## "Ligand" Binding

"The secret of life is molecular recognition; the ability of one molecule to "recognize" another through weak bonding interactions."

Linus Pauling at the 25th anniversary of the Institute of Molecular Biology at the University of Oregon

Binding is the first step necessary for a biological response. Before macromolecules can perform a function, they normally must interact with a specific ligand. In some cases like myoglobin, binding and subsequent release of the ligand might be the sole function of the macromolecule. To understanding binding, we must consider the equilbria involved, how binding is affected by ligand and macromolecule concentration, and how to experimentally analyze and interpret binding data and binding curves.

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## **Goals for this Unit**

## Understand basic ligand binding equation

- essential terms and equations
- equilibrium binding / meaning of Kd / van't Hoff plots
- When you can simply by assuming [S] ~ [So]

## Complex equilibrium binding

- Multiple sites / independent or cooperative
- Diff. Microscopic vs. Macroscopic binding constants
- Scatchard plots and Hill Plots

## Techniques to determine K<sub>d</sub>

- Simple (Equil. Dialysis; Fluor) / ITC / SPR
- How to derive Kd from Equil. Dialysis data
- How to interpret Fluor / ITC and SPR data

Summary of Key Equations / Relationships  $E + S \Leftrightarrow ES ; \quad for single site$   $K_d = k_{off} / k_{on} = [E][S] / [ES] \text{ and } K_a = 1 / K_d$   $S_o = S + ES; \quad E_o = E + ES$ define Fractional Occupancy of sites  $\theta = [ES] / [E_o] = [ES] / ([E] + [ES]) \times [S] / [S] \times [1 / ES] / [1 / ES]$ then  $\theta = [S] / (K_d + [S])$ thus when  $[S] = -K_d$ , then  $\theta = 0.50$ when  $[S] = -4K_d$ , then  $\theta = 0.80$ when  $[S] = -4K_d$ , then  $\theta = 0.91$ Note: [S] = conc. of free ligand!!

w	hat is the meaning of the dissociation constant (Kd) for binding of a single ligand to its site?	
1. <i>K</i> <sub>d</sub> has units of concentration, M or mol/liter		
2.	$K_{d}$ gives the concentration of ligand to saturate 50% of the ligand binding sites (when the total site concentration is much lower than $K_{d}$ )	
3. /	Almost all binding sites are saturated when the free ligand concentration is 10 x $K_{\rm d}$	
4. <sup>.</sup>	The dissociation constant $K_d$ is related to Gibbs free energy $\Delta G^o$ by the relation $\Delta G^o = -R T \ln K_d$	



## *K*<sub>d</sub> values in biological systems

Movovalent ions binding to proteins or DNA have  $K_d 0.1 \text{ mM}$  to 10 mM Allosteric activators of enzymes e. g. NAD have  $K_d 0.1 \mu$ M to 0.1 mM

Site specific binding to DNA K<sub>d</sub> 1 nM to 1 pM

Trypsin inhibitor to pancreatic trypsin protease K<sub>d</sub> 0.01 pM

Antibody-antigen interaction have K<sub>d</sub> 0.1 mM to 0.0001 pM









## **No Assumptions - Key Equations**

Derivation of quadratic equation--with no assumptions concerning substrate concentration.

$fraction = \theta = [ES]/[E]_0 = [ES]/([E]+[ES])$	The derivation starts the same as above
$\theta = \frac{K_a[E][S]}{[E] + K_a[E][S]} = \frac{K_a[S]}{1 + K_a[S]} = \frac{[S]}{1 / K_a + [S]}$	The Fraction of sites bound relative to [S]
$\theta = \frac{[S]}{K_d + [S]} = \frac{[S]_0 - [ES]}{K_d + [S]_0 - [ES]} = \frac{[ES]}{[E]_0}$	<sup>-⊕</sup> Substitution of [S]=[S] <sub>0</sub> -[ES]
$[ES](K_d + [S]_0) - [ES]^2 = [E]_0[S]_0 - [ES][E]_0$ $[ES]^2 - [ES](K_d + [S]_0 + [E]_0) + [E]_0[S]_0$	TForm of equation require solution as the roots of the quadratic equation

### **No Assumptions - Key Equations**

Solution is quadratic equation:

$$ax^{2} + bx + c = 0$$
solution provided by the roots of the quadratic
$$x = \frac{-b \pm \sqrt{b^{2} - 4ac}}{2a}$$

$$[ES] = \frac{(E_{0} + S_{0} + K_{d}) - \sqrt{(E_{0} + S_{0} + K_{d})^{2} - 4 \cdot E_{0} \cdot S_{0}}}{2}$$

$$\theta = \frac{(E_{0} + S_{0} + K_{d}) - \sqrt{(E_{0} + S_{0} + K_{d})^{2} - 4 \cdot E_{0} \cdot S_{0}}}{2 \cdot E_{0}}$$

This equation must be used rather than the hyperbola whenever the enzyme concentration is comparable or greater than the dissociation constant. As a general rule, if  $[E]_0$  is less than 5 times the  $K_d$ , the hyperbolic fit is probably adequate.









### Fraction of sites bound

If the equations are defined for moles of substrate bound per mole of dimers instead of per mole of active sites, the equations becomes:

$$\upsilon = \frac{K_1[S] + 2K_1K_2[S]^2}{1 + K_1[S] + K_1K_2[S]^2} \quad \text{where } \upsilon = \frac{\text{moles S bound}}{\text{mole of dimense}}$$

Accordingly, the binding equation ranges from 0-2 moles bound rather than from 0-1 fraction of sites occupied.





### **EXPERIMENTAL DETERMINATION OF Kd**

**TECHNIQUES THAT REQUIRE SEPARATION OF BOUND FROM FREE LIGAND** – Care must be given to ensure that the equilibrium of M + L <==> ML is not shifted during the separation technique.

• *gel filration chromatography* - Add M to a given concentration of L. Then elute the mixture on a gel filtration column, eluting with the free ligand at the same concentration. The ML complex will elute first and can be quantitated . If you measure the free ligand coming off the column, it will be constant after the ML elutes with the exception of a single dip near where the free L would elute if the column was eluted without free L in the buffer solution. This dip represents the amount of ligand bound by M.

• membrane filtration - Add M to radiolableled L, equilibrate, and then filter through a filter which binds M and ML. For instance, a nitrocellulose membrane binds proteins irreversibly. Determine the amount of radiolabeled L on the membrane which equals [ML].

• precipitation - Add a precipitating agent like ammonium sulfate, which precipitates proteins and hence both M and ML. Determine the amount of ML.

### **EXPERIMENTAL DETERMINATION OF Kd**

#### TECH. THAT DO NOT REQUIRE SEPARATION OF BOUND FROM FREE LIGAND

• equilibrium dialysis - Place M in a dialysis bag and dialyse against a solution containing a ligand whose concentration can be determined using radioisotopic or spectroscopic techniques. At equilibrium, determine free L by sampling the solution surrounding the bag. By mass balance, determine the amount of bound ligand, which for a 1:1 stoichiometry gives ML. Repeat at different ligand concentrations.

• *gel shift assay* – Use autoradiography and gel electrophoresis with radiolabeled macromolecule.

• *spectroscopy* - Find a ligand whose absorbance or fluorescence spectra changes when bound to M. Alternatively, monitor a group on M whose absorbance or fluorescence spectra changes when bound to L.





# **Spectroscopy Fluorescence Spectroscopy** $F = F_0 + \Delta F \cdot \theta$ This equation normalizes the observable signal to a scale that can be related to fractional occupancy, $\theta$ . and $\theta$ is defined by either: $\theta = \frac{[S]_0}{K_d + [S]_0}$ OR..... $\theta = \frac{(E_0 + S_0 + K_d) - \sqrt{(E_0 + S_0 + K_d)^2 - 4 \cdot E_0 \cdot S_0}}{2 \cdot E_0}$ **Fluorescence Anisotropy**

Definition of fluorescence anisotropy *r* 

$$r = \frac{I_{\rm II} - I_{\perp}}{I_{\rm II} + 2I_{\perp}}$$

$$\theta = \frac{\left[P_{\text{tot}}\right]}{\left[P_{\text{tot}}\right] + K_D} = \frac{r_{\text{measured}} - r_{\text{min}}}{r_{\text{max}} - r_{\text{min}}}$$





# Analysis of binding of RNAP $\cdot \sigma^{54}$ to a promoter DNA sequence by measurements of fluorescence anisotropy



### **EXPERIMENTAL DETERMINATION OF Kd**

TECH. THAT DO NOT REQUIRE SEPARATION OF BOUND FROM FREE LIGAND

• *Equilibrium dialysis* - Place M in a dialysis bag and dialyse against a solution containing a ligand whose concentration can be determined using radioisotopic or spectroscopic techniques.

• *Fluorescence spectroscopy* - Find a ligand whose absorbance or fluorescence spectra changes when bound to M. Alternatively, monitor a group on M whose absorbance or fluorescence spectra changes when bound to L.

• Other (higher tech) methods: ITC - Isothermal Titration Calorimetry SPR – Surface Plasmon Resonance Fast Kinetics









## Binding - SPR or BIA

"The secret of life is molecular recognition" "Binding is the first step necessary for a biological response"

**Biacore's SPR technology**: label-free technology for *monitoring biomolecular interactions as they occur.* 

The detection principle relies on surface plasmon resonance (SPR), an electron charge density wave phenomenon that arises at the surface of a metallic film when light is reflected at the film under specific conditions.

The resonance is a result of energy and momentum being *transformed* from incident photons into surface plasmons, and is sensitive to the refractive index of the medium on the <u>opposite side</u> of the film from the reflected light.

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Note: Many of these figures/notes were taken from on-line resources from Biacore





Measure reflected (polarized) light as function of angle.

At a certain "Magic Angle" light is not reflected ("total internal reflection") but interacts with free electrons in gold to form a resonant energy wave – or surface plasmon.

Plasmon – A plasmon is a collective oscillation of the conduction electrons in a metal - a quasiparticle that can be regarded as a hybrid of the conducting electrons and the photon.

Angle is sensitive to refractive index of dielectric which varies with concentration of molecules on the other side of gold layer!





### **Three Corner Stones of Biacore Technology**



1. The Biacore sensor chip is at the heart of the technology. Quantitative measurements of the binding interaction between one or more molecules are dependent on the immobilization of a target molecule to the sensor chip surface. Binding partners to the target can be captured from a complex mixture, in most cases, without prior purification (for example, clinical material, cell culture media) as they pass over the chip. Interactions between proteins, nucleic acids, lipids, carbohydrates and even whole cells can be studied. The sensor chip consists of a glass surface, coated with a thin layer of gold. This forms the basis for a range of specialized surfaces designed to optimize the binding of a variety of molecules.

GOLD





## The Sensorgram is Information Rich





### Same affinity but different kinetics

- All 4 compounds have the same affinity  $K_D = 10 \text{ nM} = 10^{-8} \text{ M}$
- The binding kinetic constants vary by 4 orders of magnitude





## **SPR technology**

- Non-label
- Real-time
- Unique, high quality data on molecular interactions
- · Simple assay design
- · Robust and reproducible
- · Walk-away automation
- · Small amount of sample required



## **Applications in Proteomics**

- Fast, simple and compatible with any biological sample
- Monitors binding of native proteins from crude or purified samples
- Detects even low affinity binding events
- Recovers samples for MS analysis and identification
- Confirms results from other techniques
- Provides functional (interaction)
   data



## **Summary**

- SPR detects binding events as changes in mass at the chip surface
- Real-time kinetic measurements
- Qualitative rankings
- Measurement of *active* concentration
- Information about structure-activity relationships
- Low volumes of precious samples needed

**BUT !!!** -SPR is not a true solution method (vs. ITC) Attaching receptor to surface can influence binding properties.