**Introduction**

Biolayer Interferometry (BLI) is an optical technique that measures macromolecular interactions by analyzing interference patterns of white light reflected from the surface of a biosensor tip. BLI experiments are used to determine the kinetics and affinity of molecular interactions. In a BLI experiment, one molecule (the **Load Sample**) is immobilized to a Dip and Read Biosensor and binding of a second molecule (the **Analyte Sample**) is then measured. A change in the number of molecules bound to the end of the biosensor tip causes a shift in the interference pattern that is measured in real-time. BLI can be used to measure kinetic binding constants ($k_a$, $k_d$) and equilibrium binding constants (affinity, $K_D = 1/K_d$). Response is measured as a nm shift in the interference pattern and is proportional to the number of molecules bound to the surface of the biosensor. Response is recorded and displayed on a sensogram in real time.

**Instrument Overview**

The CMI has a **BLItz** instrument from ForteBio, which measures binding kinetics and equilibrium by BLI using ForteBio dip and read biosensors in low volume. For higher sensitivity and throughput and better data fitting, users should consider using the Octet RED384.

**Applications**

- Kinetic binding: $k_a$, $k_d$
- Equilibrium binding: $K_D$
- Macromolecular interactions > 10 KDa

**Key Features**

- Disposable biosensors (sensor regeneration not required)
- Low sample volume: 4 μl sample
Required Supplies

- ForteBio Biosensors.
  - See table below for popular sensor types and part numbers. Go to the ForteBio website: [http://www.fortebio.com/biosensor-types.html](http://www.fortebio.com/biosensor-types.html), for additional sensor types, including Anti-Mouse IgG Fc, Anti-Human Fab, Anti-GST, and biosensors recommended for quantitation.
- 96-well microplate to rehydrate sensors.
- Black 0.5 ml tubes (provided by the CMI).
- An empty biosensor tray to use as a working tray for hydration (optional).
- Pipettes (recommended).

<table>
<thead>
<tr>
<th>Popular ForteBio Dip and Read Biosensors for Kinetics</th>
<th>Part Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Streptavidin (SA) biosensors</td>
<td>18-5019 (96/tray)</td>
</tr>
<tr>
<td>High Precision Streptavidin (SAX) biosensors</td>
<td>18-5117</td>
</tr>
<tr>
<td>Super-Streptavidin (SSA) biosensors (for small molecules)</td>
<td>18-5057</td>
</tr>
<tr>
<td>anti-His (HIS1K) biosensors</td>
<td>18-5120</td>
</tr>
<tr>
<td>Ni-NTA (NTA) biosensors</td>
<td>18-5101</td>
</tr>
<tr>
<td>Anti-Human IgG Fc biosensors</td>
<td>18-5010</td>
</tr>
</tbody>
</table>

Sample Preparation

Assay Buffers

- Many buffers are compatible with BLI. It’s usually a good idea to start with a buffer system in which your proteins are well behaved.
- **Addition of 0.05% Tween 20 (or other surfactant) is usually required** to prevent non-specific binding, which is a frequent problem in BLI experiments.
  - Try detergent concentrations above the CMC, typically in the range of 0.02-0.1%.
- The sample used for the association phase should be in a buffer identically matched to that used for the baseline and dissociation phase.
  - Buffer match is especially important when a buffer component has a high refractive index, such as DMSO. Immobilized load sample should also be in the same buffer, if possible.
- **0.1% BSA** can also be used to minimize non-specific binding.
  - ForteBio sells a detergent-based Kinetic Buffer (PBS + 0.02 % Tween20, 0.1 % BSA, 0.05 % sodium azide) that you might consider.
  - **NOTE:** BSA is not universally beneficial and can sometimes increase non-specific binding.
- The same tube of buffer should be used for the baseline and dissociation phase.

Samples

- All BLI experiments are setup with one molecule fixed to the biosensor surface (the Load Sample) and a second molecule in solution (the Analyte Sample).
- Concentration should be accurately measured
  - Errors in Load concentration can affect signal intensity
Errors in the Analyte concentration will directly translate to errors in the $K_D$.

- Protein aggregates will interfere with BLI.
  - Filter or centrifuge samples before use.
  - Assess protein heterogeneity via light scattering.
  - Purify protein samples with soluble aggregates by size-exclusion chromatography.

- Recommended concentration ranges:
  - Load Sample (immobilized) 10-50 µg/ml (~µM range)
  - Analyte 0.01 – 100 X $K_D$ (0.01 – 10 X $K_D$)

- Sample and reagent volumes
  - **Sample** is generally placed in the dropper but can also be placed in a black tube
  - **Buffer for baseline** is placed in a black tube
  - **Buffer for hydrating the sensor** is placed in a 96-well plate (under the sensor tray)
    - Dropper volume 4 µl
    - Tube volume 250 µl
    - Plate volume 200 µl (Hydration)

### Getting Started
A BLItz experiment involves multiple steps in which ForteBio Dip and Read Biosensor are placed in either a 4 µL drop holder or in a 0.5 mL tube (filled with 250 µL). Baselines, dissociation, and quench steps should be performed in a tube. Loading and association can be performed either in the drop holder (for steps ≤ 5 min, generally) or in a tube (for steps > 5 min).

- **Immobilization**
- **Interaction Analysis**
- **Regeneration** *(Optional)*

### Resources
Additional resources are available at the instrument, including instrument manual.
Experimental Design Tips

- Do not overload the immobilized molecule.
- The same tube containing buffer should be used for the baseline and dissociation phase.
- The BLItz analysis will only perform a single reference sample subtraction (zero concentration of analyte). Use it.
- Do a control experiment with a reference sensor to measure non-specific binding to the sensor (using the highest concentration of analyte sample). If you see non-specific binding, then:
  - Optimize buffer conditions to eliminate non-specific binding.
  - Do a reference sensor measurement for each concentration of analyte and process and fit the data manually in a 3rd party fitting software package, such as GraphPad Prism.
  - Switch to the Octet RED384 instrument to perform reference sensor subtraction.

General Care and Maintenance

Startup

1. Book time on the PPMS calendar before you start.
2. Login to the computer using your PPMS credentials (eCommons ID and password).
3. Turn on the instrument using the switch at the back.
4. Open the BLItz Pro software.
5. Choose an experiment type: almost all experiments will be Advanced Kinetics in which one molecule is immobilized and binding of another is measured.

Data Collection and Analysis

Advanced Kinetics Experiment

1. Preparation
   a. Hydrate the biosensors for at least 10 min before each measurement.
   b. Prepare protein samples in assay buffer (load 4 μL per measurement).
   c. Prepare assay buffer (250 μL in a black tube).
2. Input the Run Settings
   a. Include the molar concentration of the analyte sample for each measurement.
   b. Always enable the shaker.
   c. Set step types and duration.
      i) For a first experiment, the default values are a good start.
3. Click Next when ready to start the experiment.
4. Follow the prompts to switch between the buffer in the tube and samples in the drop holder.
   a. Drop holder can be removed to aid in cleaning or filling.
   b. Always return drop holder to the instrument.
Data Analysis
1. Select the zero concentration of analyte as the Reference sample.
2. Enable step corrections at the start of association and dissociation.
3. Choose local or global fitting.
4. Click Analyze.

Shutdown
5. Clean drop holder.
6. Remove and discard biosensors and black tubes.
7. Return borrowed empty sensor trays to the drawer under the instrument.
8. Clean up in and around the instrument.
9. Turn off the instrument power.
10. Close the control and analysis software.
11. Logoff from PPMS!

Book time and Report Problems through the PPMS system: [https://ppms.us/hms-cmi](https://ppms.us/hms-cmi)
- rates are based on booked and real-time usage.

Contact cmi@hms.harvard.edu with questions.

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