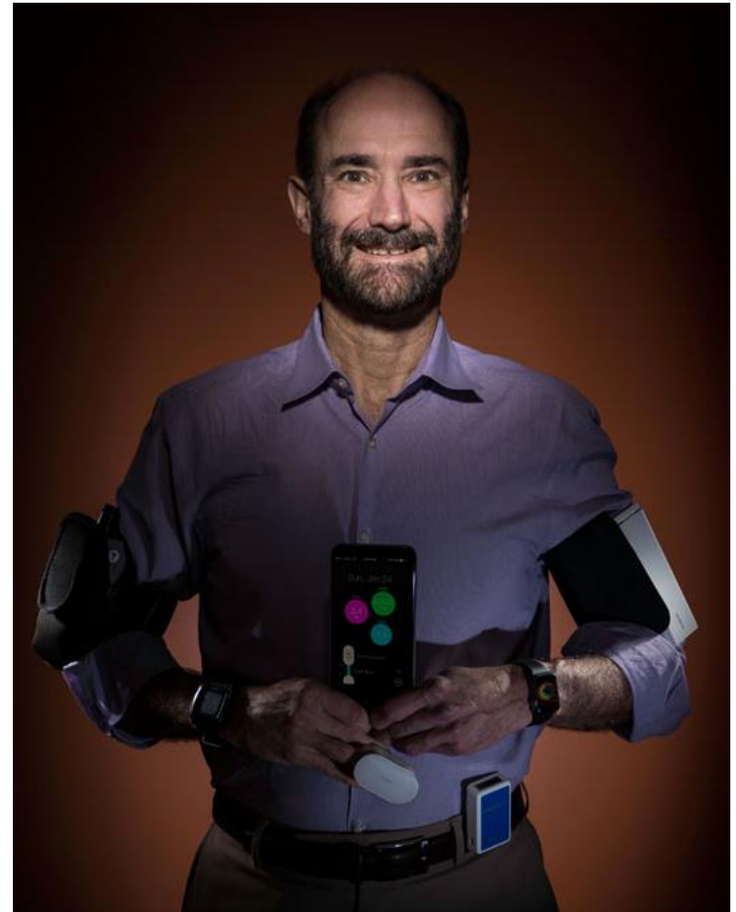
Mutated

Ancestry



Expanding personalized medicine beyond the genome.

- An integrated personal omics profile (iPOP) is an example of a more comprehensive version of personalized medicine.
- Michael Snyder had his genome sequenced and collected many other large scale datasets over an extended period of time.



Integrated personal omics profile (iPOP)

- Numerous types of data were collected, primarily from blood samples. The datasets include:
 - Transcriptomic
 - Proteomic
 - Metabolomic
 - Cytokine profiling
 - Autoantibody profiling
 - Medical exams

