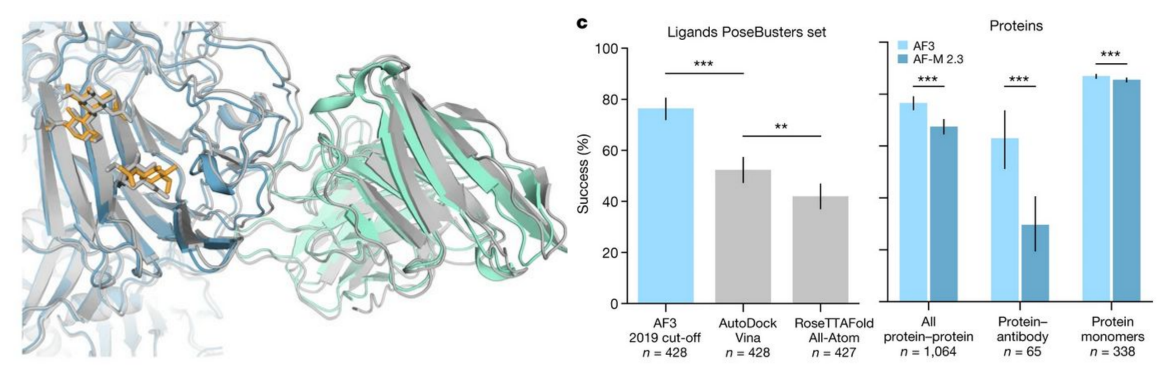
### **AMIDD 2024 Lecture 5: Protein-Ligand Interaction**



Left: human coronavirus spike protein bound to neutralization antibody. Right: Performance of AlphaFold3 for protein-ligand interaction prediction, and for protein-protein interaction. AF-M: AlphaFold Multimer. Adapted from Abramson, ..., Hassabis, Jumper, <u>Accurate structure prediction of biomolecular interactions with AlphaFold 3</u>, Nature (2024)

#### Dr. Jitao David Zhang, Computational Biologist

<sup>1</sup> Pharmaceutical Sciences, Pharma Research and Early Development, Roche Innovation Center Basel, F. Hoffmann-La Roche <sup>2</sup> Department of Mathematics and Informatics, University of Basel

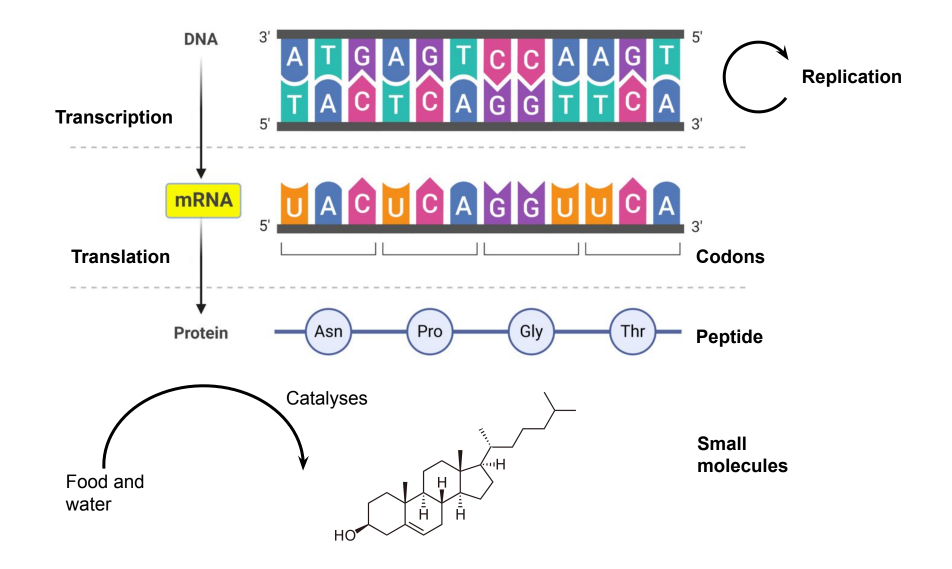


#### **Topics of lecture 5**

- Protein, ligand, and protein-ligand interaction
- ODE-based mechanistic models

#### The central dogma





### Amino acids, the building blocks of proteins, form peptide bonds

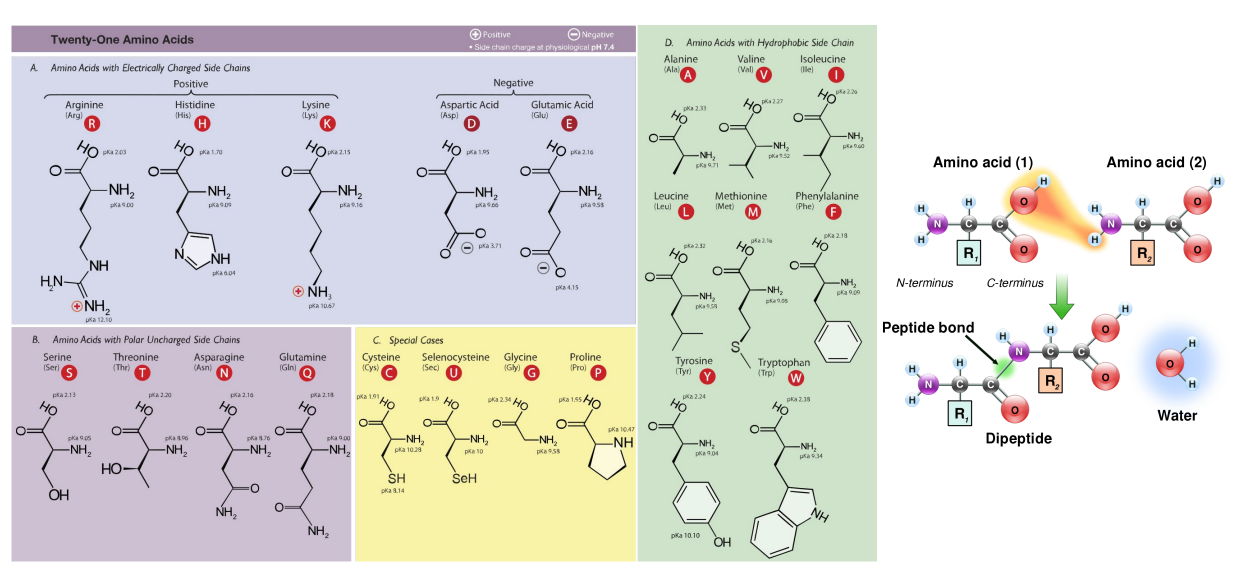


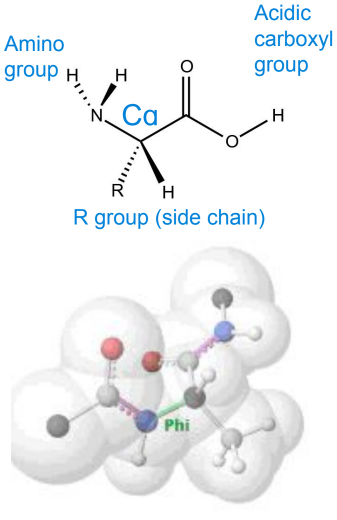
Figure by Dan Cojocari. Reused with CC license from wikimedia

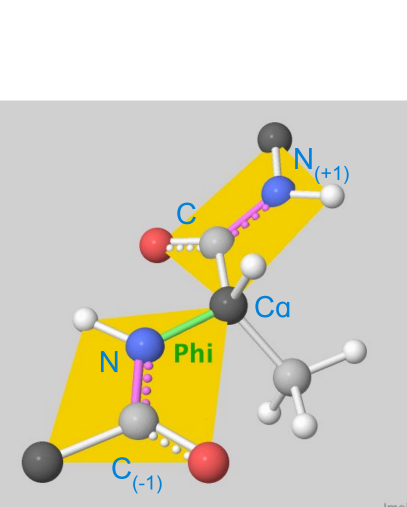
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### **Primary structure of proteins**

- (Top left) Human proteins are chains of amino acids (AAs). The backbone remains the same while the side chain varies among AAs.
- (Right) The amino group and the carboxyl group of adjacent amino acids form peptide bonds. Proteins are therefore called *polypeptides*.
- C-Ca bonds Ca-N bonds can rotate at two *dihedral angles*, Ψ (psi) and φ (phi), respectively.
- (Bottom left) Due to steric collisions, not all combinations of Phi/Psi are possible.





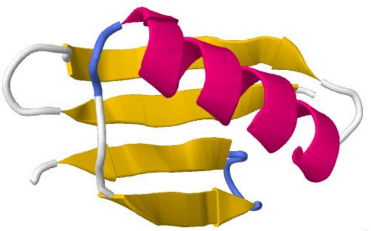


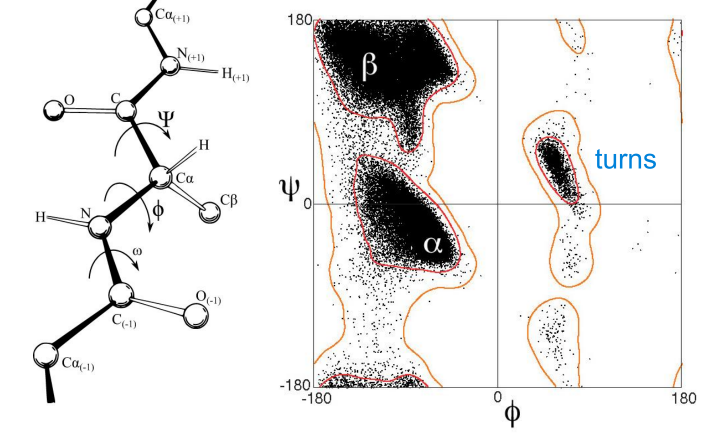
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## The Ramachandran Principle: alpha helices, beta strands, and turns are the most likely confirmations for a polypeptide

Most other conformations are impossible to due to clashes, known as *steric collisions*, between atoms.

To learn more about the topic, check out the <u>YouTube video tutorial</u> or the <u>Slides</u> by Eric Martz, and finish the <u>Quiz</u>.



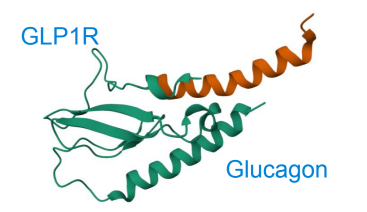


100,000 dots taken from high-resolution crystallographic structures. <u>Wikimedia Commons</u> courtesy Jane and David Richardson (<u>Proteins 50:437, 2003</u>). This plot excludes Gly, Pro, and pre-Pro.

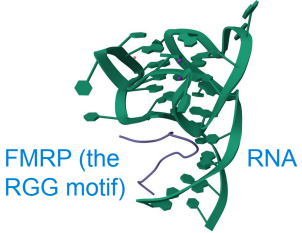
Jmol

### Proteins specifically and tightly bind to other molecules



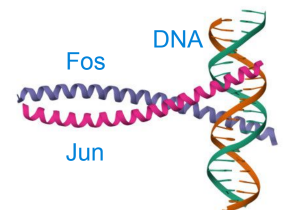


One protein binds to another protein PDB 3iol



Protein binds to RNA.

Protein FMRP is encoded by gene *FMR1*. Mutations associated with *FMR1* induce the fragile X syndrome.<u>PDB 5DE5</u>



#### **Protein complex binds to DNA.** The complex Fos:Jun is known as AP-1, a transcription factor. <u>PDB 1FOS</u>.

CYP3A4 3 Caffeine molecules

Protein binds to small molecule. Cytochrome P450 3A4 (CYP3A4) is a major drug metabolizing enzyme, which also metabolizes caffeine. PDB 8so1

### Major protein classes by functions

**Enzymes**: catalysis of chemical reactions.

To learn the basics of enzymes, watch the video How Enzymes Work.

**Transporters:** moving ions, small molecules, and proteins across membranes.

To learn the basics of transporters and other • ways cell transport material across membranes, Watch the video *Biology: Cell Transport*.

**Receptors and kinases:** signalling allows cells adapt to the environment.

To learn the basics of cellular signaling, watch • the video <u>Common cell signaling pathway</u>.

Structural proteins: stiffness, rigidity, and mechanistical forces.

#### Top: an antigen presenting cell. Bottom: a T cell. The red dot: a virus

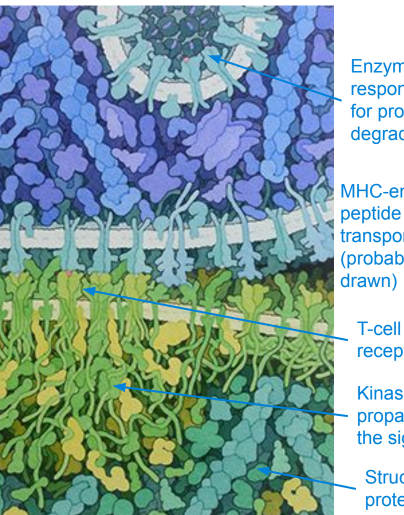


Figure: Immunological Synapse, David S. Goodsell, 2020

Enzymes responsible for protein degradation

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MHC-encoded transporter (probably not

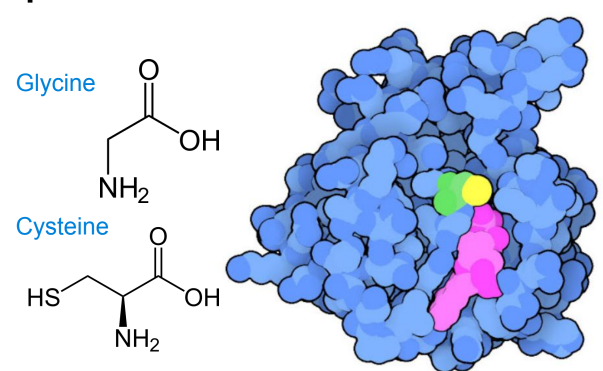
receptor

Kinases that propagate the signal

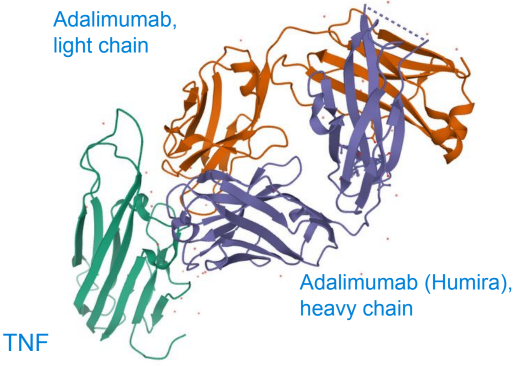
> Structural proteins



### Some diseases are caused by perturbed functions of single protein



Mutation of glycine (G) to cysteine (M) at position 12 (green, with sulfur in yellow) in Ras protein leads to a protein that is continually activated. The structure of the oncogenic mutant (PDB ID <u>4ldj</u>) reveals that the mutation modifies the interaction with GDP (magenta) and GTP, which act as the switch that turns the protein on and off.

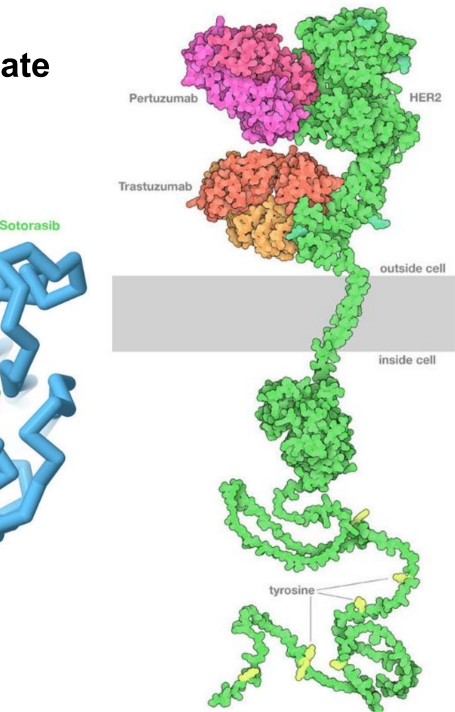


Tumor necrosis factor (TNFa) promotes the inflammatory response in autoimmune diseases, for instance inflammatory bowel disease and rheumatoid arthritis. Monoclonal antibodies against TNFa, for instance adalimumab (Humira), are used for such indications.<u>PDB 3WD5</u>

### Many drugs are ligands of proteins and modulate protein's function

Left: The drug *sotorasib* binds covalently to the sulfur atom in cysteine 12 of the *Ras* protein, blocking its action. The drug is shown with carbon atoms in green, the cysteine sulfur is in yellow, and GDP is in magenta. Image created in Jmol using PDB ID <u>60im</u>.

Right: The extracellular domain of HER2 bound to two therapeutic antibodies: *pertuzumab* and *trastuzumab*. The antibodies block the formation of active dimers of the receptor, thus blocking the growth signal (PDB 6ogi). The transmembrane domain is from PDB 2ksi. The kinase domain inside the cell is from PDB ID 3pp0, and the unstructured tail at bottom is from *AlphaFold2*.







Ill. Niklas Elmehed © Nobel Prize Outreach John J. Hopfield

Ill. Niklas Elmehed © Nobel Prize Outreach Geoffrey E. Hinton



Ill. Niklas Elmehed © Nobel Prize Outreach David Baker



Ill. Niklas Elmehed © Nobel Prize Outreach **Demis Hassabis** 

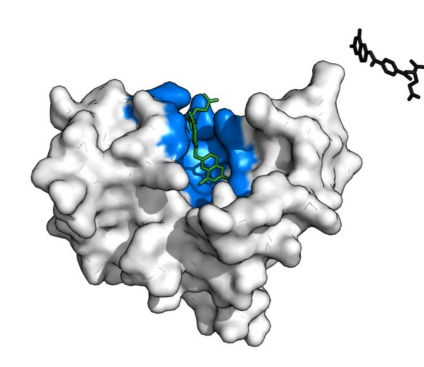


Ill. Niklas Elmehed © Nobel Prize Outreach John M. Jumper

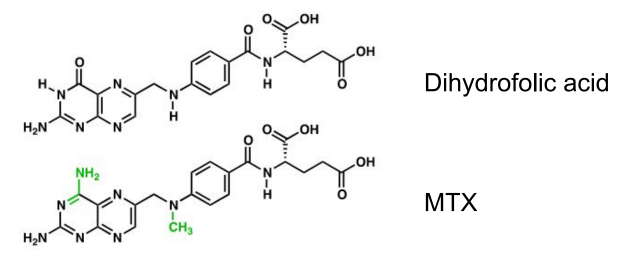
The Nobel Prize in Physics 2024: "for foundational discoveries and inventions that enable machine learning with artificial neural networks"

The Nobel Prize in Chemistry 2024 was divided, one half awarded to David Baker "for computational protein design", the other half jointly to Demis Hassabis and John M. Jumper "for protein structure prediction"

### Drugs can compete with natural ligands



Work by Thomas Shafee, Shared under CC-AS-4.0, and work by Boghog. Based on PDB record 4QI9.



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**The protein**: Dihydrofolate reductase (DHFR) converts dihydrofolic acid into tetrahydrofolate. The process is important for cell proliferation and cell growth. DHFR is a drug target for oncology (cancer) and autoimmune diseases.

**The natural substrate**: Dihydrofolic acid (vitamin B9), in black. Dihydrofolic acid is the *natural ligand* of DHFR.

**The drug:** Methotrexate (MTX), in green, is a *synthesized ligand* of DHFR, and it is a *competitive inhibitor* of DHFR.

**The binding site**: where the enzyme binds its substrate and catalyses the chemical reaction, in <u>blue</u>.



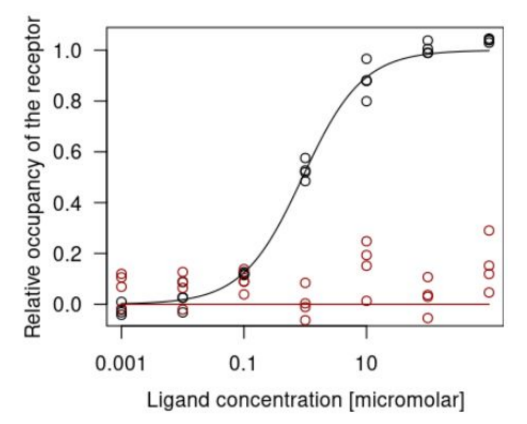
### Concentration-occupancy curves characterize protein-ligand binding

**X-axis**: ligand concentration. Common units: molar (M), micromolar (mM, 10<sup>-6</sup> M), nanomolar (nM, 10<sup>-9</sup> M), picomolar (pM, 10<sup>-12</sup> M).

**Y-axis:** relative occupancy of the receptor. Alternative values are possible, for instance response (more about that later).

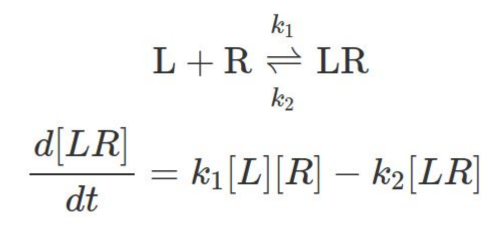
**Points**: individual measurements. In this plot: mean value of replicates with error bars indicating variability.

**Lines**: fitted sigmoidal curves using the Hill function or its variants.

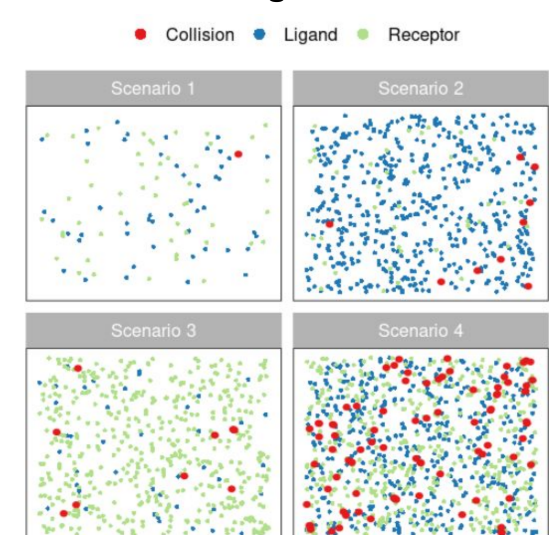




## A simple mathematical model addresses a key question: how is a receptor occupied by varying concentrations of drugs?

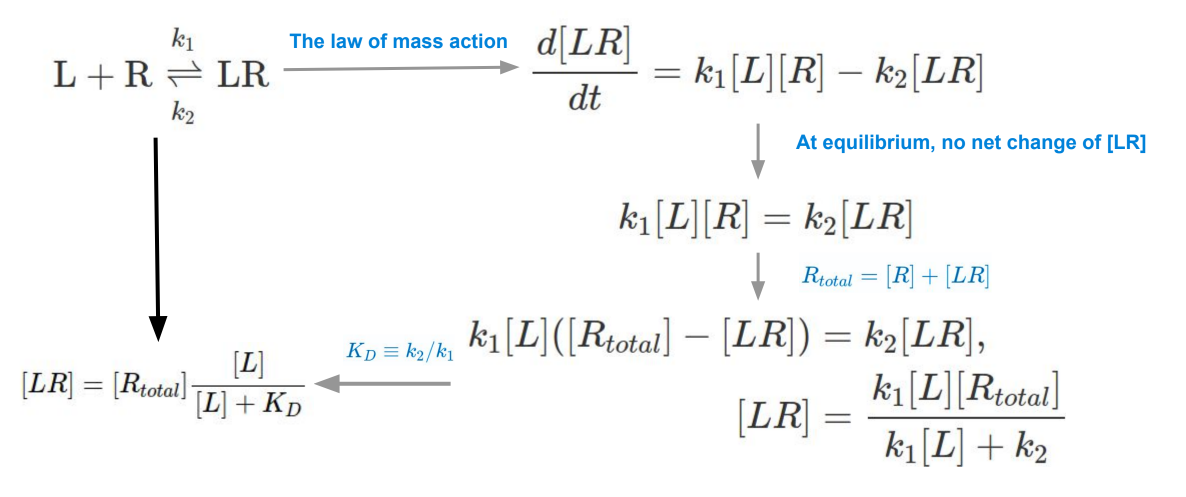


- Ligand binding to receptor is a reversible reaction.
- The law of mass action: the rate of the chemical reaction is directly proportional to the product of the activities or concentrations of the reactants. The proposition can be derived from the *collision theory*. See the right graph for an illustration.





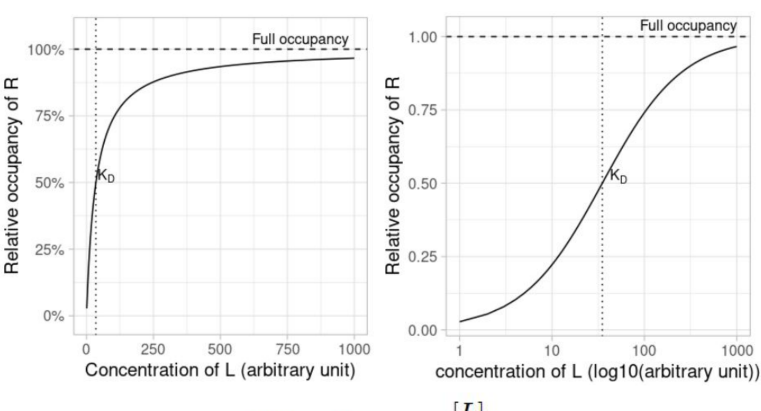
# An ordinary differential equation (ODE) model quantifies receptor occupancy by varying concentrations of ligands



## 

### The ODE model induces the simplest form of *Hill-Langmuir Equation*

- The Hill-Langmuir function describes the occupancy of receptors by natural ligands of drugs.
  - We can interpret K<sub>D</sub> in two ways: the ratio of reaction speeds, and the concentration required to occupy half of the receptors.
  - We will further discuss the Hill function again in the future.



 $\mathrm{L}+\mathrm{R}\rightleftharpoons\mathrm{LR} \qquad K_D\equiv k_2/k_1$ 

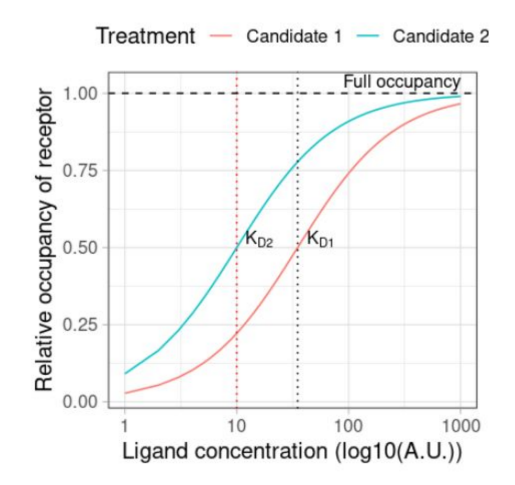
 $k_1$ 

 $k_2$ 

$$[LR] = [R_{total}] rac{[L]}{[L] + K_D}$$



## Question: all other conditions the same, which drug candidate is more favorable? Why?





### The Lotka-Volterra model of predator-prey relationships

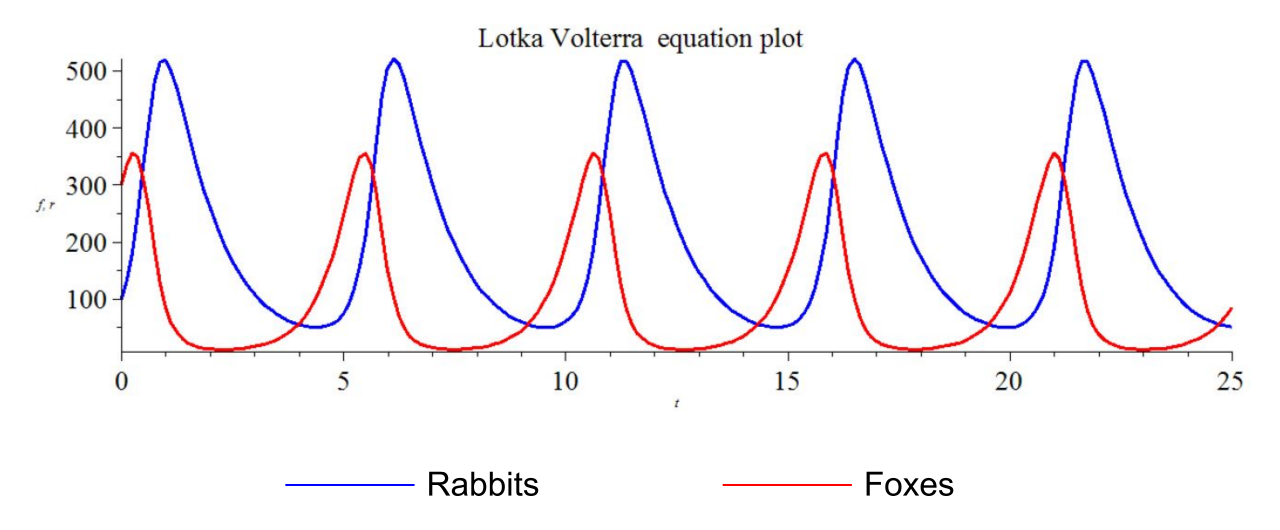
• The Lotka-Volterra equations modelling predator-prey relationships.

$$\frac{dx}{dt} = \alpha x - \beta xy,$$
(1)
$$\frac{dy}{dt} = -\gamma y + \delta xy,$$
(2)

where

- x is the number of prey (e.g. rabbits),
- y is the number of predator (e.g. foxes),
- $\frac{dx}{dt}$  and  $\frac{dy}{dt}$  represent growth rates of the two populations,
- t represents time,
- $\alpha$ ,  $\beta$ ,  $\gamma$ , and  $\delta$  are real parameters specifying the interaction of the two species.

### The Lotka-Volterra equations, visualized



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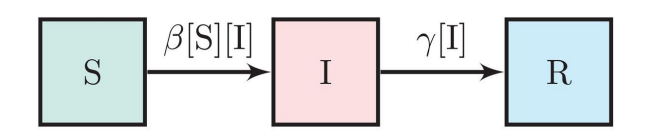
# 

# The SIR model of epidemiology models population behavior of viral infection and recovery

 $\beta IS$ 

I,

 $\beta IS$ 



dS

dt

dI

dt

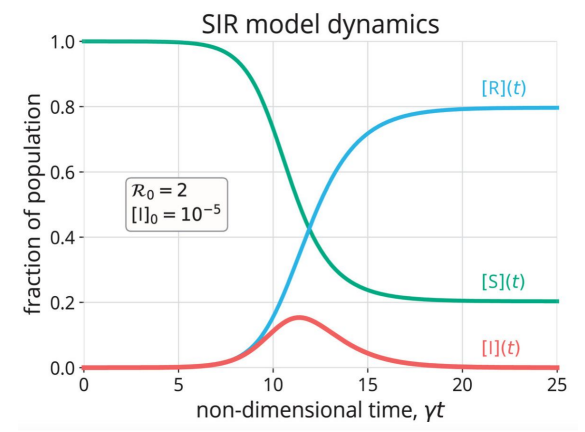
dR

dt

The SIR model of epidemiology

- S: Susceptible
- I: Infectious
- R: Removed

Simon CM. 2020. The SIR dynamic model of infectious disease transmission and its analogy with chemical kinetics. PeerJ Physical Chemistry 2:e14<u>https://doi.org/10.7717/peerj-pchem.14</u>

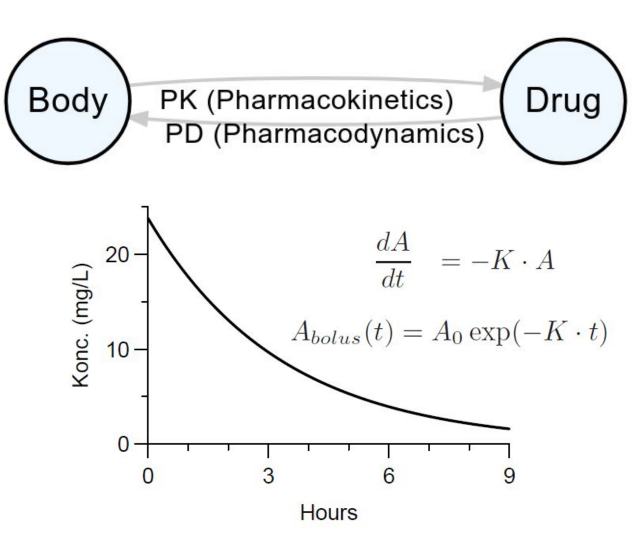


 $R_0$ , the basic reproduction number, is the number of people infected by the initial infectious individual. It is defined as  $\beta/\gamma$ .



## ODE-based mechanistic models are often used in pharmacokinetic modelling

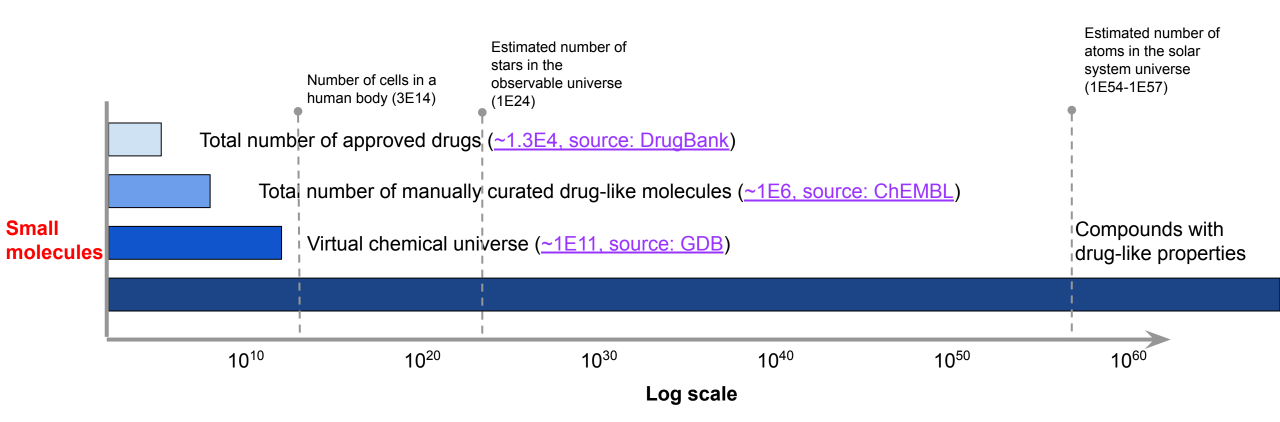
- Pharmacokinetics (PK) describes how the drug is <u>a</u>bsorbed, <u>d</u>istributed, <u>m</u>etabolised, and <u>excreted</u> by the body.
- Pharmacodynamics (PD) describes the effect of the drug to the body, mediated by drug-target interactions. PD is affected by PK, as well as other properties such as behaviour and genetics.
- A basic mathematical model of PK is a compartment model, i.e. one or more ordinary differential equations that describe the relationship between drug concentration and time. The simplest model is the decay model of bolus (injection).





### End of lecture on 18.10.2024

### Why drug *discovery*? 1. The chemical space is huge



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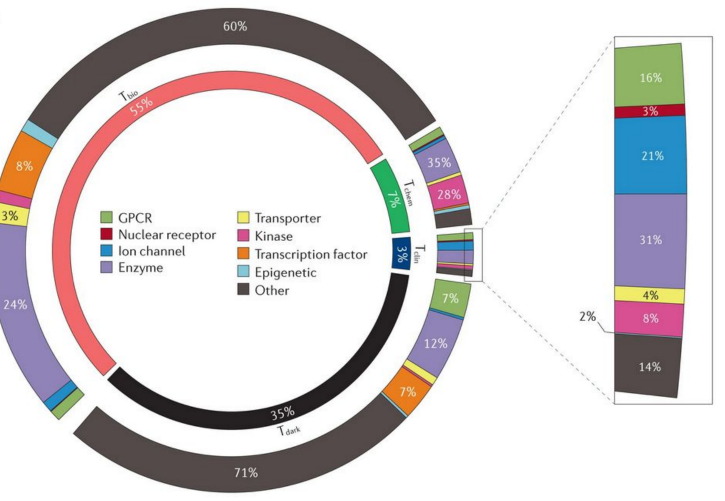
## Why drug *discovery*? 2. The druggable proteome is huge - even excluding mutations, transcriptome, genome, ...



There are about 20,000 proteins encoded by the human genome. We can classify them by (1) our knowledge of them, and (2) whether we have reliable chemical tools, biological tools, or even drugs to manipulate them.

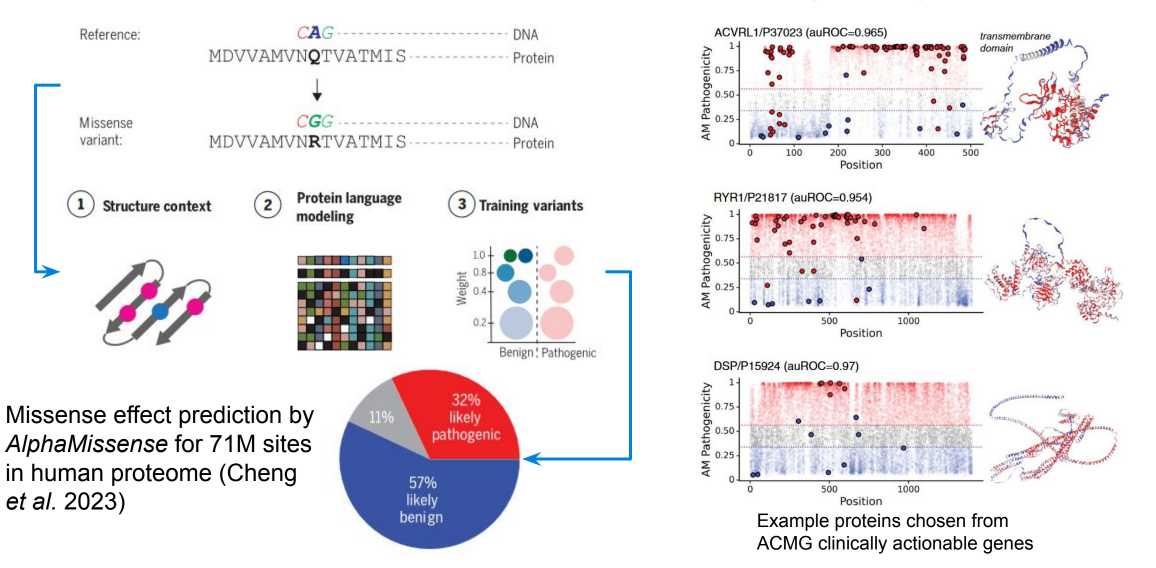
Inner ring: percentages of the whole proteome, classified by whether we have drugs ( $T_{clin}$ ), whether we have chemical tool compounds ( $T_{chem}$ ), whether we have biological compounds ( $T_{Bio}$ ), or we are in the dark ( $T_{Dark}$ ). Currently, we have only drugs for a few hundred proteins.

Outer ring: protein families.



Oprea, et al. "<u>Unexplored Therapeutic Opportunities in the Human Genome</u>." Nature Reviews Drug Discovery 17 (February 23, 2018): 317–32.

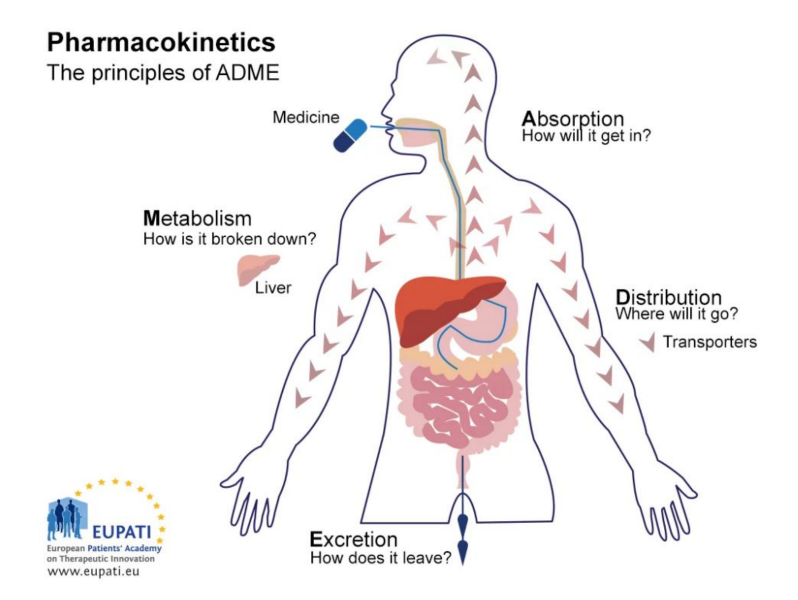
### Why drug *discovery*? 2. The druggable proteome is huge - now consider the mutations with predicted pathogenicity



#### Why drug discovery? 3. The central dogma as an information UNI BASEI channel: nodes and edges can all be targeted by drugs Small **Transcription** Catalysis **Translation** Protein DNA molecules Reverse **DNA** replication transcription Non-human Target **Example drugs** Small molecules **Dietary supplements** Enzyme inhibitors Catalysis Protein Receptor agonists/antagonists, ion channel blockers, antibodies Translation Antimicrobial protein synthesis inhibitors **RNA** Antisense oligonucleotides (ASO), vaccines **Transcription** Antimicrobials (e.g. actinomycin D and $\alpha$ -Amanitin), splicing modifiers (e.g. Risdiplam/Evrysdi) **Reverse transcription** Antivirals (e.g. reverse transcriptase inhibitors AZT/Zidovudine) DNA Gene therapies (e.g. chimeric activated receptors in T-cells, CAR-T) **DNA** replication Topoisomerase inhibitors (e.g. quinolones) and chemotherapy agents

## Why drug *discovery*? 4. The drug have to be absorbed and distributed in order to have systemic and organ-specific effects



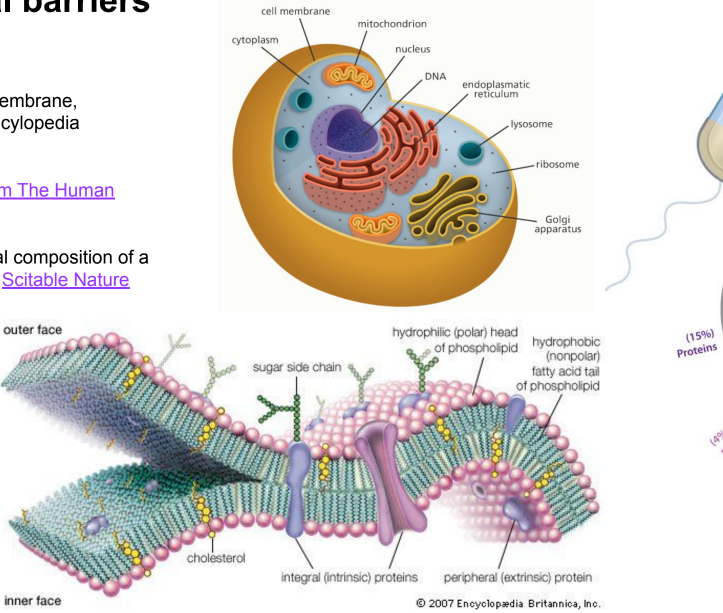


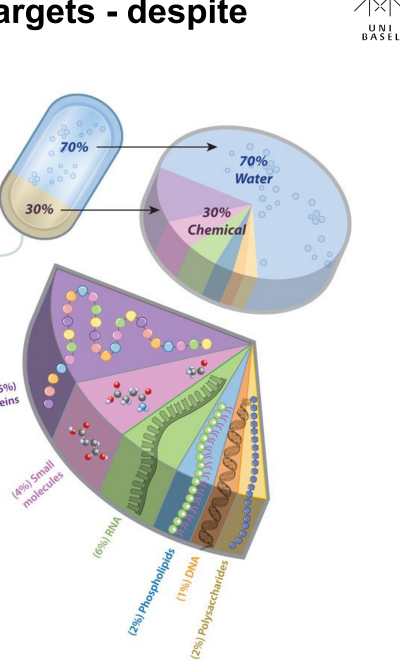
#### Why drug discovery? 5. Drugs have to reach the targets - despite physical barriers cell membrane

Bottom: Cell membrane, copyright of Encylopedia Britannica, Inc.

Top: Figure from The Human **Protein Atlas** 

Right: Chemical composition of a human cell, by Scitable Nature Education.

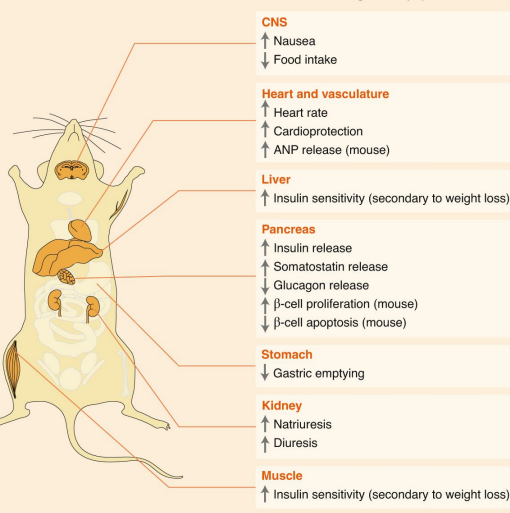




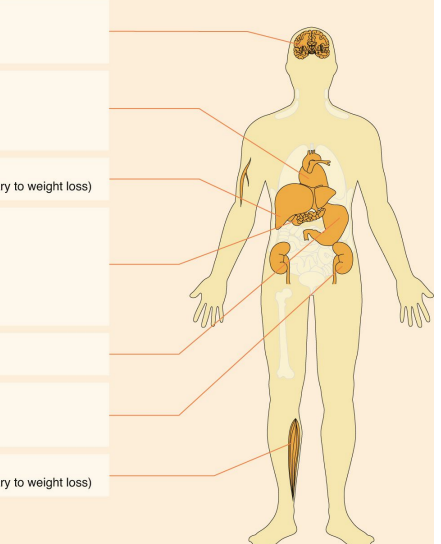
## Why drug *discovery*? 6. The drug can have organ-specific and systemic effects, causing either benefits or risks

Direct effects of Glucagon-like peptide (GLP-1) and GLP1 receptor agonists (GLP1-RA) like semaglutide.

Gribble, Fiona M., and Frank Reimann. "<u>Metabolic</u> <u>Messengers: Glucagon-like</u> <u>Peptide 1.</u>" Nature Metabolism 3, no. 2 (February 2021): 142–48.



#### Glucagon-like peptide 1



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# Why drug *discovery*? 7. Do all patients benefit from the drug, or only some of them? Learn from the story of Herceptin

#### Link to the video

#### **Questions for the video**

- 1. What is the **indication** of *Herceptin*? What is its generic (USAN, or United States Adopted Name) name?
- 2. What is the **gene target** of Herceptin?
- 3. Which class best describes the target: Enzyme, Ion channel, Receptor and Kinase, or Structural protein?
- 4. In which year was the **target** of Herceptin described? When was Herceptin **approved**?
- 5. What was the **improvement** of Herceptin compared with earlier antibodies?
- 6. Why does a **biomarker** matter besides developing drugs?
- 7. In the clinical trial of *Herceptin* for **metastatic breast cancer**, how much improvement in the **median survival** did Herceptin achieve? And how much improvement is in the **adjuvant setting** (Herceptin applied directly after operation)?

#### **Questions for further thinking**

- Susan Desmond-Hellmann summarizes successful drug development in four aspects: (1) having a deep understanding of the basic science and the characteristics of the drug, (2) targeting the right patients, (3) setting a high bar in the clinic, and (4) working effectively with key regulatory decision makers. Where do you think mathematics and computer science play a crucial role?
- She emphasized the importance of collaboration. What skill sets do we need for that?
- How do you like her presentation? Anything that you can learn from her about presentation and storytelling?

### **Conclusions and outlook**



- We reviewed the central dogma from the drug discovery's perspective.
- We learned examples of ODE-based mechanistic models.
- We considered key aspects to consider for a drug to work.
- Next time, we shall continue learning statistic and causal models.



### **Backup slides**