

## Introduction

The 8-channel Octet RED96 system from ForteBio uses biolayer interferometry (BLI) to characterize biomolecular interactions in real-time, giving insight into desirable kinetic parameters such as  $k_{on}$ ,  $k_{off}$  and  $k_d$  for a given interaction.

Additionally, the Octet can be used for applications ranging from protein quantitation, lead identification/optimization, antibody characterization and cell line development.

## Features

### Dip and Read Technology

- The biosensor tip is coated with a compatible matrix that minimizes non-specific binding while providing a uniform and non-denaturing surface for biomolecules.

### Wide range of samples

- The Octet is suitable for use with purified samples as well as unpurified samples (including whole cells) with multiple complex matrices.

### Label free study of proteins

- Samples do not need to be chemically labelled for detection, allowing for the study of samples in their native context, giving data that is more representative of conditions *in situ*.

### Fast, simplified workflows

- Workflows can be simplified to five steps that are heavily automated, allowing for high throughput analysis of 96 well plates in under 30 minutes.

### Non-destructive investigation of samples

- Given the non-invasive nature of biolayer interferometry, samples can be recovered and used for further experiments, cutting down on the amount of protein needed.

### User friendly software

- Software makes setting up runs simple, allows for modification of protocols mid run and makes data processing straightforward.



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Produced by Serban Popa

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## Octet RED96 Biolayer Interferometry

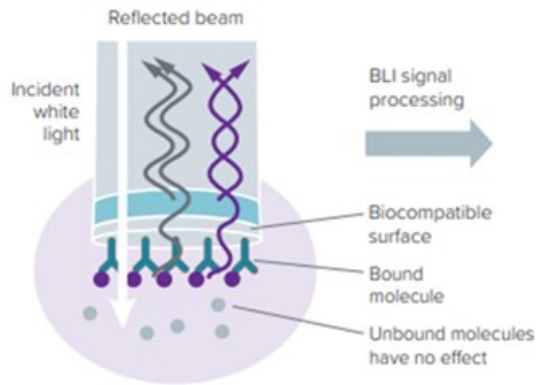
### Biomolecule Quantitation and Molecular Interaction Kinetics

- Determine  $k_{on}$ ,  $k_{off}$ ,  $k_d$  for desired biomolecular interactions in real time.
- Detection and quantitation of analytes ranging in size from hundreds of nm (bacteria/viruses) to low nm (antibodies/peptides).
- High throughput analysis of samples with 8 parallel channels.
- Wide variety of biosensor tips permitting multiple ways to immobilize target sample.

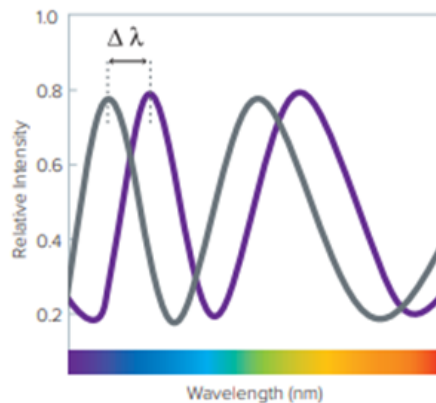
# Octet RED96 Biolayer Interferometry

## Theoretical Background

Biolayer Interferometry is an optical technique that detects changes in interference patterns from light waves. A biomolecule of interest is immobilized onto the tip of a biosensor and light interference at the tip of the biosensor is measured. The incident light reflects through two layers: a biocompatible layer and an internal reference layer.



Biological interactions will alter the thickness of the biolayer at the biosensor tip, thus altering the way the light reflects, resulting in a spectral shift that can be detected and monitored via a sensogram. The resulting spectral shift is then measured and can provide real time insight into the kinetics of the molecular interaction.

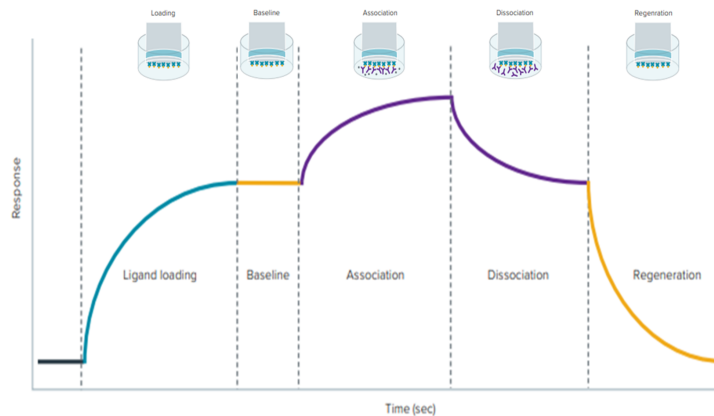


## Example workflow

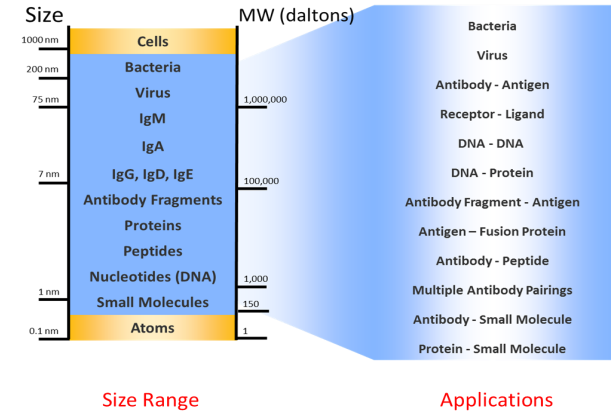
Below is a summary of a typical Octet experiment.

- 1) Loading is the immobilization of one binding partner (ligand) using the appropriate immobilization strategy.
- 2) Baseline is the washing away of any loosely bound ligand to equilibrate the biosensor tip.
- 3) Analyte association introduces the ligand's binding partner in order to obtain association kinetics.
- 4) Dissociation of the analyte from the formed complex obtains dissociation kinetics.
- 5) Regeneration of the biosensor removes bound ligand to allow for further experiments if applicable (such as when using Ni-NTA tips).

Idealized data for what one should expect to see for a typical kinetics experiment is shown below. Plotted is the response (the magnitude of the wavelength shift as a result of binding to the biosensor) as a function of time. The resulting data can then be fit to appropriate models to get the desired information.



## Applications and size ranges compatible with Octet



## Immobilization strategies

The table outlines a selection of strategies (others are available) for immobilizing samples based on sample features.

Sensor	Description	Mechanism of action
AHC	Anti-hIgG FC	Binding to human Fc-containing proteins and target analytes.
AR2G	Amine Reactive 2G	Binding to lysine residue of sample
NTA	Ni-NTA	Nickel binding to His tags
SA	Streptavidin	Binding to biotin tagged sample
SSA	Super Streptavidin	Binding of biotin tagged protein for small molecule screening

## Technical properties of the Octet

Sample volume	180-220 $\mu$ L
Analysis temperature	15-40°C
Association rate constant ( $k_a$ )	$10^2$ to $10^5$
Dissociation rate constant ( $k_d$ )	$10^{-5}$ to $10^{-2}$
Affinity constant ( $K_D$ )	0.1 mM – 0.1 nM
Molecular weight detection	>150 Da